Randomized clinical trial comparing surgery, endovenous laser ablation and ultrasound-guided foam sclerotherapy for the treatment of great saphenous varicose veins

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Background: Endovenous ablation techniques and ultrasound-guided foam sclerotherapy (UGFS) have largely replaced open surgery for treatment of great saphenous varicose veins. This was a randomized trial to compare the effect of surgery, endovenous laser ablation (EVLA) (with phlebectomies) and UGFS on quality of life and the occlusion rate of the great saphenous vein (GSV) 12 months after surgery.

Methods: Patients with symptomatic, uncomplicated varicose veins (CEAP class C2–C4) were examined at baseline, 1 month and 1 year. Before discharge and at 1 week, patients reported a pain score on a visual analogue scale. Preoperative and 1-year assessments included duplex ultrasound imaging and the Aberdeen Varicose Vein Severity Score (AVVSS).

Results: The study included 214 patients: 65 had surgery, 73 had EVLA and 76 had UGFS. At 1 year, the GSV was occluded or absent in 59 (97 per cent) of 61 patients after surgery, 71 (97 per cent) of 73 after EVLA and 37 (51 per cent) of 72 after UGFS ($P < 0.001$). The AVVSS improved significantly in comparison with preoperative values in all groups, with no significant differences between them. Perioperative pain was significantly reduced and sick leave shorter after UGFS (mean 1 day) than after EVLA (8 days) and surgery (12 days).

Conclusion: In comparison with open surgery and EVLA, UGFS resulted in equivalent improvement in quality of life but significantly higher residual GSV reflux at 12-month follow-up.

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anatomical success. Owing to variation in both the quality of the studies and reporting of the results, the quality of the evidence was graded only moderate.

The aim of the present randomized trial was to compare the efficacy of surgery, EVLA and UGFS in patients with primary symptomatic, uncomplicated great saphenous varicose veins (Clinical Etiologic Anatomic Pathophysiologic (CEAP) clinical grade C2–C4).

**Methods**

Consecutive patients with varicose veins were screened at two participating university hospitals in Finland between November 2007 and May 2010. Patients who fulfilled the inclusion criteria and were willing to participate were enrolled in a prospective randomized trial. The study was approved by the Ethics Committee of Helsinki University Central Hospital and Tampere University Hospital.

All patients provided written informed consent before participating in the study, according to the principles of the Declaration of Helsinki.

The inclusion criteria were: unilateral symptomatic, uncomplicated varicose veins (CEAP clinical classification C2–C4), duplex ultrasound-verified reflux in the GSV, mean diameter of the GSV in the thigh 5–10 mm, and age 20–70 years. Duplex ultrasound imaging was done in standing position, and reflux was measured after pneumatic compression of the calf. Incompetence was defined as a reflux of more than 0.5 s.

Exclusion criteria were: peripheral arterial disease, lymphoedema, BMI exceeding 40 kg/m², pregnancy, allergy to the sclerosant or lidocaine, severe general illness, malignancy, previous deep vein thrombosis and coagulation disorder.

**Randomization**

The patients were randomized in outpatients to receive surgery, EVLA or UGFS. This was done using block randomization with sealed envelopes.

**Procedures**

Both the surgical and EVLA procedures were performed in the day surgery unit.

In the surgical procedure, the saphenofemoral junction (SFJ) was exposed in the groin and side branches were ligated back to the femoral vein. Retrograde invagination stripping of the GSV was done, usually down to below the knee. Tumescent solution (450 ml Ringer’s solution with 50 ml 1 per cent lidocaine with adrenaline (epinephrine)) was injected into the tunnel of the stripped GSV. After stripping, hook phlebectomies were done through tiny incisions, using tumescent solution to minimize haematoma formation. The wound in the groin was closed with subcuticular sutures. Phlebectomy wounds were closed with surgical tape. Most patients had general anaesthesia. After the procedure, the leg was wrapped in bandages and covered with a class 2 graduated compression stocking. After 48 h, patients removed their bandages and then used the stocking alone during the day for up to 2 weeks. A prescription for paracetamol with, or without codeine, or ibuprofen was given on discharge with the instruction that the medication be used when necessary.

EVLA procedures were done under tumescent local anaesthesia (same solution as above) injected around the GSV under ultrasound guidance. A light sedative was administered before (diazepam) and during (alfentanil, propofol) the procedure. The laser ablation was carried out under duplex guidance. At the beginning of the study, a 980-nm diode laser (Ceralas® D 980; Biolitec, Bonn, Germany) was used, but during the study this was replaced with a 1470-nm radial laser (ELVes®; Biolitec). A pulsed mode, with a 1.5-s impulse and 12 W of energy, was used routinely, with the aim of applying 70 J/cm GSV. The EVLA catheter tip was positioned 1.5–2 cm below the SFJ using ultrasound guidance. After the laser procedure, phlebectomies were done, as in the surgical procedure. No skin sutures were used. Thereafter, the protocol was the same as that after the surgical procedure.

UGFS was undertaken in outpatients by a vascular surgeon with an assisting nurse. Patients were examined by duplex ultrasound imaging before the treatment with the patient standing, and the cause of reflux was confirmed. During the treatment the patient remained supine. The GSV was cannulated under ultrasound guidance, usually at proximal thigh level and immediately below the knee. The sclerosant foam was prepared with a double-syringe technique with a sclerosant to air ratio of 1:2. The sclerosants used were polidocanol 1 per cent (Aetoxy sclero®; Kreussler, Wiesbaden, Germany) and sodium tetradecyl sulphate (STS) 1 per cent and 3 per cent (Fibrovein™; STD Pharmaceutical Products, Hereford, UK). The injection of foam was monitored by ultrasound imaging. After treatment, a compression stocking was applied with the instruction to wear it continuously for 3 days, followed by daytime use for 11 days. At 1-month follow-up, a duplex ultrasound examination was done and, if any reflux was observed, a second treatment with foam was carried out. These patients were seen again 4 weeks after the second treatment, and the need for a possible third treatment was checked by duplex imaging.
The primary outcome measures were: 1-year occlusion (or absence) rate of GSV on routine duplex imaging, and changes in disease-specific quality of life according to the Aberdeen Varicose Vein Severity Score (AVVSS). The diameter of the GSV 20 cm below the groin was also measured and compared with preoperative values. Secondary outcome measures were: perioperative pain measured using a visual analogue scale (VAS) from 0 (no pain) to 10 (worst possible, unbearable, excruciating pain) at the time of discharge and at 1 week after the procedure; duration of sick leave; and rate of complications such as haematoma, pigmentation, thrombophlebitis and paraesthesia.

Statistical analysis
Sample size calculations indicated that to detect a 20 per cent difference in occluded or absent GSV between the groups, with \( \alpha = 0.05 \) and \( \beta = 0.8 \), 70 patients would be needed in each group.

Data were analysed according to intention to treat. The primary endpoint, occlusion or absence of GSV, was analysed in the overall study group and in three subgroups according to the size of the GSV in the mid-thigh.

Continuous variables are reported as mean(s.d.) (range). Baseline and follow-up variables were compared using the paired-samples t test and repeated-measures test. Statistical analysis was done using statistical software for Windows® (SPSS® version 19.0; IBM, Armonk, New York, USA).

Results
A total of 598 consecutive patients were screened between November 2007 and May 2010. Of these, 233 patients (233 legs) fulfilled the inclusion criteria and were willing to participate; they were randomized in the trial (Fig. 1). Nineteen randomized patients were excluded from the study before treatment. The most common reason was that the patient met an exclusion criterion that was not recognized at the time of randomization. Thus, the final study population comprised 214 patients: 65 in the surgery group, 73 in the EVLA group and 76 in the UGFS group. Owing to the operating surgeon’s preference, five patients originally randomized to EVLA were treated with surgery but, because the analysis was made according to intention to treat, these patients were analysed in EVLA group.

All 214 patients attended the 1-month follow-up, and 206 (96.3 per cent) the 1-year follow-up: 61 of 65 after surgery, all 73 patients who had EVLA and 72 of 76 who had UGFS.
Table 1  Demographics and preoperative measurements

|                        | Surgery (n = 65) | EVLA (n = 73) | UGFS (n = 76) | Total (n = 214) |
|------------------------|----------------|--------------|--------------|----------------|
| Age (years)*           | 47.3(11.3) (27–75) | 47.0(13.4) (20–73) | 48.3(12.7) (23–74) | 47.6(12.4) (20–75) |
| Sex ratio (F : M)      | 55 : 10         | 55 : 18      | 58 : 18      | 168 : 46       |
| BMI (kg/m²)*           | 25.1(3.7) (19–37) | 25.2(3.6) (19–35) | 25.8(4.6) (20–42) | 25.4(4.0) (19–42) |
| Diameter of GSV (mm)   |                |              |              |                |
| At SFJ                 | 8.7(2.0) (5–14) | 8.5(2.2) (5–15) | 8.4(1.7) (5–13) | 8.5(2.0) (5–15) |
| Below groin            | 6.6(1.3) (4–11) | 6.8(1.2) (4–10) | 6.7(1.2) (4–10) | 6.7(1.2) (4–11) |
| Mean                   | 6.2(1.1) (4–9)  | 6.3(1.1) (4–8) | 6.7(1.1) (4–9) | 6.2(1.1) (4–9) |
| Baseline CEAP class    |                |              |              |                |
| C2                     | 33             | 27           | 26           | 86             |
| C3                     | 26             | 36           | 37           | 99             |
| C4                     | 6              | 10           | 13           | 29             |
| Baseline AVVSS*        | 30.2(8-3) (16–45) | 32.4(6-7) (18–51) | 31.7(7-6) (13–52) | 31.5(6-9) (13–52) |

Values are mean(s.d.) range. EVLA, endovenous laser ablation; UGFS, ultrasound-guided foam sclerotherapy; GSV, great saphenous vein; SFJ, saphenofemoral junction; CEAP, Clinical Etiologic Anatomic Pathophysiological; AVVSS, Aberdeen Varicose Vein Severity Score.

Fig. 2 Patency of the great saphenous vein (GSV) at various diameters on duplex ultrasound imaging 1 year after a surgery, b endovenous laser ablation (EVLA) or c ultrasound-guided foam sclerotherapy (UGFS)
There were no significant differences in the basic demographics, CEAP clinical classification, AVVSS or GSV dimensions at baseline between the study groups (Table 1).

Perioperative results

The mean(s.d.) duration of treatment was 95(19) (range 62–155) min in the surgery group and 83(17) (range 50–139) min in the EVLA group ($P<0.001$). Twenty-six patients (34 per cent) in the UGFS group received two treatments; no patient required a third treatment. The sclerosants employed for the first treatment were: STS 3 per cent (64 patients), STS 1 per cent (6), polidocanol 3 per cent (4) and polidocanol 1 per cent (2). The mean volume of foam used in the GSV was 4.7(1.6) (range 2.0–9.0) ml. Some 33 per cent also had foam injected into varicose tributaries: mean volume 4.6 (range 2.0–12.0) ml. In the second treatment session, the mean volume of foam used in the GSV was 3.8(2.0–10.0) ml.

Primary outcome measures

At 1 year, the GSV was completely occluded or absent in 59 (97 per cent) of 61 patients after surgery, 71 (97 per cent) of 73 after EVLA and 37 (51 per cent) of 72 after UGFS. The GSV was partially occluded in two patients (3 per cent), none (0 per cent) and 21 patients (29 per cent) in the respective groups. The difference between UGFS and the two other treatments was significant ($P<0.001$). No patient in the surgery group and only two (3 per cent) in the EVLA group had a patent GSV after 1 year, compared with 14 (19 per cent) in the UGFS group. Of the two patients with a patent GSV in the EVLA group at 1 year, one had a tiny but patent GSV with no reflux and the other had asymptomatic reflux in a very narrow GSV. On duplex imaging at 1 year, reflux was seen in the below-knee GSV in 13, 16 and 33 per cent of patients in the surgery, EVLA and UGFS groups respectively ($P=0.008$ for UGFS versus other two procedures). Reflux in another unnamed thigh vein was present in 8, 10 and 12 per cent respectively ($P=0.471$ between groups). When GSV patency rates were analysed according to size of the GSV before treatment, there was a clear correlation between larger diameter and GSV patency, but only in the UGFS group (Fig. 2).

At baseline, there were no significant differences in median AVVSS between the groups. At 1 year, median AVVSS was significantly improved in all groups and there were no significant differences between the groups (Fig. 3).

After the 1-year follow-up visit, 16 patients had additional treatment: four patients (7 per cent) in the surgery group (UGFS for a thigh vein in 3 patients and ligation of a perforating vein in 1); one patient (1 per cent) after EVLA (foam sclerotherapy to a tributary at the ankle); and 11 patients (15 per cent) after UGFS (2 stripping of GSV, 5 EVLA of GSV and 4 repeat UGFS) ($P=0.009$).

Secondary outcome measures

Pain after treatment was significantly reduced (lower VAS score) after UGFS in comparison with the surgery and EVLA groups, both at the time of discharge, and after 1 week (Fig. 4).

The mean duration of sick leave was 12(6) (range 0–33) days after surgery, 8(5) (range 0–29) days after EVLA and 1(3) (range 0–21) days after UGFS ($P<0.001$ between UGFS and the 2 other groups). There were no major complications related to the procedures. Three patients (4 per cent) in the EVLA group and three (5 per cent) in the surgery group had a superficial wound infection. All resolved with oral antibiotics; none of the patients needed treatment in hospital.

At the 1-month follow-up, 62 per cent of patients in the surgery group had a haematoma (defined as visible localized aggregate of extravasated blood) in the operated leg, compared with 42 per cent in the EVLA group and 20 per cent in the UGFS group ($P=0.001$ between groups).
Skin pigmentation was more common after UGFS (67 per cent) than after surgery and EVLA (5 and 4 per cent respectively; \( P < 0.001 \) for UGFS versus other 2 groups). Paraesthesia was rare in all groups (2, 3 and 1 per cent respectively). Palpable lumps in the veins under the skin were present at 1 month in 54 per cent of patients after surgery, 47 per cent after EVLA and 91 per cent after UGFS (\( P < 0.001 \) for UGFS versus other two groups).

**Discussion**

The present prospective randomized trial compared three interventions for unilateral primary great saphenous varicose veins. Reflux in the GSV was extremely rare at 1 year after surgical stripping or EVLA; however, after UGFS, reflux was seen in half of the patients, and the GSV was patent and refluxing in one of five. Despite these differences, disease-specific quality of life was significantly better in all three groups after 6 months than before intervention, but worse after UGFS. The same trend was seen in other studies.\(^5\)−\(^7\)^\(^10\).

The present study also examined the effect of GSV diameter on outcome. The occlusion rate after UGFS was clearly associated with GSV diameter: less than 40 per cent in GSVs of 9 mm or larger in mid-thigh diameter, compared with 75 per cent in GSVs of less than 6 mm. This suggests that UGFS should not be recommended for veins larger than 6 mm in diameter.

There are some short-term advantages from UGFS. In all three groups, patients reported increased pain after 1 week compared with immediately after treatment; the proportion of patients with no or minimal pain decreased during the first week. The majority of patients who had UGFS had no or minimal pain at 1 week. These findings contrast with those of Rasmussen and colleagues,\(^6\) who reported that patients who had surgery or EVLA had the worst pain immediately after the procedure, which improved steadily over the first week. Some of the variation may be explained by the differences in the timing of the first measurement. In the present study, patients reported the VAS score immediately before discharge, when they may have had some remaining effects of perioperative analgesia. In the Danish study, the first measurement was completed on the first postoperative day.

Similarly, recovery from the intervention was fastest in patients who underwent UGFS. Recovery to daily work was longest after surgery; duration of sick leave was also greatest in the surgery group. These results accord with other studies,\(^2\)\(^,\)^\(^3\), which reported sick leave of 1 week.
after EVLA and 2 weeks after surgery. Rasmussen and colleagues\(^6\) reported shorter sick leave of 7-6 days after surgery, possibly owing to the use of tumescent anaesthesia in surgical procedures.

The strength of the present study is that it was randomized, with excellent follow-up, 96.3 per cent at 1 year. The main weakness is the relatively short follow-up so far. The long-term consequence of the high reflux rate after UGFS remains unclear. Another issue is that the foam used was more concentrated (air to sclerosant ratio 2:1) than in other studies. The impact of this is unknown as there are no randomized trials regarding the optimal foam recipe.

Patients with great saphenous varicose veins should understand that, although UGFS has some short-term advantages in recovery, and equivalent quality of life after 1 year, longer follow-up of the present patients may reveal that the higher rate of recurrent/residual GSV reflux at 1 year increases the long-term risk of recurrent varicose veins.

**Collaborators**

Finnish Venous Study Collaborators: V. P. Suominen (Tampere University Hospital), P. Vikatmaa, P. Aho, M. Lepäntalo, K. Halmesmäki, S. Laukontaus, E. M. Weselius, S. Vuorisalo (Helsinki University Hospital).

**Disclosure**

The authors declare no conflict of interest.

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