Towards an individualized management strategy for patients with chronic venous disease: Results of a Delphi consensus

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Abstract
Objective: To obtain consensus on management criteria for symptomatic patients with chronic venous disease (CVD; C2–C6) and superficial venous reflux.
Method: We used a Delphi method by means of 36 statements sent by email to experts in the field of phlebology across the world over the course of three rounds. The statements addressed criteria for different venous treatments in patients with different characteristics (e.g. extensive comorbidities, morbid obesity and peripheral arterial disease). If at least 70% of the ratings for a specific statement were between 6 and 9 (agreement) or between 1 and 3 (disagreement), experts’ consensus was reached.
Results: Twenty-five experts were invited to participate, of whom 24 accepted and completed all three rounds. Consensus was reached in 25/32 statements (78%). However, several statements addressing UGFS, single phlebectomies, patients with extensive comorbidities and morbid obesity remained equivocal.
Conclusion: Considerable consensus was reached within a group of experts but also some gaps in available research were highlighted.

Keywords
Varicose veins, endovascular treatment, echo-sclerotherapy

Introduction
Over the past decades, numerous effective treatments have been developed for patients with chronic venous disease (CVD). The increased availability of different treatment options has made the management of CVD more challenging. It has resulted in a large worldwide variation in management preferences for treating patients with CVD.¹² Management strategies are not only based on evidence from the literature but also on the physician’s own experience, availability (and costs) of the equipment and on national healthcare reimbursement systems. However, in clinical practice, management decisions should ideally be influenced mainly by a combination of patient characteristics, clinical findings and results of duplex ultrasound (DUS).¹³,⁴

Although there are several well-established national and international guidelines,⁵⁻⁸ their recommendations only focus on the treatment of a diseased population rather than an individual patient.⁹ Appropriate tools for management strategies incorporating the specific characteristics of an individual patient, as well as the clinical and DUS findings, are lacking in current clinical guidelines.¹⁰⁻¹² To bridge the gap between current clinical guidelines and individualized care, expert opinion methods such as a Delphi consensus¹⁰ are desired.

The aim of the present study was to achieve an international Delphi-based consensus on management criteria for patients presenting with venous symptoms, clinical signs of CVD, and superficial venous reflux confirmed by DUS.

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Methods

Expert panel

Experts were selected from a group of physicians who already participated in our recently published worldwide survey on management strategies in patients with CVD (C2–C6). Only those with at least one scientific publication in the field of phlebology in a peer-reviewed journal and at least 10 years of experience in treating patients with CVD were eligible for the present study. They had to be performing ultrasound-guided foam sclerotherapy (UGFS), endovenous thermal ablation (EVTA) and phlebectomies themselves. Two investigators (SKvdV and MGRDM) selected the expert panel. For this selection, they aimed at having a reasonable distribution between different specialties (vascular or general surgeons, dermatologists, angiologists, phlebologists) and continent or country of clinical practice. Based on the selection criteria, 25 eligible experts were invited by email. They were asked to judge on several statements in view of obtaining consensus over the course of three rounds, according to the Delphi method between March 2015 and November 2015 (Figure 1). In case of non-responding, the investigators sent two email reminders in each round.

Informed consent was not performed because no patients were involved in this study.

Delphi consensus procedure

The authors formulated the statements, using results from a worldwide survey regarding management strategies in patients with great saphenous vein (GSV) and tributary reflux. The first round contained 30 statements addressing criteria for endovenous ablation (EVA) of the refluxing saphenous trunk (ST), including both thermal and non-thermal ablation techniques (mainly for simplicity), UGFS, high ligation and stripping (HL/S), single phlebectomies without treatment of the refluxing saphenous trunk referred to as ‘ambulatory selective varicose ablation under local anaesthesia’ (ASVAL) and non-interventional measures (venotonics drugs and medical elastic compression stockings (MECS)). The statements contained scenarios with different patient characteristics, clinical class (C2–C6, according to the CEAP classification) and DUS

![Figure 1. Selection of participants for the Delphi consensus.](image-url)

*Eligibility criteria: performing varicose vein treatments for at least five years and familiar with the use of several currently used varicose veins techniques including, phlebectomies, one of the techniques for endovenous thermal ablation, and ultrasound-guided foam sclerotherapy. They were also allowed to participate if they did not perform EVTA themselves, but delegated this to a colleague when indicated. The same was true for high ligation and stripping.

**Eligibility criteria: at least one scientific publication in a phlebologic peer-reviewed journal, at least ten years of experience in treating phlebologic patients and performing ultrasound-guided foam sclerotherapy, endovenous thermal ablation and phlebectomies themselves.
findings in patients with CVD of the lower limbs and superficial venous reflux in a ST. The terminology used for the statements is summarized in Table 1.

Participants were asked to anonymously rate each statement on a 9-point scale, by ticking boxes with marks between 1 and 9, where a score of 1 denoted complete disagreement and a score of 9 indicating full agreement (Figure 2). If at least 70% of the ratings for a specific statement were between 7 and 9, it was concluded experts consented on agreeing with the statement. Vice versa, if ≥70% were between 1 and 3, it was concluded they consented on disagreeing with the statement. This cut-off level was based on previous literature. Any other distribution of ratings was valued ‘equivocal’. Participants also had the opportunity to add a comment in the ‘remark(s)’ box accompanying each statement. All this information was used for preparing the subsequent round of the Delphi consensus.

In the second round, experts received a full report of the first round, including a compilation of ratings for all statements of the first round, and all the remarks of their colleagues, provided anonymously. Then, they were asked to rate again those statements that had remained equivocal in the first round. Statements were reformulated and new statements were added based on the remarks of the experts in the previous round. For instance, in the first round, we had included older age as a patient characteristic, which might influence management. However, participating experts suggested age was not the main factor, but rather extensive comorbidities, and therefore the statement was adapted as such. Eventually, the second round contained 18 statements. In the second round, we also attached a summary of the present literature regarding each statement.

The same procedure was followed for the third (and last) round, which contained only nine statements to be rated. Again, participating experts could consider the results of the second round and the remarks of their colleagues.

In case of one or more missing items, the expert was contacted by email to retrieve the missing answer.

### Results

#### Experts

In total, 24 out of 25 contacted experts accepted to participate: 15 vascular surgeons, two general surgeons, four dermatologists, two angiologists and one phlebologist from Europe (17/25), Australia (1/25),

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**Table 1.** Definition of terminology used for the statements of the Delphi consensus.

| Venous symptoms: | ache, pain, heaviness, tightness, feeling of swelling, nocturnal cramps, itching |
|------------------|--------------------------------------------------------------------------------|
| C disease:       | clinical class according to the CEAP classification: C2: varicose veins; C3: edema; C4: skin changes (pigmentation, eczema, atrophie blanche, lipodermatosclerosis); C5: healed venous ulcer; C6: open venous ulcer |
| Saphenous trunk: | great saphenous vein, anterior accessory saphenous vein, posterior accessory saphenous vein, Giacomini vein, small saphenous vein |
| Varicose tributaries: | visible or palpable varicose veins in the subcutis |
| Reflux in a saphenous trunk: | abnormally reversed flow in a saphenous trunk, during > 500 ms at calf compression-release or Valsalva (the latter only for the SFJ) involving the terminal or preterminal valve of the SFJ or the SPJ |
| Terminal valve reflux: | reflux at the junction of the SFJ or SPJ |
| Segmental reflux: | reflux limited to a segment of a saphenous trunk, not involving the SFJ or SPJ |
| Saphenous diameter: | diameter measured in a tubular part of the refluxing vein segment, about 15 cm from the junction in standing position |
| Focal dilatation: | localized dilatation of the saphenous trunk less than 1.5–3 times the saphenous diameter above or below |
| Aneurysm: | dilatation of the saphenous vein more than three times the saphenous diameter above or below, or more than 20 mm (close to the SFJ or SPJ) |
| Morbid obesity: | BMI > 40 kg/m² or BMI > 35 kg/m² and experiencing obesity-related health conditions |
| Severe peripheral arterial disease: | ankle brachial index < 0.6 |
| Bridging: | oral anticoagulation is interrupted with bridging anticoagulation, using either heparin or low-molecular weight heparin, administered during the sub-therapeutic window |

SFJ: saphenofemoral junction; SPJ: saphenopopliteal junction; BMI: body mass index.
Figure 2. Example of a statement showing consensus on agreement (84% of marks between 7 and 9). HL/S: high ligation and stripping.

North-America (4/25) and Central- or South-America (3/25). Experts were working in private practice (13/25), general hospital (4/25) or university hospital (7/25). All these experts completed the first, second and third Delphi round (25/25). The majority of experts (16/25) had over 20 years of experience in treating patients with CVD.

Delphi consensus

After three rounds, consensus was reached in 25 out of 32 statements (78%; Table 2). All statements regarding indications for treatment with thermal or non-thermal EVA vs. treatment with HL/S reached high (>79%) consensus in favor of EVA, except for venous aneurysms. Of note, 96% of the experts agreed on that high ligation should not be added to EVA treatment. Similar level of agreement was reported about the use of MECS categorizing it 'a valuable treatment option' for ST's with small diameters <4 mm. This discrepancy illustrates that ST diameter may be used as a relevant criterion to distinguish between different treatment options,1 which is in line with the European guidelines for sclerotherapy.17

UGFS is usually considered as the second best option of the minimally invasive treatments for abolishing ST reflux.5–8 In the present consensus approach, the experts considered UGFS to be just ‘a’ treatment option if ST diameter is ≥4 mm, and only agreed on categorizing it ‘a valuable treatment option’ for ST's with small diameters <4 mm. This discrepancy illustrates that ST diameter may be used as a relevant criterion to distinguish between different treatment options,1 which is in line with the European guidelines for sclerotherapy.17

ASVAL is not yet a worldwide-accepted strategy, and this was reflected in the Delphi consensus. Contrarily to what the investigators had been suggesting, the experts could not agree on particular criteria for implementing ASVAL so far. This is somehow to be expected as only 42% of the experts had ever used ASVAL in their clinical practice (data not shown) and the evidence on the subject is still limited.11,18,19 Future research should point out whether ASVAL can be applied in all patients with varicose veins or in a selected varicose vein population only.

At the end of the first round of the Delphi consensus, older age appeared to be less relevant than we initially had hypothesized and it was therefore converted into extensive comorbidities as suggested by the participants from the second round onwards. Indeed, clinical studies demonstrated that the feasibility and safety of EVTA and tolerability of UGFS were similar in patients older or younger than 75 years.20,21 In contrast, studies regarding venous treatments in patients with (extensive) comorbidities are completely lacking. Although the field of treatment of superficial venous disease, by means of a Delphi consensus procedure. This may help physicians to make supported and individualized choices for their patients with CVD (C2–C6).

Not surprisingly, experts’ recommendations regarding EVA were in line with current guidelines on the care of patients with varicose veins and associated CVD.5–8 EVA was considered the preferred treatment for most symptomatic patients with CVD (C2–C6) and a refluxing ST >4 mm in diameter. The expert panel did not recommend an alternative treatment for specific DUS findings such as terminal valve reflux, a large (>10 mm) ST diameter or the presence of one or more focal dilatations of the ST (Table 2). EVA has now globally become the preferred treatment for ST reflux as is reflected in our results. Although EVA (thermal or non-thermal) has largely replaced HL/S in many countries, it should be acknowledged that, whenever equipment for EVA is not available, HL/S (ideally using tumescent anesthesia, in agreement with recent guidelines8) remains a valuable treatment option. While patient-reported outcomes favor EVTA, long-term efficacy of HL/S and EVTA are very similar.15,16

The experts' opinion on several statements remained equivocal after three rounds. This was the case for statements addressing different indications for UGFS, ASVAL, extensive comorbidities and morbid obesity. The experts stated that it depended on the type and extent of the patient’s comorbidities and therefore no definitive conclusion on preferred management could be made. Regarding morbid obesity, the experts would definitely prefer EVA over HL/S, but there was no agreement on limiting treatment to those patients with skin changes (C4–C6). Moreover, based on their experience, the participants were convinced that advising weight reduction unfortunately was unsuccessful in the majority of obese patients.

Throughout the different rounds of the Delphi consensus, three statements were removed, two because of redundancy after reformulating other statements and one because of lack of relevance following the experts' remarks (Supplementary Table 1).

Discussion

The present study showed that considerable consensus could be reached within a group of selected experts in the field of treatment of superficial venous disease, by means of a Delphi consensus procedure. This may help physicians to make supported and individualized choices for their patients with CVD (C2–C6).

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Table 2. Results of Delphi consensus.

| Statements | Delphi rounds |
|------------|--------------|
| EVA vs. HL/S | 1 | 2 | 3 |
| Nowadays, with the availability of endovenous treatments, HL/S is only rarely indicated. | 84% | ✔ | ✔ |
| In case of saphenous reflux, EVA is indicated rather than HL/S, even if there is C4–C6 disease. | 83% | ✔ | ✔ |
| In patients with venous symptoms and reflux in a saphenous trunk, EVA is indicated rather than HL/S, even in the presence of TV reflux. | 83% | ✔ | ✔ |
| In patients with venous symptoms and reflux in a saphenous trunk, EVA is indicated rather than HL/S, even in the presence of a large (>10 mm) saphenous diameter. | 79% | ✔ | ✔ |
| In patients with venous symptoms and reflux in a saphenous trunk, EVA is indicated rather than HL/S, even in the presence of one or more focal dilatations. | 79% | ✔ | ✔ |
| Presence of a venous aneurysm (>20 mm) within 2 cm from the SFJ or SPJ is an indication for HL/S rather than EVA. | 83% | ✔ | ✔ |
| HL should not be added to patients being treated with EVA. | 96% | ✔ | ✔ |
| UGFS | 1 | 2 | 3 |
| In the presence of C4–C6 disease in patients with reflux in a saphenous trunk >4 mm in diameter, UGFS is a treatment option. | 93% | ✔ | ✔ |
| In the presence of C4–C6 disease in patients with reflux in a saphenous trunk <4 mm in diameter, UGFS is a valuable treatment option. | 80% | ✔ | ✔ |
| In the presence of C4–C6 disease in patients with reflux in a saphenous trunk <4 mm in diameter and refluxing tributaries in a diseased skin area, UGFS of tributaries is preferred rather than phlebectomies. | 87% | ✔ | ✔ |
| In the presence of C4–C6 disease in patients with reflux in a saphenous trunk <4 mm in diameter and refluxing tributaries, UGFS and phlebectomies of tributaries at a distance from the diseased skin area, are both valuable treatment options. | 92% | ✔ | ✔ |
| In patients with venous symptoms and reflux in a saphenous trunk, UGFS (without tumescent anesthesia) is a valuable treatment option, even in the presence of a large (>10 mm) saphenous diameter. | 75% | ✔ | ✔ |
| In patients with venous symptoms and reflux in a saphenous trunk vein >10 mm in diameter, where ablation is indicated, EVA is preferred rather than UGFS. | 92% | ✔ | ✔ |
| If you have decided to ablate the saphenous trunk, in the presence of C2–C3 disease in patients with venous symptoms and reflux in a saphenous trunk vein <4 mm in diameter, UGFS is preferred rather than EVA. | 87% | ✔ | ✔ |
| EVA | 1 | 2 | 3 |
| In the presence of C4–C6 disease in patients with reflux in a saphenous trunk <4 mm in diameter, EVA is a valuable treatment option. | 71% | ✔ | ✔ |
| ASVAL | 1 | 2 | 3 |
| In the presence of C2–C3 disease in patients with venous symptoms, segmental reflux of a saphenous trunk <4 mm in diameter and large refluxing tributaries, preservation of the saphenous trunk is indicated rather than its ablation. | 83% | ✔ | ✔ |
| ASVAL is not indicated in case of reflux in a saphenous trunk and C4–C6 disease. | 83% | ✔ | ✔ |
| Non-interventional measure | 1 | 2 | 3 |
| In patients with venous symptoms, C2–C3 disease and reflux in a saphenous trunk and who are not willing to undergo any intervention or who are unfit for intervention because of extensive comorbidities, MECS should be considered. | 88% | ✔ | ✔ |
| In patients with venous symptoms, C2–C3 disease and reflux in a saphenous trunk and who are not willing to undergo any intervention or who are unfit for intervention because of extensive comorbidities, venotonic drugs should be considered. | 79% | ✔ | ✔ |
| In patients with C4–C6 disease and reflux in a saphenous trunk and who are not willing to undergo any intervention or who are unfit for intervention because of extensive comorbidities, MECS are indicated. | 96% | ✔ | ✔ |

(continued)
investigators thought that minimizing the extent of procedures in patients with comorbidities and/or morbid obesity would be good clinical practice, experts did not reach consensus on this issue. This lack of consensus could partly be explained by the fact that many different combinations of comorbidities can be thought of in these cases and the extent of comorbidities was not defined in the statements. The experts seemed to be more unanimous regarding strategies in patients under chronic anticoagulant treatment and those with severe peripheral arterial disease.

Although we found considerable consensus about treatment strategies in different clinical classes and DUS findings, statements addressing patients’ characteristics remained largely equivocal in the Delphi consensus. This clearly highlights the lack of knowledge in the field of personalized medicine and implies that more research regarding patient characteristics, and how these are influencing treatment outcomes, is needed. In this way, future guidelines may include more tools for proper stratification of patients with CVD.

Table 2. Continued

| Statements                                                                 | Delphi rounds | 1   | 2   | 3   |
|----------------------------------------------------------------------------|---------------|-----|-----|-----|
| Comorbidities                                                              |               |     |     |     |
| In patients with venous symptoms in addition to extensive comorbidities and reflux in a saphenous trunk, UGFS is indicated rather than EVA.\(^a\) |               |     |     |     |
| In patients with venous symptoms in addition to extensive comorbidities and reflux in a saphenous trunk, EVA or UGFS should only be considered in case of C4–C6 disease.\(^a\) |               |     |     |     |
| Morbid obesity                                                             |               |     |     |     |
| In patients with venous symptoms, reflux in a saphenous trunk and morbid obesity, EVA or UGFS are indicated rather than HL/S. |               |     |     | 87% |
| In patients with venous symptoms, reflux in a saphenous trunk and morbid obesity, treatment should only be considered in case of C4–C6 disease.\(^b\) |               |     | 87% |     |
| In patients with venous symptoms, morbid obesity and reflux in a saphenous trunk, weight reduction is advised prior to any venous treatment.\(^c\) |               |     |     |     |
| Anticoagulation                                                            |               |     | 92% |     |
| In patients with venous symptoms, reflux in a saphenous trunk and who are on chronic anticoagulants, EVA is indicated rather than HL/S. |               |     |     |     |
| In patients with venous symptoms, reflux in a saphenous trunk < 4 mm diameter and who are on chronic anticoagulants, UGFS is indicated rather than HL/S.\(^a\) |               |     | 91% |     |
| In patients with venous symptoms, reflux in a saphenous trunk and who are on chronic anticoagulants, EVA can be performed without bridging. |               |     | 87% |     |
| In patients with venous symptoms, reflux in a saphenous trunk and who are on chronic anticoagulants, UGFS can be performed without bridging. |               |     | 88% |     |
| In patients with venous symptoms, who are on chronic anticoagulants and who are scheduled for extensive phlebectomies, temporary discontinuation and anticoagulant treatment bridging is indicated.\(^b\) |               |     |     | 88% |
| Severe peripheral arterial disease                                           |               |     |     |     |
| In patients with C2–C3 disease, reflux in a saphenous trunk and severe peripheral arterial disease, it is preferable not to ablate the saphenous vein by means of EVA, UGFS or HL/S. |               |     | 79% |     |
| In patients with C4–C6 disease, reflux in a saphenous trunk and severe peripheral arterial disease, EVA or UGFS of the refluxing saphenous vein may be considered. |               |     | 71% |     |

EVA: endovenous ablation (including thermal and non-thermal non-tumescent techniques, excluding UGFS); HL/S: high ligation with stripping; SFJ: saphenofemoral junction; SPJ: saphenopopliteal junction; UGFS: ultrasound-guided foam sclerotherapy; ASVAL: ‘ambulatory selective varicose ablation under local anaesthesia’ (= single phlebectomies without treating the saphenous trunk); MECS: medical elastic compression stockings.

\(^a\)Statement was reformulated after first round.
\(^b\)Statement was reformulated after second round.
\(^c\)Statement was added after remarks of experts in the first round.
strategies in patients with CVD, physicians should also consider patient’s preference, as well as the impact of the disease on Health-Related Quality of Life (HRQoL), the estimated risk of deterioration of CVD and local healthcare resources.

**Limitations**

This study has several limitations. The statements may have been incomplete, but this was minimized because they were based on a prior worldwide survey among 211 specialists in CVD. Also, this study focused on the treatment of patients with venous symptoms, clinical signs of CVD, and superficial venous reflux in the ST and tributaries, and therefore our results are not generalizable to all patients with CVD (e.g. patients with a pathological perforating vein nearby a venous ulcer or patients with groin recurrence). Selection bias among the eligible experts might have occurred because the majority of the participants were working in Western Europe. However, the response rate was almost perfect. In the absence of clear guidelines for a Delphi consensus, we chose the cut-off value of 70% as acceptable, because of the relatively mild impact of CVD on HRQoL, contrarily to decision-making in more life-threatening diseases in which a higher cut-off may be desired. In addition, there is no standard method for calculating sample sizes for a Delphi study, but there is evidence that expert panels of 20 can reach a valid consensus.

**Conclusion**

In conclusion, in the present investigation about management strategies in patients with symptoms and signs of CVD and superficial venous reflux, the experts agreed on most of the strategies and their opinions largely reflected international guidelines. Nevertheless, it appeared to be more difficult to reach consensus on the influence of certain patient-related characteristics. More research is needed in the field of personalized medicine, in order to further optimize phlebologic care for all our patients presenting with CVD.

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**Contributorship**

SKvdV designed study, Delphi consultation, analysed data, drafted paper. RRvdB designed study, drafted paper. OP designed study, critically reviewed paper. TN designed study, critically reviewed paper. MGRdM designed study, Delphi consultation, analysed data, drafted paper.

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