1 PRODUCT NAME
FIBROVEIN 3.0%, 1.0%, 0.5% and 0.2% intravenous injection

2 QUALITATIVE AND QUANTITATIVE COMPOSITION
Sodium Tetradecyl Sulphate 3.0%, 1.0%, 0.5% and 0.2% intravenous injection

Contains benzyl alcohol 20mg/ml. For a full list of excipients, see section 6.1.

3 PHARMACEUTICAL FORM
Solution for injection.

Clear, colourless, sterile solution.

pH 7.5 – 7.9.

Osmolarity 247 – 273 mOsm/kg.

4 CLINICAL PARTICULARS
4.1 Therapeutic indications
Fibrovein 3% and 1%: For the treatment of varicose veins of the leg by injection sclerotherapy.

Fibrovein 0.5%: For the treatment of varicose veins and venous flares of the leg by injection sclerotherapy.

Fibrovein 0.2%: For the treatment of minor venules and spider veins (venous flares) by injection sclerotherapy.

4.2 Dose and method of administration
Fibrovein is for intravenous use only. The strength of solution required depends on the size and degree of varicosity. Spider veins should only be treated with the 0.2%, reticular veins with 0.5%, the 1% solution will be found most useful for small to medium varicosities and the 3% solution for larger varicosities. The size of non-visible varicose veins should be measured under ultrasound.

The sclerosant should be administered intravenously in small aliquots at multiple sites along the vein to be treated either as a liquid or as a sclerosant/air mixture (foam), for the treatment of larger veins with the 1% and 3% solutions. The objective is to achieve optimal destruction of the vessel wall with the minimum concentration of sclerosant necessary for a clinical result. If the concentration is too high necrosis or other adverse sequelae may occur.

Recommended doses and dosage schedules:
Adults
Fibrovein 3.0%: 0.5 to 1.0ml of 3.0% Fibrovein injected intravenously at each of 4 sites (maximum 4ml).

Fibrovein 1.0%: 0.25 to 1.0ml of 1.0% Fibrovein injected intravenously into the lumen of an isolated segment of emptied superficial vein, followed by immediate compression. A maximum of 10 sites (10ml total) may be injected during one treatment session.
Fibrovein 0.5%: 0.25 to 1.0ml of 0.5% Fibrovein injected intravenously into the lumen of an isolated segment of emptied superficial vein, followed by immediate compression. A maximum of 10 sites (10ml total) may be injected during one treatment session.

Fibrovein 0.2%: 0.1 to 1.0ml of 0.2% Fibrovein injected intravenously at each of 10 sites (maximum 10ml).

Where special caution is indicated it is recommended that a test dose of 0.25 to 0.5 ml Fibrovein should be given followed by observation of the patient for several hours before administration of a second or larger dose.

As the volume to be injected is limited per session, repeated sessions are usually needed (2 to 4 on average). To prevent a possible allergic reaction, it is recommended that a small test dose of Fibrovein should be given at the beginning of each session.

For spider veins, the smallest of needles (30 gauge) should be used to perform the injection, which should be made slowly so that the blood content of these veins is expelled. In the treatment of spider veins an air block technique may be used.

**Children**
All strengths: not recommended in children

**The Elderly**
As for adults.

**Method of administration**
Strict aseptic technique must be maintained while handling Fibrovein.

Fibrovein is a single-use parenteral product. Once the container is opened, use immediately and discard any unused portion.

Visually inspect for particulate matter before use. Solutions that contain particulate matter should not be used.

**4.3 Contraindications**
- Hypersensitivity to sodium tetradecyl sulphate or to any component of the preparation and allergic conditions.
- Patients unable to walk due to any cause, bedridden patients.
- Patients with a high risk of thrombosis e.g. patients with a congenital predisposition to blood clots or with multiple risk factors such as hormonal contraceptives or hormone replacement therapy, significant obesity, smoking or extended periods of immobility
- Recent acute superficial thrombophlebitis, deep vein thrombosis or pulmonary embolism
- Recent surgery
- Local or systemic infection.
- Varicosities caused by pelvic or abdominal tumours unless the tumour has been removed
- Uncontrolled systemic disease such as diabetes mellitus, toxic hyperthyroidism, tuberculosis, asthma, neoplasm, sepsis, blood dyscrasias and acute respiratory or skin diseases
- Evolutive cancer
- Significant valvular incompetence of the deep veins
- Occlusive arterial disease
- Huge superficial veins with wide open communications to deeper veins
- Phlebitis migrans
- Acute cellulitis
- Acute infections

### 4.4 Special warnings and precautions for use

Fibrovein should only be administered by practitioners experienced in venous anatomy and the diagnosis and treatment of conditions affecting the venous system and familiar with proper injection technique.

Emergency resuscitation equipment should be immediately available. Allergic reactions, including anaphylaxis have been reported. The possibility of an anaphylactic reaction should be kept in mind, and the physician should be prepared to treat it appropriately. Before treatment, the healthcare professional should investigate patient’s risk factors and inform them about the risks of the technique.

As a reminder, sclerotherapy is contra-indicated in patients with high risk of thromboembolic events but should also be avoided in most situations at lower risk. Sclerotherapy is notably not recommended in patients with a history of thromboembolic events.

Nevertheless, if sclerotherapy is judged necessary, preventive anticoagulation can be initiated.

**Patent foramen ovale (PFO)**

Due to the risk of circulation of product, bubbles or particulates in the right heart, the presence of a PFO may enhance the occurrence of serious arterial adverse events. In patients with history of migraine with aura, serious cerebrovascular events or pulmonary hypertension, it is recommended to search for PFO before sclerotherapy. In patients with asymptomatic but known PFO, it is recommended to use smaller volumes and avoid Valsalva manoeuvre in the minutes after injection.

Patients with a PFO have been shown to be more likely to suffer from adverse events such as temporary neurological events, visual disturbances and migraine. A symptomatic PFO is a contraindication for use of Fibrovein as a foam (see section 4.3).

**Migraine**

Previous migraine sufferers should be treated with care. Patients with previous migraine have been shown to be more likely to suffer from visual disturbances and migraine, particularly following injections with foamed sclerosant.

Use smaller volumes in patients with history of migraine.

**TIA**

Patients with a past medical history of TIA should be treated with care. Patients with previous TIA have been shown to be more likely to suffer from visual disturbances and migraine, particularly following injections with foamed sclerosant.

**Truncular varicosities**

For the treatment of truncular varicosities, there should be a minimal distance of 8 to 10 cm between the site of foam injection and the saphenofemoral junction.

**Lymphoedema**
If venous insufficiency is associated with lymphoedema, the sclerosant injection may worsen local pain and inflammation for days or several weeks. Patients should be informed of this expected phase, which does not compromise efficacy.

**Extravasation**
Severe adverse local effects, including tissue necrosis, may occur following extravasation; therefore, extreme care in intravenous needle placement and using the minimal effective volume at each injection site are important. Pigmentation may be more likely to result if blood is extravasated at the injection site (particularly when treating smaller surface veins) and compression is not used.

**Intra-arterial injection**
Sclerosants must never be injected into an artery as this can cause extended tissue necrosis and may result in loss of the extremity. Injection under duplex ultrasound is recommended in order to avoid extravasations and arterial injection.

Healthcare professional should monitor the patient during and after the administration of Fibrovein. Symptoms of hypersensitivity (redness, pruritis, cough) or neurological symptoms (scotoma, amaurosis, migraine with aura, paraesthesia, focal deficit) may happen.

**Respiratory disease**
Special care should be taken in patients with laboured breathing (bronchial asthma) or a strong predisposition to allergies (see Dosage and Administration).

**Pre-injection evaluation**
Because of the danger of thrombosis extension into the deep venous system, thorough pre-injection evaluation for valvular competency should be carried out and slow injections with a small amount (not over 2 ml) of the preparation should be injected into the varicosity. Deep venous patency must be determined by non-invasive testing such as duplex ultrasound. Venous sclerotherapy should not be undertaken if tests such as Trendelenberg and Perthes, and angiography show significant valvular or deep venous incompetence.

**Follow-up**
Healthcare professionals should see the patient again after 1 month for a control of treatment efficacy and safety, by clinical and ultrasound evaluation.

The development of deep vein thrombosis and pulmonary embolism have been reported following sclerotherapy treatment of superficial varicosities. Patients should have post-treatment follow-up of sufficient duration to assess for the development of deep vein thrombosis. Embolism may occur as long as four weeks after injection of sodium tetradecyl sulphate. Adequate post-treatment compression may decrease the incidence of deep vein thrombosis.

**Underlying arterial disease**
Extreme caution in use is required in patients with arterial disease such as severe peripheral atherosclerosis or thromboangitis obliterans (Buerger’s Disease).

**Foot and malleolar area**
Special care is required when injecting above and posterior to the medical malleolus where the posterior tibial artery may be at risk.

**4.5 Interaction with other medicines and other forms of interaction**
No interaction studies have been performed.
4.6 Fertility, pregnancy and lactation

**Fertility**
It is not known whether sodium tetradecyl sulphate affects fertility.

**Use in Pregnancy**
Safety for use in pregnancy has not been established. There are no or limited amount of data from the use of sodium tetradecyl sulphate in pregnant women. Animal studies are insufficient with respect to reproductive toxicity. Treatment should be postponed until after childbirth. Fibrovein should be used only when clearly needed for symptomatic relief and when the potential benefits outweigh the potential hazards to the foetus.

**Use in Lactation**
It is not known whether sodium tetradecyl sulphate is excreted in human milk. Caution should be exercised when used in nursing mothers.

4.7 Effects on ability to drive and use machines
A bandage and/or compression stockings may be added after treatment. This could affect the ability to drive.

4.8 Undesirable effects
The most commonly reported side effects are pain on injection, urticaria, superficial thrombophlebitis and temporary skin pigmentation after treatment. Very rarely a permanent discoloration may remain along the path of the sclerosed vein segment. Ulceration may occur following extravasation of the drug. It is important to use the lowest strength that will sclerose the vein as many of the common side effects are caused by using a concentration that is too high.

Intra-arterial injection although very rare has been reported resulting in significant tissue necrosis including loss of the extremity.

The most serious side effects are anaphylactic shock and pulmonary embolism and deaths have been reported in patients receiving sodium tetradecyl sulphate.

Adverse events are listed below by system organ class and estimated frequency from published clinical data. Frequencies are defined using the following convention:

- **Very common** ≥ 1/10
- **Common** ≥ 1/100 to < 1/10
- **Uncommon** ≥ 1/1000 to < 1/100
- **Rare** ≥ 1/10,000 to < 1/1000
- **Very rare (includes isolated reports)** ≤ 1/10,000

<table>
<thead>
<tr>
<th>Immune disorders</th>
<th>Using liquid</th>
<th>Using foam</th>
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<tbody>
<tr>
<td>Systemic allergic reactions e.g. anaphylactic shock, asthma, generalised hives.</td>
<td>Very rare</td>
<td>Very rare</td>
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<table>
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<tr>
<th>Nervous system disorders</th>
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<tr>
<td>Migraine</td>
<td>Very rare</td>
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<tr>
<td>Headache, migraine, local sensitivity disturbances (paraesthesias). Vaso-vagal reactions e.g. fainting, confusion, dizziness, loss of consciousness.</td>
<td>Very rare</td>
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<tr>
<td>Nerve damage after extravasation of the drug</td>
<td>Very rare</td>
</tr>
<tr>
<td>Weakness (hemiparesis, hemiplegia), transient ischaemic attack (TIA), palpitations.</td>
<td>Very rare</td>
</tr>
<tr>
<td>Stroke</td>
<td>Very rare</td>
</tr>
</tbody>
</table>

**Eye disorders**

| Scotoma, scintillating scotoma. | Very rare | Uncommon |

**Vascular disorders**

| Superficial thrombophlebitis, phlebitis. | Common | Very common |
| Matting (growth of very fine spider veins in treated area). | Uncommon | Common |
| Deep vein thrombosis (mostly muscular and distal). | Very rare | Uncommon |
| Pulmonary embolism, vasculitis, circulatory collapse. | Very rare | Very rare |

Distal tissue necrosis following intra-arterial injection, may lead to gangrene. Most cases have involved the posterior tibial artery above the medial malleolus. Arterial spasm can occur despite intravenous injection.

| Very rare | Very rare |

**Respiratory disorders**

| Coughing, shortness of breath, sensation of pressure/tightness in the chest. | Very rare | Rare |

**Gastro-intestinal disorders**

| Nausea, vomiting, diarrhoea, feeling of swollen/thick tongue, dry mouth. | Very rare | Very rare |

**Skin and subcutaneous tissue disorders**

| Skin discolouration (hyperpigmentation, more rarely - haematoma & ecchymosis). | Uncommon | Common |
| Local allergic and non-allergic skin reactions e.g. erythema, urticaria, dermatitis, swelling/induration. | Uncommon | Uncommon |
| Local sloughing and necrosis of skin & tissues. | Rare | Rare |

| Common | Uncommon |
| Very rare | Very rare |

**General disorders**

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicine is important. It allows continued monitoring of the benefit/risk balance of the medicine. Healthcare professionals are asked to report any suspected adverse reactions [https://nzphvc.otago.ac.nz/reporting/](https://nzphvc.otago.ac.nz/reporting/).

**4.9 Overdose**

No case of systemic overdose has been reported. Using a higher concentration than recommended in small veins may lead to pigmentation and/or local tissue necrosis.

For advice on the management of overdose please contact the National Poisons Centre on 0800 POISON (0800 764766).
5 PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties
Pharmacotherapeutic group: Sclerosing agents for local injections, ATC code: C05BB04

Sodium tetradecyl sulphate is a sclerosing agent. Intravenous injection causes intima inflammation and thrombus formation. This usually occludes the injected vein. Subsequent formation of fibrous tissue results in partial or complete vein obliteration that may or may not be permanent.

Published clinical series have shown that Fibrovein converted to a foam is very effective at treating larger varicose veins e.g. great saphenous vein and tributaries. The foam is able to displace the blood and the sclerosant has more time to act on the endothelium compared to the liquid form. Some adverse events are more frequent following foam sclerotherapy than liquid sclerotherapy e.g. headache, migraine and visual disturbances. Adverse neurological events may also occur but these are rare.

5.2 Pharmacokinetic properties

Absorption
Fibrovein containing sodium tetradecyl sulphate is administered directly into the lumen of the isolated segment of vein/venule.

Distribution
In humans, the majority (75%) of an injected dose of radiolabelled 3% sodium tetradecyl sulphate rapidly disappeared from the empty varicose vein injection site into communicating blood vessels with rapid passage into the deep calf veins.

In rats, at 72 hours after intravenous dosing of radiolabelled sodium tetradecyl sulphate, tissue levels of radiolabel found in the sampled tissues (liver, kidney, lipid and skeletal muscle) were extremely low. Although there was some evidence of radiolabel associated with the injection site, the levels were very low.

Biotransformation
The metabolism of sodium tetradecyl sulphate has not been confirmed.

Elimination
Of an intravenously administered radiolabelled dose, 70% was recovered in the urine of rats within the first 24 hours post-dosing. At the end of the 72 hour post-dose period, 73.5% of the radiolabel had been recovered from the urine and 18.2% recovered from the faeces.

Hepatic/Renal Impairment
No pharmacokinetics studies have been performed in patients with hepatic or renal impairment.

5.3 Preclinical safety data
There are no additional data of relevance to the prescriber other than those already mentioned in other sections of the datasheet.

6 PHARMACEUTICAL PARTICULARS

6.1 List of excipients
Benzyl Alcohol BP, Disodium Hydrogen Phosphate BP, Potassium dihydrogen phosphate, Water for injections BP.
6.2 Incompatibilities
This product is not compatible with heparin.

In the absence of compatibility studies, this medicinal product should not be mixed with other medicinal products.

6.3 Shelf life
Shelf life is 36 months (3 years) from manufacture.

Each 2ml glass ampoule is for single use only. The in-use period of each 5ml multi-dose vial is a single session of therapy and for use in the treatment of a single patient. Unused vial contents should be discarded immediately afterwards.

6.4 Special precautions for storage
Store below 25°C. Do not freeze. Keep the vial/ampoule in the outer carton in order to protect from light.

6.5 Nature and contents of container
Fibrovein 3.0%: 2ml ampoules and 5ml vials
Fibrovein 1.0%: 2ml ampoules
Fibrovein 0.5%: 2ml ampoules
Fibrovein 0.2%: 5ml vials.

2ml ampoule (Type 1 glass). Pack size 5 ampoules.
5ml vial (Type 1 glass) with a stopper (chlorobutyl) and aluminium seal with flip-off cap (polypropylene). Pack size of 5 vials.

6.6 Special precautions for disposal
The foam must be prepared just before use and administered by a physician appropriately trained in the correct generation and administration of foam.

Strict aseptic technique must be maintained while manufacturing the foam.

Manufacture of foam using the Tessari technique
To create the foam 1ml of liquid sclerosant is drawn into a sterile syringe and 3ml or 4ml of sterile air is drawn into another sterile syringe. The air is drawn through a 0.2 μm filter to ensure it is sterile. The syringes are then connected using a sterile three way tap/valve (Fig. 1).

The sclerosant/air mixture is then forced back and forth from one syringe to the other through the 3-way valve approximately 20 times to produce a smooth, consistent foam (Fig. 2&3).

The syringe containing the foam, is then removed and the vein is injected immediately (Fig. 4).

The sclerosant foam must be used within sixty seconds of production. After sixty seconds any remaining foam should be discarded. More foam should be prepared if required.

The quality of the foam should be checked before administration. It should appear homogeneous with no large bubbles visible to the naked eye.
The quality of foam depends on specific criteria:

1. The product concentration: Foam can only be prepared with concentrations of 1 to 3% sodium tetradecyl sulfate.

2. The proportion of liquid to air: Usually, this proportion is 1 volume of liquid for 3 volumes of air.

3. Number of backwards and forwards passes: The physician should follow precisely the number of movements defined for each technique.

4. Macroscopic consistency of the foam: The quality of the foam should be checked outside of the syringe before administration. The foam should be homogenous, soft and cohesive with no visible large bubbles. If large bubbles are visible, the foam should be thrown away and a new foam prepared.

5. The total time of preparation of the foam: The preparation should take around 10 seconds from the first to the last backwards and forwards movement.

6. The maximum time between preparation and injection: The sclerosant foam must be used within sixty seconds of production. After sixty seconds, any remaining foam should be discarded. More foam should be prepared if required.

**Disposal**

There are no special requirements for disposal.

**7 MEDICINE SCHEDULE**

Prescription medicine
8 SPONSOR
New Zealand Medical and Scientific Ltd
PO Box 24-138
Royal Oak
AUCLAND
Ph: 09 259 4062
Fax: 09 259 4067

9 DATE OF FIRST APPROVAL
31 December 1969

10 DATE OF REVISION OF THE TEXT
8 November 2021
## SUMMARY TABLE OF CHANGES

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<td>4.4</td>
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