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A Randomized Trial Comparing Treatments for Varicose Veins


BACKGROUND
Ultrasound-guided foam sclerotherapy and endovenous laser ablation are widely used alternatives to surgery for the treatment of varicose veins, but their comparative effectiveness and safety remain uncertain.

METHODS
In a randomized trial involving 798 participants with primary varicose veins at 11 centers in the United Kingdom, we compared the outcomes of foam, laser, and surgical treatments. Primary outcomes at 6 months were disease-specific quality of life and generic quality of life, as measured on several scales. Secondary outcomes included complications and measures of clinical success.

RESULTS
After adjustment for baseline scores and other covariates, the mean disease-specific quality of life was slightly worse after treatment with foam than after surgery (P=0.006) but was similar in the laser and surgery groups. There were no significant differences between the surgery group and the foam or the laser group in measures of generic quality of life. The frequency of procedural complications was similar in the foam group (6%) and the surgery group (7%) but was lower in the laser group (1%) than in the surgery group (P<0.001); the frequency of serious adverse events (approximately 3%) was similar among the groups. Measures of clinical success were similar among the groups, but successful ablation of the main trunks of the saphenous vein was less common in the foam group than in the surgery group (P<0.001).

CONCLUSIONS
Quality-of-life measures were generally similar among the study groups, with the exception of a slightly worse disease-specific quality of life in the foam group than in the surgery group. All treatments had similar clinical efficacy, but complications were less frequent after laser treatment and ablation rates were lower after foam treatment. (Funded by the Health Technology Assessment Programme of the National Institute for Health Research; Current Controlled Trials number, ISRCTN51995477.)
Ultrasonic-guided foam sclerotherapy and thermal ablation techniques such as endovenous laser ablation have become widely used alternatives to surgery for the treatment of varicose veins. Previous randomized trials and meta-analyses have shown these treatments to be effective in terms of short-term technical success and clinician-reported outcomes. Clinical practice guidelines recommend the use of patient-reported quality of life to assess the outcomes of treatment of varicose veins. Quality of life was a primary outcome measure in two small randomized trials that compared surgery and endovenous laser ablation, but to our knowledge, it has not been assessed as a primary outcome in randomized trials involving foam sclerotherapy.

We performed the Comparison of Laser, Surgery, and Foam Sclerotherapy (CLASS) trial, a large, multicenter, randomized, comparative-effectiveness trial, to assess quality of life and other outcomes of treatment of varicose veins. We compared foam sclerotherapy, laser therapy (with subsequent foam sclerotherapy for residual varicosities, if required), and surgery.

** METHODS **

**PATIENTS**

We recruited patients requiring treatment of varicose veins in 11 vascular surgery departments in the United Kingdom between November 2008 and October 2012. All patients were assessed by a vascular surgeon and underwent initial duplex ultrasonographic scanning to assess suitability for treatment and entry into the study. Inclusion criteria were an age of 18 years or older, the presence of unilateral or bilateral primary symptomatic varicose veins (grade C2 or higher according to the clinical, etiologic, anatomical, and pathophysiological [CEAP] classification system, with C0 indicating no signs of venous disease, C1 telangiectases or veins ≤3 mm in diameter, C2 varicose veins >3 mm in diameter, C3 the presence of edema, C4 skin and subcutaneous changes, C5 healed ulcers, and C6 active ulceration), and reflux of the great or small saphenous veins of more than 1 second on duplex ultrasonography. Exclusion criteria were current deep-vein thrombosis, acute superficial-vein thrombosis, a diameter of the main truncal saphenous vein of less than 3 mm or more than 15 mm, tortuous veins considered to be unsuitable for laser treatment, and contraindications to the use of foam or to general or regional anesthesia.

**RANDOMIZATION AND STUDY TREATMENT**

A computer-generated randomization system was used and was managed by the Centre for Healthcare Randomised Trials, University of Aberdeen, Aberdeen, United Kingdom. Participants underwent randomization with even assignments to all treatment options available at each investigating center and with stratification according to the number of available options (stratum A, eight hospitals offering all three treatment options; and stratum B, three hospitals offering treatment with only foam or surgery). Treatments were assigned with the use of a minimization algorithm that included center, age (<50 years or ≥50 years), sex, reflux of either the great or the small saphenous veins (or both), and the presence or absence of unilateral or bilateral varicose veins.

Details of treatment methods are described in the published protocol (available with the full text of this article at NEJM.org). Briefly, surgery consisted of proximal ligation and stripping (of the great saphenous vein only) and concurrent phlebectomies. Foam was produced with the use of the Tessari technique at a ratio of 0.5 ml of sodium tetradecyl sulfate to 1.5 ml of air (3% sodium tetradecyl sulfate for saphenous veins and 1% for varicosities, with a maximum of 12 ml of foam per session). The use of sodium tetradecyl sulfate is licensed, but the trial involved its off-license use as a foam rather than as its manufactured liquid form. Laser ablation of truncal saphenous veins performed while the patient was under local anesthesia was followed by foam sclerotherapy to residual varicosities after 6 weeks follow-up if required, with the exception that one center performed concurrent phlebectomies.

**OUTCOME MEASURES**

Outcomes were assessed at baseline and at 6 weeks and 6 months after treatment. The primary outcome measures were patient-reported disease-specific quality of life, measured with the use of the Aberdeen Varicose Veins Questionnaire (AVVQ), and patient-reported generic (i.e., general) quality of life, measured at 6 months after treatment with the use of the EuroQol Group 5-Dimension Self-Report Questionnaire (EQ-5D) and the Medical Outcomes Study 36-Item Short-Form Health Survey (SF-36). Another prespecified primary outcome for this trial — 5-year estimated cost-effectiveness, measured as cost per quality-adjusted life-year gained — is not reported here.

The AVVQ is an internationally validated...
change-responsive tool for the assessment of quality of life in patients with varicose veins.\textsuperscript{20,21-26} It consists of 12 questions and a set of mannequin legs on which participants are asked to draw their veins. Scores range from 0 to 100, with higher scores indicating a worse quality of life. The EQ-5D is a standardized index valuation for health status; it includes five dimensions (mobility, self-care, usual activities, pain or discomfort, and anxiety or depression; scores range from −0.594 to 1.000, with higher scores indicating a better quality of life) and a visual analogue scale (EQ-5D VAS; scores range from 0 to 100, with higher scores indicating better health).\textsuperscript{27} The SF-36 is a validated and reliable assessment of quality of life and is widely used for a variety of clinical conditions.\textsuperscript{28} The 36 questions assess eight domains and yield two summary scores (the physical component and the mental component), with each summary score ranging from 0 to 100 and higher scores indicating greater well-being. For all these measures of quality of life, minimal clinically important differences after treatment for varicose veins are not known.

Secondary outcomes were as follows: clinical success at 6 weeks and 6 months, as measured by the proportion of patients with residual varicose veins (assessed by the participant and the nurse), venous clinical severity score (a score composed of nine categories relating to symptoms or signs of venous disease and one category relating to the use of compression; scores range from 0 [no venous disease] to 30 [most severe venous disease]), and complications (assessed by the surgeon or nurse); quality of life according to the AVVQ, EQ-5D, and SF-36 at 6 weeks; the EQ-5D VAS and the eight SF-36 domains at 6 weeks and 6 months; and ablation rates of the main trunks of the saphenous vein according to duplex ultrasonography at 6 weeks and 6 months, assessed with the use of a standardized technique\textsuperscript{22} and reporting tool by independent, accredited vascular technologists (with the exception of one center where scanning was performed by a surgeon who had not performed the treatment). Blinding with respect to the treatment used was not feasible.

**STUDY OVERSIGHT**

The trial was approved by the research ethics committee and the Medicines and Healthcare Products Regulatory Authority. Written informed consent was obtained from all participants. The trial was overseen by a trial steering committee and an independent data and safety monitoring committee. Data analysis was performed by statisticians at the Centre for Healthcare Randomised Trials. The project management group (the first six and the last four authors) takes responsibility for the accuracy and completeness of the data, analyses, and reporting and for the fidelity of the study to the protocol.

**STATISTICAL ANALYSIS**

An intention-to-treat analysis was performed for the prespecified comparisons of treatment with foam versus surgery, involving participants from strata A and B, and treatment with laser versus surgery, involving participants from stratum A. In addition, we performed a post hoc analysis of laser therapy versus foam sclerotherapy, involving participants from stratum A. The principal analysis of the trial was performed when all participants had completed the 6-month follow-up. Study analyses were conducted according to a prespecified statistical plan (available with the protocol at NEJM.org) with the use of SAS software, version 9.3 (SAS Institute).

To analyze comparisons between groups, we used general linear models with adjustments for covariates used in the minimization algorithm and, where possible, adjustments for baseline scores (AVVQ, EQ-5D, SF-36, and venous clinical severity scores). No adjustment was prespecified for multiple comparisons. However, for the secondary outcome measures presented here, we consider differences to be significant only for P values of less than 0.005. We analyzed the continuous outcomes with mixed-model repeated-measures analysis, with a compound-symmetry covariance matrix and with center fitted as a random effect. Saphenous-vein ablation rates were analyzed with the use of ordinal logistic regression, and rates of complications were analyzed with the use of binary logistic regression. Sensitivity analyses (see Table S1 in the Supplementary Appendix, available at NEJM.org) were carried out in the case of missing AVVQ responses at 6 months.\textsuperscript{29}

The initial planned sample size was 1015 patients, which, at a two-sided 5% significance level, would provide more than 90% power to detect a difference of 0.25 SD in the AVVQ score for the comparison of foam sclerotherapy with surgery.\textsuperscript{30,31}
and 80% power to detect a difference of 0.25 SD in the AVVQ score for the comparison of laser with surgery. The data and safety monitoring committee and trial steering committee approved a revised recruitment target of 779 patients on the basis of data showing that the correlation between the AVVQ score at baseline and at 6 months was better than originally assumed. Only the data and safety monitoring committee was aware of the outcome data according to group assignment during the trial.

### Results

#### Patients and Treatment

Of 6592 participants referred from primary care for consideration of treatment, 3369 (51.1%) met the eligibility criteria, of whom 798 (23.7%) consented to participation in the trial (Fig. 1) and 785 were included in the trial. The reasons for ineligibility and the reasons that eligible patients declined randomization are summarized in Table S2 in the Supplementary Appendix. Most ineligible participants did not require treatment of the truncal saphenous veins (i.e., the patients were asymptomatic or did not have reflux) or had recurrent varicose veins. Most of the eligible participants who declined to undergo randomization had a preference for a specific treatment. Baseline demographic and clinical data for the three groups are shown in Table 1. Baseline characteristics were generally similar among the groups, with the following exceptions: the diameter of the great saphenous vein was larger in the laser group than in the foam group ($P=0.008$), the incidence of reflux below the knee was greater in the foam group than in the laser group ($P=0.03$), and the SF-36 mental component score, which was slightly higher (better generic quality of life) in the laser group than in the foam group ($P=0.04$) (see Table S3 in the Supplementary Appendix).

#### Secondary Outcomes

#### Quality of Life

Table S4 in the Supplementary Appendix shows results for secondary quality-of-life outcomes. At 6 weeks, significant between-group differences ($P<0.005$) included a lower AVVQ score (indicating a better disease-specific quality of life) in the surgery group than in the foam group (effect size, $-2.3$; 95% CI, $-3.7$ to $-0.9$) and lower SF-36 scores (indicating a worse generic quality of life) in the surgery group than in the laser group for the domains of bodily pain (effect size, $-2.7$; 95% CI, $-4.4$ to $-0.9$), vitality (effect size, $-2.3$; 95% CI, $-3.9$ to $-0.8$), role limitations due to emotional health (effect size, $-2.4$; 95% CI, $-4.0$ to $-0.8$), and role limitations due to physical health (effect size, $-3.5$; 95% CI, $-5.2$ to $-1.8$). These four SF-36 domain scores did not differ significantly (with $P<0.005$ considered to indicate statistical signifi-
cance) between groups at 6 months. For the post hoc comparisons of laser treatment versus foam treatment, only the EQ-5D score was significantly lower (indicating a worse generic quality of life) in the foam group at 6 weeks (0.044; 95% CI, 0.014 to 0.074).

Clinical Outcomes
The venous clinical severity score was moderately but significantly lower (indicating less residual venous disease) in the surgery group than in the foam group after 6 weeks, but there was no significant difference at 6 months (see Table S5 in the Supplementary Appendix). There was no significant difference in this score between the laser and the surgery groups or between the laser and the foam groups at either time.

At both 6 weeks and 6 months, there were fewer residual varicose veins, as assessed by both participants and nurses, in the surgery group than in the foam group, but the differences were small. There were fewer residual varicose veins in the surgery group than in the laser group at 6 weeks, but not at 6 months, as reported by the nurses but not the participants. There were no significant differences between the laser and the foam
Comparing Treatments for Varicose Veins

groups at 6 weeks, but there were fewer residual veins in the laser group than in the foam group at 6 months, as reported by the participants (but not the nurses) (see Table S5 in the Supplementary Appendix).

The frequency of completely successful ablation of great saphenous veins was significantly higher among participants who were randomly assigned to surgery (84.4%) or laser treatment (83.0%) than among those assigned to foam treatment (54.6%, \( P < 0.001 \) for both comparisons). There was no significant difference in success rates between the surgery and the laser groups (Table 3).

### Complications

Serious and nonserious adverse events are shown in Table 4. There were no significant differences between groups in the number of serious adverse events (see Table S6 in the Supplementary Appendix for details of serious adverse events). The frequency of any procedural complications (i.e., complications occurring during treatment) was lower in the laser group (1.0%) than in the foam group (6.2%) or the surgery group (7.1%) (\( P < 0.001 \) for both comparisons). At 6 weeks and 6 months, the frequency of overall complications (predominantly lumpiness and skin staining) was greatest in the foam group; the complication rate was significantly greater in the foam group than in the surgery group or the laser group at 6 weeks and was significantly greater in the foam group than in the surgery group at 6 months.

### Discussion

In this multicenter trial comparing foam sclerotherapy or laser treatment with surgery for the treatment of primary varicose veins, quality-of-life measures at 6 months did not differ substantially between groups. However, patients treated with foam had moderately worse outcomes on a
measure of disease-specific quality of life (AVVQ) than did those who underwent surgery. These differences were small, and their clinical importance is uncertain.

The frequency of complications (e.g., lumpiness, skin staining, and numbness) was lower after treatment with laser than after treatment with either foam or surgery; these differences are likely to have affected quality of life. However, the use of foam sclerotherapy for residual varicosities in a third of the participants in the laser group at 6 weeks may have attenuated early quality-of-life benefits associated with this treatment as compared with treatment with foam or surgery.31

The baseline scores and overall improvements in quality of life that we observed were similar to those reported in smaller European randomized trials, supporting the generalizability of our findings.2,5,6,8-10,12,32 The nature and frequency of complications were also similar to those previously reported in the literature.1-16

The three groups had similar improvements in the venous clinical severity score at 6 months. However, successful ablation of the great saphenous veins at 6 weeks occurred significantly less often after foam treatment (complete ablation, 55%; partial ablation with a patent segment and no reflux, 23%) than after either surgery (complete ablation, 84%; partial ablation, 6%) or laser treatment (complete ablation, 83%; partial ablation, 8%). Observed ablation rates, in particular for foam sclerotherapy, were lower than the rates in some previous studies, which used less strict definitions of success,1,6,10,14 but were similar to the rates reported in two randomized trials that

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<tbody>
<tr>
<td><strong>AVVQ¶</strong></td>
<td></td>
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</tr>
<tr>
<td>Baseline</td>
<td>17.8±9.1</td>
<td>17.6±9.9</td>
<td>18.2±9.1</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>6 Mo after treatment</td>
<td>7.9±8.4</td>
<td>9.1±7.9</td>
<td>7.8±7.5</td>
<td>−1.7 (−3.0 to −0.5)</td>
<td>−0.6 (−2.2 to 0.9)</td>
<td>−1.1 (−2.6 to 0.4)</td>
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<td><strong>EQ-5D</strong></td>
<td></td>
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<tr>
<td>Baseline</td>
<td>0.792±0.168</td>
<td>0.803±0.177</td>
<td>0.784±0.175</td>
<td>0.005</td>
<td>−0.015</td>
<td>0.025</td>
</tr>
<tr>
<td>6 Mo after treatment</td>
<td>0.903±0.171</td>
<td>0.895±0.174</td>
<td>0.881±0.202</td>
<td>(−0.025 to 0.035)</td>
<td>(−0.051 to 0.021)</td>
<td>(−0.010 to 0.059)</td>
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<tr>
<td><strong>SF-36 physical component††</strong></td>
<td></td>
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<tr>
<td>Baseline</td>
<td>48.6±7.8</td>
<td>48.9±8.0</td>
<td>48.2±8.6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 Mo after treatment</td>
<td>52.6±7.3</td>
<td>52.3±8.5</td>
<td>52.4±8.9</td>
<td>1.0 (−0.2 to 2.3)</td>
<td>0.1 (−1.4 to 1.6)</td>
<td>0.7 (−0.8 to 2.1)</td>
</tr>
<tr>
<td><strong>SF-36 mental component††</strong></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>51.9±9.0</td>
<td>52.4±8.7</td>
<td>51.2±9.4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 Mo after treatment</td>
<td>53.5±7.7</td>
<td>52.2±9.1</td>
<td>52.1±8.6</td>
<td>0.2 (−1.1 to 1.6)</td>
<td>−1.3 (−2.9 to 0.2)</td>
<td>1.5 (0.0 to 3.1)</td>
</tr>
</tbody>
</table>

* Plus–minus values are means ±SD.
† The comparison includes participants from stratum A (eight hospitals offering all three treatment options) and stratum B (three hospitals offering treatment with only foam or surgery).
‡ The comparison includes participants from stratum A only.
§ The post hoc comparison of laser versus foam includes participants from stratum A only.
¶ The Aberdeen Varicose Veins Questionnaire (AVVQ) consists of 12 questions and a leg diagram assessing disease-specific quality of life, with scores ranging from 0 (best) to 100 (worst).
‖ P = 0.006.
** The EuroQol Group 5-Dimension Self-Report Questionnaire (EQ-5D) consists of five questions, each with three response options, with scores ranging from −0.594 (worst) to 1 (best).
†† The Medical Outcomes Study 36-Item Short-Form Health Survey (SF-36) comprises a physical component summary score and a mental component summary score, each consisting of the sum of four individual physical domains, with scores ranging from 0 (worst) to 100 (best).
‡‡ P = 0.048.
used definitions of technical success that were similar to ours.12,15 In contrast to the assessment of ablation in previous trials, we determined whether ablation was complete or partial on the basis of duplex ultrasound scans obtained by independent, accredited vascular technologists rather than by the surgeons who had performed the treatment. The disparity between clinical measures of success and technical success observed in this study has also been observed in other trials of treatment for varicose veins.18,19

For patients undergoing laser treatment, concurrent treatment of varicosities remains controversial.16 A previous, single-center trial showed significant improvements in the AVVQ score at 6 weeks among patients who underwent phlebectomy concomitantly with laser therapy, as compared with those who underwent laser therapy alone.33 In our study, participants in the laser group received treatment directed to the main saphenous vein, without concomitant phlebectomies (except at one center). We found no significant differences in the AVVQ score among patients undergoing laser treatment versus surgery at 6 weeks, despite the use of concomitant phlebectomies in the surgery group.

The limitations of our study should be acknowledged. First, it was not feasible for the participants or assessors to be kept unaware of the assigned treatment. Second, we did not include a group of patients assigned to a sham procedure and thus cannot assess the effect of treatment relative to such a control. Third, eight comparisons involved primary outcomes, and a large number of comparisons involved secondary outcomes; thus, it is likely that some differences may have occurred by chance. We considered differences for secondary outcome measures to be significant only for $P \leq 0.01$.

In conclusion, our multicenter trial comparing the clinical effectiveness of endovenous laser ablation, foam sclerotherapy, and surgery for the treatment of varicose veins showed no clinically substantial between-group differences in quality of life. Moderate differences in disease-specific quality of life favored surgery over treatment.
with foam, and moderate differences in generic quality of life favored laser treatment over foam. All treatments had similar clinical efficacy, but there were fewer complications after laser treatment, and ablation rates were lower after treatment with foam.

The views expressed in this article are those of the authors and do not necessarily reflect the views of the National Institute for Health Research, the National Health Service, the Department of Health, or the funders that provided institutional support for this study.

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Disclosure forms provided by the authors are available with the full text of this article at NEJM.org.

We thank Janice Cruden for her secretarial support and data management; Gladys McPherson and the programming team at the Centre for Healthcare Randomised Trials; Tracey Davidson, Lynda Constable, Jackie Ellington, Laura Elliott, and Yvonne Ferrie for help with scoring the Aberdeen Varicose Vein Questionnaire; Luke Vale and Laura Ternent, our original economists in the group; members of the Project Management Group for their ongoing advice and support of the trial; members of the study team (Graeme MacLennan, Maria Prior, and Denise Bolsover) who contributed to the behavioral recovery component of the trial; the independent members of the trial steering committee (Alun Davies [chair], Ian Loftus, and Jane Nixon) and the data and safety monitoring committee (Gerry Stansby [chair], Winston Banya, and Marcus Flather); and the staff members at recruitment sites (see the Supplementary Appendix) who facilitated recruitment, treatment, and follow-up of trial participants.

Table 4. Serious Adverse Events and Other Complications According to Treatment Group.*

<table>
<thead>
<tr>
<th>Event</th>
<th>Laser Group</th>
<th>Foam Group</th>
<th>Surgery Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total number/total number (percent)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Serious adverse events overall</td>
<td>7/210 (3.3)</td>
<td>11/286 (3.8)</td>
<td>10/289 (3.5)</td>
</tr>
<tr>
<td>Serious adverse events related to treatment</td>
<td>2/210 (1.0)</td>
<td>3/286 (1.0)</td>
<td>4/289 (1.4)</td>
</tr>
</tbody>
</table>

Other complications

<table>
<thead>
<tr>
<th>Event</th>
<th>Laser Group</th>
<th>Foam Group</th>
<th>Surgery Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Any procedural complication during treatment</td>
<td>2/205 (1.0)</td>
<td>17/276 (6.2)†</td>
<td>19/267 (7.1)†</td>
</tr>
<tr>
<td>Any complication at 6 wk</td>
<td>103/193 (53.4)</td>
<td>219/265 (82.6)†</td>
<td>168/251 (66.9)‡</td>
</tr>
<tr>
<td>Any complication at 6 mo</td>
<td>89/183 (48.6)</td>
<td>144/251 (57.4)§</td>
<td>109/236 (46.2)</td>
</tr>
<tr>
<td>Numbness at 6 wk</td>
<td>22/193 (11.4)¶</td>
<td>15/265 (5.7)</td>
<td>45/251 (17.9)‖</td>
</tr>
<tr>
<td>Numbness at 6 mo</td>
<td>17/183 (9.2)¶</td>
<td>10/251 (4.0)</td>
<td>37/236 (15.6)</td>
</tr>
<tr>
<td>Persistent bruising at 6 wk</td>
<td>10/193 (5.2)</td>
<td>49/265 (18.5)†</td>
<td>32/251 (12.7)‡</td>
</tr>
<tr>
<td>Persistent bruising at 6 mo</td>
<td>25/183 (13.6)</td>
<td>38/251 (15.2)</td>
<td>40/236 (17.0)</td>
</tr>
<tr>
<td>Persistent tenderness and discomfort at 6 wk</td>
<td>41/193 (21.2)</td>
<td>122/265 (46.0)†</td>
<td>79/231 (31.5)¶</td>
</tr>
<tr>
<td>Skin loss or ulceration at 6 wk</td>
<td>0/193</td>
<td>2/265 (0.8)</td>
<td>1/251 (0.4)</td>
</tr>
<tr>
<td>Skin loss or ulceration at 6 mo</td>
<td>1/183 (0.6)</td>
<td>2/251 (0.8)</td>
<td>0/236</td>
</tr>
<tr>
<td>Lumpiness at 6 wk</td>
<td>36/193 (18.7)</td>
<td>171/265 (64.5)†**</td>
<td>83/251 (33.1)‡</td>
</tr>
<tr>
<td>Lumpiness at 6 mo</td>
<td>25/183 (13.6)§</td>
<td>67/251 (26.6)‡**</td>
<td>17/236 (7.2)‡</td>
</tr>
<tr>
<td>Development of thread vein at 6 wk</td>
<td>10/193 (5.2)</td>
<td>27/265 (10.2)‡</td>
<td>21/231 (8.4)</td>
</tr>
<tr>
<td>Development of thread vein at 6 mo</td>
<td>24/183 (13.2)</td>
<td>34/251 (13.6)</td>
<td>26/236 (11.0)</td>
</tr>
<tr>
<td>Skin staining at 6 wk</td>
<td>18/193 (9.3)</td>
<td>105/265 (39.6)†**</td>
<td>20/251 (8.0)</td>
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<tr>
<td>Skin staining at 6 mo</td>
<td>32/183 (17.4)§</td>
<td>92/251 (36.6)†**</td>
<td>24/236 (10.2)</td>
</tr>
</tbody>
</table>

* The comparison between the foam group and the surgery group includes participants from strata A and B; the comparison between the surgery group and the laser group includes participants from stratum A only. The post hoc analysis of the laser group versus the foam group includes participants from stratum A only; 31% of the patients in the laser group also underwent foam treatment of varicosities. Thread veins are small clusters of blue or red veins.

† P<0.001 for the comparison with the laser group.
‡ P<0.05 for the comparison with the laser group.
§ P<0.05 for the comparison with the surgery group.
¶ P<0.05 for the comparison with the foam group.
‖ P<0.001 for the comparison with the foam group.
** P<0.001 for the comparison with the surgery group.
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27. EQ-SD. Rotterdam, the Netherlands: EuroQol (http://www.euroqol.org).


