Established lymphoedema is associated with significant morbidity. Suboptimal management of people with lymphoedema can result in avoidable health complications, such as cellulitis with prolonged hospitalisation and poorer patient outcomes, as shown in Figure 1.

What is lymphoedema?
Lymphoedema is a chronic swelling of a limb or body region. It occurs when there is an imbalance in the transportation and/or production of fluid in the interstitial tissues resulting in the accumulation of extracellular fluid. It may arise as a result of a congenital malformation of the lymphatic system (primary lymphoedema) or due to damage and trauma to, or interference with, the lymphatic vessels or nodes (secondary lymphoedema). The main causes of both types of lymphoedema are listed in Box 1. Although lymphoedema starts as a fluid-focused condition, in some patients it can progress to the accumulation of adipose and fibrotic tissue in the affected region. This excess fluid and/or tissue reaction is associated with inflammatory mediators, resulting in altered local immunity.

In developed countries, the most common cause of lymphoedema is cancer related (Figure 2). Approximately 15.5% of cancer survivors develop lymphoedema. This varies depending on the type of cancer, surgical intervention and adjuvant therapies. Other causes of secondary lymphoedema include trauma, infection, surgery and vascular conditions. A newly emerging and increasingly frequent cause of lymphoedema is obesity. The prevalence of this is less clear and largely undocumented. Primary lymphoedema makes up the remainder of the

KEY POINTS
- GPs will see both patients at risk of lymphoedema (for whom prevention is crucial) and those who already have lymphoedema.
- Early detection of lymphoedema and intervention significantly improve patient outcomes, and early referral of patients to a qualified lymphoedema practitioner for treatment will achieve the best outcomes.
- Compression is the mainstay of lymphoedema management but must be carefully prescribed to prevent adverse effects.
- Lymphatic insufficiency may be an underlying cause for patients presenting with limb cellulitis; control of the swelling can reduce the frequency of cellulitis episodes.
- GPs can facilitate prompt antibiotic therapy for lymphoedema-related cellulitis, which can present differentially from classic cellulitis and usually requires a longer course of antibiotics.
- Simple strategies can become part of an everyday routine for patients with support from their GP and lymphoedema practitioner.
cases. There may be several contributing causes present in the one patient, called mixed lymphoedema.

Prevalence studies in the UK have reported 0.13 to 2.0 per 1000 people living with lymphoedema. There are no equivalent Australian data, but conservative estimates suggest that there might be at least 35,000 people living with lymphoedema in the country.

### Presentation

Lymphoedema should be considered in patients who have swelling of a body part that has persisted for more than three months (or earlier if it is associated with surgical removal of lymph nodes). It may be associated with heaviness and aching, and in the initial stages, diurnal fluctuation.

### Diagnosis

The diagnosis of lymphoedema is a clinical one based on history and examination. It should be holistic in its approach and encompass any physical, functional or psychosocial issues a person may be facing.

Steps in the diagnosis of lymphoedema, including the differential diagnoses to consider, are summarised in the Flowchart. Further medical review and investigations are needed to exclude other causes of swelling when the following red flags are present:

- sudden acute onset of swelling
- pain as a main presenting complaint
- skin colour changes
- prominent collateral veins in the region of swelling
- lumps, sores, ulcers
- paraesthesia, other neurological signs
- history of unmonitored cancer

Important signs to look for in the clinical examination are the Stemmer’s sign.

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**Figure 1.** The lymphoedema cycle.

Reproduced with permission from the Lymphoedema Action Alliance.

**Figure 2.** Breast cancer-related secondary lymphoedema.

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**1. MAIN CAUSES OF LYMPHOEDEMA**

**Primary lymphoedema**

- Congenital abnormalities or malformation of the lymphatic system (presenting under 2 years of age)
- Late onset due to underlying malformation (presenting over 2 years of age)
- Syndromal conditions: lymphoedema-distichiasis, Klippel-Trenaunay syndrome and Prader-Willi syndrome, among others

**Secondary lymphoedema**

- Trauma and tissue damage: lymph node excision, radiotherapy, burns, varicose vein surgery, wounds
- Malignant disease: lymph node metastases, infiltrative carcinoma, lymphoma, pressure from large tumours
- Venous disease: chronic venous insufficiency/ulceration, post-thrombotic syndrome
- Infection: cellulitis/erysipelas, lymphadenitis, filariasis
- Inflammation: rheumatoid arthritis, dermatitis, psoriasis, sarcoidosis
- Medications: including calcium channel blockers, corticosteroids, NSAIDs, pregabalin, taxanes
- Other: obesity, lipoedema, dependency oedema, paralysis

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and the pitting sign (Figures 3a and b). The Stemmer’s sign is the inability to pinch the skin at the base of the second toe or middle finger, which is considered pathognomonic of lymphoedema. The pitting sign is persistent indentation of the skin following firm thumb pressure for up to 30 seconds over the oedematous area, and is indicative of subcutaneous oedema. Lymphoedema can present many years after the triggering event, as shown in the following case.

Abbreviations: BMI = body mass index; CRP = C-reactive protein; DVT = deep vein thrombosis; ESR = erythrocyte sedimentation rate; EUC = electrolytes, urea and creatinine; FBC = full blood count; HbA1c = glycated haemoglobin; LFT = liver function test; OGTT = oral glucose tolerance test; TFT = thyroid function test.

*Modified from Lymphoedema: Guide for Diagnosis and Management in General Practice, with permission from the Lymphoedema Association of Victoria and General Practice – Victoria.
A case of secondary lymphoedema

A 53-year-old farmer presents with swelling in his right leg (Figure 4). His leg has been progressively becoming more swollen and heavy, and now he can no longer fit into his right work boot. On history, he reveals that he had a melanoma removed from his right thigh 10 years ago. He remembers having had surgery to his groin for lymph node removal. He has a large fibrotic scar on his right thigh and groin.

Key messages

• Lymphoedema can present many years after the triggering event.
• This patient has been at a high risk for lymphoedema ever since he had groin lymph node surgery. He would have benefited from regular surveillance and education on risk factors to raise awareness of the early signs and symptoms.
• It is important to exclude DVT and any recurrence of melanoma.

Investigations

Several investigations can be useful tools in the process of lymphoedema diagnosis, but not all are necessary.

• Lymphoscintigraphy is a nuclear medicine procedure used to demonstrate lymphatic morphology and function. It is particularly useful where the cause of swelling is unclear (see the case below).
• New imaging techniques are rapidly evolving, such as indocyanine green fluoroscopy (ICG), which can be used to map the superficial lymphatics. This can be useful in difficult clinical cases.
• Ultrasound, Doppler ultrasound, MRI and CT can be useful to exclude differential diagnoses and also to assess for local area tissue changes.
• Bioimpedance spectroscopy (BIS) using low frequencies can measure extracellular fluid and can indicate differences between normal and lymphoedematous limbs. It is useful in monitoring patients at high risk of lymphoedema to detect early subclinical changes.
A case of primary lymphoedema in an adolescent

A 14-year-old girl presents with a swollen right foot and ankle (Figure 5). The swelling has been present for six months, and persisted after a minor sprain. Her GP has already excluded a DVT and other differential diagnoses, but is not sure what else can be causing this problem.

Key messages
- Persistent swelling of more than three months needs a diagnosis, and primary lymphoedema should be considered.
- In this instance, lymphoscintigraphy is indicated to help with the diagnosis of primary lymphoedema (Figure 6).
- Early referral is recommended in such cases for treatment in a multidisciplinary paediatric lymphoedema clinic (if available) or alternatively to a trained lymphoedema practitioner to optimise swelling and prevent infections.

Treatment

Once the diagnosis of lymphoedema is confirmed, the patient should be referred to a lymphoedema practitioner who can provide a range of treatments, help in developing skills for lifelong self-management and prescribe suitable compression garments.

Best practice lymphoedema treatment is a multimodal regimen (Figure 7), including the components listed below.
- Skin care involves cleansing and moisturising the skin to provide a barrier and encourage lymphatic flow. It includes regular checks for fungal infections and cellulitis.
- Exercise encourages lymph flow and is important for weight management. Hydrotherapy is a recommended form of exercise due to the incorporation of hydrostatic pressure.
- Compression is used initially in the form of bandaging to reduce swelling. Compression garments can then be used to maintain this reduction (see below).
- Manual lymphatic drainage is a specific decongestive massage technique to help move fluid away from congested areas.
- Education and support is central to these principles and to GP management. Engagement of patients in their self-management regimen is crucial.

The ability of compression garments (e.g. sleeves, stockings, gloves) to maintain the gains made by intensive treatment depends on the appropriate choice of garment and the adherence of the person to the wearing regimen. For this reason, the garment should be fitted by a qualified professional, such as a trained lymphoedema practitioner or garment provider, to ensure the correct pressure and gradient distal to proximal. Numerous types of compression are available in a range of sizes, fabrics and methods of application with varying levels of compression (Table 1 and Figure 8).

Contraindications for compression include severe arterial insufficiency, uncontrolled heart failure or severe peripheral neuropathy (e.g. diabetes). Before commencing lower leg compression, vascular assessment is recommended to exclude any underlying arterial insufficiency.

Figure 6. Lymphoscintigraphy of a 14-year-old patient with primary lymphoedema of the right leg. Six hours after radioactive colloid injection between the toes, there is no transit of the colloid via the lymphatic vessels to the lymph nodes in the right groin, and reduced transit on the left.

Figure 7. Principles of lymphoedema treatment.
ankle-brachial pressure index (ABPI) can be used if there are concerns about arterial insufficiency.

Newer compression treatments are constantly emerging. These include sequential intermittent pneumatic compression, low level laser, negative pressure and lymph taping.

The GP has a role in the long-term chronic care management plan of patients with lymphoedema (Table 2). Simple treatments and discussions with patients can make a significant difference to their outcomes. Reinforcing good skin care, travel precautions, antibiotic use, exercise and massage are all part of the GP’s role. GPs can facilitate the creation of a good multidisciplinary team that could consist of a lymphoedema practitioner, podiatrist, dietitian and psychologist.

Where to find a lymphoedema practitioner?
Lymphoedema practitioners are allied health professionals (e.g. physiotherapists, occupational therapists, nurses) and remedial massage therapists who have completed Australasian Lymphology Association (ALA) accredited lymphoedema management training. They can be found in some larger public hospitals that

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**TABLE 1. LEVELS OF COMPRESSION**

<table>
<thead>
<tr>
<th>Class of compression</th>
<th>Pressure</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preventive (e.g. flight socks)</td>
<td>15–20 mmHg</td>
<td>Prevention; inappropriate for lymphoedema</td>
</tr>
<tr>
<td>Class I</td>
<td>20–30 mmHg</td>
<td>Vascular or early lymphoedema</td>
</tr>
<tr>
<td>Class II</td>
<td>30–40 mmHg</td>
<td>Established lymphoedema, particularly arm</td>
</tr>
<tr>
<td>Class III</td>
<td>40–50 mmHg</td>
<td>Established lymphoedema, particularly leg</td>
</tr>
<tr>
<td>Class IV</td>
<td>50–60 mmHg</td>
<td>Severe lymphoedema</td>
</tr>
</tbody>
</table>

* This is a guide only, and clinical factors and patient tolerability need to be considered.

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**TABLE 2. EXAMPLES OF GP CHRONIC MANAGEMENT PLANS FOR PATIENTS WITH OR AT RISK OF LYMPHOEDEMA**

<table>
<thead>
<tr>
<th>Condition</th>
<th>Goals</th>
<th>Treatment</th>
<th>Arrangements/referrals</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Example for patients with established lymphoedema</strong></td>
<td><strong>• Reduction in size of limb and better control of swelling</strong>&lt;br&gt;<strong>• Avoidance (or early detection) of episodes of cellulitis</strong></td>
<td><strong>• Regular review with lymphoedema practitioner – weekly to monthly sessions as required</strong>&lt;br&gt;<strong>• Wearing regular compression garments</strong>&lt;br&gt;<strong>• Six-monthly review with GP</strong>&lt;br&gt;<strong>• Travel with antibiotics for early initiation if required</strong>&lt;br&gt;<strong>• Self management:</strong>&lt;br&gt;  – regular exercise such as swimming&lt;br&gt;  – self-massage&lt;br&gt;  – skin care including checking for infections&lt;br&gt;  – weight management</td>
<td><strong>• GP (six-monthly) review</strong>&lt;br&gt;<strong>• Specialist review as needed</strong>&lt;br&gt;<strong>• Lymphoedema practitioner (weekly to monthly; six-monthly once stable)</strong>&lt;br&gt;<strong>• Patient self-care daily</strong>&lt;br&gt;<strong>• Podiatrist as needed</strong>&lt;br&gt;<strong>• Dietitian as needed</strong>&lt;br&gt;<strong>• Exercise physiologist as needed</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Examples for patients at risk of lymphoedema</strong></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast cancer</td>
<td><strong>• Early detection of any recurrence of cancer</strong></td>
<td><strong>• Regular follow up with oncologist, surgeon and other specialists</strong></td>
<td><strong>• Annual specialist review</strong></td>
</tr>
<tr>
<td>Axillary dissection and radiotherapy in context of breast cancer</td>
<td><strong>• Early detection and treatment of lymphoedema</strong></td>
<td><strong>• Monitoring for signs and symptoms of lymphoedema (swelling, tightness, heaviness)</strong>&lt;br&gt;<strong>• Bioimpedance and circumferential measurements at baseline and follow up</strong></td>
<td><strong>• Six-monthly check with GP and referral to lymphoedema practitioner as appropriate</strong></td>
</tr>
</tbody>
</table>

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have a lymphoedema service available, and there are also several private practitioners. The ALA website has an online register of qualified lymphoedema practitioners (http://www.lymphoedema.org.au/the-register/find-a-practitioner).8

### Lymphoedema-related cellulitis

Cellulitis presents with an acute spreading inflammation of the skin and subcutaneous tissues characterised by pain, swelling, warmth and erythema. Its onset can be highly variable in lymphoedema. Systemic symptoms often precede these signs. The decision to start antibiotics should not depend on blood test results, but rather on the patient’s clinical presentation. Prompt treatment is essential to avoid further damage to the lymphatics, which, in turn, may predispose to recurrent cellulitis.

Nonpurulent cellulitis is most likely caused by group A streptococci, which remain 100% sensitive to penicillin. Purulent cellulitis is more likely to be caused by *Staphylococcus aureus*, which has a significant resistance profile.9 If present any pus should be swabbed; however, empirical antibiotics should be commenced in the interim (Table 3).

Antibiotics should be continued for 14 days after a definite clinical response is noted. Drawing a line around cellulitic areas and reviewing 48 hours after starting antibiotics are important to establish efficacy of treatment and appropriateness of referral to hospital. Patients should be encouraged to return to wearing compression garments as soon as there is reduction in the acute symptoms of cellulitis.

The case of an elderly patient with mixed lymphoedema, a combination of secondary and dependent lymphoedema with cellulitis, is described below.

### A case of dependency oedema, ulcers and recurrent cellulitis

An 82-year-old woman presents with her niece. She has swelling in her legs bilaterally. Her right leg is significantly more swollen (Figure 9). It is red, hot and tender across her right shin. She has had a fever for 48 hours. She has a history of varicose vein surgery and hysterectomy when in her 40s. She prefers to sleep in a recliner chair.

### Key messages

- This patient has a combination of secondary oedema and dependent oedema, which has placed her at increased risk of cellulitis.
- Her cellulitis needs to be treated urgently.
- The diagnoses of congestive cardiac failure and DVT need to be excluded.
- Concurrent treatment of her comorbidities and lymphoedema can help to improve her quality of life. Treating the swelling and addressing the skin integrity will reduce her risk of recurrent cellulitis, and assist in mobilisation.

### Dispelling the myths

There are many myths in the lymphoedema literature regarding safety for patients. Some of these include: avoiding repetitive exercise; avoiding blood pressure measurements, venepuncture or injections in the affected limb; flying being a precipitating factor to lymphoedema; and avoiding all use of diuretics. Below the authors attempt to set the record straight.

- Exercise and usual activities should be encouraged in patients with lymphoedema. Recent Australian

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### Table 3. Recommended Antibiotic Treatment for Lymphoedema-Related Cellulitis*

<table>
<thead>
<tr>
<th>Condition</th>
<th>Antibiotic</th>
<th>Hypersensitivity to penicillin</th>
<th>Allergic to penicillin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute cellulitis – treat for minimum of three weeks</td>
<td>Phenoxymethylpenicillin 500 mg every six hours For suspected staphylococci (pus or folliculitis) – dicloxacillin or flucloxacillin 500 mg every six hours</td>
<td>Cephalexin 500 mg every six hours</td>
<td>Clindamycin 300 mg every eight hours</td>
</tr>
<tr>
<td>Recurrent cellulitis – consider prophylactic prescription if more than two episodes per year</td>
<td>Phenoxymethylpenicillin 500 mg daily or 250 mg twice daily</td>
<td>Cephalexin 500 mg daily or 250 mg twice daily</td>
<td>Erythromycin 250 mg daily</td>
</tr>
</tbody>
</table>

* Refer to the Australasian Lymphology Association Antibiotic Consensus Guidelines: Management of Cellulitis in Lymphoedema for more details.9
TABLE 4. RECOMMENDATIONS FOR MONITORING THOSE AT RISK OF LYMPHOEDEMA

<table>
<thead>
<tr>
<th>At-risk condition</th>
<th>Recommendation for monitoring</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cancer (e.g., breast, melanoma, gynaecological, prostate, sarcoma, head and neck)</td>
<td>Three-monthly intervals for the first two years Six- to 12-monthly review thereafter Education to facilitate self-monitoring</td>
</tr>
<tr>
<td>Cellulitis</td>
<td>Review at resolution of infection Education to facilitate self-monitoring</td>
</tr>
<tr>
<td>Venous surgeries (e.g., for varicose veins)</td>
<td>Education to facilitate self-monitoring</td>
</tr>
<tr>
<td>Deep vein thrombosis</td>
<td>Education to facilitate self-monitoring</td>
</tr>
<tr>
<td>Hereditary lymphoedema</td>
<td>Review of siblings and progeny regularly or as clinically relevant Genetic counselling as appropriate</td>
</tr>
</tbody>
</table>

research has shown that exercise may be preventive for those at risk of breast cancer-related lymphoedema.\(^{10}\)

- Blood pressure should ideally be measured in an unaffected arm. However, there is no evidence of exacerbation from taking single blood pressure readings in a lymphoedema-affected arm. Taking repetitive blood pressure readings (e.g., during surgery or 24-hour blood pressure monitoring) should be discouraged in the affected limb.

- There is no evidence that venepuncture triggers lymphoedema when performed using aseptic techniques by experienced health professionals; however, when possible it should be performed on the unaffected arm.

- Injections or vaccinations (e.g., tetanus vaccination) can induce an inflammatory reaction and, therefore, should be avoided in the affected limb.

- Flying can be a trigger for exacerbations in those with established lymphoedema. There is little evidence of this in at-risk populations. Prophylactic garments are often worn to prevent lymphoedema, but it is vital that they are appropriately fitted and trialled before travel. An ill-fitting garment can do more harm than good due to a possible tourniquet effect or poor pressure gradient.

- Diuretics do not remove lymphatic fluid in subcutaneous tissue. However, their use may be indicated where there is coexistent congestive cardiac failure. Diuretics can be used concurrently with compression in this instance.

- Bilateral cellulitis is extremely rare. Symmetrical erythema in the legs is most likely due to stasis dermatitis, particularly in the absence of systemic features.

Prevention

Effective identification of patients at risk of lymphoedema relies on an awareness of the causes of lymphoedema (see Box 1).\(^{1}\) The GP has an important role in monitoring for the early detection of lymphoedema in high-risk groups, especially after cancer therapy resulting in lymph node excision (Table 4). This monitoring should become part of a GP’s chronic care management plan.

Surgical intervention

New treatments are available for the surgical management of lymphoedema. These are available to selected patients in specialist units. Liposuction is an option for severe lymphoedema that is no longer responsive to conservative treatment. However, patients who have liposuction are still required to wear compression garments. Super-microsurgical techniques based on new imaging are an emerging area of plastic surgery. These include lymphovenous anastomosis and vascularised lymph node transfer. These specialised surgical techniques should be embedded in an integrated lymphoedema service model.\(^{11}\)

Conclusion

Lymphoedema detection and treatments are constantly evolving with new techniques in compression and surgery. Early and accurate diagnosis of lymphoedema by the GP remains the key to accessing appropriate treatment. GP co-ordination of a multidisciplinary team will ensure the best outcomes for the patient with, or at risk of, lymphoedema. A list of useful online resources for GPs is given in Box 2.

References

A list of references is included in the online version of this article (www.medicinetoday.com.au).

COMPETING INTERESTS: None.
Lymphoedema
Breaking the swollen cycle

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References


