National Clinical Guideline Centre

Varicose Veins Appendices

Varicose Veins

Appendices A–O

Appendices Methods, evidence and recommendations July 2013

Draft for Consultation

Commissioned by the National Institute for Health and Care Excellence

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Varicose Veins Full Guideline Appendices (July 2013)

Funding National Institute for Health and Care Excellence

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1 Appendices

Appendix A: Scope

NATIONAL INSTITUTE FOR HEALTH AND CLINICAL EXCELLENCE

SCOPE

1 Guideline title

Varicose veins in the legs: the diagnosis and management of varicose veins

1.1 Short title

Varicose veins in the legs

2 The remit

The Department of Health has asked NICE: 'To produce a clinical guideline on the management of varicose veins'.

3 Clinical need for the guideline

3.1 Epidemiology

- a) Varicose veins are a common condition. They are dilated, often palpable, subcutaneous veins with reversed blood flow and are most commonly located on the lower legs.
- b) The Edinburgh Vein Study (1999) showed age-adjusted prevalence rates for varicose veins of 39.7% in men and 32.2% in women. The same study found prevalence rates for chronic venous insufficiency of 9.4% in men and 6.6% in women. In contrast the BONN Vein study II (2010) found lower prevalence rates for varicose veins (25.1%) and higher rates for chronic venous insufficiency (16.0%); it did not identify gender differences.
- c) The Framingham Study (1988) conducted in the USA found that the annual incidence of varicose veins was 1.9% for men and 2.6% for women. The incidence was found not to vary within the age range (40–89 years).

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- d) The age of onset does vary and prevalence rises with age.
 Varicose veins are common during pregnancy and affect about 40% of pregnant women.
- e) The clinical presentation of varicose veins differs and some people are asymptomatic. In the majority of people varicose veins do not cause damage or threat to the limb but are associated with aching, itching, burning, cramps at night, and restless legs. In some people with varicose veins, progression of the condition may result in more severe problems such as skin pigmentation changes, eczema, infection, superficial thrombophlebitis, bleeding, loss of subcutaneous tissue, lipodermatosclerosis and venous ulceration.

3.2 Current practice

- Current management of varicose veins is controversial and there is considerable variation in clinical practice.
- b) There is a lack of consensus about optimum indications for referral and treatment. Suitability for varicose vein treatment is mainly determined by clinical examination and followed by a hand held doppler and/or duplex scan to determine whether venous reflux is present. However, there can be an inconsistent association between the symptoms of varicose veins and their severity or size on examination.
- c) There are many clinical grading systems for varicose veins, including CEAP (clinical signs, aetiologic classification, anatomic distribution and pathophysiological dysfunction). However, with most of these there is a lack of agreement as to their usefulness for clinical decision making. Although CEAP is the most widely accepted grading system for varicose veins as a way to determine treatment needs, it is not discriminatory when looking at mild forms of the disease or predicting who would benefit the most from intervention.

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- d) Treatment options include:
 - Conservative treatment this includes diet, lifestyle advice and compression therapy. These are often used as first-line treatments in primary care.
 - Pharmacological treatments for the relief of symptoms. There
 are none currently licensed for use in the UK.
 - Interventional procedures:
 - Surgical treatments these include ligation (tying off the vein), stripping and avulsion (different ways of removing the vein). These operations can be performed under general, regional or local anaesthesia, depending on the preferences of the surgeon and patient, and on the extent and the complexity of the varicose veins to be treated.
 - Sclerotherapy injecting a sclerosing (irritating) agent directly into the varicose veins. This can be either as liquid or foam. This causes an inflammatory response that closes off the vein.
 - Thermal ablation heating the vein from inside (for example using radiofrequency or laser catheters), this causes irreversible damage to the vein and its lining and closes it off.
- e) Often several of the above techniques are used in combination.
 Treatment choice depends on a number of factors; symptoms, severity, patient preference and available medical resources.
- f) The lack of clarity over assessment and the perceived similarity in outcomes from the different the interventional therapies have led to considerable variation in the management of varicose veins.

4 The guideline

The guideline development process is described in detail on the NICE website (see section 6, 'Further information').

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This scope defines what the guideline will (and will not) examine, and what the guideline developers will consider. The scope is based on the referral from the Department of Health.

The areas that will be addressed by the guideline are described in the following sections.

4.1 Population

4.1.1 Groups that will be covered

- Adults (18 and older) with primary or recurrent varicose veins in their legs.
- b) The particular needs of pregnant women will be considered.

4.1.2 Groups that will not be covered

- a) Children and young people (younger than 18).
- b) People with venous malformations.
- c) People with varicose veins in places other than their legs.

4.2 Healthcare setting

- a) NHS healthcare settings in which varicose veins are managed.
- 4.3 Clinical management

4.3.1 Key clinical issues that will be covered

- Assessment for referral and treatment, including hand held doppler, duplex scan and clinical grading systems.
- b) Conservative treatments, including
 - lifestyle advice
 - compression therapies.

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- c) Interventional therapies, for example:
 - surgical treatments
 - thermal ablation treatments.
- d) Information and support needs of patients and carers.

4.3.2 Clinical issues that will not be covered

- a) Management of leg ulcers, other than the role of ablative truncal venous interventions.
- b) Spider veins (thread veins).
- Management of pelvic varicose veins unless they are associated with primary or recurrent lower limb varicose veins.
- d) Management of varicose veins not located on the legs.
- e) Pharmacological treatment.
- f) Alternative or complementary treatment.

4.4 Main outcomes

- a) Health-related quality of life, using generic validated tools (for example, Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (for example, Chronic Venous Insufficiency Questionnaire).
- b) Patient-assessed symptoms.
- c) Physician-reported outcome (venous clinical severity score or venous disability score).
- Complications from varicose veins (skin ulcer occurrence or changes, haemorrhage, and phlebitis).
- e) Adverse events from intervention (including stroke, deep vein thrombosis and neuropraxia).

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f) Recurrent varicose veins.

g) Vein reflux and occlusion (blockage) rates.

4.5 Economic aspects

Developers will take into account both clinical and cost effectiveness when making recommendations involving a choice between alternative interventions. A review of the economic evidence will be conducted and analyses will be carried out as appropriate. The preferred unit of effectiveness is the quality-adjusted life year (QALY), and the costs considered will usually only be from an NHS and personal social services (PSS) perspective. Further detail on the methods can be found in 'The guidelines manual' (see 'Further information').

4.6 Status

4.6.1 Scope

This is the final scope.

4.6.2 Timing

The development of the guideline recommendations will begin in September 2011.

5 Related NICE guidance

5.1 Published guidance

5.1.1 NICE guidance to be incorporated

NICE interventional procedure guidance 314 (2009) 'Ultrasound-guided foam sclerotherapy for varicose veins' is being updated and we expect that guidance will be available in late 2012. If the updated guidance recommends that the procedure can be used without the need for special arrangements for clinical governance, consent or research, the interventional procedure guidance will be incorporated into the guideline.

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This guideline will also incorporate the following NICE guidance.

- Endovenous laser treatment of the long saphenous vein. NICE interventional procedure guidance 52 (2004). Available from www.nice.org.uk/guidance/IPG52
- Transilluminated powered phlebectomy for varicose veins. NICE interventional procedure guidance 37 (2004). Available from www.nice.org.uk/guidance/IPG37
- Radiofrequency ablation of varicose veins. NICE interventional procedure guidance 8 (2003). Available from <u>www.nice.org.uk/guidance/IPG8</u>

5.1.2 Other related NICE guidance

- Promoting physical activity in the workplace. NICE public health guidance 13 (2008). Available from <u>www.nice.org.uk/guidance/PH13</u>
- Smoking cessation services. NICE public health guidance 10 (2008). Available from <u>www.nice.org.uk/guidance/PH10</u>
- Physical activity and the environment. NICE public health guidance 8 (2008). Available from <u>www.nice.org.uk/guidance/PH8</u>
- Obesity. NICE clinical guideline 43 (2006). Available from <u>www.nice.org.uk/guidance/CG43</u>
- Four commonly used methods to increase physical activity. NICE public health guidance 2 (2006). Available from <u>www.nice.org.uk/guidance/PH2</u>
- Brief interventions and referral for smoking cessation in primary care and other settings. NICE public health guidance 1 (2006). Available from www.nice.org.uk/guidance/PH1
- NICE referral advice recommendations database [online]. Available from www.nice.org.uk/usingguidance/referraladvice/index.jsp

6 Further information

Information on the guideline development process is provided in:

- 'How NICE clinical guidelines are developed: an overview for stakeholders' the public and the NHS'
- 'The guidelines manual'.

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These are available from the NICE website

(<u>www.nice.org.uk/guidelinesmanual</u>). Information on the progress of the guideline will also be available from the NICE website (<u>www.nice.org.uk</u>).

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Appendix B: Declarations of interest

2 **B.1** Professor Alun Davies

CDC mosting		Action taken
GDG meeting	Declaration of Interests	Action taken
On Application	AD declared he knew of no personal pecuniary interests, personal family interests, non- personal pecuniary interests or personal non- pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting [21/09/11]	None Declared	None
Second GDG Meeting [02/11/11]	Personal pecuniary (non-specific): Attended and lectured at Turkish Vascular society, European Society of Vascular surgery, European Venous Forum, ARAB vein meeting (Saudi Arabia). The meeting organisers paid expenses but no monies were given for lecturing. The organisers would have had multiple healthcare related sponsors.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary
Third GDG Meeting [14/12/11]	Personal pecuniary (non-specific): Expenses paid for attending and lecturing at Veith symposium (USA). The meeting organisers paid expenses but no monies were given for lecturing. The organisers would have had multiple industrial sponsors.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary
Fourth GDG Meeting [25/01/12]	Personal Pecuniary (non-specific): Attended academic meeting of CACVS (Controversies and Updates in Vascular Surgery) in France and received expenses for registration, travel and accommodation from meeting organisers, who would have had industrial sponsorship.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary.
	Non-personal pecuniary (non-specific): Research department was awarded a grant from Venous Forum UK.	The grant was awarded by an academic body, not involved in the manufacture of varicose veins related equipment. NOF, KK and KH agreed no action was necessary.
Fifth GDG Meeting [07/03/12]	Personal Pecuniary (non-specific): Attended American Venous Forum (USA) with registration paid for by meeting organisers who would have had industrial sponsorship.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary
	Non-personal pecuniary (non-specific) : Met with Sapheon to investigate commencing a trial in the UK on vein ablation with a novel glue technology. A grant has been awarded from Sapheon to Imperial College for which AD is the principle investigator.	Glue is not a technology under consideration in this guideline and Sapheon do not make any other products used in the management of varicose veins. NOF, KK and KH agreed no action was necessary.

GDG meeting	Declaration of Interests	Action taken
	Non personal pecuniary (non-specific): A grant was awarded to the research department from Clarivein (novel vibrating technology for varicose vein treatment), for which AD is the principle investigator.	The technique under investigation is not a technology under consideration in this guideline and Clarivein do not make any other products used in the management of varicose veins. NOF, KK and KH agreed no action was necessary.
Sixth GDG Meeting [18/04/12]	Non personal pecuniary (non-specific):- A grant of was awarded to the research department from Geko to fund a trial looking at an electrical stimulation tool for the prevention of VTE.	The prevention of VTE is not relevant to this guideline and the Geko do not make any other products used in the management of varicose veins. NOF, KK and KH agreed no action was necessary.
	Personal pecuniary (non-specific): Organised and attended major meeting (Charing Cross Symposium) with multiple industrial sponsors and attended meal.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary.
	Personal pecuniary (non-specific): Received a bursary to cover travel and accommodation expenses from Australasian College of Phlebology (ACP) to attend and lecture at their annual meeting (Australia). The meeting organisers who would have had industrial sponsorship.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary
	Personal pecuniary (non-specific): Attended and meeting with Servier (Paris) to discuss Daflon a pharmaceutical treatment of varicose veins. Servier paid travel expenses.	Pharmacological treatments are under consideration in this guideline and Servier do not make any other products used in the management of varicose veins. NOF, KK and KH agreed no action was necessary.
	Personal non-pecuniary (non-specific): Received expenses for travel, registration and accommodation to the European Vascular Course (Belgium) from the meeting organisers. The meeting organisers who would have had industrial sponsorship.	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. NOF, KK and KH agreed no action was necessary
Seventh GDG Meeting [11/07/12]	Did not attend this meeting	No action necessary
Eighth GDG Meeting [03/10/12]	Personal pecuniary (non-specific): Attended vascular meeting in India funded by the Indian vascular society and VASCUTEK.	VASCUTEK do not make any venous products. Funding not beyond that reasonably expected. KK and KH agreed no action was necessary
Ninth GDG	No change to declarations of interest.	None

GDG meeting	Declaration of Interests	Action taken
Meeting [07/11/12]		
Tenth GDG Meeting [19/12/12]	Personal pecuniary (non-specific): Attended the Veith Vascular Symposium as a guest speaker and gave 7 talks. Fights were funded by VASCUTEK UK and the meeting was sponsored by multiple companies	VASCUTEK do not make any venous products. Funding not beyond that reasonably expected. KK and KH agreed no action was necessary
Eleventh GDG Meeting [10/04/13]	Non-personal pecuniary interest: Attended the ISVS Miami, AVF Phoenix and CX symposium – all meetings had commercial sponsorship from many organisations	No action necessary

1 B.2 Dr. Mustapha Azzam

GDG meeting	Declaration of Interests	Action taken
On Application	MA declared he knew of no personal pecuniary interests, personal family interests, non-personal pecuniary interests or personal non-pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting [21/09/11]	None	None
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	None	None
Fourth GDG Meeting [25/01/12]	None	None
Fifth GDG Meeting [07/03/12]	Personal pecuniary (non-specific): Attended a meeting with GEKO to discuss their electrical stimulation device for the prevention of DVT.	The prevention of VTE is not relevant to this guideline and the GEKO do not make any other products used in the management of varicose veins. AD, NOF, KK and KH agreed no action was necessary.
Sixth GDG Meeting [18/04/12]	Personal pecuniary (non-specific):- Attended a dinner with representatives of Sapheon.	Sapheon do not make anything under consideration in the guideline. AD, KK and KH agreed no action was necessary.
	Personal pecuniary (non-specific): Attended an advisory board meeting with First Sky Medical	First Sky Medical do not make anything under consideration in the guideline. AD, KK and KH agreed no action was necessary
Seventh GDG	Personal pecuniary (non-specific): Assisted	The prevention of VTE is not relevant to

GDG meeting	Declaration of Interests	Action taken
Meeting [11/07/12]	GEKO in setting up a Doppler protocol for a trial of their electrical stimulation device for the prevention of DVT in orthopaedic patients.	this guideline and GEKO do not make any other products used in the management of varicose veins. AD, KK and KH agreed no action was necessary.
Eighth GDG Meeting [03/10/12]	No change to declarations of interest.	None
Ninth GDG Meeting [07/11/12]	No change to declarations of interest.	None
Tenth GDG Meeting [19/12/12]	No change to declarations of interest.	None
Eleventh GDG Meeting [10/04/13]	Personal pecuniary interest : Attended a meeting about Geko Electro stimulation device – device for prevention.	No action was necessary

2 B.3 Professor Andrew Bradbury

GDG meeting	Declaration of Interests	Action taken
On Application	AB declared he knew of no personal pecuniary interests, personal family interests, non- personal pecuniary interests or personal non- pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting [21/09/11]	Did not attend this meeting	None
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	Personal pecuniary (Specific): Attended as part of the faculty the European Venous Forum (EVF) Hands-On Workshop (HOW) in Vienna in November 2011. Travel and accommodation costs were covered by the EVF and those funds were obtained from a range of different companies who manufacture materials and equipment used for the treatment of venous disease.	No direct sponsorship by any manufacturer company involved in varicose veins related products. All sponsorship not considered to be beyond what would reasonably be required for accommodation, meals and travel to attend. AD, NOF and KK agreed no action was necessary.
Fourth GDG Meeting [25/01/12]	No change to declarations of interest.	None
Fifth GDG Meeting	No change to declarations of interest.	None

GDG meeting	Declaration of Interests	Action taken
[07/03/12]		
Sixth GDG Meeting [18/04/12]	Non-personal pecuniary (Specific): Department did some research which was funded by British Biotechnology Group (BTG) Ltd, who are trying to make a commercial foam, looking at the evaluation of a novel health-related quality-of-life (HRQL) instrument administered via a Palmtop Application Device (PAD). The funds were used to pay the salary of a research nurse. No personal payment received	No personal knowledge of the intervention or matter either through his or her own work, or through direct supervision of other people's work. AD, KK and KH agreed no action was required.
Seventh GDG Meeting [11/07/12]	Personal Non-pecuniary –Had a discussion with the Chief Executive of STD pharmaceuticals (who make a chemical called STS which can be used to make foam for the treatment of varicose veins) during dinner at the European Venous Forum annual meeting in Florence in June 2012. During that dinner the application of a European licence for STS foam was discussed.	Discussed within the GDG. It was concluded that as the GDG were aware of this declaration and could take it into consideration during the GDG meetings they did not feel that AB should be excluded. AB will remain in the discussion and development of recommendations.
Eighth GDG Meeting [03/10/12]	Did not attend this meeting	None
Ninth GDG Meeting [07/11/12]	No change to declarations of interest.	None
Tenth GDG Meeting [19/12/12]	Did not attend this meeting	None
Eleventh GDG Meeting [10/04/13]	Non-personal pecuniary interest: Attended a faculty of Charing Cross Meeting – expenses paid by Charing Cross – no personal fee.	No action necessary

2 B.4 Dr. Jocelyn Brookes

GDG meeting	Declaration of Interests	Action taken
On Application	JB declared he knew of no personal pecuniary interests, personal family interests, non- personal pecuniary interests or personal non- pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting [21/09/11]	None	None
Second GDG Meeting	None	None

GDG meeting	Declaration of Interests	Action taken
[02/11/11]		
Third GDG Meeting [14/12/11]	None	None
Fourth GDG Meeting [25/01/12]	Did not attend this meeting	None
Fifth GDG Meeting [07/03/12]	None	None
Sixth GDG Meeting [18/04/12]	Did not attend this meeting	None
Seventh GDG Meeting [11/07/12]	None	None
Eighth GDG Meeting [03/10/12]	None	None
Ninth GDG Meeting [07/11/12]	Did not attend this meeting	None
Tenth GDG Meeting [19/12/12]	None	None
Eleventh GDG Meeting [10/04/13]	Personal pecuniary interest : Attended the LINC meeting sponsored by Spectrometics Ltd Personal non-pecuniary interest : Participate in private practice involving treatment of varicose vein.	No action necessary

2 B.5 Mrs. Joyce Calam

GDG meeting	Declaration of Interests	Action taken
On Application	JC declared she knew of no personal pecuniary interests, personal family interests, non- personal pecuniary interests or personal non- pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting	None	None

GDG meeting	Declaration of Interests	Action taken
[21/09/11]		
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	None	None
Fourth GDG Meeting [25/01/12]	None	None
Fifth GDG Meeting [07/03/12]	None	None
Sixth GDG Meeting [18/04/12]	None	None
Seventh GDG Meeting [11/07/12]	None	None
Eighth GDG Meeting [03/10/12]	None	None
Ninth GDG Meeting [07/11/12]	None	None
Tenth GDG Meeting [19/12/12]	None	None
Eleventh GDG Meeting [10/04/13]	None	None

2 B.6 Mr. David Evans

GDG meeting	Declaration of Interests	Action taken
On Application	DE declared he knew of no personal pecuniary interests, personal family interests, non- personal pecuniary interests or personal non- pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting [21/09/11]	None	None

GDG meeting	Declaration of Interests	Action taken
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	Personal pecuniary (non-specific) : I hold the shares with Astrazeneca and GlaxoSmithKleine.	As Astrazeneca nor GlaxoSmithKleine is involved in the manufacture of any products relevant to the varicose vein guideline AD, NO'F and KK agreed that no action is necessary.
Fourth GDG Meeting [25/01/12]	No change to declarations of interest.	None
Fifth GDG Meeting [07/03/12]	No change to declarations of interest.	None
Sixth GDG Meeting [18/04/12]	No change to declarations of interest.	None
Seventh GDG Meeting [11/07/12]	No change to declarations of interest.	None
Eighth GDG Meeting [03/10/12]	No change to declarations of interest.	None
Ninth GDG Meeting [07/11/12]	No change to declarations of interest.	None
Tenth GDG Meeting [19/12/12]	No change to declarations of interest.	None
Eleventh GDG Meeting [10/04/13]	Did not attend.	None

2 B.7 Mr Nick Hickey

GDG meeting	Declaration of Interests	Action taken
On Application	Personal pecuniary interest - I am employed by the NHS as a vascular surgeon, the role includes the management of varicose veins. I am also in private practice, including the treatment of varicose veins.	No action necessary
First GDG	No change to declarations of interest.	None

GDG meeting	Declaration of Interests	Action taken
meeting [21/09/11]		
Second GDG Meeting [02/11/11]	No change to declarations of interest.	None
Third GDG Meeting [14/12/11]	No change to declarations of interest.	None
Fourth GDG Meeting [25/01/12]	No change to declarations of interest.	None
Fifth GDG Meeting [07/03/12]	No change to declarations of interest.	None
Sixth GDG Meeting [18/04/12]	No change to declarations of interest.	None
Seventh GDG Meeting [11/07/12]	No change to declarations of interest.	None
Eighth GDG Meeting [03/10/12]	No change to declarations of interest.	None
Ninth GDG Meeting [07/11/12]	No change to declarations of interest.	None
Tenth GDG Meeting [19/12/12]	No change to declarations of interest.	None
Eleventh GDG Meeting [10/04/13]	None	None

2 B.8 Mr Keith Poskitt

GDG meeting	Declaration of Interests	Action taken
On Application	KP declared he knew of no personal pecuniary interests, personal family interests, non- personal pecuniary interests or personal non- pecuniary interests in the past 12 months or upcoming months.	No action necessary

GDG meeting	Declaration of Interests	Action taken
First GDG meeting [21/09/11]	None	None
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	Personal pecuniary (non-specific): Attendance at Veith Symposium meeting	No direct sponsorship by any manufacturer company involved in varicose veins related products. Funding not beyond that reasonably expected. AD, NOF and KK agreed no action was necessary.
Fourth GDG Meeting [25/01/12]	None	None
Fifth GDG Meeting [07/03/12]	Personal non pecuniary : Attended a breakfast meeting on chronic venous insufficiency at House of Commons on 7th March 2012	AD, NOF and KK agreed that no action was necessary.
Sixth GDG Meeting [18/04/12]	None	None
Seventh GDG Meeting [11/07/12]	Non personal pecuniary (specific): Received funding from STD pharmaceuticals to support a trip by his registrar to present research in Florence at European Venous Forum	KK and KH agreed that no action was necessary.
Eighth GDG Meeting [03/10/12]	No change to declarations of interest.	None
Ninth GDG Meeting [07/11/12]	No change to declarations of interest.	None
Tenth GDG Meeting [19/12/12]	No change to declarations of interest.	None
Eleventh GDG Meeting [10/04/13]	None	None

2 B.9 Ms. Hazel Trender

GDG meeting	Declaration of Interests	Action taken
On Application	HT declared she knew of no personal pecuniary interests,	No action necessary

GDG meeting	Declaration of Interests	Action taken
	non-personal pecuniary interests or personal non-pecuniary interests in the past 12 months or upcoming months.	
First GDG meeting [21/09/11]	None	None
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	None	None
Fourth GDG Meeting [25/01/12]	Personal pecuniary (specific): Personal non- pecuniary - Sponsorship to attend Vascular Society Annual meeting from Medi UK	Medi-UK manufactures compression stocking which may be used for varicose veins patients. However, funding not beyond that reasonably expected. AD, NOF and KK agreed no action was necessary.
Fifth GDG Meeting [07/03/12]	No change to declarations of interest.	None
Sixth GDG Meeting [18/04/12]	Personal pecuniary (specific): Personal non- pecuniary - Funding for registration and accommodation for venous forum from Activa (stocking manufacturers)	Activa manufactures compressions stockings which may be used for varicose veins patients. However, funding not beyond that reasonably expected. AD, NOF and KK agreed no action was necessary.
Seventh GDG Meeting [11/07/12]	No change to declarations of interest.	None
Eighth GDG Meeting [03/10/12]	No change to declarations of interest.	None
Ninth GDG Meeting [07/11/12]	No change to declarations of interest.	None
Tenth GDG Meeting [19/12/12]	Personal pecuniary (specific): Sponsorship from Medi UK to attend the Vascular Society Meeting.	Medi-UK manufactures compression stocking which may be used for varicose veins patients. However, funding not beyond that reasonably expected. AD and KK agreed no action was necessary.

GDG meeting	Declaration of Interests	Action taken
Eleventh GDG Meeting [10/04/13]	None	None

2 B.10 Dr. Mark Vaughn

GDG meeting	Declaration of Interests	Action taken
On Application	MV declared he knew of no personal pecuniary interests, personal family interests, non-personal pecuniary interests or personal non-pecuniary interests in the past 12 months or upcoming months.	No action necessary
First GDG meeting [21/09/11]	None	None
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	None	None
Fourth GDG Meeting [25/01/12]	None	None
Fifth GDG Meeting [07/03/12]	None	None
Sixth GDG Meeting [18/04/12]	None	None
Seventh GDG Meeting [11/07/12]	None	None
Eighth GDG Meeting [03/10/12]	None	None
Ninth GDG Meeting [07/11/12]	Did not attend this meeting	None
Tenth GDG Meeting	Personal non-pecuniary : Agreed to speak at the National Primary Care Conference in May	None

GDG meeting	Declaration of Interests	Action taken
[19/12/12]	2013. There have been no discussions about payment.	
Eleventh GDG Meeting [10/04/13]	Personal pecuniary interest : Will be speaking at Primary Care 2013; expenses to be reimbursed.	No action necessary.

2 B.11 Expert Advisors

3 B.11.1 Dr. Christine Evans

GDG meeting	Declaration of Interests	Action taken
On Application	None	None

4 B.11.2 Ms. Jenny Greenfield

GDG meeting	Declaration of Interests	Action taken
On application	None	None
First GDG meeting [21/09/11]	None	None
Second GDG Meeting [02/11/11]	None	None
Third GDG Meeting [14/12/11]	None	None
Fourth GDG Meeting [25/01/12]	None	None
Fifth GDG Meeting [07/03/12]	None	None
Sixth GDG Meeting [18/04/12]	Did not attend	None
Seventh GDG Meeting [11/07/12]	None	None
Eighth GDG Meeting [03/10/12]	Did not attend this meeting	None
Ninth GDG	None	None

GDG meeting	Declaration of Interests	Action taken
Meeting		
[07/11/12]		

2 B.11.3 Ms. Janine Elson

GDG meeting	Declaration of Interests	Action taken
On application	None	None
Ninth GDG Meeting [07/11/12]	Did not attend this meeting	None

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Appendix C: Review protocols

2 C.1 Chapter 5 – patient perceptions and expectations

Review question	What are the perceptions and expectations of people with varicose veins (e.g. natura history, treatment) and how can they be addressed?
Objectives	 To identify the perceptions and expectations of people with varicose veins with regators. Risk factors for developing varicose veins Progression of varicose veins Expectations about treatment To identify what information should be given to people with varicose veins in order to manage expectations and perceptions To identify how information should be given
Setting	Primary and secondary care
Population	Adults with leg varicose veins.
Intervention	NA
Comparison	NA
Outcomes	Any perceptions and expectations that are identified by people with varicose veins about their condition including those before and after treatment. How people with varicose veins would like to receive information on their condition.
Evaluation	Narrative summary of findings on patient perceptions and expectations related to the assessment, treatment, treatment success/failure, retreatment, adverse events and disease progression of varicose veins. Studies suggesting how such expectations can be addressed were also evaluated.
Exclusion	Studies that do not specify a varicose veins population. Opinion papers
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL and Psych Lit. Studies will be restricted to English language only. We will look for studies collecting data on patient expectations and perceptions related the assessment, treatment, treatment success/failure, retreatment, adverse events and disease progression of varicose veins including risk factors for development of varicose veins.
	We will also look for studies suggesting how such expectations can be addressed.
The review strategy	Qualitative studies and questionnaire surveys will be searched If there are no published opinions on how expectations can be addressed, that part of question will be answered solely by GDG consensus.
Key papers	Shepherd AC et al. Phlebology 2010; 25: 54-65 Darvall KA et al. Euro J Vasc Endovasc Surg 2009; 38: 642-647 Palfreyman SJ et al. J Clin Nurs 2004; 13: 332-340

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1 C.2 Chapter 6 – referral to a vascular service

2 C.2.1 Factors associated with disease progression

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Table 2: Revi	ew protocol: factors associated with disease progression
Review question	 a. In people with leg varicose veins at CEAP class C2 which signs, symptoms and/or patient characteristics are associated with disease progression to i) C3, ii) C4*, iii) C6? b. In people with leg varicose veins at CEAP class C3 which signs, symptoms and/or patient characteristics are associated with disease progression to i) C4*, ii) C6? c. In people with leg varicose veins at CEAP class C4* which signs, symptoms and/or patient characteristics are associated with disease progression to i) C4*, iii) C6? c. In people with leg varicose veins at CEAP class C4* which signs, symptoms and/or patient characteristics are associated with disease progression to C6?
	* Will separate out CEAP classes C4a and C4b where evidence exists
Population	Adults with leg varicose veins at CEAP stage C2 OR C3 OR C4 [as in parts a), b) and c) of the clinical question]
Prognostic	Clinical signs that can be assessed by a non-vascular specialist:
Factors	Location/extent of varicose veins
	 Any other aspects of physical examination
	Clinical symptoms that can be assessed by a non-vascular specialist: • Severity of pain
	 Severity of other varicose veins symptoms
	• Sevency of other varicose vents symptoms
	Patient characteristics that can be assessed by a non-vascular specialist:
	• Age
	• BMI
	Comorbidities
	 Pregnancy/no of previous pregnancies
	 Past history of deep vein thrombosis (DVT)
	Recurrent varicose veins
Outcomes	Progression to the CEAP class endpoints defined by parts a), b) or c) of the clinical question
Exclusion	Studies that do not specify a varicose veins population.
	 Stratify studies with people who have previously-treated varicose veins
	• Exclude studies where follow-up was less than 1 year
Search	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL
strategy	Studies will be restricted to English Language only
The review strategy	 Where studies based on individual patient data (pooled analysis) are available, these are reviewed and other type of evidence such as meta-analysis, systematic reviews, prospective cohorts/case-control and cross-sectional studies are not included. Hierarchy of evidence (only go down a level if there is a lack of evidence): Pooled analysis of patient level data Meta-analysis/systematic reviews Cohort Studies Other observational studies
Key papers	Bonn vein studies, Nelson, NZ data, Framingham study

1 C.2.2 Factors associated with treatment success

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Table 3: Review protocol: factors associated with treatment success

Review question	In people with leg varicose veins are there any factors (clinical signs and symptoms or patient reported outcomes) that would predict increased benefits or harms from varicose veins interventional treatments?
Population	Adults with leg varicose veins
Prognostic Factors	 Clinical signs and symptoms that can be assessed by a non-vascular specialist: Any aspects of physical examination (CEAP) Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/cosmesis, swelling (oedema), aching, heaviness.)
	Patient characteristics that can be assessed by a non-vascular specialist:
	• Age
	• BMI
	Comorbidities
	• Parity
	Recurrent varicose veins
	 Medical history (including family history)
	Patient reported outcomes that can be assessed by a non-vascular specialist:
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D)
	• disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score).
Outcomes	Patient-reported outcome:-
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score).
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/cosmesis, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score, CEAP)
	Presence of reflux:
	Within 3 months
	• >3–12 months
	• >1–5 years
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above
	Immediate: Within 3 months post intervention
	 Intermediate: >3–12 months post intervention
	 Long term: >1–5 years post intervention
	Adverse events from intervention (including venous thrombo-embolism [VTE], i.e. pulmonary embolism [PE] and deep vein thrombosis (DVT); major neurological event (i.e. stroke); local neurological events, i.e. nerve injury/damage, paraesthesia, neuralgia,

numbness; post-procedure pain; phlebitis; skin pigmentation/discolouration.

Review question	In people with leg varicose veins are there any factors (clinical signs and symptoms or patient reported outcomes) that would predict increased benefits or harms from varicose veins interventional treatments?
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis. Return to work/normal activities
Exclusion	Studies that do not specify a varicose veins population. Studies not using a multivariable analysis
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL Studies will be restricted to English Language only
The review strategy	 Where studies based on individual patient data (pooled analysis) are available, these are reviewed and other type of evidence such as meta-analysis, systematic reviews, prospective cohorts/case-control and cross-sectional studies are not included. Hierarchy of evidence (only go down a level if there is a lack of literature): Pooled analysis of patient level data Meta-analysis/systematic reviews Cohort Studies Other observational studies
Analysis	 Stratification will occur by treatment type Stripping Foam sclerotherapy Endothermal ablation.

2 C.3 Chapter 7 – assessment for treatment

3 C.3.1 Diagnostic accuracy of hand held Doppler

Table 4: Review protocol: Diagnostic accuracy of hand held Doppler

Review question	What is the diagnostic accuracy of hand held Doppler (HHD) compared to Duplex scanning when used in patients with varicose veins?
Population	Adults with leg varicose veins.
Index tests	Hand held Doppler ultrasound testing for venous reflux
Reference standard	Duplex ultrasound scanning for venous reflux
Outcomes	Main outcomes:
	 Sensitivity (%) and specificity (%), for particular threshold(s)
	 Area under the ROC curve (AUC) – measure of predictive accuracy
	Other outcomes:
	Positive/negative predictive value
	 Positive/ negative diagnostic likelihood ratios
	 Post-test probability (at a set pre-test probability)
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL.
	Studies will be restricted to English language only.
The review strategy	Diagnostic studies

Review question	What is the diagnostic accuracy of hand held Doppler (HHD) compared to Duplex scanning when used in patients with varicose veins?
Analysis	We will analyse the diagnostic accuracy of hand held Doppler ultrasound.
	We will note at what point in the patient pathway the study is done.

1 C.3.2 Assessment with duplex prior to interventional treatment

Table 5: Review protocol: Duplex vs. no duplex prior to interventional treatment

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Review question	Does the use of duplex ultrasound during assessment improve outcome after interventional treatment compared to no duplex scanning in people with leg varicose veins?
Population	Adults with leg varicose veins.
Intervention	Duplex ultrasound assessment prior to interventional treatment (surgery, endothermal ablation or foam sclerotherapy)
Comparison	No duplex ultrasound assessment prior to interventional treatment
Outcomes	Patient-reported outcome:-
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score) <n.b. only="" overall="" scores="" to="" use=""></n.b.>
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/cosmesis, swelling, aching, heaviness).
	Physician-reported outcomes (venous clinical severity score or venous disability score). <n.b. only="" overall="" scores="" to="" use=""></n.b.>
	 Presence of reflux: Within 3 months >3–12 months >1–5 years <<i>N.B. if no reflux data is available, to include incomplete impartial occlusion/incomplete stripping rates within the same analysis></i> Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: Immediate: Within 3 months post intervention Intermediate: >3–12 months post intervention Long term: >1–5 years post intervention Adverse events from intervention including: venous thromboembolism [VTE], i.e. pulmonary embolism [PE] and deep vein thrombosis
	 (DVT); major neurological event (i.e. stroke); local neurological events, i.e. nerve injury/damage, paraesthesia, neuralgia, numbness; post-procedure pain; phlebitis; skin pigmentation/discolouration. Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis. Return to work/normal activities
Exclusion	Studies that do not specify a varicose veins population. Studies that compare different interventions as well as the use/no use of duplex
Search	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL.

Review question	Does the use of duplex ultrasound during assessment improve outcome after interventional treatment compared to no duplex scanning in people with leg varicose veins?
strategy	Studies will be restricted to English language only.
The review strategy	Systematic reviews RCTs Non-randomised clinical trials
Analysis	We should stratify by the different interventional treatments used in the different studies, as the difference between use of duplex and no duplex may differ depending on which treatment is subsequently used (for example, the use of duplex may be important in optimising surgical outcomes, but may be less important with thermal ablation). Sub-grouping will occur if there is statistical heterogeneity in meta-analysis results. Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6).
	Sub-group by primary and recurrent varicose veins.

1 C.4 Chapter 8 – conservative management

2 C.4.1 Conservative treatment vs. no treatment

Table 6: Review protocol: compression vs. no treatment/lifestyle advice	
Review question	What is the clinical and cost effectiveness of compression therapy compared with no treatment or lifestyle advice in people with leg varicose veins?
Population	Adults with varicose veins in the legs
Intervention	Compression therapy, specifically compression hosiery (compression stockings)
	Both above knee and below knee compression hosiery will be included.
	[There will be no comparison between types of compression therapy].
Comparison	• no treatment, or
	non-compressive stockings, orplacebo, or
	 lifestyle advice (including advice on weight loss, exercise, smoking, occupational standing/leg elevation, etc.)
Outcomes	Patient-reported outcome
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score).
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/body image, swelling (oedema), aching, heaviness.)
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods:
	 Immediate: ≤1 month post intervention
	 Intermediate: >1month up to 12 months post intervention
	 Long term: >12 months up to 5 years post intervention
	Adverse events from intervention including:
	 manifestations of reduced arterial flow,
	skin pressure damage
	• ulceration,
	allergic reactions,
	 blistering, discomfort, a constitute of successive ticktoore
	a sensation of excessive tightness.Also non-compliance, and withdrawal from study due to adverse effects
	• Also non-compliance, and withdrawal nom study due to adverse effects
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.
Exclusion	Compression therapy applied after an interventional procedure (i.e. after sclerotherapy).
	Compression or bandaging applied for the management of venous ulcers (i.e. C6)
	Pneumatic intermittent compression.
	Studies that do not specify a varicose veins population

Review question	What is the clinical and cost effectiveness of compression therapy compared with no treatment or lifestyle advice in people with leg varicose veins?
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL.
	Studies will be restricted to English language only
Search terms	Elastic stockings
	(graduated) Compression therapy/hosiery/stockings
The review strategy	Systematic reviews
	RCTs (cross over trials will be included where the time of treatment was short enough not to result in a natural change in patient condition and the washout period long enough to negate any impact of the stockings).
	If no RCTs, then conference abstracts, and then observational studies.
Analysis	Stratification for studies focussed on pregnant women.
	A meta-analysis will be conducted on RCTs with appropriate outcome data.
	Sub-grouping will occur if there is statistical heterogeneity in meta-analysis results:
	Sub-group for disease stage (i.e. CEAP classification, C2, C3, C4, C5).
	Sub group for above and below knee hosiery

1 C.4.2 Compression vs. interventional treatment

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Table 7: Review protocol: compression vs. interventional treatment

	What is the clinical and cost effectiveness of compression therapy compared with
	a) stripping surgery; or
	b) endothermal ablation; or
Review	c) foam sclerotherapy
question	in people with leg varicose veins?
Population	Adults with varicose veins in the legs
Intervention	Compression therapy, specifically compression hosiery (compression stockings)
	Both above knee and below knee compression hosiery will be included.
	[There will be no comparison between types of compression therapy].
Comparison	Foam sclerotherapy \pm crossectomy OR
	Stripping surgery + ligation [± phlebectomy]
	OR
	Endothermal ablation [± foam sclerotherapy/phlebectomy]
Outcomes	Patient-reported outcomes
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score).
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/body image, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods:
	 Immediate: ≤1 month post intervention
	 Intermediate: >1month up to 12 months post intervention
	• Long term: >12 months up to 5 years post intervention
	Adverse events from intervention including:
	 manifestations of reduced arterial flow,
	• major vascular injury,
	• skin pressure damage,
	• ulceration,
	• allergic reactions,
	• blistering,
	• discomfort,
	 sensation of excessive tightness.
	 venous thromboembolism (pulmonary embolism [PE] and deep vein thrombosis (DVT);
	• Central neurological event (permanent (i.e. stroke, TIA) and transient i.e. migraine, transient visual disturbance);
	 local neurological events (permanent and transient) i.e. nerve injury/damage, paraesthesia, neuralgia, numbness;
	• post-procedure pain;
	• phlebitis;

Review question	 What is the clinical and cost effectiveness of compression therapy compared with a) stripping surgery; or b) endothermal ablation; or c) foam sclerotherapy in people with leg varicose veins?
	skin pigmentation/discolouration.Also non-compliance, and withdrawal from study due to adverse effects)
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis. Return to work and/or normal activities
Exclusion	Studies that do not specify a varicose veins population Compression therapy applied after an interventional procedure (i.e. after sclerotherapy). Compression or bandaging applied for the management of venous ulcers (i.e. C6) Pneumatic intermittent compression. Cryostripping
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL. Studies will be restricted to English language only
The review strategy	RCTs first. If no RCTs, then conference abstracts, and then observational studies. Systematic reviews
Analysis	A meta-analysis will be conducted on RCTs with appropriate outcome data. Stratification for studies focussed on pregnant women.
	Sub-grouping will occur if there is statistical heterogeneity in meta-analysis results: Sub-group for disease stage (i.e. CEAP classification, C2, C3, C4, C5). Sub group for above and below knee hosiery

1 C.5 Chapter 9 – interventional treatment

2 C.5.1 Stripping surgery vs. foam sclerotherapy

Review question	What is the clinical and cost effectiveness of stripping surgery compared with foam sclerotherapy in people with truncal leg varicose veins?
Population	Adults with truncal leg varicose veins.
•	-
Intervention	 Stripping surgery (including conventional stripping, invagination stripping=inverting stripping=PIN [perforation invagination], 'high-tie'=crossectomy, saphenofemoral junction disconnection, saphenopopliteal) with ligation, sequential stripping surgery. [± phlebectomy] [NOTE: Stripping surgery comes hand-in-hand with ligation, i.e. it is normal practice for ligation to occur before stripping]
Comparison	Foam sclerotherapy
	[± crossectomy (ligation)]
	[NOTE: compression therapy is applied after the procedure as part of the treatment]
Outcomes	Patient-reported outcome:-
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36 EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score).
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/body language, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Presence of reflux:
	Within 3 months
	• >3–12 months
	 >1–5 years
	<n.b. analysis="" available,="" data="" if="" impartial="" include="" incomplete="" is="" no="" occlusion="" rates="" reflux="" same="" stripping="" the="" to="" within=""></n.b.>
	 Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above Immediate: Within 3 months post intervention
	 Intermediate: >3–12 months post intervention
	• Long term: >1–5 years post intervention
	Adverse events from intervention including:
	 venous thrombo-embolism [VTE], i.e. pulmonary embolism [PE] and deep vein thrombosis (DVT);
	 major neurological event (i.e. stroke);
	• local neurological events, i.e. nerve injury/damage, paraesthesia, neuralgia, numbness;
	• post-procedure pain;
	• phlebitis;
	 skin pigmentation/discolouration.
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.

	Return to work/normal activities
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL. Studies will be restricted to English language only.
The review strategy	RCTs first. If no RCTs are available, then consider conference abstracts, and if none are available then consider observational studies. Systematic reviews
Analysis	 A meta-analysis will be conducted on RCTs with appropriate outcome data. Stratification from the outset: foam sclerotherapy ± crossectomy (i.e. ligation). primary and recurrent varicose veins Further sub-grouping will occur if there is statistical heterogeneity in meta-analysis results. Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6).
Key papers	Murad MH, Coto-Yglesias F, Zumaeta-Garcia M, Elamin MB, Duggirala MK, Erwin PJ, Montori VM, and Gloviczki P. A systematic review and meta-analysis of the treatments of varicose veins. [Review]. Journal of Vascular Surgery 2011; 53: 49S - 65S Rigby KA, Palfreyman SJ, Beverley C, Michaels JA. Surgery versus sclerotherapy for the treatment of varicose veins. Cochrane Database Syst Rev. 2004;18;(4):CD004980.

1 C.5.2 Stripping surgery vs. endothermal ablation

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Table 9: Review protocol: stripping surgery vs. endothermal ablation

Review question	What is the clinical and cost effectiveness of stripping surgery compared with endothermal ablation in people with truncal leg varicose veins?
Population	Adults with truncal leg varicose veins.
Intervention	 Stripping surgery (including conventional stripping, invagination stripping=inverting stripping=PIN [perforation invagination], 'high-tie'=crossectomy, saphenofemoral junction disconnection, saphenopopliteal) with ligation, sequential stripping surgery. [± phlebectomy] [NOTE: Stripping surgery comes hand-in-hand with ligation, i.e. it is normal practice for
	ligation to occur before stripping]
Comparison	Endothermal ablation, including:radiofrequency ablation
	(endovenous) laser ablation (EVLA)
	steam ablation
	[± foam sclerotherapy/phlebectomy (for tributaries)]
Outcomes	Patient-reported outcome:-
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score).
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/body language, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Presence of reflux:

Review	What is the clinical and cost effectiveness of stripping surgery compared with
question	endothermal ablation in people with truncal leg varicose veins?
	Within 3 months
	• >3–12 months
	• >1–5 years
	<n.b. analysis="" available,="" data="" if="" impartial="" include="" incomplete="" is="" no="" occlusion="" rates="" reflux="" same="" stripping="" the="" to="" within=""></n.b.>
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above
	Immediate: Within 3 months post intervention
	 Intermediate: >3–12 months post intervention
	 Long term: >1–5 years post intervention
	Adverse events from intervention including:
	 venous thrombo-embolism [VTE], i.e. pulmonary embolism [PE] and deep vein thrombosis (DVT);
	 major neurological event (i.e. stroke);
	 local neurological events, i.e. nerve injury/damage, paraesthesia, neuralgia, numbness;
	• post-procedure pain;
	• phlebitis;
	 skin pigmentation/discolouration.
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.
	Return to work/normal activities
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL. Studies will be restricted to English language only.
The review strategy	RCTs first. If no RCTs are available, then consider conference abstracts, and if none are available then consider observational studies. Systematic reviews
Analysis	A meta-analysis will be conducted on RCTs with appropriate outcome data.
	Stratification from the outset:
	primary and recurrent varicose veins
	Sub-grouping will occur if there is statistical heterogeneity in meta-analysis results. Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6). Sub-group by types of endothermal ablation
Key naners	van den Bos R, Arends L, Kockaert M, Neumann M, and Nijsten T. Endovenous therapies of
Key papers	lower extremity varicosities: a meta-analysis. [Review] [42 refs]. Journal of Vascular Surgery 2009; 49: 230 - 239

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Foam sclerotherapy vs. endothermal ablations 1 C.5.3

Table 10: Review protocol: foam sclerotherapy versus endothermal ablation

Table 10: Review protocol: foam sclerotherapy versus endothermal ablation		
Review	What is the clinical and cost effectiveness of foam sclerotherapy compared with	
question	endothermal ablation in people with truncal leg varicose veins?	
Population	Adults with truncal leg varicose veins.	
Intervention	Foam sclerotherapy(including ultrasound-guided foam sclerotherapy (UGFS))	
	$[\pm \text{ crossectomy (ligation)}]$	
	[NOTE: compression therapy is applied after the procedure as part of the treatment]	
Comparison	Endothermal ablation, including:	
	radiofrequency ablation	
	(endovenous) laser ablation (EVLA)	
	steam ablation	
	[foam sclerotherapy/phlebectomy (for tributaries)]	
Outcomes	Patient-reported outcome:-	
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score) 	
	• Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/body image, swelling (oedema), aching, heaviness.	
	Physician-reported outcomes (venous clinical severity score or venous disability score).	
	Presence of reflux:	
	Within 3 months	
	• >3–12 months	
	• >1–5 years	
	<n.b. analysis="" available,="" data="" if="" impartial="" include="" incomplete="" is="" no="" occlusion="" rates="" reflux="" same="" stripping="" the="" to="" within=""></n.b.>	
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above	
	Immediate: Within 3 months post intervention	
	 Intermediate: >3–12 months post intervention 	
	 Long term: >1–5 years post intervention 	
	Adverse events from intervention including:	
	 venous thrombo-embolism [VTE], i.e. pulmonary embolism [PE] and deep vein thrombosis (DVT); 	
	 major neurological event (i.e. stroke); 	
	 local neurological events, i.e. nerve injury/damage, paraesthesia, neuralgia, numbness; 	
	post-procedure pain;	
	phlebitis;	
	 skin pigmentation/discolouration. 	
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.	

Return to work/normal activities

Review	What is the clinical and cost effectiveness of foam sclerotherapy compared with
question	endothermal ablation in people with truncal leg varicose veins?
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL.e Studies will be restricted to English language only.
The review strategy	RCTs first. If no RCTs are available, then consider conference abstracts of RCTs, and if none are available then consider observational studies. Systematic reviews
Analysis	 A meta-analysis will be conducted on RCTs with appropriate outcome data. Stratification from the outset: foam sclerotherapy ± crossectomy (i.e. ligation). primary and recurrent varicose veins Sub-grouping will occur if there is statistical heterogeneity in meta-analysis results. Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6). Sub-group by types of endothermal ablation.

1 C.5.4 Tributary treatment: avulsion surgery vs. foam sclerotherapy

2

Table 11: Review protocol: avulsion surgery vs. foam sclerotherapy for tributary treatment

Review question	What is the clinical and cost effectiveness of avulsion surgery compared with foam sclerotherapy in people with tributary leg varicose veins?
Population	Adults with tributary leg varicose veins.
Intervention	Avulsion surgery (ambulatory phlebectomy, phlebectomy)
Comparison	Foam sclerotherapy to tributary veins. (including ultrasound-guided foam sclerotherapy (UGFS)) [NOTE: compression therapy is applied after the procedure as part of the treatment]
Outcomes	Patient-reported outcome:-
Outcomes	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score)
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/body image, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Presence of reflux:
	Within 3 months
	• >3–12 months
	• >1–5 years
	<n.b. analysis="" available,="" data="" if="" impartial="" include="" incomplete="" is="" no="" occlusion="" rates="" reflux="" same="" stripping="" the="" to="" within=""></n.b.>
	 Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above Immediate: Within 3 months post intervention Intermediate: >3-12 months post intervention

Review question	What is the clinical and cost effectiveness of avulsion surgery compared with foam sclerotherapy in people with tributary leg varicose veins?
	• Long term: >1–5 years post intervention
	Adverse events from intervention including:
	 venous thrombo-embolism [VTE], i.e. pulmonary embolism [PE] and deep vein thrombosis (DVT);
	 major neurological event (i.e. stroke);
	 local neurological events, i.e. nerve injury/damage, paraesthesia, neuralgia, numbness;
	 post-procedure pain;
	• phlebitis;
	 skin pigmentation/discolouration.
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.
	Return to work/normal activities
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL. Studies will be restricted to English language only.
The review strategy	RCTs first. If no RCTs are available, then consider conference abstracts, and if none are available then consider observational studies. Systematic reviews
Analysis	A meta-analysis will be conducted on RCTs with appropriate outcome data.
	Within the above sub-groups, further sub-grouping will occur if there is statistical heterogeneity in meta-analysis results.
	Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6).
	Sub-group by primary and recurrent varicose veins.

1 C.5.5 Truncal treatment and tributary treatment vs. truncal treatment alone

2

 Table 12: Review protocol: Truncal treatment and tributary treatment vs. truncal treatment alone

Review question	What is the clinical and cost effectiveness of truncal treatment accompanied by tributary treatment (avulsion or sclerotherapy) compared with truncal treatment alone in people with leg varicose veins?
Population	Adults with leg varicose veins.
Intervention	Stripping surgery ¹ accompanied by tributary treatments (avulsion ² / foam sclerotherapy ³) OR
	Endothermal ablation ⁴ accompanied by tributary treatments (avulsion ² / foam sclerotherapy ³)
	OR
	Foam sclerotherapy ³ accompanied by tributary treatments (avulsion ² / foam sclerotherapy ³)
	1: Stripping surgery (including conventional stripping, invagination stripping=inverting stripping=PIN [perforation invagination], 'high-tie'=crossectomy, sapheno-femoral junction disconnection, sapheno-popliteal) with ligation, sequential stripping surgery.[± phlebectomy]
	2: Avulsion surgery (ambulatory phlebectomy, phlebectomy)

Review question	What is the clinical and cost effectiveness of truncal treatment accompanied by tributary treatment (avulsion or sclerotherapy) compared with truncal treatment alone in people with leg varicose veins?
	 3: Foam sclerotherapy 4: Endothermal ablation, including: radiofrequency ablation (endovenous) laser ablation (EVLA, EVLT) steam ablation [± foam sclerotherapy/phlebectomy (for tributaries)]
Comparison	The comparator in each case will be the truncal intervention, but without tributary treatment (avulsion / sclerotherapy) as an adjunct
Outcomes	Patient-reported outcomes:-
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score)
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/cosmesis*, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Presence of reflux:
	Within 3 months
	• >3–12 months
	• >1–5 years
	<n.b. analysis="" available,="" data="" if="" impartial="" include="" incomplete="" is="" no="" occlusion="" rates="" reflux="" same="" stripping="" the="" to="" within=""></n.b.>
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above
	Immediate: Within 3 months post intervention
	 Intermediate: >3–12 months post intervention
	 Long term: >1–5 years post intervention
	Adverse events from intervention, including:
	 venous thromboembolism (pulmonary embolism [PE] and deep vein thrombosis (DVT – to be reported separately)
	 global neurological event (i.e. stroke, TIA);
	 local neurological events (i.e. nerve injury/damage, paresthesia, neuralgia, numbness).
	post-procedure pain
	phlebitis
	skin pigmentation/discolouration.
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.
	Return to work / normal activities
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL. Studies will be restricted to English language only.
The review	RCTs first. If no RCTs are available, then consider conference abstracts, and if none are

Review question	What is the clinical and cost effectiveness of truncal treatment accompanied by tributary treatment (avulsion or sclerotherapy) compared with truncal treatment alone in people with leg varicose veins?
strategy	available then consider observational studies. Systematic reviews
Analysis	 A meta-analysis will be conducted on RCTs with appropriate outcome data. Stratification from the outset: foam sclerotherapy ± crossectomy (i.e. ligation). Within the above sub-groups, further sub-grouping will occur if there is statistical heterogeneity in meta-analysis results. Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6). Sub-group by primary and recurrent varicose veins.

3

4

2 C.6 Chapter 10 – compression after treatment

Table 13: Review protocol: compression after interventional treatment vs. interventional treatment alone

Review question	What is the clinical and cost effectiveness of interventional treatment followed by compression compared with interventional treatment alone in people with leg varicose veins, and, if so, what type of compression, pressure of compression and/or duration of compression is optimal?
Population	Adults with leg varicose veins.
Intervention	Stripping surgery ¹ followed by compression ² OR Avulsion surgery ³ followed by compression ² OR Endothermal ablation ⁴ followed by compression ² OR Foam sclerotherapy ⁵ followed by compression ² 1: Stripping surgery (including conventional stripping, invagination stripping=inverting stripping=PIN [perforation invagination], 'high-tie'=crossectomy, saphenofemoral junction disconnection, saphenopopliteal) with ligation, sequential stripping surgery.[± phlebectomy]. Note: Short-term (up to 7 days) 'routine' elastic bandaging is allowed in both groups. 2: Compression therapy, specifically compression hosiery (compression stockings). Both above knee and below knee compression hosiery will be included. 3: Avulsion surgery (ambulatory phlebectomy, phlebectomy) Note: Short-term (up to 5 days) 'routine' elastic bandaging is allowed in both groups. 4: Endothermal ablation, including: • radiofrequency ablation • (endovenous) laser ablation (EVLA, EVLT) • steam ablation

	What is the clinical and cost effectiveness of interventional treatment followed by compression compared with interventional treatment alone in people with leg varicose
Review question	veins, and, if so, what type of compression, pressure of compression and/or duration of compression is optimal?
question	[± foam sclerotherapy/phlebectomy (for tributaries)]
	Note: Short-term (up to 5 days) 'routine' elastic bandaging is allowed in both groups.
	5: Foam sclerotherapy[\pm crossectomy (ligation)] Note: Short-term (up to 5 days) 'routine' elastic bandaging is allowed in both groups.
Comparison	For the first part of the review question, the comparator in each case will be as the intervention, but without compression as an adjunct.
	For the second part of the review question, the comparator will be as the intervention but adjunctive compression will vary in terms of:
	 another type of compression (i.e. bandaging)
	a different compression pressure
	a different duration of treatment
Outcomes	Patient-reported outcome:-
	 Health-related quality of life, using generic (e.g. Medical Outcomes Study Short Form 36, EQ-5D) and disease specific validated tools (e.g. Chronic Venous Insufficiency Questionnaire, Aberdeen Varicose Vein Symptom Severity Score)
	 Patient-assessed symptoms (including pain, discomfort, cosmetic concerns/cosmesis*, swelling (oedema), aching, heaviness.
	Physician-reported outcomes (venous clinical severity score or venous disability score).
	Presence of reflux:
	Within 3 months
	• >3–12 months
	• >1–5 years
	<n.b. analysis="" available,="" data="" if="" impartial="" include="" incomplete="" is="" no="" occlusion="" rates="" reflux="" same="" stripping="" the="" to="" within=""></n.b.>
	Need for additional/further treatment (i.e. compression therapy and/or ablative techniques) over the following time periods: (same time intervals as above
	Immediate: Within 3 months post intervention
	 Intermediate: >3–12 months post intervention
	 Long term: >1–5 years post intervention
	Adverse events from intervention, including:
	 venous thrombo-embolism (i.e. pulmonary embolism [PE] and deep vein thrombosis (DVT) to be reported separately)
	global neurological event (i.e. stroke);
	 local neurological events (i.e. nerve injury/damage, paresthesia, neuralgia, numbness). post-procedure pain
	• phlebitis
	skin pigmentation/discolouration.
	Prevention of complications from varicose veins (leg ulcer occurrence or recurrence, haemorrhage (bleeding) and thrombophlebitis.

Review question	What is the clinical and cost effectiveness of interventional treatment followed by compression compared with interventional treatment alone in people with leg varicose veins, and, if so, what type of compression, pressure of compression and/or duration of compression is optimal?
	Return to work / normal activities
Exclusion	Studies that do not specify a varicose veins population.
Search strategy	The databases to be searched are Medline, Embase, The Cochrane Library, CINAHL. Studies will be restricted to English language only.
The review strategy	RCTs first. If no RCTs are available, then consider conference abstracts, and if none are available then consider observational studies. Systematic reviews
Analysis	A meta-analysis will be conducted on RCTs with appropriate outcome data. Within the above sub-groups, further sub-grouping will occur if there is statistical heterogeneity in meta-analysis results. Sub-group by disease stage (i.e. CEAP classification C2, C3, C4, C5, C6). Sub-group by primary and recurrent varicose veins.

2

1 C.7 Economic review protocol

Table 14: Appended economic review protocol

Review	
question	All questions – health economic evidence
Objectives	To identify economic studies relevant to the review questions set out above.
Criteria	Populations, interventions and comparators as specified in the individual review protocols above. Must be a relevant economic study design (cost-utility analysis, cost-benefit analysis, cost-effectiveness analysis, cost-consequence analysis, comparative cost analysis).
Search strategy	An economic study search was undertaken using population specific terms and an economic study filter – see Appendix F
Review strategy	Each study is assessed using the NICE economic evaluation checklist – NICE (2009) Guidelines Manual.

Inclusion/exclusion criteria

If a study is rated as both 'Directly applicable' and 'minor limitations' (using the NICE economic evaluation checklist) then it should be included in the guideline. An evidence table should be completed and it should be included in the economic profile.

If a study is rated as either 'Not applicable' or 'Very serious limitations' then it should be excluded from the guideline. It should not be included in the economic profile and there is no need to include an evidence table.

If a study is rated as 'Partially applicable' and/or 'potentially serious limitations' then there is discretion over whether it should be included. The health economist should make a decision based on the relative applicability and quality of the available evidence for that question, in discussion with the GDG if required. The ultimate aim being to include studies that are helpful for decision making in the context of the guideline and current NHS setting. Where exclusions occur on this basis, this should be noted in the relevant section of the guideline with references.

Also exclude:

- unpublished reports unless submitted as part of a call for evidence
- abstract-only studies
- letters
- editorials
- reviews of economic evaluations^a
- foreign language articles

Where there is discretion

The health economist should be guided by the following hierarchies:

Setting:

- 1. UK NHS
- 2. OECD countries with predominantly public health insurance systems (e.g. France, Germany, Sweden)
- 3. OECD countries with predominantly private health insurance systems (e.g. USA, Switzerland)
- 4. Non-OECD settings (always 'Not applicable')

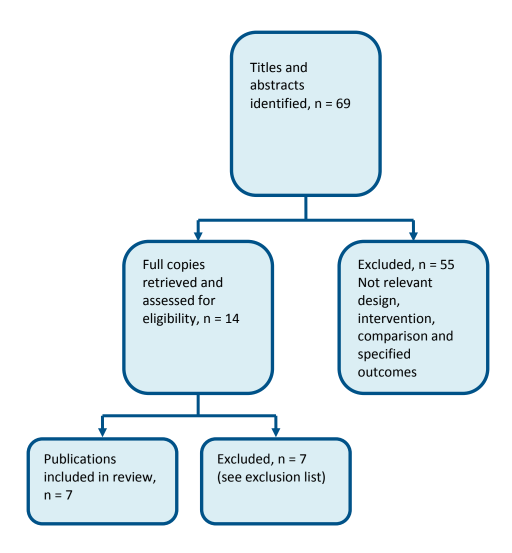
Economic study type:

Review	
question	All questions – health economic evidence
	1. Cost-utility analysis
	 Other type of full economic evaluation (cost-benefit analysis, cost-effectiveness analysis, cost-consequence analysis)
	3. Comparative cost analysis
	 Non-comparative cost analyses including cost of illness studies (always 'Not applicable')
	Year of analysis:
	The more recent the study, the more applicable it is
	Quality and relevance of effectiveness data used in the economic analysis:
	 The more closely the effectiveness data used in the economic analysis matches with the studies included for the clinical review the more useful the analysis will be to decision making for the guideline.
(a) Recent revi then be ord	iews will be ordered although not reviewed. The bibliographies will be checked for relevant studies, which will dered.

Appendix D: Clinical article selection

2 D.1 Chapter 5 – patient perceptions and expectations

Figure 1: Clinical article selection: patient perceptions and expectations

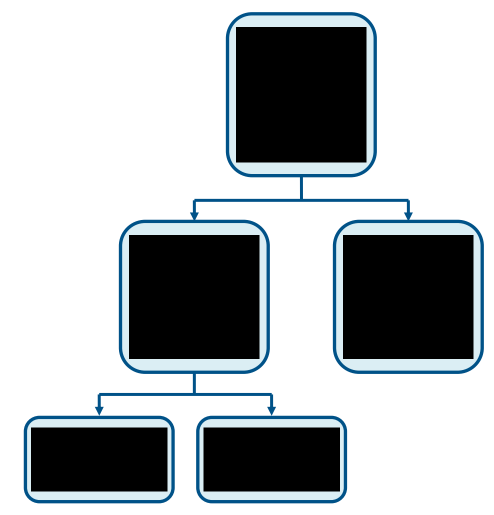


3

D.2 Chapter 6 – referral to a vascular service

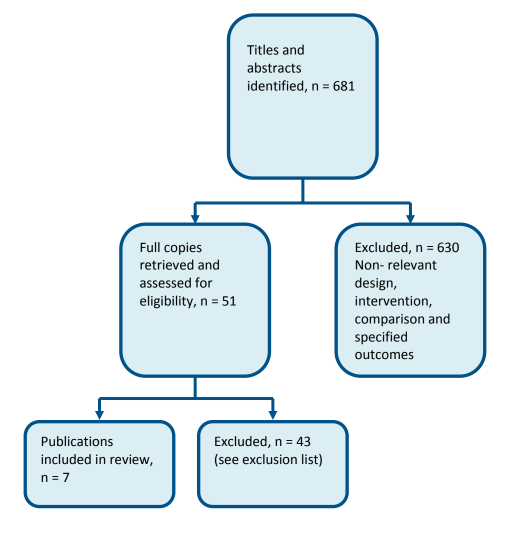
2 D.2.1 Risk factors associated with disease progression

Figure 2: Clinical article selection: factors associated with disease progression



1 D.2.2 Risk factors affecting treatment success

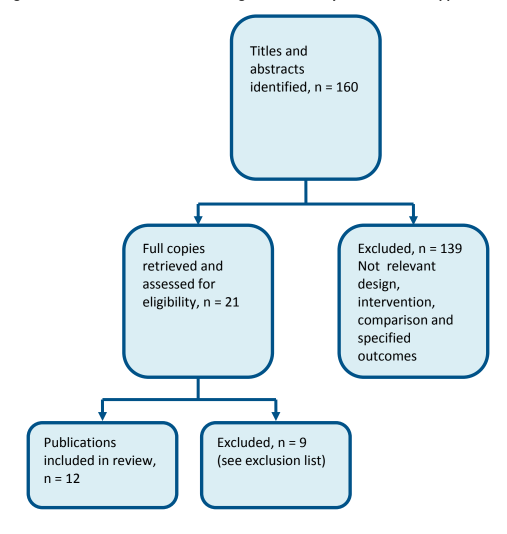
2 Figure 3: Clinical article selection: factors associated with treatment success/failure



1 D.3 Chapter 7 – assessment for treatment

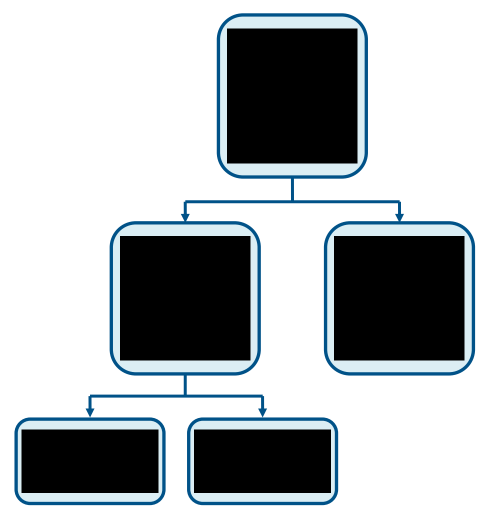
2 D.3.1 Diagnostic accuracy of hand held Doppler

3 Figure 4: Clinical article selection: diagnostic accuracy of hand held Doppler



1 **D.3.2** Assessment with duplex prior to interventional treatment

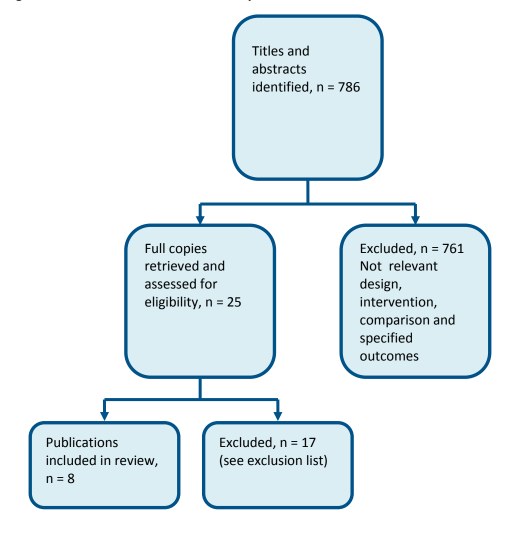
Figure 5: Clinical article selection: assessment with duplex prior to interventional treatment



1 D.4 Chapter 8 – conservative management

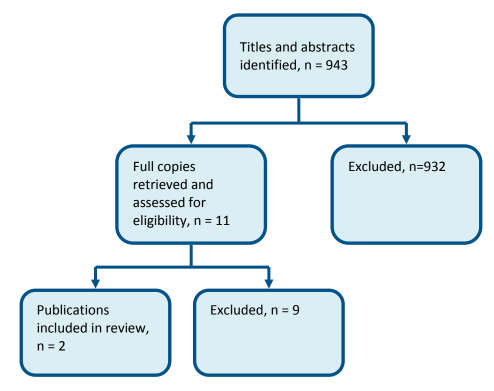
2 D.4.1 Compression vs. no treatment

3 Figure 6: Clinical article selection: compression vs. no treatment



1 D.4.2 Compression vs. interventional treatment

2 Figure 7: Clinical article selection for compression vs. interventional treatment



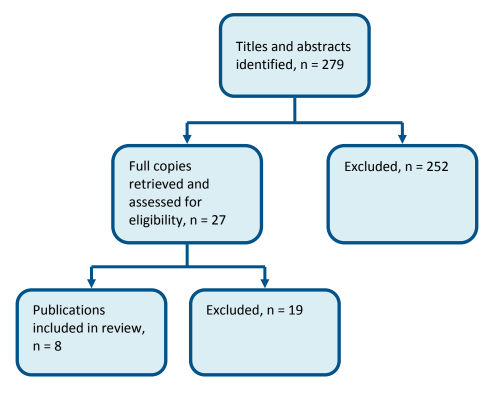
3

7

4 D.5 Chapter 9 – interventional treatment

5 D.5.1 Stripping surgery vs. foam sclerotherapy

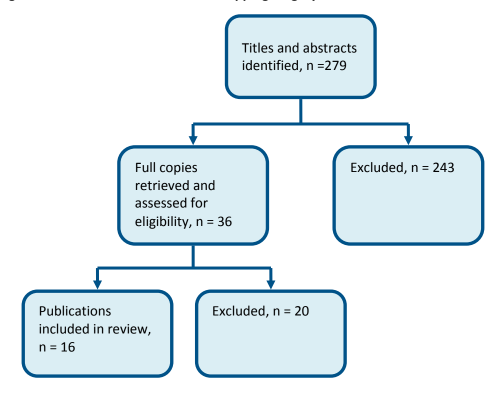
6 Figure 8: Clinical article selection: Stripping surgery vs. foam sclerotherapy review



1

2 D.5.2 Stripping surgery vs. endothermal ablation

3 Figure 9: Clinical article selection: stripping surgery vs. endothermal ablation

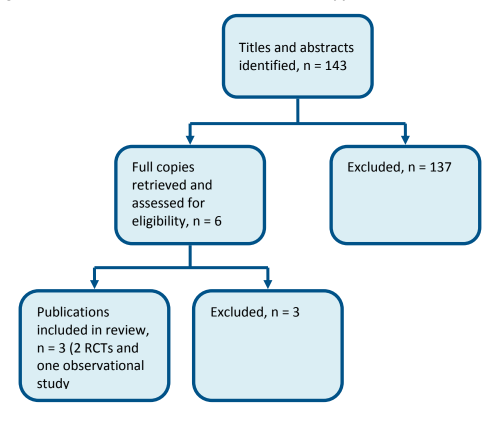


4

7

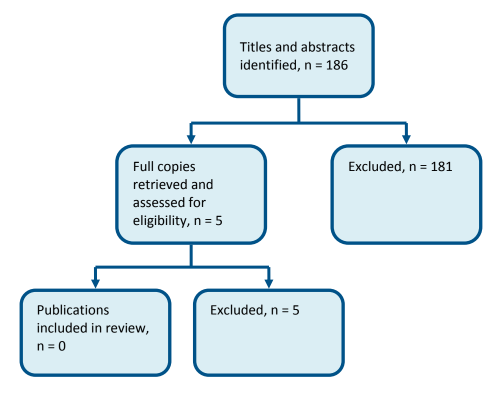
5 D.5.3 Foam sclerotherapy vs. endothermal ablation

6 Figure 10: Clinical article selection for foam sclerotherapy vs. endothermal ablation review



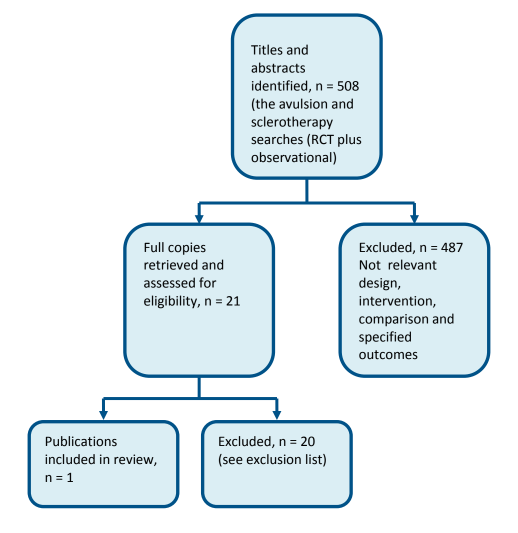
1 D.5.4 Tributary treatment: avulsion vs. foam sclerotherapy

2 Figure 11: Clinical article selection: foam sclerotherapy vs. avulsion for tributary veins



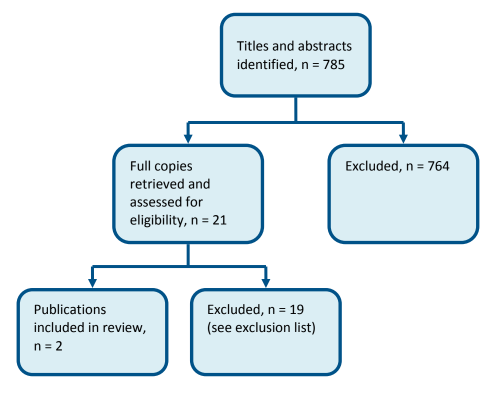
1 D.5.5 Truncal treatment and tributary treatment vs. truncal treatment alone

Figure 12: Clinical article selection: truncal with tributary treatment vs. truncal treatment alone



D.6 Chapter **10** – compression after interventional treatment

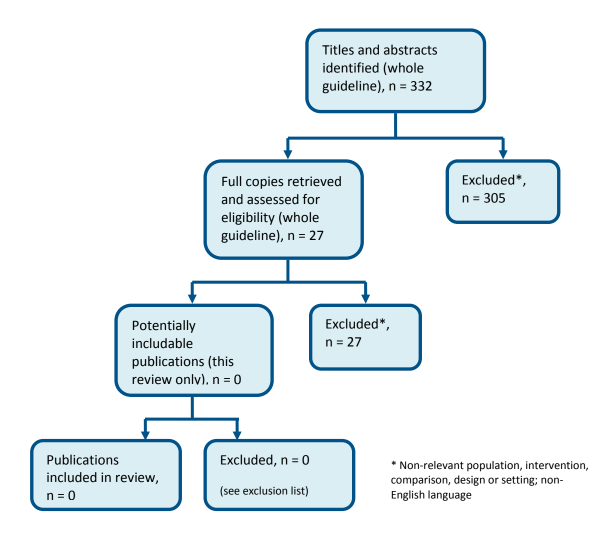
Figure 13: Clinical article selection: Interventional treatment with compression vs. interventional treatment alone



Appendix E: Economic article selection

2 E.1 Chapter 5 – patient perceptions and expectations

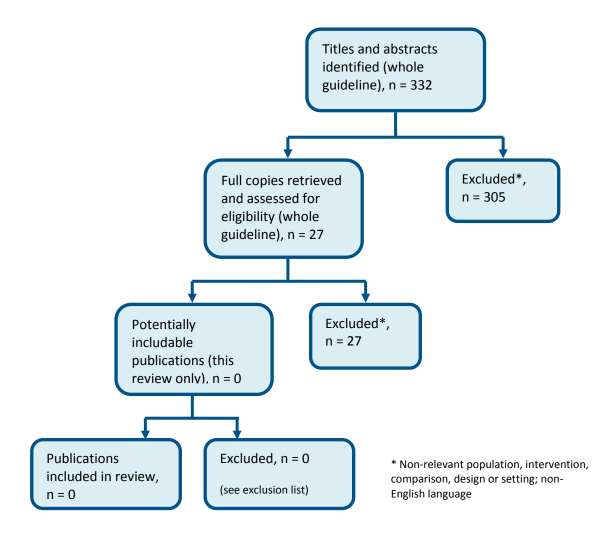
Figure 14: Economic article selection: patient perceptions and expectations



1

2 E.2 Chapter 6 – referral to a vascular service

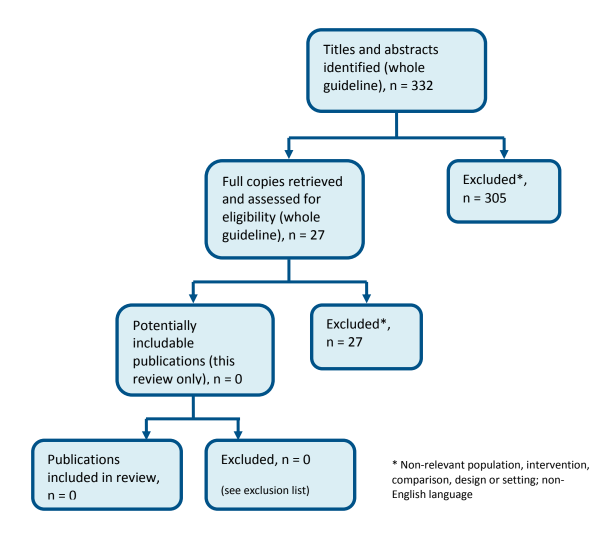
Figure 15: Economic article selection: referral to a vascular service



1 E.3 Chapter 7 – assessment for treatment

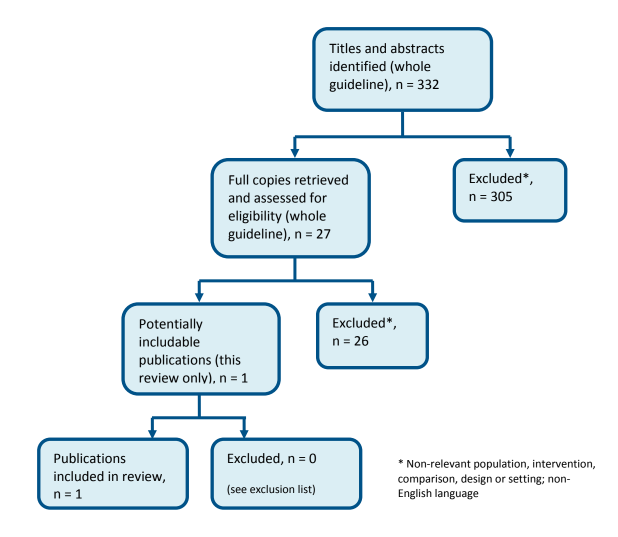
2 E.3.1 Duplex vs. Hand Held Doppler

- 3 Figure 16: Economic article selection: duplex vs. hand held doppler review
- 4



1 E.3.2 Duplex assessment prior to interventional treatment

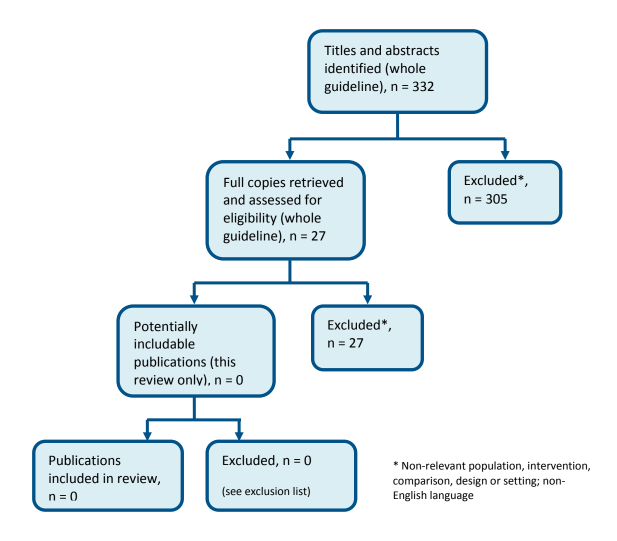
Figure 17: Economic article selection: duplex assessment prior to interventional treatment



1 E.4 Chapter 8 – conservative management

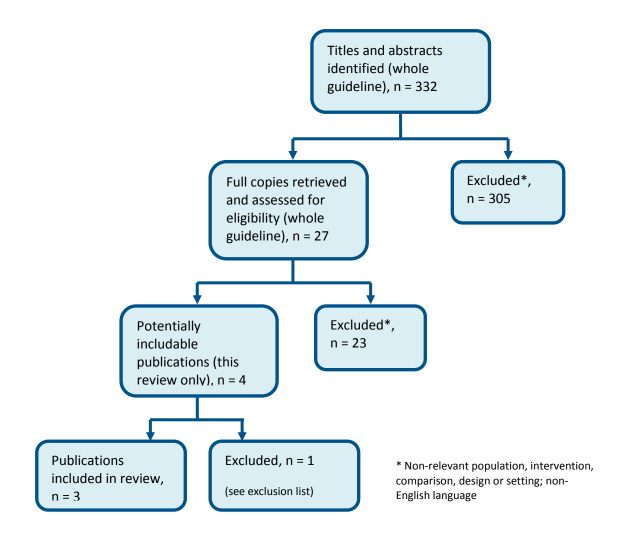
2 E.4.1 Compression vs. no treatment

Figure 18: Economic article selection: compression vs.no treatment



1 E.4.2 Compression vs. interventional treatment

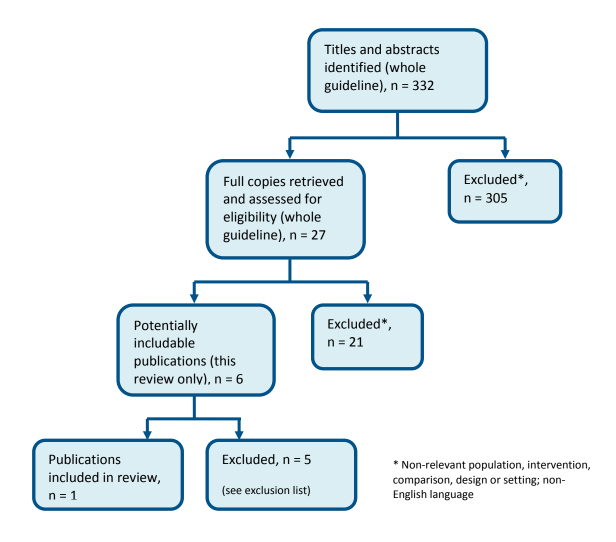
Figure 19: Economic article selection: compression vs. interventional treatment



1 E.5 Chapter 9 – interventional treatment

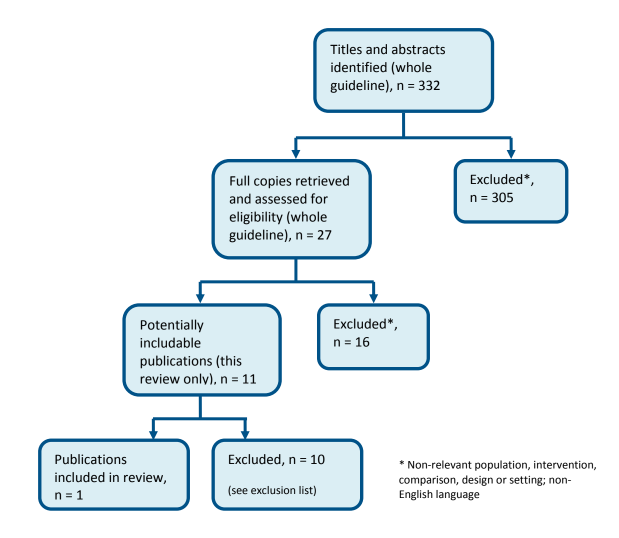
2 E.5.1 Stripping surgery vs. foam sclerotherapy

Figure 20: Economic article selection: stripping surgery vs. foam sclerotherapy



1 E.5.2 Stripping surgery vs. endothermal ablation

Figure 21: Economic article selection: stripping surgery vs. endothermal ablation



1 E.5.3 Foam sclerotherapy vs. endothermal ablation

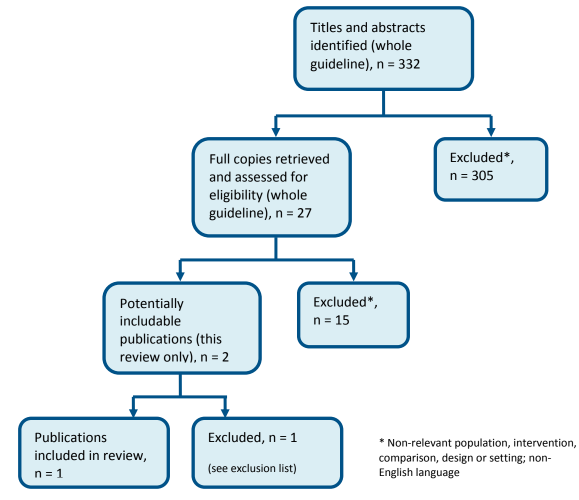
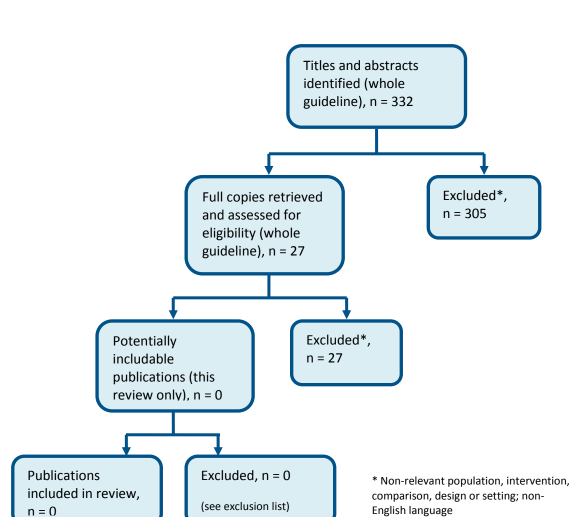


Figure 22: Economic article selection: foam sclerotherapy vs. endothermal ablation

1 E.5.4 Tributary treatment: avulsion vs. foam sclerotherapy

- 2 Figure 23: Economic article selection: avulsion vs. foam sclerotherapy
- 3



1 E.5.5 Truncal treatment and tributary treatment vs. truncal treatment alone

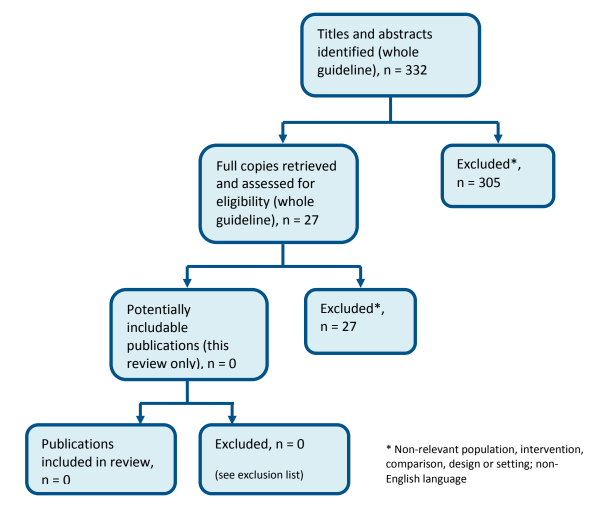
2 3 Figure 24: Economic article selection: truncal treatment and tributary treatment vs. truncal treatment alone

4

Titles and abstracts identified (whole guideline), n = 332 Excluded*, Full copies retrieved n = 305 and assessed for eligibility (whole guideline), n = 27 Excluded*, Potentially n = 26 includable publications (this review only), n = 1**Publications** Excluded, n = 0* Non-relevant population, intervention, included in review, comparison, design or setting; non-(see exclusion list) n = 1 English language

1 E.6 Chapter 10 – compression after treatment

2 Figure 25: Economic article selection: compression after treatment





Appendix F: Literature search strategies

Contents

Introduction	Search methodology
Section F.1	Study filter terms
F.1.1	Systematic reviews (SR)
F.1.2	Randomized controlled trials (RCT)
F.1.3	Observational studies
F.1.4	Economic studies
F.1.5	Quality of life studies
F.1.6	Diagnostic accuracy
Section F.2	Standard population search strategy Populations used for all search questions unless stated in F.3
Section F.3	Searches for specific questions with intervention (and population where different from F.2)
F.3.1	Patient information
F.3.2	Assessment for referral
F.3.3	Assessment for treatment
F.3.4	Conservative management
F.3.5	Interventional treatment
F.3.6	Compression post treatment
Section F.4	Economic searches
F.4.1	Economic reviews
F.4.2	Quality of life reviews

4 Introduction

Search strategies used for the Varicose Veins guideline were run in accordance with the Guidelines Manual (NICE, 2009). All searches were run finally on **17 October 2012** unless otherwise stated. Any studies added to the databases after this date were not included unless specifically stated in the text.

8 Clinical searches

Searches for **clinical reviews** were run in Medline (OVID), Embase (OVID), and the Cochrane Library (Wiley). Typically, searches were constructed in the following way:

- A PICO format was used for intervention searches. **Population** (P) terms were combined with **Intervention** (I) and sometimes **Comparison** (C) terms (as indicated in the tables under each individual question in Section F.3). An intervention can be a drug, a procedure or a diagnostic test. **Outcome** (O) terms are rarely used in search strategies for interventions. Study type filters (F.1) were added where appropriate.
- 17In addition to the databases outlined above, searches F.3.1 and F.3.4 were run in Cinahl (EBSCO), and18F.3.1 in PsycINFO (OVID).

1 Economic searches

2 Searches for **economic evidence** were run in Medline (Ovid), Embase (Ovid), the NHS Economic 3 Evaluations Database (NHS EED), the Health Technology Assessment (HTA) database and the Health 4 Economic Evaluation Database (HEED). NHS EED and HTA were searched via the Cochrane (Wiley) 5 interface. For Medline and Embase an economic filter (see F.1.4) was applied to the standard 6 population. All other searches were conducted using only population terms.

7 F.1 Study design search terms

8 F.1.1 Systematic review (SR) search terms

9 Medline search terms

wieunne	edime search terms	
1.	Meta-analysis/	
2.	Meta-analysis as topic/	
3.	(meta analy* or metanaly* or metaanaly*).ti,ab.	
4.	((systematic* or evidence*) adj2 (review* or overview*)).ti,ab.	
5.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.	
6.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.	
7.	(search* adj4 literature).ab.	
8.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.	
9.	cochrane.jw.	
10.	or/1-9	

Embase search terms

LIIIbase	search terms
1.	Systematic review/
2.	Meta-analysis/
3.	(meta analy* or metanaly* or metaanaly*).ti,ab.
4.	((systematic or evidence) adj2 (review* or overview*)).ti,ab.
5.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.
6.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.
7.	(search* adj4 literature).ab.
8.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.
9.	((pool* or combined) adj2 (data or trials or studies or results)).ab.
10.	cochrane.jw.
11.	or/1-10

11 F.1.2 Randomised controlled studies (RCTs) search terms

12

Medline search terms

Randomized controlled trial.pt.	
Controlled clinical trial.pt.	
randomi#ed.ab.	
placebo.ab.	
randomly.ab.	

6.	Clinical trials as topic.sh.
7.	trial*.ti.
8.	or/1-7

Embase search terms

1.	random*.ti,ab.
2.	factorial*.ti,ab.
3.	(crossover* or cross over*).ti,ab.
4.	((doubl* or singl*) adj blind*).ti,ab.
5.	(assign* or allocat* or volunteer* or placebo*).ti,ab.
6.	Crossover procedure/
7.	Single blind procedure/
8.	Randomized controlled trial/
9.	Rouble blind procedure/
10.	or/1-9

2 F.1.3 Observational studies search terms

Medline search terms

weatine se	viedline search terms	
1.	Epidemiologic studies/	
2.	exp Case control studies/	
3.	exp Cohort studies/	
4.	Cross-sectional studies/	
5.	case control.ti,ab.	
6.	(cohort adj (study or studies or analys*)).ti,ab.	
7.	((follow-up or observational or uncontrolled or non randomi#ed or nonrandomi#ed or epidemiologic*) adj (study or studies)).ti,ab.	
8.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort*)).ti,ab.	
9.	or/1-8	

4

3

Embase search terms

Empase s		
1.	Clinical study/	
2.	exp Case control study/	
3.	Family study/	
4.	Longitudinal study/	
5.	Retrospective study/	
6.	Prospective study/	
7.	Cross-sectional study/	
8.	Cohort analysis/	
9.	Follow-up/	
10.	cohort*.ti,ab.	
11.	9 and 10	
12.	case control.ti,ab.	
13.	(cohort adj (study or studies or analys*)).ti,ab.	
14.	((follow-up or observational or uncontrolled or non randomi#ed or nonrandomi#ed or epidemiologic*) adj (study or studies)).ti,ab.	

15.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort*)).ti,ab.
16.	or/1-8,11-15

1 F.1.4 Health economic search terms

Medline search terms

1.	Economics/
2.	Value of life/
3.	exp "Costs and cost analysis"/
4.	exp Economics, Hospital/
5.	exp Economics, Medical/
6.	exp Resource allocation/
7.	Economics, Nursing/
8.	Economics, Pharmaceutical/
9.	exp "Fees and charges"/
10.	exp Budgets/
11.	budget*.ti,ab.
12.	cost*.ti,ab.
13.	(economic* or pharmaco?economic*).ti,ab.
14.	(price* or pricing*).ti,ab.
15.	(financ* or fee or fees or expenditure* or saving*).ti,ab.
16.	(value adj2 (money or monetary)).ti,ab.
17.	resourc* allocat*.ti,ab.
18.	(fund or funds or funding* or funded).ti,ab.
19.	(ration or rations or rationing* or rationed).ti,ab.
20.	ec.fs.
21.	or/1-20

Embase search terms

1.	Health economics/
2.	exp Economic evaluation/
3.	exp Health care cost/
4.	exp Fee/
5.	Budget/
6.	Funding/
7.	Resource allocation/
8.	budget*.ti,ab.
9.	cost*.ti,ab.
10.	(economic* or pharmaco?economic*).ti,ab.
11.	(price* or pricing*).ti,ab.
12.	(financ* or fee or fees or expenditure* or saving*).ti,ab.
13.	(value adj2 (money or monetary)).ti,ab.
14.	resourc* allocat*.ti,ab.
15.	(fund or funds or funding* or funded).ti,ab.
16.	(ration or rations or rationing* or rationed).ti,ab.

1 F.1.5 Quality of life search terms

2 Medline search terms

1.	Quality-adjusted life years/
2.	Sickness impact profile/
3.	(quality adj2 (wellbeing or well being)).ti,ab.
4.	sickness impact profile.ti,ab.
5.	disability adjusted life.ti,ab.
6.	(qal* or qtime* or qwb* or daly*).ti,ab.
7.	(euroqol* or eq5d* or eq 5d*).ti,ab.
8.	(qol* or hql* or hqol* or h qol* or hrqol* or hr qol*).ti,ab.
9.	(health utility* or utility score* or disutilit*).ti,ab.
10.	(hui or hui1 or hui2 or hui3).ti,ab.
11.	health* year* equivalent*.ti,ab.
12.	(hye or hyes).ti,ab.
13.	rosser.ti,ab.
14.	(willingness to pay or time tradeoff or time trade off or tto or standard gamble*).ti,ab.
15.	(sf36 or sf 36 or short form 36 or shortform 36 or shortform36).ti,ab.
16.	(sf20 or sf 20 or short form 20 or shortform 20 or shortform20).ti,ab.
17.	(sf12 or sf 12 or short form 12 or shortform 12 or shortform12).ti,ab.
18.	(sf8 or sf 8 or short form 8 or shortform 8 or shortform8).ti,ab.
19.	(sf6 or sf 6 or short form 6 or shortform 6 or shortform6).ti,ab.
20.	or/1-19

Embase search terms

1.	Quality adjusted life year/	
2.	"Quality of life index"/	
3.	Short form 12/ or Short form 20/ or Short form 36/ or Short form 8/	
4.	Sickness impact profile/	
5.	(quality adj2 (wellbeing or well being)).ti,ab.	
6.	sickness impact profile.ti,ab.	
7.	disability adjusted life.ti,ab.	
8.	(qal* or qtime* or qwb* or daly*).ti,ab.	
9.	(euroqol* or eq5d* or eq 5d*).ti,ab.	
10.	(qol* or hql* or hqol* or h qol* or hrqol* or hr qol*).ti,ab.	
11.	(health utility* or utility score* or disutilit*).ti,ab.	
12.	(hui or hui1 or hui2 or hui3).ti,ab.	
13.	health* year* equivalent*.ti,ab.	
14.	(hye or hyes).ti,ab.	
15.	rosser.ti,ab.	
16.	(willingness to pay or time tradeoff or time trade off or tto or standard gamble*).ti,ab.	
17.	(sf36 or sf 36 or short form 36 or shortform 36 or shortform36).ti,ab.	
18.	(sf20 or sf 20 or short form 20 or shortform 20 or shortform20).ti,ab.	
19.	(sf12 or sf 12 or short form 12 or shortform 12 or shortform12).ti,ab.	
20.	(sf8 or sf 8 or short form 8 or shortform 8 or shortform8).ti,ab.	
21.	(sf6 or sf 6 or short form 6 or shortform 6 or shortform6).ti,ab.	

22. or/1-21

1 F.1.6 Diagnostic accuracy

2

3

5

Medline search terms

1.	exp "Sensitivity and specificity"/
2.	(sensitivity or specificity).ti,ab.
3.	((pre test or pretest or post test) adj probability).ti,ab.
4.	(prognos* or predict*).ti,ab,hw.
5.	(PPV or NPV).ti,ab.
6.	Likelihood function/
7.	(ROC curve* or AUC).ti,ab.
8.	(diagnos* adj2 (performance* or accurac* or utilit* or value* or efficien* or effectiveness)).ti,ab.
9.	gold standard.ab.
10.	(improve* adj3 (outcome* or result*)).ti,ab.
11.	Treatment outcome/
12.	or/1-11

Embase search terms

Empase	imbase search terms	
1.	exp "Sensitivity and specificity"/	
2.	Diagnostic accuracy/	
3.	Diagnostic test accuracy study/	
4.	Treatment outcome/	
5.	(prognos* or predict*).ti,ab,hw.	
6.	(sensitivity or specificity).ti,ab.	
7.	((pre test or pretest or post test) adj probability).ti,ab.	
8.	(PPV or NPV).ti,ab.	
9.	likelihood ratio*.ti,ab.	
10.	(ROC curve* or AUC).ti,ab.	
11.	(diagnos* adj2 (performance* or accurac* or utilit* or value* or efficien* or effectiveness)).ti,ab.	
12.	(improve* adj3 (outcome* or result*)).ti,ab.	
13.	gold standard.ab.	
14.	or/1-13	

4 F.2 Standard population search strategy

Medline search terms

INEUIIII	Medime search terms	
1.	exp Varicose veins/	
2.	(varicos* adj5 vein*).ti,ab.	
3.	Saphenous vein/	
4.	(sapheno* adj3 (vein* or junction* or incompet* or reflux or insufficien*)).ti,ab.	
5.	exp Coronary artery bypass/	
6.	((coronary or bypass) and graft).ti,ab.	
7.	(3 or 4) not (5 or 6)	
8.	1 or 2 or 7	

9.	Venous insufficiency/
10.	((venous or vein* or varico* or truncal or valvular) adj3 (insufficien* or incompet* or disorder* or reflux)).ti,ab.
11.	((venous or varico*) adj disease*).ti,ab.
12.	((perforator or superficial or tortuous) adj3 vein*).ti,ab.
13.	(varix or varices or varicosi* or ceap).ti,ab.
14.	((varico* or venous or vein*) adj3 ulcer*).ti,ab.
15.	or/9-14
16.	exp Lower extremity/
17.	(lower adj2 extremit*).ti,ab.
18.	(leg* or limb* or calf or calves or thigh* or groin* or ankle* or foot or feet or pelvis or pelvic or vulva* or vulvo* or ovari* or ovary or vagina* or uterus or uterin*).ti,ab.
19.	or/16-18
20.	8 or (15 and 19)
21.	limit 20 to english language
22.	Letter/
23.	Editorial/
24.	News/
25.	exp Historical article/
26.	Anecdotes as topic/
27.	Comment/
28.	Case report/
29.	(letter or comment*).ti.
30.	or/22-29
31.	30 not (Randomized controlled trial/ or random*.ti,ab.)
32.	Animals/ not Humans/
33.	exp Animals, Laboratory/
34.	exp Animal experimentation/
35.	exp Models, Animal/
36.	exp Rodentia/
37.	(rat or rats or mouse or mice).ti.
38.	or/31-37
39.	21 not 38

Embase search terms

1.	Varicosis/	
2.	Leg varicosis/	
3.	(varicos* adj5 vein*).ti,ab.	
4.	Saphenous vein/	
5.	(sapheno* adj3 (vein* or junction or incompet* or reflux or insufficien*)).ti,ab.	
6.	exp Coronary artery bypass graft/	
7.	((coronary or bypass) and graft).ti,ab.	
8.	(4 or 5) not (6 or 7)	
9.	1 or 2 or 3 or 8	
10.	exp Vein insufficiency/	
11.	((venous or vein* or varico* or truncal or valvular) adj3 (insufficien* or incompet* or disorder*	

	or reflux)).ti,ab.
12.	((venous or varico*) adj disease*).ti,ab.
13.	((perforator or superficial or tortuous) adj3 vein*).ti,ab.
14.	(varix or varices or varicosi* or ceap).ti,ab.
15.	((varico* or venous or vein*) adj3 ulcer*).ti,ab.
16.	or/10-15
17.	exp Leg/
18.	(lower adj2 extremit*).ti,ab.
19.	(leg* or limb* or calf or calves or thigh* or groin* or ankle* or foot or feet or pelvis or pelvic or vulva* or vulvo* or ovari* or ovary or vagina* or uterus or uterin*).ti,ab.
20.	or/17-19
21.	9 or (16 and 20)
22.	limit 21 to english language
23.	Letter.pt. or Letter/
24.	Note.pt.
25.	Editorial.pt.
26.	Case report/ or Case study/
27.	(letter or comment*).ti.
28.	or/23-27
29.	28 not (randomized controlled trial/ or random*.ti,ab.)
30.	Animal/ not Human/
31.	Nonhuman/
32.	exp Animal experiment/
33.	exp Experimental animal/
34.	Animal model/
35.	exp Rodent/
36.	(rat or rats or mouse or mice).ti.
37.	or/29-36
38.	22 not 37

Cinahl search terms

(MH "Varicose Veins")		
varicos* n5 vein*		
(MH "Saphenous Vein")		
sapheno* and (vein* or junction* or incompet* or reflux or insufficien*)		
(MH "Coronary Artery Bypass+")		
(S3 or S4) not S5		
S1 or S2 or S6		
(MH "Venous Insufficiency")		
(venous or vein* or varico* or truncal or valvular) and (insufficien* or incompet* or disorder* or reflux)		
(perforator n3 vein*) or (superficial n3 vein*) or (tortuous n3 vein*)		
varix or varices or varicosi* or ceap		
(venous n1 disease*) or (varico* n1 disease*)		
(varico* n3 ulcer*) or (vein* n3 ulcer*) or (venous n3 ulcer*)		
S8 or S9 or S10 or S11 or S12 or S13		

S15	(MH "Lower Extremity+")
S16	lower n2 extremit*
S17	leg* or limb* or calf or calves or thigh* or groin* or ankle* or foot or feet or pelvis or pelvic or vulva* or vulvo* or ovari* or ovary or vagina* or uterus or uterin*
S18	S15 or S16 or S17
S19	S7 or (S14 and S18)

Cochrane search terms

MeSH descriptor Varicose Veins explode all trees
(varicos* NEAR/5 vein*):ti,ab
MeSH descriptor Saphenous Vein, this term only
(sapheno* NEAR/3 (vein* or junction* or incompet* or reflux or insufficien*)):ti,ab
MeSH descriptor Coronary Artery Bypass explode all trees
((coronary or bypass) and graft):ti,ab
((#3 or #4) and not (#5 OR #6))
(#1 or #2 or #7)
MeSH descriptor Venous Insufficiency, this term only
((venous or vein* or varico* or truncal or valvular) NEAR/3 (insufficien* or incompet* or disorder* or reflux)):ti,ab
((perforator or superficial or tortuous) NEAR/3 vein*):ti,ab
(varix or varices or varicosi* or ceap):ti,ab
((varico* or vein* or venous) NEAR/3 ulcer*):ti,ab
((venous or varico*) NEXT disease*):ti,ab
(#9 or #10 or #11 or #12 or #13 or #14)
MeSH descriptor Lower Extremity explode all trees
(lower NEAR/2 extremit*):ti,ab
(leg* or limb* or calf or calves or thigh* or groin* or ankle* or foot or feet or pelvis or pelvic or vulva* or vulvo* or ovari* or ovary or vagina* or uterus or uterin*):ti,ab
(#16 or #17 or #18)
(#8 or (#15 and #19))

2 F.3 Searches by specific questions

3 F.3.1 Patient information

4 5

6

7

Q. What are the perceptions and expectations of people with varicose veins (e.g. natural history, treatment) and how can they be addressed?

Search constructed by combining the columns in the following table using the AND Boolean operator

Varicose veins Patient	information Nor	ne All yea	ars - 17/10/2012

Medline search terms

1.	((client* or patient* or user* or carer* or consumer* or customer* or health) adj3 (information* or educat* or knowledge or literacy or belief* or perception* or understanding or expectation* or prefer* or satisfaction or acceptance or compliance or adherence or concordance)).ti,ab,hw.
2.	(information* adj3 (need* or requirement* or support* or seek* or access* or disseminat*)).ti,ab,hw.

3.	((patient* or user* or carer* or consumer* or customer* or health) adj3 (literature or leaflet* or booklet* or pamphlet* or questionnaire* or survey* or handout* or internet or website* or consult* or interview*)).ti,ab.
4.	Telemedicine/
5.	Interview/
6.	Telephone/
7.	Publications/
8.	Pamphlets/
9.	Internet/
10.	or/1-9

Embase search terms

1.	((client* or patient* or user* or carer* or consumer* or customer* or health or medical) adj3 (information* or educat* or knowledge or literacy or belief* or perception* or understanding or expectation* or attitude* or prefer* or satisfaction or acceptance or compliance or adherence or concordance or advocacy)).ti,ab,hw.
2.	(information* adj3 (need* or requirement* or support* or seek* or access* or disseminat*)).ti,ab,hw.
3.	((patient* or user* or carer* or consumer* or customer* or health) adj3 (literature or leaflet* or booklet* or pamphlet* or questionnaire* or survey* or handout* or internet or website* or consult* or interview*)).ti,ab.
4.	exp Telehealth/
5.	exp Interview/
6.	Telephone/
7.	Publication/
8.	Internet/
9.	or/1-8

Cinahl search terms

eins")
Vein")
ein* or junction* or incompet* or reflux or insufficien*)
rtery Bypass+")
ufficiency")
or varico* or truncal or valvular) and (insufficien* or incompet* or disorder*
in*) or (superficial n3 vein*) or (tortuous n3 vein*)
r varicosi* or ceap
se*) or (varico* n1 disease*)
) or (vein n3 ulcer*) or (venous n3 ulcer*)
r S11 or S12 or S13
emity+")
t*
calf or calves or thigh* or groin* or ankle* or foot or feet or pelvis or pelvic or or ovari* or ovary or vagina* or uterus or uterin*

information* n2 support* or ation* n2 disseminat* nt* n3 educat*
ation* n2 disseminat*
ation* n2 disseminat*
nt* n3 educat*
educat* or health* n3
booklet* or patient* n3 pamphlet* ent* n3 handout* or patient* n3
)
raphy or PT book or PT book PT computer program or PT or PT interview or PT letter or PT or PT pamphlet chapter or PT d answers" or PT response or PT

Cochrane search terms

#1.	MeSH descriptor Patient Acceptance of Health Care, this term only
#2.	MeSH descriptor Patient Compliance, this term only
#3.	MeSH descriptor Patient Education as Topic, this term only
#4.	MeSH descriptor Patient Preference, this term only
#5.	MeSH descriptor Patient Satisfaction, this term only
#6.	MeSH descriptor Consumer Health Information, this term only
#7.	MeSH descriptor Consumer Satisfaction, this term only
#8.	MeSH descriptor Health Literacy, this term only
#9.	MeSH descriptor Health Knowledge, Attitudes, Practice, this term only
#10.	MeSH descriptor Telemedicine, this term only
#11.	MeSH descriptor Access to Information, this term only
#12.	MeSH descriptor Information Dissemination, this term only
#13.	MeSH descriptor Information Seeking Behavior, this term only
#14.	MeSH descriptor Pamphlets, this term only
#15.	MeSH descriptor Internet, this term only
#16.	MeSH descriptor Interviews as Topic explode all trees
#17.	MeSH descriptor Telephone, this term only
#18.	MeSH descriptor Telemedicine, this term only
#19.	((client* or patient* or user* or carer* or consumer* or customer* or health) NEAR/3 (information* or educat* or knowledge or literacy or belief* or perception* or attitude* or understanding or expectation* or prefer* or satisfaction or acceptance or compliance or adherence or concordance)):ti,ab
#20.	(information* NEAR/3 (need* or requirement* or support* or seek* or access* or

	disseminat*)):ti,ab
#21.	((patient* or user* or carer* or consumer* or customer* or health) NEAR/3 (literature or leaflet* or booklet* or pamphlet* or questionnaire* or survey* or handout* or internet or website* or consult* or interview*)):ti,ab
#22.	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15 or #16 or #17 or #18 or #19 or #20 or #21)

PsycINFO search terms

1.	(varicos* adj5 vein*).ti,ab.	
2.	((venous or vein* or varico* or truncal or valvular) adj3 (insufficien* or incompet* or disorder* or reflux)).ti,ab.	
3.	((venous or varico*) adj disease*).ti,ab.	
4.	((perforator or superficial or tortuous) adj3 vein*).ti,ab.	
5.	(varix or varices or varicosi* or ceap).ti,ab.	
6.	((varico* or venous or vein*) adj3 ulcer*).ti,ab.	
7.	(leg* or limb* or calf or calves or thigh* or groin* or ankle* or foot or feet or lower extremit*).ti,ab.	
8.	(2 or 3 or 4 or 5 or 6) and 7	
9.	1 or 8	

2 F.3.2 Assessment for referral

3 F.3.2.1 Disease progression

- 4Q.a) In people with leg varicose veins at CEAP class C2 which signs, symptoms and/or5patient characteristics are associated with disease progression to i) C3, ii) C4* iii) C6?607b) In people with leg varicose veins at CEAP class C3 which signs, symptoms and/or8patient characteristics are associated with disease progression to i) C4* ii) C6?91010c) In people with leg varicose veins at CEAP class C4* which signs, symptoms and/or11patient characteristics are associated with disease progression to C6?
- 12

1

Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Disease classification AND disease progression	Observational studies* [Medline and Embase only]	All years - 17/10/2012

13 *Observational search filter in F.1.3 expanded for this question

Medline search terms

1	4

Observational search fill	Lei III F.1.3	expanded for	this question

1.	(CEAP or C2 or C3 or C4 or C5 or C6 or C-2 or C-3 or C-4 or C-5 or C-6).ti,ab.	
2.	(class* or stage* or staging).ti,ab.	
3.	(skin adj2 (discol* or change* or pigment*)).ti,ab.	
4.	(ulcer* or oedem* or edem* or lipoderm* or eczema or atroph*).ti,ab.	
5.	or/1-4	
6.	Disease progression/	
7.	Natural history/	
8.	Risk factors/	
9.	(risk* or course* or predict* or incidence or prognos* or progress* or natural history).ti,ab.	

10.	or/6-9
11.	5 and 10
12.	Epidemiologic studies/
13.	exp Case control studies/
14.	exp Cohort studies/
15.	Cross-sectional studies/
16.	case control.ti,ab.
17.	((follow-up or observational or uncontrolled or non randomi#ed or nonrandomi#ed or epidemiologic*) adj (study or studies)).ti,ab.
18.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort*)).ti,ab.
19.	(cohort* or group* or subgroup* or participant*).ti,ab.
20.	or/12-19
21.	11 and 20

Embase search terms

1.	Disease classification/			
2.	(CEAP or C2 or C3 or C4 or C5 or C6 or C-2 or C-3 or C-4 or C-5 or C-6).ti,ab.			
3.	(class* or stage* or staging).ti,ab.			
4.	(skin adj2 (discol* or change* or pigment*)).ti,ab.			
5.	(ulcer* or oedem* or edem* or lipoderm* or eczema or atroph*).ti,ab.			
6.	or/1-5			
7.	Risk factor/			
8.	Disease course/			
9.	Disease exacerbation/			
10.	Predictive value/			
11.	(risk* or course* or predict* or incidence or prognos* or progress* or natural history).ti,ab.			
12.	or/7-11			
13.	6 and 12			
14.	Clinical study/			
15.	exp Case control study/			
16.	Family study/			
17.	Longitudinal study/			
18.	Retrospective study/			
19.	Prospective study/			
20.	Cross-sectional study/			
21.	Cohort analysis/			
22.	Follow-up/			
23.	case control.ti,ab.			
24.	((follow-up or observational or uncontrolled or non randomi#ed or nonrandomi#ed or epidemiologic*) adj (study or studies)).ti,ab.			
25.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort*)).ti,ab.			
26.	(cohort* or group* or subgroup* or participant*).ti,ab.			
27.	or/14-26			
28.	13 and 27			

1 F.3.2.2 Prediction of treatment outcomes

2 3

4

5

Q. In people with leg varicose veins are there any factors (clinical signs and symptoms or patient reported outcomes) that would predict increased benefits or harms from interventional treatments for varicose veins?

Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Interventional treatments AND risk factors	RCTs, SRs and Observational studies* [Medline and Embase only]	All years - 17/10/2012

6 *Observational search filter in F.1.3 expanded for this question

7

8

Medline search terms 1. Vascular surgical procedures/ 2. Ablation techniques/ 3. Laser therapy/ 4. Sclerotherapy/ 5. Sclerosing solutions/ 6. Sodium tetradecyl sulfate/ 7. (scleros* or sclerotherap* or sclero therap* or UGFS).ti,ab. 8. (polidocanol or POL or sodium tetradecyl or STS or sotradecol or fibrovein or sclerovein).ti,ab. 9. (foam or microfoam or liquid).ti,ab. 10. (strip* or cryostrip* or saphenect* or disconnect*).ti,ab. 11. (avuls* or phlebectom* or microphlebectom* or miniphlebectom* or trans illuminate* or transilluminate* or trivex).ti,ab. 12. surg*.ti. 13. (ablat* or endoluminal or laser or radiofrequency or radio frequency or endovenous or RFA or EVLT).ti,ab. 14. or/1-13 15. Risk factors/ 16. Risk assessment/ 17. exp Treatment outcome/ 18. Prognosis/ 19. Disease progression/ 20. (risk* or benefit* or harm* or predisp* or pre dispos* or prognos* or course* or progress* or predict* or characteristic* or factor*).ti,ab. 21. (age or gender or sex or BMI or heredity or weight or body mass or family history or obes* or pregnan* or birth or childbirth or lifestyle or occupation or contracept* or mobility or smoking or drinking or co-morb* or comorb* or reflux).ti,ab. 22. or/15-21 23. 14 and 22

Embase search terms

1.	Phlebectomy/	
2.	Sclerotherapy/	
3.	Sclerosing agent/	
4.	Tetradecyl sulfate sodium/	
5.	Polidocanol/	

6.	Vein stripping/		
7.	Laser surgery/		
8.	Radiofrequency ablation/		
9.	Endovenous laser ablation/		
10.	(avuls* or phlebectom* or microphlebectom* or miniphlebectom* or trans illuminate* or transilluminate* or trivex).ti,ab.		
11.	(scleros* or sclerotherap* or UGFS).ti,ab.		
12.	(polidocanol or POL or sodium tetradecyl or STS or sotradecol or fibrovein or sclerovein).ti,ab.		
13.	(foam or microfoam or liquid).ti,ab.		
14.	(strip* or cryostrip* or saphenect* or disconnect*).ti,ab.		
15.	(ablat* or endoluminal or laser or radiofrequency or radio frequency or endovenous or RFA or EVLT).ti,ab.		
16.	surg*.ti.		
17.	or/1-16		
18.	risk.hw.		
19.	predict*.hw.		
20.	Treatment outcome/		
21.	Treatment failure/		
22.	Treatment response/		
23.	Prognosis/		
24.	Disease course/		
25.	(risk* or benefit* or harm* or predisp* or pre dispos* or prognos* or course* or progress* or predict* or characteristic* or factor*).ti,ab.		
26.	(age or gender or sex or BMI or heredity or weight or body mass or family history or obes* or pregnan* or birth or childbirth or lifestyle or occupation or contracept* or mobility or smoking or drinking or co-morb* or comorb* or reflux).ti,ab.		
27.	or/18-26		
28.	17 and 27		

Cochrane search terms

#1.	MeSH descriptor Vascular Surgical Procedures, this term only		
#2.	MeSH descriptor Ablation Techniques, this term only		
#3.	MeSH descriptor Laser Therapy, this term only		
#4.	MeSH descriptor Sclerotherapy, this term only		
#5.	MeSH descriptor Sclerosing Solutions, this term only		
#6.	MeSH descriptor Sodium Tetradecyl Sulfate, this term only		
#7.	(scleros* or sclerotherap* or "sclero therapy" or UGFS):ti,ab		
#8.	(polidocanol or POL or "sodium tetradecyl" or STS or sotradecol or fibrovein or sclerovein):ti,ab		
#9.	(foam or microfoam or liquid):ti,ab		
#10.	(strip* or cryostrip* or saphenect* or disconnect*):ti,ab		
#11.	(avuls* or phlebectom* or microphlebectom* or miniphlebectom* or "trans illuminated" or transilluminate* or trivex):ti,ab		
#12.	(ablat* or endoluminal or laser or radiofrequency or "radio frequency" or endovenous or RFA or EVLT):ti,ab		
#13.	surg*:ti		
#14.	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13)		

	-		
#15.	MeSH descriptor Risk Factors, this term only		
#16.	MeSH descriptor Risk Assessment, this term only		
#17.	MeSH descriptor Treatment Outcome explode all trees		
#18.	MeSH descriptor Prognosis, this term only		
#19.	MeSH descriptor Disease Progression, this term only		
#20.	(risk* or benefit* or harm* or predisp* or "pre disposed" or "pre disposition" or prognos* or course* or progress* or predict* or characteristic* or factor*):ti,ab		
#21.	21. (age or gender or sex or BMI or heredity or weight or "body mass" or "family history" or obes or pregnan* or birth or childbirth or lifestyle or occupation or contracept* or mobility or smoking or drinking or co-morbidity or comorbidities or comorb* or reflux):ti,ab		
#22.	(#15 or #16 or #17 or #18 or #19 or #20 or #21)		
#23.	(#14 and #22)		

1 **F.3.3** Assessment for treatment

- 2 The following two questions were searched using a single strategy:
 - Q1. Does the use of duplex ultrasound during assessment improve outcome after interventional treatment compared to no duplex scanning in people with leg varicose veins?
 - Q2. What is the diagnostic accuracy of hand held Doppler (HHD) compared to gold standard of duplex scanning when used in patients with varicose veins?
 - Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Duplex ultrasound	Diagnostic accuracy [Medline and Embase only]	All years – 17/10/2012

9 Medline search terms

1.	exp Ultrasonography, Doppler/
2.	Ultrasonography, Interventional/
3.	(ultraso* or echograph* or sonogra* or Doppler or duplex or DU or DUS).ti,ab.
4.	or/1-3

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Embase search terms

1.	Echography/		
2.	Doppler echography/		
3.	Endoscopic echography/		
4.	Color ultrasound flowmetry/		
5.	Diagnostic imaging/		
6.	(ultraso* or echograph* or sonogra* or Doppler or duplex or DU or DUS).ti,ab.		
7.	or/1-6		

Cochrane search terms

#1.	MeSH descriptor Ultrasonography, Doppler explode all trees		
#2.	MeSH descriptor Ultrasonography, Interventional, this term only		
#3.	(ultraso* or echograph* or sonogra* or Doppler or duplex or DU or DUS):ti,ab		
#4.	(#1 or #2 or #3)		
#5.	MeSH descriptor Sensitivity and specificity explode all trees		

#6.	MeSH descriptor Prognosis, this term only		
#7.	MeSH descriptor Treatment outcome, this term only		
#8.	MeSH descriptor Likelihood functions, this term only		
#9.	(sensitivity or specificity):ti,ab		
#10.	(("pre test" or pretest or "post test") NEXT probability):ti,ab		
#11.	(prognos* or predict*):ti,ab		
#12.	(PPV or NPV):ti,ab		
#13.	("ROC curve*" or AUC):ti,ab		
#14.	(diagnos* NEAR/2 (performance* or accurac* or utilit* or value* or efficien* or effectiveness)):ti,ab		
#15.	"gold standard":ab		
#16.	(improve* NEAR/3 (outcome* or result*)):ti,ab		
#17.	(#5 or #6 or #7 or #8 or #9 or #10 or #11 or #12 or #13 or #14 or #15 or #16)		
#18.	(#4 and #17)		

1 F.3.4 Conservative management

2 The following four questions were searched using a single strategy:

- Q1. What is the clinical and cost effectiveness of compression therapy compared with no treatment or lifestyle advice in people with leg varicose veins?
 - Q2. What is the clinical and cost effectiveness and safety of compression therapy compared with foam sclerotherapy in people with leg varicose veins?
- Q3. What is the clinical and cost effectiveness and safety of compression therapy compared with stripping surgery in people with leg varicose veins?
- 9 Q4. What is the clinical and cost effectiveness and safety of compression therapy compared 10 with endothermal ablation in people with leg varicose veins?

Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Compression therapy	RCTs, SRs and Observational studies [Medline and Embase only]	All years - 17/10/2012

Medline search terms

1.	Pressure/	
2.	Bandages/	
3.	exp Compression bandages/	
4.	Intermittent pneumatic compression devices/	
5.	((compressi* or pressure) and (hosiery or stocking* or bandag*)).ti,ab.	
6.	((compressi* or pressure or hosiery or stocking* or bandag*) adj2 (therap* or treatment* or device* or eccentric or pneumatic)).ti,ab.	
7.	((elastic or system* or support) adj2 (hosiery or stocking* or bandag*)).ti,ab.	
8.	(external* adj2 compression).ti,ab.	
9.	conservative treatment*.ti,ab.	
10.	or/1-9	

Embase search terms

1.

Compression/

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2.	exp Compression therapy/	
3.	Compression bandage/	
4.	*Bandage/	
5.	Compression garment/	
6.	Intermittent pneumatic compression device/	
7.	((compressi* or pressure) and (hosiery or stocking* or bandag*)).ti,ab.	
8.	((compressi* or pressure or hosiery or stocking* or bandag*) adj2 (therap* or treatment* or device* or eccentric or pneumatic)).ti,ab.	
9.	((elastic or system* or support) adj2 (hosiery or stocking* or bandag*)).ti,ab.	
10.	(external* adj2 compression).ti,ab.	
11.	conservative treatment*.ti,ab.	
12.	or/1-11	

Cinahl search terms

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S1	(MH "Compression therapy")		
S2	(MH "Compression garments")		
S3	(compressi* or pressure) and (hosiery or stocking* or bandag*)		
S4	(compressi* n2 therap*) or (compressi* n2 treatment) or (compressi* n2 device*) or (compressi* n2 eccentric) or (compressi* n2 pneumatic) or (pressure n2 therap*) or (pressure n2 treatment)		
S5	(elastic n2 hosiery) or (elastic n2 stocking*) or (elastic n2 bandag*) or (system n2 hosiery) or (system n2 stocking*) or (system n2 bandag*) or (support n2 hosiery) or (support n2 stocking*) or (support n2 bandag*)		
S6	external* n2 compression		
S7	conservative treatment*		
S8	S1 or S2 or S3 or S4 or S5 or S6 or S7		

Cochrane search terms

#1	MeSH descriptor Pressure, this term only	
#2	MeSH descriptor Bandages, this term only	
#3	MeSH descriptor Compression Bandages explode all trees	
#4	MeSH descriptor Intermittent Pneumatic Compression Devices, this term only	
#5	((compressi* or pressure) and (hosiery or stocking* or bandag*)):ti,ab	
#6	((compressi* or pressure or hosiery or stocking* or bandag*) NEAR/2 (therap* or treatment* or device* or eccentric or pneumatic)):ti,ab	
#7	((elastic or system* or support) NEAR/2 (hosiery or stocking* or bandag*)):ti,ab	
#8	(external* NEAR/2 compression):ti,ab	
#9	(conservative NEXT treatment*):ti,ab	
#10	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9)	

3 F.3.5 Interventional treatment

4 F.3.5.1 Stripping surgery

5 The following two questions were searched using a single strategy:

6 Q1. What is the clinical and cost effectiveness and safety of stripping surgery compared with 7 foam sclerotherapy in people with truncal leg varicose veins?

Q2. What is the clinical and cost effectiveness and safety of stripping surgery compared with endothermal ablation in people with truncal leg varicose veins?

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Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Stripping surgery	RCTs and SRs [Medline and Embase only]	All years - 17/10/2012

4 Medline search terms

1.	Vascular surgical procedures/	
2.	(strip* or cryostrip* or saphenect*).ti,ab.	
3.	surg*.ti.	
4.	or/1-3	

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Embase search terms

1.	Vein stripping/
2.	(strip* or cryostrip* or saphenect*).ti,ab.
3.	surg*.ti.
4.	or/1-3

Cochrane search terms

#1	MeSH descriptor Vascular Surgical Procedures, this term only
#2	(strip* or cryostrip* or saphenect*):ti,ab
#3	(#1 or #2)

7 F.3.5.2 Sclerotherapy

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Q. What is the clinical and cost effectiveness of foam sclerotherapy compared with endothermal ablation in people with truncal leg varicose veins?

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Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Sclerotherapy	RCTs, SRs and Observational studies [Medline and Embase only]	All years – 17/10/2012

11 Medline search terms

1.	Sclerotherapy/	
2.	Sclerosing solutions/	
3.	Sodium tetradecyl sulfate/	
4.	(scleros* or sclerotherap* or UGFS).ti,ab.	
5.	(polidocanol or POL or sodium tetradecyl or STS or sotradecol or fibrovein or sclerovein).ti,ab.	
6.	(foam or microfoam or liquid).ti,ab.	
7.	or/1-6	

Embase search terms

1.	Sclerotherapy/
2.	Sclerosing agent/
3.	Tetradecyl sulfate sodium/
4.	Polidocanol/
5.	(scleros* or sclerotherap* or UGFS).ti,ab.

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6.	(polidocanol or POL or sodium tetradecyl or STS or sotradecol or fibrovein or sclerovein).ti,ab.	
7.	(foam or microfoam or liquid).ti,ab.	
8.	or/1-7	

Cochrane search terms

Cochrane search terms		
MeSH descriptor Sclerotherapy, this term only		
MeSH descriptor Sclerosing solutions, this term only		
MeSH descriptor Sodium tetradecyl sulfate, this term only		
(scleros* or sclerotherap* or UGFS):ti,ab		
(polidocanol or POL or sodium tetradecyl or STS or sotradecol or fibrovein or sclerovein):ti,ab		
(foam or microfoam or liquid):ti,ab		
(#1 or #2 or #3 or #4 or #5 or #6)		

2 F.3.5.3 Avulsion surgery

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- 3 The following two questions were searched using a single strategy:
 - Q1. What is the clinical and cost effectiveness of avulsion surgery compared with sclerotherapy in people with tributary leg varicose veins?
 - Q2. What is the clinical and cost effectiveness of truncal treatment accompanied by tributary treatment (avulsion or sclerotherapy) compared with truncal treatment alone in people with leg varicose veins?
 - Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filters used	Date parameters
Varicose veins	Avulsion surgery	RCTs, SRs and Observational	All years - 17/10/2012
		studies [Medline and Embase only]	

10 Medline search terms

1.	(avuls* or phlebectom* or microphlebectom* or miniphlebectom* or trans illuminate* or transilluminate* or trivex).ti,ab.
2.	(stab or hook*).ti,ab.
3.	((ambula* or minim* invasive) adj3 surg*).ti,ab.
4.	(branch* adj2 (venous or vein or varicos*)).ti,ab.
5.	tributar*.ti,ab.
6.	or/1-5

Embase search terms

Phlebectomy/
(avuls* or phlebectom* or microphlebectom* or miniphlebectom* or trans illuminate* or transilluminate* or trivex).ti,ab.
(stab or hook*).ti,ab.
((ambula* or minim* invasive) adj3 surg*).ti,ab.
(branch* adj2 (venous or vein or varicos*)).ti,ab.
tributar*.ti,ab.
or/1-6

C

Cochrane search terms #1 (avuls* or phlebectom* or microphlebectom* or miniphlebectom* or trans illuminate* or

#2	(stab or hook*):ti,ab	
#3	((ambula* or minim* invasive) NEAR/3 surg*):ti,ab	
#4	(branch* NEAR/2 (venous or vein or varicos*)):ti,ab	
#5	tributar*:ti,ab	
#6	(#1 or #2 or #3 or #4 or #5)	

1 F.3.6 Compression post treatment

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Q. What is the clinical and cost effectiveness of compression post-ablative treatment compared with ablative treatment alone in people with truncal leg varicose veins?

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Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Intervention	Study filter used	Date parameters
Varicose veins	Compression therapy	RCTs, SRs and Observational studies [Medline and Embase only]	All years - 17/10/2012

Medline search terms

1.	Pressure/
2.	Bandages/
3.	exp Compression bandages/
4.	Intermittent pneumatic compression devices/
5.	((compressi* or pressure) and (hosiery or stocking* or bandag*)).ti,ab.
6.	((compressi* or pressure or hosiery or stocking* or bandag*) adj2 (therap* or treatment* or device* or eccentric or pneumatic)).ti,ab.
7.	((elastic or system* or support) adj2 (hosiery or stocking* or bandag*)).ti,ab.
8.	(external* adj2 compression).ti,ab.
9.	conservative treatment*.ti,ab.
10.	or/1-9

Embase search terms

1.	Compression/
2.	exp Compression therapy/
3.	Compression bandage/
4.	*Bandage/
5.	Compression garment/
6.	Intermittent pneumatic compression device/
7.	((compressi* or pressure) and (hosiery or stocking* or bandag*)).ti,ab.
8.	((compressi* or pressure or hosiery or stocking* or bandag*) adj2 (therap* or treatment* or device* or eccentric or pneumatic)).ti,ab.
9.	((elastic or system* or support) adj2 (hosiery or stocking* or bandag*)).ti,ab.
10.	(external* adj2 compression).ti,ab.
11.	conservative treatment*.ti,ab.
12.	or/1-11

Cinahl search terms

S1	(MH "Compression therapy")	
S2	(MH "Compression garments")	
S3	(compressi* or pressure) and (hosiery or stocking* or bandag*)	
S4	(compressi* n2 therap*) or (compressi* n2 treatment) or (compressi* n2 device*) or	

	(compressi* n2 eccentric) or (compressi* n2 pneumatic) or (pressure n2 therap*) or (pressure n2 treatment)
S5	(elastic n2 hosiery) or (elastic n2 stocking*) or (elastic n2 bandag*) or (system n2 hosiery) or (system n2 stocking*) or (system n2 bandag*) or (support n2 hosiery) or (support n2 stocking*) or (support n2 bandag*)
S6	external* n2 compression
S7	conservative treatment*
S8	S1 or S2 or S3 or S4 or S5 or S6 or S7

Cochrane search terms

cocinan		
#1	MeSH descriptor Pressure, this term only	
#2	MeSH descriptor Bandages, this term only	
#3	MeSH descriptor Compression Bandages explode all trees	
#4	MeSH descriptor Intermittent Pneumatic Compression Devices, this term only	
#5	((compressi* or pressure) and (hosiery or stocking* or bandag*)):ti,ab	
#6	((compressi* or pressure or hosiery or stocking* or bandag*) NEAR/2 (therap* or treatment* or device* or eccentric or pneumatic)):ti,ab	
#7	((elastic or system* or support) NEAR/2 (hosiery or stocking* or bandag*)):ti,ab	
#8	(external* NEAR/2 compression):ti,ab	
#9	(conservative NEXT treatment*):ti,ab	
#10	(#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9)	

2 F.4 Economic searches

3 F.4.1 Economic reviews

Economic searches were run in Medline and Embase by combining the standard population with an
economic filter, and limiting by date range (see table below). Economic searches were executed in
the HEED and Cochrane (NHS EED and HTA) databases by running a standard population without
date limitation. Search terms for the HEED database are given below (for Cochrane population see
F.2).

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Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Study filter used	Date parameters
Varicose veins	Economic [Embase and Medline]	 2009 - 17/10/2012 (Medline and Embase) All years - 17/10/2012 (NHS EED, HTA and HEED)

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HEED search terms

TILLD Scare	
1.	AX=varicos* AND vein*
2.	AX=sapheno* AND vein*
3.	AX=venous AND insufficien*
4.	CS=1 OR 2 OR 3

11 F.4.2 Quality of life reviews

- 12 Quality of life (QOL) searches were run in Medline and Embase by combining the standard 13 population with the QOL filter (F.1.5) without date limitation.
- 14 Search constructed by combining the columns in the following table using the AND Boolean operator

Population	Study filter used	Date parameters
Varicose veins	QOL	All years – 17/10/2012

Appendix G: Evidence tables clinical studies

G.1 Chapter 5 – patient perceptions and expectations

Table 15: Bobridge 2011²⁷

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
Bobridge A, Sandison S, Paterson J, Puckridge P, Esplin M. A pilot study of the development and implementation of a 'best practice' patient information booklet for patients with chronic venous insufficiency. Phlebology 2011; 26: 338- 343	Observational before-after study. Because of the lack of a control group the findings of this study are prone to considerable threats to internal validity. Setting: Australian General Hospital.	30 originally recruited. 3 withdrew due to significant health problems and one withdrew with time management issues. 26 thus started the study. At the 6 month follow-up a further 3 withdrew due to significant health problems and 3 were lost to follow- up: 20 were thus left for analysis at 6 months. This was a per-protocol analysis.	 15 women, 11 men; mean (range) age: 71.8 (38-90); CEAP stages: C3: 11.5%, C4: 38.5%, C5: 30.8%, C6: 19.2%; Current treatment: compression stockings - 69.2%, compression bandages – 30.8%, moisturising skin – 26.1%, leg exercises – 15.4%, leg elevation – 3.8%; median 24 month duration of chronic venous insufficiency (CVI) (range 0.25-684 months); most common causes of CVI were superficial and deep perforator incompetence, and superficial great saphenous and deep incompetence (15.4% each); co-morbidities: hypertension (50%), type II diabetes (15.4%), 	Patients given baseline questionnaires: health Education Impact Questionnaire (HEIQ), measuring the participants chronic venous insufficiency (CVI) knowledge; they were also given the CIVIQ. The patients were then given an information booklet, which had been developed on the best available evidence from the literature; it contained lay term information on the pathophysiology of CVI and the importance of skin care, leg elevation, exercise, diet and compression garments. The information booklet had been amended after consultations with vascular clinicians. A vascular nurse specialist gave the booklet, and explained its contents. The participant was asked to take the information booklet home, to read through the information, and to undertake the recommended best practice activities in their home environment over the next 6 month period. No other intervention was given, though it is assumed (unclearly reported) that	None reported

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
			 thyroid dysfunction (15.4%); BMI range 20-35.4 (mean 30.8). Inclusion: Duplex evidence of reflux, with a CAEP of C3-C6. Exclusion: Leg swelling due to cardiac, renal or hepatic dysfunction, lymphodema, lipoedema, DVT, cellulitis, cancer or post-op swelling; diminished mental cognition; physical disability. 	patients were allowed to continue their current treatment regimens (as described in patient characteristics column). Further measurements (HEIQ and CIVIQ) were taken at 1 and 6 months post-booklet allocation via the telephone.	

Evidence tables clinical studies

Results: The paper gives descriptive results only. P values given but no event rates.

<u>1 month post booklet implementation (n=26)</u>

There were significant improvements in participants performing at least one activity to improve their CVI (p=0.01), monitoring their CVI (p=0.045), knowing things which could trigger their CVI and make it worse (p=0.005), having effective ways to prevent their CVI symptoms from limiting what they can do (p=0.045). There were also improvements in the ability to travel by car or bus (p=0.05), undertaking social activities (p=0.03) and feeling less embarrassed about showing their legs (p=0.025).

6 months post booklet implementation (n=20)

During the time between 1 and 6 months, there was a significant reduction in the number of people worrying about CVI (p=0.012) and feeling embarrassed about showing their legs (p=0.005), as well as being able to climb several flights of stairs (p=0.008)

At 6 months, there were also significant improvements in performing at least one activity to improve CVI (p=0.003), knowing things which could trigger CVI and make it worse (p=0.016), having effective ways to prevent CVI symptoms limiting what they can do (p=0.008), worrying about their CVI (p=0.03) and feeling a sense of hopelessness about their CVI (p=0.007). there was also a significant improvement in leg and ankle pain (p=0.038), ability to do domestic chores (p=0.017), feeling nervous and tense (p=0.026) and feeling embarrassed about showing their legs (p=0.008).

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
Campbell WB, Decaluwe H, MacIntyre JB, Thompson JF, Cowan AR. Most patients with varicose veins have fears or concerns about the future, in addition to their presenting symptoms. Eur J Vasc Endovasc Surg 2006; 31: 332-334.	Quantitative cross- sectional questionnaire study. Setting: unclear but likely to be a vascular unit in an NHS secondary care trust.	203 patients initially sent in their questionnaires – a 62% response rate. 13 were later excluded due to 8 not having varicose veins, 3 having ulcers and 2 having phlebitis. Hence 190 participated in the study.	Patients referred to a vascular unit with "uncomplicated varicose veins". 75% female; median age 51 (range 20-83).	 Patients who were due to attend the vascular clinic were sent questionnaires beforehand, and asked to bring them in completed for clinic. No reminders were given, and no patient was asked to complete a questionnaire after receiving advice in clinic. The questionnaire contained 13 questions about symptoms and future expectations. The questions relevant to future expectations were: Have you any other concerns, worries or fears about your varicose veins? Yes/No If yes, what are they? Are you worried that your varicose veins might cause you medical harm? Yes/No If yes, what exactly are you worried about? 	None reported

Table 16: Campbell 2006⁴¹

Results:

Negative expectations about prognosis

- 150/190 (79%) reported that they had concerns, worries or fears about their varicose veins:
- 59/190 (31%) feared future thrombosis
- 30/190 (16%) feared future trauma/bleeding
- 28/190 (15%) feared future ulcers
- 22/190 (12%) feared future circulatory disease
- 8/190 (4%) feared future phlebitis
- 57/190 (30%)had general concerns about the future, particularly if there was a family history of varicose veins.

These worries contrasted with the answers to another question on the questionnaire, which asked about their reason for seeking medical advice: only 27/173 (16%) stated concerns about the future.

Evidence tables clinical studies

		Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
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How can these expectations be addressed?

The authors suggested that "good explanation (both verbal and written) about the nature and prognosis of varicose veins should be a routine part of good patient management. Reassurance against the likelihood of a benign prognosis leads many to decide against treatment, especially if they understand they can return in future. We do not know which patients will go on to develop skin problems or ulceration, but clinical experience suggests that the proportion is small and patients should be told the warning signs of eczema or darkening of the skin at the ankle."

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
Darvall KAL, Bate GR, Sam RC, Adam DJ, Silverman SH, Bradbury AW. Patients' expectations before and satisfaction after ultrasound guided foam sclerotherap y for varicose veins. Eur J Vasc Endovasc Surg 2009; 38: 642-647.	Quantitative cross- sectional questionnaire study. Setting: Large NHS secondary care trust	351 patients (464 legs). 80% response rate for the expectations questionnaire.	Consecutive patients undergoing foam sclerotherapy for symptomatic varicose veins. Patients had been referred for treatment by their GPs to NHS vascular surgeons at a single hospital. 35% male; 25% had recurrent disease; 67% had CEAP 2-3; 33% had CEAP 4-6; all had symptomatic primary varicose veins; 97% had superficial venous reflux only and 3% had both superficial and deep reflux; all were secondary to reflux not obstruction. Treatment was for the great saphenous vein (76% of patients), small saphenous vein (10%) and a combination of great saphenous vein, small saphenous vein and anterior accessory saphenous vein (14%).	A questionnaire was given one week prior to foam sclerotherapy treatment. The questionnaire responses were on a 5 point Likert scale (an awful lot, a lot, quite a bit, a little, not at all), which was later collapsed to 3 categories: "a significant improvement", "quite a bit" and "not at all" (the paper uses the term "moderate" in the results section, which presumably means "quite a bit"). The patient could also indicate if a question was not applicable. The questionnaire was given for each leg to be treated (so a bilaterally affected patient would do two questionnaires). Section 1 asked about the expected improvements in symptoms (pain or aching, itching, tingling, restless legs, cramps, swelling and heaviness). Section2 asked about expected improvements in appearance, lifestyle (choice of clothes, work performance, social and leisure activities and relationships. [<i>In addition a post treatment questionnaire was given 6 months after treatment to ascertain actual subjective improvements in all these areas, using the same response categories. Integration of responses from the pre and post treatment questionnaires allowed estimation of whether expectations were met.</i>]	None reported

11 200065 47.

Results:

Symptom expectations

A significant improvement in symptoms was expected in approximately 33% of legs, and a moderate improvement in 67%.

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
		between 49% and 6 ere met or exceeded		ement in symptoms at 6 month post-tre	eatment and about 10% had no improvement
The detailed	expectations dat	a for individual syn	nptoms are given below (the perc	entage figures are approximate as extra	polated from low resolution table)
Symptom		Expectatio	on of significant improvement (%	Expectation	n of moderate improvement (%)
pain	37	,		63	
itch	32			68	
tingling	24			76	
cramp	30			70	
restless legs	29			71	
swelling	37			63	
heaviness	37			63	
Percentages	where pre-opera	tive expectations	were not met 6 months post-ope		
	Fa	actor		•••••	[n=281] where expectations were not met
Symptoms		ain		20%	
	Ite			21%	
		ngling		18%	
		amp		23%	
		estless legs		22%	
	SI	velling		27%	
		eaviness		18%	
Other factors	A	opearance of the le	gs	12%	
	Cl	noice of clothes tha	t can be worn	25%	
	Pe	erformance at work	< compared with the second sec	25%	
	Re	elationships		14%	
	Er	njoyment of leisure	activities	30%	

Cosmetic expectations

Over 60% of patients expected a significant improvement in the appearance of their legs (a further 30% expected a moderate improvement).

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
[96% actually	noticed a signific	ant improvement d	at 6 months.]		
Life-style ben	efits expectation	s			
Approx. 30% of	of patients expect	ted significant imp	rovement in the choice of clothes	s, and a further 40% expected moderate improvement.	
[75% of patier	nts met or exceed	led these expectati	ons]		
Approx. 27% of	of patients expect	ted significant imp	rovement in performance at wor	k, and a further 40% expected moderate improvement.	
[75% of patier	nts met or exceed	led these expectati	ons]		
Approx. 27% o	of patients expect	ted significant imp	rovement in leisure activities, and	d a further 40% expected moderate improvement.	
[75% of patie	ents met or exceed	ded these expectat	ions]		
Relationships	expectations				
10% of patien	ts expected signif	ficant improvemen	t in relationships, and a further 1	.5% expected moderate improvement.	
[>50% actuall	y experienced suc	ch improvements.]			
Factors affect	ing expectations				
each related to	o higher expectat	ions of cosmetic in		evious varicose veins surgery. Younger age (<55yrs) and CEAP st tients had higher expectations in terms of clothes choice. Work,	

Table 18: Dillon	2005~				
Reference	Study type	No of patients	Patient characteristics	Methodology	Source of funding
Dillon MF, Carr CJ, Feeley TMF, Tierney S. Impact of the informed consent process on patients' understanding of varicose veins and their treatment. Irish Journal of medical science 2005; 174: 23- 27	Quantitative questionnaire study carried out in Republic of Ireland. Setting: randomly selected vascular clinics in Republic of Ireland.	82 given the original questionnaire, and all 82 completed it. 67/82 completed the telephone interview 2 weeks post- information provision (pre surgery) and reasons for drop-out are not given.	Patients with newly diagnosed varicose veins referred to randomly selected vascular clinics for surgery. 57 females; median age (range) of 46 (17-72) years; 37/82 had completed secondary education.	The initial written questionnaire was given at the first vascular clinic appointment. It is unclear if this was given before or after the consultation, but, given the questionnaires purpose of evaluating initial expectations, it appears likely it was before the consultation. This because the consultation included an in-depth discussion of the nature and consequences of surgery; furthermore, immediately after the consultation the patient was given a leaflet reiterating this information. This questionnaire assessed the expectations of the outcome of surgery and the perception of threats to health from varicose veins. Two weeks later (but before the surgery itself) the patients were given a repeat of the initial questionnaire (but in telephone interview form in all but one case) to assess the effects of the discussion and leaflet on expectations	None reported

Results:

Expectations about varicose vein risks

- 46/82 initially believed that they were at "high risk" of developing ulcers. Two weeks after information giving this figure was 40/67.
- 41/82 initially believed that they were at "high risk" of developing DVT. Two weeks after information giving this figure was 33/67.
- 26/82 initially believed that they were at "high risk" of bleeding from minor injuries. Two weeks after information giving this figure was 45/67.
- 27/82 initially believed that they were at "high risk" of developing gangrene. Two weeks after information giving this figure was 19/67.

34/82 initially stated that their varicose veins caused them "significant personal anxiety".

Expectations of surgery

(% given for results of telephone questionnaire in paper, which cannot be converted to a fraction due to uncertainty in the value of the denominator)

- 66/82 initially believed that surgery will improve appearance. Two weeks after information giving this figure was 90%.
- 63/82 initially believed that surgery will improve pain. Two weeks after information giving this figure was 84%

Table 18: Dillon 2005⁸²

Reference	Study type	No of patients	Patient characteristics	Methodology	Source of funding
 62/82 initially 	v believed that surger	y will improve itch. Two	weeks after information	giving this figure was 80%.	
• 63/82 initially	v believed that surger	y will improve heaviness	. Two weeks after inform	ation giving this figure was 86%.	
• 55/82 initially	v believed that surger	y will improve flares. Two	o weeks after informatio	n giving this figure was 31%.	
Other expectatio	ns				
• 57/72 initially	believed that recove	ery after surgery would ta	ake < 2 weeks. Two week	s after information giving this figure was 44/62.	
• 15/72 initially	v believed that return	to work after surgery wo	ould take a month. Two v	veeks after information giving this figure was 17/62.	
		ne interview was poor – o e likely in those who reca	-	ny adverse effects. 50/67 remembered getting the educ	ational leaflet, and recall

Author summary: Patients attending varicose veins clinics have an unrealistic expectation of the benefits of surgery and fail to understand the benign nature of their condition. The outpatient [information-giving] process has little effect on patient-held beliefs.

Palfreyman SJ, Drewery- Carter K, Rigby K, Michaels JA, Tod AM. varicose events a qualitative scondary care trust in Sheffield.16 recruited by a research nurse.Patients who had been referred for varicose vein specialist investigation by their GPs. Reasons for seeking relief of symptoms.Interviews were conducted 5-14 days after attendance at a surgical outpatient clinic. Interviews were semi- structured and conducted in a quiet room within the hospital setting. The interviews were tape-recorded and lasted 30-45 minutes.Northern General Nursing and PAMs Research Grant – no conflict of interest likely.Patients with varicose veins sa puposive sand reasons for seeking treatment. Journal of Clinical Nursing 2004; 13: 332-340.16 recruited by a research and a range of and a range of ages.Patients who had been referred for varicose vein specialist investigation by their GPS. Reasons for seeking relief of symptoms.Interviews were conducted 5-14 days after attendance at a surgical outpatient clinic. Interviews were tape-recorded and lasted 30-45 minutes.Northern General hursing and asted 30-45 minutes.Purposive sand reasons for seeking treatment. Journal of Clinical Nursing 2004; 13: 332-340.Interviews and reasons of time with varicose veins 4-51 years; previous treatments were non (n=7), compression (n=6), treatment (n=3).Interviews the anonymous transcripts and key themes were shown to a subset ofPatients of both genders, and a range of UltrastInterview of the reating for and conservative treatment (n=13) and conservative treatment (n=3).Interviews were tested by feedback to the patient during<
interviewees to check validity of findings

Table 10. Dalf 2004199

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
Positive expe	ctations of treat	ment effects on the	e level current symptoms		
Patients gene	erally had an expe	ectation that somet	hing could be done about their symptor	15.	
"more than	anything is that	it won't be as it is n	ow, so that the pain factor, the heavine	s, everything that goes with it hopefully will have gone	"
Although an o	expectation of co	smetic improveme	nt was also present, this expectation wa	s not the main reason for seeking treatment.	
Positive expe	ectations of prog	nosis if treated			
-			vent future deterioration of symptoms a	nd extent of varicose veins.	
Patients eithe	er had the expect	ation of no possibil	ity of recurrence, or that even a short sy	mptom free period would be worth it.	
Even those w	ith previous surg	ery expected that t	heir surgery this time would work better	, and that even a short symptom free period would be w	orth it.
Nogativo ovo	octations				
Negative exp	ectations				
Negative exp	ectations of prog	gnosis if untreated			
An important	motivation for t	reatment was that	OVT and ulceration could occur later bed	ause of their varicose veins. A particular concern was that	t varicose veins mig
exacerbate th	ne risks of develo	ping a DVT whilst fl	ying.		
Negative exp	ectations about	adverse events of s	urgery		
Fear of surge	ry was common:				
"I'm in the middle now. I'm frightened of having them done and I'm frightened of having them"					
How can the	se expectations b	be addressed?			
Consideratio	n of patient expe	ctations should influ	uence the nature of the nurses' assessm	ent and information giving.	
Author's sum	marv: "thev h	ad actively sought t	reatment from the health service, with	he expectation that they will gain symptom reliefthis w	ish for their
	• •			ce suggests that surgery in particular may not have any effective surgery s	

Evidence tables clinical studies

symptoms to be relieved by treatment might be an unrealistic expectation, as the evidence suggests that surgery in particular may not have any effect on the symptoms experienced in the leg with varicose veins......The patients, in seeking to relieve their symptoms, were after an immediate benefit. This belief meant that they disregarded the potential risks of treatment... the participants were also being unnecessarily anxious about the complications of varicose veins.....such worries were not supported by the evidence....patients over-estimate the extent to which the appearance of their legs can be improved by treatment....."

Table 20: Shepherd 2010²⁴⁷

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
Shepherd AC, Gohel MS, Lim CS, Hamish M, Davies AH. The treatment of varicose veins: an investigation of patient preferences and expectations. Phlebology 2010; 25: 54-65.	Cross-sectional questionnaire survey. Setting: vascular clinic in an NHS secondary care trust.	111. 83 gave complete responses and the remaining 28 gave partial responses.	Consecutive patients referred to one consultant vascular surgeon with symptomatic varicose veins; 73% of patients were female; 43% were unemployed, and 17% were part-time employees; age range 18- 83; reported co- morbidities were: • hypertension - 16% • previous deep vein thrombosis – 7% • asthma – 5% • diabetes – 4% • epilepsy - 2% • Chronic obstructive pulmonary disease (COPD) – 2% • Ischeamic heart disease (IHD) – 2% • Transient ischaemic attacks (TIAs) – 1% • No co-morbidities – 61%	Patients were invited to complete an anonymous questionnaire prior to their consultation (and prior to any information being given out). Questions related to occupation, physical symptoms and impact of the varicose veins, patient knowledge of existing treatments, concerns about complications and recurrence, preferred treatment options and factors that might influence decisions regarding treatment.	None reported

Results:

Negative expectations about varicose veins treatments

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding

The main concerns that patients had about treatment were presented in a low resolution figure, and so exact data are unclear. However, it appears that about 35% were "extremely concerned" about recurrence, and about 16% were "extremely concerned" about discomfort after treatment

Awareness of treatment options

- 86% aware of surgery as an option
- 32% were aware of laser ablation
- 22% were aware of sclerotherapy
- 18% were aware of radiofrequency ablation.
- 10% were unaware of any treatments.

24/103 expressed a preference for endovenous treatments (i.e. endothermal ablation or foam sclerotherapy) over surgery. Of the endovenous treatments, laser was the most popular (first choice of 11%). Most patients (74/103) stated that they didn't know enough to express a treatment preference.

Table 21: Zubilewicz 2009²⁹⁰

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding
Zubilewicz R, Chmiel- Perzynska I, Derkacz M, Schabowski J. The women's span of knowledge about chronic venous disease. Family Medicine and Primary Care Review 2009; 11: 919-922.	Cross-sectional survey. Setting: Poland but no other details provided.	156	Polish women with chronic venous disease (CVD) who had never been treated. Average age was 44.5 (16) years. 19% were <30 years old, 68% were between 31-65 years old and 13% were over 65 years old. 14% had primary education, 47% secondary education or vocational training and 39% had a university degree.	Participants were given a multiple choice questionnaire, which was aimed to assess knowledge concerning modifiable risk factors for chronic venous disease and the presence of symptoms. No other details given.	None reported

Results:

Expectations / preconceived ideas about modifiable risk factors

The following were the most often suggested risk factors for chronic venous disease. The figures given are the percentage of participants believing it was a risk factor:

- Overweight and obesity (85%)
- High heeled footwear (73%)
- Standing and sitting postures at work (71% and 61% respectively)
- Pregnancy (58%)
- Crossing legs (51%)
- Long journeys by car or plane (40%)
- Oral contraceptives (30%)
- Use of depilatory wax (17%)
- Under-floor heating (11%)
- Physical activity (20%)

Reference	Study type	No. of patients	Patient characteristics	Methodology	Source of funding			
>50% of respondents of <65 years assessed CVD as a severe disorder which lessened Quality of Life (QoL). Approximately 70% of women >65 years considered CVD as especially serious.								
	hat CVD was a risk factor for ulce	eration.						
Approximately 70%	of women under 30 years regard	ed CVD as primarily a cosmetic	problem.					

The perception of CVD as a serious disease was higher in those with lower educational attainment.

G.2 Chapter 6 – referral from primary care

G.2.1 Risk factors associated with disease progression

Table 22: Boccalon 1997²⁸

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
Boccalon H, Janbon C, Saumet JL, Tafani A, Roux T, Vilain C. Characteristic s of chronic venous insufficiency in 895 patients followed in general practice. International Angiology 1997; 16: 226-34	Cross- sectional, but contained potentially useful gender and existence of previous thrombus aetiology findings which will also be effectively case- control.	895, drawn from all regions of France. These included 229 who were asymptomati c and without any detectable signs. These 229 are not included in this review analysis.	Chronic venous insufficiency of lower limbs, all of which had been treated with 2 months of daily 1g microflavanoid fractions. Inclusion: >18; at least one symptom from heaviness, pain or night cramps attributable to CVi for at least 1 year; worsened by prolonged standing or sitting, warmth, and improved by elevation, activity or compression; functional discomfort had to be at least 40/100 on a VAS. Exclusion: arteriopathy or neuropathy of the lower limbs.	Gender Age Secondary aetiology	Development of skin changes or ulcerative changes, in terms of being categorised in the three groups: Group 1: no skin changes; group 2: hyper-pigmentation with no ulceration; group 3: more severe skin changes (included "pre-ulcerative*" changes or ulceration). *not explained further.	NA	Not stated
Results: Gender							

Group 1 had 33 men (age 51) and 278 women (age 44); group 2 had 25 men (age 54) and 269 women (age 53) and group 3 had 12 men (age 59) and 49 women (age

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outc	ome measures	Length of follow-up	Source of funding		
67).										
For each group, the % of men and women were:										
	Gro	oup 1 (<c4)< th=""><th>Group 2 (skin cl ulceration or ulc</th><th>nanges not including pre- ceration)</th><th></th><th colspan="2">Group 3 (more severe skin changes including ulceration or ulceration)</th><th>cluding pre-</th></c4)<>	Group 2 (skin cl ulceration or ulc	nanges not including pre- ceration)		Group 3 (more severe skin changes including ulceration or ulceration)		cluding pre-		
men	47.	1%	35.7%			17.1%				
women	46.	6%	45.1%			8.2%				

Secondary aetiology

The percentage with secondary CVI were: group 1: 3.2%; group 2: 9.9%; group 3: 27.9%. The paper reported that a previous episode of DVT was more commonly reported in the history of patients with the most severe objective signs and that this was significant (p<0.001).

<u>Age</u>

The severity increased with age. Gp 1 were 45 (14) yrs, group 2 were 53 (15) yrs and Group 3 were 65 (13) years.

Other factors

Other factors were considered but they were cross-sectional and so do not indicate prognosis for progression.

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
Pannier F, Rabe E. progression of Chronic /enous Disorders: Results from the Bonn /ein Study. ournal of /ascular Surgery 2011; 53: 254-255	Prospective cohort study	3072 enrolled at baseline. 1978 remained in the study at follow-up. (These were a cross-section of all people, as varicose veins were not an inclusion criterion). The relevant figure is 290, however, as this represents the number with C2 at baseline (and who also attended at follow-up). 432 had C2 at baseline, indicating very high attrition of 142 participants. Reasons for attrition not reported.	Participants were sampled randomly from the population aged 18-79 years living in Bonn and two rural townships. 3072 represented a response rate of 59%. The age and gender were representative of the general German population. Inclusion: 18-79 years; German nationality. [Note varicose veins or CVI were not an inclusion criterion] Exclusion: hemiparesis/leg amputations; severe illness; moribund patients; systemic inactivating disease. Baseline Characteristics: For those attending at baseline, • 56.2% were male, • 33.6% were 18-39 years, • 37.4% were 40-59 years and • 29% were 60-79 years. • 43.9% had a BMI of <25. • 9.6% were C0, • 59% were C1, • 14.3% were C2,	A standardised questionnaire was used to collect information at baseline on the following risk factors: • sociodemographic status • smoking • alcohol • physical activity • blood pressure • medical history • quality of life • hormonal intake • contraceptive pill • professional stress/ work strenuousness • BMI • Heaviness • Feeling of tension • Swelling feeling • Pain during prolonged walking • Itching	Progression from C2 to C3- 6 over the 6.6 years	6.6 years	Not stated

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding			
			13.55 were C3,2.9% were C4 and							
			• 0.7% were C5-6.							
			• 62.4% lived in urban areas.							
			No information reported on the characteristics of the 290							
			relevant C2 participants remaining at 6.6 year follow-up.							
Results: ** Th	ese data has b	een removed as it is ac	ademic in confidence							
lisk factor			RR (95% CI) of the progression from C2 to C3-6							
** <mark>These data </mark> confidence	has been remov	ved as it is academic in								

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
-	but cross-sectional analyses included a well (see risk factor studied section). Potential confounders such as socioeconomic status were	participate, leaving 240. This represents a respons rate of 63%. 120 were C6 or C5*[cases]. Of these, 24 had had previous surgery Mean age of first developing an ulcer was 56 (15.5) years [approx 8 years prior to study, on average] – this means any retrospective recollection of <8 years previously would be unlikely to be representative of true "causes". Median (IQR) of 2(1-3) active episodes, each of a mean (sd) duration of 7(13) years.	recruited from the register of venous patients scanned in a vascular laboratory a a large Scottish NHS trust, as well as GP practices in a Scottis region. The cases were to have an open or healed ulcer (C5/6), and the ulcer was to have been active for at least 8 weeks.	gender, height and age (these could no	versus no ulceration.	Unclear, as patients were simply asked to recollect activity data when aged 35-45 years – the time duration back to this would have varied widely, and some patients mannot have even reached this age range.	None

Table 24: Robertson 2009²³⁰

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
	Poor attempt to ensure direction of any cause-effect was unambiguous. No reports of blinding of assessors. Overall, very low quality.	bilaterally affected, only	but were kept in the analysis on the basis that they formed a small percentage	ages of 35 and 45. However, a small proportion (probably <25%) of patients were within those age groups at the time of assessment and so, for those patients, these measures were cross- sectional. The cause-effect status of smoking history was fairly clear given that the mean pack years were around 16. Pre-ulcer weight was another informative factor. Cause-effect was unclear for previous history of DVT/PE or phlebitis as it was very unclear whether these were antecedents of ulceration, or merely an uncongenial accompaniment.			

Results. <u>ONLY results that pertain to risk factors that are likely to have preceded ulceration are included</u>. Cross-sectional data (except for those where the possible direction of cause-effect is fixed, such as gender) are not included as they are of no relevance to the issue of prognosis. Relevant univariable results (only adjusted for age and sex) are given below.

Multivariable results are not given as no potentially prognostic factors remained in the model after stepwise removal. All the results presented here are univariate results

Risk factor	mean (sd) RF in case / % with RF	mean (sd) of RF in controls/ % with RF	P value
Age	64.1(13.4)	59.9(11.7)	0.01
% male	55%	43%	0.07
Risk factor	mean (sd) RF in case	mean (sd) of RF in controls	OR (CI) [univariable in terms of no adjustment for other RF, except age and sex] ^b

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
height		1.67(0.11)	1.67(0.1)	0.54(0.02-19.41)			
Smoking pack	k years	4.47 (3.16-6.32)	4.1(2.45-5.48)	1.08(0.9-1.29)			
Risk factor		%with the RF in case	% with RF in controls	OR (CI) [univariable in terms of no ad	justment for a	other RF, except age	and sex] ^b
Physical exer	cise in past year ^a						
Nil	. ,	28.8	14.9	Reference			
Light		35.6	42.1	0.44(0.21-0.91)			
Mod		28.2	36.8	0.43(0.20-0.93)			
Strenuous		6.8	6.1	0.70(0.20-2.41)			
Physical exer	cise aged 35-45						
Nil	-	15.3	14.0	Reference			
Light		28	28.9	0.86(0.37-2.01)			
Moderate		39.8	44.7	0.76(0.34-1.68)			
Strenuous		16.9	12.3	1.29(0.48-3.49)			
Daily activity	in past year ^a						
sitting		35.6	17.5	Reference			
walking		48.3	60.5	0.43 90.22-0.820			
light loads		7.6	15.8	0.29(0.11-0.77)			
heavy work		8.5	6.1	0.99(0.29-3.35)			
Daily activity	aged 35-45						
sitting		16.1	14.9	Reference			
walking		47.5	44.7	1.09(0.49-2.41)			

Reference	Study type	No. of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
light loads		14.4	18.4	0.79(0.31-2.03)			
heavy work		22.0	21.9	0.86(0.35-2.10)			
Risk factor (R	F)	%with the RF in case	% with RF in controls	P value			
History of phl	ebitis	37	28	NS			
History of leg	fracture	18	11	NS			
History of art	hritis	40	35	NS			
Ever smoked		63.6	45.6	0.009			

(a) Unlikely to have preceded ulceration but included for completeness(b) The OR is the odds ratio of ulceration for every additional increment of the continuous variable [adjusted for age and sex]

Reference	Study type	No. of patients	Patient cha	aracteristics		Risk factors studied	Outcome measures	Length of follow-up	Source of funding
Scott TE, LaMorte WW, Gorin DR, Menzoian JO. Risk factors for chronic venous insufficience y: a dual case- control study. J Vasc Surg 1995; 22:622-8	components. Potential confounders such as socioeconomic status, age, BMI etc were not matched. A multivariable analysis was performed	129 with varicose veins and 93 with chronic venous insufficiency "CVI". All those with CVI had ulceration. There was also a group of subject with no venous disease (used to assess risks for <i>initially</i> <i>developing</i> venous disease, so not relevant to this review question) that is not included here. Rates of refusal were described as <5% and to be the same across the groups.	English; un Cases: Pati cared for ir medical ce ulcers, but described a based on v Controls: F	n vascular surge ntre in the USA. elsewhere in th as having ulcera isible appearance ratients with var the same clinic a CVI (with ulceration) 59(1.6) 30.4(1) 58% 51.6% 25990 (1080)	ormed consent. II or class III CVI, ry clinics at a large Unclear if all had the paper they are tion. Diagnosis ce.	Retrospective information on potential risk factors was done via a structured interview, by an interviewer blinded to the status of the patients. These included medical history, years of smoking, standing at work and exercise levels. Of these, standing at work and exercise were cross- sectional and so not included in this review.	Existence of ulceration	Unclear. The only retrospective questions were medical history and years smoked, and the distance back into the past these variables occupied was unspecified.	None stated

Table 25: Scott et al. 1995²⁴²

Reference	Study type	No. of patients	Patient ch	aracteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
	blinded to the group statue of the subjects.							
	Overall, very low quality.							

Results: <u>ONLY results that pertain to variables that are likely to have preceded ulceration are included</u>. Cross-sectional data (except for those where the direction of cause-effect is fixed, such as gender) are not included as they are of no relevance to the issue of prognosis. Relevant univariable results (only adjusted for age and sex) are given below.

Risk factor	CVI %	Varicose vein %	Significant difference? (p values not stated)
History of heart disease	22.6%	4.6%	Y
History of diabetes mellitus	22.6%	2.3%	Y
History of hypertension	49.5%	16.3%	Y
History of kidney disease	4.4%	2.3%	Ν
History of arthritis	19.7%	13.9%	Ν
History of leg injury	54.8%	17.8%	Υ
History of phlebitis/clot	45.6%	24.2%	Υ
History of oral contraceptive use	5.1%	20.7%	Υ
years smoked	17 (1.7)	8.8(1.0)	Υ
Multivariable analysis results			
Risk factor	OR for ulceration		
age	1.07/yr (1.04-1.1)		
male sex	8 (3.5-18.3)		
BMI	1.07/kg/m ² (1.01-1.13)		
no health insurance*	3.2 (1.3-7.7)		
history of leg injury	4.7 (2.1-10.5)		
Diabetes mellitus	4.3 (0.99-18.7)		

G.2.2 Factors associated with response to treatment

Table 26: Fischer 2006⁹⁸

Reference	Study type	No of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
Fischer R, Chandler JG, Stenger D, Puhan MA, De Maeseneer MG, Schimmelpfennig L. Patient characteristics and physician- determined variables affecting saphenofemoral reflux recurrence after ligation and stripping of the great saphenous vein. J Vasc Surg 2006; 43: 81-7	Prospective observation al study. Main aim was to evaluate modifiers of treatment success.	n=1261 patients /1638 limbs. Unspecified attrition, but sophisticate d imputation used.	Patients undergoing SFJ ligation and GSV stripping, from 1978 to 2003. Inclusion: Primary operations Exclusion: History of DVT, or serious trauma to the affected leg; procedures involving crossectomy but not GSV stripping. Baseline Characteristics: Mean age 49.7(12) at the time of operation	BMI, prior parity, interim pregnancy, deep venous insufficiency, age, gender, side affected, diabetes mellitus.	Saphenofemoral reflux recurrence, using duplex, but, in earlier cases continuous wave doppler. Reflux had to last >0.5 seconds.	Variable. All follow-ups were after 1991. Categorised as 2- 6 yrs, 7-12 yrs and > 12 years. follow-up duration was normalised through adjustment in the multivariable analysis. Mean was 6.6(4.3)yrs.	

Results:

BMI>29, prior parity, and interim pregnancy were all associated with an increased odds of reflux recurrence. The table below shows the results of the multivariable logistic regression, with odds for recurrence of SFJ reflux recurrence at a mean of 6.6 years shown for relevant patient-related variables.

Variable	OR (95% Cls)
BMI >29 at baseline (compared to <29)	1.65(1.12,2.43)
Prior parity (compared to none)	2.69(1.45,4.97)
Interim pregnancy (compared to not)*	4.74(2.47, 9.12)

*not a variable that can predict treatment efficacy at the pre-treatment stage, so excluded from results in review.

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
Gibson KD, Ferris BL, Polissar N, Neradilek B, Pepper D. Endovenous laser treatment of the short saphenous vein: efficacy and complications . J Vasc Surg 2007; 45: 795-803	Prospective consecutive enrolment of patients. Main aim was to evaluate treatment success and modifiers of AEs.	n=187 patients/210 legs. High (40%) attrition by the stage of the final follow-up (2- 11 months).	C2-6 patients undergoing EVLA	Anatomic patterns of the SSV. Type A was a SPJ with no significant branches; type B was a SPJ with a large extension Giacomini vein; type C was a SPJ or SFJ with no direct termination into a deep vein, and the SSV continued as a Giacomini vein above the popliteal fossa. Author's own classification system. Also: gender, leg side, preoperative presence of ulcer, pre-op presence of stasis, pre-op presence of pain, and age.	Incidence of the adverse event of DVT at 2-4 days (but unclear) Recanalisation at 2-11 months (but unclear)	4-10 months	None stated

Table 27: Gibson et al. 2007¹⁰³

Results:

DVT risk factors

SSV anatomy had an association with DVT incidence. The risks of DVT for each group were as follows: Group A: 10/88 (11.4%); Group B: 2/69 (2.9%); Group C: 0/52 (0%). Specifically, a SPJ with no significant branches (Type A) carried a trend (p=0.07) for a higher risk than type B [type B compared to type A, for risk of DVT: OR:0.23(0.05, 1.10)]. There were no DVT cases in type C, so no ORs could be produced, but the Fisher exact test showed that type C had a significantly lower risk than type A (p=0.013).

No multivariable results are given, but this is because the only variable that had a p<0.1 on univariate testing was SSV anatomy type. Hence no variables other than SSV

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding			
	ould have been p Its are given belo		iable model – hence the un	ivariable results for SSV anato	my type are the full results.	. For completeness,	all			
Risk factor for DVT (reference given in brackets)			OR (95% CI) for DVT	OR (95% CI) for DVT at variable time (adjusted for time)						
right side (comp	right side (compared to left)			0.64(0.20, 2.09)						
stasis (compare	d to no stasis)		0.46 (0.1, 2.16)	0.46 (0.1, 2.16)						
Age (per 10 year	r increment)		0.99(0.62,1.57)	0.99(0.62,1.57)						
		pe A) [there were ould be produced]								
Gender			0/28 DVTs in men, 12	0/28 DVTs in men, 12/182 DVTs in women, p=0.4*						
Pre-op ulcer			0/11 DVTs in those w	0/11 DVTs in those with ulcers, 12/199 DVTs in those with no ulcers, p=0.5*						
Pain			0/13 DVTs in those w	0/13 DVTs in those with pain, 12/197 DVTs in those with no pain, $p=0.5^*$						
ulcer, stasis or pain			0/11 DVTs in those w	0/11 DVTs in those with ulcers, stasis or pain $12/199$ DVTs in those with no ulcers, stasis or pain , p=0.5*						

Recanalisation risk factors

A logistic regression analysis using the same risk factors was carried out to evaluate their effects on the odds of recanalisation. No results were reported, other than that none of the variables had a significant relationship with recanalisation.

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
Gonzalez- Zeh R, Armisen R, Barahona S. Endovenous laser and echo-guided foam ablation in great saphenous vein reflux: one year follow-up results. J Vasc Surg 2008; 48: 940-6	Non-randomised trial with main aim of comparing 2 treatments, but with logistic regression analysis included to assess effects of potential treatment modifiers. Only one limb per patient was included and treated in this study. A single surgeon with experience of 800 EVLA s and 2000 foam sclerotherapies did both interventions. Patients were not allowed to mix, to avoid contamination of patient expectations.	98. No patients dropped out and all followed up.	C2-6 patients undergoing EVLA and foam sclerotherapy. Patients were allowed to choose between foam sclerotherapy and EVLA, and they were told the efficacy of each was equivalent. Inclusion: Primary incompetence of the GSV and SFJ insufficiency with a reflux time of 0.5 seconds measured over a distance of at least 20cm in the upper leg. Exclusion: pregnancy; active thrombophlebitis, clotting disturbances; thrombophilia or coagulation disorders; History of DVT; history of malignancies. Baseline characteristics: Despite the lack of randomisation the groups were well matched.	Clinical grouping (C1- 6), pre-op VCSS, age, pre-op GSV diameter	Presence of reflux, as measured by duplex.	1 week, 1 month, 6 months and 1 year.	Not state

Table 28: Gonzalez-Zeh et al. 2008¹⁰⁷

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
	ultrasound follow- ups done by an assessor blinded to treatment, but probably not to baseline predictors.						

Results:

Subgroups analysis showed that a larger pre-op GSV diameter was associated with reflux in both the foam and EVLA treatments. Veins <6.5cm have a 90% success rate with foam, and veins <12mm have a 90% success rate with laser.

Logistic regression analysis showed that for each treatment, pre-op GSV diameter (?>12mm, unclear) was the only factor significantly predicting reflux. The multivariable results for each treatment separately are given below. The OR(95% CIs) are for the odds of reflux. The analysis is unclearly reported. The reference values for categorical variables (Clinical groups, GSV diameter) are unclear. It is likely that the reference value of GSV diameter is <12mm (therefore the variable below is given as GSV >12mm). Though not stated it is likely that the ORs for the continuous variables (age, VCSS) are per increment increase in the variable.

Variable	Foam sclerotherapy	laser
clinical groups C1-6	0.89(0.39-2.20)	2.87(0.33-24.77)
VCSS	0.97(0.44-2.15)	0.31(0.03-3.12)
Age	0.99(0.91-1.08)	0.94(0.79-1.09)
GSV diameter (>12mm?)	1.68(1.24-2.27)	1.91(1.02-3.59)

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
Islamoglu F. An alternative treatment for varicose veins: ligation plus foam sclerotherapy . Dermatol surg 2011; 37: 470-479	Prospective non- randomised study. Patients were allowed to choose treatments. The main aim was the comparison of stripping versus foam sclerotherapy and crossectomy, but in the absence of a differential treatment effect most of the results sections focus on the non-treatment predictors of treatment success/failure.	372. No mention of drop-outs. Unclear if the sample were defined by completers only.	C2-6 patients undergoing foam sclerotherapy with crossectomy or classic stripping. All done by the same surgeon. Mean age 48.6(10.1). 159/372 male. 156/372 in sclerotherapy group. All symptomatic. Bilateral in 51 subjects. Inclusion: GSV reflux; C2-6; primary aetiology. Exclusion: pregnancy; sclerosant allergy; acute thrombophlebitis; acute DVT; local infection; immobility.	Unilateral/bilateral, pre-operative CEAP, employment, familial predisposition, gender, DVT, age, pre-operative deep venous insufficiency (DVI), pre-operative perforator incompetence (PI).	Symptom recurrence, post- operative CEAP, post-operative Perforator incompetence.	6 months, and at further 6 month intervals (mean follow-up was 10.2 (5.1) months.	Not stated

Table 29: Islamoglu 2011¹²²

Results

Multivariable results only, all adjusted for treatment type (always NS in all analyses) as well as other variables. The time of follow-up is unclear, but presumably 6-12 months.

Post-op symptom recurrence

These results were poorly reported by the paper. The directions of the ORs in the text do not tally with the raw data for the unilateral/bilateral variable. The direction of effect given below is that determined by the raw data. ORs are for the existence of post-operative symptom recurrence.

Variable	OR (95% Cls)
unilateral (versus bilateral)	2.376 (1.682-3.356)

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding	
Pre-op CEAP <u>></u> 3	3 (versus <3)	3.298(1.897-5.731)						
No job (versus	a job)	0.133(0.073-0.243)						
No family histo history)	ry (versus a family	0.357(0.198-0.643)						
Post-op CEAP <	<u>: 3</u>							
Again, poorly r	eported. ORs are fo	r post op CEAP <3						
Variable OR (95% CIs)								
unilateral (vers	us bilateral)	2.497(1.337-4.663)						
Pre-op CEAP <3	3 (versus <u>></u> 3)	1.445(0.368-4.818)						
male (versus fe	emale)	1.542(0.201-3.355)						
No previous D\ previous DVT)	/T (versus	2.827(0.831-9.619)						
Age <60 (versu	s >60)	1.215(0.262-4.012)						
Post-op perfor	ator incompetence							
Age >60 (comp	ared to <60)	23.618(8.423-66.223)						
Pre-op CEAP >3 <3)	3 (compared to	2.741(1.174-6.401)						
No job (compa	red to employed)	0.112(0.039-0.317)						
Family history (none)	(compared to	2.927(1.020-8.398)						
Pre-op PI (com	pared to none)	6.102(2.214-16.815)						

Evidence tables clinical studies

There was also a multivariable analysis evaluating which factors were associated with earlier (<1 year) or later symptom recurrence, amongst the sub-group with symptom recurrence. Results of this have not been included in this review because they are outside the scope of the review question.

Reference	Study type	No. of patients	Patient characteristics	Patient-related risk factors studied	Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
AJ, Paisley A et al.	consecutive unselected patients aiming to look for factors influencing disease-specific quality of life. This included patient factors, as well as operative and	•	undergoing GSV, SSV o SEPS surgery. GSV surgery comprised flus SPJ ligation, stripping o	age, gender, pre-operative AVVSSS (high = worse), CEAP grade, first time/recurrent, History of DVT. In patients with bilateral disease, the factors entered into the analysis were those for the worst affected leg.	Post-operative AVVQ	6 months/ 2 yea	None stated.

Table 30: McKenzie et al. 2002¹⁵⁶

Results:

6 months multivariable

A higher baseline AVVQ, baseline recurrent disease and baseline CEAP 4 disease predicted higher (worse) AVVQ at 6 months. This model explained 60% of the total variation in AVVQ at 6 months. Square root used to normalise the distribution of baseline AVVQ (Log not possible as raw scores included zero)

Factor	Parameter estimate	SE	t	р
square root of baseline AVVQ	0.57	0.07	7.78	< 0.001
primary/recurrent procedure	0.45	0.17	0.15	0.009
CEAP 4	0.39	0.17	0.14	0.026

2 years multivariable

A higher baseline AVVSS and baseline CEAP 5 disease predicted higher (worse) AVVQ at 2 year. In contrast, previous GSV surgery predicted a lower AVVQ. This model explained 47% of the total variation in AVVQ at 2 years. Square root used to normalise the distribution of baseline AVVQ (Log not possible as raw scores included zero)

Factor	Parameter estimate	SE	t	p
square root of baseline AVVQ	0.47	0.08	6.16	<0.001

Reference	Study type	No. of patients	Patient ch	aracteristics			Outcome measures	Length of prospective follow-up / retrospective recollection	Source of funding
GSV surgery		-0.73		0.31	-2.35	0.02			
CEAP 5		0.62		0.28	2.19	0.030			

Reference	Study type	No of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
Myers KA, Jolley D, Clough A, Kirwan J. Outcome of ultrasound- guided sclerotherapy for varicose veins: medium-term results assessed by ultrasound surveillance. Eur J Vasc Endovasc Surg 2007; 33: 116-121	Prospective observational study. Main aim was to evaluate modifiers of treatment success.	489 patients (677 limbs). Time to event study so attrition catered for in analysis.	Inclusion: C2-6 patients undergoing ultrasound guided sclerotherapy (mainly foam but some liquid). Some of these were given over 3-4 separate sessions. Exclusion: Previous EVLA Baseline Characteristics: Age range 19-92 (median 53); women: 401/489; C2-3 in 90%; 115 limbs were recurrent and the rest were first-time.	Type of vein, age, gender, diameter of GSV, side, CEAP grade.	Time to failure was the outcome. Treatment success defined as persistent occlusion or absence of reflux in treated veins – assessed by ultrasound (unclear if duplex). Time to failure was therefore the duration between the first treatment session (out of the 1-4) achieving full success and the first follow-up when reflux was noted.	Every 3-5 days after each of the 1-4 sclerotherapy sessions; then at 6 weeks; and then at 6 months for 2 years; and then annually.	Not stated

Results: A multivariable cox-regression analysis was carried out for factors influencing failure in all saphenous veins. The table below summarises the results, with a higher HR indicating a greater risk of failure at any point in time compared to the reference category. Younger age and larger (>6mm) diameter GSV were associated with a worse outcome.

Variable (and reference category)	Index category of variable	n	Hazard Ratio (95% CI)
Age (compared to 50-59)	<40	93	2.16(1.27,3.66)
	40-49	121	1.11(0.69,1.78)
	60-69	118	1.22(0.79,1.89)
	70+	87	0.63(0.35,1.14)
Sex (compared to female)	Male	112	1.31(0.88,1.94)

Reference	Study type	No of patients	Patie	nt characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
Side (compared	l to left)	Right		313	1.19 (0.89, 1.57)			
Vein (compared to GSV)		SSV		174	1.58(1.11, 2.24)			
CEAP (compare	CEAP (compared to C2/3)			62	1.57(0.91, 2.73)			
vein diameter (vein diameter (compared to			152	1.27((0.79,2.03)			
<5mm)		6mm		152	2.07(1.35, 3.18)			
		>6mm		112	2.22(1.4, 3.5)			

Reference Stud type	-	No of patients	Patient characteristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding
	ospec e hort idy.	116 patients, having 126 procedures. Appears to have very high attrition, as only 116/235 eligible patients attended follow-up. These could have included the worst (or best) responders. But how many of these eligible patients were actually recruited in the first place? Very unclear. Unclear if analysis was by procedure (n=126) or patient (n=116), but likely to be the latter, as there would probably only have been one outcome assessment per person, and thus one analysis (for example if a patient had two UGFS procedures, the second would be a top up and a single outcome would relate to both).	53 men, 63 women. Median age was 55 (range 18-80). Target veins were the GSV (n=75), SSV (n=13), and accessory GSV (n=8). Others involved other veins or more than a single target vein (n=30).	Gender Previous surgery Sites of injection Maximum concentration of sclerosant Pre-procedure CEAP Compliance with post treatment compression Age Volume of sclerosant	Successful outcome – complete occlusion of the target vein on duplex analysis on follow-up. Existence of any complications Existence of each complication analysed separately (superficial thrombophlebitis, pain, skin staining, DVT, allergy and skin blistering)	3 months minimum	None.

Evidence tables clinical studies

Table 32: Thomasset 2010²⁶⁸

Results: Analysis was poorly reported though it seems univariate analyses for the 8 risk factors were performed. Although this study did therefore not meet the inclusion criterion of having a multivariable analysis, because only one risk factor was significant on univariate testing, a multivariable analysis would have been an unnecessary next step anyway, so this study has been included.

For the outcome of complete occlusion of the target vein, the only risk factor associated was compliance with post-procedure compression hosiery (p<0.05). No effect sizes were presented. This is not a factor that could be ascertained pre-treatment (although it is conceivable that patients could be asked if they thought they'd be compliant with stockings after treatment) and so has little value in making a pre-treatment prediction about which patients will do well.

For the outcome of any complication, female gender was associated with a greater risk (p<0.05). No effect size was reported. For each complication considered separately, female gender was associated with skin staining (P<0.05). No effect sizes were given. There were no associations between female gender and any other complications considered singly.

G.3 Chapter 7 – assessment for treatment

.1 Diagnostic accuracy of hand held Doppler ultrasound

Table 33: Campbell 1997⁴⁴

Reference	Study type	No of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Campbell WB, Niblett PG, Ridler BMF, Peters AS, Thomson JF. Hand held doppler as a screening test in primary varicose veins. British Journal of Surgery 1997; 84: 1541- 1543	Diagnosti c review study.	85 (122 legs)	Patients referred to the vascular outpatient clinic with primary and previously un- operated varicose veins. Gender: 52 women Age:range18-89 (median 53).	Hand held doppler. Performed in standing. Tourniquet test used for the GSV. Positioning not given. >1 second reflux regarded as significant. Carried out by consultant(103 legs), trainee (in 17) and unknown (2 legs).	Duplex, using a Diasonics VST masters scanner with a 5MHz linear array probe. Positioning not given. >1 second reflux regarded as significant. Duplex operator not reported.	Blinding NOT stated Test interval <u>not</u> clear: "another visit". Expertise comparability <u>not</u> clear. No previous treatments. CEAP status not reported.	Sensitivity and specificity	Not reported

Results: Raw data only available for the popliteal fossa (percentages given for the GSV, but not possible to convert these to raw numbers due to lack of data on the numbers with duplex-confirmed reflux).

popliteal fossa	+ve on duplex	-ve on duplex			
+ve onHHD	28	8			
-ve on HHD	11	74			

Popliteal fossa +ve and –ve predictive values, and all CIs, derived from raw data. No raw data given for groin or 10cm below groin.

Site examined	sensitivity	Specificity	Positive predictive value	Negative predictive value
Great Saphenous Vein	0.86	0.82	-	-
10cm below groin (alternative GSV)	0.81	0.85	-	-
Popliteal fossa	0.72 (0.55-0.85)	0.90(0.82-0.96)	0.78(0.62-0.88)	0.87(0.78-0.93)

Table 34: Darke	≥ 1997 ⁶⁴							
Reference	Study type	No of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Darke SG, Vetrivel S, Foy DMA, Smith S, Baker S. A comparison of duplex scanning and continuous wave doppler in the assessment of primary and uncomplicated varicose veins. Eur J Vasc Endovasc Surg 1997; 14: 457- 461	Diagnostic accuracy study	73 patients (100 legs)	73 patients referred to a consultant vascular surgeon with primary uncomplicated varicose veins. Gender: 55 females; Age: mean 47.5 (range 22-74);	continuous wave Doppler, using a Huntleigh dopplex 500 probe at 8MHz. Positioning not given. Reflux definition not described in terms of duration. Carried out by a "single observer".	Duplex, using an Acuson 128/10 colour duplex scanner with a 7MHz linear array probe. This was carried out blind to the doppler findings. Reflux defined as >0.5 secs. Carried out by a medical technologist.	Blinding carried out Test interval not stated Expertise comparability <u>not</u> clear. Stage of disease and previous treatment history not given.	Sensitivity and specificity	Not stated

Results: No CIs provided in the paper. The raw data below were gathered from the paper, and the CIs were calculated.

Great saphenous vein	+ve on duplex	-ve on duplex	Short saphenous vein	+ve on duplex	-ve on duplex	
+ve on HHD	83	0	+ve on HHD	19	5	
-ve on HHD	4	13	-ve on HHD	2	74	

In paper only sensitivity and specificity provided, but +ve and –ve predictive values have been calculated from the raw values.

Site examined	sensitivity [TP/TP+FN]	Specificity [TN/TN+FP]	+ve predictive value [TP/TP+FP]	-ve predictive value
				[TN/TN+FN]

Evidence tables clinical studies

Reference	Study type	No of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Great saphenous vein	[<i>83/83+4</i>] 0.95(0.89-0.9	99)	[<i>13/13+0</i>]. 1.00(0.75-1.00)		[<i>83/83+0]</i> 1.00(0.95-1.00))		[13/13+4]. 0.75(0.52-0.89	9)
Short saphenous vein	[19/19+2] 0.90(0.70-0.9	99)	[74/74+5] 0.94(0.86-0.98)				[74/74+2] 0.97(0.91-0.99)	

Table 35: DePal	1111 1995							
Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
DePalma RG, Hart MT, Zanin L and Massarin EH. Physical examination, doppler ultrasound and colour flow duplex scanning: guides to therapy for primary varicose veins. Phlebology 1993; 8: 7-11.	Diagnostic review study.	40 (80 legs)	Symptomatic patients presenting with primary varicosities in the great saphenous distribution. Gender: 31 women; Age: 27-64 yrs; All had mild- moderate symptoms. Typical symptoms were aching in the evening. 22/80 limbs had had prior stripping, but were still symptomatic.	Hand held 9.1 MHz CW Doppler pencil probe at an acute angle of 30-45 deg. Patient positioning not described. No definition of reflux duration threshold. Carried out by senior author, who was probably a vascular surgeon, but unclear.	Duplex, with a QUAD-1 colour flow scanner, with 5MHz probe. Carried out in standing and supine. No definition of reflux duration threshold. Carried out by 2 vascular technical observers.	Blinding carried out Test interval <u>not</u> stated Expertise comparability <u>not</u> clear. 28% with prior stripping. CEAP status not reported	Sensitivity and specificity, positive predictive value, negative predictive value	Not reported
Results: Raw data	а							
SFJ	+ve on duplex	-ve on duplex		SFJ in sub-group stripping n=22	with previous	+ve on duplex	-ve on duplex	
+ve onHHD	24	5		+ve onHHD		8	1	
-ve on HHD	26	25		-ve on HHD		9	4	

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
No CIs given in pa	per. Cls calculated	from raw values.						
Site examined	sensitivity [TP/TP	+FN]	Specificity [TN/TN	N+FP]	Positive predictiv [TP/TP+FP]	e value	Negative predicti [TN/TN+FN]	ve value
SFJ n=80 limbs	[24/24+26] 0.48(0).34-0.63)	[25/25+5] 0.83(0.	65-0.94)	[24/24+5] 0.83(0.	66-0.92)	[25/25+26] 0.49(0	0.36-0.62)
SFJ in sub-group with previous stripping n=22	[8/8+9] 0.47(0.26	8/8+9] 0.47(0.26-0.69)		[4/4+1] 0.80(0.38-0.96)		-0.98)	[4/4+9] 0.31(0.13-0.58)	

Reference	, ,,	No. of Datients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Cent PJ, Weston AJ. Duplex canning may be used electively in batients with primary varicose veins. Ann R Coll Surg Engl 1998;80: 888-393	study. (1	72 patients 108 imbs)	People with primary varicose veins, who had not undergone previous injection sclerotherapy or surgical treatment. Gender: 20 males and 52 females. Median age: 44.5 years (range 19-73 years). CEAP stage (limbs): C1: 1/108 C2: 96/108 C3: 0/108 C4: 9/108 C5; 0/108 C6: 2/108	Hand held Doppler, with 8MHz probe (Multi-Duplex). Carried out by one consultant vascular surgeon. Measurement performed in the standing position, with the affected limb slightly flexed at hip and knee. The probe placed over the sapheno- femoral junction and the calf compressed. Reflux lasting longer than 0.5 seconds was regarded as significant. This was then repeated at the great saphenous vein.	Duplex (with guided pulse wave spectral doppler), using a Siemens Q2000 machine, with a 5 MHz curvilinear probe. Patient measured in standing with weight off the affected limb. Reversed flow of over 1 second was considered abnormal. Carried out immediately after hand held Doppler scanning. This was carried out by another consultant radiologist who was unaware of the results of the HHD assessment	Blinding carried out Tests followed each other immediately Expertise of operators comparable No previous treatment and mostly CEAP stage 2	Sensitivity, specificity. Positive predictive value and negative predictive value of hand held Doppler.	None stated

Table 26. Kent 1000131

Results: HHD diagnostic accuracy compared to gold standard of Duplex. <u>This study did not report the raw data</u>. The data below is all that was presented. (* with tourniquet)

(* with tournique	:()					
Site examined	sensitivity	specificity	Positive predictive	negative predictive		
			value	value		
SFJ	0.93	0.91	0.96	0.86		

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
GSV	0.95	0.68	0.91	0.81				
MTP*	0.87	0.26	0.16	0.92				
SPJ	0.82	0.80	0.44	0.96				
PV*	0.50	0.90	0.44	0.92				

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Kim J, Richards S, Kent PJ. Clinical examination of varicose veins – a validation study. Ann Royal College Surgery Engl 2000; 82: 171- 175.	Diagnostic accuracy study.	44 patients (70 limbs)	Primary and previously untreated varicose veins presenting for Duplex scanning were tested. Secondary varicose veins and previous surgery patients excluded. CEAP stages: C1: 2/70, C2: 67/70 C3: 1/70.	Hand held Doppler with 8MHz probe (Huntleigh technologies). Patient stood on unaffected leg. Probe placed on sapheno-femoral junction. Calf squeezed, and subsequent reflux of <0.5 sec was deemed significant. Then repeated over the GSV and SPJ. Carried out by house officer	Duplex (with guided pulse wave spectral doppler), using a Diagnostic US systems 3535 machine (B&K Medical, Denmark) machine, with a 5 MHz curvilinear probe. Reversed flow of over 1 second was considered abnormal. Carried out immediately after hand held Doppler scanning. This was carried out by a vascular technologist who was unaware of the results of the hand held Doppler assessment	Blinding carried out Tests followed each other immediately Expertise of operators <u>not</u> comparable. No previous treatment and mostly C2	Sensitivity, specificity. Positive predictive value and negative predictive value of hand held Doppler	None stated
		ort the raw d		is all that was presente				
Site examined	sensitivity		specificity		Positive predictive val	ue	Negative predicti	ve value
SFJ	0.97		0.73		0.80		0.96	
GSV	0.82		0.92		0.84		0.74	
SPJ	0.80		0.90		0.57		0.97	

Evidence tables clinical studies

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Mercer KG, Scott DJA, Berridge DC. Pre-operative duplex imaging is required before all operations for primary varicose veins. British journal of Surgery 1998; 85: 1495-1497.	Diagnostic accuracy study.	61 patients (81 legs)	Primary varicose veins.	Hand held doppler, with 8MHz probe (Multi-Dopplex). Carried out by one consultant vascular surgeon. In standing with the affected leg slightly flexed, reflux looked for at sapheno- femoral junction, great saphenous vein and sapheno-popliteal junction. Reflux >0.5 sec regarded as significant.	At a separate appointment (time after not described) Colour flow duplex using a Siemens Quantum 2000 or B&K 3535 (with Acuson 128 5MHz curvilinear probe). Positioning unclear. Carried out by a consultant vascular radiologist. Reflux >0.5 sec regarded as significant.	Blinding definitely NOT carried out (duplex operator reported as having access to hand held Doppler results) Test interval unclear, but described as at a separate appointment Expertise of operators comparable. Treatment history and stage of disease unclear	Sensitivity, specificity of hand held Doppler	Not reported
Results: Raw res	ults:							
SFJ	+ve on duplex	-ve on duplex			SPJ	+ve on duplex	-ve on duplex	
+ve onHHD	43	2			+ve onHHD	20	4	
-ve on HHD	16	28			-ve on HHD	6	59	
Thigh Perforators	+ve on duplex	-ve on duplex						
+ve onHHD	18	8						

Table 38: Mercer 1998¹⁶⁷

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-ve on HHD

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Positive and negative predictive values not given in paper, but calculated from raw values.								
Site examined	sensitivity [TP/TP+FN]		Specificity [TN/TN+FP]		Positive predictive value [TP/TP+FP]		Negative predictive value [TN/TN+FN]	
SFJ	[43/43+16]		<i>[28/28+2]</i>		[43/43+2]		<i>[28/28+16]</i>	
	0.73 (0.60-0.84)		0.93 (0.78-0.99)		0.96 (0.85-0.99)		0.64 (0.50-0.76)	
SPJ	[20/20+6]		<i>[59/59+4]</i>		[20/20+4]		[59/59+6]	
	0.77 (0.56-0.91)		0.94 (0.85-0.98)		0.83 (0.64-0.93)		0.91 (0.81-0.96)	
Thigh	<i>[18/18+17]</i>		[46/46+8]		<i>[18/18+8]</i>		[46/46+17]	
perforator	0.51 (0.34-0.69)		0.85 (0.73-0.93)		0.69 (0.5-0.84)		0.73 (0.61-0.82)	

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Rautio T, Perala J, Eiik H, Haukipuro K, Iuvonen T. nfluence of preoperative duplex ultrasonograph y on the operative procedure for primary varicose vein surgery. Phlebology 2002; 16: 149- 153	Diagnostic accuracy study.	49 patients (62 legs).	Patients with primary, previously untreated and uncomplicated varicose veins, referred for surgery. Exclusion: previous history of DVT. Median age: 45.5 years (range 19- 66). Gender: 5 male and 44 female patients. Venous disability score was 0-1 in all cases. Superficial reflux was detected in 55/62 limbs. No deep vein/perforator reflux detected.	Hand held Doppler with an 8MHz probe (Hadeco minidoppler ES- 100X). Patients tested in a semi- supine position. The sapheno- femoral junction, and the great saphenous vein at three separate points, were insonated. An audible flow signal lasting for > 1 sec was significant. The Sapheno-popliteal junction and short saphenous vein were also insonated if there were clinical evidence suggesting involvement. Done by an experienced General Surgeon.	Duplex scanning with a 5MHz probe (Toshiba Power Vision 8000, Japan). Patients supine with slight truncal elevation. Reverse flow of >1 second regarded as pathological. Done by a consultant vascular radiologist blinded to the hand held Doppler results.	Blinding carried out Tests followed each other at same appointment Expertise of operators comparable. No previous treatment and venous disability score of 0-1	Sensitivity, specificity. Positive predictive value and negative predictive value of hand held Doppler.	None stated

Table 39: Rautio 2002B²²⁶

Results: Raw data:

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
SFJ	+ve on duplex	-ve on duplex			GSV1 (mid thigh) [these findings were used for the report]	+ve on duplex	-ve on duplex	
+ve onHHD	31	1			+ve onHHD	24	1	
-ve on HHD	17	13			-ve on HHD	25	12	
GSV2 (popliteal fossa)	+ve on duplex	-ve on duplex			GSV3 (calf)	+ve on duplex	-ve on duplex	
+ve onHHD	22	3			+ve onHHD	15	3	
-ve on HHD	19	18			-ve on HHD	17	27	
95% CIs are also i	ncluded in rou	nd brackets. <i>Ro</i>	w data in italics and s	quare brackets.				
Site examined	sensitivity [T	P/TP+FN]	Specificity [TN/TN+	-FP]	+ve predictive value	e [TP/TP+FP]	-ve predictive va	lue [TN/TN+FN
SFJ	[31/31+17] 0.65 (0.49-0.7	78)	[13/13+1] 0.93 (0.66-1.00)		[31/31+1] 0.97 (0.84-0.99)		[13/13+17] 0.45 (0.29-0.62)	
GSV1 (mid thigh)	[24/24+25] 0.49 (0.34-0.6	54)	[12/12+1] 0.92 (0.64-1)		[24/24+1] 0.96 (0.81-0.99)		[12/12+25] 0.32 (0.20-0.49)	
GSV2 (popliteal fossa)	[22/22+19] 0.54 (0.39-0.6	58)	[18/18+3] 0.86 (0.65-0.95)		[22/22+3] 0.88 (0.70-0.96)		[18/18+19] 0.47 (0.33-0.64)	
GSV3 (calf)	[15/15+17] 0.47 (0.30-0.6	54)	[27/27+3] 0.90 (0.74-0.97)		[15/15+3] 0.83 (0.61-0.94)		[27/27+17] 0.61 (0.47-0.74)	

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Rautio T, Perala J, Biancari F, Wiik H, Ohtonen P, Haukipuro K, Juvonen T. Accuracy of hand held doppler in planning the operation for primary varicose veins. Eur J Vasc Endovasc Surg 2002; 24: 450- 455	Diagnostic review study. Handheld Doppler and Duplex done on the same day by different people.	111 patients (142 limbs)	Patients referred for surgical treatment of varicose veins with primary, uncomplicated and previously untreated varicose veins Exclusion: History of lower limb venous thrombosis Gender: 96 females Mean age (range): 42(23- 76) mean BMI (range): 25.6(18.3-52.8); Venous disability score 0: 14/111, 1: 85/111, 2: 12/111; CEAP stage: C1 (5/142), C2 (67/142), C3 (59/142), C4 (11/142).	Hand held doppler using a 8MHz probe (Hadeco mini- doppler ES- 100X). Patients were examined in a semi-supine position with the upper body elevated at 45 degrees. Audible flow signal of >1 sec was taken as the threshold of significant reflux. Carried out by consultant general surgeon.	Duplex scanning with a 7.5MHz probe (Toshiba Power Vision 8000). Positioning as for the hand held Doppler examination. Reflux >1 second was regarded as significant. Carried out by consultant vascular radiologist.	Blinding carried out Test interval within the same day Expertise of operators comparable Mostly C2-3, and had no previous treatments.	Sensitivity and specificity, positive predictive value, negative predictive value and kappa co- efficient.	Not reported.
Results:								
SFJ	+ve on duplex	-ve on dupl	ex	GSV1 (upper thigh)	+ve on duplex	-ve on duplex		
+ve onHHD	59	1		+ve onHHD	54	8		

Table 40: Rautio 2002A²²⁵

Reference	Study type	No. of patients	Patient chara	cteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
GSV2 (lower thigh)	+ve on duplex	-ve on dupl	ex		GSV3 (calf)	+ve on duplex	-ve on duplex		
+ve onHHD	53	10			+ve onHHD	46	14		
-ve on HHD	33	59			-ve on HHD	23	59		
SPJ	+ve on duplex	-ve on dupl	ex						
+ve onHHD	3	4							
-ve on HHD	10	95							
95% CIs are also	included in round	brackets. Rav	/ data in italics (and square brac	ckets.				
Site examined	sensitivity [TP/TP+FN]	Specificity [TN/TN+FP]	+ve predictive	e value [TP/TP+FP]	-ve predictive valu	ie [TN/TN+FN]	kappa co- efficient	
SFJ n=142	[59/59+46] 0.56(0.46-0.66)	[36/36+1]0	.97(0.86-100)	[59/59+1] 0.9	8(0.91-1)	<i>[36/36+46]</i> 0.44(0 .55)	.34-0	38(24-53)	
GSV1 (upper thigh) n=142	[54/54+39] 0.58(0.47-0.68)	[41/41+8]0	.84(0.70-0.93)	[54/54+8]0.87	7(0.77-0.93)	[41/41+39] 0.51(0	41-0.62)	36(21-51)	
GSV2 (lower thigh) n=142	[53/53+33] 0.62(0.51-0.71)	[<i>59/59+10</i>] 0.90)	0.82(0.70-	[53/53+10]0.8	34(0.73-0.91)	[59/59+33] 0.58(0	47-0.69)	41(26-56)	
GSV3 (calf) n=142	[46/46+23] 0.67(0.55-0.77)	[59/59+14] 0.88)	0.81(0.70-	[46/46+14]0.7	7(0.65-0.86)	[59/59+23] 0.72(0	61-0.81)	48(33-62)	
SPJ n=112	<i>[3/3+10]</i> 0.23(0.05-0.54)	[95/95+4] 0	.96(0.90-0.99)	[3/3+4]0.43(0	.16-0.75)	[95/95+10] 0.91(0	83-0.95)	24(-14–61)	

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Salaman RA, Fligelstone LJ, Wright N, Pugh KG, Harding KG, Lane IF. Hand held bi- directional doppler versus colour duplex scanning in the pre-operative assessment of varicose veins. J Vasc Invest 1995; 1:183-6	Diagnostic accuracy study	42(72)	Patients awaiting varicose vein surgery or attending the vascular outpatient clinic with symptomatic varicose veins.	Hand held Doppler with a Dopplex MD2 bi-directional hand-held Doppler unit with an 8MHz probe. Reflux duration threshold not stated. Done by an experienced vascular research fellow.	Duplex done with a Toshiba SPA270A scanner with a 5MHz linear array probe. Done by a vascular medical scientist. Reflux defined as >0.5 secs of retrograde flow.	Blinding unclear – reported that "both investigations were reported independently" Test interval not reported. Expertise of operators probable. Surgical history unclear and disease severity unclear.	Sensitivity and specificity, Positive predictive value, Negative predictive value	Not stated

Table 41: Salaman 1995²³⁶

Results: NB: these data are extracted from data provided, in a different form, in the paper. Note how the total n in each grid varies, from 72 (the expected value) to 77. This must be due to errors in the data on the paper.

SFJ	+ve on duplex	-ve on duplex	SPJ	+ve on duplex	-ve on duplex	
+ve onHHD	49	1	+ve onHHD	10	6	
-ve on HHD	4	18	-ve on HHD	8	50	
Thigh perforator	+ve on duplex	-ve on duplex	calf/ankle perforator	+ve on duplex	-ve on duplex	
-	+ve on duplex	-ve on duplex		+ve on duplex	-ve on duplex	

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
common femoral	+ve on duplex	-ve on duplex		popliteal	+ve on duplex	-ve on duplex		
+ve onHHD	0	13		+ve onHHD	2	1		
-ve on HHD	1	58		-ve on HHD	3	68		
Site examined	sensitivity [TP/TF	P+FN]	specificity [TN/TN+FP]		Positive predictiv [TP/TP+FP]	ve value	Negative predicti [TN/TN+FN]	ive value
SFJ	[49/49+4] 0.92(0.	.82-0.98)	[18/18+1] 0.95(0.	.74-1.00)	[49/49+1] 0.98(0	.90-0.99)	[18/18+4] 0.82(0.	.62-0.93)
SPJ	[10/10+8] 0.56(0.	.31-0.78)	[50/50+6]0.89(0.7	78-0.96)	[10/10+6] 0.63(0	.39-0.82)	[50/50+8] 0.86(0.	75-0.93)
Thigh perforators	[2/2+5] 0.29(0.04	I-0.71)	[54/54+13] 0.81(0	0.69-0.89)	[2/2+13] 0.13(0.0	04-0.38)	[54/54+5] 0.92(0.	.82-0.96)
Calf/ankle perforators	[2/2+4] 0.33(0.10	/2+4] 0.33(0.10-0.70)		[67/67+4] 0.94(0.86-0.98))-0.70)	[67/67+4] 0.94(0.	.86-0.98)
Popliteal	[2/2+3] 0.4(0.05-	0.85)	[68/68+1] 0.99(0.	.92-1)	[2/2+1] 0.67(0.21	-0.94)	[68/68+3]0.96(0.8	88-0.99)

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Schultheiss R, Billeter M, Bollinger A, Franzeck UK. Comparison between clinical examination, cw-Doppler Ultrasound and Colour-duplex sonography in the diagnosis of incompetent perforating veins. Eur J vasc Endovasc surg 1997; 13: 122- 126.	Diagnostic accuracy study.	19 patients (19 limbs)	Patients with chronic venous insufficiency. 2 described as C3, 14 as C4 and 3 as C5. Exclusion: C6 disease, PAD, cardiac problems, diabetes mellitus, nephropathy. Age: Mean age of the women was 62.8 years (range 44-79 years) and of the men was 56.3 years (range 32-76).	Hand held cw doppler ultrasound carried out by experienced medical doctor. 8.2 or 5.3 MHz pencil probe (Parks Electronics Lab model 10110). Testing carried out in standing over areas of marked fascial defect. No definition of reflux given in terms of duration.	Duplex carried out by another medical doctor blinded to HHD results. Linear 5 and 7 MHz probes were used (Acuson 128 XP/10). Done in standing. Reflux defined as reverse flow of >0.5 sec.	Blinding carried out Test interval not stated Expertise comparability not clear. Mostly C4. Previous treatment status not given.	Sensitivity, specificity of hand held Doppler.	Swiss Phlebology Society.

Site examined	sensitivity	specificity			
perforating veins	0.29	0.15			

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
van der Heijden FHWM, Bruyninckx CMA. Preoperative colour-coded duplex scanning in varicose veins of the lower extremity. Eur J Surg 1993; 159: 329-333	Diagnostic study	48 (68 legs)	Patients with leg varicose veins. Gender: 35 women; Age: mean age 48 years (range 16-77); Previous treatment: 10 had had previous stripping.	Continuous wave doppler done by a vascular technician. No other details given of positioning or definition of reflux in terms of duration.	Duplex carried out by a surgical resident. Toshiba SSA- 270A machine used, with 5MHz linear array transducer. Patients examined upright. Reflux of 0.5 seconds regarded as significant.	Blinding carried out Test interval <u>not</u> stated, but appears to be same day Expertise comparability probable. 21% with prior stripping. CEAP status not reported	No diagnostic outcomes presented by the paper, but some raw data allowed calculations.	None stated

Table 43: van der Heijden 1993²⁷⁶

Results: These were based on interpretation of the data in the paper which was presented (the numbers with duplex signs of incompetence were given, and also specific information given where there was discordance between HHD and duplex). In some cases a false negative result was not due to failure to observe reflux, but an incorrect identification of the source of reflux. Accuracy of these data is suspect.

SFJ	+ve on duplex	-ve on duplex	Great saphenous ve	+ve on duplex in	-ve on duplex	
+ve onHHD	45	1	+ve onHHD	41	1	
-ve on HHD	2	20	-ve on HHD	4	22	
short saphenous vein	+ve on duplex	-ve on duplex	Perforating veins	+ve on duplex	-ve on duplex	
+ve on HHD	16	0	+ve on HHD	10	1	
-ve on HHD	2	50	-ve on HHD	9	17	

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
SPJ	+ve on duplex	-ve on duplex						
+ve on HHD	17	0						
-ve on HHD	0	51						
Site examined	sensitivity [TP/TP	P+FN]	Specificity [TN/TI	N+FP]	Positive predictiv [TP/TP+FP]	ve value	Negative predict	ive value
SFJ	[45/45+2] 0.96(0.	85-0.99)	[20/20+1] 0.95(0.	.76-1)	[45/45+1] 0.98(0	.89-0.99)	[20/20+2] 0.91(0.	72-0.98)
Great saphenous vein	[41/41+4] 0.91(0.79-0.98)		[22/22+1] 0.96(0.78-1)		[41/41+1] 0.98(0.88-0.99)		[22/22+4] 0.84(0.67-0.94)	
Short saphenous vein	[16/16+2] 0.89(0.65-0.99)		[50/50+0] 1(0.93-1)		[<i>16/16+0</i>] 1(0.77-1)		[<i>50/50+2]</i> 0.95(0.86-0.99)	
Perforating	[10/10+9]		[17/17+1]		[10/10+1]		[17/17+9]	
veins	0.53(0.29-0.76)	.53(0.29-0.76)			0.91(0.62-0.98)		0.65(0.46-0.81)	
SPJ	[17/17+0] 1(0.8-1	.99)	[51/51+0] 1(0.93-	-1)	[17/17+0] 1(0.78-	-1)	[51/51+0] 1(0.91-	-1)

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
Wills V, Moylan D, Chambers J. The use of routine duplex scanning in the assessment of varicose veins. Aust NZ J Surg 1998; 68: 41- 44	Diagnosti c accuracy study	188 patients (315 legs)	Patients with varicose veins who had been referred to a vascular surgeon. Gender: 142 female Mean age 54.1 yrs (range 21-79 years) Previous treatment:122/315 legs Of these, 86 legs had had high ligation <u>+</u> other treatment, 8 had stab avulsions and 29 had sclerotherapy. 16/315 legs were thought to have secondary varicose veins (15 previous DVT and 1 arteriovenous malformation); skin changes present in 99/315 legs C4: 69 C6: 30 No disease, or only	Hand held doppler combined with clinical assessment. Parks hand-held doppler probe (8MHz) used. This was combined with clinical assessment, involving trendelenburg testing with a tourniquet. Patient position not described. Reflux definition not described in terms of duration. Done by a specialist vascular surgeon.	Duplex, using a Toshiba 270 scanner with a 5 MHz probe and colour flow imaging. Leg being examined was in a dependent position. Reflux defined as retrograde flow of >1 sec after the release of manual calf compression. Done by a trained vascular technician.	 Blinding NOT stated. Time interval not stated. Delay likely as stated that patient were 'referred' for duplex. Expertise of operators probably comparable. 39% had had previous treatment and 31% had skin changes 	Sensitivity and specificity	None reported

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Other issues of importance	Outcome measures	Source of funding
			superficial tributaries, with a normal great					
			saphenous vein, was seen in 35 legs.					
Results: <u>Insuff</u> Site examined		i ta given. The	e data below is all that w					
Sapheno-femo			Sensitivity 71.2%	Specificity 70.9%				
•	•							
Sapheno-popli			36.1%	92.1%				
Perforating vei	Perforating veins		43.6%	78.7%				
Deep Venous			29.2%	94.8%				
'uncomplicated	SFJ of a sub-group of legs with uncomplicated' varicose veins (no skin changes and not recurrent)		80.2%	52.2%				

G.3.2 Duplex assessment prior to interventional treatment

Table 45: Blomgren 2006A²⁵

Blomgren L, Johansson G, Bergdyis D. Quality of life after surgery for varicose veins and the impact of pre-operative duplex, reflux based on a nadomised surgery 2006; 20: 30-34.RCT. Sealed envelope system used unclear). 237Valuation: Primary varicose veins. randomised though to mention of how the randomised surgery 2006; 20: 30-34.No mention of how the randomised surgery 2006; 20: surgers 2005; 20: 30-34.No mention of how the randomised surgery 2006; 20: surgers 2005; 20: surgers 20: surgers	Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Results:	Johansson G, Bergqvist D. Quality of life after surgery for varicose veins and the impact of pre-operative duplex; reflux based on a randomised trial. Annals of vascular surgery 2006; 20: 30-34. (NB – Same study as Blomgren 2011 ²⁶ and Blomgren 2005 ²⁴)	envelope system used for allocation concealmen t. No mention of how the randomised sequence was drawn up. Study involved 20	randomised (though unclear). 237 reported to have given full follow-up data. 250 patients attended 2 year follow- up. Unclear how many legs were involved in the study. No reports of any who did not complete	Exclusion: pure cosmetic complaints, previous venous surgery or sclerotherapy, history of suspected or manifest deep venous thrombosis, active or healed leg ulcer, peripheral arterial disease, previous significant trauma to the leg, general illness and drug or alcohol abuse. Baseline characteristics: Poorly described (but available in Blomgren 2005). Overall mean (range) age was 47 (22-73) and 71% were women. 45 with bilateral surgery, 16 in duplex group and 29 in no duplex group (p=0.030. Skin changes present in 18%, with no differences between the groups. An important confounder was the surgery used, as this differed between groups. The duplex group had more patients than the non-duplex group with removal of the GSV and SSV, and less patients in the duplex group than the non-	Duplex scan. Surgical procedures that followed were removal of GSV/SSV, extrafascial ligation of perforators, and stab avulsions of	operative duplex scan. Surgical procedures that followed were removal of GSV/SSV, extrafascial ligation of perforators, and stab avulsions of	2 years	life Patient assessed symptoms Rates of recurrence and reoperatio n (dealt with in detail in Blomgren	None
Duplex No duplex	Results:								

Reference	Study type	No. of patients	Patient characteris	tics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
patient assessed sy operated limbs u	•	rse at 2 years com	pared to baseline	15/130		19/120				
Rates of recurrence	e and reoperation	n		significantly higher in no duplex group (more details given in Blomgren 2005)						
Quality of life – SF-	Quality of life – SF-36 domains				erence between t ame when patien parately. No data	ts with bilateral	and unilatera			

Reference	Study type	No. of patients	Patient char	acteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Blomgren L, Johansson G, Bergqvist D. Randomised clinical trial of routine preoperative duplex imaging before varicose vein surgery. British Journal of Surgery 2005; 92: 688-694 (NB – Same study as Blomgren 2011 and Blomgren 2006)	RCT. Sealed envelope system used for allocation concealment. If both legs included, both given the same randomisatio n (i.e. randomised by patient). No mention of how the randomised sequence was drawn up. Study involved 20 surgeons.	308 randomised but 15 initially excluded because of refusal, pregnancy and remote residency. In the duplex group 8 (8 legs) were excluded (2 patient request, 2 inclusion criteria violation, 2 moved to remote region, 2 pregnancy). In the no duplex group; 7 (7 legs) were excluded (4 patient request, 2 inclusion criteria violation, 1 moved to remote region). This left, by the point of the duplex intervention, 148 patients (166 legs) in the duplex group and 145 patients (177 legs) in the no duplex	an indication of the surger Exclusion: p previous ver sclerotherap manifest der active or hea arterial dise	n for surger on). ure cosmeti nous surger y, history o ep venous t aled leg ulce ase, previou ne leg, gene hol abuse.	f suspected or hrombosis, er, peripheral is significant ral illness and	Pre-operative Duplex scan, using a colour flow duplex machine (Acuson XP128 and Acuson Sequioa 512). Reflux with a duration of >0.5 seconds was regarded as significant. (Some surgeons also did a pre- operative hand held Doppler scan). Surgical procedures that followed were removal of GSV/SSV, extrafascial	No pre- operative duplex scan. (But some surgeons did a pre- operative hand held Doppler scan). Surgical procedures that followed were removal of GSV/SSV, extrafascial ligation of perforators, and stab avulsions of tributaries. Most done under general	2 years	Reflux at 2 months Reflux at 2 years	None

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Reference	Study type	No. of patients	Patient characteristi	s Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
		group. Loss to follow-up; 2 months: Duplex – 5 people (6 legs); No duplex – 8 people (11 legs). 2 years: Duplex – 35 people (39 legs); No duplex – 39 people (48 legs).		ligation of perforators, and stab avulsions of tributaries. Most done under general anaesthetic. Importantly, in the duplex group the duplex assessment led to the alteration of surgery from the pre- determined course in 44/166 legs.	anaesthetic. Naturally, in the absence of duplex assessment, the predetermi ned course based on clinical examinatio n was adhered to.			
Results: Analy	sis was done by le	gs. For reflux, intentior	to treat results given					
				Duplex			No dupl	
SFJ reflux at 2	months			10/160			37/166	
SPJ reflux at 2	months			4/160			9/166	
SFJ and/or SPJ	reflux (i.e. reflux a	anywhere!) at 2 month	S	14/160			44/166	5
SFJ reflux at 2	Vears			14/127			44/129)

Reference	Study type	No. of patients	Patient characteristi	cs	Intervention	Comparison	Length of follow-up		Source of funding		
SPJ reflux at 2 ye	ears				7/127			13/129)		
SFJ and/or SPJ r	eflux (i.e. reflux a	nywhere!) at 2 years			19/127			53/129			
	r operation in first coms, or patients'	t 2 years (indication w wish)	as persistent or	3/145 (includ	ing patient with another hospit		at	14/147			
Adverse events											
DVT at 2 years					0/145			0/147			
Proportion with	improvement in	CEAP category at 2 years	ars		104/145			86/147	7		
	• •	ges (C4+) at 2 years (baseline difference, but ne, so does not affect validity of result on rigl		15/145				19/147	7		
	ith oedema (C3) at 2 years (baseline difference, with mor Iseline, which threatens the validity of the result on right)			25/145				38/147	7		

Reference	Study type	No. of patients	Patient char	acteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Blomgren L, Johansson G, Emanuelsson L, Dahlberg- Akerman, Thermaenius P, Bergqvist D. Late follow-up of a randomised trial of routine duplex imaging before varicose vein surgery. British Journal of surgery 2011; 98: 1112- 1116.(NB – Same study as Blomgren 2005 and Blomgren 2006)	RCT. Sealed envelope system used for allocation concealment. If both legs included, both given the same randomisatio n (i.e. randomised by patient). No mention of how the randomised sequence was drawn up. Study involved 20 surgeons.	308 randomised but 15 initially excluded because of refusal, pregnancy and remote residency. In the duplex group 8 (8 legs) were excluded (2 patient request, 2 inclusion criteria violation, 2 moved to remote region, 2 pregnancy). In the no duplex group 7 (7 legs) were excluded (4 patient request, 2 inclusion criteria violation, 1 moved to remote region). This left, by the point of the duplex intervention, 148 patients (166 legs) in the duplex group and 145 patients (177 legs) in the no duplex group. Loss to follow-up; 2 months: Duplex – 5 people (6 legs); No	Inclusion: Previous of the view of the vie	cation for su the surgeon ure cosmeti previous ve clerotherapy r manifest d active or he neral arteria nificant trau illness and c se.	rgery (in). c nous n, history of eep venous aled leg l disease, ma to the lrug or	Pre- operative Duplex scan, using a colour flow duplex machine (Acuson XP128 and Acuson Sequioa 512). Reflux with a duration of >0.5 seconds was regarded as significant. (Some surgeons also did a pre- operative HHD scan).	No pre- operative duplex scan. (But some surgeons did a pre- operative HHD scan). Surgical procedures that followed were removal of GSV/SSV, extrafascial ligation of perforators, and stab avulsions of tributaries. Most done under general anaesthetic. Naturally, in the absence of duplex	7 years	Reflux at 7 years	None

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Reference	Study type	No. of patients	Patient characteris	tics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
		duplex – 8 people (11 legs). 2 years: Duplex – 35 people (39 legs); No duplex – 39 people (48 legs). 7 years: Clinical examination: Duplex: 62 people (70 legs); No duplex: 56(88). Interview and info from patient notes: Duplex: 34 people (42 legs); No duplex: 32(43).			were removal of GSV/SSV, extrafascial ligation of perforators, and stab avulsions of tributaries. Most done under general anaesthetic. Importantly, in the duplex group the duplex assessment led to the alteration of surgery from the pre- determined course in 44/166 legs.	assessment, the predetermin ed course based on clinical examination was adhered to.			
Results:				Duplex		No Duplex			
SFJ reflux at 7 y	vears			11/95		38/99			
SPJ reflux at 7 y	/ears			2/95		9/99			

Reference	Study type	No. of patients	Patient characteri	stics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding		
SFJ and/or SPJ	reflux (i.e. reflux a	anywhere!) at 7 years		13/95		46/99					
Condition of tro at 7 years	eated leg compar	ed to before surgery (Unc	hanged or worse)	16/123		28/126					
Quality of life -	- SF-36 at 7 years			No data given, a variable betwee			were no differ	ences in any S	SF-36		
Reoperation or	r scheduled for re	operation at 7 years		15/124		38/134	ias).				
Complications	of varicose veins a	at 7 years					4				
Venous ulcer				0/70		0/88					
Hyper-pigme	ntation or eczema	а		3/70		9/88					

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Smith JJ, Brown L, Greenhalgh RM, Davies AH. Randomised trial of pre-operative colour duplex marking in primary varicose vein surgery: outcome is not improved. Eur j Vasc Endovasc Surg 2002; 23: 336-343.	RCT. Randomisatio n done with sealed envelope system with allocation by a third party. Randomised by patient not leg.	149 patients randomised . None lost to follow-up and none discontinue d interventio n.	Inclusion: patients with primary varicose veins without venous ulceration. Baseline characteristics: No details on demographic characteristics. However, clear information on the surgery each group received. The groups were very similar for the number of "HSL/strip/phlebectomy" procedures, and "phlebectomy alone" procedures, but the duplex group had more SPJ and phlebectomies, and more "short and long saphenous system together" procedures. However the latter two classes only comprised a very small proportion of all procedures and so broadly the surgical procedures were comparable. Quality of life was described as comparable for all quality of life measures.	Pre-surgical duplex assessment. Duplex carried out by an experienced vascular technologist. Accuson 2000 scanner with 7.5 MHz linear array probe was used. Hand held HD assessment also carried out, using an Imax continuous wave doppler with 8MHz probe. Common procedures: operative procedures were done with general anaesthetic. For long saphenous system, flush sapheno- femoral transfixion with division of surrounding branches and removal of the great saphenous vein was carried out. For small saphenous vein popliteal transfixion and removal of the small saphenous vein was carried out.	No pre- surgical duplex assessment. Unclear, but it is likely these patients <i>did</i> have hand held Doppler assessment. N=97	12 months	Quality of life Reflux	None stated

Table 40. Smith 2002251

Reference	Study type	No. of patients	Patient cl	naracteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding		
					Tributary varicosities removed with phlebectomy hooks or mosquito clips. N=92						
Results:											
				Duplex		No Dup	No Duplex				
Reflux SFJ 6weeks				1/92		1/97	1/97				
Reflux GSV 12 mor	V 12 months			8/92		9/97*					
Reflux SSV 6 week				4/92		6/97					
Reflux SSV 12 mon	iths			6/92			8/97				
Reflux perforators	6 weeks			1/92			5/97				
Reflux perforators	12 months			4/92		15/97					
Development of ne	ew branch varic	osities at 12 mo	nths	8/92		9/97					
Aberdeen Questio GIVEN)	nnaire (AVVQ) a	t 6 weeks (NO V	ARIANCE	10.85		15.85 (15.85 (P=0.034)				
Aberdeen Questio	nnaire (AVVQ) a	t 12 months		No difference repor	ted [p=0.187] (data in low	resolution figure	, no data in tex	t)			
SF 36 6 weeks	5 6 weeks			No diff reported p>0	0.38 (all domains)						
SF 36 12 months					0.15 (all domains)						

* paper reports a total of 17 having GSV at 12 months, and then "of which 8 were in the duplex group and 19 in the no duplex group". This was assumed to be a typographical error, and that it should have been 9 rather than 19.

G.4 Chapter 8 – conservative management

1 Compression vs. no treatment

Table 49: Anderson 1990⁷

Reference	Study type	No. of patients	Patient characteristics	Interventio n	Compariso n	Length of follow-up	Outcome measures	Source of funding
Anderson JH, Geraghty JG, Wilson YT, Murray GD, McArdle CS, Anderson JR. Paroven and graduated compression hosiery for superficial venous insufficiency. 1990; 5: 271- 276.	Randomised cross-over trial. Four groups were involved – Paroven alone; hosiery and placebo; paroven and hosiery; and placebo alone. The subjects were randomised to start in one of these 4 groups, and treatment sequences were balanced within groups of 12 in three Latin squares. Only the results from placebo alone and hosiery and placebo are included in this evidence table.	72. 6 did not complete the trial. Not possible to determin e the numbers for each treatment group. No ITT reported.	 Mean age 40 years (range 20-61 years). 39 patients spent at least 2/3 of their time at work standing. Inclusion: patients on waiting list for varicose veins surgery (mean of 6 months on list) who indicated the presence of at least 2 of the following symptoms: leg pain, heaviness, itch, cramps, swelling. Exclusion: If the only complaint was cosmetic distress. Age >65 years; clinical evidence of peripheral arterial disease (PAD), concurrent treatment with diuretics, Ca²⁺ antagonists, NSAIDs, vasodilators, or corticosteroids; history of DVT. Baseline characteristics: Not given for the 4 randomised cross-over groups. 	Full length hosiery fitted to give a pressure at the ankle of 30-40 mmHg. Hosiery removed in bed. Used for 4 weeks	Placebo is not described. It is likely it was a sham pill, but unclear.	Length of each treatment – 4 weeks.	Patient assessed symptoms: (using visual analogue scale (VAS).	not stated

Reference	Study type	No. of patients	Patient characteristics	Interventio n	Compariso n	Length of follow-up	Outcome measures	Source o funding
	periods described							
	 but patients 							
	were supposed to							
	attend for post-							
	test outcome							
	assessment after							
	50 days, so this							
	implies a wash-							
	out of 22 days.							
	Details of							
	randomisation							
	and allocation							
	concealment not							
	given. No							
	evidence of							
	blinding.							

Results: VAS scores at the end of the 4 week treatment period are given. The risk of bias from order effects (carry-over) minimised by Latin squares method of ensuring balanced ordering of treatments.

VAS (higher the worse the severity)	Compression [mean(SE)]	Placebo [mean(SE)]	p value (post-hoc)
Pain	34.7 (3.6)	37.6 (3.6)	0.06
heaviness	34.1(3.8)	36.3 (3.5)	0.39
itch	32.0 (3.8)	30.5 (3.9)	0.56
swelling	28.2 (3.6)	35.3 (3.7)	0.13
night cramps	22.4(3.1)	24.9 (3.0)	0.24
body image concerns	43.2(4.6)	41.1 (4.7)	0.43
Author's conclusions: No conclusions made	for bosiony along		

Author's conclusions: No conclusions made for hosiery alone.

Reference	Study type	No. of patients	Patient characteristics	Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Benigni JP, Sadoun S, Allaert FA, Vin F. Efficacy of Class 1 elastic compression stockings in the early stages of chronic venous disease. International Angiology 2003; 22: 383- 392	RCT. Multi-centre cross- over trial, with 7 day washout period. Randomised, but method not mentioned. No allocation concealment mentioned. Double blinded, but few details given. Withdrawal: 8 by day 14, further 3 by day 21 and further 3 by day 21 and further 3 by day 35 (total=14). Ignoring losses in the wash-out period the placebo group lost 7 over the two periods, but the intervention group lost only 4. NB: For all outcomes (except mood and daily work activity) the detailed results given in the paper are ONLY from the first phase, PRIOR to cross-over. Hence this is not truly a cross-over study. Full cross-over results were given for mood and daily work activity but no reasons	125. ITT analysis used. Used on those patients who had worn "study stockings at least once" and who had been evaluated at least once.	Inclusion: female patients aged 18-75 years, with early stage Chronic Venous Disease (CVD) of the legs. Thread veins, non-saphenous varicose veins (<3mm) or ankle oedema without skin changes. Symptoms including pain, heavy legs, cramps, paraesthesia or ankle swelling. Global painful leg discomfort lasting >8 days, and with a visual analogue scale (VAS) of 4/10 or more on the day of testing. Competent deep venous trunks, competent greater saphenous veins (<5mm), competent lesser saphenous veins (<4mm), competent calf perforating veins, shown by a venous refilling time of >24 seconds and an ankle diameter of 20-26cm and a maximum calf diameter of 33-43cm. Exclusion : male patients, suffering from chronic or severe disease. Symptoms of signs in the legs due to pathology of cardiac, renal, hepatic, metabolic, neurological, osteo-articular or traumatic origin. BMI>30. Any risk factors for worsening CVD: recent venous thrombosis, pregnancy or childbirth within 6 months. Past history of DVT. Skin changes, permanent ankle oedema. Ultrasound evidence of valvular incompetence in the sapheno-femoral or sapheno-	Class 1 knee-high graduated compressi on stockings (13-20 hPa). Given for 14 days, and worn for a minimum of 6 hours per day. Cross over study, so half the participan ts were randomis ed to receive this treatment first, prior to the 7 day washout period.	Placebo: regular knee-high stocking used as a "referenc e" stocking, providing <10hPa. Very similar in appearan ce. Given for 14 days, and worn for a minimum of 6 hours per day. Cross over study, so half the participan ts were randomis ed to receive this	At end of double cross-over treatment period (35 days)	Patient assessed symptoms: global discomfort in legs, pain, heavy legs, cramps, swelling in ankles, mood, daily work activity. Adverse events.	not stated

Table 50: Benigni 2003¹⁹

Reference Stu	tudy type	No. of patients	Patient characteristics	Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
ful the op pu red the glo he sw en	re given for the lack of all cross-over results for he other variables. This pens up the risk of ublication bias and educed confidence in he validity of the resented findings. All hat is stated is that the lobal discomfort, pain, eavy legs, cramps and welling measured at the nd of the cross-over tage were "similar".		 popliteal junctions, obstacles or reflux in the deep venous network. Use of Calcium channel blockers, anti-coagulants, diuretics, anti-inflammatory drugs, Vitamin C, recent hormonal treatment, recent phlebotonic or pain medications, recent elastic compression and indications for sclerotherapy or surgery. Baseline characteristics: No significant differences between the two randomised groups (control first versus intervention first) for age, weight, height, professional status, risk factors or past medical history. No baseline differences in outcome variables. 		treatment first, prior to the 7 day washout period.			

Results: NOTE: Full cross-over results are given for mood and daily work activity only. For all others, results pertain to those recorded at the end of the first phase (before cross-over). All that is stated is that the group differences in global discomfort, pain, heavy legs, cramps and swelling measured at the end of the cross-over stage between the two treatment groups were "similar" to those at day 14.

	Compression [mean VAS (sd)]	Placebo [mean VAS (sd)]	p value
Global painful discomfort in the legs during days 7-14	1.8 (1.7)	3.1 (2.1)	<0.05
Global painful discomfort in the legs at day14	1.4 (1.8)	2.9 (2.1)	<0.01
Mood (unclear if a high score represents good or bad mood)	1.1 (1.7)	1.5 (1.9)	
Daily work activity (unclear if a high score represents high or low activity)	1.1 (1.6)	1.6 (1.8)	
	Compression [count with no change or a deterioration]	Placebo [count with no change or a deterioration]	

Reference	Study type	No. of patients	Patient character	ristics	Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Pain at day 14		27/61		37/53		0.0215			
Heavy legs at da	ay 14	20/59		35/54		0.0025			
Cramps at day 1	.4	37/61		44/55		0.0379			
Ankle swelling d	lay 14	35/61		43/53		0.0240			

No outcome reporting for paresthesia, leg volume, changes in venous refilling time or venous pump power, except that no difference between compression and placebo.

Adverse events were reported as being significantly worse for those using the placebo stockings – slipping sensation on the leg, warming sensation, a feeling of pressure on the legs.

Author's conclusions: The wearing of class 1 graduated compression knee-high stockings (10-15 mmHg at the ankle) for a 15 day treatment period results in a significant improvement in the symptomatology and in the quality of life criteria in patients presenting with early-stage CVD of the lower extremities.

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Junger M, Galler S, Klscz T, Steins A, Hahn M. Improvement of cutaneous microangiopa thy by compression therapy in chronic venous insufficiency. Phlebology Suppl. 1996; 1: S10-S13.	Cohort study.	20	Inclusion: Chronic Venous Insufficiency (CVI) class I (n=7) and II (n=13), according to Widmer's classification; Baseline characteristics: Gender: 10 men; 10 women; mean age 54.9 years(9.5); 3 with previous DVT; all with ankle brachial index>1.0; all had reduced venous refill time (20(12) seconds); Doppler showed incompetence of: • deep veins in 7 patients • superficial veins in 12 patients • perforating veins in 6 patients.	2 weeks of short- stretch bandaging, followed by 2 more weeks with class II compression stockings (the 3 with previous DVT used class III).	Before and after design.	2 weeks and 4 weeks.	Patient assessed symptoms: pain, tautness, swelling, itching, and feelings of cold, heat and restriction (on a scale of 0-3 (max complains)	None stated

Results: Reported that subjective treatments in all patients decreased during treatment, except for the feeling of coldness, which increased again during the second part of the study using compression stockings. There were no complaints by patients about feelings of constriction during the second part of therapy. No numerical data presented.

Author's conclusions: No relevant conclusion with reference to patient symptoms.

Reference	Study type	No. of patients	Patient characteristics	Interventi on	Compariso n	Length of follow-up	Outcome measures	Source of funding
Krijnen RMA, de Boer EM, Ader HJ, Osinga DSC, Bruynzeel DP. Compression stockings and rubber floor mats: do they benefit workers with chronic venous insufficiency and a standing profession? JOEM 1997; 9: 889-894.	Quasi-randomised controlled trial. The truly random part was the allocation to treatment and control groups. No details of randomisation method used. No evidence given of allocation concealment. There was a further, non- random splitting of the treatment patients to the two treatments – compression stockings or the use of rubber mats to stand on. This was decided by the safety and hygiene conditions of the factory concerned. In 10 factories compression stockings were used, and in 4 rubber mats were used. It is conceivable that this could cause bias by the compression group being from a certain type of factory (specific hygiene and safety conditions) and the placebo group being from any type of factory (perhaps those working in a certain type of factory – i.e. heavy industry - would have different risk	 114 in total. 101 in the control (n=50) and compression (n=51) groups. In the compression group one refused to wear hosiery and a further 5 were lost to follow-up, for "unrelated reasons". A further 15 stopped wearing the stockings every day during the stockings every day during the study, for reasons including poor fit or skin problems. 16 were lost to follow-up from the control group. 	All male factory workers with a predominantly standing job from 14 factories. All with evidence of chronic venous insufficiency (CVI). 40 had complications including trunk varicosis, lipodermatosclerosis, hyperpigmentation, atrophy blanche or dermatitis. None had leg ulcers. Inclusion: Evidence of CVI by physical examination, Doppler ultrasound investigation and light reflective rheography. Standing factory job. Exclusion: Individuals with only intracutaneous or only a few small varicose veins. Baseline characteristics: No comparison of anthropometric baseline, the compression and control groups were similar for proportion having pain, with 10/30 in pain in the intervention group and 13/34 in the control group [estimated from graph] (no statistical analysis done), and the	Below knee class II (30-32 mmHg) seamless compressi on stockings. Used during working hours only.	Not described, but appears to be no treatment.	3 months	Patient assessed symptom s Adverse events	not stated

Reference	Study type	No. of patients	Patient characteristics	Interventi on	Compariso n	Length of follow-up	Outcome measures	Source o funding
	factors). However this would probably be a small effect as only 13 subjects out of 64 in the treatment group were put in the mat group. Only results for compression versus control are included in this evidence table.	No ITT performed so analysis restricted to the 30 who wore the stockings almost every day, and the 34 in the control group who attended follow-up.	control group [19/34, estimated from the graph] had a slightly lower proportion of people than the compression group [21/30] with a tired feeling at baseline. This slight difference will have favoured the control group, and thus does not invalidate the post- intervention finding that the compression group had a lower proportion of people with tired legs than the control group.					

Results: Post-test results given. Despite the lack of confirmation that the groups were similar at baseline for the variables below, the differences seen are unlikely to have led to a bias favouring compression. In particular, for the tired feeling, less control were in pain at baseline which would favour the control group.

	Compression	Control	p value
Patients with complaints of tired legs (proportion of subjects) at 3 months	8/30	18/34 (estimated from graph)	<0.005
Patients with complaints of pain (proportion of subjects) at 3 months	2/30	12/34 (estimated from graph)	<0.05
Patients with overall decrease in complaints at 3 months	17/30 For the 15 not wearing stockings every day: 4/15	4/50 (16 were reported lost to follow- up, and no ITT was done, so this may be a typographical error, and the correct result may be 4/34).	
Patients in favour of continuing stockings beyond study duration.	26/45		

Reference Study type		No. of patients	Patient characteristi	Interventi cs on	Compariso n	Length of follow-up	Outcome measures	Source o funding
Adverse events Reasons for non-compliance only the 15 who did not wear the stockings everyday were asked. Each person could give only one reason each) itch red and swollen skin too tight	2/15 2/15 5/15							

Author's conclusions: Compression stockings appeared to be superior [....] with regard to applicability, [and] diminishing subjective complaints....

Reference	Study type	No. of patients	Patient characteristics	Intervention	Compari son	Length of follow-up	Outcome measures	Source of funding
Lurie F, Kistner RL. Trends in patient reported outcomes of conservative and surgical treatment of primary chronic venous disease contradict current practices. Annals of Surgery 2011; 254: 363-367.	Observatio nal single group before- after study.	150 were originally selected. These were divided into two groups who both initially had compression therapy (one group later had endovenous radiofrequency ablation whilst the other continued with conservative treatment, and the results of these later treatments will not be included here). However, since the data were continuous it is not possible to combine the results for the initial compression phase, so data from the larger group of 121 patients is given.	Consecutive patients with primary CVD were selected according to the criteria below. Inclusion: confirmed primary aetiology; unilateral involvement; great saphenous vein reflux; C2-C4; no use of compression for at least one year. Exclusion: CEAP stages C5-6; small saphenous vein involvement; current or recent use of compression; non-compliance with compression therapy; difficulty completing quality of life form; problems with English language comprehension. Baseline characteristics: Gender: 38% male; Age: mean age 54.4(11.7); CEAP stage: C2: 32.2%; C3: 24%; C4: 43.8%;	Compression therapy by 20- 30mmHg knee-high graduated compression stockings given for 2 to 6 weeks. Lifestyle advice (weight loss, exercise and frequent leg elevation) ALSO given.	Pre versus post (2-6 weeks of treatmen t).	2-6 weeks.	Patient reported quality of life: Disease specific SQOR-V form Patient assessed symptoms: Symptom score	None stated

Table 53: Lurie 2011¹⁵⁴

Results: mean (sd) given. N=121

	pre-compression	post-compression	p value
Symptom score (this is made up of part of the SQOR-V form, comprising severity of pain, heaviness, itching, night cramps, heat or burning, tinglir		6.3(5.8)	not given

Reference	Study type	No. of patients	Patient characteristi	cs	Intervention	Compari son	Length of follow-up	Outcome measures	Source of funding
throbbing, restl scores of these indicates worse	9 symptoms, e								
	•	o of several domains, witl fic QoL; 190 is the maxin	•	62.5(20.6)		48.9(17.9)		not given	

Author's conclusions: compression therapy selectively improves some symptoms....the QOL outcomes of compression therapy were better than the symptom response.

Reference	Study type	No. of patients	Patient characteristics	Intervention	Compariso n	Length of follow-up	Outcome measures	Source of funding
Motykie GD, Caprini JA, Arcelus JI, Reyna JJ, Overom E, Mokhtee D. Evaluation of therapeutic compression stockings in the treatment of chronic venous insufficiency. Dermatol Surg 1999; 25: 116-120	Before-after design observational trial, without control group. Therefore subject to uncontrolled threats to internal validity, such as time effects, placebo effects etc.	112. Those with bilateral symptoms included, but unclear how many.	Inclusion: Patients with chronic venous insufficiency (CVI). Exclusion: patients currently wearing compression stockings. Baseline characteristics Gender: 95 females, 17 males; Age: range 27-85 years (mean 46.8); No prior CVI or varicose veins treatment: 95/112. Prior treatment 17/112 • 11/17 sclerotherapy • 6/17 stripping surgery. Flawed statistics provided for CEAP class. Authors gave percentages of participants with the main symptom characteristic of each CEAP class which is not helpful as some symptoms will span multiple CEAP grades (i.e. instead of "swelling" indicating those with swelling but NOT pigmentation or ulceration, which would be equivalent to CEAP3, the swelling statistic included any of those also with pigmentation and ulceration, which was therefore no longer equivalent to CEAP3).	30-40 mmHg compression stockings for 16 months. Hours per day and night-use unclear. The stockings varied • 36% thigh lengt, • 17% mid-thigh length • 47% knee or calf length	Post treatment compared to pre- treatment.	1 month and 16 months (treatment continued to end of follow-up).	Patient assessed symptoms Adverse events	None stated

data points there we Patient assessed sym with 1=minimal prob problem) swelling pain discolouration	vere. mptoms (1-5 sc	ale, pre		iven were given bilateral sto 1 month post-compress		vith bilateral sy	mptoms unkno [.]	wn. Hence uncle	ear how many		
with 1=minimal prob problem) swelling pain discolouration	•			1 month post-compress							
pain discolouration					ion 16 months pos	t-compression		Vilcoxon signed pite the present			
discolouration		2.4	2.45(1.25) 1.47(0.83)		1.13(0.51)	1.13(0.51) 1.38(0.69)		P<0.001 for comparison between baseline and 1 month for all variables.			
	pain		94(1.29)	1.77(1.09)	1.38(0.69)						
cosmetic problems		2.7	'6(1.29)	2.23(1.22)	1.81(0.99)	1.81(0.99)		P<0.0001 for comparison between baseline and 16 months for all variables			
	cosmetic problems		3(1.41)	2.50(1.41)	1.98(0.99)						
activity tolerance		2.3	3(1.35)	1.71(1.19)	1.38(0.73)	1.38(0.73)					
depression		1.7	2(1.12)	1.42(0.87)	1.29(0.81)						
sleep problems		2.0	00(1.25)	1.46(0.99)	1.24(0.63)						
Adverse events (scale patient assessed sym		for									
numbness		NA		1.41(1.20)	1.20(0.92)		NA	NA			
Sweating Itchiness new pain				1.45(1.00)	1.23(0.83)						
				1.40(0.97)	1.14(0.78)	1.14(0.78) 1.12(0.80)					
				1.44(1.20)	1.12(0.80)						
Compliance (still wea	earing stockings)		92/112	78/112						

Reference	Study type	No. of patients	Patient characteristics	Intervention	Compariso n	Length of follow-up	Outcome measures	Source of funding
Pannier F, Hofffmann B, Stang A, Jockel KH, Rabe E. Prevalence and acceptance of therapy with medical compression stockings. Phlebologie 2007; 36: 245-249.	Cross- sectional questionnaire/ interview study.	3072	Inclusion: Randomly recruited from a German city and environs; included ALL residents, regardless of health status; urban to city ratio of 2:1; 59% response rate; demographics representative of the general German population; Baseline characteristics: mean BMI: 25.6(4.8); 890 described as having varicose veins. 961 had history of chronic venous sufficiency (CVI) at CEAP stages C2-6: • C2: 439/961 • C3: 412/961 • C4: 88/961 • C5: 19/961 • C6: 3/961.	Patients with a history of varicose veins were asked about their use of medical compression stockings (MCS).	ΝΑ		Patient assessed symptoms Adverse events	None stated

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Results: 10.3%% (n=316) of all sample with CEAP stages C2-6 had used MCS (32.9% of those with C2-C6). At the time of interview, 210/316 (66.5%) who had ever received MCS were not wearing them at the time of interview, and had not worn them in the last 4 weeks, indicating a compliance figure of <u>33.5%</u>.

The groups who had used them in the last 4 weeks and not used them in the last 4 weeks differed in terms of the proportions from each CEAP class, with the most severely affected tending to be more compliant:

CEAP	Medical compression stockings used, but not in last 4 weeks	Medical compression stockings used in last 4 weeks
C2	96/210 (21.9%)	26/106 (5.9%)
C3	78/210 (18.9%)	50/106 (12.1%)

Reference	Study type	No. of patients	Patient characteristics	Intervention	Compariso n	Length of follow-up	Outcome measures	Source of funding
C4		2	8/210 (31.8%)		20/106 (22.7	'%)		
C5-C6		8	/210 (36.4%)		10/106 (45.5%)			

For those who used MCS currently, patients usually wore their MCS 5 or more days per week (73%) and for 8 or more hours per day (89.4%).

71.3% of the interviewed participants using MCS said their medical condition had improved with MCS therapy. This included:

- reduction in swelling (84.2%)
- reduction in heaviness (89.4%)
- reduction in leg pain after prolonged standing (60.9%)
- reduction in tension in the legs (78.9%)

Most patients could not remember the compression class, but available evidence suggested:

- class 1: 13
- class II: 149
- class III: 26

The types were:

- compression tights: 34%
- thigh compression stockings: 23%
- lower leg compression stockings: 41.1%

Adverse events were reported as:

- pruritis (8.4%)
- eczemas (1.6%)
- constrictions under the MCS (8.4%)
- slipping of stockings (3.6%)

Author's conclusions: An improvement of their condition was attributed to [MCS] by 80% of patients.

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Raju S, Hollis K, Neglen P, Mississippi F. Use of compression stockings in chronic venous disease: patient compliance and efficacy. Ann Vasc Surg 2007; 21: 790-795	Case series. Extremely limited methodology in terms of being open to multiple risks of bias.	3144	Inclusion: Stated that "new" chronic venous disease (CVD) cases, but then also stated that they had been under care with GP or other specialists for variable periods of time; Baseline Characteristics: • Age: median 58 (range 17- 92); • Gender: Male: Female =1:2 • CEAP stages • CEAP 0-2: 67% • CEAP 3: 22% • CEAP 4: 4% • CEAP 4: 4% • CEAP 5: 4% • CEAP 6: 3% • Aetiology was primary in 58% and post-thrombotic in 42%.	None. This was an observational study of CVD patients, and only 37% were using stockings.	NA		Compliance	None stated

200-217

Results:

21% of patients reported full "compliance". 12% used them most days and 4% some days. The other 63% did not use the stockings at all or had abandoned them after a trial period in the past. Compliance did not differ according to CEAP class, gender or previous DVT. Compliance did improve with longer duration of treatment Reviewer's comment: As not all the patients in the study had been prescribed stockings, these daily use figures of 21%, most days use figures of 12% and occasional use values of 4% do not really equal compliance, as compliance must make use of the number prescribed them as the denominator. Clearly a patient never prescribed a treatment cannot be described as non-compliant. As only 75% of the patients had been recommended stockings by a doctor, the true full compliance figure would be 21/0.75= 28%; full and partial compliance would be (21+12)/0.75=44% and full, partial and minimal compliance would be 21+12+4)/0.75=49.33%.

Primary reasons for non-use of stocking, of those that were recommended stockings by their doctor

unable to state a reason

40%

. 201

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
lack of efficacy		20%						
poor fit/cut off	fcirculation	17.3%						
too hot		9.3%						
soreness		2.67%						
needs applicat assistance	ion	2.67%						
cosmetic reaso	ons	2.67%						
itching/dermat	titis	2.67%						
worsening of s	ymptoms	1.33%						
lack of self-disc	cipline	0.67%						
Cost		0.53%						
Work-related		0.27%						
Author's concl	usions: Non-con	npliance is very h	nigh in patients with CVD regardles	s of age, sex, aetiology o	f CVD, duration	of symptoms of	or disease severity	

Table 57: Michaels 2006¹⁷⁰

Reference	Study type	No. of patients	Patient ch	naracteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Michaels JA, Brazier JE, Campbell WB, MacIntyre JB, Palfreyman SJ, Ratcliffe J. Randomised clinical trial comparing surgery with conservative treatment for uncomplicated varicose veins.	RCT. Computer randomisati on and group allocation by telephone. Thus allocation concealmen t very likely. No blinding reported	246 randomised. Although there were a number of bilateral cases, the total number of legs is not reported. In the surgery group there were 18.1% with bilateral surgery. However it appears that randomisation was only by the worst	to vascula hospitals Inclusion: with saph popliteal Exclusion disability complicat veins < 5m thigh.	primary vario eno-femoral o reflux. co-existing o precluding sur ions of varico nm in diameto comparison. F	large cose veins or sapheno- disease or rgery; se veins; er in lower	Use of compression hosiery, alongside lifestyle advice relating to exercise, leg elevation, and weight / diet management. Duration of treatment	Stripping surgery, done under general anaesthetic and usually as a day case. For patients with affected great saphenous veins(GSV): flush ligation at the saphenofemo	1 and 2 years.	Quality of life Patient assessed symptoms Patient satisfaction Adverse events	NHS HTA programme.
British Journal of Surgery	for patient	leg, and so results relating to the		conservati ve	surgery	unclear.	ral junction, with stripping			
2006; 93: 175-	researchers	better leg are not	F:M	87:35	83:41		of the GSV to			
181	, or	included here.	Age	49.5	49		knee level,			
AND	assessment of the	Of the 122	height	168	167.8		with multiple phlebectomie			
AND	outcomes	randomised to	BMI	26.9	26.4		S.			
Michaels JA,	relevant to	conservative	smokers	21.3%	26.6%		For patients			
Campbell WB, Brazier JE, MacIntyre JB,	the review question. Intention to	treatment, all received treatment. 21 lost	Family history of VV.	70.5%	73.4%		with affected short saphenous			
Palfreyman SJ, Ratcliffe J, Rigby K.	treat carried out in terms of	to follow-up at 1 year, leaving 101 for analysis. A	Family history of leg	7.4%	16.1%		veins(SSV): sapheno- popliteal			

Reference	Study type	No. of patients	Patient ch	aracteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Randomised clinical trial, observational study and assessment of cost- effectiveness of the treatment of varicose veins (REACTIV trial). Health technology assessment 2006; vol 10: number 13. This second article covers the same trial with the same patients, and contains the same information as the first study, with some small additions (in red).	those refusing the randomised treatment being kept in that group and analysed. However no imputation carried out for those failing to attend follow-ups.	further 63 also discontinued compression over the following 3 years, opting for surgery. However these were kept in the analysis in the conservative treatment group, as per ITT. Of the 124 allocated to surgery, 109 received surgery. 9 refused surgery and had conservative treatment instead, and 6 deferred. These 15 were kept in the group, and analysed, with ITT. 43 lost to follow-up at 1 year, and so there were 81 available for analysis at 1 year. Some of the 43 lost to follow-up were contacted and reported that their	ulcers Previous pregnan cies (mean) EQ-5D	2.1 0.74(0.11) 0.77(0.18)	2.1 0.73(0.1) 0.76(0.19)		ligation at the sapheno- femoral junction, with stripping of the SSV in some patients, with multiple phlebectomie s.			

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
		withdrawal was not due to lack of efficacy or adverse events, but mostly because they were well.						
Results:								
Outcome			Conservative		Surgery			
SF-6D 1 yr mear	n(sd)		0.73(0.11) n=98		0.77(0.10) r	i=75		
SF-6D 2 years m	nean(sd)		0.72(0.13) n=47		0.78(0.10) r	= 44		
EQ-5D 1 yr mea	n(sd)		0.78(0.18) n=101		0.87(0.14) r	i=78		
EQ-5D 2 years r	nean(sd)		0.85 (0.17) n=44		0.84 (0.21)	n=34		
SF-36			no overall scores given – o	only sub-scales given				
Aching (proport	ion same or wo	orse) at 1 yr	72/97		15/75			
heaviness (prop	ortion same or	worse) at 1 yr	52/97		9/75			
itching (proporti	ion same or wo	orse) at 1 yr	42/97		10/75			
swelling (propor	tion same or w	vorse) at 1 yr	31/97		8/75			
cosmetic concer	ns (proportion	same or worse) at 1 yr	75/97		13/75			
Patient dissatisfa	action at ?1 yea	ar (follow-up point unclea	r) 53/107		3/65			
Adverse events								
neural damage	e (footdrop reso	olving in 8/52)	0/122		1/124			
post-op pain			0/122		3/124			
phlebitis			3/122		0/124			

Evidence tables clinical studies

1

G.5 Chapter 9 – interventional treatment

.1 Stripping surgery vs. foam sclerotherapy

Table 58: Abela 2008²

Reference	Study type	No of patients	Patient	characto	eristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
Abela R et al. Reverse foam sclerotherap y of the great saphenous vein with sapheno- femoral ligation compared to standard and invagination stripping: a prospective clinical series. EUROPEAN JOURNAL OF VASCULAR & ENDOVASCU LAR	RCT, UK. Allocation concealment via sealed envelopes. Randomisation method unclear. [Technical failures occurred in 2 patients in the standard stripping group, 4 patients in the invagination group, and 3 patients in the foam sclerotherapy group. [BUT no. of people who withdrew from study was not described]	90 consecutive limbs of 82 patients with incompeten ce of the GSV resulting from varicose veins. (74 of the 82 patients had unilateral saphenous incompeten ce and 8 had bilateral incompeten ce)	old wit sympto varicos and GS by diag ultraso Exclusi	h CEAP 2 pmatic pr e veins (i V reflux, nostic du und asse on: Not c e charac Stand ard stripp ing 46 (18- 66)	and 3 imary .e. with confirn plex ssment lisclose	n SFJ ned d	Stripping surgery via 1) Standard stripping (using a Babcock- type flexible stripper) or 2) Invagination technique. Tumescent anaesthesia applied along the length of the GSV prior to stripping. All legs dressed post- operatively with foam strip padding applied externally over the length of the GSV track, which was secured using an elastic adhesive bandage. 1 day post op, drains removed, legs dressings taken down and replaced	Reverse foam sclerotherapy: 3ml of 1% sodium tetradecyl sulphate (Fibrovein®) mixed with 3ml air resulting in 6ml foam; injected into collapsed vien via the angiography catheter as this was withdrawn along the length of the vein (hence 'reverse foam'). Proximal GSV tied 5cm distal to its cut end and redundant few cm of vein excised. Complete filling of vein	2 weeks post treatment.	Post-op thigh bruising (reported by patients and observers). Adverse events, including post- procedure pain (as indicated by no. of patients using analgesia).	Not stated	

Reference	Study type	No of patients	Patien	t characto	eristics	5	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
SURGERY. 2008;36(4):4 85-490.			F:M ratio	17:13	15: 15	22:	by Class II graduated compression stockings worn continuously until follow-up at Day 15.	checked by ultrasonography. [NOTE: to ensure uniformity between the procedures, tumescent anaesthesia applied along the length of the GSV prior to stripping.] All legs dressed post- operatively with foam strip padding applied externally over the length of the GSV track, which was secured using an elastic adhesive bandage. 1 day post op, drains removed, legs dressings taken down and replaced by Class Il graduated compression stockings worn continuously until			

Reference	Study type	No pati	of ients	Patient	characteristic	S	Interve	ntion	Comparison	Length of follow-up	Outcome measures	Source of funding
									follow-up at day 15.			
Results												
Outcome		Standard st (n= 30 legs)			Invagination (n= 30 legs)	strippin	g	Reverse foam (n= 30 legs)	sclerotherapy			
Post-op bruisin	ng	Patient (%)	Observe	er (%)	Patient (%)	Observ	er (%)	Patient (%)	Observer (%)			
None		13	20		13	13		67	77			
Moderate		50	73		70	80		30	23			
Significant		37	7		17	7		3	0			
Post-procedur	e pain	Use of anal	gesic post-c	op (% pa	tients)							
No		5/30 (17%)		7/3	0 (23%)		23/	30 (77%)				
Occasional		25/30 (83%)	19/3	30 (63%)		7/3	0 (23%)				
Regular		0		4/3	0 (13%)		0					
Adverse events		-			e events attrib uring the follo			of foam sclerot	herapy were			

Author's conclusions: Standard stripping of the GSV and invagination stripping are not associated with major discomfort and problems in the early post-op period. SFJ ligation and GSV reverse foam sclerotherapy yielded greater patient satisfaction with less post-op bruising and discomfort and reduced analgesic requirements. (NOTE: data not reported in paper for the outcomes in bold)

Reference	Study type	No. of patients	Patient ch	naracteristics		Intervent ion	Comparis on	Length of follow-up	Outcome measures	Source of funding
Bountouroglo u DG, Azzam M, Kakkos SK, Pathmarajah M, Young P, Geroulakos. Eur J Vasc Surg 2006; 31: 93-100.	RCT. Allocation decided by random drawing of sealed envelopes. With this form of randomisation, allocation concealment likely initially, but less likely as study goes on (see notes for Liu et al. 2011). No mention of blinding.	60 patients	varicose va incompete for varicos case surge Exclusion both the G surgery or or risk fact PVD; relev pregnancy	primary varice SV and SSV; pr sclerotherapy; cors for DVT; Co rant allergies; n	v us treatment ility for day osities involving rev. var. Veins history of DVT oagulopathy; nalignancy;	Ligation performe d at SFJ. GSV stripped from SFJ to a level just below the knee. General anaesthet ic used.	Ligation performe d as for stripping under LA. Varicositi es injected with 6mL of a 3% STD sclerosant (foam), then	3 weeks and 3 months	AVVQ VCSS Adverse events Treatment failure	None
				Stripping	Sclero	Multiple	compressi on			
			Age	20-76	21-72	phlebecto mies also	applied using			
			Female	60%	47%	performe d.	foam			
			C2	8/28	11/30		pads and a class II			
			C3	14/28	8/30	6	compressi			
			C4	6/28	7/30	Compress ion	on stocking			
			C5	1/28	3/30	bandages applied	for 2 weeks.			
			C6	1/28	1/30	post op	WEEKS.			
			VCSS	2-16	2-13	and then				

Table 59: Bountouroglou 2006³⁰

Reference	Study type	No. of patients	Patient characteristics	Intervent ion	Comparis on	Length of follow-up	Outcome measures	Source of funding
Reference	Study type	patients		replaced by a class 1 elastic stocking for 3 weeks. Randomis ed n=30; 2 did not receive treatment at all (1 moved, 1 moved out of area). 28 attended 3 week assessme nt. 5 lost to 3 month assessme	on Randomis ed n=30; All received treatment . 30 attended 3 week follow-up. 1 lost to 3 month follow-up	follow-up	measures	funding
				nt (no reasons given).				

Evidence tables clinical studies

Stripping surgery (n=28)

Foam sclerotherapy (n=30)

Reference	Study type		No. of patients	Patient characteristics		Intervent ion	Comparis on	Length of follow-up	Outcome measures	Source of funding
HRQoL – AVVS	[median	pre:2	6.1		pre:15.4					
		post:1	14.1		post:9.3					
VCSS [median (range)]	3(0-4))		1(0-5)					
CEAP [median (range)]	1(0-5))		1(0-5)					
Adverse events	(time not stated)									
DVT		0/28			0/30					
PE		0/28			0/30					
phlebitis		0/28			3/30					
skin pigmentati	ion	1/28			2/30					
neural injury		2/28			0/30					

Author's conclusions: US guided sclerotherapy combined with sapheno-femoral ligation was less expensive, involved a shorter treatment time and resulted in more rapid recovery.

Reference	Study type	No. of patients	Patient cha	racteristics	1	Intervent ion	Comparis on	Length of follow-up	Outcome measures	Source of funding
Figuerido M, Araujo S, Barros N, Miranda F. Results of surgical treatment compared with ultrasound- guided foam sclerotherapy in patients with varicose veins: a prospective randomised study. Eur J Vasc Surg 2009;	RCT. Allocation decided by random drawing of papers from a box. With this form of randomisation, allocation concealment likely initially, but less likely as study went on (see notes for Liu et al. 2011). No mention of blinding.	60 patients	vascular sur, Inclusion: N varicose veit Exclusion: H thrombophi bronchial as syndrome; s immobility; arterial insu	gery outpa lo previous ns; age 18- listory of D lia, allergy thma, post severe syste pregnancy; fficiency; A abetic foot; nocardiogra	treatment of 70; C5; VT, to polidocanol, -thrombotic emic disease; ; peripheral BI<0.8); LL patent foramen aphy.	Saphenof emoral or saphenop opliteal ligation combined with saphenou s stripping and phlebecto my for varicose saphenou s tributarie s and ligation of incompet ent perforatin g veins. All surgery done in one session. Regional anaesthes ia used. Inelastic	Injections in standing. Injections of foam made into the saphenou s trunk. Accessory veins cannulate d using 25 gauge butterfly needles. Foam was polidocan ol and air in a ratio of 1:4. The GSV received 8-10 ml with a polidocan ol concentra tion of 3%, the small	1,2 and 6 months post interventi on.	VCCS Adverse events treatment failure	None

Table 60: Figueiredo 2009⁹⁷

Reference	Study type	No. of patients	Patient characteristics	Intervent	Comparis on	Length of follow-up	Outcome measures	Source of funding
				bandages	saphenou			
				2 days	s vein 5ml			
				post op	at a			
				and then	concentra			
				30-40	tion of 1			
				mmHg	or 3%, the			
				below	accessory			
				knee .	veins 5ml			
				compressi	at a			
				on for 2	concentra			
				months.	tion of 1% and			
					perforatin			
					g veins 1-			
				30	2 ml at a			
				randomis	concentra			
				ed.	tion of			
				Unclear	1%. Foam			
				how	progress			
				many	along			
				received	veins			
				surgery,	imaged			
				but 1 lost	with US.			
				to follow-	Maximum			
				up	bolus of			
				(reasons	10ml in			
				not given)	one			
					session.			
					Sessions			
					repeated			
					as needed			
					every 30			
					days up to			

Reference	Study type	No. of patients	Patient characteristics	Intervent ion	Comparis on	Length of follow-up	Outcome measures	Source of funding
					а			
					maximum			
					of 3			
					(average			
					sessions			
					per			
					patient			
					were 2.1).			
					After 15			
					mins of			
					compressi			
					on of the			
					SFJ or			
					SSV, the			
					limb			
					bandaged			
					using an			
					inelastic			
					bandage			
					for 3-5			
					days.			
					Then 30-			
					40 mmHg			
					below			
					knee			
					compressi			
					on for 3			
					months.			
					30			
					randomis			
					ed.			
					Unclear			

Reference	Study type	No. of patients	Patient characteristics		Intervent ion	Comparis on	Length of follow-up	Outcome measures	Source of funding
						how many received surgery, but 3 lost to follow- up (reasons not given)			
Results: mean	(sd)								
		Stripp	ing surgery (n=29)	Foam so	lerotherapy	(n=27)			
VCSS pain 30 d	ays	0.93(0.		0.89(0.5					
VCSSpain 60 da	ays	0.79(0.	49)	0.59(0.5	0)				
VCSS pain 180	days	0.72(0.	53)	0.56(0.5	1)				
VCSS oedema	30 days	0.69(0.	60)	0.70(0.5	4)				
VCSS oedema 6	60 days	0.59(0.	63)	0.56(0.6	4)				
VCSS oedema1	.80 days	0.55(0.	63)	0.48(0.6	4)				
VCSS inflamma	ition 30 days	0.76(0.	44)	0.89(0.3	2)				
VCSS inflamma	ition 60 days	0.72(0.	45)	0.89(0.3	2)				
VCSS inflamma	ition180 days	0.72(0.	45)	0.89(0.3	2)				
Adverse events	5								
neurological (s	ubjective)	6/29		0/27					
Reflux/recanali	isation	3/29		6/27					

Author conclusions: US guided foam sclerotherapy is a safe and effective option for patients with chronic venous disorders.

Reference	Study type	No. of patients	Patient chara	octeristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Kalodiki E, Lattimer C, Azzam M, Shawish E, Bountouroglo u D, Geroulakos G. Long term results of a randomised controlled trial on ultrasound- guided foam sclerotherapy combined with saphenofemor al ligation vs. standard surgery for varicose veins. Journal of vascular surgery; 2011 (in press). NB This article in press has many typos and errors.	RCT. Patients selected by drawing sealed envelopes. For impact of this on likelihood of adequate allocation concealment please see notes on Liu et al. 2011. Initially only the most symptomatic leg was randomised in bilateral patients. However if varicose veins developed in the contralateral limb this was included and given the same randomisation.	73 patients (82 legs).	the GSV and s previous surg varicosities, p PVD, relevant pregnancy.	varicosities previous tr ay case surg imary varic saphenous ery or scler bast DVT, Co c allergies, r	involving the eatment and gery. osities involving vein (???), otherapy for	Conventio nal high DFL stripping surgery combined with multiple phlebecto mies using Muller hooks. General anaesthes ia used. Immediat ely post- op compressi ve bandages applied, which were changed for a compressi ve stocking before	Foam sclerother apy with 6mL of 3% Sodium tetra decyl sulphate (STS) (1.2 mL of STS mixed with 4.8 mL of air) injected directly into the vein under US guidance. Post procedur e a 18-24 mmHg thigh high graduated elastic compressi on stocking was	3 weeks, 3,6 and 12 months and yearly thereafter . Median follow-up was 5 years.	HRQoL – SF36 and AVVs. Physician reported disease severity – CEAP, VCSS, VSDS. Adverse events Reflux	None

Table 61: Kalodiki 2011¹²⁹21

Reference	Study type	No. of patients	Patient chara	cteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source o funding
			SF-36	unclear) No report on similarity. No data given but appear very similar on figure		discharge from the day ward. n=43 limbs (39 subjects)	applied. Patients worse this for 2 weeks continuou sly, and then 1 week in the day only for another week. Patients also told to walk for 2 miles / 2 hours daily n=39 limbs (34 subjects)			
Results: Mediar	n (IQR) given.									
		Stripping surgery	r (n=43 legs)	Foams	clerothera	py (n=39 legs) р			

Stripping surgery (n=43 legs)Foam sclerotherapy (n=39 legs)pAVVQ at 3 years8.94 (IQR unclear)4.97(IQR unclear)NS??AVVQ at 5 years5.45(IQR unclear)7.345(IQR unclear)0.015SF-36Data only given in figures, and p values
only given in text. Reported that noFoam sclerotherapy (n=39 legs)p

Reference	Study type	No. of patients	Patient characteristics		Interventi on	Comp on	paris	Length of follow-up	Outcome measures	Source of funding
			en groups in changes =0.724) and mental							
years VCSS increase fr	om baseline to 3 om baseline to 5	1(0-9) very unclearly rep	orted	1(0-9) very unclearly rep	orted		0.504			
years VCSS absolute so VCSS absolute so		reported no diff b reported no diff b		0.313 0.104						
VSDS at 3 years		0.5(IQR unclear)		1.0(IQR unclear)			0.780			
VSDS at 5 years Adverse events		1.0(IQR unclear)		0.25(IQR unclear)			0.388			
thromoboemb	olism (DVT and PE)	0/43		0/39						
major neurolo	gic event (i.e. stroke)	0/43		0/39						
skin pigmentat		2/43		1/39						
thrombophleb		0/43		3/39						
saphenous ner	rve injury	2/43		0/39						
hematoma		1/43		0/39						
skin ulcer Reflux at 3 years	SABOVE KNEE*	1/43 7/26		0/39 11/33						
Reflux at 5 years	S ABOVE KNEE	9/26		13/33						

Reference Study type	No. of patients	Patient characteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Reflux at 3 years BELOW KNEE	14/26		19/33					
Reflux at 5 years BELOW KNEE	9/26		14/33					

*cannot sum above and below knee data as some may be from the same subjects

Author's conclusions: At 3 and 5 years of follow-up, the treatment was equally effective in the surgical and foam groups, as demonstrated with VCSS, VSDS and the SF-36...at 5 years the AVVQ was significantly better in the surgical group.

Reference	Study type	No of patients	Patient characteristics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Liu X, Jia X, Guo W, Xiong H, Zhang M, Liu X, Du X, Zhang MH. Ultrasound- guided foam sclerotherap y of the great saphenous vein with sapheno- femoral ligation compared to standard stripping: a prospective clinical study. International Angiology 2011; 30: 321-6.	RCT; China. Allocations placed in 60 sealed envelopes and then shuffled. Each recruited patient given one. Not stated if there was any patient blinding. Also no mention of assessor blinding. Although the method of allocation meant that group allocation was automatically concealed from the recruiter at first, it is possible that towards the end of recruitment it might become possible to predict the allocation of the next patient to be recruited (i.e. if after 50 patients it	60 patients randomis ed; 59 treated.	Patients undergoing treatment for varicose veins in a vascular surgery clinic; 26 men and 34 women; CEAP ranged from C2-C6 (no breakdown given). Median age 49 (range 37-66). Inclusion: symptomatic primary varicose veins with primary SFJ and GSV reflux, as shown by duplex (reflux duration >0.5 secs after calf compression-release manoeuvres). Exclusion: None given Baseline characteristics: Demographics not given. No breakdown of any characteristics done by group. However, groups matched for CEAP median (range): both 4(2-6). AVVQ median (range) was similar: Sclerotherapy: 15(11-26); surgery: 19(14-29).	N=30 Stripping surgery, using a flexible intraluminal stripper to strip from groin to knee. Preceded by flush ligation, division of tributaries. Varicosities also treated by phlebectomy. Done under GA. All patients with residual varicose veins in both groups received additional foam sclerotherapy as outpatients (not stated when). 30 randomised; 29 received treatment (patient changed mind); 1 loss to follow-up at 3 months (didn't	N=29 Ultrasound-guided foam obliteration of the GSV. After SFJ ligation, 6mL sclerosing foam (1 part of 1% Lauromacrogol [Polidocanol] and 4 parts of air) injected into GSV proximal cut end via 10mL syringe connected to a 21 gauge butterfly. Sclerosant foam flow monitored via US. Remnant trunks not filled with foam were punctured and filled with additional sclerosant. Done under GA . All patients with residual varicose veins in both groups received additional foam	3 months and 6 months post-op.	AVVQ Physician reported Adverse events treatment failure	None

Table 62: Liu2011¹⁴⁸

Reference	Study type	No of patients	Patient characteristics		Intervention	Cor	nparison	Length of follow- up	Outcome measures	Source of funding
	was known that 28/30 stripping places had been already taken up, then it would be known that the probability of the next patient being a control patient would be 80%).				attend follow-up); No further loss to follow-up at 6 months.	30 f rec 2 lo at 3 atto furt 6 m	erotherapy as ipatients (not ted when). randomised; 29 eived treatment; oss to follow-up 8 months (didn't end follow-up); 2 ther patients t to follow-up at nonths (didn't end follow-up).			
Results:										
			Stripping surgery (n=30)	Foa	m sclerotherapy (n=29	Ð)	р			
HRQL										
AVVQ media	in (range) 3 months		12(8-17)	9(5	-16)		no intergroup p	value given		
Physician rep	orted outcomes									
CEAP median	(range) 3 months		1(0-4)	1(0	-3)		no intergroup p	value given		
Adverse ever	nts									
groin hemato	oma		1/30	0/2	9					
neural injury			2/30	0/2	9					

Reference	Study type	No of patients	Patient characteristics		Intervention	Con	nparison	Length of follow- up	Outco meas	Source of funding
thrombophlet	pitis		1/30	3/2	9					
skin pigmenta	tion		1/30	2/2	9					
DVT			0/30	0/2	9					
PE			0/30	0/2	9					
Post op use of	fanalgesia (any)		24/30	8/2	9					
No full obliter session of scle	ation 3 months (requiri ero)	ing second	3/28	3/2	8					
No full obliter session of scle	ation 6 months (requiri ero)	ing second	3/26	5/2	5					

Author conclusions: US guided sclerotherapy, combined with sapheno-femoral ligation involved a shorter treatment time, less post-operative discomfort and resulted in more rapid recovery compared to conventional GSV stripping.

Reference	Study type	No of patients	Patient char	acteristic	S		Stripping surgery	Foam sclerotherap Y	Endotherm al ablation	Length of follow-up	Outcome measures	Source of funding
Rasmussen LH, Lawaetz M, Bjoern L, Vennits B, Blemings A, and Eklof B. Randomize d clinical trial comparing endovenou s laser ablation, radiofrequ ency ablation, foam sclerothera py and surgical stripping for great saphenous varicose veins. British Journal of Surgery 2011;	RCT, Denmark. 2 private surgical centres. Randomisatio n in blocks, but method not clearly described. Allocation concealment adequate via sealed envelopes. No blinding reported.	500 patients were randomise d 125 in each group. The results for endotherm al laser ablation (n=125) and radio frequency ablation (n=125) have been combined in these results. All received interventio n except one patient (and 1 leg) in the	Inclusion: 18 varicose veir Saphenous V defined by re Bilateral treat in same grout recurrent va if GSV preser Exclusion: D saphenous t anterior acco small sapher DVT; history brachial prese axial deep vei GSV. Baseline chat (range). Auth well-matchet N subjects n legs Age	ns; CEAP 2 Vein(GSV) eflux >0.5 atment al up. Patien ricose vei rved to the uplication runk or a essory sap nous vein of arteria ssure inde ein insuffi	2-4; Gre i incomp isecs or lowed s its with ins also ne groin n of the n incom phenou reflux; al insuff ex <0.9, iciency;	eat petence, duplex; so long as included n. petent s vein; previous ficiency: or both; tortuous	N=125 (143 legs). Flush ligation of the GSV and division of all tributaries . Use of a PIN stripper to strip GSV to just below the knee. Common procedure s: Phlebecto mies to remove varicositie s in all groups. Compressi on applied post operativel	N=125 (145 legs) Ultrasound- guided foam sclerotherapy [UGFS] (in the reversed Trendelenbur g position); 3% polidocanol (Aethoxyscler ol®); 2ml solution mixed with 8ml air. Retreatment allowed within 1 month. Common procedures: Phlebectomie s to remove varicosities in all groups. Compression applied via a 30 mmHg	N=250 (292 legs) Endovenou s Laser ablation (EVLA): Under duplex guidance with 980nm diode laser for 1 st 17 pts and then 1470 diode laser for the rest. Pulse mode was used in one centre but continuous used in the other. Cannulation just below the knee. The laser fibre was advanced	3 days, 1 month, 1 year. It was intended to continue follow-up yearly for 5 years.	Quality of life: SF36 score and AVVQ. Physician- reported outcomes: VCSS. Treatment failure: defined as a patent GSV with reflux, or GSV not stripped successfully Recurrence rates Adverse events (including complicatio ns e.g. DVT and PE, post- interventio n superficial	Public health Insuranc e Research Foundati on of Denmark

Reference	Study type	No of patients	Patient chara	acteristic	S		Stripping surgery	Foam sclerotherap Y	Endotherm al ablation	Length of follow-up	Outcome measures	Source of funding
Reference 98:1079– 1087.	Study type		Patient chara %Female CEAP2-3 (legs) CEAP 4-6 (legs) Previous surgery GSV (diam) (mm) Number of phlebecto mies surgeon's time (min)	(19- 72) 77 97 3 8 7.8 (3-14) 15 (1- 48) 32 (15- 80)	s (18- 75) 76 96 4 4 4 8.7 (3- 20) 15 (1- 43) 19 (5- 145)	(18-75) 77 97 3 3 8 7.8 (3- 14) 15 (1- 48) 32 (15- 80)		sclerotherap	al ablation until 2cm below the sapheno- femoral junction, and then the GSV was ablated during withdrawal. Endovenou s radiofreque ncy ablation (EVRF): catheter advanced under US guidance to 2cm below the sapheno- femoral junction.	-		of
		endotherm al group and n=27 (34 legs) in the stripping group.							junction. Then withdrawn, with temperatur es maintained			

Reference	Study type	No of patients	Patient characteristics	Stripping surgery	Foam sclerotherap y	Endotherm al ablation	Length of follow-up	Outcome measures	Source of funding
						degrees for 20 seconds per segment via a thermostat. Common procedures: Phlebectom ies to remove varicosities in all groups. Compressio n applied post operatively for all – at			

Results

	Stripping surgery (n=124 patients)	Foam Sclerotherapy (n=124 patients)	Endovenous Laser Ablation (n=125)	Radiofrequency ablation (n=125)
Treatment failure at 3 days ^a	4/141 legs	3/143 legs	0/143	0/146
Treatment failure at 1 month	3/135 legs	2/144 legs	1/144	0/141
Recurrent varicose veins at 1 year	16/108	17/123	7/121	6/124

Reference	Study type	No of patients	Patient characteristic	cs	Stripping surgery	Foam sclerotherap Y	Endotherm al ablation	Length of follow-up	Outcome measures	Source of funding
Adverse eve	ents during 1 st n	nonth								
		Stripping surg	ery	Foam Sclerother	ару	Endovenous Las	er Ablation	Radiofre	equency ablat	ion
Major:										
DVT (requiri	ng treatment)	1/135		1/144		0/144		0/141		
PE (requiring	g treatment)	0/135		1/144		0/144		0/141		
Minor:										
Phlebitis		5/135		17/144		4/144		12/141		
Infection		1/135		4/144		0/144		1/141		
Paraesthesia	1	5/13		2/144		3/144		6/141		
hyper pigme	entation	6/135		8/144		3/144		8/141		
Haemorrhag	ge	1/135		1/144		1/144		0/141		
	n first 10 days VAS, 0-10, 10	2.25 (2.23)		1.60 (2.04)		1.21 (1.72)		2.25 (2.)	23)	
SF-36 score	(mean [SD]) at	1yr								
		Stripping surg	ery	Foam Sclerother	ару	Endovenous Las	er Ablation	Radiofr	equency ablat	ion
Physic	cal functioning		92.82 (13.35)	91.	.33 (14.93)		92.02 (11.6	1)	g	92.22(12.62)
	Role physical		93.41(16.32)	90.	.36 (20.56)		93.51(14.7	8)	ç	4.65(10.64)
	Bodily pain		88.77(17.11)	85	.11(23.45)		88.43(19.5	5)	8	9.92(16.85)
(General health		66.02 (14.00)	63.	.36 (18.31)		64.90(11.9	9)	e	7.08(11.82)
	Vitality		76.99 (15.54)	73.	.20 (22.67)		77.74(14.0	3)		76(17.51)
Soc	ial functioning		95.19 (11.60)	93.	.10 (16.51)		96.51(11.2	2)	g	7.11(14.45)

91.92 (17.11)

84.58 (15.77)

95.95(10.15)

87.70(10.51)

94.20 (14.02)

85.92 (12.18)

Role – emotional

Mental health

94.5(11.02)

87.08(11.94)

Reference	Study type	No of patients	Patient characteristic	S	Stripping surgery	Foam sclerotherap Y	Endotherm al ablation	Length of follow-up	Outcome measures	Source of funding
Summary Sc	ores									
PHYSICAL 4 v	weeks	48.14(7.21)		49.2(7.56)		47.68(6.95)		49.88(7)		
PHYSICAL 1 y	vear	53.33(5.9)		51.94(7.66)		52.62(5.98)		53.23(5.	32)	
MENTAL 4 w	eeks	55.15(7.81)		56.1(7.51)		55.55(8.21)		55.57(7.	38)	
MENTAL 1 ye	ear	55.83(6.31)		54.73(8.89)		56.74(5.44)		56.52(6.	17)	
VCSS		NOTE that onl	y graphs were presente	ed, and no actual o	data were pr	ovided.				
AVVSSS		NOTE that onl	y graphs were presente	ed, and no actual o	data were pr	ovided.				
Time to retur activities (da median(rang	ys)	4 (0-30)		1 (0-30)		2(0-25)		1(0-30)		
Time to retur (days) media	in (range)	4.3 (0-42)	SV not strinned successfu	2.9 (0-33)		3.6(0-46)		2.9(0-14		

(a) Failure defined as patent GSV with reflux OR GSV not stripped successfully (unfair comparison as unstrapped GSV does not equal incompetent GSV, whereas unblocked GSV has to occur with reflux.

Reference	Study type	No. of patients	Patient ch	aracteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Shadid N, Ceulen R, Nelemans P. et al. Randomised clinical trial of ultrasound- guided foam sclerotherapy versus surgery for the incompetent great saphenous vein.	Multicentre randomised controlled trial. Computer generated randomisation with permuted blocks of eight.	 460 randomised (233 to foam and 227 to surgery). 3 did not receive foam (2 declined and 1 not feasible) and 27 did not receive surgery (24 declined, 1 CVA, 1 surgeon not co-operating, 1 pregnancy). 17 lost to follow-up in Foam; 23 lost to follow-up in stripping. Resaons did not appear to be related to outcome. 	incompeter symptoms incompeter reflux time venous sy Exclusion contraind	active ulcerati	2-5EpAsPr or more n with and GSV, with ormal deep on;	Stripping surgery. Day case under GA or spinal. Ligation of SFJ and GSV divided and stripped to just below knee. Phlebecto mies done as needed. Class II elastic stockings worn for 6 weeks.	Foam sclerother apy. 3% policodan ol in a foam (1:4 ratio of sclerosant to air). Class II elastic stockings worn for 6 weeks. NO CROSSECT OMY	2 years	Recurrence (clinical symptoms plus reflux) at 2 years Reflux (regardless of symptoms) at 2 years EQ-5D Symptoms Adverse events	None stated

Table 64: Shadid2012²⁴⁴

Reference	Study type	No. of patients	Patient characteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Results:									
			Stripping surgery	Foam sclerothe	rapy				
Clinical recurr reflux)at 2 yea	ence (symptoms with ars	accompanying	16/177	24/213					
Clinical recurr reflux)at 1yea	ence (symptoms with r	accompanying	13.8%	11.9%					
Clinical recurr reflux)at 3 mo	ence (symptoms with onths	accompanying	8.56%	10.8%					
Reflux at 2 yea	ars		18.2%	21.3%					
Reflux at 1yea	r		23.1%	29.16%					
Reflux at 3 mc	onths		18.2%	21.3%					
Change in VCS	S from baseline at 2 γ	/ears	-1.75 (2.135)	-1.49 (2.135)			2 (no variance each calculate	es reported bu ed as: 2.135)	t common
Mean change	in EQ-5D from baseli	ne at 2 years	0.061 (0.211)	0.064 (0.211)			9 (no variance each calculate	es reported bui ed as: 0.211)	t common
More pain at 2	2 years		6/177	14/213					
More tired/he	eavy feeling at 2 years		5/177	6/213					
More cramps	at 2 years		8/177	8/213					
More restless	legs at 2 years		21/177	29/213					
Patient not sa	tisfied – aesthetic at 2	2 years	23/177	31/213					
Patient not sa	tisfied – functional at	2 years	17/177	17/213					
More pain at 2	1 year		14/188	20/221					
More tired/he	eavy feeling at 1 year		9/188	5/221					
More cramps	at 1 year		9/188	10/221					
More restless	legs at 1 year		26/188	34/221					
Patient not sa	tisfied – aesthetic at 2	1 year	32/188	33/221					

Reference	Study type	No. of patients	Patient characteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Patient not satis	sfied – functional at 1	-	28/188	22/221					Ū
More pain at 3 r	months		10/176	12/217					
More tired/hear	vy feeling at 3 month	IS	2/176	8/217					
More cramps at	: 3 months		6/176	9/217					
More restless le	egs at 3 months		16/176	27/217					
Patient not satis	sfied – aesthetic at 3	months	19/176	39/217					
Patient not satis	sfied – functional at 3	3 months	15/176	20/217					
Mean change of to 2 years	f EQ-%D "health state	e" from baseline	-1.8(25.4)	-0.36(25.4)			77 (no varianc each calculate	es reported bu ed as: 25.4)	t common
Complete satisfa complaints at 2	action with reductior years	n in venous	117/177	127/213					
Need for >1 trea	atment		10/200	40/230					
			[2 for re-surgery, 8 for foam]	[35 one extra se >1 extra session		ı, 5			
Adverse events	(within 1 week)								
Groin infection			4/200	0/230					
Haematoma			3/200	0/230					
Parasthesia			6/200	0/230					
Pain at injection	n site		0/200	6/230					
Thrombophlebit	tis		0/200	17/230					
Headache/migra	aine		0/200	3/230					
DVT			0/200	1/230					
PE			0/200	1/230					
Later adverse ev	vents (at 2 yrs)								
Hyper-pigmenta	ation		2/200	12/230					
Telangiectatic m	natting		2/200	6/230					

Evidence tables clinical studies

1

Reference	Study type	No. of patients	Patient cha	aracteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
Wright D, Gobin JP, Bradbury AW, Coleridge- Smith P, Spoelstra H, Berridge D, Wittens CHA, Sommer A, Nelzen O, and Chanter D. Varisolve polidocanol microfoam compared with surgery or sclerotherapy in the management of varicose veins in the presence of trunk vein incompetence : European randomized controlled trial. Phlebology 2006;21:180 – 190.	RCT, open-label, international, multicentre. Randomisation not described. Allocation concealment unclear. No blinding.	2 cohorts but only looking at the Varisolve cohort/ surgery (n=311); 210 patients randomis ed to Varisolve and 101 randomis ed to Surgery. Withdrew from study after treatment : 32 patients from Varisolve group and 7 from the	years with veins (C2-C incompeter blood flow demonstra saphenous 10cm proxi incompeter system on of occlusion of reflux wa Exclusion: of major su venographi evidence o immobility, contraindic including so asthma and	moderate-to (4); SFJ or SPJ nce or both v for >1s and < ted by duples incompetent incompetent imal trunk ve nce; normal of duplex scann n or incompetent as acceptable C5 and C6 ex- uperficial thro ic or ultrason f current or p ; BMI >32kg/fic cations for po evere hypertor	vith retrograde 75 4 scanning; great 2 ce and/or short 2 c; minimum of 3 in 3 deep venous 3 ing, no evidence 4 tence. Evidence 4 tence. Evidence 5 cluded; history 9 mbophlebitis; 9 ographic 1 revious DVT; m ² ; 1 lidocanol ension, diabetes, eriosclerosis.	N=94 patients treated with Surgery: high ligation performe d in 91.5% of patients, stripping in 88.3% and avulsion phlebeco my in 53.2%.	N=178 patients treated with Varisolve [®] polidocan ol microfoa m; a uniform foam of physicolo gic gases, principally oxygen and carbon dioxide, combined with a 1% aqueous solution of polidocan ol. Maximal dose initially set at 60mL and subseque	Days 7 and 28, and at months 3 and 12.	Post- procedure pain at Day 6 (Visual Analogue Scale (VAS) 1-100mm scale) Response defined as: occlusion (or for surgery, absence) of the treated vein AND elimination of junctional reflux. Adverse events: common treatment- related adverse events (i.e. contusion, skin	Acknowl edgement t of Sigvaris Ltd for the provision of compress ion stockings , but none declared for the provision of Varisolve

Reference	Study type		No. of patients	Patient cha	aracteristics		Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
			surgery group.	M:F	34:60	66:112		ntly reduced		discolouratio n, pain in	
				CEAP C3	11 (11.7%)	14 (7.9%)		to 30mL. The		limb, headache	
				CEAP C4	10 (10.6%)	20 (11.2%)		Varisolve technique		and haematoma)	
				Primary VV	/			carried		and serious	
				GSV	77 (81.9%)	141 (79.2%)		out under ultrasoun		adverse events (i.e.	
				SSV	8 (8.5%)	12 (6.7%)		d guidance.		DVT, PE)	
				GSV+SSV	0	8 (4.5%)		30- 40mmHg			
					Recurrent VV			thigh length			
			GSV	6 (6.4%)	11 (6.2%)	comp	compressi				
				SSV	2 (2.1%)	1 (0.6%)	on stocki	stocking	stocking		
				GSV+SSV	1 (1.7%)	3 (1.7%)		worn 14 days post treatment			
Results:											
		Surgery (n=9	-		foam sclerot	herapy (n=178)	р				
Response (i.e. oc	clusion of tru	unk vein and e	limination of	reflux)							
at 3 months		82/94 (87.2%	6)	120/176 (68.2%)						
at 12 months		81/94 (86.2%	6)	111/176 (63.1%)						
Post-procedure p 6 (Median score		9		2			<0.001				

Reference	Study type		No. of patients	Patient characteristics	Interventi on	Comparis on	Length of follow-up	Outcome measures	Source of funding
100mm VAS sca			•						Ū
Most common t	reatment-rela	ted adverse ev	vents						
Contusion		80 (85.1%)		122 (68.5%)	-				
Skin discolourati pigmentation)	ion (= hyper	39 (41.5%)		98 (55.1%)	-				
Pain in limb		39 (41.5%)		73 (41%)	-				
Headache		20 (21.3%)		41 (23%)	-				
Haematoma		1 (1.1%)		11 (6.2%)	-				
Serious adverse	events								
Deep vein throm	nbosis	0		9 (4.5%)	-				
Pulmonary emb	olism	0		0	-				

Author's conclusions: Varisolve was non-inferior to alternative treatment. Surgery was more efficacious, but Varisolve caused less pain. The Varisolve technique is a useful additional treatment for varicose veins and trunk vein incompetence.

G.5.2 Stripping surgery vs. endothermal ablation

Table 66: Carradice201147

Reference	Study type	No. of patients	Patient ch	naracterist	ics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Carradice D, Mekako AI, Mazari FAK, Samuel N, Hatfield J and Chetter IC. Randomised clinical trial of endovenous laser ablation compared with conventional surgery for great saphenous varicose veins. British Journal of Surgery 2011A; 98:	RCT. Non blinded. "Randomisation" through choice of sealed envelopes by the patient. This method, though unorthodox, ensures allocation concealment provided all envelopes are identical and have been properly shuffled. However, towards the end of the trial, it may become	280 randomised (140 from each group). In endovenous laser ablation (EVLA) group, 1 did not receive intervention; in surgery group, 3 did not receive intervention (all withdrew from trial). Over one year, in EVLA group 24	symptoma veins, with incompete GSV. Incorreflux of a Doppler. Exclusion for ipsilate deep vein obstruction pregnancy pulses. Baseline o significant SF36 men Continuou	atic unilate h isolated S	reflux in the defined as ec on treatment se veins, ence or 8 years, ole foot stics: Only e was for (p=0.03). mean (sd)	Stripping surgery. Flush SFJ ligation followed by ligation of all tributaries to the 2 nd branch, then inversion stripping of the GSV to the knee. GA used. Common procedures: Multi- phlebectomies given to both groups as needed.	EVLA. GSV cannulated at the lowest point of reflux. EVLA at 810nm wavelength and power 14W applied during withdrawal. LA used.	1 week, 6 weeks, 3 months and 1 year.	SF-36 EuroQoL 5D AVVQ CEAP VCSS post-op pain Satisfaction with cosmetic result.	None
501-510.	easier to guess	lost to		EVLA	Surgery					
	to which group recruited subject will be allocated,	follow-up and in surgery	Age	49 (14)	49 (13)					
	if it is known that there has	group 15 lost to	M:F ex smoker	54:85 35/132	47:90 37/130					

Reference	Study type	No. of patients	Patient ch	aracteristi	cs	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding																			
	been an imbalance of	follow-up.	current smoker	35/132	30/130																								
	allocations up to that point. In		BMI	26.6 (5.0)	26.0 (4.3)																								
	that sense, the risk of allocation concealment		GSV diam (groin) mm	8.7 (2.7)	8.2 (2.7)																								
	breaking down is		VCSS*	4 (3-5)	4 (3-5)																								
	possible.		CEAP 2	P 2 95/138 96/137																									
			CEAP 3-6 43/138 41/137																										
			AVVQ*	12.6 (9.6- 17.2)	13.7 (9.9- 18.2)																								
			SF36*																										
			physical functioning	90 (75- 100)	90 (80-100)																								
			role physical	100 (50- 100)	100 (75-100)																								
			bodily pain	74 (52- 100)	74 (52-100)																								
																						gen health	77 (62- 92)	77 (67-87)					
																			Vitality	70 (55- 80)	70 (53-80)								
			social functioning	100 (75- 100)																									
			role emotional	100	100																								
			mental	84 (68-	80 (68-90)																								

Reference	Study type	No. of patients	Patient characteristics			Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
			health	92)							
			EQ5d*	0.848 (0.796- 1.00)	0.841 (0.796- 1)						
			SF6D*	0.804 (0.744- 0.856)	0.795 (0.717- 0.847)						
Results:											
			EVLA			Surgery	р	р			
Health related	d QoL (SF36) High	er score is bette	r outcome).	Median (IQ	R)						
physical functioning -1week			88 (70-95)			80(65-90)	0	0.012			
role physical -1week			100 (25-100)			50(0-100)	0	0.005			
bodily pain -1week			74(54-84)			62(41-74)	62(41-74)		0.031		
gen health -1week			81(67-92)			82(72-92)	Ν	NS			
vitality -1week			70(60-80)			65(55-800	0.049				
social functioning -1week		100 (75-100)			75(63-100)		0.004				
role emotional -1week			100			100(67-100)		0.027			
mental health -1week			88(76-92)			84(68-92)		NS			
physical functioning -1year			95(85-100)			95(80-100)					
role physical -1year			100			100		IS			
bodily pain -1year			100(72-100)			94(72-100)		NS			
gen health -1year			82(67-92)			82(72-92)		NS			
vitality -1year			75(60-85)			75(65-85)	5-85) NS				
social functioning -1year			100(88-100)			100(75-100)	Ν	NS			
role emotional -1year			100			100	Ν	NS			
mental health -1year			88(74-92)			88(76-92)	Ν	NS			
AVVQ 1 week			16.6 (12	.4-21.1)		16.5(12.2-22.7)	Ν	NS			
AVVQ 1 year			2.0(0-5.3	3)		2.0(0-5.3)	Ν	NS			

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
EQ5D - 1 week EQ5D – 1 year			0.796(0.760-1) 1(0.877-1)	0.801(0.691-0.895) 1(0.841-1)		NS NS		
SF-5D – 1 wee SF-5D – 1 year			0.796(0.735-0.838) 0.843(0.773-0.876)	0.759(0.672-0.830) 0.835(0.777-0.878)		0.003 NS		
pain at day 1			Less pain from day 1 to day 6 i no data given except in low re	e	001). But			
VCSS at 1 year			1(0-1)	1(0-1)		NS		
Adverse event	S							
sensory distur	bance		4/137	13/133		0.020		
haematoma			1/137	11/133		0.003		
infection			2/137	8/133	(0.048		
phlebitis			4/137	6/133		0.536		
persistent pair	ı		1/137	5/133		0.116		
pigmentation			4/137	1/133		0.371		
anaesthetic co	mplication		0/137	3/133		0.118		
persistent bru	sing		1/137	2/133	(0.618		
allergy			0/137	1/133	(0.493		
thromboembo	lism		0/137	0/133		1		
Cosmetic satis	faction at 1 year		EVLA higher satisfaction (p=0.0	034) but no data given.				
Overall satisfa	ction at 1 year		No difference between groups	for overall satisfaction (NS	5) but no data	given in paper		

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Carradice D, Mekako AI, Mazari FAK, Samuel N, Hatfield J, Chetter IC. Clinical and technical outcomes from a randomised clinical trial of endovenous laser ablation compared with conventional surgery for great saphenous varicose veins. British Journal of Surgery 2011; 98: 1117- 1123.	This is the same study as Carradice 2011A, with some additional outcomes. See Carradice 20011A	None	Carradice D, Mekako AI, Mazari FAK, Samuel N, Hatfield J, Chetter IC. Clinical and technical outcomes from a randomised clinical trial of endovenous laser ablation compared with conventional surgery for great saphenous varicose veins. British Journal of Surgery 2011; 98: 1117-1123.	This is the same study as Carradice 2011A, with some additional outcomes. See Carradice 20011A	None	Carradice D, Mekako Al, Mazari FAK, Samuel N, Hatfield J, Chetter IC. Clinical and technical outcomes from a randomise d clinical trial of endoveno us laser ablation compared with convention al surgery for great saphenous varicose veins. British Journal of Surgery 2011; 98: 1117-	This is the same study as Carradice 2011A, with some additional outcomes. See Carradice 20011A	None

Table 67: Carradice2011A⁴⁸

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
						1123.		
Results								
				Surgery	EVLA	р		
	-		lly remove the GSV and all groin SFJ (EVLA)] – data collected within 6	10/132	1/137	0.005		
(Surgery) or re), or the develo	development of neovascularisation opment of new segments of new segments of ors.	23/113	5/124	0.001		

Reference	Study type	No. of patients	Patient cha	racteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of fundin g
Darwood RJ, Theivacumar N, Dellagrammatica s D, Mavor AID, Gough MJ. Randomised clinical trial comparing endovenous laser ablation with surgery for the treatment of primary great saphenous varicose veins. British Journal of Surgery 2008; 95: 294-301	RCT, UK. Allocation concealment likely through the use of sealed envelopes. Randomisatio n method unclear. No ITT used. Drop out: EVLA1:3/49 legs EVLA2: 9/42 legs SURGERY: 11/45 legs Reasons: unclear	136 legs from 118 patients – if bilateral symptoms, both legs were used in the study, and each leg was given the same treatment. This will have artificially reduced variance within groups, but as data were analysed non- parametricall y this will not have mattered.	varicose vei saphenofem confirmed v Exclusion: tu unsuitable f large incom saphenous v morbidity p anaesthesia Baseline cha IQR shown. sample size (very sensib	ns and priman noral incompe- vith duplex US aking warfarin for EVLA (twis petent anteri- vein) or surge rohibiting ger). aracteristics: No statistical calculation ta- ile as lack of p -confidence i	And the standard stan	Saphenofemora I ligation, GSV stripping to knee level with perforation invagination stripper, and multiple phlebectomies of varicosities as a day case under GA. Prophylactic heparin given pre-surgery. LA applied to the wound area but not the GSV tract. Non stretch bandaging used for 1 day, followed by full length grade II compression stocking for 2 weeks.	Endovenous laser ablation (EVLA) using a 810nm diode laser source. The GSV was cannulated near the knee under US- guidance and LA applied to the vein. A 600 micron laser fibre inserted and drawn back through the vein. Two different laser protocols were used: EVLA1: 12W power with 1 sec pulses and 1 sec intervals. Catheter withdrawn 2-3 mm on each	1, 6, 12 and 52 weeks post treatment	Disease specific QoL [Aberdeen VV symptom score (AVVSS)] Abolition of reflux with Duplex imaging Patient symptoms and cosmesis Patient satisfactio n	not state

Reference Stud	Study type	No. of patients	Patient characteristics I			Intervention	Comparison	Length of follow-up	Outcome measures	Source of fundin g
up (thos droj wer up): EVL legs EVL legs SUR 2/34 Rea	opping out ere followed 1): 7LA1:5/46 gs 7LA2: 4/33		CEAP5 CEAP unknown VCSS bilateral AVVSS	3/47 1/47 4 (3-5) 9/47 11.76 (9.8- 19.4)	0/33 2/33 4 (3-5) 6/33 14.3 (8.9- 19.6)		rest interval. EVLA2: 14W power continuously with constant 2-3 mm/sec withdrawal. Non stretch compression bandage applied postoperativel y for 1 week, followed by a grade II compression stocking for a further week. Injection sclerotherapy would be performed at 6 weeks for residual varicosities if requested by the patient. This partially controls for the		Physician outcomes – venous clinical severity score (VCSS) Adverse events, including post- procedure pain.	

Reference	Study type	No. of patients		Patient characterist	ics	Intervention		Comparison	Length of follow-up	Outcome measures	Source of fundin g
								phlebectomies given to the surgery group, but there is still a possibility that the differences between sclerotherapy and phlebectomies would have confounded results.			
Results: Median (I	QR)										
Outcome			EVLA	1	EVLA2		Sur	gery	р		
QoL – AVVSS 3mor 100 worst symptor	•	100 <i>,</i> with	5.6 (1.45-8.2); n=34	4.2 (1.7-7.9); r	1=20	5.3	2 (1.0-7.7); n=26	not s	tated	
QoL – AVVSS 12mc	onths		1.8 (0.1-5.9); n=22	2.5 (0-5.6); n=	15	3.9	(0-10.3); n=9	not s	tated	
QoL – AVVSS impro 3months	ovement from bas	eline at	9.4 (4	4.5-14.9)	10.3(5.0-15.0)		8.4	(4.5-13.2)	0.694	l	
Abolition of GSV re	eflux at 12 weeks		41/4	2	26/29		28/	/32	0.22	7	
Abolition of GSV re only those with ab			24/2	8	19/21		11/	/12			
Abolition of SFJ ref	lux at 12 weeks		39/4	2	27/29		32/	/32	0.30	7	
Abolition of SFJ ref only those with ab			23/2	8	19/21		11/	/12			
VCSS			0 (0-	1)	0 (0-1)		0 (0	D-1)	NS		

Reference	Study type	No. of patients		Patient characteristics		Intervention		Comparison	Lengt follov		Outcome measures	Source of fundin g
Patient satisfaction 100 = high satisfact)mm VAS;	95 (8	31-95)	91 (84-97)		91	(81-95)		0.267		
Cosmesis at 12 wee high satisfaction)	eks (100mm VAS;	100 =	92 (8	30-95)	92 (77-95)		93	(76-98)		0.980		
Adverse events			EVLA	\1	EVLA2		Sur	gery				
Post-operative pair medians	n (mean of the firs	st 7 days	11		18		14					
symptomatic phleb	oitis		6/42		3/29		0					
paraesthesiae/ nur	nbness		0		1/29		4/3	2				
pruritis at cannulat	ion site		0		1/29		0					
upper thigh discolo	ouration/bruising		0		1/29		2/3	2				
wound infections r	equiring antibioti	cs	0		0		2/3	2				
ARDS, leading to 7	days ventilation i	n ICU	0		0		1/3	2				
Treatment failure			see a	abolition of reflux results								
return to work in o	ne week		29/3	4	20/24		14/	25				

Reference	Study type	No. of patients	Patient charact	teristics		Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
ElKaffas KH, ElKashef O, ElBaz W. Great Saphenous Vein Radiofrequency Ablation versus standard dtripping in the management of primary varicose veins – a randomised clinical trial. Angiology 2011; 62: 49-54	RCT. Randomisation method was where patients "blindly choose an assignment card that would put them in either group": this has clear scope for unconscious bias from the researcher. For example, the card relating to the preferred allocation could be placed in a certain position aimed at encouraging its choice. Allocation	180 patients. 90 in EVRF group and 90 in surgery group. Drop out data is unclear, as 90 are reported to have had treatment in the EVRF group, but the drop-out data for that group are reported with a denominator of 88. Drop- outs over 24 months were 7 for the EVRF group and 9 for the surgery group. No ITT	Inclusion: SFJ a duplex, either of compression/re manoeuvres. Exclusion: Patie superficial veno patients on ant pacemakers, se disease, pregna diameter > 18 m twisted veins. Baseline charae group older and diameter. All el Age M:F discomfort oedema skin changes	on standing elease or Va ents with de bus thromb cicoagulants erious syste ancy. Also C nm in the th cteristics: s d with large	r manual alsalver eep or osis; 5; PAD, mic GSV lumen high or very urgery er GSV	SF high ligation and GSV stripping at ankle (n=40) or knee (n=50). GA used for all, all were managed as inpatients. Stripping performed after wrapping elastic bandage to reduce postoperative haematoma, with operating table tilted 30 deg foot up. Common procedures: Both groups had stab phlebectomies as needed (15/90 in EVRF gp and 39./90 in	Radiofrequency ablation (EVRF) used. Closure system (VNUS medical technologies) used. two operators involved. Vein cannulated at point of most distal reflux and tip of RF catheter placed at least 2cm distal to the saphenofemoral junction or just distal to the superficial epigastric vein orifice. LA used only. RF catheter used temperatures of 80-85 deg. One operator only.	1 week, 1 month, 6 months, 12 months 18 months, 24 months. Mean FOLLOW- UP 20.9 (6.8) months.	Treatment failure Adverse events Recurrence	None

Table 69: Elkaffas 2011¹¹³

Reference	Study type	No. of patients	Patient charact	eristics		Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
	concealment is inherent in the design, as the allocation cannot be known by the researcher until it is made by the patient; however the problems with unconscious bias remain.	(imputation) was reported. This would only be relevant to the Kaplan Meir analysis, which will hopefully have censored the	C3 C4 C5 GSV diameter (mm) [mean(range)] op duration (mins)	27 9 3 7.8 (4.5- 12) 40 (12)	27 12 6 8.6 (4- 14) 45 (13)	surgery group).	24/90 also needed later postoperative sclerotherapy for small veins was needed (none in surgery group).			
	No evidence of blinding.	data.								

Results:

	EVLA	Surgery	р
Treatment failure at completion of intervention (no full occlusion or failure to remove the GSV)	6/90	0/90	
Adverse events (imm. Post op)			
focal paresthesia	9/90	3/90	
Thrombophlebitis	6/90	0/90	
severe pain requiring analgesics	12/90	12/90	
hematoma formation	1/90	12/90	
iliofemoral DVT	0/90	1/90	
severe groin infection	0/90	3/90	

Reference	Study type	No. of patients	Patient characteristics		Intervention	Comparis	son	Length of follow- up	Outcome measures	Source of funding
pulmonary embo	olism			0/90			0/90			
Recurrence over	24 months			12/90			9/90			
Average (?media given)	n) time to recurre	ier graph provided but no data	23.3 m	onths (95%CI: 22.5-	24.1)	23.0 m 21.3-24	onths (95%Cl 1.6)	: 0.4		
return to normal	activity			3 (3)			7 (2.6)			

Reference	Study type	No. of patients	Patient chara	octeristics		Intervention	Comparison	Lengt h of follow -up	Outcome measure s	Source of fundin g
Flessenkamper I, Hartmann M, Stenger D, Roll S. Endovenous laser ablation with and without high ligation compared with high ligation and stripping in the treatment of great saphenous varicose veins: initial results of a multicentre randomised controlled trial. Phlebology 2012; DOI: 10.1258/phleb.2011.01114 7	RCT. Three arm RCT, but one arm not included in this review (EVLA with high ligation) as it is not "standard" treatment. Non computer randomisatio n but clear allocation concealment. No assessor or other blinding.	301 randomised to the 2 arms under consideration . No attrition at 2 months follow-up	Inclusion: 18- symptoms of insufficiency of GSV; life expec- Exclusion: pro- All had insuffi- diameter of t the junction of Baseline char comparable: Baseline char comparable: Comparable: Comparable: Comparable: Comparable: Co	superficial ve with proven r ectancy > 5 ye evious surger icient termina he GSV 5cm c was <u><</u> 16mm.	nous eflux into ars; y of the GSV. Il valve, listal from	High ligation of the SFJ and stripping using invagination technique. Open surgery. Anaesthetic unclear. Miniophlebe ctomies carried out as required	980nm laser (EVLA) at 30W in continuous mode. Performed under duplex guidance. Tumescent anaesthesia. Miniophlebe ctomies carried out as required.	2 month s	AEs GSV reflux	Germa n society of phlebo ogy

Evidence tables clinical studies

Table 70: Flessenkamper 2012⁹⁹

Results:

Reference	Study type	No. pati	of ents	Patient characteristics		Intervention	Comparison	Lengt h of follow -up	Outcome measure s	Source of fundin g
			Stripping		EVLA		р			
Adverse Events										
DVT			1/159		1/142					
Lymphoedema			4/159		2/142					
Neurological sensory deficit	S		2/159		3/142					
Ecchymosis			100/159		72/142					
Post op pain day 1			6/159		4/142					
Post op pain 2-5 days			23/159		20/142					
Post op pain >5 days			50/159		57/142					
Saphenous nerve damage			1/159		5/142					
SFJ reflux at 2 months			0/159		38/142		P<0.0002	L		
Work disability (days)			9.2		9.3					
VDS - asymptomatic			77/159		84/142					

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Hinchliffe RJ, Ubhi J, Beech A, Ellison J, Brathwaite BD. A prospective randomised controlled trial of VNUS Closure versus surgery for the treatment of recurrent long saphenous varicose veins. Eur J Vasc Endovasc Surg. 2006; 31: 212-218.	Randomised within-subject design. The researchers used a non-specified random method to decide which side was treated with what. Allocation concealment not applicable, as the decision on the side would be made after the recruitment. Patients reported to be blinded as to the treatments used on each side, assisted by the use of opaque dressings over the groin. Assessor photographing patients' legs for assessment of bruising was blinded to	16 patients. No reports of loss to follow- up.	Inclusion: recurrent varicose veins previously treated by sapheno-femoral ligation; all CEAP 2 and above; persistent and incompetent GSV suitable for treatment with EVRF; >18 yrs; Exclusion: pregnancy, twisted GSV, GSV <3mm, > 12mm; thrombotic scarring of GSV; no GSV present. Baseline Characteristics: 12 women, 4 men; median age 54 (44-66 yrs); median CEAP: 3 (class 2, n=1, class 2, n=14, class 4, n=1).	Ligation and inversion stripping to just below the knee. to just below the knee. Common procedures: Multiple stab avulsions, general anaesthetic, compression bandages for 2 weeks.	EVRF with VNUS closure system. The probe was retracted in 1cm increments.	6 weeks, 1 year		None

Table 71:Hinchcliffe 2006114

Reference	Study type	No. of patients		ent characteristics		Intervention	Со	mparison	Length of follow-up	Outcome measures	Source of funding
	patient group.										
Results: No va	iriances given for co	ntinuous v	variables	i.							
		S	Surgery		EVLA			р			
Post Ix pain (ti median (IQR)	me point not given)	[VAS]; 3	3.8 (0.6-6	5.3)	1.7 (0.	2-4)		0.02			
	ost Ix bruising (time point not given) /AS]; median (IQR)		5.2 (2.6-7.0)		1.7 (0.4	1.7 (0.4-4.4)		0.03			
-	of bruising covering legs (time point ot given); median (IQR)		21.8 (15.	7-28.5)	11.9 (8	8.9-18.3)		0.02			
Adverse event	S										
DVT		C	0/16		0/16						
vessel perfor	ration	C	0/16		0/16						
PE		C	0/16		0/16						
skin burns		C	0/16		0/16						
lymphatic lea	ak	C	0/16		0/16	0/16					
post operativ	ve neuralgia	C	0/16		2/16						
thigh discom	nfort	C	0/16		2/16						
wound infect	tion	1	1/16		0/16						
numbness		Э	3/16		0/16						
thrombophle	ebitis	1	1/16		0/16						
leg oedema		1	1/16		0/16						
non-fully strip weeks	ped / non- occlusion	6 2	2/16		3/16						
persistent inco truncal veins 6	ompetence in accesso weeks	ory 2	2/16		3/16						

Reference	Study type	No. of patients	Patient characte	ristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding										
Lurie F, Creton D, Eklof B, Kabnick LS, Sites in USA, Kistner RL, Pichot O, 	group and 40 to the surgery group. 3 patients immediate ly withdrew	Symptomatic var incompetence, co Inclusion: Revers standing; age 21- classification; am vein < 1.2 cm in s reflux permissible Exclusion: vein d cm. Duplication co incompetent acco branch; small SV prev. DVT; ABI<0 twisted GSV segre Baseline characte	onfirmed with e flow lasting 80 yrs; C2-4 bulatory; sap supine; segme e. iameter >1.2 of saphenous essory sapher reflux; thigh .9; axial DV re nent to be tre	a duplex. a .0.5 secs in henous ental deep cm or <0.2 trunk or hous varices; iflux;	Physicians followed their standard practice using either an olive- tipped device or a PIN stripper. Ligation in the femoral triangle. Common procedures: Adjunctrive procedure on varices and perforator vessels limited to below-knee.	EV radio- frequency obliteration. The Closure catheter and system (VNUS med technologies) was used according to established methods.	72 hours, 1 wk, 3 wks, and 4 months.	Disease specific QoL Occlusion and reflex rates patient reported symptoms Adverse events	VNUS medical technologie s, inc. Clean conflict of interest.											
stripping in a selected	on unclear: "via	surgery	surgery	surgery		surgery	surgery	surgery	surgery	surgery	surgery	surgery	surgery group. A	surgery	surgery EVRE surgery					
patient population	internet". Allocation	further	Age	49 (4)	47 (4)															
(EVOLVeS study). J Vasc	VOLVeS concealment did not	VCSS	4.8 (0.34)	4.39 (0.38)																
Surg 2003; 38: 207-14 Blinding unclear. Bilat treatment of	ding lear. Bilat tment of	Female	32 (74.4)	26 (72.2)																
		Marking 25/44 25/26																		
	one patient in the EVRF		CEAP2	36/44	28/36															

Table 72: Lurie2003¹⁵²

Reference	Study type	No. of patients	Patient character	istics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
	group, thus artificially	EVRF patient	CEAP4	4/44	4/36					
	decreasing variance in	was withdrawn	GA used	12/44	19/44					
	that group.	after discovery of previous surgery in the same vein. One further EVRF patient	Adjunctive phlebectomies done	42/44	36/36					
			mean avulsions/extre mity	8.6 (2.6)	9.8 (2.8)					
			mean length of 37 (2) 40 (2) treated segment							
		was withdrawn	site 1	20	14					
		from follow-up	site 2	6	9					
		after he was found	site 3	7	7					
		to have C6, a protocol	site 4	9	3					
			site 5	2	3					

Results: Comparisons of QoL scores were adjusted for the type of anaesthesia and number of adjunctive procedures.

	EVRF	Surgery
CIVIQ2 Disease specific QoL (0-100, 100 worst) global score (mean (<u>se</u>)). Change from baseline to 72 hrs.	-3 (2.7)	13.3 (3.1)
CIVIQ2 Disease specific QoL (0-100, 100 worst) global score (mean (<u>se</u>)). Change from baseline to 1 wk	-9.2 (2.3)	3.7 (2.5)
CIVIQ2 Disease specific QoL (0-100, 100 worst) pain dimension	-1.77 (0.6)	2.9 (0.7)

Reference	Study type	No. of patients	Patient characteristic	S	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
(mean (<u>se</u>)). C	hange from base									
	e specific QoL (0 hange from base) pain dimension	-2.4 (0.6)			1.2 (0.7)			
	e specific QoL (C ean (<u>se</u>)). Chang			0.82 (0.69)			4.85 (0.79)			
	e specific QoL (C ean (<u>se</u>)). Chang			-0.97 (0.65)			2.02 (0.72)			
NB: The other	two dimension	s of the CIVIQ2	are not reported exce	ot the fact that they	vwere NS. Data at la	ater follow-ups n	ot reported!			
Occlusion and	l reflux rates									
reflux at 72 l	hrs			5/43			0/36			
reflux at 1 w	eek			unclear		0/36				
complete occ	clusion of GSV at	72 hours		36/43		0/36				
complete occ	clusion of GSV at	1 wk		41/43		0/36				
complete occ	clusion of GSV at	t 3 wks		41/43		0/36				
complete occ	clusion of GSV at	4 months		42/43			0/36			
patient report	ed symptoms				y detail, except that y but no reports of a		a clear advantage for "pain". The VCSS is t at baseline.			
Adverse even	ts at 72 hrs									
none				19/44			6/36			
infection				0/44			2/36			
superficial ve	enous thrombos	is		0/44			1/36			
tenderness				2/44			9/36			
lymphocele				0/44			0/36			
bleeding from	m stab wound			3/44			3/36			
eccymosis				12/44			19/36			
erythema				6/44			3/36			
hematoma				7/44			14/36			

Reference	Study type	No. of patients	Patient characteristic	S	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
paresthesia				5/44			2/36		
hyper pigme	ntation			0/44			0/36		
Adverse event	ts at 1 week								
none				15/44			5/36		
infection				0/44			1/36		
superficial ve	enous thrombosis	5		1/44			2/36		
tenderness				5/44			10/36		
lymphocele				0/44			1/36		
bleeding from	m stab wound			0/44			0/36		
eccymosis				14/44			23/36		
erythema				2/44			1/36		
hematoma				6/44			18/36		
paresthesia				10/44			5/36		
hyper pigme	ntation			0/44			0/36		
Adverse event	s at 3 weeks								
none				31/44			14/36		
infection				0/44			1/36		
	enous thrombosis	5		2/44			1/36		
tenderness				4/44			9/36		
lymphocele				0/44			0/36		
-	m stab wound			0/44			0/36		
eccymosis				1/44			7/36		
erythema				1/44			3/36		
hematoma				1/44			12/36		
paresthesia				7/44			2/36		
hyper pigme	ntation			1/44			0/36		

Reference	Study type	No. of patients	Patient characteristic	S	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Adverse events	s at 4 months								
none				36/43			26/34		
ecchymosis				0/43			1/34		
erythema				0/43			2/34		
hematoma				0/43			3/34		
Treatment fail	ure			2/44			0/36		
return to norm	nal activity			1.15 (1.5)			3.89 (1.5)		
	·							lculated from t	the p value

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Lurie F, Creton D, Eklof B, Kabnick LS, Kistner RL, Pichot O, Sessa C and Schuller- Petrovic S. Prospective randomised study of endovenous radiofrequen cy obliteration (closure) versus ligation and vein stripping (EVOLVeS): Two-year follow-up. Eur J Vasc Surg 2005; 29, 67-73	RCT. Follow up to Lurie 2003. Please see table for Lurie 2003.	As Lurie 2003. This study involved follow-up at 2 further time points: 1 year and 2 years. At one year, data were missing for 21 limbs in the EVLA group and 20 limbs from the surgery group. At 2 years, data were missing for 10 limbs in the EVLA group and 11 limbs from the surgery	See Lurie 2003. In addition CEAP scores at baseline were reported as equivalent, with 82% and 78% at C2 in EVLA and surgery respectively, 9% and 11% at C3 and 9% and 11% at C4 (chi sq. 0.2, P=0.9).	see Lurie 2003	see Lurie 2003	As Lurie 2003, plus 1 year and 2 years.	Some outcomes missing from the Lurie 2003 paper are included in this, as well as new 1 yr and 2 year data. QOL Reflux rates Recurrence rates VCSS	not stated

Table 73: Lurie2005¹⁵³

Reference	Study type	No. of patien group.	ts	Patie	nt characteris	tics	Inter	vention		Comparison	Lengt follow		Outcome measures		Source of funding
Results:															
			EVRF		Surgery										
reflux at 4 mo	nths		4/43												
reflux at 2 yea	rs		2/36		3/29										
	tes of recurrent v ined 1 and 2 year		14.3%		20.9%	long ra test NS									
CIVIQ2 Disease year and 2 year	e specific global C ars	QoL at 1		tion in	-			-		nbiguity about v group had a sigr					
CIVIQ2 Diseas of QoL at all ti	e specific pain din me points	nension	Howev	ver rep	-	e EVLA g	roup h	-) variance indica v higher improve				•	•
VCSS			extract	tion in	npossible). Ho	wever re	eporte	d that the	EVLA g	nbiguity about v group had a sigr ficantly differen	nificant	ly lowe	r (better) VCS		
			EVRF							Surgery					
						%						%			
	t 4 mths (analysed vith a chi square	d	C0			34.1				CO		41.2			

Reference	Study type	No. of patients	Patient characterist	tics Int	ervention	Comparison	Length of follow-up		Source of funding
		C1		38.6		C1	32.	4	
		C2		18.2		C2	20.	6	
		C3		0		C3	2.9		
		C4		9.1		C4	2.9		
				%			%		
	at 1 yr (analysed with a chi square	CO		28		CO	15		
		C1		56		C1	40		
		C2		8		C2	30		
		C3		0		C3	5		
		C4		8		C4	10		
				%			%		
	at 2 yrs (analysed with a chi square	CO		33.3		CO	27.	6	
		C1		41.7		C1	31		
		C2		22.2		C2	31		
		C3		0		C3	3.4		
		C4		2.8		C4	6.9		
Treatment fai	ilure	See L	urie 2003						

Reference	Study type	No. of patients	Patient characteristic	s	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Perala J, Rautio T, Biancari F, Ohtonen P, Wiik H, Heikkinen T, Juvonen T. Radiofrequency endovenous obliteration versus stripping of the long saphenous vein in the management of primary varicose veins: 3 year outcome of a randomised study. Ann Vasc Surg 2005; 19: 669-672.	3 year follow- up of Rautio 2002.	see Rautio 2002. No further drop out from 50 days to 3 years.	See Rautio 2002		Stripping surgery. Groin dissected to expose the SFJ. Side branches of the GSV at the SFJ were divided and ligated. After local phlebectomy, the GSV was stripped from just below the knee to the groin with the venostrip with a 9mm olive.	Radiofrequency endovenous obliteration carried out with the VNUS Closure system (see Lurie studies).	3 years.	Decrease in VCSS VDS at 3 years VSDS at 3 years Satisfaction with cosmetic result. Recurrence Treatment failure	not stated
Results:									
		EVRF		Surgery		р			
Average decrease in baseline to 3 years				4.0 (1.2)		0.7			
VDS at 3 years (med	dian (range))	an (range)) 0 in all except for 1		0 in all e	xcept for 1	1			
VSDS at 3 years		0 in all except for 3		0 in all except for 1		0.6			
Lack of satisfaction result at 3 years	of satisfaction with cosmetic 1/15 at 3 years			2/13					
Would not recomm friend	end to a	0/15		0/13					

Table 74: Perala2005 204

Reference	Study type	No. of patients	Patient characteristics	s	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Recurrence of varico determined by surge		5/15		2/13		0.4??			
Recurrence of varico determined by patient		4/15		2/13		0.065??			
Reoperation for recu varicose veins	ırrent	1/15		1/13					
Adverse events symptom relating t nerve injury superficial thrombo		1/15 1/15		5/15 0/15					
Treatment failure		see Rautio 2002							
Reflux		3/15		0/15					

Reference	Study type	No. of patients	Patient cha	aracteristics		Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Pronk P, Gauw SA, Mooij MC, Gaastra MTW, Lawson JA, van Goethem AR, van Vlijmen-van Keulen CJ. Randomised controlled trial comparing saphenofemoral ligation and stripping of the great saphenous vein with endovenous laser ablation (980nm) using local tumescent anaesthesia: one year results. Eur J Endovasc Surg 2010; 40: 649- 656.	RCT. Non blinded. Computer randomisation used. No reporting of allocation concealment.	130 legs in 122 patients (EVLA n=62; surgery n=68). Patients with bilateral VV were randomised only once. All had treatment. At 6 weeks, 2 lost to follow- up from surgery group, none from the EVLA group. The 2 lost to follow-up did complete their post-op questionnaires. There was further loss to follow-up at 1 year (7 further lost in surgery group and 6 further lost in the EVLA group)	veins. Inclusion: 3 >0.5 secs o between 0 Exclusion: GSV; intraf <15cm mea downward intolerance superficial VT; deep ve	>18 years; CE n duplex; GS .3 and 1.5cm Previous surg ascial GSV re asured from s s; pregnancy e of lidocaine	gery of the flux length SFJ ; immobility; ; active evious or deep ciency.	High ligation of GSV and ligation of all tributaries via groin incision, followed by PIN stripping, with access via incision below the knee. Common procedures: Tumescent anaesthesia given to all. Sclerotherapy of superficial varicose veins also given to all. Short stretch bandages applied to the whole leg for 1 week.	EVLA. Proximal 10cm of the incompetent GSV treated with an energy does of 100 J/min followed by a targeted energy dose determined by the diameter of the GSV (0.3-0.4 – 50 J/cm up to >0.6 cm – 80 J/cm).	1 week, 6 weeks, 6 months, 12 months.	Post op pain Symptoms CEAP score Patient satisfaction	None

Table 75: Brenk 2010213

Reference	Study type	No. of patients	Patient ch	aracteristic	5	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
			itching	26/68	20/62					
			cosmetic	13/68	13/62					
			pain	13/68	9/62					
			restless legs	6/68	11/62					
			calf cramps	8/68	8/62					
			CEAP 2	26/68	29/62					
			CEAP3	36/68	29/62					
			CEAP4	5/68	4/62					
			CEAP5	1/68	0/62					
Results:										
		Surgery			EVLA		Ρ			
Post-op pain du	ring op	3.39(2.57)		2.21(2.40)					
Post-op pain da	/1	4.00(2.34	.)		3.58(2.600					
Post-op pain da	/2	3.12(2.38)		3.05(2.48)					
Post-op pain da	/3	2.38(2.11	.)		2.76(2.53)					
Post-op pain da	7	1.78(1.94	.)		3.74(2.72)					
Post-op pain da	/10	1.18(1.49)		2.65(2.21)					

Reference Study	type	No. of	patients	Patient characteristic	S	Intervention	Com	parison	Length of follow- up	Outcome measures	Source of funding
Post-op pain day14			0.77(1.46)	1.66(2.04)						
Tired legs 1 year			8/62		5/56			0.49			
oedema 1 year			10/62		6/56			0.39			
itching 1 year			6/62		3/56			0.50			
cosmetic 1 year			8/62		4/56			0.31			
pain 1 year			6/62		1/56			0.12			
restless legs 1 year	stless legs 1 year		4/62		7/56			0.43			
calf cramps 1 year			2/62		5/56			0.25			
CEAP 0 1 year			21/61		19/56			0.96			
CEAP 1 1 year			22/61		20/56			0.97			
CEAP 2 1 year			11/61		9/56			0.78			
CEAP 3 1 year			6/61		7/56			0.65			
CEAP 4 1 year			0/61		1/56						
CEAP 5 1 year			1/61		0/56						
QoL – cosmetic concerns (months [10 best]	(VAS) at 6		7 (4-10)		7.5 (3-10)						
Willing to do the same pro	ocedure ag	gain	53/67		47/61						

Reference St	tudy type	No. o	fpatients	Patient characteristic	S	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Would recommend to relative	to a friend or		59/67		51/61					
Adverse events										
wound infection			0/67		0/61					
DVT			0/67		0/61					
post op bleeding			2/67		0/61					
Thrombus at SFJ			0/67		3/61					
paresthesia			1/67		2/61					
persistent neurolog	gical injury (1 ye	ear)	1/67		0/61					
Recurrence at 1 year	r (presence of re	eflux)	5/56		5/49					
Recanalisation of GSN	Recanalisation of GSV				3/49					
return to normal acti	ivities		3.2 (4)		3.2 (4.3)					

Reference	Study type	No. of patients	Patient characteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Rasmussen LH, Bjoern L, Lawaetz M, Blemings A, Lawaetz B, Ekolof B. Randomised trial comparing endovenous laser ablation of the great saphenous vein with high ligation and stripping in patients with varicose veins: short term results. J Vasc Surgery 2007; 46: 308-15	RCT. Randomisatio n method unclear, but blocks of 10 were used. Allocation concealment achieved through sealed envelopes. Two experienced surgeons involved, but unclear whether these did both or just one kind of intervention. Two centres also involved, but unclear how these were distributed across groups. Bilateral treatment allowed, with	121 patients (137 legs). ITT reported as used, but unclear about how the loss to follow-up were managed. No evidence of data imputation. Loss to follow-up (cumulative):	Inclusion: varicose veins, CEAP C2-4 EpAsPr. Aged 18-80 yrs. GSV incompetence defined by reflux >0.5 sec on duplex. Previous high ligation was permitted. Exclusion: Duplication of the saphenous trunk, incompetent anterior accessory GSV, previous DVT, arterial insufficiency and/or ankle-brachial index <0.9, axial deep venous insufficiency and twisted GSV. Baseline characteristics: (n represents number of legs). For baseline values of outcomes, variance given is range. Authors stated no group differences. But note the numerical difference in C4 disease.	EVL done under duplex guidance with a 980mm diode laser, with pulse mode, 1.5 sec impulse, 1.5 sec pause and 12W. GSV accessed percutaneously. Catheter advanced until 1- 2 cm below saphenofemoral junction. Common procedures: Tumescent anaesthesia administered with a syringe with US guidance to the GSV. Post surgery, compressive bandaging applied, and then replaced with a class 1 compressive stocking left in situ for 2 weeks	High ligation and perforate invagination stripping performed through a groin incision of 4-6 cm, with flush division of the GSV and division of all tributaries behind the second level of the division. If the vein broke then attempts were made to remove it from a more distal position below the knee. Common procedures: See intervention column.	12 d, 1,3 and 6 months.	Physician outcomes - VVSS QOL - disease specific AVVSS QOL - general SF-36 Adverse events Treatment failure	Public health insurance research foundatic of Denmark

Table 76: Rasmussen2007²¹⁹

Reference	Study type	No.	of pati	ents	Patient ch	aracteristic	S	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
Reference	Study type the same treatment given to both legs (risk of artificial reduction in variance). No blinding achieved.	12 d 1 mo 3 mo 6	ef pati EVL A 2 4 6 15		Patient ch prev high lig M:F CEAP 2 CEAP4	aracteristic Surgery 8/59 16:43 51/59 3/59	EVLA 8/62 21:41 50/62 9/62	Intervention (not entirely clear this was used for EVL patients, however). Both groups also had multiple phlebectomies as needed. Diclofenac given as pain relief.	Comparison				
	mo	mo	GSV diam (mm) reflux	7.6 (2.1)	7.9 (2.7) 2.6 (1.1)								
						time (s) AVVSS	16.1 (4.4- 34.3)	18.6 (3.6- 40.2)					
						VCSS	VCSS 2.4 (2- 12)	2.8 (1-8)					
					SF36 – physical function	89.3 (25-100)	87 (25- 100)						
					SF36 – role- physical	89.3 (25-100)	87 (25- 100)						
					SF36 – bodily pain	77.1 (22-100)	76.6 (22- 100)						

Reference	Study type	No. of patients	Patient c	haracteristi	cs	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding		
			SF36 – gen health	67.6 (28-80)	65.2 (32- 80)							
			SF36 – vitality	73.1 (12.5- 100)	69 (12.5- 100)							
Results: For a	continuous variat	oles, only mean and ra	nges given.									
All outcomes	s measured at 3 r	months	Strip	ping surgery	Endove	enous laser ablatio	n					
AVVSS 3 (sco	re out of 100, wit	th 100 worst symptom	s) 8.2 (0)-31.2)	6.9 (0-4	13.8)						
VCSS			0.2 (0)-2)	0.1 (0-2	2)						
SF36 – physic	cal function		92.2	92.2 (43.7-100)		93.9 (56.2-100)						
SF36 – role-p	hysical		92.2	92.2 (43.7-100)		93.9 (56.2-100)						
SF36 – bodily	6 – bodily pain		89.5	(31-100)	89.1 (3	2-100)						
SF36 – gen h	ealth		66.7	66.7 (20-80)		2-80)						
SF36 – vitalit	У		79 (3	79 (37.5-100)		8.7-100)						
SF36 – social	functioning		97.1	(12.5-100)	94.5 (3	7.5-100)						
SF-36- role e	motional		95.8	(58.3-100)	94.4 (3	3.3-100)						
SF-36 – ment	tal health		89.2	(60-100)	84.3 (2	5-100)						
All outcomes	s measured at 6 r	nonths	Strip	oing surgery	Endove	enous laser ablatio	n					
AVVSS (score	e out of 100, with	100 worst symptoms)	5.3 (0-33.1)	7.1 (0-3	38.7)						
VCSS			0.2 (0)-2)	0.4 (0-7	7)						
SF36 – physic	cal function		92.6	(50-100)	93.9 (4	3.7-100)						
SF36 – role-p	hysical		92.6	950-100)	93.9 (4	3.7-100)						
SF36 – bodily	ı pain		86.5	(20-100)	90.9 (5	1-100)						
SF36 – gener	al health		67 (3	3.6-80)	67.9 (4	0-80)						
SF36 – vitalit	У		82.9	(56.2-100)	77 (18.	7-100)						
SF36 – social	functioning		98.8	(62.5-100)	98.2 (6	2.5-100)						

Reference	Study type	No. of patients	Patient characteristics	1	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
SF-36- role e	motional		95.7 (50-100)	95 (58.3-:	100)				
SF-36 – ment	al health		90.2 (70-100)	86.2 (40-3	100)				
Adverse eve	nts		Stripping surgery	Endoven	Endovenous laser ablation				
major compl	ication – infection	n of groin (12days)	1/68	0/67					
phlebitis (12	days)		2/68	2/67					
phlebitis (1m	ionth)		2/66	2/65					
bruising (12d	ays)		15/68	7/67					
hematoma (1	L2days)		5/68	3/67					
hematoma (1	Lmonth)		1/66	0/65					
parasthsia (1	month)		0/66	1/65					
paraesthesia	e (6months)		1/50	0/54					
NOT stripped	l /occluded 12 day	ys	0/67	2/68					
NOT stripped	l/occluded 1 mon	th	0/65	2/66					
NOT stripped	l/occluded 3 mon	1/63	0/63						
NOT stripped	l/occluded 6 mon	3/53	1/50						
time to resur	ne normal activity	y (days)	7.7 (6.1)	6.9 (7)					
Author's con	clusions: The trea	atments were equally	safe and efficient at elimir	nating GSV r	eflux.				

Rasmussen LH, Bjoern L, Lawaetz M, Lawaetz B, Blemings A, Eklof B. Randomised clinical trial comparing endovenous laser ablation withRCT. Follow up to Rasmussen 2007Follow up to Rasmussen 2007Follow up to Rasmussen 2007Follow up to Rasmussen 2007Follow up to Rasmussen comparing endovenous comparing endovenousFollow up to Rasmussen 2007Follow up to Rasmussen 2007Follow up to Rasmussen comparing endovenousClinical severity scorePublic Resea Resea research Found of	Table 77: Rasmussenzoit	,							
Lawaetz M, Lawaetz B, Blemings A, Eklof B. Rasmussen Randomised clinical trial comparing endovenous laser ablation with stripping of the great saphenous vein: clinical outcome and recurrence after 2 years. Eur J Vasc Surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040yp to Rasmussen 2007Rasmussen 2007Rasmussen Rasmussen 2007Clinical Resear Found of Denmi Stripping of the great saphenous vein: clinical outcome and recurrence after 2 years. Eur J Vasc Surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040yp to Rasmussen RasmussenRasmussen 2007 Rasmussen Rasmussen 2007Rasmussen 2007 Rasmussen 2007Rasmussen Stripping of the great saphenous vein: clinical outcome and recurrence after 2 years. Eur J Vasc Surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040Point the great subscript of the great saphenous vein: clinical saphenous vein: clinical outcome and recurrence after 2 years. Eur J Vasc Surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040Point the great subscript of the great saphenous vein: clinical surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040Point the great subscript of the great surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040Point the great subscript of the great surg 2010: in press. Doi: 10.1016/j.ejvs.2009.11.040Point the great subscript of the great <br< th=""><th>Reference</th><th>Study type</th><th></th><th>Patient characteristics</th><th>Intervention</th><th>Comparison</th><th>-</th><th></th><th>Source of funding</th></br<>	Reference	Study type		Patient characteristics	Intervention	Comparison	-		Source of funding
AVVSS, VCSS and domains of the SF36 were reported to to differ at 2 years, but no data given (except in very low resolution figures)EVLASurgerypRecurrence (legs) at 2 years [note the denominators do not reflect the drop outs that occurred earlier in the study, so ITT? But no mention of ITT or imputation].18/6925/68ns	Lawaetz M, Lawaetz B, Blemings A, Eklof B. Randomised clinical trial comparing endovenous laser ablation with stripping of the great saphenous vein: clinical outcome and recurrence after 2 years. Eur J Vasc Surg 2010: in press. Doi:	up to Rasmussen	to Rasmussen		•	Rasmussen	2 yrs	Clinical severity score	Public health Insurance Research Foundation of Denmark
EVLASurgerypRecurrence (legs) at 2 years [note the denominators do not reflect the drop outs that occurred earlier in the study, so ITT? But no mention of ITT or imputation].18/6925/68ns	Results:								
Recurrence (legs) at 2 years [note the denominators do not reflect the drop outs that occurred earlier in the study, so ITT? But no mention of ITT or imputation].	AVVSS, VCSS and domains of	f the SF36 were i	reported to no	ot differ at 2 years, but no data	given (except in very	y low resolution	figures)		
not reflect the drop outs that occurred earlier in the study, so ITT? But no mention of ITT or imputation].				EVLA	Surgery	р			
Reflux into the AAGSV 6/69 3/69	not reflect the drop outs that	ot reflect the drop outs that occurred earlier in the		18/69	25/68	ns			
	Reflux into the AAGSV			6/69	3/69				

3/69

9/69

6/69

6/69

2/69

4/69

3/69

9/69

Table 77: Rasmussen2010²²⁰

Reflux in the groin

Reflux in thigh perforators

Reflux in lower leg perforators

Retreatment (mainly for cosmetic reasons)

Table 78: Rasmussen2011²²¹

See Table 63 for evidence table.

Table 79: Rass2011²²²

Reference	Study type	No. of	Patient characteristics		Intervention	Comparison	Length of	Outcome	Source of
		patients					follow-up	measures	funding
Rass K, Frings N, Glowacki P, Hamsch C, Graber S, Vogt T, Tilgen W. Comparable effectiveness of endovenous laser ablation and high ligation with stripping of the great saphenous vein. Arch Dermatol 2011; ecopy published online	RCT. Randomisation method unclear. Allocation concealment likely through use of an independent remote centre for treatment allocation. Only one limb used per patient. If bilaterally affected the worst limb was used for the study. Each treatment given in a	400 randomised - 200 initially allocated to each group. 15 did not receive EVLT as they declined to participate. Of the 185 who received EVLT, 12 were lost to follow-up (11 refused or unavailable and 1 died). 39 did not	Inclusion: GSV insufficiency incompetence and reflux to level; CVI and/or symptoms by GSV incompetence and/ clinical findings at risk of va vein bleeding, thrombophle DVT; age 18-65 years; perfor status, according to the Am Society of Anaesthesiologis 1-11. Exclusion: Previous groin su except inguinal herniotomy post accessory saphenous v incompetence; small saphe insufficiency requiring treat thrombophilia; PAD; Maligr disease diagnosed in past 5 pregnancy or lactation. Baseline characteristics: Th baseline characteristics of t whom outcome data is ava not shown. The baseline dat includes those lost to follow for whom no ITT was under	o knee o knee is caused /or severe aricose ebitis and formance nerican sts of class urgery y; ant or vein enous vein otment; nant 5 years; he those for allable are ata below w-up, and rtaken!	Endovenous Laser therapy done with a 810mm diode laser, using Seldinger's technique, and a 20W laser power. Common procedures: Tumescent anaesthesia administered. Both groups also had multiple phlebectomies as needed. Analgosedation using intravenous midazolam allowed at the surgeon's discretion given as pain relief.	Flush ligation of SFJ and invagination stripping of the GSV just below the knee. Common procedures: See intervention column.	Post op, 3 months, 1 year and 2 years.	Reflux Advers e events HVVSS (higher means worse) CIVIQ- 2(higher means worse) Patient assessed symptoms	None reported
	separate site,	55 dia not	EVLT Su	urgery					

Reference	Study type	No. of patients	Patient characteristics			Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
	leading to risk that site effects, rather than treatment effects, could explain study results. Blinding not reported.	receive stripping as they declined to participate. Of the 161 who received stripping, 18 were lost to follow-up (18 refused or were unavailable). No ITT done – 173 evaluated in EVLT group and 143 evaluated in the stripping group.	Age	47.9(10.9)	48(10.7)					
			F	67%	70%					
			BMI	26.2(4.1)	26.3(4.9)					
			CEAP 2	53/185	47/161					
			CEAP3	95/185	76/161					
			CEAP4	36/185	35/161					
			CEAP5	1/185	2/161					
			CEAP6	0/185	1/161					
			HVVSS	16.1 (4.4- 34.3)	18.6 (3.6- 40.2)					
			GSV diameter at SFJ (mm)	8.7(2.8)	8.7(2.2)					
			CIVIQ-2	28.6(19)	29.4(16)					
		High risk of attrition bias due to differential loss to follow-up								

Reference	Study type	No. of patients	Patient characteristics			Intervention	Comparison		Length of follow-up	Outcome measures	Source of funding	
		rates and no ITT.										
Results: For c	ontinuous variab	les, only mean ar	nd ranges give	n.								
					EVL				Surgery			
Reflux at 2 years				31/173				2/143				
Adverse even	ts											
DVT					1/185				1/161			
Phlebitis					20/185				4/161			
Pain at 1 week					118/185				91/161			
Pain (VAS) mean (sd) [higher worse]					1.6(0.8)				1.3(0.6)			
	Neural damage/injury 3 months				17/105				22/161			
Hyper-pigmentation 3 months				17/185 57/185				22/161				
					57/185			19/16	51			
HVSS 3 month	HVSS 3 months				3.9(3)				3.8(3)			
HVSS1 year					2(2)				2.1(3)			
HVSS2 years					2.1(3)			1.9(3)				
CIVIQ-2 [note only done on last 100 patients enrolled] – 3 months				ths	12.8(14) [43]				18(16) [37]			
CIVIQ-2 [note only done on last 100 patients enrolled] – 1 yr					10.5(14)[40]				11.1(14)[32]			
CIVIQ-2 [note only done on last 100 patients enrolled] – 2 yrs					10.8(13)[41]				9.5(11)[33]			
Pt reported pain from varicose veins (taken from CIVIQ) at 3 months				onths	3(3)				4.6(4)			

Author's conclusions: Both EVLT and HLS are comparably safe and effective procedures to treat GSV incompetence.

Reference	Study type	No. of patients	Patient characteri	istics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Rautio T, Ohinmaa A, Perala J, Ohtonen P, Heikkinen T, Wiik H, Karjalianan P, Haukipuro K, Juvonen T. Endovenous obliteration versus conventional stripping operation in the treatment of primary varicose veins: a randomised controlled trial with comparison of the costs. JVasc Surg 2002; 35: 958-965.	RCT. Randomisation method unclear, but allocation concealment likely through sealed envelopes. No evidence of blinding. All treatments performed by the same single surgeon.	36 enrolled. 3 withdrew because of an unsuitable schedule and so 33 were randomised. After randomisation, 4 withdrew from the stripping group due to disappointment in being assigned to that group. A further patient withdrew from the EVRF group because of pregnancy. Hence drop out was surgery:4, EVRF 1. Ultimately 28 participated, 15 in the EVRF group and 13 in	Patients scheduled treatment of prim Inclusion: A valsal induced reversal of lasting > 2 second the threshold for it suitable for day ca symptomatic, prev and complicated O varicosis and isola and GSV trunk inst eligible for the stu Exclusion: Coagula tortuous and large mm) GSV trunks w Veins with a curve Baseline characte was deemed diffe groups at baseline numbers in each g to chance different baseline. mean (sd) unless se EVR age 33 (F:M 14:1	ary variose veins. ver manoeuvre- of blood flow s was considered nclusion. Patients ase surgery with viously untreated GSV tributary ted unilateral SFJ ufficiency were dy. opathy, multiple, e diameter (>12 vere excluded. e >90 degrees. ristics: Only age rent between e. Hence the small group did not lead aces of note at stated. F Surgery 6.7) 38 (6.8)	Stripping surgery. Groin dissected to expose the SFJ. Side branches of the GSV at the SFJ were divided and ligated. After local phlebectomy, the GSV was stripped from just below the knee to the groin with the venostrip with a 9mm olive. Common procedures: General anaesthetic given. Local phlebectomy and microsclerotherapy performed as necessary. Knee and groin anti- embolism stockings used for first 7 days.	Radiofrequency endovenous obliteration carried out with the VNUS Closure system (see Lurie studies). The catheter with sheathed electrode was inserted percutaneously with US scan guidance into the GSV at the ankle level, and then passed up to the SFJ. The entire length of the GSV was exsanguinated with compression and elevation, and the probe was then slowly withdrawn (3	7, 14, 28 and approx 49-56 days. Mean 50 days.	Physician reported outcomes (VSDS, VCSS, VDS). QoL (RAND 36) Post- operative pain Adverse events Treatment failure	not stated

Evidence tables clinical studies

Reference	Study type	No. of patien	ts Patient char	acteristics		Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding			
		the surgery	BMI	23.3(5.3)	24.0 (1.7)		cm/min). To						
		group. A further patier "retired" [declined any	diam.	6.4 (1.7)	6.1 (1.3)		avoid damage to the saphenous						
		treatment?] but was not	Office/light work	14/15	12/13		nerve, treatment was kept above the						
		withdrawn from the stuc	heavy dy. work	1/15	0/13		medial tibial condyle.						
		It is not	retired	0/15	1/13								
		reported from which group this patient was.	VCSS (median and range)	5 (4-9)	4 (4-6)								
			VSDS	1 (1-1)	1 (1-1)								
			VDS	1 (1-2)	1 (1-1)								
			operation time (mins)	75 (16.6)	57 (11)								
		Rand – 8				Rand – 8 dimensions	All were similar across groups						
Results:													
			EVRF		Surge	ry	р						
VSDS (post op	erative). Median				0 (0-1)	-	1						
Decrease in V point.	CSS [mean(sd)]. U	nclear time	5.1 (1.5)		4.4 (1.	1)	0.19						

0 (0-1)

1

VDS (post operative). Median (range)

0 (0-1)

Reference Study type	No. of patients	Patient characteristics		Intervention	Comp	parison	Length of follow- up	Outcome measures	Source of funding
Adverse events (intraoperative)									
groin hematoma	0/1		1/13						
thermal skin injuries	3/1	5	0/13						
Adverse events (post-operative)									
saphenous nerve paraesthesiae	2/1	5	3/13						
clinical thrombophlebitis	3/1		0/13						
local heamatoma	1/1		4/13						
thermal skin injury	1/1		0/13						
total	7/1		7/13	-)					
Post-op Pain at rest (VAS) average first 14 days [mean(sd)]	ed over 0.7	(0.5)	1.7 (1.	3)		0.017			
Post-op Pain on standing (VAS) av over first 14 days [mean(sd)]	eraged 1.3	(0.7)	2.6 (1.	9)		0.026			
Post-op Pain when walking (VAS) a over first 14 days [mean(sd)]	averaged 1.8	(0.8)	3.0 (1.	8)		0.036			
RAND Physical functioning 1 week [median(IQR)] for all	x 30 (21-48)	50 (35	-65)		0.07			
RAND Physical functioning 4week	s 0 (-	5-4)	5 (0-10))		0.11			
RAND role functioning physical 1	week 75 ((38-100)	75 (25	-100)		0.8			
RAND role functioning physical 4v	veeks 0 (0	I-O)	0 (-25	- 0)		0.9			
RAND bodily pain 1 week	23 ((5-25)	38 (20	-45)		0.05			
RAND bodily pain 4weeks	-23	(-28 -0)	-10 (-3	3-0)		0.6			
RAND general health perception 1	L week 0 (0	-8)	0 (-5 –	10)		0.7			
RAND general health perception 4	1weeks -5 (-	-8 - 0)	-5 (-5-	10)		0.7			
RAND energy 1 week	10 ((-3 – 20)	0 (-10	- 15)		0.5			

Reference	Study type	No. of patie	nts	Patient characteristics		Intervention	Com	parison	Length of follow- up	Outcome measures	Source of funding
RAND energy 4	1weeks		-10	(-10-0)	-10 (-2	5-10)		0.13			
RAND social fu	nctioning 1 week		23 (0-31)	25 (13	-50)		0.4			
RAND social fu	nctioning 4weeks		0 (-1	13-0)	0 (0-0)	l .		0.3			
RAND role fund	ctioning emotiona	l 1 week	0 (0	-0)	0 (0-0)	i -		0.9			
RAND role fund	ctioning emotiona	l 4weeks	0 (0	-0)	0 (0-0)	l .		0.5			
RAND emotion	al well being 1 we	ek	-4 (-	6 – 4)	-4 (-8	-4)		0.7			
RAND emotion	al well being 4we	eks	-4 (-	8-4)	-8 (-8-	0)		0.4			
Reflux			0/1	5		reflux in an accessory n of the GSV					
analgesic use (tabs/day	(number of 600mg	; ibuprofen	0.4	(0.49)	1.3 (1.	09)					

Reference	Study type	No. of patients	Patient character	istics			Interventio	on	Comparison	Length of follow- up	Outcome measures	Source of funding
Stotter L, Schaaf I, Bockelbrink A. Comparative outcomes of radiofrequency endoluminal ablation,	RCT. Randomisation carried out by a statistical department, but no other details given, and no reports	60 patients (60 limbs).20 allocated to cryostripping (NA). 40 (20+20) allocated to the two	Inclusion: Primary varicose v reflux on duplex. Exclusion: requiring avulsion phlebectomies in the thigh. Baseline Characteristics: (al not different):		ion h.		Ligation an invaginatio stripping to below the l to just belo the knee.	n o just knee. ow	EVRF with VNUS closure system. Aim was to increase temp. To 85 deg C and	24hrs, 1 week, 6 weeks, 1 year.	Treatment failure Patient satisfaction	None
invagination stripping, and cryostripping	of allocation concealment.	groups of relevance. 1 lost from		Surger (n=)	У	EVLA (n=)	procedures General Anaestheti		the probe was retracted at		Physician global	
in the		each group	M:F	5:15		6:14			3cm/min.		assessments	
treatment of great saphenous vein		at 1 year follow-up.	age	54 me 51 woi		41 men; 44 women						
insufficiency. 2006; 21: 60- 64.			duration of pathology	7.1 yrs		9.6 years						
			GSV diam. Distal to saphenofemoral ostium	7.1 (5-	14)	6.8(4.5- 12)						
Results: No varia	ances given for co	ntinuous varial	oles.									
		:	Surgery	1	EVLA							
Physician global weeks?)[lower b	impression score	(6	0.86	(0.62							

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Reference	Study type	No. of patients	Patient characteristics		Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Treatment failur open segments)	e (immediate in te	erms of	0/20	1/20					
Hematoma surfa	ice area at 1 week	k (cm2)	109	55					
Numbers with he	eamatoma 24 hou	ırs	19/20	14/20					
Numbers with he	eamatoma 1 weeł	k	18/20	11/20					
Cumulative up to	o 6 weeks impairn	nent score	7.9	2.8					
Cumulative up to	o 6 weeks pain sco	ore	7.5	4.6					
Adverse events u	up to 6 weeks								
DVT			0/20	0/20					
PE			0/20	0/20					
Saphenous ner	ve injury		0/20	0/20					
Treatment failur neovascularisatio	e (Recanalisation on)	or		2/19					
Patient satisfacti year – very satisf	ion with appearan fied	nce of leg 1	7/19 11/19	17/19 2/19					
Patient satisfacti year –satisfied	on with appearan	nce of leg 1	1/19	0/19					
Patient satisfacti year – not satisfi	on with appearan ed	nce of leg 1							

Reference	Study type	No. of patients	Patient characteristics		Interven	tion	Comparison	Length of follow- up	Outcome measures	Source of funding
very satisfied	ion with treatmen	·	7/19 12/19	17/19 1/19						
Patient satisfacti satisfied	ion with treatme	nt 1 year –	0/19	1/19						
Patient satisfacti not satisfied	ion with treatme	nt 1 year –								

	amonia2010B ²⁵⁸									
Reference	Study type	No. of patients	Patient c	haracteris	tics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
Subramonia S, Lees T. Randomised clinical trial of radiofrequency ablation or conventional high ligation and stripping for great saphenous varicose veins. British Journal of Surgery 2010; 97: 328- 336. NB notes that its companion paper, the HE paper, is excluded and should go in exclusion list: Subramonia S, Lees T. Radiofrequenc	RCT. Randomisation using a web based randomisation method stratified for age and sex. No evidence of allocation concealment. No blinding.	93. 48 randomised to EVRF and 45 to surgery. No treatment: EVRF: 1; surgery 4 (not related to trial or treatment allocation). No further loss to follow-up. No ITT (i.e. those withdrawing from treatment not assessed, and no imputations made).	symptoma (CEAP 2-6 recurrent duplex; pa ambulato Exclusion or deep sa incompet above kne or >12mm pacemake defibrillat <0.9]; pre	i); primary GSV reflu atient fit for ry. aphenous ence; twis ee; GSV dia h; GSV thro- ers or inte cor; PAD [A gnancy. character	se veins or x on or GA; phenous ited GSV am. <3 ombus; rnal ABPI istics:	Tributaries of the GSV ligated. GSV ligated (high ligation). PIN stripper used. Common procedures: Multiple phlebectomies. Both groups had general anethetic. Above knee graduated compression stockings worn for 2 weeks. Activity advice given.	EVRF: GSV accessed percutaneously, and VNUS Closure catheter introducing, with tip just below superficial epigastric vein. With a target temperature of 85C the probe was withdrawn at a rate of 1.5- 2 cm/min for the first 3cm and then 1-3cm per min for the rest of the GSV.	1 week, 5 weeks.	CEAP Micheales classification TCSS VDS AVVQ VEINES- QoL/Sym	None

Table 82: Subramonia2010B²⁵⁸

Reference	Study type	No. of patients	Patient ch	naracterist	tics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
y ablation vs. conventional surgery for varicose veins – a comparison			CEAP3 CEAP 4- 6	9/47 1/47	7/41 1/41					
of treatment costs in a randomised			TCSS 0 TCSS 1	25/47 12/47	27/41 7/41					
rial. European ournal of vascular and endovascular			TCSS 2 TCSS 3	4/47 1/47	1/41 1/41					
urgery 2010; 9: 104-111.			TCSS <u>></u> 4 VDS 0 VDS 1	2/47	2/41					
			VDS 2	44/47 1/47	35/41 4/41					
			ASA 1 ASA II	36/47 11/47	34/41 7/41					
			Recurre nt	3/47	2/41					
Results:										
Immediate treati EVRF, not clear if	ment failure (? As			e case of		occlusion failure, detected on o duplex, corrected after	Surgery 7/41 (complete stripping not po		Ρ	

Reference	Study type	No. of patients	Patient characterist	tics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
				immed	iate retreatment)	for these)	_		
1 week treatme GSV)	nt failure (defined	by duplex imag	ining of reflux in	0/47		5/41			
Adverse events									
cutaneous sen	sory abnormalities	s 1 week		9/47		20/41		0.003	
cutaneous sen	sory abnormalities	s 5 weeks		7/47		19/41		0.003	
parasthesia 1	week			5/47		11/41		0.049	
parasthesia 5 v	weeks			6/47		5/41		0.936	
groin wound p	oroblems 1 week			0/47		7/41			
non tender pa	lpable GSV with ov	verlying pigmen	tation 1 week	5/47		0/41			
pain level (VAS	5) during first weel	k post op (med[l	QR])	1.7 (0.	5-4.3)	4 (2.35-6.05)		95% CI= -2.75,	-0.79
numbers with	pain requiring ana	esthesia post o	р	30/47		40/41			
satisfaction (VAS	S)(med[IQR])			10 (8.4	l-10)	8.5 (7.5-10)		95% CI= 0.15, 1.44	
Numbers compl	etely satisfied (VA	S score of 10)		27/47		11/41		0.004	
numbers unwilli	umbers unwilling to recommend the procedure to others					9/41		0.005	
TCCS and VDS				TCSS ar	plete outcome reporting; "more t nd VDS after surgery. Two patien fter treatment"	•		•	
	mprovement in AVVQ QoL score (mean; variance for each group not iven, but CIs for the main difference given) 5 weeks [negative score			-9.12		-8.24		-3.64, 1.89; p=	0.532

Evidence tables clinical studies

Reference	Study type	No. of patients	Patient characterist	ics	Intervention	Comparison	Length of follow- up	Outcome measures	Source of funding
better]									
-	ysis improvement ot given, but CIs fo		ore (mean; variance erence given) 5	12.62		9.94		-1.65, 7.01; p=	0.220
	ysis improvement viven, but CIs for th		ean; variance for ice given) 5 weeks	12.80		7.83		0.80, 9.14; p=0).02
	AVVQ QoL score (the main differen		for each group not ks	(upper conver within	ed by using the equation SE= – lower CI)/3.92, and then ting SE OF THE MEAD DIFF to gp sd for each. ly /4.042 as used t dist for small	-8.24 (6.405)			

G.5.3 Foam sclerotherapy vs. endothermal ablation

Table 83: Rasmussen2011²²¹

See Table 63 for evidence table

Fable 84: Lattim							-		-	
Reference	Study type	Number of patients	Patient	character	istics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Lattimer CR et al. Cost and effectiveness of laser with phlebectomies compared with foam sclerotherapy in superficial venous insufficiency. Early results of a randomised controlled trial. Eur J Vasc and Endovasc Surg 2012; 43: 594- 600	Random ised controll ed trial. Number ed sealed envelop es used so probabl e allocati on conceal ment. Blinding not carried out. Strictly 1 leg per particip ant- if bilateral the worse leg was studied.	110 randomised – 56 EVLA and 54 FS. 6 did not receive EVLA or switched groups and 4 did not receive FS or switched groups. Per protocol analysis used as only those completing treatment (50 and 50) analysed at 3 week follow- up. Then 4 and 5 (respectively) lost to follow- up at 3 month follow-up. Overall, the loss was comparable across groups so minimal risk of selection bias.	sympto reflux of for bot Exclusio incomp >12mm of stud DVT; ar disease maligna known	on: Primary pomatic VV; on duplex; : h techniqu on: SPJ petence, GS i; prev surg y leg; histo terial occlu e (ABPI<0.8 ancy; pregr relevant al e Characte EVLA 47.4 (21-74) 62% 56% 44% 14%	SFJ suitable es. SV gery or FS ry of usive); active hancy; ilergies.	Outpatient FS – max 12ml foam. 1% STDS used. Injected into saphenous trunk at knee level. Tributaries treated as required on a subsequent occasion using a 21 gauge needle.	EVLA to GSV in day surgery theatre, using ELVes painless diode laser; 1470 nm; tumescent anaesthesia used. Access point near knee. Incompetent saphenous tributaries treated concurrently.	3 months	AVVQ VCSS Reflux Post procedure pain Return to normal activities	STD pharmaceut icals

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Reference	Study type	Number of patients	Patient	character	istics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
			4	30%	48%					
			5/6	12%	16%					
			VCSS	6 (2-20)	7 (3-17)					
			AVVQ	20 (1- 53)	25 (4- 50)					
			VFI	4.7 (0.9- 17.8)	5.9(1.1- 15.5)					
			GSV diam	7 (4-12)	8 (5-12)					
			BK GSV reflux	58%	74%					

Results

	EVLA	Foam Sclerotherapy	р
Median (IQR) pain for 7 days after treatment (VAS score with 100 worst pain)	33(18-54)	14(6-34)	P=0.005 Man Whitney U test
Median (IQR) time to return to normal activities	7.5(2-15)	3(1-10)	P=0.011 Man Whitney U test
Median days requiring analgesia tablets	2(0-21)	0(0-14)	
Reflux (Above knee) 3 weeks	1/50	8/50	
Reflux (Above knee) 3 months	9/46	9/45	
Reflux (Below knee) 3 weeks	7/50	24/50	

	Study Sype	Number patients	-	Patient characteristic	s	Intervention	Comparison	Length of follow-up		Source of funding
Reflux (Below knee)	3 months	5	21/46		19/45	5				
Need for further trea	atment		3/50) (but this may be beca ailable for FS group, in a				
Median (IQR) AVVQ	at 3 mont	ths	5.8(2.5-	12.2)	12.4(6-21.9)				
Median (IQR) VCSS a	it 3 montl	hs	1(0-3)		2(1-4)				
Median (IQR) VFI at 3	3 months		1.5(1.1-	2.4)	1.9(1	.3-2.7)				
Change from baselin months	e in AVV	Q by 3	12		9			0.	.062	
Change from baselin weeks	ie in VCSS	by 3	3		3			0.	.721	
Change from baselin months	ie in VCSS	by 3	5		4			0.	.817	
Change from baselin 3 months	ie in VCSS	VFI by	2.6		3.1			0.	.791	
Adverse event: DVT			1/50		0/50					
Adverse event: thror	mbosis		2/50		8/50					

Reference	Study type	No of patients	Patien charac	t teristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Gonzales- Zeh R, Armisen R, Barahona S. Endovenous aser and echo-guided foam ablation in great saphenous vein reflux: one year follow-up results. ournal of vascular surgery 2008; 48: 940-6.	Prospective cohort study (effectively a non- randomised trial). Patients were told each treatment option was equivalent and were allowed to choose their own group [NB this probably carries less of a bias risk than physician allocation, as less risk, albeit in the internet age, of allocation based on prognostic characteristics]. Only one limb per patient was included and treated in this study. A single surgeon with experience of 800 EVLA procedures and 2000 foam sclerotherapy	98. No patients dropped out and all followed up.	incom GSV ar insuffic reflux second over a least 2 upper Exclus active throm clottin throm clottin throm coagul history of mal Baselin charac	ciency wi time of 0 ds measu distance 0cm in the leg. ion: pregetion disturb bophlebi g disturb bophlebi g disturb the the stics: ce the lacc g were wo	of the ith a .5 red of at ne mancy; tis, ances; or orders; history s. k of the	US guided foam sclerotherapy. Policodanol was used with a sclerosant to air ratio of 1:4. Single injection using a venflon of 3% foam applied with the patient in supine. Injection at the point of most distal reflux in GSV. Volume of injection decided by surgeon. Immediate 2 minute compression applied afterward. Common procedures: Full length class II compression stocking applied 10 minutes post procedure with the patient lying down. The stockings were to be continued for 7 days and nights continuously and for 7 additional days where usage in the day only was required. Patients advised to walk for 30	Endovenous laser ablation. GSV at the point of most distal reflux was punctured with a 21 gauge needle under US guidance, and a guide wire passed up the GSV, followed by a 5 Fr introducer sheath to a point 1cm below the SFJ. 600 micron laser filament passed through the sheath to a point 1 cm below the SFJ. Then it was withdrawn at 1-2 mm/sec in continuous mode, with energy delivered by a 980 nm diode laser at a power of 15 W. Common procedures: See	1 week, 1 month, 6 months and 1 year.	Reflux	Not stated

Table OF. - la - **7** - h 2000¹⁰⁷

Reference	Study type	No of patients	Patien charao	t teristics		Intervention	Comparisor		Length of follow-up	Outcome measures	Source of funding
	procedures did both interventions. Patients were not allowed to mix, to avoid contamination of		3 16/4 4 5 5 9/45 6 3/45 2/45		16/53 10/53 6/53 5/53	post procedure and to walk daily for 30 minutes. Simple non- prescription analgesia allowed for pain.	column.				
	patient expectations. Clinical and		supe rficia I	45/4 5	45/53						
	ultrasound follow-ups done by a blinded assessor.		VCSS (med ian[I QR])	3 (3- 5)	3 (3- 5)						
Results											
						EVLA	Foam sc	lerothera	ару	р	
Reflux at 7 day	ys (not all reflux, inclu	ides open ar	d flux)			0/53	4/45			not given	
Reflux at 6 mc recanalisation	onths (not strictly refl)	ux, but refer	red to a	s partial		1/53	2/45			not given	
Reflux at 1 yea	ar (true reflux)					1/45	8/45			0.0360	
Non occlusion	at one year					3/45	12/45			0.04650	
VCSS at 1 year year) [median	· (only includes those (IQR)]	with no reca	inalisati	on at on	9	3(3-2)	2(3-2)			No p for between gro	oup effects
Adverse even	ts										
pain (VAS 1-10), 10 worst)	4.9 (1.5); n=45	2	1.0 (1.5)	n=53		0.0082			
phlebitis		10/45		2	22/53			0.0529			
paraesthesia		2/45		1	/53			0.5923			
DVT		0/45		2	2/53			0.4982			

G.5.4 Truncal and tributary treatment vs. truncal treatment alone

Table 86: Carradice 2009⁴⁶

Reference	Study type	No. patients	Patient cha	aracteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Carradice D, Mekako AI, Hatfield J, Chetter IC. Randomised clinical trial of concomitant or sequential phlebectomy after endovenous laser therapy for varicose veins. British Journal of Surgery 2009; 96: 369-375	RCT. "rando mised to one of two groups using sealed envelop es". Thus allocati on conceal ment likely, but method of random isation unclear.	50, 25 randomised to each group. Per- protocol analysis used. In the truncal +tributary group one was lost to follow-up by 6 weeks, and all received intervention. In the trunk only group, one was withdrawn after not receiving intervention, and none others lost to follow-up by 6 weeks. Thus 24 in each group analysed (4% loss in each – unlikely to cause bias) at 6 weeks. At 3 months, further losses to follow-up led to 23 in the truncal	unilateral, s saphenous incompeter duplex; per > 4mm. Exclusion: s saphenous incompeter Baseline Ch	Patients with symptomatic varicose veiu nce and GSV igenicular ve Saphenopop or deep ven nce on duple haracteristic: e for all base truncal + tributary 51.1(14.3) 8:17 4(2.25-5) 85(70- 99) 74(51- 84)	c great ns; SFJ reflux on ein diameter liteal, small ous ex. s:	Endovenous laser therapy - cannulation at the GSV. 600nm laser fibre introduced, delivering 14 W continuous 810 nm laser. Target energy delivery was 80-100 J/cm. Concomitant ambulatory phlebectomy of varicosities also carried out. Stab incisions of 1- 2mm made over varicose tributaries, and veins avulsed. This is not clear, but looks likely that	Endovenous laser therapy as for intervention group, but with no concomitant procedures. Sequential ambulatory phlebectomi es allowed after 6 weeks if required.	6 weeks (after 6 weeks, sequential ambulatory phlebectomi es allowed after this time in the comparison group). Further follow-ups were used at 3 months and 1 year.	AVVQ SF36 & EQ- SD VCSS Return to normal activity and work Post procedure pain Obliteration Need for further	None specified

Reference	Study type	No. patients	Patient c	haracteristics		Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
		+ tributary group and 22 in the trunk only group, and at 1 year the	AVVQ	13.29(11. 09- 15.50)	13.75(11.67 -15.82)	BOTH groups had an elastic bandage applied to the			procedures	
		analysed figures	EQ-5D			leg for 1 week, and replaced by a class II (20-30 mmHg) full length graduated support stocking for a further 5 weeks.				
Results:			-			T	L.			
$\Lambda \setminus \Lambda \cup \Omega$ 6 weeks (lowe	ar bottor) [median(IOR)]		Frucal + tributa 7.9 (4.1 – 10.7)	-	Truncal on 13.5 (10.9 ·	-		p <0.001	
AVVQ 0 weeks (lowe				2.0 (0.4-7.7) [2:		9.6(2.2 – 1			0.015	
SF-36 EQ-5D						n low resolution g			0.010	
Reflux at 1 week			()/24		0/24				
SFJ Reflux at 1 year			:	L/20		2/21				
GSV reflux at 1 year			2	2/20		1/21				
VCCS at 1-6 weeks (I	ower bette	er)	I	no data		no date				
VCSS at 3 months			(0(0-1) [23]		2(0-2) [22]			<0.001	
VCSS at 1 year			(0(0-1) [20]		1(0-1) [21]			0.433	
Adverse events										
phlebitis			(0/24		1/24				
nigmentation	mentation		2/24		0/24					

Reference	Study type	No. patients	Patient	Patient characteristics		rvention	Comparison	Length of follow-up	Outcome measures	Source of funding	
thigh "neuralgia"				1/24		0/24					
post op pain				no data but recorded no diffe	rence	between gro	oups at days 1,3	and 7			
return to work (day				10 (4-21) 3(1-14)				0.054			
return to normal ac	tivity (days)		8(1-14)		2(1-5)			0.166		
need for subsequer	nt ambulato	ory phlebectomy at 6	weeks	1/25		16/24					
Patient satisfaction would have it again	satisfaction (would recommend to a friend or nave it again)		20/20		19/21						

G.6 Chapter 10 – compression after interventional treatment

Table 87: Hamel-Desnos 2010¹¹¹

Reference	Study type	No. of patients	Patient characteristics	Intervention	Compariso n	Length of follow-up	Outcome measures	Source of funding
Hamel-Desnos CM, Guias BJ, Desnos PR, Mesgard A. Foam sclerotherapy of the saphenous veins: randomised controlled trial with or without compression. Eur J Vasc Endovasc Surg 2010; 39: 500-507	RCT. Method of randomisation unclear. It is stated that randomisation was done 5-10 minutes after sclerotherapy using a randomisation list provided by the statistician. It is therefore likely that there was no allocation concealment. There were two study centres and two regions of treatment (great or small saphenous vein) and randomisation was stratified for these. No	60 (31 in combination, 29 in sclerotherapy only). No drop-outs reported.	Inclusion: Patients_presenting for treatment of symptomatic varicose veins; aged >18; incompetence of the GSV or SSV; trunk diameter >8mm for GSV and 6mm for SSV; venous reflux lasting at least 1 sec; C2- 6. Exclusion: Any factors limiting the ability to participate in an informed manner; isolated SFJ incompetence; post-surgical recurrence of varices without trunk recurrence; chronic liver or renal disease; pregnancy/lactation; malignancy; history of DVT; cardio-vascular/respiratory problems; Coagulopathy; alcohol intolerance; allergies; patent foramen ovale; previous migraine or CNS disturbance after sclerosing therapy; lycra allergy; inability to apply compression. Only difference was for age(p=0.018); overall mean	Foam sclerotherapy using one volume of aetoxisclerol and 4 volumes of sterile air. Up to 3 sessions were permitted. 5-10 minutes after the first sclerotherapy session, class 2 French standard 15- 20 mmHg stockings (thigh length for GSV and knee length for SSV) were applied, to be worn during the day for 3 weeks following	Foam sclerother apy using one volume of aetoxiscler ol and 4 volumes of sterile air. Up to 3 sessions were permitted. No compressi on given.	1 month	reflux at 1 month after treatment. QoL Patient assessed symptoms Adverse events	Some funding (for stats) by compressio n stocking company, as well as free stockings.

Reference	Study type	No. of patients	Patient	character	istics	Intervention	Compariso n	Length of follow-up	Outcome measures	Source of funding
	mention of		CEAP cla	ass was 2.	6 (range 2-6)	treatment.				
	blinding.		Baseline	e Characte	eristics:					
				Sclero + compre ssion	sclero only					
			age	61(11)	53(14)					
			%men	13	3					
			GSV affected	19/31	17/29					
Results:										
			Sclerothera	py plus co	ompression	Sclerotherapy	/ only	р		
Reflux at 28 days			0/31			0/29				
-	al score – change fro ge = improvement)	om baseline	-5.5 (10) [22	2]		-9 (9.9) [21]				
-	al score – change fro ge = improvement)	om baseline	-9.4 (10) [23	3]		-11 (14) [24]				
rate of improvem	nent at day 28							reportedly	no difference b	etween group
heavy legs			(20/30) 67%	6		(16/29) 55%				
pain			(21/30) 70%	6		(17/29) 59%				
oedema			(15/30) 50%	/ 0		(15/29) 52%				
paraesthesia			(17/30) 57%	6		(13/29) 45%				
cramp			(11/30) 37%	6		(16/29) 55%				
	ator of 30 for the co n in the paper (no d	-			only group led	to the best agreer	ment with the			
Patient satisfaction	on with sclerotherag	ру								
"very effective"	' day 14		15/30			20/29				
"very effective"	' day 28		22/30			19/29				

Reference	Study type	No. of patients	Patient characteristics	Intervention	Compariso n	Length of follow-up	Outcome measures	Source of funding
Patient satisfaction "very effective" da		Ν		7/30				
Adverse events								
major neurological	events <24 hrs	()/31	0/29				
visual disturbance (mins	scotoma) resolving	within 15	0/31	1/29				
moderate pain day	28	:	1/30	3/29				
pigmentation		:	2/30	1/29				
thrombophlebitis		:	3/30	3/29				
compliance with co	mpression							
number wearing e	every day	:	12/30					
mean number of o	days use (max 21da	ays)	11					

Reference	Study type	No. patients	Patient char	racteristic	S	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Houtermans- Auckel JP, van Rossum E, Teijink JAW, Dahlmans AAHR, Eussen EFB, Nicolai SPA, Welton RJTJ. To waer or not to wear compression stockings after varicose vein stripping: a randomised controlled trial. Eur J Endovasc Surg 2009; 38: 387-391.	RCT. Random ised using comput er generat ed random isation list. Closed envelop es used, with pre- random isation allocati on conceal ment. No blinding after random isation.	104 randomised – 52 allocated to each group. 2 patients in the comparison group and 6 patients in the intervention group dropped out 3 days post- operatively (prior to starting the 4 weeks compression). These drop-outs were thus not due to lack of efficacy or adverse events of compression as an adjunct, and this unlikely to cause bias. All unavailable for follow-up. Available case analysis done, with 46 analysed in the intervention group and 50 in the comparison group.	Inclusion: Produce to GSV mincompetenduplex US; C Excluson: Pawear compression wear compression with ulcers. Baseline Cha Described as Men% age R leg % Muller phlebecto my done as well	reflux;_cor ce of the C2-C3; atients un ession sto had alrea n stocking aracterist	mplete GSV on able to ockings; ady used s; patients ics:	Crossectomy and short GSV inversion stripping, done as day surgery, 90% of which was under spinal anaesthetic. Standard elastic bandaging with a rolled gauze over the proximal part of the GSV applied for 3 days. Then fitted with class 2 medical compression stockings (measured fit), at 23-32 mmHg for 4 weeks, day and night for the first 2 weeks and then day only for the final 2 weeks.	Crossectomy and short GSV inversion stripping, done as day surgery, 90% of which was under spinal anaesthetic. Standard elastic bandaging with a rolled gauze over the proximal part of the GSV applied for 3 days. No further compression given.	4 weeks post-op	post –op pain (VAS) Post operative adverse events Return to full activity	None

Table 88: Houtermans-Auckel 2009¹¹⁸

Reference	Study type	No. patients		Patient chara	cteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding
Results:										
			-	ry plus ression	Surgery only					
Adverse events										
post op pain (\	/AS) at 3 days		2.5(2.	8)[46]	1.8(2.2)[50]					
post op pain (\	/AS) at 2 week	s	2.2(2.	3)[46]	2.2(2.4)[50]					
post op pain (\	/AS) at 4 week	s	0.8(1.	5)[46]	0.5(0.8)[50]					
Numbness 3 da	ays		4/46		3/50					
Numbness 3 da	ays		0/46		2/50					
Numbness 3 da	ays		0/46		0/50					
Bleeding 3 day	'S		0/46		0/50					
Infection 3 day	/S		2/46		1/50					
Seroma 3 days			2/46		1/50					
Return to work (measured at 4	4 weeks)	15(8.4)[46]	11(7.5)[50]					

G.7 Chapter 11 - Pregnancy

Table 89: Mota-Capitao 1995¹⁷⁵

Reference	Study type	No. of patients	Patient charac	teristics	Risk factors studied	Outcome measures	Length of follow-up	Source of funding	
Mota-Capitao L, Menezes JD, Gouveia- Oliveira A. Clinical predictors of the severity of chronic venous insufficiency of the lower limbs: a multivariable analysis. Phlebology 1995; 10: 155- 159.	Cross- sectional for many variables, but effectively a case-study for the potentially prognostic variables of family history and past medical history.	474 patient presenting to 18 different GPs with CV symptoms.	4% were class (asymptomatio 1 (mild), 33% c	0 CVI c), 42% class class 2 d 21class 3 had a ces. 3% had rT. 90% had s for > 1 d to	Many "risk factors" in this study were measured cross-sectionally, but potentially prognostic variables are age, sex, pregnancy, parity, hormones, family history, medication and past medical history.	A linear relationship between the risk factors and the ordinal outcome variable (class of CVI) was taken as evidence of an association of the risk factor with progression of CVI. However, only retrospective risk factors could be said to have a causative effect.	NA	Not stated	
Results:									
After multivariable	After multivariable analysis of factors having a linear relationship with CVI class, the potentially prognostic variables that remained in the model were as follows:								
Factor			co-efficient	p					
age			0.036	<0.001					

age	0.036	<0.001
CVI in both parents	0.568	0.026
History of thrombophlebitis	0.775	0.019
History of post-thrombotic syndrome	1.627	0.028
History of lymphoedema	1.712	0.026

Evidence tables clinical studies

Table 90: Fischer 2006⁹⁸

See Table 26 for evidence table.

Table 91: Zubilewicz 2009²⁹⁰

See Table 21 for evidence table.

Table 92: Thaler 2001²⁶⁰

Reference	Study type	No. of patients	Patient charac	cteristics			Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
Thaler et al. Compression stockings prophylaxis of emergent varicose veins in pregnancy: a prospective randomised controlled study. Swiss Med Weekly 2001; 131: 659-662.	Randomise d controlled study. No blinding for patients or HCP but blinding of assessors. No mention of randomisati on method nor of allocation concealmen t. Stratified randomisati on for venous status at entry (slight varicose changes present/abs	45 randomised. 3 drop outs from treatment in group 1 of the intervention group (see Intervention column) due to miscarriage, relocation abroad and failure to re- attend. No drop outs in the other groups. There were 9 losses of final follow-up data (controls: 3, group 1: 2; group 2: 4) but all were included in the analysis using earlier follow-up measures. Attrition bias	Inclusion: Preguncomplicated gestation; abso Exclusion (post compression; f Baseline chara Maternal age parity Numbers with varicose veins at entry	d pregnai ence of G st entry): miscarria	ncies <12 w SSV reflux a intoleranc ge	at SFJ.	Two treatment groups: group 1 wore class I compression tights on the left leg and class II on the right leg; group 2 wore class II compression tights on the left leg and class I on the right leg; This mirror division was to exclude a laterality bias in varicose vein emergence.	No stocking control group	Up to 6-8 weeks post- partum	Emergence of any varicose veins (including reticular veins) Duplex evidence of GSV reflux (>2 secs).	Ganzioni, a stocking manufactu rer, provided stockings and "logistic support".	

Reference	Study type	No. of patients	Patient chara	cteristics	Intervention	Comparison	Length of follow-up	Outcome measures	Source of funding	
	ent).	minimal as there was only a 10% differential in drop out between combined group 1,2 and the controls.			the purposes of the review the results for group 1 and 2 have been merged.					
Results:										
			controls	Group 1	Group 2	р				
Patients with	emergent varico	ose veins at 1 week	7/14	5/12	8/14 0.94					
post-partum				13/26	13/26					
Patients with emergent third trimester GSV reflux at the SFJ during 3rd trimester		4/15	1/27		0.047					
Leg symptoms at one week postpartum – better than baseline		0/14	3/12	4/14	0.03	0.03				
				7/26						

Appendix H: Evidence tables economic studies

Chapter 7 – assessment for treatment

Table 93: Blomgren 2006A

L. Blomgren, N. Zethraeus, G. Johansson, B. Jonsson, and D. Bergqvist. Cost consequences of preoperative duplex examination before varicose vein surgery: a randomized clinical trial. *Phlebology* 21 (2): 90-95, 2006.

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CC Study design: RCT – Within trial analysis Approach to analysis: Comparison of costs arising in the first 2 years of varicose vein treatment Perspective: Care- giver in Sweden (direct costs only) Time horizon: 2 years Discounting: Costs = 3%;	Population: Patients aged 20-75 admitted to hospital with varicose veins. Excluding those with pure cosmetic complaints, previous venous surgery or sclerotherapy, history of suspected or manifest deep venous thrombosis, active or healed leg ulcer, peripheral arterial disease, previous significant trauma to the leg, general illness and drug or alcohol abuse. Mean age: Duplex = 48 years, No duplex = 45 years. Gender: Duplex = 37% male, No duplex = 33% male Intervention 1: Duplex for pre-operative examination on top of participating surgeons' standard procedure for	Mean cost per patient: Intervention 1: SEK 13,051 (£900) Intervention 2: SEK 11,193 (£772) Incremental (2-1): SEK 1,858 (£128) Currency & cost year: 2004 Swedish krona (presented here as 2004 UK pounds ^a) Cost components incorporated: Costs for staff, physicians, colour flow duplex imagers and overhead costs, operating room costs (including salaries for anaesthetic and theatre staff, drugs, materials for cleaning and draping, gowns and	Quality of Life: No significant difference (no further data reported). ²⁵	ICER: NR Analysis of uncertainty: Uncertainty was not explored.

	clinical examination (varied by surgeon, sometimes included hand-held Doppler testing). ²⁴ Intervention 2: Participating surgeons' standard procedure for clinical examination (varied by surgeon, sometimes included hand-held Doppler testing) only.	gloves), extra operative costs (ie for preoperative mapping), and admission costs if the patient was required to stay overnight.	
ources			

Cost sources: All costs were taken from the hospital accounting system (Capio St Göran's Hospital, Stockholm, Sweden).

Comments Source of funding: NR. **Limitations:** The time horizon was restricted to two years and thus may not fully capture cost differences between the different assessment strategies; costs of re-treatment post 2 years which are likely to favour use of duplex will not have been captured. Uncertainty is not formally explored, but the authors note that with a longer follow-up the use of duplex could be cost-saving. QALYs are not considered, therefore no ICER can be presented. Finally, unit costs and resource

use estimates are obtained from the trial only, rather than via a systematic procedure.

Overall applicability^b: Partially applicable Overall quality^c: Potentially serious limitations

Abbreviations: CC = cost comparison; ICER = incremental cost-effectiveness ratio; NR = not reported; QALYs = quality-adjusted life years; SEK = Swedish Krona

(a) Converted using 2004 purchasing power parities¹⁹⁵

(b) Directly applicable / Partially applicable / Not applicable

(c) Minor limitations /Potentially serious limitations / Very serious limitations

H.2 Chapter 8 – conservative management

Table 94: GOHEL2010¹⁰⁶ See Table 97: GOHEL2010

Table 95: MICHAELS2006¹⁷⁰

J. A. Michaels, W. B. Campbell, J. E. Brazier, J. B. Macintyre, S. J. Palfreyman, J. Ratcliffe, and K. Rigby. Randomised clinical trial, observational study and assessment of cost-effectiveness of the treatment of varicose veins (REACTIV trial). *Health Technol.Assess.* 10 (13):1-196, 2006.

Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA Study design: Decision-analytic Markov models were built for three different patient groups. Approach to analysis: Cost effectiveness modelling was over a period of 10 years (120 cycles). The analysis was not based solely on data from the 2- year randomized controlled trial because of small	Population & interventionsPopulation:Patients with primary varicose veins. Subgroups: 'moderate' varicose veins with reflux; and 'severe' varicose veins.Cohort settings: Start age = 46 years Female = 90%Intervention 1: Conservative treatmentIntervention 2: Standard surgery (saphenofemoral ligation, stripping and multiple phlebectomies)	Mean cost per patient: Moderate varicose veins Intervention 1: £473 Intervention 2: £920 Incremental (2-1): £447 Severe varicose veins Intervention 1: £0 Intervention 2: £880 Incremental (2-1): £880 Currency & cost year: 2003 UK pounds Cost components incorporated: Initial costs of treatment	Health outcomes Primary outcome measure: QALYs (mean per patient) Moderate varicose veins (Group 2)Intervention 1: 6.589 QALYS Intervention 2: 6.803 QALYS Incremental (2-1): 0.214 QALYS Severe varicose veins (Group 3)Intervention 1: 6.341 QALYS Intervention 2: 6.795 QALYS Incremental (2-1): 0.454 QALYS	Cost effectiveness Moderate varicose veins (Group 2) Intervention 2 versus Intervention 1: ICER: £2,089 per QALY gained (d/a) Severe varicose veins (Group 3) Intervention 2 versus Intervention 1 ICER: £1,938 per QALY gained (d/a)† Analysis of uncertainty: Univariate sensitivity analysis performed on: costs of surgery, costs of major complications after surgery, probability of residual veins after surgery, probability of minor complications after surgery and difference in the progression rate of reflux versus no reflux.
Markov models were built for three different patient groups. Approach to analysis: Cost effectiveness modelling was over a period of 10 years (120 cycles). The analysis was not based solely on data from the 2- year randomized controlled trial	<pre>varicose veins. Cohort settings: Start age = 46 years Female = 90% Intervention 1: Conservative treatment Intervention 2: Standard surgery (saphenofemoral ligation, stripping and multiple</pre>	Intervention 2: £920 Incremental (2-1): £447 Severe varicose veins Intervention 1: £0 Intervention 2: £880 Incremental (2-1): £880 Currency & cost year: 2003 UK pounds Cost components incorporated: Initial costs of treatment	Intervention 2: 6.803 QALYs Incremental (2-1): 0.214 QALYs Severe varicose veins (Group 3)Intervention 1: 6.341 QALYs Intervention 2: 6.795 QALYs Incremental (2-1): 0.454	Severe varicose veins (Group 3) Intervention 2 versus Intervention 1 ICER: £1,938 per QALY gained (d/a)† Analysis of uncertainty: Univariate sensitivity analysis performed on: costs of surgery, costs of major complications after surgery, probability of residual veins after surgery, probability of minor complications after surgery and difference in the progression rate of reflux versus no
sample size. A separate within-trial economic analysis was also carried out; results are as in	Liquid sclerotherapy was also included as a comparator; results are not presented here.	(surgery, sclerotherapy), costs of retreatment, hospital admission/visits, visits to the GP, practice nurse and other healthcare professionals		Generally, the cost-effectiveness results are fairly robust to the univariate and multivariate sensitivity analyses. All ICERs fall below £20,000 per QALY.

J. A. Michaels, W. B. Campbell, J. E. Brazier, J. B. Macintyre, S. J. Palfreyman, J. Ratcliffe, and K. Rigby. Randomised clinical trial, observational study and assessment of cost-effectiveness of the treatment of varicose veins (REACTIV trial). *Health Technol.Assess.* 10 (13):1-196, 2006.

	Radcliffe et al (2006)	(e.g., visits to the A&E and anticoagulation units).		
	Perspective:	Also included are costs of		
	UK NHS	developing major or minor		
	Time horizon: 10 yrs	surgical complications, and		
	Treatment effect	costs of co-morbidity.		
	duration: 10 yrs			
	Discounting: Costs =			
	3.5%; Outcomes =			
l	3.5%			
	- ·			

Data sources

Health outcomes: Some outcomes (for example, risk of complications following surgery or sclerotherapy, and rate of progression/recurrence of varicose veins) were taken from the randomized controlled trial contained in the report. Other outcomes (for example, probability of progression with reflux and progression without reflux) were informed by systematic reviews including Rigby et al. 2004²²⁸.

Quality-of-life weights: Derived from SF-6D and EQ-5D scores. SF-6D scores were calculated from SF-36 data using an algorithm developed by Brazier et al. 2002³⁴.

Cost sources: Unit costs for all resources used by patients in the randomized controlled trial were obtained from the data sources in the UK including the NHS Reference costs, the Personal Social Services Research Unit and the British National Formulary (BNF). Data on resources use collected from the randomized controlled trial.

Comments

Source of funding: NHS R&D Health Technology Assessment (HTA) programme; **Limitations:** The retreatment options and rates of retreatment modelled are based on expert opinion, although no detail is given on the expert(s) or how this information was elicited. The clinical pathway is based on strict assumptions of who can receive which treatment, and may not fully reflect what happens in current practice. Utility data is based on an average of SF-36 and EQ-5D data; no reason is given.

Overall applicability*: Directly applicable **Overall quality**:** Minor limitations

Abbreviations: CUA = cost-utility analysis; d/a deterministic analysis ICER = incremental cost-effectiveness ratio; NR = not reported.

⁺The within trial analysis was conducted for this group – results are as presented in Ratcliffe et al 2006 (Table 96)

^ Surgery shows extended dominance over sclerotherapy in that a blend between conservative treatment and surgery offers better value for money than sclerotherapy;

* Directly applicable / Partially applicable / Not applicable; ** Minor limitations /Potentially serious Limitations / Very serious limitations

Table 96: RATCLIFFE2006²²³

J. Ratcliffe, J. E. Brazier, W. B. Campbell, S. Palfreyman, J. B. Macintyre, and J. A. Michaels. Cost-effectiveness analysis of surgery versus conservative treatment for uncomplicated varicose veins in a randomized clinical trial. *Br.J.Surg.* 93 (2):182-186, 2006.

Study details	Population & interventions	Costs ⁺	Health outcomes ⁺	Cost effectiveness ⁺
Economic analysis: CUA Study design: A randomized controlled trial conducted at two vascular units within the NHS. Approach to analysis based on the 2-year data from the randomized controlled trial Perspective: UK NHS Time horizon: 2 years Treatment effect duration: 2 yrs Discounting: Costs = 3.5%; Outcomes = 3.5%	Population:Patients with uncomplicated varicose veins and evidence of saphenopopliteal reflux.Patients with recurrent varicose veins were excluded.Cohort settings: Start age = NR Male/ Female = NRIntervention 1: Conservative treatment (compression therapy plus lifestyle advice) N=124Intervention 2: Stripping surgery N=122	Mean per patient: Intervention 1: £345 Intervention 2: £733 Incremental (2-1): £389 (95% CI: 282 to 506; p < 0.05) Currency & cost year: 2002-2003 UK pounds Cost components incorporated: Hospital inpatient admissions, surgical treatments, outpatient visits, other NHS visits (to the A&E, anticoagulation clinics, GP or practice nurse), retreatment costs, compression hosiery and treatment of complications.	Primary outcome measure: QALYs per patient (using SF-6D scores, n=94) Intervention 1: 1.420 QALYs Intervention 2: 1.503 QALYs Incremental (2-1): 0.083 QALYs (95% CI: 0.005 to 0.162; p < 0.05) Other outcome measures (mean): QALYs per patients (using EQ- 5D values, n=91) Incremental (2-1): 0.133 QALYs	Intervention 2 versus Intervention 1 (using SF-6D scores): ICER: £4,682 per QALY gained (pa) 95% CI for ICER: £2,039 to £20,830 per QALY gained Probability cost-effective: With a threshold of £20,000 per QALY and QALY estimates based on SF-6D scores, the probability that surgery is cost- effective was 70%. At a £30,000 per QALY threshold, the probability increases to 76%. Analysis of uncertainty: Uncertainty around cost-effectiveness was assessed using bootstrap methods. The percentiles from the bootstrap replications were used to derive the cost- effectiveness acceptability curve. Sensitivity analysis showed that the economic results are fairly robust. Using EQ-5D values (instead of SF-6D scores) gives an ICER of £3,299 per QALY. Using NHS Reference Costs for surgical treatment (instead of local unit costs) gives an ICER of £5,708 per QALY.

Varicose Veins Full Guideline A

J. Ratcliffe, J. E. Brazier, W. B. Campbell, S. Palfreyman, J. B. Macintyre, and J. A. Michaels. Cost-effectiveness analysis of surgery versus conservative treatment for uncomplicated varicose veins in a randomized clinical trial. *Br.J.Surg.* 93 (2):182-186, 2006.

Health outcomes: This was taken from the results of the randomized controlled trial reported by Michaels et al. 2006¹⁶⁹.

Quality-of-life weights: SF-36 questionnaire scores at 1, 6, 12 and 24 months of follow-up were translated into preference-based SF-6D scores using the algorithm developed by Brazier et al. 2002³⁴. EQ-5D scores were also considered.

Cost sources: NHS Reference Costs and Personal Social Services Research Unit. Where national data was not available, local unit costs were obtained from the finance departments of two hospitals.

Comments

Source of funding: NHS R&D Health Technology Assessment (HTA) programme; **Limitations:** No decision analytic model was conducted to capture long-term costs and health outcomes. The short 2-year time horizon may underestimate the cost-effectiveness of surgical treatment as the clinical benefits of surgery including improvements in health-related quality of life would be expected to endure beyond 24 months. Including long-term costs and health outcomes may still give lower ICERs.

Overall applicability*: Directly applicable **Overall quality**:** Minor limitations

Abbreviations: CI = confidence interval; CUA = cost-utility analysis; d/a deterministic analysis; ICER = incremental cost-effectiveness ratio; NR = not reported; pa = probabilistic analysis * Directly applicable / Partially applicable / Not applicable; ** Minor limitations /Potentially serious Limitations / Very serious limitations †These results are also presented in Michaels 2006¹⁷⁰

Table 97: GOHEL2010

M. S. Gohel, D. M. Epste	in, and A. H. Davies. Cost-effect	tiveness of traditional and endo	M. S. Gohel, D. M. Epstein, and A. H. Davies. Cost-effectiveness of traditional and endovenous treatments for varicose veins. Br.J.Surg. 97 (12):1815-1823, 2010.								
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness							
Economic analysis: CUA Study design: Decision-analytic Markov model Approach to analysis: The model considers in the first 3 months the following outcomes: (1) initial intervention was successful and patient had no residual varicosities (2) veins were occluded but there remain residual varicosities, and (3) there is residual reflux or incomplete occlusion. Thereafter, the model considers the recurrence of vein reflux but not the recurrence of varicosities.	Population: Patients with unilateral symptomatic primary saphenous varicose veins Cohort settings: Start age = NR Male/Female = NR Intervention 1: Conservative care Intervention 2: Ultrasound-guided foam sclerotherapy Intervention 3: Endovenous laser ablation (local anaesthesia) Intervention 4: Radiofrequency ablation (local anaesthesia) Intervention 5: Surgery (day case) Intervention 6: Endovenous laser ablation (general anaesthesia) Intervention 7:	Mean cost per patient: Intervention 1: £0 Intervention 2: £429 Intervention 3: £1,031 Intervention 4: £1,110 Intervention 5: £1,242 Intervention 6: £1,915 Intervention 7: £1,964 Intervention 8: £2,000 Currency & cost year: 2008 UK pounds Cost components incorporated: Costs of catheter and generator, staff, ultrasonography, outpatient visits and sclerosant	Primary outcome measure: QALYs (mean per patient) Intervention 1: 3.522 QALYs Intervention 2: 3.836 QALYs Intervention 3: 3.940 QALYs Intervention 4: 3.944 QALYs Intervention 5: 3.951 QALYs Intervention 6: 3.954 QALYs Intervention 8: 3.954 QALYs	ICERs Intervention 2 versus Intervention 1: £1,366 per QALY gained (d/a) Intervention 3 versus Intervention 2: £5,799 per QALY gained (d/a) Intervention 4 versus Intervention 3: £17,350 per QALY gained (d/a) Intervention 5 versus Intervention 4: £19,012 per QALY gained (d/a) Intervention 7 versus Intervention 5: £100,451 per QALY gained (d/a) Intervention 6 was extendedly dominated and intervention 8 was dominated. Intervention 5 was the cost-effective strategy with a probability of 0.29. Intervention 3 had a probability of 0.29. Intervention 3 had a probability of 0.24. Analysis of uncertainty: One-way sensitivity analysis was conducted by varying: (1) the costs of treatments (2) estimates of relative treatment effectiveness with regards to saphenous vein reflux and residual varicosities and (3) the correlation							

IVI. S. Gonel, D. IVI. Epste	ein, and A. H. Davies. Cost-effect	iveness of traditional and endov	enous treatments for varicose	veins. Br.J.Surg. 97 (12):1815-1823, 2010.
Time horizon: 5 years Treatment effect duration: 5 years	Radiofrequency ablation (general anaesthesia) Intervention 8:			between the risks of incomplete vein occlusion after treatment and residual varicosities.
Discounting: Costs =3.5% ; Outcomes = 3.5%	Surgery (in patient)			The results changed significantly from the base case in the following instances. If the odds ratio for re-intervention for residual varicosities after sequential versus concomitant phlebectomy was 5.50, radiofrequency ablation (under local anaesthesia) and endovenous laser ablation (under local anaesthesia) are equally likely to be cost effective and day-case surgery is

Data sources

Health outcomes: Some outcomes were taken from clinical trials whilst other outcomes were informed by meta-analytic studies. The probability of complete/ successful occlusion following surgical ligation and stripping, for instance, was informed by the results of the clinical trial van den Bos et al. 2008 ²⁷⁵; and the relative risks of retreatment for residual varicosities after sequential versus concomitant phlebectomy was taken from results of the randomized controlled study Carradice et al. 2009 ⁴⁶. The odds ratio of incomplete occlusion following stripping surgery versus sclerotherapy, on the other hand, was informed by the meta-analysis Wright et al. 2006 ²⁸⁶.

dominated.

Quality-of-life weights: EQ5D-derived HRQoL scores and profile based on Rasmussen et al. 2007 ²¹⁹, Rautio et al. 2002 ²²⁴ and Michaels et al. 2006 ¹⁷⁰. **Cost sources:** 2008-2009 UK NHS Reference costs, published drug and device manufacturer's list prices (for 2008-2009).

Comments

Source of funding: European Venous Forum Group, which is partly funded by the pharmaceutical industry; **Limitations:** Modelling was undertaken over a 5 year horizon, yet the costs and health outcomes associated with recurrence of varicosities are not considered beyond the first 3 months. All treatments of residual varicosities with ultrasound-guided foam sclerotherapy at 3 months are assumed to be successful.

Overall applicability*: Directly applicable **Overall quality**:** Potentially serious limitations

Abbreviations: CI = confidence interval; CUA = cost-utility analysis; d/a deterministic analysis ICER = incremental cost-effectiveness ratio; NR = not reported; pa = probabilistic analysis; * Directly applicable / Partially applicable / Not applicable; ** Minor limitations /Potentially serious Limitations / Very serious limitations

1 Appendix I: Forest plots

2 I.1 Chapter 7 – assessment for treatment

3 I.1.1 Diagnostic accuracy of hand held doppler

Figure 26: Diagnostic accuracy of hand held Doppler (threshold 0.5 seconds) vs. Duplex (threshold 1 second): Sapheno-femoral junction

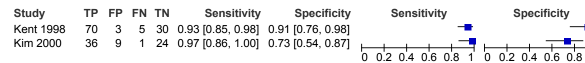


Figure 27: Diagnostic accuracy of hand held Doppler (threshold 1 second) vs. Duplex (threshold 1 second): Sapheno-femoral junction

Study	TP	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
Rautio 2002A	59	1	46	36	0.56 [0.46, 0.66]	0.97 [0.86, 1.00]		
Rautio 2002B	31	1	17	13	0.65 [0.49, 0.78]	0.93 [0.66, 1.00]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 28: Diagnostic accuracy of hand held Doppler (threshold 0.5 second) vs. Duplex (threshold 0.5 second): Sapheno-femoral junction

Study	TP	FP	FN	TN	Sensitivity	Specificity	Sensitivity	Specificity
Mercer 1998	43	2	16	28	0.73 [0.60, 0.84]	0.93 [0.78, 0.99]		

Figure 29: Diagnostic accuracy of hand held Doppler (threshold unknown) vs. Duplex (threshold unknown): Sapheno-femoral junction

Study	ΤР	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
DePalma 1993	24	5	26	25	0.48 [0.34, 0.63]	0.83 [0.65, 0.94]		
Salaman 1995	49	1	4	18	0.92 [0.82, 0.98]	0.95 [0.74, 1.00]		
Van der Heijden 1993	45	1	2	20	0.96 [0.85, 0.99]	0.95 [0.76, 1.00]		

Figure 30: Diagnostic accuracy of hand held Doppler (threshold 0.5 second) vs. Duplex (threshold 1 second): Sapheno-popliteal junction

Study	TP	FP	FN	TN	Sensitivity	Specificity	Sensitivity	Specificity
Kent 1998	14	18	3	73	0.82 [0.57, 0.96]	0.80 [0.71, 0.88]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 31: Diagnostic accuracy of hand held Doppler (threshold 1 second) vs. Duplex (threshold 1 second): Sapheno-popliteal junction

Study	ТР	FP	FN	TN	Sensitivity	Specificity	Sensitivity	Specificity
Rautio 2002A	3	4	10	95	0.23 [0.05, 0.54]	0.96 [0.90, 0.99]		

Figure 32: Diagnostic accuracy of hand held Doppler (threshold 0.5 second) vs. Duplex (threshold 0.5 second): Sapheno-popliteal junction

Study	TP	FP	FN	TΝ	Sensitivity	Specificity	Sensitivity	Specificity
Mercer 1998	20	4	6	59	0.77 [0.56, 0.91]	0.94 [0.85, 0.98]		

Figure 33: Diagnostic accuracy of hand held Doppler (threshold unknown) vs. Duplex (threshold unknown): Sapheno-popliteal junction

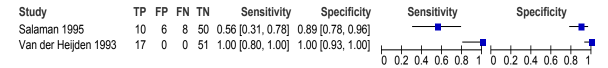


Figure 34: Diagnostic accuracy of hand held Doppler (threshold 0.5 second) vs. Duplex (threshold 1 second): Great Saphenous Vein

Study	TP	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
Kent 1998	79	8	4	17	0.95 [0.88, 0.99]	0.68 [0.46, 0.85]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 35: Diagnostic accuracy of hand held Doppler (threshold 1 second) vs. Duplex (threshold 1 second): Great Saphenous Vein

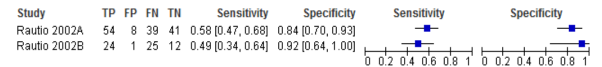


Figure 36: Diagnostic accuracy of hand held Doppler (threshold unknown) vs. Duplex (threshold unknown): Great Saphenous Vein

Study	TP	FP	FN	TN	Sensitivity	Specificity	Sensitivity	Specificity
Darke 1997	83	0	4	13	0.95 [0.89, 0.99]	1.00 [0.75, 1.00]	-	
Van der Heijden 1993	41	1	4	22	0.91 [0.79, 0.98]	0.96 [0.78, 1.00]		

Figure 37: Diagnostic accuracy of hand held Doppler (threshold unknown) vs. Duplex (threshold unknown): Short Saphenous Vein

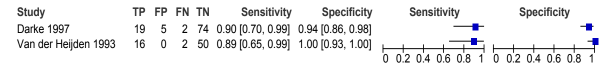


Figure 38: Diagnostic accuracy of hand held Doppler (threshold 0.5 second) vs. Duplex (threshold 1 second): Perforators

Study	TP	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
Kent 1998	13	69	2	24	0.87 [0.60, 0.98]	0.26 [0.17, 0.36]	0 0.2 0.4 0.6 0.8 1	

Figure 39: Diagnostic accuracy of hand held Doppler (threshold 0.5 second)vs. Duplex (threshold 0.5 second): Perforators

Study	ΤР	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
Mercer 1998	18	8	17	46	0.51 [0.34, 0.69]	0.85 [0.73, 0.93]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 40: Diagnostic accuracy of hand held Doppler (threshold unknown) vs. Duplex (threshold unknown): Perforators

Study	TP	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
Salaman 1995	2	13	5	54	0.29 [0.04, 0.71]	0.81 [0.69, 0.89]		
Van der Heijden 1993	10	1	9	17	0.53 [0.29, 0.76]	0.94 [0.73, 1.00]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 41: Diagnostic accuracy of hand held Doppler (threshold 0.5 second) vs. Duplex (threshold 1 second): Popliteal veins

Study	TP	FP	FN	TN	Sensitivity	Specificity	Sensitivity	Specificity
Kent 1998	7	9	7	84	0.50 [0.23, 0.77]	0.90 [0.82, 0.95]		

Figure 42: Diagnostic accuracy of hand held Doppler (threshold unknown) vs. Duplex (threshold unknown): Popliteal veins

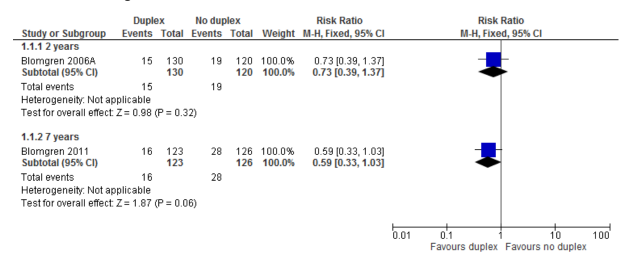
Study	TP	FP	FN	TN	Sensitivity	Specificity	Sensitivity	Specificity
Salaman 1995	2	1	3	68	0.40 [0.05, 0.85]	0.99 [0.92, 1.00]	0 0.2 0.4 0.6 0.8 1	

Figure 43: Diagnostic accuracy of hand held Doppler (threshold 1 second) vs. Duplex (threshold 1 second): Popliteal fossa

Study	TP	FP	FN	ΤN	Sensitivity	Specificity	Sensitivity	Specificity
Campbell 1997	28	8	11	74	0.72 [0.55, 0.85]	0.90 [0.82, 0.96]		

1 I.1.2 Duplex assessment prior to interventional treatment

Figure 44: Duplex prior to treatment vs. no duplex: patient-assessed symptoms - Operated limbs unchanged or worse



	Duple	x	No dup	olex		Risk Ratio	Risk	Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixe	ed, 95% Cl
1.2.1 6-8 weeks								
Blomgren 2005	10	160	37	166	97.4%	0.28 [0.14, 0.54]		
Smith 2002	1	92	1	97	2.6%	1.05 [0.07, 16.61]		-
Subtotal (95% CI)		252		263	100.0%	0.30 [0.16, 0.57]	•	
Total events	11		38					
Heterogeneity: Chi ² =				0%				
Test for overall effect:	Z = 3.69 (I	⊃ = 0.0	002)					
1.2.2 2 years								
Blomgren 2005	14	127	44	129	100.0%	0.32 [0.19, 0.56]		
Subtotal (95% CI)		127		129	100.0%	0.32 [0.19, 0.56]	•	
Total events	14		44					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z = 4.03 (I	⊃ < 0.0	001)					
1.2.3 7 years								
Blomgren 2011	11	95	38	99	100.0%	0.30 [0.16, 0.55]		
Subtotal (95% CI)		95		99	100.0%	0.30 [0.16, 0.55]		
Total events	11		38					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z = 3.86 (I	⊃ = 0.0	001)					
							FF	ļ
							0.01 0.1	11010
							⊢avours duplex	Favours no dup

Figure 45: Duplex prior to treatment vs. no duplex: SFJ reflux

1

Figure 46: Duplex prior to treatment vs. no duplex: SPJ reflux

	Duple	x	No dup	olex		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.3.1 8 weeks							
Blomgren 2005	4	160	9	166	100.0%	0.46 [0.14, 1.47]	
Subtotal (95% CI)		160		166	100.0%	0.46 [0.14, 1.47]	
Total events	4		9				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 1.31 (F	^o = 0.1	9)				
1.3.2 2 years							
Blomgren 2005	7	127	13	129	100.0%	0.55 [0.23, 1.33]	
Subtotal (95% CI)		127		129	100.0%	0.55 [0.23, 1.33]	
Total events	7		13				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 1.34 (F	P = 0.1	8)				
1.3.3 7 years							
Blomgren 2011	2	95	9	99	100.0%	0.23 [0.05, 1.04]	
Subtotal (95% CI)		95		99	100.0%	0.23 [0.05, 1.04]	
Total events	2		9				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 1.90 (F	P = 0.0	6)				
							0.01 0.1 1 10 10

Favours duplex Favours no duplex

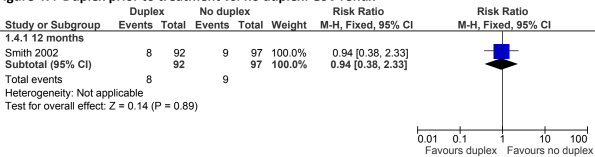


Figure 47: Duplex prior to treatment vs. no duplex: GSV reflux

Figure 48: Duplex prior to treatment vs. no duplex: SSV reflux

	Duple		No dup			Risk Ratio	Risk R		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed	, 95% CI	
1.5.1 6 weeks									
Smith 2002	4	92	6	97	100.0%	0.70 [0.20, 2.41]		_	
Subtotal (95% CI)		92		97	100.0%	0.70 [0.20, 2.41]			
Total events	4		6						
Heterogeneity: Not app	olicable								
Test for overall effect:	Z = 0.56 (P = 0.58	8)						
1.5.2 12 months									
Smith 2002	6	92	8	97	100.0%	0.79 [0.29, 2.19]			
Subtotal (95% CI)		92		97	100.0%	0.79 [0.29, 2.19]	\bullet	►	
Total events	6		8						
Heterogeneity: Not app	olicable								
		D - 0 6	5)						
Test for overall effect:	Z = 0.45 (I	0.03	5)						
Test for overall effect:	Z = 0.45 (0.03	5)						

Favours experimental Favours control

Figure 49: Duplex prior to treatment vs. no duplex: Perforators reflux

	Duple	x	No dup	olex		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% Cl	
1.6.1 6 weeks								
Smith 2002 Subtotal (95% CI)	1	92 92	5	97 97	100.0% 100.0%	0.21 [0.03, 1.77] 0.21 [0.03 , 1.77]		
Total events Heterogeneity: Not app	1 licable		5					
Test for overall effect: 2	Z = 1.43 (I	P = 0.1	5)					
1.6.2 12 months								
Smith 2002 Subtotal (95% Cl)	4	92 92	15	97 97	100.0% 100.0%	0.28 [0.10, 0.82] 0.28 [0.10, 0.82]		
Total events Heterogeneity: Not app Test for overall effect: 2		⊃ = 0.0	15 2)					
							0.01 0.1 1 10	100

0.1 10 100 Favours duplex Favours no duplex

, ca.							
	Duple	ЭX	No dup	olex		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Smith 2002	8	92	9	97	100.0%	0.94 [0.38, 2.33]	
Total (95% CI)		92		97	100.0%	0.94 [0.38, 2.33]	•
Total events Heterogeneity: Not app Test for overall effect: :		P = 0.8	9)				
		. 0.0	<i>c</i> ,				Favours duplex Favours no duplex

Figure 50: Duplex prior to treatment vs. no duplex: Development of new branch varicosities at 1 vear

Figure 51: Duplex prior to treatment vs. no duplex: Need for, or actual, re-operation

	Duple	x	No dup	olex		Risk Ratio		Risk Ra	atio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C		M-H, Fixed,	95% CI
1.8.1 2 years									
Blomgren 2005 Subtotal (95% CI)	3	145 145	14	147 147	100.0% 100.0%	0.22 [0.06, 0.74] 0.22 [0.06, 0.74]			
Total events	3		14						
Heterogeneity: Not ap	plicable								
Test for overall effect:	•	⊃ = 0.0	1)						
			,						
1.8.2 7 years									
Blomgren 2011	15	124	38	134	100.0%	0.43 [0.25, 0.74]			
Subtotal (95% CI)		124		134	100.0%	0.43 [0.25, 0.74]		•	
Total events	15		38						
Heterogeneity: Not ap	plicable								
Test for overall effect:	•	⊃ = 0.0	02)						
	(- /						
							<u> </u>		
							0.01	0.1 1	10 100

Favours duplex Favours no duplex

Figure 52: Duplex prior to treatment vs. no duplex: Adverse events - DVT

	Duplex	No du	plex		Risk Ratio	Risk Ratio
Study or Subgroup	Events Tot	al Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Blomgren 2005	0 14	45 0	147		Not estimable	
Total (95% CI)	14	15	147		Not estimable	
Total events	0	0				
Heterogeneity: Not app Test for overall effect:						0.01 0.1 1 10 100 Favours duplex Favours no duplex

Figure 53: Duplex prior to treatment vs. no duplex: Complications of varicose veins at 7 years – venous ulcer

	Duple	x	No dup	olex		Risk Ratio	Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixe	d, 95% Cl	
Blomgren 2011	0	70	0	88		Not estimable			
Total (95% CI)		70		88		Not estimable			
Total events	0		0						
Heterogeneity: Not app	olicable						0.01 0.1 1	10 1	100
Test for overall effect:	Not applic	able					Favours duplex		

Figure 54: Duplex prior to treatment vs. no duplex: Complications of varicose veins at 7 years – pigmentation/eczema

	Duplex	ĸ	No dup	lex		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% Cl
Blomgren 2011	3	70	9	88	100.0%	0.42 [0.12, 1.49]	
Total (95% Cl)		70		88	100.0%	0.42 [0.12, 1.49]	
Total events	3		9				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 1.34 (P	9 = 0.18	8)				Favours duplex Favours no duplex

I I.2 Chapter 8 – conservative management

2 I.2.1 Compression vs. no treatment/lifestyle advice

Figure 55: Compression vs. no treatment/lifestyle advice: numbers with pain or no improvement at end of treatment

	Compres	ssion	Contr	rol	Risk Ratio Risk Ratio				
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Rand	dom, 95% Cl	
Benigni 2003	27	61	37	53	64.6%	0.63 [0.45, 0.88]	-	ŀ	
Krijnen 1997	2	30	12	34	35.4%	0.19 [0.05, 0.78]			
Total (95% CI)		91		87	100.0%	0.41 [0.12, 1.40]		-	
Total events	29		49						
Heterogeneity: Tau ² =				= 0.08);	l² = 67%		0.01 0.1	1 10	100
Test for overall effect:	: Z = 1.42 (P	^r = 0.15)				Fav	ours compression	Favours cont	rol

Figure 56: Compression vs. no treatment/lifestyle advice: pain levels (VAS) at the end of treatment (better indicated by lower values). SMD used as scale of VAS unclear in both studies.

Juan										
	Com	pressi	on	0	Control			Std. Mean Difference	Std. Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% C	I IV, Random, 95% CI	
Anderson 1990	34.7	29.25	66	37.6	29.25	66	50.7%	-0.10 [-0.44, 0.24]] —	
Benigni 2003	1.4	1.8	62	2.9	2.1	55	49.3%	-0.77 [-1.14, -0.39]] —	
Total (95% CI)			128			121	100.0%	-0.43 [-1.08, 0.23]		
Heterogeneity: Tau ² =				1 (P = 0	.01); I² =	= 85%			+ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$ $+$;
Test for overall effect:	Z=1.28	I (P = 0.	20)						Favours compression Favours cor	ntrol

Figure 57: Compression vs. no treatment/lifestyle advice: numbers with heavy or tired legs or no improvement in heavy or tired legs at end of treatment

-	Compres	sion	Contr	ol	-	Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Benigni 2003	20	59	35	54	68.4%	0.52 [0.35, 0.79]	
Krijnen 1997	8	30	18	34	31.6%	0.50 [0.26, 0.99]	
Total (95% CI)		89		88	100.0%	0.52 [0.36, 0.73]	•
Total events	28		53				
Heterogeneity: Chi ² =	0.01, df = 1	(P = 0.9	92); l² = 0	%			0.01 0.1 1 10 100
Test for overall effect:	Z = 3.70 (P	= 0.000	02)			Fa	vours compression Favours control

Figure 58: Compression vs. no treatment/lifestyle advice: heavy or tired legs (VAS) at end of treatment (better indicated by lower values)

	Com	pressi	on	C	ontrol			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C	CI IV, Fixed, 95% CI
Anderson 1990	34.1	30.9	66	36.3	28.4	66	100.0%	-2.20 [-12.33, 7.93]	
Total (95% CI)			66			66	100.0%	-2.20 [-12.33, 7.93]	-
Heterogeneity: Not app Test for overall effect: 2		(P = 0	.67)					1	-50 -25 0 25 50 Favours compression Favours control

Figure 59: Compression vs. no treatment/lifestyle advice: numbers with no improvement in cramps at end of treatment

	Compres	sion	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Benigni 2003	37	61	44	55	100.0%	0.76 [0.60, 0.97]	
Total (95% CI)		61		55	100.0%	0.76 [0.60, 0.97]	•
Total events	37		44				
Heterogeneity: Not app	plicable					_	
Test for overall effect:	Z = 2.25 (P	= 0.02)				Favo	0.5 0.7 1 1.5 2 burs compression Favours control

Figure 60: Compression vs. no treatment/lifestyle advice: night cramps level (VAS) at end of treatment (better indicated by lower values)

	Com	pressi	on	Control				Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Anderson 1990	22.4	25.2	66	24.9	24.4	66	100.0%	-2.50 [-10.96, 5.96]	
Total (95% CI)			66			66	100.0%	-2.50 [-10.96, 5.96]	
Heterogeneity: Not ap Test for overall effect:		(P = 0	.56)					F	-20 -10 0 10 20 avours compression Favours control

Figure 61: Compression vs. no treatment/lifestyle advice: numbers of patients reporting no improvement in ankle swelling at end of treatment

	Compres	sion	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% Cl
Benigni 2003	35	61	43	53	100.0%	0.71 [0.55, 0.91]	
Total (95% CI)		61		53	100.0%	0.71 [0.55, 0.91]	\bullet
Total events	35		43				
Heterogeneity: Not app	olicable						0.5 0.7 1 1.5 2
Test for overall effect:	Z = 2.69 (P	= 0.007	")			Fa	avours compression Favours control

Figure 62: Compression vs. no treatment/lifestyle advice: self-reported swelling levels (VAS) at end of treatment (better indicated by lower values)

	Con	pressi	on	С	ontrol			Mean Difference		Me	an Differe	nce	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95%	CI	IV,	Fixed, 95%	% CI	
Anderson 1990	28.2	29.25	66	35.3	30.1	66	100.0%	-7.10 [-17.23, 3.03	8]				
Total (95% CI)			66			66	100.0%	-7.10 [-17.23, 3.03]				
Heterogeneity: Not ap Test for overall effect:		(P = 0.	17)						-100 Favours	-50 compress	0 ion Favo	50 50 ours contr	100 ol

Figure 63: Compression vs. no treatment/lifestyle advice body image dissatisfaction (VAS) at end of treatment (better indicated by lower values)

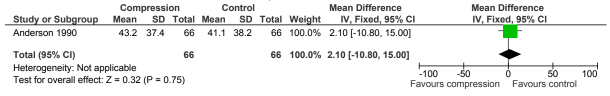


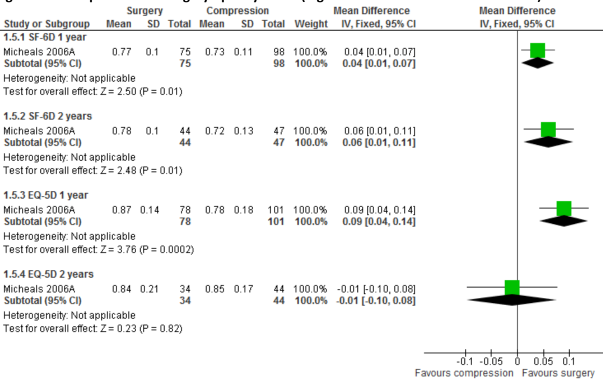
Figure 64: Compression vs. no treatment/lifestyle advice: numbers reporting fewer complaints

	compression		control			Risk Ratio	Risk Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixe	d, 95% Cl		
Krijnen 1997	17	30	4	34	100.0%	4.82 [1.82, 12.73]				
Total (95% CI)		30		34	100.0%	4.82 [1.82, 12.73]				
Total events	17		4							
Heterogeneity: Not ap Test for overall effect:	•	P = 0.00	2)				0.1 0.2 0.5 1 Favours control	2 5 10 Favours compress		

2 I.2.2 Compression vs. interventional treatment

3 I.2.2.1 Compression vs. surgery

Figure 65: Compression vs. surgery: quality of life (higher score indicates better outcome)



	Compres	-	Surge			Risk Ratio	ortion the same or worse Risk Ratio
Study or Subgroup	Events		0		Weight		M-H, Fixed, 95% Cl
1.6.1 aching at 1 yr	Lvento	Total	Lvento	Total	Weight		
Micheals 2006A	72	97	15	75	100.0%	3.71 [2.33, 5.92]	│ . <mark></mark>
Subtotal (95% CI)		97		75	100.0%	3.71 [2.33, 5.92]	│ ₹
Total events	72		15				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 5.50 (P	< 0.000	01)				
1.6.2 heaviness at 1 y	/ear						
Micheals 2006A	52	97	9	75	100.0%	4.47 [2.36, 8.47]	- <mark>-</mark> -
Subtotal (95% CI)		97		75	100.0%	4.47 [2.36, 8.47]	
Total events	52		9				
Heterogeneity: Not app							
Test for overall effect:	Z = 4.58 (P	< 0.000	01)				
1.6.3 itching at 1 year	r						
Micheals 2006A	42	97	10		100.0%	3.25 [1.75, 6.04]	
Subtotal (95% CI)		97		75	100.0%	3.25 [1.75, 6.04]	
Total events	42		10				
Heterogeneity: Not app			-				
Test for overall effect:	Z = 3.72 (P	= 0.000	2)				
1.6.4 swelling at 1 years	ar						
Micheals 2006A	31	97	8		100.0%	3.00 [1.46, 6.13]	
Subtotal (95% CI)		97		75	100.0%	3.00 [1.46, 6.13]	
Total events	31		8				
Heterogeneity: Not app							
Test for overall effect:	Z = 3.00 (P	= 0.003)				
1.6.5 cosmetic conce	erns at 1 ye	ar					
Micheals 2006A Subtotal (95% CI)	75	97 97	13	75 75	100.0% 100.0%	4.46 [2.69, 7.40] 4.46 [2.69, 7.40]	
Total events	75	•1	13				•
Heterogeneity: Not app							
Test for overall effect:	Z = 5.79 (P	< 0.000	01)				
						0.01	0.1 1 10 1
							s compression Favours surgery

Figure 66: Compression vs. surgery: Patient assessed symptoms (proportion the same or worse)

Figure 67: Compression vs. surgery: Adverse events – neural injury/damage

	Favours compres	Favours compression				Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% (CI M-H, Fixed, 95% CI
Micheals 2006A	0	122	1	124	100.0%	0.34 [0.01, 8.24	
Total (95% CI)		122		124	100.0%	0.34 [0.01, 8.24]	
Total events	0		1				
Heterogeneity: Not app Test for overall effect:						1	0.01 0.1 1 10 100 Favours compression Favours surgery

Figure 68: Compression vs. surgery: Patient dissatisfaction at 1 year

	compres	sion	surge	ry		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95%	CI M-H, Fixed, 95% CI
Micheals 2006A	53	107	3	65	100.0%	10.73 [3.50, 32.94	4] — <mark>—</mark> —
Total (95% CI)		107		65	100.0%	10.73 [3.50, 32.94	
Total events	53		3				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 4.15 (P	< 0.000)1)				Favours compression Favours surgery

I.3 Chapter 9 – interventional treatment

2 I.3.1 Stripping surgery vs. foam sclerotherapy

Figure 69: Stripping surgery vs. foam sclerotherapy:SF-36 Physical 4 weeks



Figure 70: Stripping surgery vs. foam sclerotherapy:SF-36 Physical 1 year

	surgery			sclerotherapy				Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C	I IV, Fixed, 95% CI
Rasmussen 2011	53.33	5.9	125	51.94	7.66	125	100.0%	1.39 [-0.30, 3.08]	
Total (95% CI)			125			125	100.0%	1.39 [-0.30, 3.08]	
Heterogeneity: Not ap Test for overall effect:	•		0.11)						-4 -2 0 2 4 Favours sclerotherapy Favours surgery

Figure 71: Stripping surgery vs. foam sclerotherapy: SF-36 mental 4 weeks

	SL	irgery		sclerotherapy				Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C	I IV, Fixed, 95% CI
Rasmussen 2011	55.15	7.81	125	56.1	7.51	125	100.0%	-0.95 [-2.85, 0.95	
Total (95% CI)			125			125	100.0%	-0.95 [-2.85, 0.95]	ı 🔶
Heterogeneity: Not ap Test for overall effect:).33)						-10 -5 0 5 10 Favours sclerotherapy Favours surgery

Figure 72: Stripping surgery vs. foam sclerotherapy: SF-36 mental 1 year

	SL	rgery sclerotherapy				ару		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Rasmussen 2011	55.83	6.31	125	54.73	8.89	125	100.0%	1.10 [-0.81, 3.01]	
Total (95% CI)			125			125	100.0%	1.10 [-0.81, 3.01]	
Heterogeneity: Not ap Test for overall effect:	•		0.26)					F	-4 -2 0 2 4 avours sclerotherapy Favours surgery

Figure 73: Stripping surgery vs. foam sclerotherapy: Patient- assessed outcomes: EQ-5D change from baseline to 2 years (higher better)

	S	tripping		Scle	Sclerotherapy			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C	I IV, Fixed, 95% CI
Shadid 2012	0.061	0.211	177	0.064	0.211	213	100.0%	-0.00 [-0.05, 0.04]]
Total (95% CI)			177			213	100.0%	-0.00 [-0.05, 0.04]	
Heterogeneity: Not a Test for overall effect			89)						-0.1 -0.05 0 0.05 0.1 Favours sclerotherapy Favours stripping

Figure 74: Stripping surgery vs. foam sclerotherapy: Patient-assessed symptoms: Pain due to varicose veins (subscale from SF-36)

	Strippin			Scle	rothera	ру		Mean Difference	Mean Difference
or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
io crossectomy us	ed with	scleroti	herapy						
ussen et al, 2011 t al (95% Cl)	85.11	23.45	124 124	88.77	17.11		100.0% 100.0%	-3.66 [-8.77, 1.45] - 3.66 [-8.77, 1.45]	-
igeneity: Not applic: ir overall effect: Z =		: 0.16)							
								-	-10 -5 0 5 10 Stripping Sclerotherapy

Figure 75: Stripping surgery vs. foam sclerotherapy: Patient-assessed symptoms: worsening of symptoms at 2 years

	Stripp	-	Sclerothe			Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl	
2.2.1 more pain								
Shadid 2012	6	177	14		100.0%	0.52 [0.20, 1.31]		
Subtotal (95% CI)		177		213	100.0%	0.52 [0.20, 1.31]		
Total events	6		14					
Heterogeneity: Not ap	pplicable							
Test for overall effect:	: Z = 1.39 ((P = 0.1	7)					
2.2.2 more heavy/tire	ed legs							
Shadid 2012	5	177	6	213	100.0%	1.00 [0.31, 3.23]		
Subtotal (95% CI)		177		213	100.0%	1.00 [0.31, 3.23]	-	
Total events	5		6					
Heterogeneity: Not ap	pplicable							
Test for overall effect	: Z = 0.00 ((P = 1.0	0)					
2.2.3 more cramps								
Shadid 2012	8	177	8	213	100.0%	1.20 [0.46, 3.14]		
Subtotal (95% CI)		177		213	100.0%	1.20 [0.46, 3.14]	-	
Total events	8		8					
Heterogeneity: Not ap	pplicable							
Test for overall effect	: Z = 0.38 ((P = 0.7	1)					
							0.01 0.1 1 10	10
Toot for outgroup dif	×	0 K 12 - 7	07 46-0	(D – O /		<i>v</i>	favours Stripping favours Sclero	tnera

Test for subgroup differences: $Chi^2 = 1.67$, df = 2 (P = 0.43), $l^2 = 0\%$

Figure 76: Stripping surgery vs. foam sclerotherapy: Patient-assessed symptoms: worsening of	f
symptoms at 1 year	

Sympt	lonis at	тусс	21				
	Stripp	ing	Sclerothe	егару		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% CI
2.3.1 more pain							
Shadid 2012	14	188	20	221	100.0%	0.82 [0.43, 1.58]	
Subtotal (95% CI)		188		221	100.0%	0.82 [0.43, 1.58]	-
Total events	14		20				
Heterogeneity: Not ap							
Test for overall effect:	Z=0.58 ((P = 0.5	i6)				
2.3.2 more heavy/tire	ed legs						
Shadid 2012	9	188	5	221	100.0%	2.12 [0.72, 6.20]	+
Subtotal (95% CI)		188		221	100.0%	2.12 [0.72, 6.20]	-
Total events	9		5				
Heterogeneity: Not ap	oplicable						
Test for overall effect:	Z=1.37 ((P = 0.1	7)				
2.3.3 more cramps							
Shadid 2012	9	188	10	221	100.0%	1.06 [0.44, 2.55]	
Subtotal (95% CI)		188		221	100.0%	1.06 [0.44, 2.55]	•
Total events	9		10				
Heterogeneity: Not ap	oplicable						
Test for overall effect:	Z=0.13 ((P = 0.9	10)				
							0.01 0.1 i 10 100
Test for subgroup diff	ferences.	Chi ž – 1	716 df= 2	P = 0.2	$(4) \mathbf{F} = 7$	496	Favours Stripping Favours Sclerothera
rearier adequade an	ici chicca.	vin = .	2. i 0, ui – Z	$y_1 = 0.5$	$a_{0} = c_{0}$	1.79	

Test for subgroup differences: $Chi^2 = 2.16$, df = 2 (P = 0.34), $I^2 = 7.4\%$

Figure 77: Stripping surgery vs. foam sclerotherapy: Patient-assessed symptoms: worsening of symptoms at 3 months

sympt	oms at	3 mo	ntns				
	Strippi	ing	Sclerothe	егару		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
2.10.1 more pain							
Shadid 2012 Subtotal (95% CI)	10	176 176	12	217 217	100.0% 100.0%	1.03 [0.45, 2.32] 1.03 [0.45, 2.32]	
Total events	10		12				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 0.07 (P = 0.9	5)				
2.10.2 more heavy/tir	red legs						_
Shadid 2012 Subtotal (95% CI)	2	176 176	8	217 217	100.0% 100.0%	0.31 [0.07, 1.43] 0.31 [0.07, 1.43]	
Total events Heterogeneity: Not ap	2 Indicable		8				
Test for overall effect:	•	D – 0 1	2)				
restion overall ellect.	2 - 1.50 (r – 0.1	3)				
2.10.3 more cramps							
Shadid 2012 Subtotal (95% CI)	6	176 176	9	217 217	100.0% 100.0%	0.82 [0.30, 2.27] 0.82 [0.30, 2.27]	
Total events Heterogeneity: Not ap	6 Inlicable		9				
Test for overall effect:	•	P = 0.7	0)				
			-,				
							0.01 0.1 1 10 100
							Favours Stripping Favours Sclerotherapy
Test for subgroup diff	erences:	Chi ^z = 1	1.84, df = 2	(P = 0.4	0), I ^z = 09	6	· · · · · · · · · · · · · · · · · · ·

2 3

Figure 78: Stripping surgery vs. foam sclerotherapy: Physician-reported outcomes: overall VCSS score change from baseline by 2 years

	St	tripping		Scle	rothera	ру		Mean Difference	Mean Difference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI		
Shadid 2012	-1.75	2.135	177	-1.49	2.135	213	100.0%	-0.26 [-0.69, 0.17]			
Total (95% CI)			177			213	100.0%	-0.26 [-0.69, 0.17]	•		
Heterogeneity: Not ap Test for overall effect:	•		23)						-2 -1 0 1 2 Favours stripping Favours sclerotherapy		

Figure 79: Stripping surgery vs. foam sclerotherapy: Physician-reported outcomes: VCSS pain

•		ripping	-	-	othera			Mean Difference	Mean Difference
or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1 month									
ido 2009	0.93	0.53	29	0.89	0.51	27	100.0%	0.04 [-0.23, 0.31]	
al (95% CI)			29			27	100.0%	0.04 [-0.23, 0.31]	
igeneity: Not ap	plicable	9							
r overall effect:	Z = 0.29	9 (P = 0).77)						
0									
2 months									
ido 2009	0.79	0.49	29 29	0.59	0.5	27		0.20 [-0.06, 0.46]	
al (95% Cl)			29			27	100.0%	0.20 [-0.06, 0.46]	
geneity: Not ap			1400						
r overall effect:	Z= 1.51	(F = 0).13)						
6 months									
ido 2009	0.72	0.53	29	0.56	0.51	27	100.0%	0.16 [-0.11, 0.43]	
al (95% CI)			29			27		0.16 [-0.11, 0.43]	
igeneity: Not ap	plicable								
r overall effect:	Z=1.15	5 (P = 0).25)						
									-0.5 -0.25 0 0.25 0.5 Favours stripping Favours sclerothera

1

2

Figure 80: Stripping surgery vs. foam sclerotherapy: Physician-reported outcomes: VCSS oedema

	Sti	ripping	g	Scler	othera	ару		Mean Difference	Mean Difference
or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1 month									
do 2009	0.69	0.6	29	0.7	0.54	27	100.0%	-0.01 [-0.31, 0.29]	
al (95% CI)			29			27	100.0%	-0.01 [-0.31, 0.29]	
geneity: Not app	plicable								
overall effect:	Z = 0.07	(P=0	0.95)						
2 months									
do 2009	0.59	0.63	29	0.56	0.64	27	100.0%	0.03 [-0.30, 0.36]	
al (95% CI)			29			27	100.0%	0.03 [-0.30, 0.36]	
geneity: Not app	plicable								
overall effect:	Z = 0.18	6 (P = 0	0.86)						
6 months									
do 2009	0.55	0.63	29	0.48	0.64	27	100.0%	0.07 [-0.26, 0.40]	
al (95% CI)			29			27	100.0%	0.07 [-0.26, 0.40]	
geneity: Not ap	plicable								
overall effect:	Z = 0.41	(P = 0	0.68)						
									-0.5 -0.25 0 0.25 0.5
									Favours stripping Favours sclerotherapy

Varicose Veins Full Guideline Appendices (July 2013)

Figure 81: Stripping surgery vs. foam sclerotherapy: Physician-reported outcomes: VCSS inflammation

	IIIIIaII	IIIIa	lion						
	St	ripping	9	Scler	othera	ару		Mean Difference	Mean Difference
or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1 month									
ido 2009	0.76	0.44	29	0.89	0.32	27	100.0%	-0.13 [-0.33, 0.07]	_
al (95% CI)			29			27	100.0%	-0.13 [-0.33, 0.07]	
geneity: Not a	pplicable	!							
r overall effect	t: Z = 1.27	' (P = (0.20)						
2 months									_
ido 2009	0.72	0.45	29 29	0.89	0.32	27 27		-0.17 [-0.37, 0.03]	
al (95% CI)			29			21	100.0%	-0.17 [-0.37, 0.03]	
geneity: Not a			140						
r overall effect	L. Z = 1.04	+ (F = (5.10)						
6 months									
ido 2009	0.72	0.45	29	0.89	0.32	27	100.0%	-0.17 [-0.37, 0.03]	
al (95% CI)			29			27		-0.17 [-0.37, 0.03]	
geneity: Not a	pplicable	!							
r overall effect	t: Z = 1.64	(P = (0.10)						
		-							
									-0.5 -0.25 0 0.25 0.5
									-0.5 -0.25 0 0.25 0.5

Favours stripping Favours sclerotherapy

1

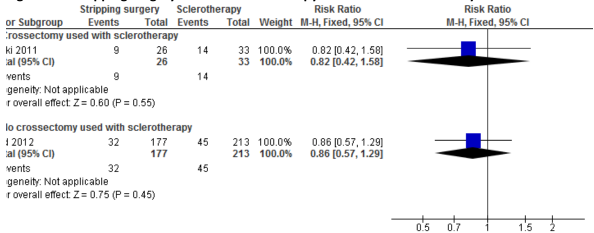
Figure 82: Stripping surgery vs. foam sclerotherapy: Presence of reflux within 3 months

	Favours stri	oping	Scleroth	erapy		Risk Ratio	Risk Ratio
or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% CI
rossectomy used v	with sclerothe	егару					
uroglou 2006	4	23	4	29	3.6%	1.26 [0.35, 4.50]	! •
11	3	28	3	28	3.1%	1.00 [0.22, 4.54]	
al (95% CI)		51		57	6.7%	1.14 [0.43, 3.02]	
vents	7		7				
geneity: Chi² = 0.05	, df = 1 (P = 0.3	32); I ² = I	0%				
r overall effect: Z = 0	0.27 (P = 0.79)						
o crossectomy use	ed with sclero	therapy					
ussen et al, 2011	3	135	2	144	2.0%	1.60 [0.27, 9.43]	
12012	32	176	56	217	51.4%	0.70 [0.48, 1.04]	-8-1
et al, 2006	12	94	56	176	39.9%		
al (95% CI)		405		537	93.3%	0.59 [0.43, 0.81]	•
vents	47		114				
geneity: Chi² = 3.76	, df = 2 (P = 0.1	15); I² = -	47%				
r overall effect: Z = 3	3.27 (P = 0.001)					
95% CI)		456		594	100.0%	0.63 [0.47, 0.85]	◆
vents	54		121				
geneity: Chi² = 5.28	, df = 4 (P = 0.3	26); I² = 3	24%				0.01 0.1 1 10 10
r overall effect: Z = 3	3.05 (P = 0.002	l)					Favours stripping Favours sclerothera
r subgroup differen	ces: Chi ² = 1.5	7, df = 1	(P = 0.21)	, I ² = 36.	.3%		ravours suppling Tavours scierourers

	Stripping su	rgery	Sclerothe	егару		Risk Ratio	Risk Ratio
or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
rossectomy used	d with scleroth	егару					
1	3	26	5	25	11.0%	0.58 [0.15, 2.16]	
al (95% CI)		26		25	11.0%	0.58 [0.15, 2.16]	
vents	3		5				
geneity: Not appli	cable						
r overall effect: Z =	= 0.82 (P = 0.41)	I					
o crossectomy u	sed with sclere	otherapy	1				
do 2009	3	29	6	27	11.5%	0.47 [0.13, 1.68]	
issen et al, 2011	4	108	20	123	15.2%	0.23 (0.08, 0.65)	
2012	43	188	64	221	34.4%	0.79 [0.57, 1.10]	
et al, 2006	13	94	65	176	27.9%	0.37 [0.22, 0.64]	
al (95% CI)		419		547	89.0%	0.46 [0.25, 0.84]	\bullet
vents	63		155				
geneity: Tau ² = 0.2 r overall effect: Z =			= 0.03); I ^z	= 68%			
95% CI)		445		572	100.0%	0.48 [0.29, 0.81]	•
vents geneity: Tau ² = 0.1 r overall effect: Z = r subgroup differe	= 2.77 (P = 0.00)	5)			,		0.05 0.2 1 5 20 Favours stripping Favours sclerotherap

Figure 83: Stripping surgery vs. foam sclerotherapy: Presence of reflux >3-12 months

Figure 84: Stripping surgery vs. foam sclerotherapy: Presence of reflux >1-5 years



Favours stripping Favours sclerotherapy

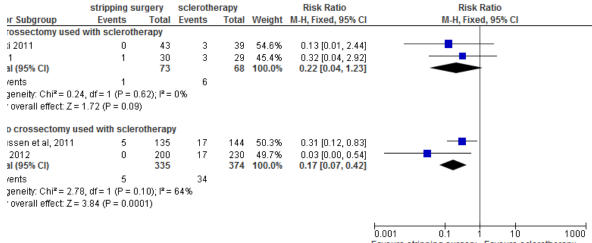
Figure 85: Stripping surgery vs. foam sclerotherapy: Need for further treatment from >3-12

	months						
	Stripping su	urgery	Sclerothe	erapy		Risk Ratio	Risk Ratio
or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% Cl
rossectomy us	sed with sclere	otherapy	,				
uroglou 2006 al (95% Cl)	2	28 28	4	30 30	100.0% 100.0%	0.54 [0.11, 2.70] 0.54 [0.11, 2.70]	
vents geneity: Not ap r overall effect:).45)	4				
							Image: Construction of the structure Image: Constructure Image: C

Figure 86: Stripping surgery vs. foam sclerotherapy: Major neurological event

	stripping su	urgery	sclerothe	erapy		Risk Ratio	Risk Ratio
or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI
rossectomy us	sed with scler	otherapy	y				
ti 2011 al (95% CI)	0	43 43	0	39 39		Not estimable Not estimable	
vents geneity: Not approver and a second sec			0				
							0.01 0.1 1 10 100 Favours stripping Favours sclerotherap

Figure 87: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: Phlebitis



Favours stripping surgery Favours sclerotherapy

Figure 88: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: PE

or Subgroup rossectomy used at 2011 1 at (95% CI) vents	Events with scleroth 0 0		Events 0 0	Total 39 29	Weight	M-H, Fixed, 95% Cl		Ν	∕I-H, Fixe	ed, 95% CI		
ti 2011 1 al (95% CI)	0 0	43 30				Not estimable						
1 al (95% CI)	0	30				Not estimable						
	-		0	29						1		
	0	73		20		Not estimable						
rents	0			68		Not estimable						
			0									
geneity: Not applic:	able											
overall effect: Not	applicable											
o crossectomy us	sed with scler	otherapy	y									
issen et al, 2011	0	135	1	144	51.0%	0.36 [0.01, 8.65]					_	
2012	0	200	1	230	49.0%	0.38 [0.02, 9.35]			-		_	
et al, 2006	0	94	0	178		Not estimable						
al (95% CI)		429		552	100.0%	0.37 [0.04, 3.53]						
/ents	0		2									
geneity: Chi² = 0.00	0, df = 1 (P = 0	.97); l² =	0%									
r overall effect: Z =	0.87 (P = 0.39)										
							0.01	0.1		1	10	100

Favours stripping surgery Favours sclerotherapy

	stripping su	irgery	scleroth	егару		Risk Ratio		Risk F	Ratio	
r Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixed	I, 95% CI	
ossectomy used	with scleroth	erapy								
i 2011	0	43	0	39		Not estimable				
1	0	30	0	29		Not estimable				
al (95% CI)		73		68		Not estimable				
ents	0		0							
eneity: Not applic	able									
overall effect: Not	t applicable									
crossectomy us	od with color	othoran								
-	seu with scier		y		40.000	4 07 10 07 40 001				
ssen et al, 2011	1	135	1	144	10.8%	1.07 [0.07, 16.88]				
2012	0	200	1	230	15.6%	0.38 [0.02, 9.35]				
t al, 2006	0	94	9	178	73.6%	0.10 [0.01, 1.69]			_	
I (95% CI)		429		552	100.0%	0.25 [0.05, 1.21]				
ents	1		11							
eneity: Chi² = 1.5	4, df = 2 (P = 0	.46); I² =	0%							
overall effect: Z =	1.73 (P = 0.08)								
							0.001	0.1 1	10	100
							E-	rinning ourgood	Environ a stand	

Figure 89: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: DVT

Favours stripping surgery Favours sclerotherapy

Figure 90: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: Nerve injury/damage - with crossectomy



2

1

Figure 91: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: Nerve injury/damage - no crossectomy

	aamage		0000000	,						
	stripping s	urgery	scleroth	erapy		Risk Ratio		Risk Rat	tio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixed, 9	95% CI	
5.6.2 no crossectomy us	ed with scler	rotherapy	y							
Figuerido 2009	6	29	0	27	17.7%	12.13 [0.72, 205.60]			-	
Rasmussen et al, 2011	5	135	2	144	66.3%	2.67 [0.53, 13.51]				
Shadid 2012 Subtotal (95% CI)	6	200 364	0	230 401	15.9% 100.0%	14.94 [0.85, 263.56] 6.30 [1.87, 21.20]		-		
Total events Heterogeneity: Chi² = 1.63 Test for overall effect: Z =		~ ~ ~	2 0%							
							L	0.1 1	10	100

Favours Stripping Surgery Favours Sclerotherapy

Figure 92: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: Skin discolouration/hyper pigmentation

r Subgroup	stripping su Events with scleroth	rgery Total	sclerothe	егару		Risk Ratio		Risk F	Ratio	
		Total	Evente						tutio -	
ossoctomy used	with scleroth		Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixed	l, 95% Cl	
ossectomy used		erapy								
2011	2	43	1	39	34.0%	1.81 [0.17, 19.23]			-	
1	1	30	2	29	66.0%	0.48 [0.05, 5.05]				
l (95% CI)		73		68	100.0%	0.94 [0.19, 4.53]				
ents	3		3							
eneity: Chi ² = 0.6	1, $df = 1$ (P = 0.	44); l² =	0%							
overall effect: Z =	0.08 (P = 0.93))								
crossectomy us	sed with scler	otherapy	,							
ssen et al, 2011	6	135	8	144	8.9%	0.80 [0.28, 2.25]				
2012	2	200	12	230	12.9%	0.19 [0.04, 0.85]		• <u> </u>		
et al, 2006	39	94	98	178	78.2%	0.75 [0.57, 0.99]				
I (95% CI)		429		552	100.0%	0.69 [0.53, 0.89]		•		
ents	47		118							
eneity: Chi² = 3.3	7, $df = 2$ (P = 0.	19); I² =	41%							
overall effect: Z =	2.78 (P = 0.00	5)								
							0.05 (0.2 1	5	20
							Favours stripp	ing surgery	Favours scler	otherapy

Figure 93: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: Post procedure pain

	procedu	i e pai					
	stripping su	irgery	scleroth	erapy		Risk Ratio	Risk Ratio
or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
rossectomy u	ised with scle	rothera	ру				
et al, 2008	48	60	7	30	53.4%	3.43 [1.77, 6.64]	- ∎ -
1	24	30	8	29	46.6%	2.90 [1.57, 5.37]	-∎ -
al (95% CI)		90		59	100.0%	3.18 [2.01, 5.03]	•
vents	72		15				
geneity: Chi ² =	: 0.14, df = 1 (F	P = 0.71	I²=0%				
r overall effect:	Z = 4.96 (P <	0.00001)				
o crossectom	ny used with s	clerothe	rapy				
et al, 2006	39	94	73	178	100.0%	1.01 [0.75, 1.36]	
al (95% CI)		94		178	100.0%	1.01 [0.75, 1.36]	
vents	39		73				
geneity: Not ap	pplicable						
r overall effect:	Z = 0.08 (P =	0.94)					
							0.001 0.1 1 10 1000 Favours stripping surgery Favours sclerotherapy
							Favours surpring surgery Favours scierourerapy

Figure 94: Stripping surgery vs. foam sclerotherapy: Adverse events from intervention: Post procedure pain VAS 1-10

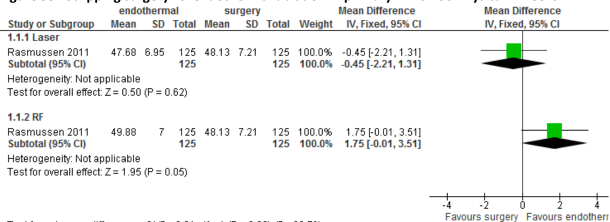
	p. 000		, pan			•								
	stripp	ing sur	gery	scler	othera	ру		Mean Difference			Mean Dif	ference		
r Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C			IV, Fixed	l, 95% Cl		
crossectomy u	used with	sclerot	herapy	,										
sen et al, 2011 Il (95% CI)	2.25	2.23	135 135	1.6	2.04		100.0% 100.0%	0.65 [0.15, 1.15] 0.65 [0.15, 1.15]					-	
eneity: Not appli overall effect: Z		= 0.01)												
									—					
									-2	-1	Ċ)	1	2

Favours stripping surgery Favours sclerotherapy

1 I.3.2 Stripping surgery vs. endothermal ablation

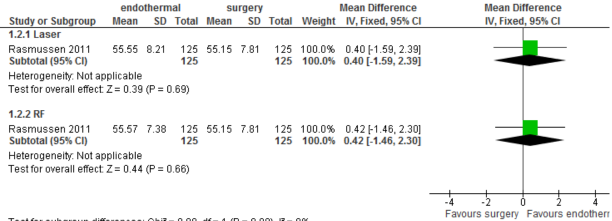
2 I.3.2.1 Primary varicose veins

Figure 95: Stripping surgery vs. endothermal ablation in primary VV: SF-36 Physical 4 weeks



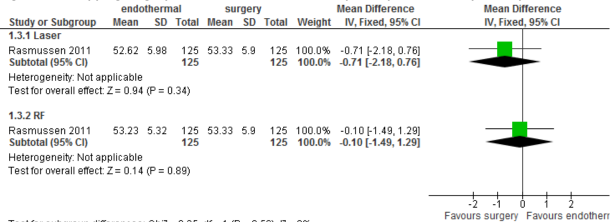
Test for subgroup differences: Chi² = 3.01, df = 1 (P = 0.08), l² = 66.7%

Figure 96: Stripping surgery vs. endothermal ablation in primary VV: SF-36 mental 4 weeks



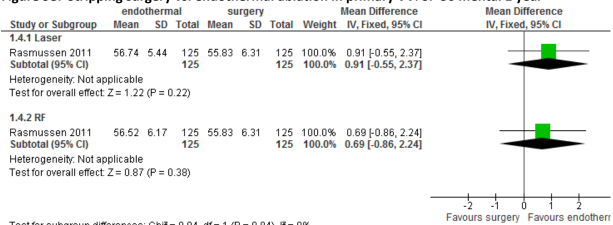
Test for subgroup differences: Chi² = 0.00, df = 1 (P = 0.99), l² = 0%

Figure 97: Stripping surgery vs. endothermal ablation in primary VV:SF-36 Physical 1 year



Test for subgroup differences: Chi² = 0.35, df = 1 (P = 0.56), l² = 0%

Figure 98: Stripping surgery vs. endothermal ablation in primary VV: SF-36 mental 1 year



Test for subgroup differences: Chi² = 0.04, df = 1 (P = 0.84), l² = 0%

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Figure 99: Stripping surgery vs. endothermal ablation in primary VV: Global quality of life – followup 1-12 weeks, 1 year and 2 years

			•		-				
	End	lotherma	d i	S	tripping			Std. Mean Difference	Std. Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
1.1.1 1-12 weeks									
Lurie 2003	-9.2	15.088	43	3.7	15	36	32.1%	-0.85 [-1.31, -0.39]	e
Rass 2011	12.8	14	43	18	16	37	33.3%	-0.34 [-0.79, 0.10]	
Subramonia 2010 Subtotal (95% CI)	-9.12	6.405	47 133	-8.24	6.405	41 114	34.7% 100.0%	-0.14 [-0.56, 0.28] - 0.43 [-0.84, -0.02]	
Heterogeneity: Tau² = Test for overall effect: .				(P = 0.0	08); I² =	61%			
1.1.2 1 year									
Rass 2011 Subtotal (95% CI)	10.5	14	40 40	11.1	14	32 32	100.0% 100.0%	-0.04 [-0.51, 0.42] - 0.04 [-0.51, 0.42]	-
Heterogeneity: Not ap Test for overall effect:	•		6)						
1.1.3 2 years									
Rass 2011 Subtotal (95% CI)	10.8	13	41 41	9.5	11	33 <mark>33</mark>	100.0% 100.0%	0.11 [-0.35, 0.56] 0.11 [-0.35, 0.56]	-
Heterogeneity: Not ap Test for overall effect:	•		5)						

Favours endothermal Favours stripping

[Note that Subramonia 2010 used AVVQ, whilst Rass 2012 and Lurie 2003 used CIVIQ -2 – hence the use of standardised mean differences]

Figure 100: Stripping surgery vs. endothermal ablation in primary VV: Patient reported symptoms – oedema and pain (dichotomous)

Sympt		uema	i anu p			nousj	
	Endothe	rmal	Stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% (CI M-H, Fixed, 95% CI
1.9.2 oedema							
Pronk 2010	6	56	10	62	100.0%	0.66 [0.26, 1.71	
Subtotal (95% CI)		56		62	100.0%	0.66 [0.26, 1.71]	
Total events	6		10				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 0.85 (P	9 = 0.40)				
1.9.4 pain							
Pronk 2010	1	56	6	62	100.0%	0.18 [0.02, 1.49]	
Subtotal (95% CI)		56		62	100.0%	0.18 [0.02, 1.49]	
Total events	1		6				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 1.59 (P	9 = 0.11)				
	,						
							0.01 0.1 1 10 100 Favours endothermal Favours stripping
							i avouis chaothermai i avouis stripping

Figure 101: Stripping surgery vs. endothermal ablation in primary VV: dissatisfaction with body image

	Endothe	rmal	Stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Perala 2005	1	15	2	13	19.1%	0.43 [0.04, 4.25]	
Pronk 2010	4	56	8	62	67.6%	0.55 [0.18, 1.74]	∎ ∔_
Stotter 2006	0	19	1	19	13.3%	0.33 [0.01, 7.70]	• • • • • • • • • • • • • • • • • • •
Total (95% CI)		90		94	100.0%	0.50 [0.19, 1.32]	-
Total events	5		11				
Heterogeneity: Chi ² =	0.11, df = 3	2 (P = 0	.95); l² = l	0%			
Test for overall effect:	Z=1.40 (F	° = 0.16)				Favours endothermal Favours stripping

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Figure 102: Stripping surgery vs. endothermal ablation in primary VV: Physician reported disease severity (post-test; continuous)

	Endo	thern	nal	Str	ippin	g		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1.6.1 1 year									
Rass 2011	2	2	173	2.1	3	143	100.0%	-0.10 [-0.67, 0.47]	
Subtotal (95% CI)			173			143	100.0%	-0.10 [-0.67, 0.47]	
Heterogeneity: Not ap	plicable								
Test for overall effect	Z = 0.34	(P = ().73)						
1.6.2 2 years									
Rass 2011	2.1	3	173	1.9	3	143	100.0%	0.20 [-0.46, 0.86]	
Subtotal (95% CI)			173			143	100.0%	0.20 [-0.46, 0.86]	
Heterogeneity: Not ap	plicable								
Test for overall effect	Z = 0.59	(P = 0).56)						
									-2 -1 0 1 2

Favours endothermal Favours stripping

Figure 103: Stripping surgery vs. endothermal ablation in primary VV: Physician reported disease severity (change from baseline; continuous)

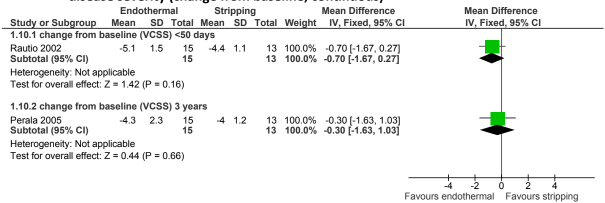


Figure 104: stripping surgery vs. endothermal ablation in primary VV: Physician reported disease severity (dichotomous)

								Dist Dist.	
	Endothe		Strippi			Risk Ratio		Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixed, 95% Cl	
1.12.1 1 week FU									
Lurie 2005	8	43	6	36	100.0%	1.12 [0.43, 2.92]			
Subtotal (95% CI)		43		36	100.0%	1.12 [0.43, 2.92]			
Total events	8		6						
Heterogeneity: Not ap	plicable								
Test for overall effect:	Z = 0.22 (P	9 = 0.82))						
1.12.4 1-2 year FU									
Lurie 2005	9	36	12	29	43.5%	0.60 [0.30, 1.23]		_ _	
Pronk 2010	17	56	18	61	56.5%	1.03 [0.59, 1.79]			
Subtotal (95% CI)		92		90	100.0%	0.84 [0.55, 1.30]		•	
Total events	26		30						
Heterogeneity: Chi ² =	1.33. df = 1	(P = 0.3)	25): l² = 2	5%					
Test for overall effect:	,	·							
		,							
								-++	-+
							0.05	0.2 1 5	20

Favours endothermal Favours stripping

Figure 105: Stripping surgery vs. endothermal ablation in primary VV: Physician reported disease severity (dichotomous) – asymptomatic according to VDS

	,						0
	endothe	rmal	stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Flassenkamper 2012	84	142	77	159	100.0%	1.22 [0.99, 1.51]	
Total (95% CI)		142		159	100.0%	1.22 [0.99, 1.51]	•
Total events	84		77				
Heterogeneity: Not appl							0.01 0.1 1 10 10
Test for overall effect: Z	= 1.86 (P =	= 0.06)					Favours stripping Favours endothern

Figure 106: Stripping surgery vs. endothermal ablation in primary VV: GSV reflux

1.9.1 0-12 weeks Carradice 2011 1 137 10 132 15.0% 0.10 [0.01, 0.74] Darwood 2008 4 71 4 32 19.7% 0.45 [0.12, 1.69] ElKaffas 2011 6 90 0 90 10.8% 13.00 [0.74, 227.39] Lurie 2003 5 43 0 36 10.8% 9.25 [0.53, 161.82] Rasmussen 2007 0 63 1 63 9.5% 0.33 [0.01, 8.03] Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rasmussen 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1.3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] 1.4 Lurie 2005 2 36 3 29		Endothe		Stripp	-		Risk Ratio	Risk Ratio
Carradice 2011 1 137 10 132 15.0% 0.10 [0.01, 0.74] Darwood 2008 4 71 4 32 19.7% 0.45 [0.12, 1.69] ElKaffas 2011 6 90 0 90 10.8% 13.00 [0.74, 227.39] Lurie 2003 5 43 0 36 10.8% 9.25 [0.53, 161.82] Rasmussen 2007 0 63 1 63 9.5% 0.33 [0.01, 8.03] Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subtramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtoal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: $Z = 1.18$ (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2017 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]		Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
Darwood 2008 4 71 4 32 19.7% 0.45 [0.12, 1.69] ElKaffas 2011 6 90 0 90 10.8% 13.00 [0.74, 227.39] Lurie 2003 5 43 0 36 10.8% 9.25 [0.53, 161.82] Rasmussen 2007 0 63 1 63 9.5% 0.33 [0.01, 8.03] Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subtrati (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); l ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1.3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2017 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]								
ElKaffas 2011 6 90 0 90 10.8% 13.00 [0.74, 227.39] Lurie 2003 5 43 0 36 10.8% 9.25 [0.53, 161.82] Rasmussen 2007 0 63 1 63 9.5% 0.33 [0.01, 8.03] Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subtramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	Carradice 2011	1	137	10	132	15.0%	0.10 [0.01, 0.74]	
Lurie 2003 5 43 0 36 10.8% 9.25 [0.53, 161.82] Rasmussen 2007 0 63 1 63 9.5% 0.33 [0.01, 8.03] Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtatal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	Darwood 2008	4	71	4	32	19.7%	0.45 [0.12, 1.69]	
Rasmussen 2007 0 63 1 63 9.5% 0.33 [0.01, 8.03] Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subtramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00]	ElKaffas 2011	6	90	0	90	10.8%	13.00 [0.74, 227.39]	
Rasmussen 2011 1 285 3 135 13.8% 0.16 [0.02, 1.50] Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Preala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 1.43 [0.48, 4.29]	Lurie 2003	5	43	0	36	10.8%	9.25 [0.53, 161.82]	
Rautio 2002 0 15 1 13 9.7% 0.29 [0.01, 6.60] Subramonia 2010 0 47 5 41 10.8% 0.08 [0.00, 1.40] Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 1.43 [0.48, 4.29] 4 Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] 4	Rasmussen 2007	0	63	1	63	9.5%	0.33 [0.01, 8.03]	
Subramonia 2010 0 47 5 41 10.8% 0.08 0.00, 1.40 Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); P = 52% Test for overall effect: $Z = 1.18$ (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 1.43 [0.48, 4.29] 4 Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	Rasmussen 2011	1	285	3	135	13.8%	0.16 [0.02, 1.50]	
Subtotal (95% CI) 751 542 100.0% 0.48 [0.14, 1.64] Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] •	Rautio 2002	0	15	1	13	9.7%	0.29 [0.01, 6.60]	
Total events 17 24 Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: $Z = 1.18$ (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.36, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	3ubramonia 2010	0		5	41	10.8%	0.08 [0.00, 1.40]	
Heterogeneity: Tau ² = 1.57; Chi ² = 14.63, df = 7 (P = 0.04); I ² = 52% Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	Subtotal (95% CI)		751		542	100.0%	0.48 [0.14, 1.64]	-
Test for overall effect: Z = 1.18 (P = 0.24) 1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] •	Total events	17		24				
1.9.2 1-3 years Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] •	Heterogeneity: Tau ² =	- 1.57; Chi ^z	= 14.63	3, df = 7 (8	P = 0.0	4); I ^z = 52	%	
Darwood 2008 6 49 1 12 8.1% 1.47 [0.19, 11.08] Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] •	fest for overall effect:	Z=1.18 (F	P = 0.24)				
Lurie 2005 2 36 3 29 11.2% 0.54 [0.10, 3.00] Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] •	1.9.2 1-3 years							
Perala 2005 3 15 0 15 4.0% 7.00 [0.39, 124.83] Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24] •	Darwood 2008	6	49	1	12	8.1%	1.47 [0.19, 11.08]	+-
Pronk 2010 5 49 5 56 23.9% 1.14 [0.35, 3.71] Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% Cl) 513 342 100.0% 1.26 [0.71, 2.24] Image: Content of the second s	_urie 2005	2	36	3	29	11.2%	0.54 [0.10, 3.00]	
Rasmussen 2007 1 50 3 53 6.7% 0.35 [0.04, 3.29] Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% Cl) 513 342 100.0% 1.26 [0.71, 2.24] Image: Constraint of the second	°erala 2005	3	15	0	15	4.0%	7.00 [0.39, 124.83]	
Rasmussen 2010 6 69 3 69 18.4% 2.00 [0.52, 7.68] Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	Pronk 2010	5	49	5	56	23.9%	1.14 [0.35, 3.71]	_ _
Rasmussen 2011 13 245 4 108 27.6% 1.43 [0.48, 4.29] • • • • • • • • • • • • • • • • • • •		1	50	3	53	6.7%	0.35 [0.04, 3.29]	
Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]	Rasmussen 2007		60	3	69	18.4%	2.00 [0.52, 7.68]	- +
Subtotal (95% CI) 513 342 100.0% 1.26 [0.71, 2.24]		6	09			27.00		
Total events 36 19	Rasmussen 2010	-		4	108	27.0%	1.40 [0.40] 4.20]	
	Rasmussen 2010 Rasmussen 2011	-	245	4				+
Heterogeneity: Tau ² = 0.00; Chi ² = 4.12, df = 6 (P = 0.66); I ² = 0%	Rasmussen 2010 Rasmussen 2011 Subtotal (95% CI)	13	245					+
Test for overall effect: Z = 0.78 (P = 0.44)	Rasmussen 2010 Rasmussen 2011 Subtotal (95% CI) Fotal events	13 36	245 <mark>513</mark>	19	342	100.0%		•

0.001 0.1 1 10 100 Favours endothermal Favours stripping

op pain	(dichoto	mous					
	Endothe	rmal	Strippi	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
1.11.1 Pain or tendern	ess within	72 houi	rs				
ElKaffas 2011	12	90	12	90	6.9%	1.00 [0.47, 2.11]	_
Flassenkamper 2012	4	142	6	159	3.3%	0.75 [0.21, 2.59]	
Lurie 2003	2	44	9	36	5.7%	0.18 [0.04, 0.79]	
Subramonia 2010 Subtotal (95% CI)	30	47 323	40	41 326	24.7% 40.7%	0.65 [0.52, 0.82] 0.65 [0.51, 0.85]	+ ♦
Total events	48		67				
Heterogeneity: Chi ² = 4	.21, df = 3 (P = 0.24	4); I ^z = 29 ⁴	%			
Test for overall effect: Z	:= 3.24 (P =	0.001)					
1.11.2 Post operative	pain at 7 da	iys					
Rass 2011	118	185	91	161	56.4%	1.13 [0.95, 1.34]	
Subtotal (95% CI)		185		161	56.4%	1.13 [0.95, 1.34]	•
Total events	118		91				
Heterogeneity: Not app							
Test for overall effect: Z	:= 1.36 (P =	: 0.17)					
1.11.3 persistent tend	erness (fol	low up i	not repor	ted)			
CarradiceA 2011	1	137	5	133	2.9%		
Subtotal (95% CI)		137		133	2.9%	0.19 [0.02, 1.64]	
Total events	1		5				
Heterogeneity: Not app	licable						
Test for overall effect: Z	:= 1.51 (P =	: 0.13)					
Total (95% CI)		645		620	100.0%	0.91 [0.79, 1.05]	•
Total events	167		163				
Heterogeneity: Chi ² = 2	1.28, df = 5	(P = 0.0	0007); I² =	: 77%			0.05 0.2 1 5 20
Test for overall effect: Z	:= 1.32 (P =	: 0.19)					Favours endothermal Favours stripping
Test for subgroup differ	rences: Chi	² = 13.9	19, df = 2 ((P = 0.0)009), I ^z =	: 85.7%	arears endoarennar i avoars suppling

Figure 107: Stripping surgery vs. endothermal ablation in primary VV: Adverse events – post op pain (dichotomous

Figure 108: Stripping surgery vs. endothermal ablation in primary VV: Adverse events - Post op pain (continuous variable)

	End	othern	nal	St	ripping	9		Mean Difference		Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C	:1	IV, Fixed, 95% CI	
1.25.1 1 day laser											
Pronk 2010 Subtotal (95% CI)	3.58	2.6	62 62	4	2.34	68 68	100.0% 100.0%				
Heterogeneity: Not ap	plicable										
Test for overall effect:	Z = 0.96	(P = 0).33)								
1.25.5 10-14days lase	er										
Pronk 2010	1.66	2.04	62	0.77	1.46	68	45.1%	0.89 [0.27, 1.51]		 −∎−	
Rasmussen 2011 Subtotal (95% CI)	2.58	2.41	143 205	2.25	2.23	123 191					
Heterogeneity: Chi ² =	1.75, df :	= 1 (P	= 0.19)	; l² = 43	8%						
Test for overall effect:	Z = 2.76	(P = 0	0.006)								
1.25.7 10-14 days RF											
Rasmussen 2011	1.21	1.72	146	2.25	2.23	123	70.7%	-1.04 [-1.52, -0.56]			
Rautio 2002	0.7	0.5	15	1.7	1.3	13		-1.00 [-1.75, -0.25]			
Subtotal (95% CI)			161			136	100.0%	-1.03 [-1.43, -0.62]		◆	
Heterogeneity: Chi ² = Test for overall effect:	,		,	,	6						
	2 4.00			,							
									++		
									-4 -2		
									Favours endo	othermal Favours strippin	ıg

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phiebi	us/unron	npobr	nepitis				
	Endothe	rmal	Stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
1.13.1 0-12 days							
CarradiceA 2011	4	137	6	133	43.3%	0.65 [0.19, 2.24]	
Darwood 2008	9	71	0	32	4.9%	8.71 [0.52, 145.20]	
ElKaffas 2011	6	90	0	90	3.6%	13.00 [0.74, 227.39]	
Rasmussen 2007	2	67	2	68	14.1%	1.01 [0.15, 7.00]	+
Rass 2011	20	185	4	161	30.4%	4.35 [1.52, 12.47]	— —
Rautio 2002 Subtotal (95% CI)	3	15 565	0	13 497	3.8% 100.0%	6.13 (0.35, 108.58) 2.86 [1.55, 5.29]	
Total events	44		12				_
Heterogeneity: Chi ² =	= 9.16. df=	5 (P = 0	10): I ² = -	45%			
Test for overall effect	•						
1.13.2 1 month to 3	10310						
Perala 2005	1	15		40	5 7 W	2 6 2 10 4 2 60 401	
Rasmussen 2007	2	15 65	0 2	13 66	5.7% 21.3%	2.63 [0.12, 59.40]	
Rasmussen 2007	16	285	2 5	135	72.9%	1.02 [0.15, 6.99] 1.52 [0.57, 4.05]	
Subtotal (95% CI)	10	365	5		100.0%	1.47 [0.64, 3.41]	
Total events	19	000	7	214	1001070	1111 [0101] 0111]	
Heterogeneity: Chi ² =		2 (P = 0	•	0%			
Test for overall effect				0.00			
		0.01	/				
							0.01 0.1 1 10 10
							Favours endothermal Favours stripping

Figure 109: Stripping surgery vs. endothermal ablation in primary VV: Adverse events – phlebitis/thrombophlebitis

Figure 110: Stripping surgery vs. endothermal ablation in primary VV: Adverse events – sensory deficits/neural injury

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	Endothe	rmal	Stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% Cl
1.14.1 0-12 weeks							
CarradiceA 2011	4	137	13	133	15.4%	0.30 [0.10, 0.89]	
Darwood 2008	1	71	4	32	6.4%	0.11 [0.01, 0.97]	<
ElKaffas 2011	9	90	3	90	3.5%	3.00 [0.84, 10.72]	+
Flassenkamper 2012	3	142	2	142	2.3%	1.50 [0.25, 8.84]	
Lurie 2003	10	44	5	36	6.4%	1.64 [0.61, 4.36]	- +
Pronk 2010	2	61	1	67	1.1%	2.20 [0.20, 23.62]	
Rasmussen 2007	1	65	0	66	0.6%	3.05 [0.13, 73.42]	
Rasmussen 2011	9	285	5	135	7.9%	0.85 [0.29, 2.50]	
Rass 2011	17	185	22	161	27.5%	0.67 [0.37, 1.22]	
Rautio 2002	2	15	3	13	3.8%	0.58 [0.11, 2.94]	
Stotter 2006	0	20	0	20		Not estimable	
Subramonia 2010	9	47	20	41	25.0%	0.39 [0.20, 0.76]	
Subtotal (95% CI)		1162		936	100.0%	0.71 [0.53, 0.97]	•
Total events	67		78				
Heterogeneity: Chi ² = 18	3.54, df = 1	0 (P = 0	1.05); I ^z = 1	46%			
Test for overall effect: Z :	= 2.18 (P =	: 0.03)					
1.14.2 >6 months							_
Perala 2005	1	15	5	13	64.2%	0.17 [0.02, 1.30]	
Pronk 2010	0	61	1	67	17.1%	0.37 [0.02, 8.81]	
Rasmussen 2007	0	54	1	50	18.7%	0.31 [0.01, 7.42]	
Subtotal (95% CI)		130		130	100.0%	0.23 [0.05, 1.02]	
Total events	1		7				
Heterogeneity: Chi ² = 0.1			1); I² = 0%	5			
Test for overall effect: Z	= 1.94 (P =	: 0.05)					
							0.05 0.2 1 5 20

0.05 0.2 1 5 20 Favours endothermal Favours stripping

Figure 111: Strip	oping su	rgery	vs. enc	lothe	rmal at	plation in prima	ry VV: Adverse events - DVI
	Endothe	rmal	Strippi	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
CarradiceA 2011	0	137	0	133		Not estimable	
ElKaffas 2011	0	90	1	90	27.0%	0.33 [0.01, 8.08]	
Flassenkamper 2012	1	142	1	159	17.0%	1.12 [0.07, 17.74]	
Pronk 2010	0	61	0	67		Not estimable	
Rasmussen 2011	0	285	1	135	36.7%	0.16 [0.01, 3.87]	← ■
Rass 2011	1	185	1	161	19.3%	0.87 [0.05, 13.80]	
Stotter 2006	0	20	0	20		Not estimable	
Total (95% CI)		920		765	100.0%	0.51 [0.13, 2.01]	
Total events	2		4				
Heterogeneity: Chi ² = 1.0	04, df = 3 (P = 0.79	9); I ² = 0%	5			
Test for overall effect: Z =	= 0.97 (P =	: 0.33)					Favours endothermal Favours stripping

Figure 111: Stripping surgery vs. endothermal ablation in primary VV: Adverse events - DVT

Figure 112: Stripping surgery vs. endothermal ablation in primary VV: Adverse events – limb discolouration

	Endothe	rmal	Stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
1.16.1 0-1 month							
CarradiceA 2011	4	137	1	133	28.7%	3.88 [0.44, 34.29]	
Darwood 2008	0	71	2	32	19.0%	0.09 (0.00, 1.86)	← ■
Lurie 2003	0	44	0	36		Not estimable	
Rasmussen 2011 Subtotal (95% CI)	11	285 537	6	135 336	52.2% 100.0%	0.87 [0.33, 2.30] 0.87 [0.18, 4.14]	
Total events	15		9				
Heterogeneity: Tau² = Test for overall effect:	•		4	= 0.14); I² = 49%	5	
1.16.2 3-4 months							
Lurie 2003	0	43	3	34	39.3%	0.11 [0.01, 2.13]	← ■ / · · · · · · · · · · · · · · · · · ·
Rass 2011 Subtotal (95% Cl)	57	185 228	19	161 195	60.7% 100.0%	2.61 [1.62, 4.20] 0.76 [0.04, 16.04]	-
Total events	57		22				
Heterogeneity: Tau ² = Test for overall effect:	•			= 0.04); I² = 77%	5	

0.01 0.1 1 10 10 Favours endothermal Favours stripping

Figure 113: Stripping surgery vs. endothermal ablation in primary VV: Return to normal activities by endothermal type

activit	les by	enu	other	mai t	ype					
	Endo	otherm	nal	Sti	ripping)		Mean Difference	Mean Dif	ference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% C	I IV, Fixed,	, 95% CI
1.20.1 RF										
ElKaffas 2011	3	3	90	7	2.6	90	62.4%	-4.00 [-4.82, -3.18] 📕	
Lurie 2003	1.36	0.92	44	6.65	6.76	36	8.5%	-5.29 [-7.51, -3.07		
Subtotal (95% CI)			134			126	70.8%	-4.15 [-4.92, -3.38]	♦	
Heterogeneity: Chi ² =	1.14, df	= 1 (P	= 0.29)); I² = 1 2	%					
Test for overall effect:	Z = 10.5	i8 (P <	0.0000	01)						
1.20.2 Laser										
Pronk 2010	3.2	4.3	62	3.2	4	68	20.5%	0.00 [-1.43, 1.43] –	-
Rasmussen 2007	6.9	7	69	7.7	6.1	68	8.7%	-0.80 [-3.00, 1.40] _+	_
Subtotal (95% CI)			131			136	29.2%	-0.24 [-1.44, 0.96	i 🔶	>
Heterogeneity: Chi ² =	0.36, df	= 1 (P	= 0.55)); I ^z = 0%	6					
Test for overall effect:	Z = 0.39) (P = 0	0.70)							
Total (95% CI)			265			262	100.0%	-3.01 [-3.66, -2.36]	↓ ♦	
Heterogeneity: Chi ² =	30.50, d	lf = 3 (l	P < 0.0	0001); P	'= 90%	6				
Test for overall effect:	Z= 9.12	(P < 0	0.00001	0					-10 -5 Ó	, 0 10
Test for subgroup diff	erences	Chi ≇⊧	= 29.00), df = 1	(P < 0.	00001), I ² = 96.6	i%	Favours endothermal	ravours surpping

Figure 114: Stripping surgery vs. endothermal ablation in primary VV: Return to work by endothermal type

endo	tnerm	аі тур	e						
	End	otherm	al	St	tripping			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1.22.1 RF									
Lurie 2003	4.7	11.86	44	12.4	11.59	36	4.7%	-7.70 [-12.86, -2.54]	
Rautio 2002	6.5	3.3	15	15.6	6	13	9.3%	-9.10 [-12.76, -5.44]	
Subtotal (95% CI)			59			49	14.1%	-8.63 [-11.62, -5.64]	◆
Heterogeneity: Chi ² =	: 0.19, df	= 1 (P =	0.66);	I² = 0%					
Test for overall effect	: Z = 5.66	6 (P ≺ 0.	00001)						
1.22.2 Laser									
Pronk 2010	4.4	5.4	62	4.2	3.7	68	48.6%	0.20 [-1.41, 1.81]	+
Rasmussen 2007	7	6	69	7.6	4.9	68	37.3%	-0.60 [-2.43, 1.23]	
Subtotal (95% CI)			131			136	85.9%	-0.15 [-1.36, 1.06]	◆
Heterogeneity: Chi ² =	: 0.41, df	= 1 (P =	0.52);	l² = 0%					
Test for overall effect	: Z = 0.24	(P = 0.	81)						
Total (95% CI)			190			185	100.0%	-1.34 [-2.46, -0.22]	◆
Heterogeneity: Chi ² =	: 27.22, c	lf = 3 (P	< 0.00	001); P=	= 89%				
Test for overall effect									-10 -5 0 5 10
Test for subaroup dif			r .	df = 1 (F	- ⊂ 0.00	1001), P	² = 96.2%		Favours endothermal Favours stripping

2

3 I.3.2.2 Recurrent varicose veins

Figure 115: Stripping surgery vs. endothermal ablation in recurrent VV: GSV reflux – 6 weeks

U	Endotherma	al (RF)	Stripp	ing		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Hinchcliffe 2006	3	16	2	16	100.0%	1.50 [0.29, 7.81]	
Total (95% CI)		16		16	100.0%	1.50 [0.29, 7.81]	
Total events	3		2				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 0.48 (P = 1	0.63)					Favours endothermal Favours stripping

Figure 116: Stripping surgery vs. endothermal ablation in recurrent VV:Adverse events – thrombophlebitis (6 weeks)

	Endotherma	Stripp	ing		Risk Ratio		Risk Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H	l, Fixed, 95	% CI	
Hinchcliffe 2006	0	16	1	16	100.0%	0.33 [0.01, 7.62]				
Total (95% CI)		16		16	100.0%	0.33 [0.01, 7.62]				
Total events	0		1							
Heterogeneity: Not ap	pplicable						0.01 0.1			10
Test for overall effect	: Z = 0.69 (P =	0.49)				Far	vours endothermal	(RF) Favo		

Figure 117: Stripping surgery vs. endothermal ablation in recurrent VV:Adverse events – sensory deficits / neural injury (neuralgia and numbness at 6 weeks)

561156	.,	/	· • · · · · · · · · · · · · · · · · · ·				
	Endotherma	Stripp	ing		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI
Hinchcliffe 2006	2	16	3	16	100.0%	0.67 [0.13, 3.47	
Total (95% CI)		16		16	100.0%	0.67 [0.13, 3.47]	
Total events	2		3				
Heterogeneity: Not ap Test for overall effect:	•	0.63)				F	0.01 0.1 1 10 10 avours endothermal (RF) Favours stripping

Figure 118: Stripping surgery vs. endothermal ablation in recurrent VV: Adverse events – infection (6 weeks)

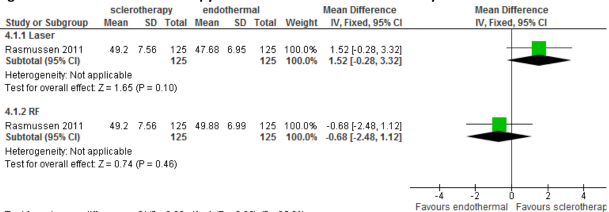
		,							
	Endotherma	Stripp	ing		Risk Ratio	Risk Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixe	ed, 95% CI	
Hinchcliffe 2006	0	16	1	16	100.0%	0.33 [0.01, 7.62]			
Total (95% CI)		16		16	100.0%	0.33 [0.01, 7.62]			
Total events	0		1						
Heterogeneity: Not ap Test for overall effect:	•	0.49)				Fa	0.01 0.1 vours endothermal (RF)	1 10 Favours stripping	10 g

Figure 119: Stripping surgery vs. endothermal ablation in recurrent VV: Adverse events -

oeden	าล								
	Endotherm	al (RF)	Stripp	ing		Risk Ratio	R	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	М-Н,	Fixed, 95% CI	
Hinchcliffe 2006	0	16	1	16	100.0%	0.33 [0.01, 7.62]			
Total (95% CI)		16		16	100.0%	0.33 [0.01, 7.62]			
Total events	0		1						
Heterogeneity: Not ap	plicable						0.01 0.1		0 10
Test for overall effect: .	Z = 0.69 (P =	0.49)				Fa	avours endothermal (f		

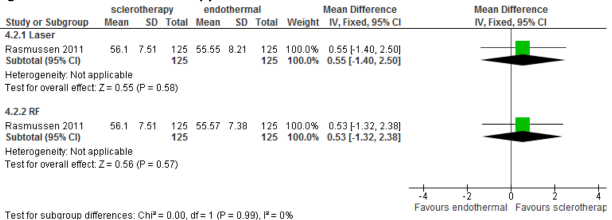
1.3.3 Foam sclerotherapy vs. endothermal ablation 1

Figure 120: Foam sclerotherapy vs. endothermal ablation: SF-36 Physical 4 weeks



Test for subgroup differences: $Chi^2 = 2.86$, df = 1 (P = 0.09), $l^2 = 65.0\%$

Figure 121: Foam sclerotherapy vs. endothermal ablation: SF-36 mental 4 weeks



Foam sclerotherapy vs. endothermal ablation: SF-36 Physical 1 year Figure 122:

-	scler	othera	ару	endo	otherm	nal		Mean Difference	Mean Difference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI		
4.3.1 Laser											
Rasmussen 2011 Subtotal (95% CI)	51.93	7.66	125 125	52.62	5.98	125 125		-0.69 [-2.39, 1.01] - 0.69 [-2.39, 1.01]			
Heterogeneity: Not a	pplicable										
Test for overall effect	: Z = 0.79	(P = 0	.43)								
4.3.2 RF									_		
Rasmussen 2011 Subtotal (95% CI)	51.93	7.66	125 125	53.23	5.32			-1.30 [-2.93, 0.33] - 1.30 [-2.93, 0.33]			
Heterogeneity: Not a	pplicable										
Test for overall effect	:Z=1.56	i (P = 0	.12)								
									-2 -1 0 1 2		
									Favours endothermal Favours sclerother		

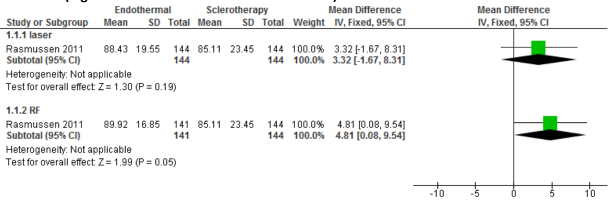
Test for subgroup differences: $Chi^2 = 0.26$, df = 1 (P = 0.61), l² = 0%

Figure 123:	Foam	scier	othe	rapy v	/s. er	ndotr	hermai	ablation: SF-36	o mental 1 year
	scler	rothera	ару	endo	otherm	nal		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
4.4.1 Laser									
Rasmussen 2011 Subtotal (95% CI)	54.73	8.89	125 125	56.74	5.44		100.0% 100.0%	-2.01 [-3.84, -0.18] - 2.01 [-3.84, -0.18]	
Heterogeneity: Not a	applicable								
Test for overall effec	t: Z = 2.16	6 (P = 0	1.03)						
4.4.2 RF									
Rasmussen 2011 Subtotal (95% CI)	54.73	8.89	125 125	56.52	6.17	125 125	100.0% 100.0%	-1.79 [-3.69, 0.11] - 1.79 [-3.69, 0.11]	
Heterogeneity: Not a	applicable	1							
Test for overall effec	t: Z = 1.85	5 (P = 0	1.06)						
									-4 -2 0 2 4
Test for subgroup d	ifforoncoc	- Chiž-	- 0 0 2	df = 1 /0		7) 12 - 1	nov.		Favours endothermal Favours sclerotherap

Figure 123: Foam sclerotherapy vs. endothermal ablation: SF-36 mental 1 year

Test for subgroup differences: $Chi^2 = 0.03$, df = 1 (P = 0.87), $I^2 = 0\%$

Figure 124: Foam sclerotherapy vs. endothermal ablation: Pain due to varicose veins (1 year) (higher better as taken from SF-36 sub-scale)



Favours sclerotherapy Favours endothermal

Figure 125: Foam sclerotherapy vs. endothermal ablation: Reflux above knee at 3 days

0									,
	Sclerotherapy		Endothe	rmal		Peto Odds Ratio		Peto	Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% Cl		Peto, Fixed, 95% CI	
1.2.1 laser									
Rasmussen 2011 Subtotal (95% CI)	3	143 143	0	143 143	100.0% 100.0%	7.49 [0.77, 72.62] 7.49 [0.77, 72.62]			
Total events Heterogeneity: Not ap Test for overall effect: .	•	= 0.08)	0						
1.2.2 RF									
Rasmussen 2011 Subtotal (95% CI)	3	143 143	0	146 146	100.0% 100.0%	7.65 [0.79, 74.17] 7.65 [0.79, 74.17]			
Total events Heterogeneity: Not ap Test for overall effect: .	•	= 0.08)	0						
							L	0.1	

Favours sclerotherapy Favours endothermal

Figure 126: Foam sclerotherapy vs. endothermal ablation (Laser):Reflux above knee at 3-4 weeks

	Sclerothe	erapy	Endothe	rmal		Risk Ratio	Risk Ratio		
Study or Subgroup	Events	nts Total Events Total		Weight	Weight IV, Fixed, 95% CI		IV, Fixed, 95% CI		
1.3.1 laser									
Lattimer 2012	8	50	1	50	57.8%	8.00 [1.04, 61.62]			
Rasmussen 2011	2	144	1	144	42.2%	2.00 [0.18, 21.81]			
Subtotal (95% CI)		194		194	100.0%	4.46 [0.94, 21.04]			
Total events	10		2						
Heterogeneity: Chi ² =	= 0.75, df = 1	(P = 0.)	39); I^z = 09	6					
Test for overall effect	t: Z = 1.89 (P	= 0.06)							
							+		<u> </u>
							0.05	0.2	1 5

320

Favours sclerotherapy Favours endothermal

Varicose Veins Full Guideline Appendices (July 2013)

Figure 127: Foam sclerotherapy vs. endothermal ablation (RF): Reflux above knee at 1 month

	Sclerothe	erapy	Endothe	rmal		Peto Odds Ratio	Peto Odds Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI		
1.4.2 RF									
Rasmussen 2011 Subtotal (95% CI)	2	144 144	0	141 141	100.0% 100.0%	7.29 [0.45, 117.10] 7.29 [0.45, 117.10]			
Total events Heterogeneity: Not ap Test for overall effect:	•	= 0.16)	0						
							0.01 0.1 1 10 10 Favours sclerotherapy Favours endothermal		

Figure 128: Foam sclerotherapy vs. endothermal ablation(Laser): Reflux above knee at 3 months

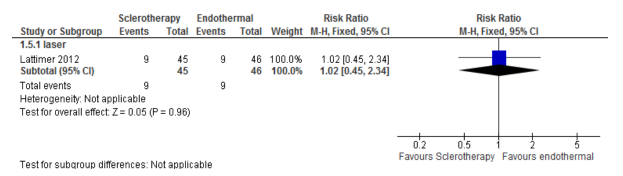


Figure 129: Foam sclerotherapy vs. endothermal ablation: Reflux above knee 1 year

	Sclerothe	erapy			Risk Ratio		Risk Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fix	ed, 95% Cl	
1.6.1 laser										
Rasmussen 2011 Subtotal (95% CI)	20	123 123	7	121 121	100.0% 100.0%	2.81 [1.23, 6.40] 2.81 [1.23, 6.40]				
Total events Heterogeneity: Not ap Test for overall effect:	•	= 0.01)	7							
1.6.2 RF										
Rasmussen 2011 Subtotal (95% CI)	20	123 123	6	124 124	100.0% 100.0%	3.36 [1.40, 8.08] 3.36 [1.40, 8.08]				
Total events Heterogeneity: Not ap Test for overall effect:	•	= 0.007	6 7)							
							0.01	0.1	 110	10

Favours sclerotherapy Favours endothermal

	Sclerothe	rapy	Endothe	rmal		Risk Ratio	Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixe	ed, 95% Cl	
Gonzalez-Zeh 2008	8	53	1	45	100.0%	6.79 [0.88, 52.27]	-		
Total (95% CI)		53		45	100.0%	6.79 [0.88, 52.27]			
Total events	8		1						
Heterogeneity: Not ap Test for overall effect:	•	= 0.07)					0.01 0.1 Favours sclerotherapy	1 10 Favours enothe	10 ermal

Figure 130: Foam sclerotherapy vs. endothermal ablation: Reflux at 1 year observational data

Figure 131: Foam sclerotherapy vs. endothermal ablation: Reflux below knee 3 weeks

	scleroth	егару	Controlendot	hermal		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
1.7.1 laser							
Lattimer 2012 Subtotal (95% CI)	24	50 50	7	50 <mark>50</mark>	100.0% 100.0%	3.43 [1.63, 7.22] 3.43 [1.63, 7.22]	-
Total events Heterogeneity: Not a Test for overall effect		P = 0.001	7				
Test for subgroup dif	fferences: N	lot appli	cable				0.01 0.1 1 10 10 Favours sclerotherapy Favours endotherma

Figure 132: Foam sclerotherapy vs. endothermal ablation: Reflux below knee 3 months

	sclerothe	егару	endothe	ermal		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% CI
1.8.1 laser							
Lattimer 2012 Subtotal (95% CI)	19	45 45	21	46 46	100.0% 100.0%	0.92 [0.58, 1.47] 0.92 [0.58, 1.47]	
Total events Heterogeneity: Not aj Test for overall effect	•	P = 0.74)	21				
Test for subgroup dif	ferences: N	lot appli	cable				0.5 0.7 1 1.5 2 Favours sclerotherapy Favours endotherma

Figure 133: Foam sclerotherapy vs. endothermal ablation: Adverse events – pain (VAS) observational data

	Scierc	othera	ру	Endo	therm	nal		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Gonzalez-Zeh 2008	4	1.5	53	4.9	1.5	45	100.0%	-0.90 [-1.50, -0.30]	
Total (95% CI)			53			45	100.0%	-0.90 [-1.50, -0.30]	· · · · · · · · · · · · · · · · · · ·
Heterogeneity: Not ap Test for overall effect:	•	(P = 0	.003)						-4 -2 0 2 4 Favours sclerotherapy Favours endotherma

Figure 134:Foam sclerotherapy vs. endothermal ablation: Adverse events – post op pain 10days

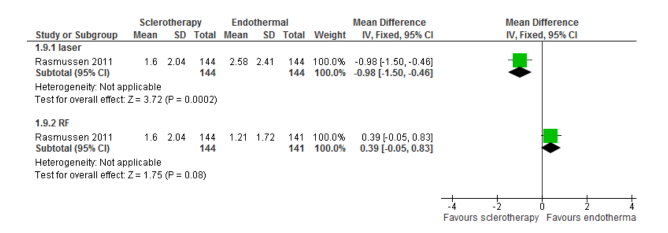


Figure 135: Foam sclerotherapy vs. endothermal ablation: Adverse events – DVT observational data

	Sclerothe	егару	Endothe	ermal		Peto Odds Ratio	Peto Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% Cl	Peto, Fixed, 95% Cl
Gonzalez-Zeh 2008	2	53	0	45	100.0%	6.48 [0.40, 106.04]	
Total (95% CI)		53		45	100.0%	6.48 [0.40, 106.04]	
Total events	2		0				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z=1.31 (P	= 0.19)					Favours sclerotherapy Favours endothermal

Figure 136: Foam sclerotherapy vs. endothermal ablation: Adverse events – neural iniurv/damage

i i i jui y	7 uuniugt	•					
	Sclerothe	erapy	Endothe	rmal		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% Cl
1.11.1 laser							
Rasmussen 2011 Subtotal (95% CI)	2	144 144	3	144 144	100.0% 100.0%	0.67 [0.11, 3.93] 0.67 [0.11, 3.93]	
Total events	2		3				
Heterogeneity: Not a	pplicable						
Test for overall effect	• •	= 0.65)					
		,					
1.11.2 RF							
Rasmussen 2011	2	144	6	141	100.0%	0.33 [0.07, 1.59]	
Subtotal (95% CI)		144		141	100.0%	0.33 [0.07, 1.59]	
Total events	2		6				
Heterogeneity: Not a	pplicable						
Test for overall effect	: Z = 1.39 (P	= 0.17)					
							0.01 0.1 1 10 10
							Favours sclerotherapy Favours endothermal

Figure 137: Foam sclerotherapy vs. endothermal ablation: Adverse events – paraesthesia observational data

	Sclerothe	erapy	Endothe	rmal		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Gonzalez-Zeh 2008	1	53	2	45	100.0%	0.42 [0.04, 4.53]	
Total (95% CI)		53		45	100.0%	0.42 [0.04, 4.53]	
Total events	1		2				
Heterogeneity: Not ap Test for overall effect:	•	= 0.48)					0.01 0.1 1 10 100 Favours sclerotherapy Favours endothermal

Figure 138: Foam sclerotherapy vs. endothermal ablation: Adverse events PE

	Sclerothe	егару	Endothe	rmal		Peto Odds Ratio	Peto Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI
1.12.1 RF							
Rasmussen 2011 Subtotal (95% Cl)	1	144 <mark>144</mark>	0	141 141	100.0% 100.0%	7.24 [0.14, 364.79] 7.24 [0.14, 364.79]	
Total events	1		0				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z=0.99 (P	= 0.32)					
1.12.2 laser							
Rasmussen 2011 Subtotal (95% CI)	1	144 <mark>144</mark>	0	144 144	100.0% 100.0%	7.39 [0.15, 372.38] 7.39 [0.15, 372.38]	
Total events	1		0				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z=1.00 (P	= 0.32)					
							Favours sclerotherapy Favours endotherma

Figure 139: Foam sclerotherapy vs. endothermal ablation: Adverse events – Phlebitis

	Sclerothe	rapy	Endothe	rmal		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
1.13.1 laser							
Rasmussen 2011 Subtotal (95% CI)	17	144 144	4	144 144	100.0% 100.0%	4.25 [1.47, 12.32] 4.25 [1.47, 12.32]	
Total events	17		4				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z=2.66 (P	= 0.008	i)				
1.13.2 RF							
Rasmussen 2011 Subtotal (95% CI)	17	144 144	12	141 141	100.0% 100.0%	1.39 [0.69, 2.80] 1.39 [0.69, 2.80]	
Total events	17		12				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z=0.91 (P	= 0.36)					
							Favours sclerotherapy Favours endothern

Figure 140: Foam sclerotherapy vs. endothermal ablation: Adverse events – Phlebitis observational data

	Sclerothe	егару	Endothe	ermal		Risk Ratio	Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fix	ed, 95% CI	
Gonzalez-Zeh 2008	22	53	10	45	100.0%	1.87 [0.99, 3.52]			
Total (95% CI)		53		45	100.0%	1.87 [0.99, 3.52]		◆	
Total events	22		10						
Heterogeneity: Not ap Test for overall effect:		= 0.05)					0.01 0.1 Favours sclerotherapy	1 10 Favours end	10 Iotherma

Figure 141: Foam sclerotherapy vs. endothermal ablation: Adverse events – hyperpigmentation

pigme	entation										
	Scieroth	erapy	Endothe	rmal		Risk Ratio			Risk Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H	I, Fixed, 95%	CI	
1.14.1 laser											
Rasmussen 2011 Subtotal (95% CI)	8	144 144	3	144 144	100.0% 100.0%	2.67 [0.72, 9.85] 2.67 [0.72, 9.85]					
Total events	8		3								
Heterogeneity: Not a	pplicable										
Test for overall effect	: Z=1.47 (F	9 = 0.14)									
1.14.2 RF											
Rasmussen 2011	8	144	8	141	100.0%	0.98 [0.38, 2.54]					
Subtotal (95% CI)		144		141	100.0%	0.98 [0.38, 2.54]					
Total events	8		8								
Heterogeneity: Not a	pplicable										
Test for overall effect	: Z = 0.04 (F	e 0.97)									
							0.01	0.1		10	10

Favours sclerotherapy Favours endothermal

Figure 142: Foam sclerotherapy vs. endothermal ablation(Laser): Need for further treatment

	scleroth	егару	endothe	ermal		Risk Ratio	Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixe	d, 95% Cl	
Lattimer 2012	28	50	3	50	100.0%	9.33 [3.03, 28.73]			
Total (95% CI)		50		50	100.0%	9.33 [3.03, 28.73]			
Total events	28		3						
Heterogeneity: Not ap Test for overall effect:	•	P < 0.000	01)				0.01 0.1 Favours sclerotherapy	1 10 Favours endothe	100 rmal

1 I.3.4 Truncal treatment with tributary treatment vs. truncal treatment alone

Figure 143: Truncal + tributary vs. truncal alone: Reflux at one week endo + tributary endo alone Risk Ratio **Risk Ratio** Total Events Total Weight M-H, Fixed, 95% Cl M-H, Fixed, 95% Cl Study or Subgroup Events 1.1.1 SFJ 1 week Carradice 2009 0 24 0 Not estimable 24 Subtotal (95% CI) 24 24 Not estimable Total events 0 0 Heterogeneity: Not applicable Test for overall effect: Not applicable 1.1.2 GSV 1 week Carradice 2009 0 Not estimable 24 **24** 0 24 **24** Subtotal (95% CI) Not estimable 0 Total events 0 Heterogeneity: Not applicable Test for overall effect: Not applicable 0.01 100 0.1 10 Favours endo + tributary Favours endo alone

Figure 144: Truncal + tributary vs. truncal alone: Adverse events - phlebitis

	endo + trib	utary	endo al	one		Risk Ratio	Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fix	ed, 95% Cl	
Carradice 2009	0	24	1	24	100.0%	0.33 [0.01, 7.80]			
Total (95% CI)		24		24	100.0%	0.33 [0.01, 7.80]			
Total events	0		1						
Heterogeneity: Not ap Test for overall effect:		0.49)				F	0.01 0.1 avours endo + tributary	1 10 Favours endo alone	100 e

Figure 145: Truncal + tributary vs. truncal alone: Adverse events - pigmentation

-	endo + trib	utary	endo al	lone		Peto Odds Ratio		Peto Od	ds Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% Cl		Peto, Fixe	ed, 95% Cl	
Carradice 2009	2	24	0	24	100.0%	7.72 [0.47, 127.14]				
Total (95% CI)		24		24	100.0%	7.72 [0.47, 127.14]				
Total events	2		0							
Heterogeneity: Not a	•						0.005	0.1 1	10	20
Test for overall effect	: Z = 1.43 (P =	= 0.15)					Favours en	do + tributary	Favours endo	alone

Source: <Insert Source text here>

endo + tributary endo alone Peto Odds Ratio Peto Odds Ratio Total Events Total Weight Study or Subgroup Events Peto, Fixed, 95% CI Peto, Fixed, 95% CI Carradice 2009 1 24 0 24 100.0% 7.39 [0.15, 372.38] Total (95% CI) 24 100.0% 7.39 [0.15, 372.38] 24 Total events 0 1 Heterogeneity: Not applicable 0.01 100 0.1 10 Test for overall effect: Z = 1.00 (P = 0.32) Favours endo + tributary Favours endo alone

Figure 146: Truncal + tributary vs. truncal alone: Adverse events - thigh neuralgia

Truncal + tributary vs. truncal alone: Need for ambulatory phlebectomy at 6 weeks Figure 147:

	endo + trib	utary	endo al	one		Risk Ratio			Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C			M-H, Fix	ed, 95% C		
Carradice 2009	1	25	16	24	100.0%	0.06 [0.01, 0.42]	•					
Total (95% CI)		25		24	100.0%	0.06 [0.01, 0.42]						
Total events	1		16									
Heterogeneity: Not ap Test for overall effect:		0.005)				F	0.01 avours	0.1 endo +	tributary	l 1 Favours	10 endo a	100 Ilone

1

Chapter 10 – compression after interventional treatment 1.4 2

3 1.4.1.1 Compression after surgery vs. surgery alone

Figure 148: Compression after interventional treatment vs. interventional treatment alone: adverse events - post operative pain (higher worse)

		P			~ ~	···· /·			
	surg v	vith co	mp	surg	g alon	ie		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1.1.1 3 days									
Houtermans-Auckel 2009	2.5	2.8	46	1.8	2.2	50	100.0%	0.70 [-0.31, 1.71]	-+
Subtotal (95% CI)			46			50	100.0%	0.70 [-0.31, 1.71]	-
Heterogeneity: Not applicab	le								
Test for overall effect: Z = 1.3	35 (P = 0.	18)							
1.1.2 2 weeks									
Houtermans-Auckel 2009	2.2	2.3	46	2.2	2.4	50	100.0%	0.00 [-0.94, 0.94]	
Subtotal (95% CI)			46			50	100.0%	0.00 [-0.94, 0.94]	-
Heterogeneity: Not applicab	le								
Test for overall effect: Z = 0.0	00 (P = 1.	00)							
1.1.3 4 weeks									
Houtermans-Auckel 2009	0.8	1.5	46	0.5	0.8	50	100.0%	0.30 [-0.19, 0.79]	
Subtotal (95% CI)			46			50	100.0%	0.30 [-0.19, 0.79]	
Heterogeneity: Not applicab	le								
Test for overall effect: Z = 1.3	21 (P = 0.	23)							

-2 ά 2 -1 ł Favours surg + comp Favours surg alone

Figure 149: Compression after interventional treatment vs. interventional treatment alone: adverse events - numbness

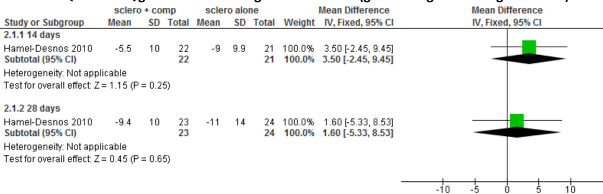
	surg with c	omp	surg al	one		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
1.2.1 3 days							
Houtermans-Auckel 2009	4	46	3		100.0%	1.45 [0.34, 6.13]	
Subtotal (95% CI)		46		50	100.0%	1.45 [0.34, 6.13]	
Total events	4		3				
Heterogeneity: Not applicable	Э						
Test for overall effect: Z = 0.5	0 (P = 0.61)						
1.2.2 2 weeks							
Houtermans-Auckel 2009	0	46	2	50	100.0%	0.22 [0.01, 4.40]	
Subtotal (95% CI)		46		50	100.0%	0.22 [0.01, 4.40]	
Total events	0		2				
Heterogeneity: Not applicable	Э						
Test for overall effect: Z = 0.9	9 (P = 0.32)						
1.2.3 4 weeks							
Houtermans-Auckel 2009	0	46	0	50		Not estimable	
Subtotal (95% CI)	_	46	_	50		Not estimable	
Total events	0		0				
Heterogeneity: Not applicable							
Test for overall effect: Not app	olicable						
							0.01 0.1 1 10 10
							Favours surg with comp Favours surg alone

Figure 150: Compression after interventional treatment vs. interventional treatment alone: return to work (days)

		•							
	surg w	vith co	mp	surg	g alor	ne		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Houtermans-Auckel 2009	15	8.4	46	11	7.5	50	100.0%	4.00 [0.80, 7.20]	
Total (95% CI)			46			50	100.0%	4.00 [0.80, 7.20]	
Heterogeneity: Not applicab Test for overall effect: Z = 2		0.01)						Fi	-4 -2 0 2 4 avours surg with comp Favours surg only

1 I.4.1.2 Compression after foam sclerotherapy vs. foam sclerotherapy alone

Figure 151: Compression after interventional treatment vs. interventional treatment alone: QoL – CIVIQ global score – change from baseline (greater negative change is better)



Favours sclero plus comp Favours sclero alone

Figure 152: Compression after interventional treatment vs. interventional treatment alone: reflux at 28 days

	····/	-								
	sclero + (comp	sclero a	lone		Risk Ratio		Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixe	d, 95% Cl	
Hamel-Desnos 2010	0	31	0	29		Not estimable				
Total (95% CI)		31		29		Not estimable				
Total events	0		0							
Heterogeneity: Not app Test for overall effect: N		le					0.01 Favours	0.1 sclero + comp	10 Favours scle	10 ro alone

Figure 153: Compression after interventional treatment vs. interventional treatment alone: adverse events – maior neurological events

auterst			or nea	. 0.08.	cui cre					
	sclero + (comp	sclero a	lone		Risk Ratio		Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixe	d, 95% Cl	
Hamel-Desnos 2010	0	31	0	29		Not estimable				
Total (95% CI)		31		29		Not estimable				
Total events	0		0							
Heterogeneity: Not app Test for overall effect: N		le					0.01 Favours s	0.1 clero + comp	10 Favours scle	10 ero alone

Figure 154: Compression after interventional treatment vs. interventional treatment alone:

adverse events – visual disturbance (scotoma) resolving within 15 minutes

	sclero + o	comp	sclero a	lone		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% CI
Hamel-Desnos 2010	0	31	1	29	100.0%	0.31 [0.01, 7.38]	
Total (95% CI)		31		29	100.0%	0.31 [0.01, 7.38]	
Total events	0		1				
Heterogeneity: Not ap Test for overall effect:		= 0.47)					0.01 0.1 1 10 100 Favours sclero + comp Favours sclero alone

Figure 155: Compression after interventional treatment vs. interventional treatment alone: adverse events – moderate pain day 28

		•• ••					
	sclero +	comp	sclero a	lone		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% CI
Hamel-Desnos 2010	1	30	3	29	100.0%	0.32 [0.04, 2.92]	
Total (95% CI)		30		29	100.0%	0.32 [0.04, 2.92]	
Total events	1		3				
Heterogeneity: Not app	plicable						0.01 0.1 1 10 100
Test for overall effect:	Z = 1.01 (P	= 0.31)					0.01 0.1 1 10 100 Favours sclero + comp Favours sclero alone

Figure 156: Compression after interventional treatment vs. interventional treatment alone: adverse events - pigmentation

auver	Se even	ιs - μι	ginenta	ation			
	sclero + o	comp	sclero a	lone		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Hamel-Desnos 2010	2	30	1	29	100.0%	1.93 [0.19, 20.18]	
Total (95% CI)		30		29	100.0%	1.93 [0.19, 20.18]	
Total events	2		1				
Heterogeneity: Not app	olicable						0.01 0.1 1 10 100
Test for overall effect:	Z = 0.55 (P	= 0.58)					Favours sclero + comp Favours sclero alone

Figure 157: Compression after interventional treatment vs. interventional treatment alone: adverse events - thrombophlebitis

				•			
	sclero + o	comp	sclero a	lone		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Hamel-Desnos 2010	3	30	3	29	100.0%	0.97 [0.21, 4.41]	
Total (95% CI)		30		29	100.0%	0.97 [0.21, 4.41]	
Total events	3		3				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 0.04 (P	= 0.97)					0.01 0.1 1 10 100 Favours sclero + comp Favours sclero alone

Figure 158: Compression after interventional treatment vs. interventional treatment alone:

-	nt assess sclero + o	-	sclero al			Risk Ratio	Risk Ratio
Study or Subgroup	Events		Events		Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% Cl
2.8.1 heavy legs							
Hamel-Desnos 2010 Subtotal (95% CI)	20	30 30	16	29 29	100.0% 100.0%	1.21 [0.80, 1.83] 1.21 [0.80, 1.83]	
Total events	20		16				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 0.90 (P	= 0.37)					
2.8.2 pain							
Hamel-Desnos 2010 Subtotal (95% CI)	21	30 30	17		100.0% 100.0%	1.19 [0.81, 1.76] 1.19 [0.81, 1.76]	
Total events	21		17				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 0.90 (P	= 0.37)					
2.8.3 oedema							
Hamel-Desnos 2010 Subtotal (95% CI)	15	30 30	15		100.0% 100.0%	0.97 [0.59, 1.60] 0.97 [0.59, 1.60]	
Total events	15		15				
Heterogeneity: Not ap	plicable						
Test for overall effect:	Z = 0.13 (P	= 0.89)					
2.8.4 paraesthesia							
Hamel-Desnos 2010	17	30	13	29	100.0%	1.26 [0.76, 2.11]	
Subtotal (95% CI)		30		29	100.0%	1.26 [0.76, 2.11]	
Total events	17		13				
Heterogeneity: Not app							
Test for overall effect:	Z = 0.90 (P	= 0.37)					
2.8.5 cramp							_
Hamel-Desnos 2010 Subtotal (95% CI)	11	30 30	16	29 29	100.0% 100.0%	0.66 [0.37, 1.18] 0.66 [0.37, 1.18]	
Total events	11		16				
Heterogeneity: Not app	plicable						
Test for overall effect:	Z = 1.40 (P	= 0.16)					
	,	,					

0.5 0.7 1 1.5 2 Favours scler + comp Favours scler alone

Appendix J: Excluded clinical studies

1 2

J.1 Chapter 5 – patient perceptions and expectations

4

3

Table 98: Studies excluded from the clinical review for chapter 5

Reference	Reason for exclusion
Bradshaw 1999 ³¹	Does not match the review question
Ching 2010 ⁵⁶	Only covered new Zealand-based websites and so not relevant to the UK
Davies 199568	Does not match the review question
Murphy 2001 ¹⁷⁹	This paper was deemed too old for its survey of internet information to have any continued relevance.
Sains 2005 ²³⁵	Does not match the review question
Scurr 2008 ²⁴³	Does not match the review question
Srilekha 2005 ²⁵⁴	Does not match the review question

5

6 J.2 Chapter 6 – referral to a vascular service

7 J.2.1 Risk factors associated with disease progression

8 9

Table 99: Studies excluded from the clinical review: risk factors associated with disease progression

Reference	Reason for exclusion
Bernardini, 2010 ²²	No relevant outcomes and also does not match review question
Brand, 1988 ³²	No relevant outcomes and also does not match review question
Carpentier, 200845	Abstract. Authors contacted for more information but no reply received.
Christenson, 2012 ⁵⁸	Abstract. Authors contacted for more information but no reply received.
Cushman, 2010 ⁶³	No relevant outcomes and also does not match review question
Diamond, 2012 ⁷⁸	No relevant outcomes and also does not match review question Abstract only
Dzieciuchowicz, 2011 ⁹¹	Incorrect study design (cross-sectional)
Gasparis, 2008 ¹⁰¹	No relevant outcomes and also does not match review question Abstract only
Kostas, 2010 ¹³⁸	No relevant outcomes and also does not match review question
Labropoulos, 2012 ¹⁴²	No relevant outcomes and also does not match review question
Mackenzie, 2003 ¹⁵⁵	No relevant outcomes and also does not match review question
McLafferty, 2009 162	Abstract. Authors contacted for more information but no reply received.
Moore, 2011 ¹⁷²	Incorrect study design (cross-sectional) Abstract only
Pannier, 2012 ²⁰³	No relevant outcomes and also does not match review question Abstract only
Mota-Capitao, 1995 ¹⁷⁵	Incorrect study design - a cross-sectional study rather than a case control or

Reference	Reason for exclusion
	prospective cohort study
Rabe, 2010 ²¹⁶	Incorrect study design (commentary)
Robertson, 2011 ²²⁹	No relevant outcomes and also does not match review question Abstract only
Treiman, 2001 ²⁷¹	No relevant outcomes and also does not match review question
Uhl, 2005 ²⁷⁴	Incorrect study design - a cross-sectional study rather than a case control or prospective cohort study

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2 J.2.2 Factors associated with better/worse response to treatment

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Table 100: Studies excluded from the clinical review: Factors associated with better/worse response to treatment

Ref ID	Reason for exclusion
Ali, 2007 ⁵	No relevant outcomes and does not match review question
Allegra, 2007 ⁶	Incorrect study design - no multivariable analysis
Atkin, 2007 ¹⁰	Incorrect study design - no multivariable analysis
Barrett, 2004 ¹²	Incorrect study design - no multivariable analysis
Bush, 2012 ³⁸	Abstract only
Calcagno, 2009 ³⁹	Incorrect study design - no multivariable analysis
Carradice, 2011 ⁴⁸	No relevant outcomes and does not match review question
Chaar, 2011 ⁵¹	Incorrect study design - retrospective
Chi, 2011 ⁵⁵	Abstract only
Ciostek, 2004 ⁵⁹	Incorrect study design - no multivariable analysis
Corbett, 2011 ⁶¹	Incorrect study design - no multivariable analysis Abstract only
Defty, 2008 ⁷⁴	Incorrect study design - no multivariable analysis
Einarsson, 1993 ⁹⁴	Intervention does not match protocol (liquid sclerotherapy)
Goode, 2009 ¹⁰⁸	No relevant outcomes and does not match review question
Hartmann, 2006 ¹¹²	Incorrect study design - no multivariable analysis
Hingorani, 2009 ¹¹⁵	Intervention does not match protocol
Jagtman, 2003 ¹²³	Intervention does not match protocol - liquid sclerotherapy
Kakkos, 2006 ¹²⁸	Incorrect study design - no multivariable analysis for the analysis frelevant to the review question
Kim, 2006 ¹³³	Incorrect study design - no multivariable analysis
Magnusson, 2006 ¹⁵⁷	Incorrect study design – retrospective study
Marsh, 2010 ¹⁶⁰	Incorrect study design - no multivariable analysis
Marston, 2008 ¹⁶¹	Incorrect study design - no multivariable analysis
Meissner, 2012 ¹⁶⁴	Abstract only
Miyazaki, 2005 ¹⁷¹	Incorrect study design – retrospective study
Mouton, 2008 ¹⁷⁷	Population does not match protocol - inguinal varices
Nash, 1991 ¹⁸¹	Incorrect study design - no multivariable analysis
Nelzen, 2010 ¹⁸³	Abstract only
Oguzkurt, 2010 ¹⁹²	Abstract only

Ref ID	Reason for exclusion
Ozsvath, 2010 ¹⁹⁶	Abstract only
Pittaluga, 2009 ²¹¹	Incorrect study design - no multivariable analysis
Pittaluga, 2012 ²¹⁰	No relevant outcomes and does not match review question
Puggioni, 2005 ²¹⁴	Incorrect study design - no multivariable analysis
Puggioni, 2009 ²¹⁵	Incorrect study design - no multivariable analysis
Sarvananthan, 2012 ²³⁷	Incorrect study design - no multivariable analysis
Shepherd, 2009 ²⁴⁶	Abstract only
Shepherd, 2010 ²⁴⁵	Multivariable analysis, considering confounders such as sex, BMI or clinical disease severity, but no report of the independent effects of each confounder.
Stother, 1974 ²⁵⁵	intervention does not match protocol (liquid sclerotherapy)
Tzilinis, 2005 ²⁷³	Incorrect study design - no multivariable analysis
Van Neer, 2006 ²⁷⁷	Incorrect study design - no multivariable analysis
Vandy, 2011 ²⁷⁸	Abstract only
Ward, 2011 ²⁸⁰	Abstract only
Wright, 2006 ²⁸⁵	Incorrect study design - no multivariable analysis
Zamboni, 2010 ²⁸⁹	Incorrect study design - no multivariable analysis

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2 J.3 Chapter 7 – assessment for treatment

3 J.3.1 Diagnostic assessment of hand held Doppler

Table 101: Studies excluded from the clinical review: diagnostic assessment of hand held Doppler

Author/title	Reason for exclusion
Antoch 2002 ⁸	Does not address diagnostic accuracy
Campbell 200543	No diagnostic data presented
Dixon 1996 ⁸⁸	Does not address diagnostic accuracy
Engel 1991 ⁹⁵	Does not address diagnostic accuracy
Olivienca 1998 ¹⁹⁴	Does not address diagnostic accuracy
Phillips 1995 ²⁰⁵	No HHD diagnostic data
Pierik 1997 ²⁰⁹	No HHD diagnostic data
Singh 1997 ²⁵⁰	No usable diagnostic data presented
Yamaki 2002 ²⁸⁸	Does not address diagnostic accuracy

5 J.3.2 Duplex assessment prior to treatment

6

Table 102: Studies excluded from the clinical review

Ref ID	Reason for exclusion
Campbell 1996 ⁴²	No relevant outcomes and does not match review question
Cavezzi 200049	Comparator does not match protocol - no comparator
Cavezzi 200750	No relevant outcomes and does not match review question
Coleridge-Smith 2007 ⁶⁰	No relevant outcomes and does not match review question

Ref ID	Reason for exclusion
De Maeseneer 2011 ⁶⁹	No relevant outcomes and does not match review question
Krodowicz 2010 ¹³⁶	No relevant outcomes and does not match review question
Levi 1995 ¹⁴⁵	Population does not match protocol - population were patients undergoing infrainguinal bypass procedures; Comparator does not match protocol - no comparator
Makris 2006 158	No relevant outcomes and does not match review question
Pachler 1998 ¹⁹⁷	Comparator does not match protocol - no comparator
Pichot 2000 ²⁰⁸	Comparator does not match protocol - no comparator
Pleass 1996 ²¹²	No relevant outcomes and does not match review question
Safar 2004 ²³⁴	Comparator does not match protocol - no comparator
Somjen 1996 ²⁵³	No relevant outcomes and does not match review question
Campbell 1996 ⁴²	No relevant outcomes and does not match review question
Oinonen 2007 ¹⁹³	No relevant outcomes and does not match review question - both Duplex and hand held doppler were used for pre-op marking, with some patients only having been examined with HHD. Also, all patients had a duplex/doppler assessment, the difference between groups being that in one group the duplex/doppler findings were only communicated to the surgeons in written form, whereas in the other group the surgeon was allowed to do a pre-op marking, by utilising HHD and the previous duplex/doppler findings.

1 J.4 Chapter 8 – conservative management

2 J.4.1 Compression vs. no treatment

Table 103: List of excluded studies

Reference	Reason for exclusion
Ahfmr 1997 ⁴	Incorrect study design - review
Bachoo 2009 ¹¹	Incorrect study design - review
Brown 1992 ³⁷	Comparator does not match protocol - other form of compression
Chant 1989 ⁵²	Comparator does not match protocol - other form of compression
Chant 1985 ⁵⁴	Comparator does not match protocol - other form of compression
Diehm 1996 ⁸¹	Incorrect outcomes reported
Geraghty 1989 ¹⁰²	Probably the same study as Anderson 1990.
Jones 1980 ¹²⁵	Comparator does not match protocol - other form of compression
Kakkar 1982 ¹²⁷	Comparator does not match protocol - other form of compression
Labropoulos 1994 ¹⁴¹	Incorrect study design – laboratory study
Lippmann 1994 ¹⁴⁷	Intervention does not match protocol - Unna's boot
Michaels 2006 ¹⁷⁰	Comparator does not match protocol – liquid sclerotherapy).
Mosti 2011 ¹⁷⁴	Comparator does not match protocol- other form of compression
Murad 2011 ¹⁷⁸	Incorrect study design – systematic review
Palfreyman 2009 ²⁰⁰	Incorrect study design – systematic review
Thaler 2001 ²⁶⁰	No relevant outcomes and does not match review question
Tisi 2011 ²⁶⁹	Incorrect study design – systematic review

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1 J.4.2 Compression vs. interventional treatment

Table 104: Studies excluded from the clinical review

Ref ID	Reason for exclusion
Abramowitz 1973 ³	Intervention does not match protocol - liquid sclerotherapy only
Barwell 2004 ¹³	Population does not match protocol - treatment of chronic ulceration
Gohel 2007 ¹⁰⁵	Population does not match protocol - treatment of chronic ulceration
Gohel 2005 ¹⁰⁴	Population does not match protocol - treatment of chronic ulceration
Guest 2003 ¹⁰⁹	Population does not match protocol - treatment of chronic ulceration
Leu 1993 ¹⁴⁴	Intervention does not match protocol – foam sclerotherapy and compression applied concomitantly
Palfreyman 2009 ²⁰⁰	Incorrect study design - systematic review.
Schul 2011 ²⁴⁰	Population does not match protocol - CEAP1 only – not a true varicose veins population.
Shingler 2011 ²⁴⁸	Incorrect study design - systematic review.

2

J.5 Chapter 9 – interventional treatment

2 J.5.1 Stripping surgery vs. foam sclerotherapy

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Table 105: Studies excluded from the clinical review

Reference	Reason for exclusion
Belcaro2003 ¹⁶	Intervention does not match protocol – avulsion
Belcaro2000 ¹⁸ .	Intervention does not match protocol – flush ligation and avulsion
Beresford1978 ²⁰	Comparator does not match protocol – liquid sclerotherapy
Chant197253	Comparator does not match protocol – liquid sclerotherapy
Doran1975 ⁸⁹	Comparator does not match protocol – liquid sclerotherapy
Eifell2006 ⁹³ .	Incorrect study design - study protocol.
Einarsson1993 ⁹⁴	Comparator does not match protocol – liquid sclerotherapy
Haas2006 ¹¹⁰ .	Non English language publication
Hobbs1968 ¹¹⁶	Comparator does not match protocol – liquid sclerotherapy
Hobbs1974 ¹¹⁷	No relevant outcomes and does not match review question
Jakobsen1979 ¹²⁴	Comparator does not match protocol – liquid sclerotherapy
Liamis2005 ¹⁴⁶ .	Abstract only
Michaels2006A ¹⁶⁹	Comparator does not match protocol – liquid sclerotherapy
Miyazaki2005 ¹⁷¹	Incorrect study design - retrospective observational study and not an RCT.
Murad2011 ¹⁷⁸	No relevant outcomes and does not match review questions
Neglen1993 ⁹⁴	No relevant outcomes and does not match review questions
Rigby2004 ²²⁸ .	No relevant outcomes and does not match review questions
Rutgers1994 ²³²	Comparator does not match protocol – liquid sclerotherapy
Van Den Bos 2009 ²⁷⁵ .	Incorrect study design - systematic review.

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5 J.5.2 Stripping surgery vs. endothermal ablation

Table 106: Studies excluded from the clinical review

Reference	Reason for exclusion
Anon2010 ¹⁶³	Incorrect study design - systematic review
Basel2012 ¹⁴	Incorrect study design - probably non-randomised. Although stated in the abstract that this paper was randomised, there is no mention of randomisation in the paper itself. Instead, the methods section states that "the study was planned as a retrospective study".
Brar2010 ³³	Incorrect study design - systematic review
Christenson2010 ⁵⁷	Some patients with bilateral symptoms had each leg randomised to a different group, making QoL and pain results rather meaningless (as perception will be global not per leg).
Darwood200966	Incorrect study design - systematic review
De Medeiros2005 ⁷¹	Intervention does not match protocol - laser group had high tie of SFJ – thus "non-standard" laser
De Medeiros2006 ⁷⁰	Same study as de Medieros 2005, with identical outcomes
Disselhoff2008 ⁸⁵	intervention does not match the protocol - cryostripping
Disselhoff2008 ⁸⁷	intervention does not match the protocol - cryostripping

Reference	Reason for exclusion
Disselhoff2009 ⁸⁴	Comparator does not match protocol - cryostripping instead of standard stripping
Disselhoff2011 ⁸⁶	intervention does not match the protocol - cryostripping
Kalteis2008 ¹³⁰	Intervention does not match protocol - laser was combined with SFJ ligation – thus "non standard" laser
Kundu2011 ¹⁴⁰	Incorrect study design - systematic review
Luebke2008 ¹⁴⁹	Incorrect study design - systematic review
Nesbitt2011 ¹⁸⁴	Incorrect study design - systematic review
Subramonia2010 ²⁵⁷	Economic analysis and otherwise same study as Subramonia 2010B
Tellings2011 ²⁵⁹	Incorrect study design - systematic review
Theivacumar2009 ²⁶¹	Incorrect study design - cohort study
Vuylsteke2006 ²⁷⁹	Incorrect study design - not randomised
Xenos2009 ²⁸⁷	Incorrect study design - systematic review

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2 J.5.3 Foam sclerotherapy vs. endothermal ablation

Table 107: Studies excluded from the clinical review

Reference	Reason for exclusion
King2009 ¹³⁵	Endovenous ablation and sclerotherapy applied concomitantly – not a comparison.
Koroglu2011 ¹³⁷	Endovenous ablation and sclerotherapy applied concomitantly – not a comparison.
Luebke2008 ¹⁵⁰	Incorrect study design - systematic review

4 J.5.4 Tributary treatment: avulsion vs. foam sclerotherapy

5

Table 108: Studies excluded from the clinical review

Reference	Reason for exclusion
Belcaro 2000 ¹⁸	Intervention does not match protocol - did not include avulsion surgery
Belcaro 2003 ¹⁶	Unclear if the foam sclerotherapy was a tributary treatment.
Belcaro 2003B ¹⁵	Identical report to the included Belcaro 2003 study.
Brethauer 2001 ³⁵	Comparator does not match protocol - liquid sclerotherapy
De Roos 2003 ⁷³	Comparator does not match protocol - liquid sclerotherapy

6

1 J.5.5 Truncal and tributary treatment vs. truncal treatment alone

Table 109: Studies excluded from the clinical review

Ref ID	Reason for exclusion
Belcaro 1998 ¹⁷	Non English speaking publication
Belcaro 2000 ¹⁸	Intervention does not match protocol - liquid sclerotherapy
Belcaro 2003B ¹⁵	Foam sclerotherapy was done up to 6 months after the surgery
Kim ¹³²	Comparator does not match protocol - included tributary treatments given with endovenous ablation.
King 2009 ¹³⁵	Incorrect study design - no comparator
Koroglu ¹³⁷	Did not address the review question.
Mekako 2006 ¹⁶⁵	Incorrect study design - no comparator
Merchant 2005 ¹⁶⁸	No relevant outcomes and does not match review question
Nicolini 2005 ¹⁸⁸	Intervention does not match protocol - those receiving tributary treatments included those receiving such treatments during the follow-up period (i.e. not just those who had tributary treatments at the same time as the truncal treatment)
Sadick 2007 ²³³	Incorrect study design - no comparator
Schanzer 2010 ²³⁸	Study comparator does not match protocol
Theivacumar 2006 ²⁶⁴	Abstract only
Theivacumar 2007 ²⁶⁵	Contained information that 40% of patients undergoing laser ablation to GSV required delayed tributary treatment. However this paper did not address the review question.
Theivacumar 2008 ²⁶⁶	No relevant outcomes and does not match review question
Theivacumar 2008A ²⁶⁷	This study had one group where sclerotherapy was used concomitantly with EVLA, but this was not to treat tributaries (it was to treat below knee truncal reflux).
Theivacumar 2009 ²⁶¹	No relevant outcomes and does not match review question
Theivacumar 2009A ²⁶³	No relevant outcomes and does not match review question
Theivacumar 2009B ²⁶²	Contained information that 44% of patients undergoing laser ablation to GSV required delayed tributary treatment. However this paper did not address the review question.
Welch 2006 ²⁸³	No relevant outcomes and does not match review question
Weiss RA, Weiss MA. ²⁸²	No relevant outcomes and does not match review question

2

2

J. J.6 Chapter 10 – compression after interventional treatment

Reference	Reason for exclusion
Biswas 2007 ²³	Compared different durations of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Bond 1999 ²⁹	Compared different types of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
de Roos 2008 ⁷²	Incorrect study design - no comparator
De Jode 1970 ⁷⁵	Intervention does not match protocol - liquid sclerotherapy used
Din 1992 ⁸³	Intervention does not match protocol - liquid sclerotherapy used
Fraser 1985 ^{73,100}	Intervention does not match protocol - liquid sclerotherapy used
Lugli 2009 ¹⁵¹	Compared different forms of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Mariani 2011 ¹⁵⁹	Compared different forms of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Melrose 1979 ¹⁶⁶	No relevant outcomes and does not match review question
Mosti 2009 ¹⁷³	Compared different levels of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
O'Hare 2010 ¹⁹⁰	Compared different durations of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Raraty 1999 ²¹⁸	Compared different durations of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Rhodes 1972 ²²⁷	Intervention does not match protocol - liquid sclerotherapy used
Rodrigus 1991 ²³¹	Compared different durations of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Shingler 2011 ²⁴⁸	Incorrect study design - systematic review
Shouler 1989 ²⁴⁹	Compared different levels of compression; this would have met the inclusion criteria for part b) of the research question if part a) had shown a reasonable level of efficacy for compression as an adjuvant therapy.
Travers ²⁷⁰	No relevant outcomes and does not match review question
Tretbar 1970 ²⁷²	Intervention does not match protocol - liquid sclerotherapy used
Weiss 1999A ²⁸¹	Intervention does not match protocol - liquid sclerotherapy used

Table 110: Studies excluded from the clinical review

Appendix K: Excluded economic studies

2 K.1 Chapter 8 – conservative management

3 K.1.1 Compression vs. interventional treatment

4 Table 111: Studies excluded from the economic review

Reference	Reason for exclusion
Eskelinen et al. 2009 ⁹⁶	This is an economic analysis based on a cohort study. The calculated QALYs appear to be inconsistent with the reported utility data and it is unclear whether the 15-dimension quality-of-life instrument has been validated.

5 K.1 Chapter 9 – interventional treatment

6 K.1.1 Stripping surgery vs. foam sclerotherapy

7 Table 112: Studies excluded from the economic review

Reference	Reason for exclusion
Beresford et al 1978 ²⁰	Dated article
Bountouroglou et al 2006 ³⁰	This study did not report QALYs. More applicable evidence was available for this comparison.
Piachaud & Weddell 1972 ²⁰⁷	Dated article
Piachaud & Weddell 1972A ²⁰⁶	Dated article
ZonMw 2005 ¹⁸⁵	Study results are not published. It is stated that a full report is available from the authors. Attempts to make contact with the authors, however, failed.

8

9 K.1.2 Stripping surgery vs. endothermal ablations

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Table 113: Studies excluded from the economic review

Reference	Reason for exclusion
Disselhoff et al 2009 ⁸⁴	The study is an economic analysis carried out alongside a randomized controlled comparing cryostripping with endovenous laser ablation.
Eidson et al 2011 ⁹²	This is a retrospective cohort study comparing the relevant treatments over a 6 month period.
Medical Advisory Secretariat 2010 ¹⁶³	This is a costing and budget impact study to identify resource and cost differences between the two interventions compared.
Ontario Health Technology Advisory Committee ¹	This is a costing and budget impact study to identify resource and cost differences between the two interventions compared.
Rasmussen et al 2007 ²¹⁹	This study was excluded as more recent results are reported in Rasmussen et al. 2011 ²²¹ .
Rasmussen et al 2011 ²¹⁹	This study was based on the Danish healthcare system and does not report QALYs. It has a lower applicability than other evidence for this comparison.

Reference	Reason for exclusion
Rautio et al 2002 ²²⁴	This study was based on the Finish healthcare system and does not report QALYs. It has a lower applicability than other evidence for this comparison.
Solis et al 2009 ²⁵²	Actual cost estimates are not provided but rather symbols are used to indicate that endovenous laser ablation is more expensive than stripping surgery.
Subramonia and Lees 2010 ²⁵⁷	This study does not report QALYs. It has a lower applicability than other evidence for this comparison
Vuylsteke et al 2006 ²⁷⁹	This study was based on the Belgian healthcare system and does not report QALYs. It has a lower applicability than other evidence for this comparison.

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2 K.1.3 Foam sclerotherapy vs. endothermal ablation

Table 114: Studies excluded from the economic review

Reference	Reason for exclusion
Rasmussen et al 2011 ²²¹	This study was based on the Danish healthcare system and does not report QALYs. It has a lower applicability than other evidence for this comparison.

Appendix L: Cost-effectiveness analysis of interventional treatments and conservative care

L.1 Introduction

Interventional treatments for varicose veins (surgery, foam sclerotherapy and endothermal ablation techniques) are likely to have significant resource implications, and as such it is important to identify which of these treatments represents a cost-effective use of NHS resource. Two published UK economic evaluations were identified in this area^{106,170}Michaels and colleagues (2006) compared surgery, liquid sclerotherapy and conservative care in various groups with differing stages of disease and found surgery to be the most cost-effective option. However this analysis did not look at endothermal techniques or foam sclerotherapy, and is therefore not complete for our purposes. The second evaluation, Gohel and colleagues (2010), was subject to several limitations; in particular, the costs were not thought to be representative of common practice as they were based on modified reference costs, and the authors appear to assume that recurrence rates do not differ after different modalities of treatment. The conclusions of the model must therefore be interpreted with caution, and are of limited value in informing recommendations.

We are also aware of an on-going NHS Health Technology Assessment (HTA) project to investigate the cost effectiveness of minimally invasive techniques.²³⁹ There is overlap between the HTA project and the analysis presented here, however we are aware of a few key differences: in contrast to our model, the aforementioned project does not include conservative care as a comparator, the clinical outcomes included in the network meta-analysis (NMA) are different, and as such the model is based on different clinical data. The HTA project is still being revised, therefore no further comment can be made here. Overall, the GDG did not deem the existing literature to be sufficient on which to base recommendations. Interventional treatments were therefore identified as a priority for original economic analysis.

Initially it was planned that analyses would also look at the cost-effectiveness of these treatments at different stages of the disease, thereby addressing the question of the optimal timing of intervention. However, a lack of relevant data meant this was not possible. This analysis therefore focuses on the cost effectiveness of the various interventional treatments for varicose veins, in comparison to non-invasive conservative care, in the general primary varicose veins population.

Analysis of patients with bilateral disease was carried out separately as a cost-comparison, with the results and their implications discussed thoroughly by the GDG.

L.2 Methods

L.2.1 Model overview

A cost-utility analysis was undertaken where costs were considered from a UK NHS and personal social services perspective and health outcomes expressed as quality adjusted life years (QALYs). Costs and QALYs were both discounted at 3.5% per annum, in accordance with the NICE reference case¹⁸².

L.2.1.1 Comparators

Four treatments were considered in the base case:

- Surgery (stripping and ligation) with or without tributary treatment, carried out as a day case procedure under general anaesthetic
- Endothermal techniques (RFA & EVLA) with concurrent phlebectomy carried out as an outpatient procedure under local anaesthetic
- Foam sclerotherapy with or without tributary treatment, carried out as an outpatient procedure under local anaesthetic
- Conservative care (compression therapy)

Endothermal techniques without concurrent phlebectomy were evaluated as a sensitivity analysis.

L.2.1.2 Population

Adults with primary unilateral great saphenous vein (GSV) incompetence, who are potentially suitable for treatment by any of the four treatment options.

In some cases, particular treatments may not be suited to an individual patient. Decision models are designed to identify the optimal choice between two or more alternative strategies; the choice between the comparators only applies to people for whom all of these are a possibility.

Sensitivity analyses included unilateral patients receiving concurrent treatment of the small saphenous vein (SSV), and a cost comparison to inform GDG discussion around treatment of patients with bilateral varicose veins.

L.2.1.3 Time horizon

The time horizon of the model was five years in the base case. This was chosen as the appropriate horizon as there is no differential mortality effect between treatment options, and no reliable evidence was found to document differences in costs and health related quality of life past this time horizon. Extrapolation of follow-up data (data is limited to a 3 year follow-up) past the 5 year time horizon would have been subject to a great deal of uncertainty and was not deemed to be appropriate in this instance.

Sensitivity analyses included evaluation over a 3 year time horizon, in line with the maximum followup time of the clinical data.

L.2.2 Approach to modelling

Interventional treatments for varicose veins are used to occlude, obliterate or strip varicose veins, thereby reducing patient symptoms and improving quality of life. When these outcomes are not achieved, top-up treatment can be given to supplement the initial treatment. For the purpose of the model, this combination of initial treatment and top-up treatment was considered to be one treatment episode. All patients in the model have an initial treatment episode (either outpatient or day case depending on the treatment option), with different treatments leading to different proportions of individuals requiring top-up treatment, resulting in a difference in costs and QALYs associated with the initial treatment episode.

Patients in the model experience an increase in quality of life (QoL) once the initial treatment episode is complete. The difference in QALYs was driven by the length of time this increase in QoL was sustained for, before a patient experienced recurrent varicose veins. The probability of having recurrent varicose veins differed by treatment and for the purpose of the model was taken from a network meta-analysis described in section L.2.3.2.2 below. The NMA was based on clinical recurrence data, which was used to capture the development of symptoms of varicose veins in a treated limb. Clinical recurrence was chosen over other possible definitions of recurrence because

symptoms (as opposed to reflux, recanalisation, or any other definition) have a direct impact on QoL. Patients could undergo a second treatment episode in the model, to alleviate symptoms of recurrent varicose veins and improve QoL again. Further information and technical details are given in the subsequent sections.

L.2.2.1 Key definitions

A **treatment episode** consists of a treatment for every patient, and a top-up treatment for the proportion of individuals who require it. There is potential for two treatment episodes in the model; an **initial treatment episode** which all patients in the model receive, and a **second treatment episode** which is given to a proportion of individuals following clinical recurrence. The second treatment episode is distinct from top-up treatment, which is considered to be part of a treatment episode.

Top-up treatment is given as part of a treatment episode (within 2 months of the initial treatment) if treatment is not deemed to be complete (i.e. if the vein undergoing treatment has not been occluded or obliterated, or if additional treatment of residual varicosities is needed). Top-up treatment was assumed to always be foam (see Table 115). In some cases the need for top-up treatment could be identified by a follow-up appointment if one is given, or the top-up treatment could have been planned before the initial treatment; in other cases, the patient may present with a need for top-up treatment. The inclusion of top-up treatment in the model is not intended to imply a recommendation of routine follow-up appointments. The concept of top-up treatment is adapted from the aforementioned HTA project,²³⁹ in which it is used to refer to residual varicose veins, and does not include unsuccessful occlusion.

Clinical recurrence is defined as development of symptoms of varicose veins in a treated limb. For the purpose of the network meta-analysis, papers which report clinical recurrence as an outcome were taken at face value.

L.2.2.2 Model structure

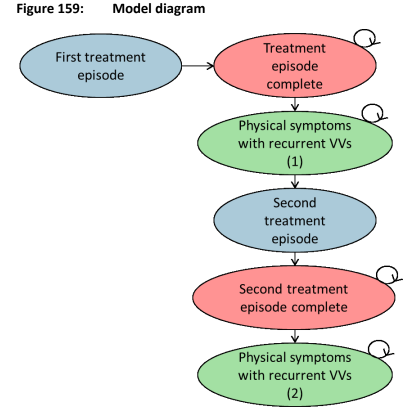
A Markov model was constructed to calculate costs and QALYs for each comparator; the key health states and transitions can be seen in Figure 159. The simplified diagram does not show death, but patients could die of all-cause mortality at any point during the model's five year time horizon.

Patients enter the model through the 'First treatment episode' state. Following completion of the treatment episode, patients move to a state of 'treatment episode complete', where they do not require any further treatment. They remain in this state until they experience clinical recurrence, at which point they transition to the state 'Physical symptoms with recurrent VVs (1)'. Patients cannot experience clinical recurrence in the first three cycles post treatment, to allow time for all top-up treatments to take place.

Only a proportion of patients with clinical recurrence go on to have further interventional treatment, whilst the rest receive conservative care. For those who do receive further interventional treatment, a delay of 6 months is assumed between the onset of clinical recurrence and the second treatment episode. This delay is based on clinical opinion and captured in the model through a series of tunnel states (these states are omitted from Figure 159 for simplicity). Following the second treatment episode, a patient can experience clinical recurrence again, but will not receive further treatment; instead they remain in the state of 'physical symptoms with recurrence VVs (2)' and receive conservative care. Conservative care received at this point is assumed not have an impact on QoL

Conservative care was modelled separately to the other three interventions, as the outcomes of completed treatment and clinical recurrence are not clinically meaningful when considering this management technique. Instead, the difference in quality of life between patients undergoing surgery and conservative care (as reported in Michaels2006¹⁷⁰) was used to calculate the difference

in QALYs over time between these two treatments. This information was then used to calculate the QALYs expected from conservative care, relative to the QALYs computed by the model for surgery. Costs were calculated by applying an annual cost to all those individuals in the model and receiving conservative care.



Schematic diagram of the Markov model designed to compare the cost-effectiveness of treatments for varicose veins. The Markov modelling approach involves a transition between different health states over time. The model is divided into monthly cycles. At the end of each cycle a transition to another health state is possible, unless people enter into an 'absorbing state' from which they cannot transition. In this model, the absorbing state is 'Physical symptoms with recurrent VVs (2)'.

The model was built with a one month cycle length as this was deemed to be the minimum clinically meaningful time interval to detect differences between interventions. All the probabilities, costs and health utilities were converted to reflect the one-month cycle length.

L.2.2.3 Key assumptions

The model employed the following key assumptions:

Table 115. Key assumptions							
Assumption	Comment						
Rates of top-up treatment are the same in the initial and second treatment episode (ie after retreatment)	The GDG deemed this to be a reasonable assumption						
Top-up treatment is always foam sclerotherapy	As the modality of top-up treatment does not affect the rate of recurrence (see assumption below), this will only effect the cost of top-up treatment. The GDG deemed this to be a reasonable simplifying assumption.						

Table 115: Key assumptions

Assumption	Comment
Patients who have had top-up treatment have the same probability of recurrence as those who haven't had top-up	The GDG deemed this to be a reasonable simplifying assumption
Constant hazard of recurrence	This was deemed to be a reasonable simplifying assumption as the time horizon of the model is relatively short
There is a 6 month delay between the onset of clinical recurrence and the second treatment episode	This is included to reflect the time between the onset of symptoms and subsequent interventional treatment.
A patient can only receive two treatment episodes in total	This is a simplifying assumption for the model but is expected to be a fair reflection of routine clinical practice
Proportions of patients having each modality of second treatment is independent of the modality of their initial treatment	The method of retreatment is more likely to be based on individual patient characteristics and the nature of the recurrence, rather than the modality of initial treatment. As the model cannot capture these factors for individual patients, the GDG deemed this to be a reasonable assumption.

L.2.2.4 Uncertainty

The model was built probabilistically to take account of the uncertainty surrounding each input parameter. In order to characterise uncertainty, a probability distribution was defined for each parameter based on error estimates from the data sources (e.g. standard errors or confidence intervals). The way in which distributions are defined reflects the nature of the data (see Table 116). When the model was run, a value for each input was randomly selected from its respective distribution. The model was run repeatedly (10, 000 times) to obtain mean cost and QALY values.

Various sensitivity analyses were also undertaken to test the robustness of model assumptions and data sources. In these analyses, one or more inputs were changed and the analysis was rerun in order to evaluate the impact of these changes on the results of the model.

Parameter	Type of distribution	Properties of distribution
Cost	Gamma	Bounded at 0, positively skewed. Derived from mean and standard error
Pre-treatment utility	Beta	Bounded on 0 – 1 interval. Derived from mean and sample size
Utility improvements and decrements	Lognormal	Bounded at 0. Derived from log of mean utility change and standard error of log of utility change
Utility difference between conservative care and surgery	Normal	Derived from mean and variance
Baseline risk and relative effects	Distribution estim	ated by sampling from network meta-analysis output

L.2.3 Model Inputs

L.2.3.1 Summary table of model inputs

Model inputs were based on clinical evidence identified in the systematic review undertaken for the guideline, supplemented by additional data sources as required. All inputs were checked for face

validity by the clinical members of the GDG. A summary of the model inputs used in the base-case analysis is provided in Table 117 Table 118 below. More details on sources, calculations and rationale for selection can be found in subsequent sections.

	Input	Source
Comparators	Surgery, foam sclerotherapy, endothermal with phlebectomies, conservative care	GDG consensus
Population	Adults with primary unilateral GSV incompetence	GDG consensus
Initial cohort settings	Age: 50 Female: 65%	Weighted average across relevant $RCTs^1$
Perspective	NHS and PSS	NICE reference case ¹⁸²
Time horizon	5 years	GDG consensus
Discount rate	Costs: 3.5% QALYs: 3.5%	NICE reference case ¹⁸²

Table 117: Summary of base-case model inputs and cohort settings

GSV = great saphenous vein

 $^{\scriptscriptstyle 1}$ the RCTs included in the network meta-analysis for clinical recurrence

Initial cohort settings

A starting age of 50 was used in the model to represent the average age of people undergoing treatment for varicose veins, and the cohort was assumed to be 65% female, based on the characteristics of patients in the included RCTs. These cohort characteristics were validated against HES data¹²¹ which confirms that the average age of patients undergoing day case treatment for varicose veins is approximately 50, and that roughly two thirds of these patients are female.

	Point	Probability Distribution						
Parameter description	estimate	distribution	parameters	Source				
Utility weights								
Primary varicose veins	0.764	Beta	α = 37600, β = 12800	PROMs ¹¹⁹				
Change in utility (from baseline) post treatment	+0.091	Lognormal	μ = -2.397, σ = 0.0007	PROMs ¹¹⁹				
Change in utility (from baseline) due to recurrent varicose veins	-0.093	Lognormal	μ = -2.388, σ = 0.0162	Beresford 2003 ²¹				
Conservative care (relative to surgery at 1 year)	-0.101	Normal	μ = 0.0004, σ = 0.0198	Michaels 2006 ¹⁷⁰				
Transition probabilities								
Probability of requiring top-	up treatment	(within 2 month	s post treatment)					
Surgery	5%	Deterministic S	SA only	GDG estimate				
Endothermal	5%	Deterministic S	SA only	GDG estimate				
Foam Sclerotherapy	20%	Deterministic S	SA only	GDG estimate				
Conservative care	NA							
Probability of recurrence (per month)								
Surgery	0.008331	Point estimate and uncertainty from NMA						
Endothermal	0.005833	Point estimate and uncertainty from NMA						
Foam Sclerotherapy	0.009141	Point estimate and uncertainty from NMA						

Parameter description	Point estimate	Probability distribution	Distribution parameters	Source	
Conservative care	NA				
Cost (£)					
Surgery	£908	Gamma	See Table 126	See Table 126	
Endothermal	£624	Gamma	See Table 128	See Table 128	
Foam Sclerotherapy	£315	Gamma	See Table 129	See Table 129	
Conservative care ^a	£234	Deterministic SA only			
Additional cost associated with retreatment	£417	Gamma	See Table 131	See Table 131	

SA = Sensitivity analysis; NMA=network meta-analysis

^athis is an annual cost (first year incurs and additional £15)

L.2.3.2 Baseline event rates and relative treatment effects

L.2.3.2.1 Top-up treatment rates

Limited data on treatment failure was available from the randomised trials, often based on different definitions of failure and very short follow-up of just a few days. In addition, treatment failure, however defined, is not the only reason that top-up treatment may be undertaken. For example, further treatment could be necessary to eradicate residual varicosities which were not treated initially (this may or may not have been planned at the time of the initial treatment). The data on treatment failure from the trials was therefore not considered to be relevant to the need for top-up treatment as defined in our model. For this reason, the proportions of patients requiring top-up after each treatment are based on GDG estimate (see Table 118).

L.2.3.2.2 Clinical recurrence

The results of conventional meta-analyses of direct evidence alone make it difficult to determine which intervention is the most effective treatment. The challenge of interpretation has arisen for two reasons:

- In isolation, each pair-wise comparison does not fully inform the choice between all the possible treatments, and having a series of discrete pair wise comparisons can be disjointed and difficult to interpret
- There are overlapping comparisons that could potentially give inconsistent estimates of effect.

This is particularly problematic for probabilistic analysis. To overcome these problems, a Bayesian network meta-analysis (NMA)⁴⁰ was conducted in WinBUGS.

Conventional meta-analysis assumes that, for a fixed-effect analysis, the relative effect of one treatment compared to another is the same across an entire set of trials. In a random-effects model, it is assumed that the relative effects are different in each trial but that they are from a single common distribution and that this distribution is common across all sets of trials.

Network meta-analysis requires an additional assumption over conventional meta-analysis. The additional assumption is that intervention A has the same relative effect across all trials of intervention A compared to intervention B as it does across trials of intervention A versus intervention C, and so on. Thus, in a random-effects network meta-analysis, the assumption is that intervention A has the same effect distribution across all trials of A versus B, A versus C and so on.

The aim of the NMA was to calculate treatment-specific probabilities of clinical recurrence following each of the different treatments. Clinical recurrence was chosen over other possible definitions of recurrence because symptoms (as opposed to reflux, recanalisation, or any other definition) are most

likely to have an impact on QoL. The GDG did not think it was appropriate to combine different measures of recurrence into one measure of effect. The definition of clinical recurrence as used in this analysis was given in section L.2.2.1.

Statistical analysis

When modelling an outcome such as clinical recurrence, it is important to consider the different follow-up times of the various trials, as longer follow-up is likely to result in more reported recurrences. To account for this, an underlying Poisson process with a constant event rate was assumed for each trial arm, and a complementary log-log (cloglog) link function used to model the event rate. The following logic was used to calculate hazards and hazard ratios:

Let *BH* and *HR* denote the baseline hazard (from the surgery arms) and treatment-specific hazard ratio for clinical recurrence; let θ represent the cloglog of the probability of clinical recurrence, *p*, and let *time* represent the duration of follow-up. Then:

$$\theta = Ln (time) + Ln (HR) + Ln (BH)$$

And:

$$p = 1 - \exp\{-\exp\theta\}$$

Surgery was chosen as the baseline comparator as it featured in the most trials. The baseline hazard was estimated on the clog-log scale through a meta-analysis of the surgery arms of the included trials. The resulting predictive distribution was inputted to the NMA for adjustment by the treatment specific hazard ratios to calculate the probability of clinical recurrence for each treatment. The codes for both the baseline and relative effects models were adapted from that provided on the NICE decision support unit website, and run in WinBUGS 14.

The baseline and relative effects models were run for 50,000 iterations with burn in periods of 50,000. Vague uninformative priors were combined with the data-driven likelihood functions to produce posterior probability estimates. Convergence was assessed by examining the history and kernel density plots.

Fixed and random effects NMAs were run, and goodness of fit estimated by calculating the total residual deviance and deviance information criteria (DIC) for each of the models. A total residual deviance close to the number of unconstrained data points (the number of trial arms in the analysis) indicated a model explaining the data at a satisfactory level. The DIC provides a measure of goodness of fit which penalises model complexity,⁷⁹ which is useful for comparing models. The choice of a fixed or random effects model can therefore be made by comparing their goodness-of-fit to the data.

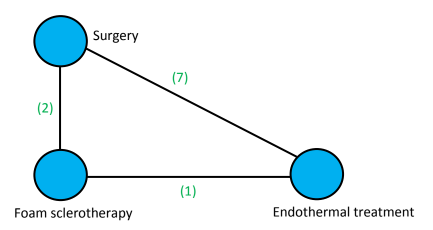
Network and Data

A total of eight studies included in the clinical reviews of the relevant treatments included clinical recurrence as an outcome. The trials included a variety of patients at differing levels of severity of varicose veins, further information on the trials can be found in chapters X and X of the full Surgery featured in all eight of the trials, endothermal treatment featured in seven, and foam sclerotherapy in two. One trial included all three comparators. The network of trials compared in Table 119. In

Figure 160 the number of trials included for each pair-wise comparison is noted in parentheses.

Note that the comparison between sclerotherapy and endothermal treatment includes only one trial. This single included study is a three arm trial which includes all of our comparators. Three arm trials are internally consistent, and as such there is no potential for inconsistency within our network, only for between-trial heterogeneity. This is discussed further by Dias and colleagues in technical support document 4,⁸⁰ in which the authors explain that 'loops of evidence that are potentially inconsistent can only arise from structures in which there are three distinct trials or sets of trials'.

Figure 160: Network of trials compared in the network meta-analysis



	Follow up	Clinical	Clinical recurrence		Total number of patients		Treatments compared			
Study name (months)	Arm 1	Arm 2	Arm 3	Arm 1	Arm 2	Arm 3	Arm 1	Arm 2	Arm 3	
Shadid 2012 ²⁴⁴	24	16	24	-	177	213	-	S	FS	-
Rasmussen 2011 ²²¹	12	16	17	23	108	123	245	S	FS	E
Carradice 2011 ⁴⁶	12	23	5	-	113	124	-	S	E	-
El Kaffas 2011 ¹¹³	24	9	12	-	90	88	-	S	E	-
Perala 2010 ²⁰⁴	36	2	5	-	13	15	-	S	Е	-
Pronk 2010 ²¹³	12	3	3	-	56	49	-	S	E	-
Rasmussen 2010 ²²⁰	24	25	18	-	68	69	-	S	E	-
Rass 2011 ²²²	24	33	28	-	143	173	-	S	E	-

Table 119: Clinical recurrence trial data for network meta-analysis

Abbreviations: S = surgery; E = endothermal treatment; FS = foam sclerotherapy

Network meta-analysis results

The total residual deviance was 25.3 for the fixed effects model and 18.6 for the random effects model which, when compared to 17 unconstrained data points, shows that the random effects model fitted the data reasonably well. DIC statistics of 105.5 and 103.4 were calculated for the fixed effects and random effects models respectively which, although the difference is small, suggests that the random effects model is the preferred option. Results are therefore presented for the random effects model only.

The final treatment-specific probability estimates and their associated confidence intervals can be seen in Table 120.

	Clinical recurrence (probability per month)						
Treatment	Mean	Standard deviation	Median	Confidence interval			
Surgery	0.008818	0.00306	0.008331	0.004284 - 0.0161			
Endothermal treatment	0.006532	0.003448	0.005833	0.002424 - 0.01472			
Foam sclerotherapy	0.01115	0.009929	0.009141	0.002795 - 0.03093			

Table 120: Network meta-analysis results – probability of clinical recurrence

As shown in Table 121, endothermal treatment was associated with the lowest probability of recurrence per month. These estimates were used to parameterise treatment effects in the decision model; deterministic point estimates were based on median values, with PSA values sampled from the WinBUGs CODA output.

A posterior estimate of heterogeneity - the between trial standard deviation - was found to be 0.58. An estimate of this magnitude indicates a large amount of variation in treatment effects calculated from different trials.

L.2.3.2.3 Retreatment

Not all patients are retreated after experiencing clinical recurrence; for some patients this is because they do not wish undergo further treatment, whereas for others it is because they are not deemed suitable for further treatment. The GDG estimated that 75% of patients would receive further interventional treatment, and it was assumed that the remaining 25% would receive conservative care. This estimate was subject to wide ranging deterministic sensitivity analysis.

For those individuals who do undergo a second treatment episode, the mode of treatment is likely to depend on the nature of their recurrence, alongside further patient characteristics. Based on their experience in practice, the GDG estimated that the following proportions of patients would have each type of retreatment (Table 121).

Table 121: Method of retreatment

Second treatment	% patients receiving each method of retreatment
Surgery	12%
Foam sclerotherapy	42%
Endothermal techniques	46%

These proportions represent an average over the 5 year time horizon, and were the same irrespective of the modality of the initial treatment. There is substantial uncertainty surrounding these estimates, however due to the nature of the model they are unlikely to drive the results. Nevertheless, these proportions were subject to extensive deterministic sensitivity analysis (seesection L.2.4).

L.2.3.2.4 Adverse events

Evidence on adverse events due to treatment identified by the clinical review was weak; different trials report different outcomes, and measure them in different ways. For example, pain is measured by VCSS at 1, 2 and 6 months in Figueiredo 2009,⁹⁷, as a dichotomous outcome at one year in Pronk 2011,²¹³ and by SF-36 in Rasmussen 2011.²²¹ In addition, the GDG members felt that the adverse

event profiles of the different treatments were similar to the extent that their inclusion would not benefit the model; therefore adverse events were not included in the analysis.

Whilst factors such as time to return from work, and time to return to usual activities may differ between treatments, these do not fall within the model's perspective of the NHS and PSS, and are therefore outside the remit of this analysis.

L.2.3.2.5 Mortality

The treatments considered in the analysis are not assumed to have any differential effect on mortality, yet patients can die at any point in the model. Age-specific all-cause mortality, weighted for the gender split of the cohort population, was based on the most recent available life tables for England and Wales (2008-20010).¹⁹¹

L.2.3.3 Utilities

In cost-utility analyses, measures of health benefit are valued in terms of quality adjusted life years (QALYs). The QALY is a measure of a person's length of life weighted by a valuation of their health related quality of life (QoL) over that period. The weight used is called a utility value, which is a measurement of the preference for a particular health state, with a score ranging from 0 (death) to 1 (perfect health). Questionnaires such as the SF-36 and SF-12 provide generic methods of describing QoL, while the EQ-5D, HUI, and SF-6D also include preference-based valuations of each health state, allowing calculation of utility scores.

The preferred method for determining utilities for NICE economic evaluations is the EuroQoL (EQ-5D) questionnaire¹⁸². The EQ-5D comprises five dimensions of health: mobility, ability to self-care, ability to undertake usual activities, pain and discomfort, and anxiety and depression. For the NICE reference case, preferences from the general public should be used. In keeping with this preference, EQ-5D data was collected from the RCTs included in the clinical review. Only four studies provided EQ-5D data, all of which are shown in Table 122. Studies which reported SF-36 data or disease specific quality of life measures (such as the AVVQ or CIVIQ2) without EQ-5D are not included here.

	Relevant	Utility value	Utility values					
Study	comparators	Baseline	3 months	6 months	12 months	24 months		
Carradice 2009 ⁴⁶¹	EVLA + phlebectomy	0.81 (0.79- 1.0) ²	1.0	NR	1.0 (1.0 – 1.0) ²	NR		
	EVLA without phlebectomy	0.83 (0.75 - 1.0) ²	0.82	NR	1.0 (0.89 – 1.0)²	NR		
Carradice 2011 ⁴⁸	Surgery	0.84 (0.8 – 1.0) ²	NR	NR	1.0 (0.84 – 1.0) ²	NR		
	EVLA	0.85 (0.8 – 1.0) ²	NR	NR	1.0 (0.87 – 1.0) ²	NR		
Michaels	Surgery	0.76 (0.19)	NR	0.89 (0.13)	0.87 (0.14)	0.84 (0.21)		
2006 ¹⁷⁰ (Group 3 only: severe varicose veins)	Conservative care	0.77 (0.18)	NR	0.80 (0.17)	0.78 (0.18)	0.85 (0.17)		
Shadid 2012 ²⁴⁴	Shadid 2012 ²⁴⁴ Surgery		Change from baseline at 2 years: +0.064 (NR)					
	Foam sclerotherapy	Change from	n baseline at 2	2 years: +0.061	l (NR)			

Table 122: EQ-5D data from clinical trials

Abbreviations: NR = not reported

Values are mean (SD) EQ-5D scores unless otherwise stated

¹ Mean utility values estimated from low resolution graph. Graph also reports data at 1 week and 6 week follow-up ² median (interquartile range)

EQ-5D is not consistently reported in the trials. It is clear from Table 122 that this outcome is reported at different follow-up times for the different comparators, thus the evidence does not lend itself to an accurate comparison of the quality of life after each treatment individually. A search of the economic and quality of life literature was therefore carried out to supplement the EQ-5D data found in the trials. The search identified two economic analyses which included EQ-5D data^{106,170}: Gohel and colleagues employed the baseline and post treatment EQ-5D scores from the surgery arm in Michaels, and the modelling section of Michaels used a combination of SF-6D and EQ-5D from the same trial. Two additional economic evaluations^{84,223} were found which used utility data calculated from the SF-6D. None of these economic analyses were considered beneficial in informing utility inputs for our model.

The search also identified two further randomised trials which included EQ-5D data.^{90,189} Neither of these studies were included in the clinical review for this guideline because the treatments compared in these trials were not relevant to the clinical questions included in this guideline. These studies were therefore not considered to be useful in informing inputs for the model.

An additional source of utility data is the Patient Reported Outcome Measures (PROMs), collated by the Department of Health (DH). Since 2009, the DH has required providers of varicose veins surgeries in England to collect and report PROMs. In practice, this means that all providers of NHS-funded varicose vein surgeries are expected to invite patients to complete a pre-operative questionnaire. Post-operative questionnaires are then sent to patients at least 3 months following their operation. The questionnaires completed by the patient, record self-reported health status assessed through a mixture of generic (EQ-5D and EQ-VAS) and condition-specific (AVVQ) questions. Where EQ-5D data is collected, this can be used to calculate the mean pre- and post-treatment utility scores of individuals receiving these treatments across England.

As of October 2012, finalised data are available for April 2010- March 2011. 8,624 records are available from varicose veins patients with valid EQ-5D responses in both pre- and post-operative questionnaires.¹²⁰ The mean utility score pre- and post-treatment, is available on the HES website, however this data does not specify results by varicose veins procedure. In theory, a dataset can be purchased from the Department of Health which would allow the data to be analysed by varicose veins procedure. However, given the likely population biases and computational time associated with analysis of such a large, incomplete data set, it was not thought that the benefits of purchasing this data set would justify the cost.

The PROMs data available from the HES website¹¹⁹ is documented in Table 123.

	Mean EQ-5D	SD	Number of completed questionnaires
Baseline	0.746	0.234	14533
Health gain post treatment	+0.096	0.256	8624 ¹

Table 123: PROMs data

¹ all valid post-operative questionnaires, for which there is a valid pre-operative questionnaire

Neither the data from the clinical review, nor the PROMs data provide reliable differential figures on the increase in utility following the different types of treatment. Therefore in the model patients receive the same increase in utility after treatment, regardless of treatment type. The PROMs data was used in preference to the clinical trial data in the model, as it reflects the mean change in utility for individuals undergoing treatment for varicose veins in routine clinical practice.

The baseline value was used in the model to represent the utility of a patient with primary varicose veins, i.e. when a patient first receives treatment. As PROMs data is measured at a minimum of 3 months after treatment, the health gain was applied 3 months after completed treatment (either

initial or secondary). The increase in utility over the 3 months immediately after treatment was assumed to be linear, and was applied in the model through a series of tunnel states until the 3 month post utility value was achieved. For the probabilistic analysis, the baseline value was modelled with a Beta distribution, and the health gain was modelled with a Lognormal distribution, as specified in Table 118.

Utility decrement associated with recurrent varicose veins

We conducted a search to investigate whether recurrent varicose veins were associated with a different level of QoL to primary varicose veins. Two studies^{21,198} were identified in this area, although neither reported utility values. One study mentioned SF36 data but was only available in abstract form,¹⁹⁸ and the other reported SF-36 data.²¹ Both of these papers were co-authored by GDG members, who we approached for further information, yet unfortunately no further data was available.

In 2008, Ara and Brazier published a method of predicting mean EQ-5D preference based index score using published mean cohort statistics from the eight dimensions of the SF-36 health profile.⁹ Therefore, in the absence of any utility data, we mapped the SF-36 data from Beresford 2003²¹ to the EQ-5D. In order to use the mapping algorithms, values for each of the eight dimensions of the questionnaire are required. These values were only reported in graphical format, and were therefore estimated using Grab it!, a programme which can be used to digitise graphs. The estimation was made 3 times, and a mean value taken. The resulting values for each of the SF-36 domains are documented in Table 124.

Ara and Brazier present several different equations to predict EQ-5D from SF-36, the choice of which depends on the outcome to be mapped. Ara and Brazier state 'when comparing incremental differences between study arms or changes over time, Equation 4 is the preferred choice'; the outcome of interest here was the difference between the utility of people with primary and recurrent varicose veins, thus Equation 4 was chosen. No measure of uncertainty was provided in the graph, so the mapping algorithm was applied deterministically. The results of the mapping exercise, including the difference in utility between individuals with primary and recurrent varicose veins, are provided in Table 124. In the model, the utility of individuals with recurrent varicose veins was calculated by subtracting the difference from the primary varicose veins utility weight, and was modelled probabilistically using a Lognormal distribution (Table 118).

	PF	SF	RP	RE	МН	VT	ВР	GH	EQ-5D	Difference
Primary	82.1	87.1	78.8	87.4	77.5	64.0	71.7	74.8	0.907	
Recurrent	70.7	75.1	63.8	75.2	65.8	53.7	62.2	64.5	0.814	0.093

Table 124: SF-36 and EQ-5D data for primary and recurrent varicose veins

Abbreviations: PF = physical functioning; SF = social functioning; RP = role – physical; RE = role – emotional; MH = mental health; VT = vitality; BP = bodily pain; GH = general health

Utility for conservative care

As mentioned previously, conservative care was modelled separately to the main analysis. The difference in utility between patients undergoing surgery and conservative care was used to calculate the difference in QALYs over time between these two treatments. The difference in utility between these two treatments was taken from Michaels and colleagues¹⁷⁰ (see Table 122), as this was the only paper found to report such data. Utility values are given at baseline, 1 month, 6 months, 12 months and 24 months post treatment; however by the 24 month follow-up, a large proportion of individuals had been lost to follow-up, and an unexpected large jump in utility is reported. This data was included in the base case analysis, and sensitivity analyses investigated the impact of omitting this 24 month data and extrapolating from the 12 month follow-up. The 1 month data was not included, as the GDG did not consider data collected within 3 months post treatment to be reliable,

and because short term follow-up utility data was not included for the other treatment modalities. The difference in utility was adjusted for the difference at baseline, and changes in utility over time (for example between baseline and 6 months) were assumed to be gradual and linear. For the probabilistic analysis the difference between utility following conservative care and surgery was modelled using a Normal distribution to allow positive and negative differences.

L.2.3.4 Resource use and costs

Costs were associated with the following health states: initial treatment episode, physical symptoms with recurrent VVs (1), second treatment episode and physical symptoms with recurrent VVs (2). The cost of the initial and second treatment episodes included the cost of a main treatment, as well as top-up treatment where applicable. The costs borne in the recurrent VVs states when no interventional treatment was being delivered were due to the on-going costs of conservative care given to people in those states.

Cost of interventional treatments

NHS reference costs do not distinguish between the different varicose vein treatments, but rather an overall cost is given for primary unilateral varicose veins procedures (differs whether the procedure is conducted as a day case, outpatient procedure etc.). Consequently NHS references costs could not be used to capture the different costs of the treatments.

A review of existing economic literature was conducted in order to identify the costs of the various treatments. Five UK studies^{30,106,143,170,257} were identified (See Table 125).

Study	Surgery	Endothermal	FS	СС	Costing technique
Bountouroglou 2006 ³⁰	£1,120.64	NA	£672.97	NA	Costs collected alongside RCT
Gohel 2010 ¹⁰⁶	£980	EVLA £1,524 RFA £776	£202	£O	Based on NHS reference costs, adapted with additional information from manufacturers and list prices.
Lattimer 2012 ¹⁴³	NA	£724.72 (£676.74 - £773.85) ¹	£126.39 (NR) ¹	NA	Costs collected alongside RCT
Michaels 2006 ¹⁷⁰ and Ratcliffe 2006 ²²³	£642.66 (236.39) ²	NA	NA	£267.52 (350.91) ^{2,3}	Costs collected alongside RCT
Subramonia 2010 ²⁵⁷	£559.13	£1,275.90	NA	NA	Costs collected alongside RCT

Table 125: UK relevant cost estimates from existing economic literature

All costs are mean initial treatment costs unless specified. Abbreviations: FS = foam sclerotherapy; CC = conservative care ¹Median (interquartile range)

²Mean costs from group 3; severe varicose veins randomised to surgery or conservative care (SD). ³Total undiscounted cost to NHS over 24 month period

It is clear from Table 125 that cost estimates obtained from the literature varied considerably, and as such the GDG did not think these costs to be a reliable representation of UK practice. GDG members attempted to gather cost information from their trusts, but there was inconsistency in how these estimates were derived, so the GDG decided to construct cost estimates using a bottom up approach.

Resource use was therefore based on GDG estimates. Where possible, unit costs for these resources were collected from nationally available lists such as the NHS reference costs, or the PSSRU. However, it was not always possible to find such costs, and in such cases unit costs were based on GDG estimates. The cost estimates for the model are presented in the subsequent sections. The estimates were intended to capture the differences between the costs, and therefore some aspects (for example the cost of the initial appointments), have been omitted, as these are assumed not to differ greatly between treatments. The cost of compression following treatment was limited to bandages (applied immediately) and one pair of stockings, in line with the recommendation that prolonged compression should not routinely be provided.

The majority of the unit costs provided below do not have an associated measure of uncertainty and were therefore not modelled probabilistically. Probabilistic modelling was possible where unit costs were taken from the NHS reference costs; a gamma distribution was fitted by manually adjusting the standard error of the mean until the interquartile range of the distribution best matched that reported for the unit cost. A gamma distribution was chosen so that the distribution was constrained at zero (to avoid negative costs) and reflect the positive skew normally seen in cost data. Total costs were subject to extensive deterministic sensitivity analyses.

Surgery

The breakdown of costs for surgery is provided in Table 126.

The GDG noted that greater perioperative care would be needed with surgery than with the other treatments, thus a perioperative care estimate was included. No reliable figure was available to reliably cost a few extra hours spent on a ward, thus the Band 5 time which would be spent looking after a surgery patient was used as a proxy to capture the difference in perioperative care between different treatments.

		Hours/units	Point		
Components	Unit cost	Required	estimate	Distribution	Source
Pre-op assessment (Band 5)	£82.00	0.25 hours	£20.50	NA	PSSRU ⁶²
Band 5	£82.00	1 hours	£82.00	NA	PSSRU ⁶²
Band 5 (anaesthetic assistant)	£82.00	1 hours	£82.00	NA	PSSRU ⁶²
Healthcare Assistant Band 3	£20.00	1 hours	£20.00	NA	PSSRU ⁶²
Consultant: surgical	£136.00	1 hours	£136.00	NA	PSSRU ⁶²
Consultant anaesthetist	£136.00	1 hours	£136.00	NA	PSSRU ⁶²
Disposables ^a	£250.00	1	£250.00	NA	GDG estimate
Duplex	£52.84	1	£52.84	Gamma	NHS reference costs ⁷⁷
Stockings	£5.99 ³	1	£5.99	NA	Cost of TED stockings: NHS supply chain catalogue ¹⁸⁷
Perioperative care					
Band 5 ^b	£82.00	1.5 hours	£123.00	NA	PSSRU ⁶²
Total			£908.33		

Table 126: Costs - Surgery

^a Includes gowns, surgical instruments, drapes, bandages and other disposable items

^bBased on 90mins pre-op where patient is looked after by ¼ nurse, 30 minutes post-op where patient is looked after by 1 nurse, additional 150mins post op ¼ nurse time

Endothermal techniques

The majority of the cost components for the two types of endothermal treatment (RFA & EVLA) were considered to be the same as each other, with the only differences being the cost of the generators, catheters and the controlled laser area required for EVLA. The cost of the catheter and generator vary, as commercial companies have individual contracts with different trusts. The costs in Table 127 have been provided by commercial companies; due to the business sensitive nature of this information individual company names have been removed.

Procedure	Company	Generator cost	Catheter cost		
EVLA	Company A	Provided on long term loan free of charge (List price is £15,000 - £22,000 but very rarely bought)	£180 - £245		
	Company B	Provided on long term loan free of charge (List price is £12,500 but very rarely bought)	£200		
RFA	Company C	Provided on long term loan free of charge	£300		
Company D		Provided on long term loan free of charge (List price is £10,000 but very rarely bought)	£250 - £300		

Table 127: Catheter costs for EVLA and RFA

As indicated in Table 127, in the vast majority of cases the generator is loaned free of charge, usually on the condition that the hospital carries out a certain amount of procedures per year. The cost of the generator was therefore not considered in the analysis. The costs of the catheters were approximated based on various estimates provided by commercial companies. Deterministic sensitivity analyses will investigate the impact of changes in cost of catheters.

The GDG decided not to explicitly include the cost of the controlled laser area needed for EVLA in the analysis, as the room would be used for a variety of laser procedures, and the cost per treatment would be highly dependent on the number of procedures undertaken.

In practice, the two endothermal treatments (EVLA and RFA) compete directly with each other, and this guideline assumes they have equal clinical effectiveness. The implication of this is that, in this analysis, whichever of these two treatments is the cheapest will be cost-effective. However, due to uncertainty around the costs, specifically that the RFA catheter is more expensive, but the EVLA requires a laser controlled area, it is not straight forward to identify which of these treatments is cheaper. For the purpose of this analysis it was assumed that once the laser controlled area had been accounted for, EVLA would cost no more than RFA; the cost of RFA was therefore assumed to be the maximum cost of endothermal treatment. This maximum cost was used in the model base case, and costs were explored thoroughly through sensitivity analysis. The breakdown of the costs is provided in Table 128.

Components	Unit cost	Hours/units Required	Point estimate	Distribution	Source
RFA catheter	£300	1	£300	NA	Table 127
EVLA catheter	£200	1	£200	NA	Table 127
Disposables ^a	£86.00	1	£86.00	NA	GDG estimate
Normal Saline £8.50 box of 20	£0.43	1	£0.43	NA	GDG estimate
Band 5	£82.00	0.75 hours	£61.50	NA	PSSRU ⁶²
Healthcare Assistant Band 3	£20.00	0.75 hours	£15.00	NA	PSSRU ⁶²
Consultant	£136.00	0.75 hours	£102.00	NA	PSSRU ⁶²
Duplex	£52.84	1	£52.84	Gamma	NHS reference costs ⁷⁷
Stockings	£5.99	1	£5.99	NA	NHS supply chain catalogue ¹⁸⁷
RFA Total			£623.76		
EVLA Total			£523.76 + laser controlled area		

Table 128: Costs – Endothermal treatment

(a) includes gowns, procedure pack, surgical instruments, drapes, bandages, syringes and other disposable items

Foam sclerotherapy

The breakdown of costs for Sclerotherapy is provided in Table 129.

Components	Unit cost	Hours/units Required	Point estimate	Distribution	Source
Consultant time	£136.00	0.75 hours	£102.00	NA	PSSRU ⁶²
Clinical nurse specialist time	£91.00	0.75 hours	£68.25	NA	PSSRU ⁶²
Disposables ^a	£50.00	1	£50.00	NA	GDG estimate
Stockings (class II)	£42.30	1	£42.30	NA	NHS drug tariff ¹⁸⁶
Duplex	£52.84	1	£52.84	Gamma	NHS reference costs ⁷⁷
Total			£315.39		

Table 129: Costs - Sclerotherapy

(a) Includes gown, needles, bandages, syringes and other disposable items

Conservative care

The breakdown of costs for conservative care is provided in Table 130.

Based on clinical opinion, it was assumed that half of the patients who receive conservative care return to their GP in the first year for further advice and reassurance. After the first year, the annual costs were based on the assumption that the patient visits the practice nurse for a few routine

appointments over the course of the year, for advice and to be re-measured for stockings. We acknowledge that in practice some of this measuring may be done by a pharmacist.

Components	Unit cost	Hours/units required	Point estimate	Source
First year costs				
GP visits	£30.00	0.5	£15.00	PSSRU ⁶²
Annual costs				
Practice nurse time	£43.00	1.5 hours	£64.50	PSSRU ⁶²
Stockings (class II)	£42.30	4 ^a	£169.20	NHS drug tariff ¹⁸⁶
First year total			£248.70	
Annual total			£233.70	

Table 130: Costs – Conservative care

(a) based on an estimated lifespan of three months per stocking

Additional costs of retreatment

There are likely to be additional costs associated with re-treatment, over and above the cost of the second treatment itself. The additional costs associated with a second treatment episode were based on clinical opinion, and are provided in Table 131.

Table 131: Additional costs for retreatment

Components	Unit cost	Hours/units required	Point estimate	Distribution	Source
GP visit	£30.00	2.5	£75.00	NA	PSSRU ⁶²
OP 1st attendance vascular surgery	£165.49	1	£165.49	Gamma	NHS reference costs ⁷⁷
OP 2+ attendance vascular surgery	£123.28	1	£123.28	Gamma	NHS reference costs ⁷⁷
Duplex scan	£52.84	1	£52.84	Gamma	NHS reference costs ⁷⁷
Total			£416.61		

Note the resource components here have been replicated from the aforementioned on-going HTA project²³⁹

L.2.4 Sensitivity analyses

The sensitivity analyses described in Table 132 were undertaken to explore the effect of different parameter inputs and assumptions on the results of the model. The results of all sensitivity analyses are presented in section L.3.1.

Table 132: Alternative values and descrip	otions for deterministic sensitivity analyses

SA1 Baseling rate	ne recurrence	Some members of the GDG felt that the baseline	0.00204	
	Baseline recurrence rate	Some members of the GDG felt that the baseline recurrence rate following surgery, to which the relative effects from the NMA were applied, was too high. The baseline rate was calculated from clinical recurrence reported in randomised trials, and could be higher than those observed in UK practice for several reasons. Sensitivity analyses employ different baseline rates of recurrence	0.00384	Recurrence rate from Shadid2012 ²⁴⁴ (lowest recurrence rate from included trials)
			0.01548	Recurrence rate from Carradice2011 ⁴⁸ (highest recurrence rate from included trials)
SA2 Endoth treatme concurr phlebeo	ent without rent	The need for top up treatment and cost of procedure is likely to be different if concurrent phlebectomy is not carried out. This sensitivity analysis evaluates the cost effectiveness of endothermal treatment without phlebectomies compared to the other treatments	10% require top up treatment Cost of procedure: £272.27 + catheter	The need for top up treatment will be higher, and the cost of procedure will be slightly lower; clinical evidence does not distinguish between endothermal with/without phlebectomy thus probability of clinical recurrence remains unchanged
SA3 Utility f	for vative care	The data used for utility of conservative care includes a sharp increase in utility at 2 years. This increase is dramatic and unexpected, thus in this SA we omit the two year data	1 year adjusted difference between utility of conservative care and surgery: -0.1	The adjusted value reported at one year is extrapolated over the 5 year time horizon
SA4 GSV + S	SSV	Treating an additional truncal vein will extend procedure time and have an impact on total procedure cost	S: £1,119.12 F: £329.84 E: £691.27	Cost increase compared to base case due to an extra 15 minutes treatment time
SA5 Time ho	orizon	The time horizon is shortened to avoid extrapolation past the maximum follow-up time found in the data	3 years	3 years is the longest follow-up of the trials included in the NMA
SA6 Costs		Various SAs to investigate how robust the model is to the costs of treatment, around which there is great uncertainty	Relative costs manipulated and costs for conservative care reduced	The GDG had no strong indication of what plausible ranges for treatment costs were; therefore threshold analyses conducted within an arbitrary but wide interval
SA6b Cost of	f catheters for	These SAs investigate how robust the model is to	EVLA catheters:	Maximum and minimum values (as at

Analysis	Parameter	Description of sensitivity analysis	Values	Comment
	endothermal treatment	the costs of the endothermal catheters, around which there is great variability	£180-£245 RFA catheters: £250- £300	October 2012) provided by commercial companies
	Top-up treatment	These SAs explore the impact of the GDG	S: 0-5%	Threshold sensitivity analyses within plausible
SA7	rates	estimate of the proportion of patients who will	F: 10-100%	range suggested by GDG members
		need top up treatment	E: 0-5%	
	Proportions receiving	The proportion receiving a conservative care 7	75%	Arbitrary, wide ranging values.
SA8	conservative care following clinical recurrence (instead of re-treatment)	following clinical recurrence is varied (the remainder receive a second treatment episode)	50%	
SA9	Proportions receiving	The type of retreatment a patient would receive	S:20%; F:10%, E:70%	Sensitivity analyses use alternative
	each type of treatment during the second treatment episode	would be highly dependent on the nature of the recurrence and further patient characteristics. This SA investigates the impact of the assumptions around the proportions of patients	S:5%; F:5%, E:90%	proportions suggested by individual GDG
			S:10%; F:45%, E:45%	members.
			S:15%; F:80%, E:5%	
		receiving each type of retreatment	S:10%; F:60%, E:30%	
			S:10%; F:50%, E:40%	

Abbreviations: F = Foam; S= Surgery; E= endothermal

1 L.2.5 Bilateral treatment

The model base case only considered unilateral patients, yet consideration should also be given to treatment of bilateral patients. The model does not lend itself to bilateral analysis, as significant assumptions would have to be made around whether top-up treatment was complete in one or two legs, whether clinical recurrence was experienced in one or two legs, and whether both legs were retreated. Furthermore, utility increases and decrements as used in the unilateral model would no longer be applicable. The GDG therefore decided that a cost-comparison was the preferred method to analyse the treatment of bilateral patients.

9 In order to calculate costs of bilateral surgery and endothermal treatment, a proportional increase 10 was applied to the unilateral costs documented in Table 126 and Table 128. A variety of scenarios were presented in which this proportional increase was varied, in order to capture uncertainty. The 11 maximum that bilateral treatment could be expected to cost would be 200% of the cost of unilateral 12 13 treatment, as would be the case if both legs were to be treated completely separately. Therefore the maximum cost of bilateral treatment was assumed to be 200% of the costs specified in Table 126 14 Table 128. The NHS reference costs⁷⁷ indicate that for day case procedures, bilateral treatment costs 15 16 112% of the cost of unilateral treatment; this was taken as the minimum proportional increase in 17 costs.

- 18 The bilateral cost of foam sclerotherapy was assumed to be twice the cost of unilateral treatment. 19 This is because there are consensus recommendations on the maximum amount of sclerosant foam 20 which should be given per session.³⁶ The recommended maximum volume per session is 10ml, and the recommended average is lower, between 2 and 8ml of sclerosant foam. In many cases this would 21 22 prevent treatment of both legs in one sitting; indeed the recommendations add that it is advisable to 23 limit the amount of sclerosant foam given per session, even if this means the patient requires more 24 than one treatment. As the costs of initial appointment have been omitted, treating the legs 25 separately can be considered equivalent to two unilateral cases from a costing point of view.
- The cost of conservative care for bilateral treatment was calculated by doubling the number of
 stockings required. The number of GP appointments and practice nurse time was assumed to stay
 the same as with unilateral treatment.
- 29 The results of the cost comparison are documented in section L.3.2.

30 L.2.6 Computations

The model was constructed in Microsoft Excel and was evaluated by cohort simulation.

32 L.2.6.1 Calculating cost effectiveness

- The widely used cost-effectiveness metric is the incremental cost-effectiveness ratio (ICER). This is calculated by dividing the difference in costs associated with two alternatives by the difference in QALYs. The decision rule then applied is that if the ICER falls below a given cost per QALY threshold the result is considered to be cost effective. If both costs are lower and QALYs are higher the option is said to dominate and an ICER is not applicable.
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 $ICER = \frac{Costs(B) - Costs(A)}{QALYs(B) - QALYs(A)}$

- Cost-effective if: ICER < Threshold
- Where: Costs/QALYs(X) = total costs/QALYs for option X

When there are more than two comparators, as in this analysis, options must be ranked in order of increasing cost then options ruled out by dominance or extended dominance before calculating ICERs for the remaining options.

It is also possible, for a particular cost-effectiveness threshold, to re-express cost-effectiveness results in term of net monetary benefit (NMB). This is calculated by multiplying the total QALYs for a comparator by the threshold cost per QALY value (for example, £20,000) and then subtracting the total costs (formula below). The decision rule then applied is that the comparator with the highest NMB is the most cost-effective option at the specified threshold. That is the option that provides the highest number of QALYs at an acceptable cost.

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Net Benefit(X) = $(QALYs(X) \times \lambda) - Costs(X)$	Cost-effective if:
Where: Costs/QALYs(X) = total costs/QALYs for option X; λ = threshold	highest net benefit

11Both methods of determining cost effectiveness will identify exactly the same optimal strategy. For12ease of computation NMB was used to identify the optimal strategy in the probabilistic analysis13simulations.

14 The probabilistic analysis was run for 10,000 simulations. Each simulation, total costs and total QALYs 15 were calculated for each strategy. Net benefit was also calculated and the most cost-effective option 16 identified (that is, the one with the highest net benefit), at a threshold of £20,000 per QALY gained. 17 The results of the probabilistic analysis were summarised in terms of mean costs, mean QALYs and 18 mean net benefit for each treatment option, where each was the average of the simulated estimates. 19 The option with the highest mean net benefit (averaged across the simulations) was the most cost-20 effective at the specified threshold. The percentage of simulations where each strategy was the most 21 cost-effective gives an indication of the strength of evidence in favour of that strategy being cost-22 effective.

Results are also presented graphically where mean total costs and mean total QALYs for each
 treatment option is plotted. Comparisons not ruled out by dominance or extended dominance are
 joined by a line on the graph where the slope represents the incremental cost-effectiveness ratio, the
 magnitude of which is labelled.

27 L.2.7 Model validation

The model was developed in consultation with the GDG; model structure, inputs and results were
 presented to and discussed with the GDG for clinical validation and interpretation.

The model was systematically checked by the health economist undertaking the analysis; this included inputting null and extreme values and checking that results were plausible given inputs. The model was peer reviewed by an experienced health economist who had not been involved in the guideline; this included systematic checking of the model calculations.

34 L.2.8 Interpreting results

NICE's report 'Social value judgements: principles for the development of NICE guidance' sets out the
 principles that GDGs should consider when judging whether an intervention offers good value for
 money. In general, an intervention was considered to be cost effective if either of the following
 criteria applied (given that the estimate was considered plausible):

The intervention dominated other relevant strategies (that is, it was both less costly in terms of resource use and more clinically effective compared with all the other relevant alternative strategies), or

• The intervention costs less than £20,000 per quality-adjusted life-year (QALY) gained compared with the next best strategy.

3 L.3 Results

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Detailed results are presented over the next few pages for the base case and various sensitivity
analyses. As the results of the deterministic and probabilistic analysis were comparable, all results
reported below are means from the probabilistic analysis unless otherwise specified.

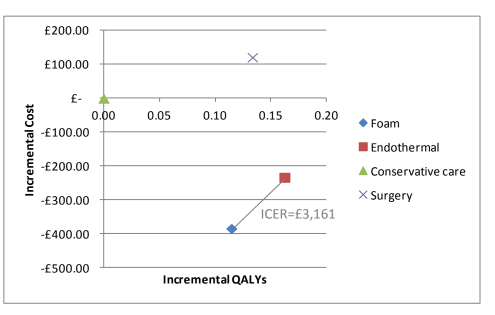
7 Table 133 and Figure 161 show the base case results. Both conservative care and surgery were
8 dominated, as they provided less QALYs at increased cost when compared to endothermal
9 treatment. As these strategies are dominated, they are not further considered in the incremental
10 analysis and the ICER is not calculated.

Mean per patient NMB at threshold **Rank at threshold Probability of** QALYs Cost of £20,000 of £20,000 being CE^a Treatment Conservative care 3.55 £1,102 £69,965 4 4% 3 £1,222 Surgery 3.69 £72,554 3% 2 Foam 3.67 £718 £72,681 23% sclerotherapy Endothermal 3.72 £869 £73,484 71% 1

11 Table 133: Mean base case results (probabilistic)

(a) For interpretation of the probability of being cost-effective see section L.2.6.1.

Figure 161: Cost effectiveness plane showing incremental cost and QALYs per patient expected with each strategy (Base case, probabilistic analysis)



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In the base case analysis, the strategy which provided the most QALYs was endothermal treatment. However, this came at an additional cost compared to foam sclerotherapy. Using the mean costs and QALYs generated over the probabilistic sensitivity analysis, the ICER of endothermal treatment compared to foam was £3,161 which is below the NICE threshold of £20,000 per QALY gained. Endothermal treatment had a probability of being cost-effective of 71%, followed by foam which had a lower chance of being the most cost-effective option of 23%. Disaggregating the results of the analysis by cost and QALYs allows us to examine the impact of key components of the model on the overall results. The QALYs associated with the initial treatment episode are the same for each treatment, therefore we know that that the difference in total QALYs is driven by the reduction in QoL associated with recurrence. Endothermal treatment has the lowest probability of recurrence per cycle, thus the results of the model align with our expectation that this treatment would lead to the highest total QALYs.

Table 134 provides the breakdown of total cost (the probabilistic costs of the initial treatment
episode are comparable to the deterministic estimates in Table 126 – Table 130). It shows that whilst
the costs due to recurrent treatment do differ (Note – this is the cost of retreatment averaged across
all patients), the difference in total costs between treatment methods is mainly due to the initial
treatment costs. Sensitivity analyses explored the impact of changes in the treatment costs – see
below.

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Table 134: Breakdown of total costs (probabilistic base case)

Treatment	Cost of initial treatment episode	Cost of recurrent treatment ^a	Total cost
Conservative care ^b	N/A	N/A	£1,102
Surgery	£924	£299	£1,222
Foam sclerotherapy	£378	£340	£718
Endothermal	£639	£230	£869

(a) This is the average cost of treatment and management of recurrent varicose veins weighted by the proportion of individuals who will require this, therefore this represents the expected per person cost of recurrence.

(b) Initial and recurrent treatment costs are not applicable for conservative care as this was modelled separately as an ongoing management technique

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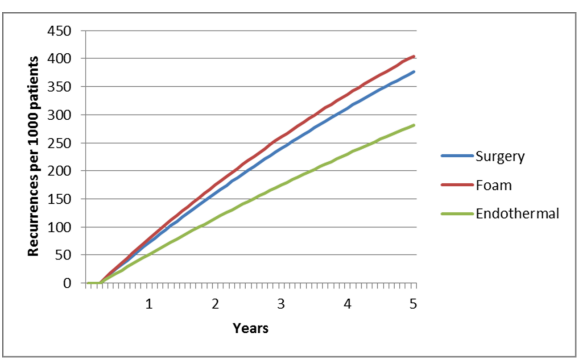
The number of clinical recurrences over time is shown in Figure 162 for each treatment. The GDG felt these values were acceptable, but noted that it was difficult to judge face validity of these results, as the majority of the GDG members do not see all clinical recurrence cases, only those patients who are to be retreated. Sensitivity analyses investigated the impact of changing the level of clinical recurrence.

24 L.3.1 Sensitivity analyses

25 Sensitivity analyses were run probabilistically unless otherwise stated. In all analyses endothermal 26 treatment was recorded as the optimal strategy. Table 136 summarises the results of these analyses. 27 Throughout all of the probabilistic sensitivity analyses, neither the probability of conservative care or 28 surgery being the optimal strategy rose above 5%. Overall, the sensitivity analyses demonstrated that 29 the results of this analysis were robust to changes in key assumptions, recurrence rates, and 30 substantial changes in relative costs.

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Figure 162: **Clinical recurrence over time**



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Bilateral treatment 3 L.3.2

The results of the cost comparison are presented in Table 135. The GDG discussed the figures in this table alongside the bilateral results, and concluded that endothermal treatment is likely to be the cost-effective treatment strategy for bilateral treatment.

Table 135: Bilateral treatment cost comparison (deterministic)

Cost of unilateral treatment	Multiplier	Cost of bilateral treatment				
Conservative care						
£234ª	NA	£403ª				
	Surgery					
£924	112%	£1,035				
	120%	£1,109				
	140%	£1,294				
	160%	£1,479				
	180%	£1,663				
	200%	£1,848				
	Endothermal					
£640	112%	£716				
	120%	£767				
	140%	£895				
	160%	£1,023				
	180%	£1,151				
	200%	£1,279				
	Foam sclerotherapy					
£378	200%	£757				

(a) represents annual cost

	Mean QALYs per patient				Mean costs per patient				Optimal	Probability CE at
Sensitivity analysis	сс	S	FS	E	сс	S	FS	E	strategy	£20,000 threshold
				SA1 : Baselin	e recurrence	e rate				
SA1a: Lowest baseline recurrence	3.62	3.75	3.74	3.77	£1,102	£1,067	£548	£746	Endothermal	74%
SA1b: Highest baseline recurrence:	3.48	3.61	3.59	3.66	£1,102	£1,402	£901	£1,015	Endothermal	74%
				SA2 – SA5	: Assumptio	ns				
SA2: Endothermal treatment does not include phlebectomy	3.55	3.69	3.67	3.72	£1,102	£1218	£713	£822	Endothermal	73%
SA3: Utility for conservative care extrapolated from 1 year value	3.25	3.69	3.67	3.72	£1,102	£1224	£722	£871	Endothermal	75%
SA4: GSV + SSV	3.55	3.69	3.67	3.72	£1,102	£1355	£805	£943	Endothermal	72%
SA5: 3 year time horizon	2.19	2.32	2.31	2.33	£699	£1085	£569	£761	Endothermal	62%
				SA6: Costs	(determinis	tic)				
SA6i: Surgery, sclerotherapy, conservative care costs reduced by 50%, cost of endothermal remains as base case	3.56	3.69	3.68	3.73	£558	£687	£432	£789	Endothermal	N/A
SA6ii: Below knee standard stockings for conservative care (£11 instead of £42 per pair)	3.56	3.69	3.68	3.73	£511	£1,179	£653	£828	Endothermal	N/A
SA6iii: Below knee standard stockings and no practice nurse time for conservative care	3.56	3.69	3.68	3.73	£211	£1,162	£635	£816	Endothermal	N/A
SA6iv: Threshold cost analysis of increase in cost of				t would have tive. In this in		-			atments remain	the same in order to

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	Mean OA	LYs per patie	ent		Mean cost	ts per patien	t			
Sensitivity analysis	СС	S	FS	E	СС	S	FS	E	Optimal strategy	Probability CE at £20,000 threshold
endothermal treatment						-				
			SA6b: Cost	of catheters	for endothe	ermal treatm	ent			
£180	3.55	3.69	3.67	3.72	£1,102	£1,209	£703	£738	Endothermal	74%
£395	3.55	3.69	3.67	3.72	£1,102	£1,233	£730	£970	Endothermal	67%
		SA7	: Top-up trea	tment rate -	threshold an	nalyses (dete	rministic)			
Surgery 0-10%				Ν	/A				Endothermal	N/A
Foam sclerotherapy 10-100%				N	/A				Endothermal	N/A
Endothermal 0-10%				N	/A				Endothermal	N/A
	SA8: Pr	oportions re	ceiving cons	ervative care	following fi	rst recurrenc	e instead of	retreatment		
SA8a: 75%	3.53	3.66	3.64	3.70	£1,102	£1,159	£648	£820	Endothermal	72%
SA8b: 50%	3.54	3.68	3.66	3.71	£1,102	£1,191	£682	£844	Endothermal	72%
	SA9: Propor	tions receivi	ng each type	of treatmen	t during the	second treat	tment episod	de (determin	istic)	
S:20%; F:10%; E:70%	3.56	3.69	3.68	3.73	£1,102	£1,235	£714	£869	Endothermal	N/A
S:5%; F:5%; E:90%	3.56	3.69	3.68	3.73	£1,102	£1,227	£706	£863	Endothermal	N/A
S:10%; F:45%;E:45%	3.56	3.69	3.68	3.73	£1,102	£1,208	£685	£849	Endothermal	N/A
S:15%; F:80%; E:5%	3.56	3.69	3.68	3.72	£1,102	£1,192	£668	£837	Endothermal	N/A
S:10%; F:60%; E:30%	3.56	3.69	3.68	3.72	£1,102	£1,200	£676	£843	Endothermal	N/A
S:10%; F:50%; E:40%	3.56	3.69	3.68	3.73	£1,102	£1,206	£682	£847	Endothermal	N/A

Cost-effectiveness analysis of interventional treatments and conservative care

1 L.4 Discussion

2 L.4.1 Summary of results

This analysis found that endothermal treatment is the most clinically and cost effective treatment strategy for people with varicose veins (note - EVLA and RFA were considered together in the model, and the results do not distinguish between these two endothermal techniques). This conclusion was robust to a wide range of sensitivity analyses, demonstrating that although uncertainty surrounds model inputs, variation within reasonable ranges does not change the results.

8 An area of particular uncertainty was the costs, yet sensitivity analyses revealed that the model is 9 robust to changes in relative costs. For example, even if the differences in costs have been 10 underestimated, endothermal treatment would remain the optimal strategy even if the costs of all 11 the other treatments are half of what we estimated in the base case. If the costs of surgery, 12 sclerotherapy and conservative care remain as specified in the base case, endothermal treatment 13 remains cost effective even with increases in cost of up to £681.

14 L.4.2 Limitations and interpretation

The clinical review was not designed to distinguish between different types of endothermal treatment and as such the results presented here do not make any distinction between RFA and EVLA, or any further variation within these treatment modalities. Whilst the decision to treat the various endothermal treatments as one combined treatment was based on GDG consensus, this could be considered a limitation of the analysis.

A further limitation of the model is the specific population to which it applies. The interventions considered are only true comparators when considering patients for whom all four treatments are a possibility, and in practice this may only be a small proportion of the varicose veins population. If endothermal treatment is not suitable for a patient then foam sclerotherapy is the cost-effective option, and if foam is not suitable either, surgery is the optimal strategy. Further issues of generalizability are discussed in section L.4.3.

- An additional drawback of this analysis is that the estimates of rates of top-up treatment were based on GDG estimates, but the clinical recurrence data was based on trial outcomes. Depending on how clinical recurrence was reported, it is likely that in some instances the trials recorded what would be deemed here as a need for top-up treatment as clinical recurrence. The implication of this is that some recurrences may have been double counted. This said, sensitivity analyses revealed that the model was robust to changes in top-up rates and in clinical recurrence rates, therefore this drawback represents only a minor limitation.
- Assumptions were made around top-up treatments, as well as modality of retreatment, which could
 potentially be considered as limitations to the model. However sensitivity analyses revealed that
 reasonable changes in these assumptions did not impact the results.
- The assumptions of the network meta-analysis model necessitated a constant hazard of clinical recurrence over time. This represents a restriction of the analysis, yet this assumption was deemed reasonable over the relatively short time horizon of the model. Ideally, utility data would have been included which reflected treatment specific improvements in quality of life, however as discussed earlier, reliable data to reflect this could not be found. Use of the PROMs data brings its own limitations, such as the potential for sampling bias.
- Finally, this analysis does not attempt to answer the questions of the optimal timing of intervention, or the optimal choice of treatment at each stage of the disease. We initially hoped to address these

questions, but reliable data were not available. Consequently, conclusions are applicable to the general varicose veins population, with no separate consideration of subgroups. Input data were collected from individuals at various stages of varicose veins severity, and we cannot be certain that interventional treatment is cost-effective in each subgroup.

5 L.4.3 Generalisability to other populations / settings

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The conclusions of this analysis are expected to apply to the UK adult varicose veins population.
Whilst varicose veins are only rarely seen in children, the results of this analysis are unlikely to be generalisable to this population.

Endothermal treatment and foam sclerotherapy were assumed to take place in an outpatient setting
 and surgery as a day case procedure. The analysis has not considered different settings of treatment,
 for example endothermal treatment as a day case procedure or surgery as an inpatient procedure.
 Nevertheless, sensitivity analysis did show that the optimal strategy was fairly robust to increases in
 the cost of endothermal treatment and so if outpatient endothermal treatment was not considered
 suitable for a patient, day case endothermal treatment may represent a cost-effective alternative.

15 L.4.4 Comparisons with published studies

16Gohel and colleagues (2010)¹⁰⁶ present the only analysis published at present which compares all the17treatments included in this analysis in a UK setting. Whilst day case surgery offers the highest net18benefit, the authors conclude that RFA or EVLA performed as an outpatient procedure, or surgery19performed as a day case procedure, are likely to be cost-effective treatments, as differences in costs20and QALYs are small. The suggestion that RFA and EVLA are cost-effective aligns with our findings21that endothermal treatment is the optimal treatment strategy.

22 L.4.5 Conclusion = evidence statement

According to the results of this original economic model based on the current clinical evidence review and GDG input, it is likely that endothermal treatment is the cost effective strategy for people in whom all treatments are suitable. When endothermal treatment is not deemed suitable for a patient, foam sclerotherapy is likely to be the optimal strategy. Surgery represents the optimal choice if neither endothermal treatment nor foam sclerotherapy are thought suitable. This evidence is directly applicable, with minor limitations.

29 L.4.6 Implications for future research

A major issue which remains to be addressed is the question of which patients should be treated, which is closely related to the question of the optimal timing of intervention. To answer such a question would require data on the natural progression of varicose veins, i.e. what happens to patients who are not treated. Such data is unlikely to emerge from future research due to ethical considerations.

Future research into the effectiveness of each intervention at each stage of the disease would be a step towards solving this issue. If this data was available, future analyses could investigate whether different treatment strategies are optimal at different stages of the disease, and potential efficiencies could be realised.

Appendix M: Network meta-analysis

2	M.1	Network meta-analysis code					
3		# Binomial likelihood, cloglog link					
4		# Random effects model for multi-arm trials					
5		model{ # *** PROGRAM STARTS					
6		for(i in 1:ns){ # LOOP THROUGH STUDIES					
7		w[i,1] <- 0 # adjustment for multi-arm trials is zero for control arm					
8		delta[i,1] <- 0 # treatment effect is zero for control arm					
9		mu[i] ~ dnorm(0,.0001) # vague priors for all trial baselines					
10		for (k in 1:na[i]) { # LOOP THROUGH ARMS					
11		r[i,k] ~ dbin(p[i,k],n[i,k]) # Binomial likelihood					
12		# model for linear predictor					
13		cloglog(p[i,k]) <- log(time[i]) + mu[i] + delta[i,k]					
14		<pre>rhat[i,k] <- p[i,k] * n[i,k] # expected value of the numerators</pre>					
15		#Deviance contribution					
16		dev[i,k] <- 2 * (r[i,k] * (log(r[i,k])-log(rhat[i,k]))					
17		+ (n[i,k]-r[i,k]) * (log(n[i,k]-r[i,k]) - log(n[i,k]-rhat[i,k]))) }					
18		# summed residual deviance contribution for this trial					
19		resdev[i] <- sum(dev[i,1:na[i]])					
20		for (k in 2:na[i]) { # LOOP THROUGH ARMS					
21		# trial-specific LOR distributions					
22		delta[i,k] ~ dnorm(md[i,k],taud[i,k])					
23		# mean of LOR distributions, with multi-arm trial correction					
24		md[i,k] <- d[t[i,k]] - d[t[i,1]] + sw[i,k]					
25		# precision of LOR distributions (with multi-arm trial correction)					
26		taud[i,k] <- tau *2*(k-1)/k					
27		# adjustment, multi-arm RCTs					
28		w[i,k] <- (delta[i,k] - d[t[i,k]] + d[t[i,1]])					
29		# cumulative adjustment for multi-arm trials					
30		sw[i,k] <- sum(w[i,1:k-1])/(k-1)					

1	}
2	}
3	totresdev <- sum(resdev[]) #Total Residual Deviance
4	d[1]<-0 # treatment effect is zero for reference treatment
5	# vague priors for treatment effects
6	for (k in 2:nt){ d[k] ~ dnorm(0,.0001) }
7	sd ~ dunif(0,5) # vague prior for between-trial SD
8	tau <- pow(sd,-2) # between-trial precision = (1/between-trial variance)
9	# Provide estimates of treatment effects T[k] on the natural (probability) scale
10	# Given a Mean Effect, meanA, for 'standard' treatment A,
11	# with precision (1/variance) precA, over a time period timeA
12	A ~ dnorm(meanA,precA)
13	for (k in 1:nt) { cloglog(T[k]) <- log(timeA) + A + d[k] }
14	# pairwise HRs and LHRs for all possible pair-wise comparisons, if nt>2
15	for (c in 1:(nt-1)) {
16	for (k in (c+1):nt) {
17	lhr[c,k] <- (d[k]-d[c])
18	log(hr[c,k]) <- lhr[c,k]
19	}
20	}
21	# ranking on relative scale
22	for (k in 1:nt) {
23	rk[k] <- rank(d[],k) # assumes events are "bad"
24	best[k] <- equals(rk[k],1) #calculate probability that treat k is best
25	}
26	} # *** PROGRAM ENDS
27	

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1 M.2 Baseline model code (meta-analysis of surgery trial arms)

Z	
3	# Random effects probability model
4	model{
5	for(i in 1 : 8) {
6	r[i] ~ dbin(p[i],n[i]) #Likelihood
7	cloglog(p[i]) <- b[i] + log(time[i]) #cloglog of response
8	b[i] ~ dnorm(d,prec) #Random effects model
9	rhat[i] <- p[i] * n[i] # expected value of the numerators
10	#Deviance contribution
11	dev[i] <- 2 * (r[i] * (log(r[i])-log(rhat[i]))
12	+ (n[i]-r[i]) * (log(n[i]-r[i]) - log(n[i]-rhat[i])))
13	}
14	b.new~dnorm(d,prec) #predictive dist. (log-odds)
15	resdev <- sum(dev[])
16	d ~ dnorm(0.0,1.0E-6) #vague prior for mean effect
17	
18	cloglog(T1) <- b.new +log(1)
19	cloglog(T12) <- b.new +log(12)
20	cloglog(T24) <- b.new +log(24)
21	cloglog(T36) <- b.new +log(36)
22	
23	#logit(T.new) <- b.new
24	<pre>#sd <- 1/sqrt(prec) # gamma prior for RE precision</pre>
25	#prec ~ dgamma(0.001,0.001)
26	sd ~ dunif(0,5) # uniform prior for RE st dev
27	prec <- pow(sd,-2)
28	}
29	
30	

Appendix N: Research recommendations

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The Guideline Development Group has made the following recommendations for research, based on its review of evidence, to improve NICE guidance and patient care in the future.

5 N.1 Key future research recommendations (FRR)

6 N.1.1 Natural history of varicose veins

In people with varicose veins at CEAP (Clinical, etiological, anatomical and pathophysiological) stage
C2 or C3, what are the factors that influence progression of the disease to CEAP stages C5 or C6?

9 Why this is important

The evidence review for the guideline showed a lack of high-quality evidence on the progression of 10 varicose veins from CEAP stage C2 or C3 to more serious varicose veins disease. A large observational 11 12 prospective cohort study, similar to the Framingham or Bonn veins studies, should be undertaken. 13 The study should recruit patients with C2 and C3 disease and follow the progress of their disease 14 over a 5-year time horizon. Consideration should be given to including a genetic component in the study because genetic factors have not been studied on a large scale. The results of such a study 15 16 should help to more accurately identify which patients are at risk of developing more serious disease 17 so that interventions can be offered at an early stage to those who will benefit most.

18 Criteria for selecting high-priority research recommendations

Criterion	Explanation
Importance to patients or the population	If the research were to identify clear factors which were indicators for progression to more serious disease, those patients who are most at risk could be identified at an earlier stage and either monitored more closely or treated at an earlier stage. This would then potentially lead to more efficient use of resources.
Relevance to NICE guidance	This research is of medium relevance to the NICE guidelines. The research is relevant but is not key to future updates as a recommendation was able to be produced in its absence.
Relevance to the NHS	The research is relevant to the NHS, depending on the results it may allow the identification of people at risk, and hence may change the allocation of resources.
National priorities	This research is not relevant to a national priority area.
Current evidence base	The evidence found from the systematic reviews in the guideline were mainly case control and small cohort studies. These are not sufficient to identify all the risk factors. In addition, no large scale study has looked to determine if there is a genetic component involved in the progression of varicose veins.
Equality	The research would not discriminate against any group.
Feasibility	The proposed research is a long term project and does have a substantial cost associated

Criterion	Explanation
	with it. However, it was the opinion of the GDG that the potential benefits of the research would outweigh the costs. The issue of treatment needs to be considered. It would not be ethical to prevent patients from having treatment as their disease worsened. Treatment should be taken into account in the research design.
Other comments	None
Study design	Prospective cohort study

2 N.1.2 Optimal interventional and conservative treatments at different stages of disease

What is the optimal treatment (compression, surgery, endothermal ablation or foam sclerotherapy)
 for varicose veins at each of the CEAP stages, i.e. CEAP stages 2–3, CEAP stage 4 and CEAP stages 5–
 6?

6 Why this is important

Much of the research into the optimum treatment for varicose veins has involved patients with
varicose veins in CEAP stages C2 and C3, so little is known of the relative efficacies of treatment at
the more severe stages of disease. Furthermore, some studies have included patients with varicose
veins at a range of stages without subgrouping, which may conceal important differences in efficacy
between different treatments at different stages of disease. Hence current treatment
recommendations, which do not differentiate between patients with varicose veins at different
stages, may not be equally effective to all patients.

14 A large-scale RCT that compares the 4 main treatments (compression, surgery, endothermal ablation 15 and foam sclerotherapy) in subgroups with varicose veins at different stages is needed. The use of 16 CEAP to categorise the disease stages is not ideal because higher CEAP stages do not necessarily 17 indicate greater severity. However, other methods of categorisation are even more problematic. Quality-of-life measures are unlikely to reflect severity of disease because of variations in perception 18 of symptoms. In addition, measuring the degree of venous reflux would necessitate a method of 19 20 quantifying reflux in the superficial venous system in a way that adequately reflects disease severity, 21 and such a method does not currently exist. Hence the CEAP categorisation question may be the best 22 choice with a refined quality of life measure.

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Criteria for selecting high-priority research recommendations

Criterion	Explanation
Importance to patients or the population	If this trial shows that differing levels of disease severity have different optimal treatments, this will lead to more appropriate and cost-effective treatments for all patients.
Relevance to NICE guidance	This research is highly relevant to NICE guidance. The answer to this question may change the guidance and the way that varicose veins are treated.
Relevance to the NHS	The impact of this research on the NHS is likely to decrease burden on the NHS as it will mean that the most cost-effective option is used for each disease stage.
National priorities	This research is not relevant to any known national priority areas.
Current evidence base	There have been no studies to date evaluating the optimal treatment strategies for different levels of disease severity.

Research recommendations

Researen recomme	nations
Relevance to NICE guidance	Results would influence recommendations regarding best management of severe venous dysfunction and with the ESCHAR study would influence the planning of venous services.
Relevance to the NHS	Leg ulceration due to superficial venous reflux is a major cause of morbidity and a huge drain on NHS resources. Improved management techniques may influence service delivery and ultimately strategic planning.
National priorities	None
Current evidence base	The RCT ESCHAR study compared surgery and compression with compression alone but it might now be suitable to consider endovenous interventional techniques which being a minimally invasive procedure is more acceptable to patients and may be delivered without delay.
Other comments	This study has recently been submitted in the form of The Early Venous Reflux Ablation (EVRA) ulcer trial to the National Institute for Health Research. In the ESCHAR study, no improvement in ulcer healing rates was seen, but operative intervention was delayed for a median time of 7 weeks. Recruitment period: 24 months. Study duration: 48 months.
Equality	The research does not address equality issues as all people will be able to access the intervention.
Feasibility	The research is expected to be able to be carried out within a realistic timescale and acceptable cost. It is not expected that there would be any ethical or technical issues.
Other comments	None
Study design	Randomised controlled trial
Criterion	Explanation
Importance to patients or the population	This study is important to those patients with CEAP stage C6 disease whose quality of life is substantially reduced by discomfort and social isolation resulting from odour and wound discharge. The social and personal impact of chronic venous leg ulceration is therefore considerable.
Equality	None identified.
Feasibility	The proposed research could be carried out within a realistic timescale and at an acceptable cost. No ethical or technical issues?
Study design	Multi-centred randomised controlled trial.

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2 N.1.3 Truncal treatment with or without concurrent tributary treatment

3 Research Question

What is the clinical and cost effectiveness of concurrent phlebectomies or foam sclerotherapy for varicose tributaries during truncal endothermal ablation for varicose veins compared with:

- truncal endothermal ablation without concurrent phlebectomies or foam sclerotherapy.
- truncal endothermal ablation with phlebectomies or foam sclerotherappy, if needed, 6–12 weeks later.

9 Why this is important

Conventional truncal stripping under general anaesthetic involves synchronous phlebectomies of
 varicose tributaries, and in ultrasound-guided foam sclerotherapy truncal and tributary veins are
 treated concurrently. In contrast, endothermal ablation may be performed alone to obliterate
 truncal incompetence, or synchronously with phlebectomies or foam sclerotherapy, and current
 practice varies.

1Synchronous tributary treatment ensures a single treatment episode, and the removal of all2symptomatic varicosities leads to a better immediate quality of life, but this takes longer and thus3may be associated with increased morbidity. Deferred tributary treatment may reduce morbidity,4and also mean that some patients do not need tributary treatment (or need fewer tributary5treatments on smaller veins). However, it involves 2 interventions for patients who need tributary6treatment. Omitting tributary treatments entirely ensures a single treatment episode, but it is7unclear whether remaining varicosities will persist and impair quality of life.

8 At present there is limited evidence from 1 small-scale (n=50) study on the use and timing of 9 tributary treatments after truncal endothermal ablation. There is a need for practice to be based on 10 empirical evidence from a large and sufficiently powered RCT comparing all 3 main intervention 11 options (no tributary treatment, concurrent tributary treatment and delayed tributary treatment).

Criterion	Explanation
Importance to patients or the population	If the trial showed a benefit of tributary treatment during or after treatment this would lead to altered guidance recommending tributary treatment during or after interventional treatment. This would improve patients' quality of life.
Relevance to NICE guidance	This research is of medium importance to the NICE guideline, as the research is relevant to the recommendations in the guideline but the research recommendations are not key to future updates.
Relevance to the NHS	The impact of this research on the NHS is minimal. If the research identified a benefit with tributary treatment during or after treatment a change in recommendation would be required which would lead to a change in practice in the NHS and could reduce costs.
National priorities	This research is not relevant to any known national priority areas.
Current evidence base	The evidence for this section was reviewed in chapter 9e. Only 1 randomised controlled trial was identified in this area. This was not prone to serious bias (unblinded, but clear allocation concealment and no attrition bias) but was possibly underpowered with high levels of imprecision for some outcomes, leading to these outcomes being graded as very low. In addition, this RCT had a short follow-up of only 6 weeks for concurrent tributary treatments vs. no tributary treatment.
Equality	The research does not address equality issues as all people will be able to access the intervention.
Feasibility	The research is expected to be able to be carried out within a realistic timescale and acceptable cost. It is not expected that there would be any ethical or technical issues.
Other comments	None
Study design	Randomised controlled trial

Criteria for selecting high-priority research recommendations

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1 N.1.4 Compression as a management option

2 What is the clinical and cost effectiveness of compression hosiery versus no compression for the 3 management of symptomatic varicose veins?

4 Why this is important

- 5 Compression hosiery is widely used as first-line treatment for symptomatic varicose veins. In some 6 areas of the UK a period of hosiery use is a precursor to referral to secondary care.
- Discomfort and difficulty in application may cause people to stop wearing compression hosiery or
 wear it only occasionally. The current evidence for the benefit of compression hosiery is weak. There
 is little evidence of an impact on symptom relief or an improvement in quality of life. It is therefore
 not possible to calculate the cost effectiveness of compression hosiery.
- A multicentre trial randomising compression hosiery versus no compression in patients with
 symptomatic varicose veins is needed. The trial should evaluate quality of life, including symptom
 reduction, and measure adherence with compression hosiery. In addition the trial should investigate
 the impact of compression on disease progression and the need for subsequent intervention.

15 Criteria for selecting high-priority research recommendations

Criterion	Explanation
Importance to patients or the population	The research is important to patients and, if results showed a benefit for compression, clinicians would be confident they were being offered a clinically proven treatment option.
Relevance to NICE guidance	This research is highly relevant to NICE guidance. The answer to this question may change the guidance and the way that varicose veins are treated. The poor quality current evidence and relatively high costs mean that compression has not been recommended, but if the research were to identify that symptoms of varicose veins were substantially reduced the recommendation may change. Compression might then provide an effective non-interventional management strategy.
Relevance to the NHS	What would be the impact on the NHS and (where relevant) the public sector of any new or altered guidance (for example, financial advantage, effect on staff, impact on strategic planning or service delivery)?
	The research could potentially reduce the variation in practice within the NHS. If the results showed that compression hosiery was effective for the management of varicose veins this would provide a non-interventional management strategy. Conversely, if the research was clear that compression stockings showed little benefit, the NHS could then stop prescribing an ineffective treatment benefit. In addition, if there was evidence concerning who benefited from compression hosiery,
	treatment could be better targeted.
x	This research is not likely to have an impact on national priorities.
Current evidence base	The systematic review of the evidence identified 3 low or very low quality RCTs, two of which were completed more than 15 years ago (see section 8.1). These investigated patient assessed symptoms and adverse events but did not look at patients' quality of life. In addition, 5 observational studies were identified which provided some further information. The GDG felt strongly that the nature of the evidence created much uncertainty about the results for all relevant outcomes.
Equality	Equality issues are not particularly relevant to this research question, although the ability to put on and take off compression hosiery and whether that had an impact in the adherence to the treatment strategy should be considered as part of the research
Feasibility	The proposed research is considered by the GDG to be feasible and able to be carried out

Criterion	Explanation
	within a realistic timescale and at an acceptable cost. There are no known ethical or technical issues.
Other comments	None
Study design	Randomised Controlled Trial

2 N.1.5 Compression after interventional treatment

What is the clinical and cost effectiveness of compression bandaging or hosiery after interventional
treatment for varicose veins compared with no compression? If there is benefit, how long should
compression bandaging or hosiery be worn for?

6 Why this is important

7 The benefit of compression after interventional treatment for varicose veins is unclear. A well-8 conducted multicentre randomised controlled trial (RCT) of compression after interventional 9 treatments would help determine whether compression is beneficial, and if so, what type is best and 10 how long it should be worn for. The trial should include patients who have 1 of the 3 main interventional treatments: endothermal ablation, ultrasound-guided foam sclerotherapy and surgery. 11 12 The patients should be divided into 3 groups based on the type of intervention they have had. There should be 6 RCT arms, 1 arm with compression and 1 arm without in each of 3 patient groups. Each 13 14 arm should have subgroups for compression type and duration. Adherence to compression 15 treatment and the impact of adherence on effectiveness should also be evaluated. A cost-16 effectiveness analysis should be performed. If compression is beneficial, such a trial should help 17 improve quality of life for people with varicose veins and reduce the longer-term need for 18 retreatment.

19 Criteria for selecting high-priority research recommendations

Criterion	Explanation
Importance to patients or the population	If the trial showed a benefit of compression hosiery after treatment this would lead to altered guidance recommending compression after interventional treatment. This would improve patients' quality of life.
Relevance to NICE guidance	This research is of medium importance to the NICE guideline, as the research is relevant to the recommendations in the guideline but the research recommendations are not key to future updates.
Relevance to the NHS	The impact of this research on the NHS is minimal. If the research identified a benefit with compression after interventional treatment a change in recommendation would be required which would lead to a change in practice in the NHS.
National priorities	This research is not relevant to any known national priority areas.
Current evidence base	The evidence for this section was reviewed in chapter 10. Only 2 low / very low quality randomised controlled trials were identified in this area which were both prone to serious bias. In most cases the imprecision of the point estimate was too large to be able to confidently judge the magnitude/direction of the true population effect.

Criterion	Explanation
Equality	The research does not address equality issues as all people will be able to access the intervention. Patient compliance with compression should be assessed in the research to determine if there are any factors which meant that compression was not suitable for specific groups.
Feasibility	The research is expected to be able to be carried out within a realistic timescale and acceptable cost. It is not expected that there would be any ethical or technical issues.
Other comments	None
Study design	Randomised controlled trial

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Appendix O: References

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