Surfer wipe out by predator fish
Kerkhoffs, G.M.M.J.; op den Akker, J.W.; Hammacher, E.R.

Published in:
British journal of sports medicine

DOI:
10.1136/bjsm.37.6.537

Citation for published version (APA):

General rights
It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations
If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: http://uba.uva.nl/en/contact, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible.
Surfer wipe out by predator fish

G M M J Kerkhoffs, J W op den Akker and E R Hammacher

Br. J. Sports Med. 2003;37;537-539
doi:10.1136/bjsm.37.6.537

Updated information and services can be found at:
http://bjsm.bmjjournals.com/cgi/content/full/37/6/537

These include:

References

1 online articles that cite this article can be accessed at:
http://bjsm.bmjjournals.com/cgi/content/full/37/6/537#otherarticles

Rapid responses

You can respond to this article at:
http://bjsm.bmjjournals.com/cgi/eletter-submit/37/6/537

Email alerting service

Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article

Topic collections

Articles on similar topics can be found in the following collections

Orthopedic and Trauma Surgery (343 articles)
Injury (834 articles)
Sports Medicine (1331 articles)

Notes

To order reprints of this article go to:
http://www.bmjjournals.com/cgi/reprintform

To subscribe to British Journal of Sports Medicine go to:
http://www.bmjjournals.com/subscriptions/
A 17 year old female athlete presented with Achilles tendinopathy. A protocol of water and land based rehabilitation was designed to achieve non-weight bearing and pain free activity, so that she could rapidly return to her event (400 m hurdles). After three weeks, she returned to regular training, and after a further three weeks she successfully competed in a 400 m hurdle event. She has been able to compete at national level symptom free for the last 18 months.

The Achilles tendon is the largest and strongest tendon in the human body and experiences high loads in all sporting activities. It has been shown to have a high capacity to withstand tensional forces.1 Chronic painful conditions located in the Achilles tendon are relatively common, especially in runners.2 The cause of such conditions is not known, but they are often associated with repetitive loading and overuse.1 Achilles tendinopathy is difficult to treat.2 Most recommend a conservative regimen as the initial strategy.4–6 Non-surgical treatments include a combination of rest, anti-inflammatory drugs, correction of malalignments, and stretching and strengthening exercises.

There is sparse scientific evidence that these treatments are successful, and surgical treatment is required in about 25% of patients.2 In a recent prospective study, treatment with heavy load, eccentric calf muscle training showed very promising results and may reduce the need for surgical treatment of tendinopathy located in the midportion of the Achilles tendon.1 The short term results of surgery are often very good, but in the few studies with long term follow up there are signs of deterioration with time.1 In addition, over the last two decades there has been a considerable increase in the incidence of Achilles tendon rupture.6

Many athletic trainers propose an active land based rehabilitation training programme for Achilles tendinopathy, which includes strength and stretching exercises and balance exercises on unstable boards. However, many physiotherapists and aquatic exercise specialists have been using water to treat conditions of the lower extremity. They maintain that water is the ideal environment for the rehabilitation of injuries for which weight bearing needs to be limited. Range of motion (ROM), strength, and functional activities performed in deep water are used to prepare the limb for weight bearing. The combination of buoyancy induced skeletal unloading and muscular relaxation can enhance ROM and flexibility. Land based tasks should be initiated before the water programme is finished to facilitate a smooth transition from pool to land.8 Over the last two decades, many athletes have routinely exercised in the pool when recovering from injury.

This study focuses on a specific rehabilitation programme (combination of aquatic and land based exercises) prescribed for a young female runner with early stage Achilles tendinopathy.

CASE REPORT

A 17 year old female athlete presented with gradually increasing pain located in the Achilles tendon area of her left foot. The clinical diagnosis was Achilles tendinopathy. She was prescribed anti-inflammatory drugs to alleviate the mildly painful symptoms. The symptoms decreased and she was able to compete in the preliminary rounds of the 400 m hurdles and gained a place in the finals to take place six weeks later. The following day, the painful symptoms in the Achilles tendon increased rapidly and spread around the posterior ankle area, so that even walking was painful. More specifically, the point tender pain and swelling was over the Achilles tendon (the middle section proximal to the calcaneal insertion). The tendon was sore on rising in the morning and stiff when she started to run. In addition, crepitus was present. She could not stand on tip toe, to go up or down stairs. On examination, movement from the neutral position to dorsiflexion (supine) was painful. After a clinical examination, she was advised to stop training. The athletic training team (an orthopaedic surgeon, a physiotherapist, and an athletic trainer) designed an aquatic rehabilitation programme to achieve early non-weight bearing and pain free activity.

REHABILITATION PROGRAMME

The goals were gait normalisation, to increase pain free dorsiflexion from 0° to 20°, proprioception ability, calf muscle strength, and strength throughout the lower extremity, and to maintain cardiovascular endurance (table 1).

To maintain her cardiovascular condition, the athlete followed a daily swimming programme. The ankle joint was taped for protection and to keep the heel in a neutral position.

The early phase (1st week) of rehabilitation consisted nearly exclusively of exercises in the pool. Passive stretching exercises in the pool to increase ROM of the ankle joint (dorsiflexion and plantar flexion) paralleled stretching exercises on the land. Mild static balance and isometric exercises (after the 5th session) were performed in the water. Gait normalisation activities, including walking at different stride lengths in different directions at various depths and different speeds in the pool, were performed. Single-leg balance and wobble board exercises in the water were included. Strengthening exercises in this phase were primarily open kinetic chain because of the pain and substitution that occurred with closed kinetic chain activities. Dynamic strength exercises for the other lower limb muscle groups were also performed with specific equipment in the pool.

The intermediate phase focused on increasing gait endurance. Mild dynamic exercises for the gastrocnemius and soleus muscles were initiated in the pool in gradually decreasing depths of water. Full passive ROM was achieved. During this week, the pain decreased considerably. Land based exercises progressed to full weight bearing activities as tolerated, with the focus on stress to the Achilles tendon.
During the third week (the late rehabilitation phase), the time spent on land based exercises was increased and the amount of time spent in the pool was decreased. Impact exercises were initiated in the pool and progressed to land, first on grass and then to the traditional track and field surface. Passive and active stretching for the ankle and calf was performed. Over the next three weeks, the athlete started a progressive training programme in order to compete in the 400 m hurdle event. She won the race, achieving a new national record (61 seconds). She has had no further symptoms.

**DISCUSSION**

On initial presentation, the athlete was treated with anti-inflammatory drugs. Although the symptoms decreased, the Achilles tendinopathy was not resolved. Kader et al. maintain that Achilles tendinitis is a degenerative, not an inflammatory, condition. Although the present case was an early stage of Achilles tendinopathy, it also was not an inflammatory condition.

In our case study, a water based exercise programme was devised for the Achilles tendinopathy because it limits weight bearing using buoyancy. During the rehabilitation period, it is very important to protect the muscle-tendon unit from excessive torque or damaging vibrational forces. Premature loading of a damaged Achilles tendon can cause a number of adverse chemical, metabolic, and vascular changes. Rehabilitation progresses consistently when inflammation is kept to a minimum. Therefore, the aquatic rehabilitation programme provided the early initiation of specific exercises (strengthening, stretching, and balance exercises of the ankle joint) to prevent atrophy, motion loss, and abnormal movement pattern development.

The cardiovascular swimming programme had two positive effects. Firstly, it maintained the athlete’s cardiovascular condition, which is also an important aspect of early intervention. Secondly, as supported by many studies, it had positive effects on the healing tissues. These effects include increased blood flow and neurological stimulation, with minimisation of adjacent tissue weakness and adverse psychological effect. However, it is important that the athlete works within the constraints of his/her limitations.

The wobble board exercises are good for stimulating proprioception. Water is an ideal environment for proprioceptive training because of its viscosity. Viscosity provides a slow motion, three dimensional resistive environment which facilitates proprioceptive feedback through functional movement patterns. This proprioceptive activity enables the athlete to incorporate safely advanced levels of dynamic stabilisation earlier in the rehabilitation schedule by decreasing both the demands of deceleration and joint excursion.

During the early phase (first week), static and isometric exercises were performed. Dynamic activity was contra-indicated because of the joint compression, vibration, and torsional forces that accompany these techniques, resulting in increased pain and inflammation. The athlete in this study was able to perform mild dynamic exercises (second week) earlier in the pool because of minimisation of some of these forces in this environment. Concentric and eccentric loading in single and then multiple planes (late phase) encouraged normal gastrocnemius and soleus function.

**CONCLUSION**

Aquatic rehabilitative exercises for Achilles tendinopathy provide an opportunity for the athlete to train in a gravity minimised environment while being immersed in a resistive medium. The unloading is critical as rehabilitation time can be minimised because safe and functional rehabilitation can

<table>
<thead>
<tr>
<th>Rehabilitation phase</th>
<th>Rehabilitation goals</th>
<th>Rehabilitation programme</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early phase (1st week)</td>
<td>Maintain cardiovascular endurance</td>
<td>Pool</td>
</tr>
<tr>
<td>Sessions 1–4 (40 min)</td>
<td>Dorsiplantar flexion</td>
<td>Swimming interval training</td>
</tr>
<tr>
<td>Sessions 5–7 (50 min)</td>
<td>Gait normalisation</td>
<td>Pool</td>
</tr>
<tr>
<td></td>
<td>Proprioception ability</td>
<td>Walking (in different stride length, depths, speeds)</td>
</tr>
<tr>
<td>Intermediate phase (2nd week)</td>
<td>Maintain cardiovascular endurance</td>
<td>Pool</td>
</tr>
<tr>
<td>Sessions 8–14 (60 min)</td>
<td>Increase gait endurance</td>
<td>Swimming interval training</td>
</tr>
<tr>
<td>Late phase (3rd week)</td>
<td>Maintain cardiovascular endurance</td>
<td>Pool</td>
</tr>
<tr>
<td>Sessions 15–21 (60–80 min)</td>
<td>More land based programme</td>
<td>Swimming interval training</td>
</tr>
<tr>
<td>Return to sport phase (4th–6th week) (60–90 min)</td>
<td>Back to training</td>
<td>Swimming interval training, impact exercises, land exercises in grass and gradually in track and field flooring, passive and active ankle calf stretching</td>
</tr>
</tbody>
</table>
be started immediately. This can minimise the injury and reduce recovery time.

The aquatic and land based programme reported here offered an effective rehabilitation protocol for the female athlete. However, a case study does not allow generalisation of the conclusions. Further investigation is needed to estimate the effectiveness of this early intervention in achieving rapid return to unlimited activity.

**Authors’ affiliations**
A G Beneka, P C Malliou, G Benekas, Democritus University of Thrace, Komotini, Greece

Correspondence to: Assistant Professor Malliou, TEFAA, 7th Km Komotini Xanthi, Komotini 69100, Greece; malliou2002@yahoo.gr

Accepted 5 November 2002

**REFERENCES**
The wound was dressed with wet gauze and flushed twice daily. The patient was kept in hospital for two days, and immediate improvement was seen. There were no complications. Four weeks after the accident, the patient had resumed his professional activities.

**DISCUSSION**

Needlefish belong to the Beloniform order which is composed of two families, the Belonids and the Hemiramphids. These are found in (sub) tropical waters of all oceans. The fish are long and slender, ranging in length from 0.5 m to more than 1.5 m. The long, narrow “beak” is comprised of jaws filled with sharp, little teeth. The colour, bluish green on top and silver on the bottom, is adapted for surface dwelling. Needlefish are surface carnivore predators.

Needlefish injury should not be confused with a stingray or catfish injury; all three fish produce penetrating injuries. However, stingray and catfish are bottom dwellers. Injury by a stingray typically occurs when it is stepped on or handled. Catfish injuries generally occur only while handling the fish. Both stingray and catfish cause injury with envenomation, which should be treated by immersion in hot water. Management of the wound is similar for all three.

A needlefish injury should be treated like a stab wound. The small diameter of the fish’s snout allows it to penetrate between bony structures of the thorax, spinal canal, and skull. Abdominal, ocular, and articular injuries have also been described. After penetration of the skin, the beak can break into several fragments and inflict severe damage at a distance from the entry point. Standard radiographs are recommended to determine the presence of retained beak. The appearance of two semiparallel lines of opacity is typical of a needlefish beak. Radiographic and clinical evaluation lead to diagnosis and early surgical treatment.

There are no prospective studies providing evidence for optimal antibiotic selection for marine acquired infections or prophylaxis. Immunocompetent patients do not always require prophylaxis. A surgical debridement is the mainstay of treatment. Tetanus prophylaxis is required if not up to date. Wounds generally should be left to heal by secondary intention.

**Authors’ affiliations**

G M M J Kerkhoffs, Academic Medical Center, Amsterdam, the Netherlands

G M M J Kerkhoffs, J W op den Akker, E R Hammacher, Department of General Surgery, Sint Antonius Hospital, Nieuwegein, the Netherlands

Correspondence to: Dr Kerkhoffs, Academic Medical Center, Orthopedic Surgery, Meibergdreef 9, Amsterdam 1100 DD, the Netherlands; ginokerkhoffs@hotmail.com

Accepted 5 November 2002

**REFERENCES**


Two cases of suprascapular neuropathy in a family

M Ravindran

Suprascapular entrapment neuropathy is well known in certain athletes, especially volleyball players. A brother and sister presented with right shoulder pain and wasting of the scapular muscles, particularly the infraspinatus. They had played volleyball for over six years and were forced to retire because of disability. Investigations showed involvement of the rhomboid muscles also, suggesting a probable extension of this syndrome to other nerves in the region such as the dorsal scapular nerve.

Suprascapular nerve entrapment with scapular and shoulder pain, associated with weakness and wasting of the spinati muscles, is well documented in sports people, especially volleyball players. Several mechanisms have been proposed to explain the symptoms, but excessive traction or stretching of the nerve is the most plausible pathomechanism. When this condition develops with an identical clinical profile in two siblings, both volleyball players of some repute, then some anatomical peculiarity seems logical. A brother and sister presented with weakness and wasting of the right scapular muscles and the details are reported here.

CASE REPORTS

Case 1
A 35 year old right handed male volleyball player developed right shoulder and scapular pain after a strenuous match in 1986. He was forced to rest for several weeks but the pain remained unabated. Before the onset of pain, he played volleyball regularly for almost seven years, often representing the state.

Within 2–3 months, wasting of the scapular muscles was observed. He was briefly investigated by radiography and electrophysiology, when neurogenic changes were observed in the right infraspinatus muscle. A year later, he had surgical decompression of the right suprascapular nerve at two levels: the suprascapular notch and the spinoglenoid notch. The surgical notes did not indicate the presence of any nerve atrophy, tight ligaments, or ganglions. However, he continued to deteriorate with persistent pain and progressive wasting, even after decompression, while he continued to play volleyball. He had to retire from active sport around 1990. Since then the condition has stabilised and the pain lessened. However, the pain recurred when he became active again as a volleyball coach. He has no history of right shoulder injury or dislocation.

Physical examination showed an athletic man with no systemic disease or skeletal deformities. There was a surgical scar extending from the right scapula to the upper arm.

All cranial nerves were intact; no abnormality of the facial muscles, sternomastoids, or trapezius was found. He had considerable wasting of the right infraspinatus muscle with weakness of external rotation of the right arm (fig 1). Right shoulder abduction was weak from initiation to 30°. He had mild winging of the right scapula but the power of the serratus anterior was near normal. There was no weakness or wasting of the right deltoid muscle. Other muscles were normal. No muscle fasciculations or myotonia was found.

Deep tendon reflexes were normal and symmetrical in both upper and lower limbs, and plantar responses were flexor bilaterally. All sensations were intact. The right shoulder showed a full range of passive movements. There was mild tenderness over the right scapular spine and acromioclavicular joint.

Routine haemogram, erythrocyte sedimentation rate, and blood chemistry were normal, including normal muscle enzymes. Collagen screening was negative. A radiograph of the right shoulder and cervical spine was normal.

Figure 1 Wasting of the right infraspinatus muscle, with a surgical scar over the right scapula extending to the arm and mild winging of the scapula.
Electromyography of the right shoulder girdle muscles showed chronic neurogenic changes from the right infraspinati and supraspinati and rhomboid muscles. Normal response was found from the right deltoid, trapezius, and serratus anterior.

Case 2
The 27 year old sister of the patient in case I, a right handed volleyball player of about seven years duration, also developed shoulder and scapular pain on the right side about five years earlier, which worsened after vigorous games or matches. Wasting slowly became evident over the right scapula with some prominence of the bone. Because of persistent pain, she too had to retire from the game about two years earlier, and since then had no further progression.

On examination, there were no skeletal abnormalities, thyromegaly, or lymphadenopathy. Demonstrable abnormality was confined to the right scapular region. There was no weakness of the right trapezius or deltoid muscle. Weakness and wasting of the right infraspinatus with weakness of the supraspinatus and rhomboid muscles was demonstrable. There was mild winging of the right scapula at rest (fig 2), with no further increase on pushing or raising the arm (serratus anterior action). There were no myotonia or muscle fasciculations. She had normal and symmetric deep tendon reflexes and flexor plantar bilaterally. She had no sensory dysfunction or muscle tenderness. Passive movements of the right shoulder showed full range and no tenderness.

Blood chemistry including muscle enzymes, haemogram, erythrocyte sedimentation rate, and thyroid function were normal. Collagen screening was negative.

Radiographic and magnetic resonance imaging of the right shoulder, scapula, and cervical spine were normal.

Electromyographic investigation showed chronic neurogenic changes from the right supraspinati and infraspinati and rhomboid muscles. Trapezius, deltoid, and serratus anterior were normal.

She refused further investigations and surgical exploration.

DISCUSSION
Suprascapular nerve palsy due to entrapment is common in several sports activities, particularly volleyball.1,4,6,8,10,11 Most of the reported cases are in professional volleyball players, and, in some, even asymptomatic atrophy of the infraspinatus muscle has been documented.1

The suprascapular nerve is a motor sensory peripheral nerve (C5) arising as a solitary branch from the superior trunk of the brachial plexus in the posterior triangle of the neck.6 It passes through the suprascapular foramen beneath the transverse scapular ligament, supplies the supraspinatus muscle, and sends sensory fibres to the acromioclavicular joint and subacromial bursa. Then it descends through the spinoglenoid notch to supply the infraspinatus muscle.

A common proximal site of entrapment of the nerve is at the suprascapular foramen, and hypertrophy or calcification of the transverse scapular ligament and other congenital malformations are among the probable causes.7,8 Distally, the nerve may become compressed at the fibro-osseous tunnel formed by the spine of the scapula and the spinoglenoid ligament.5 Suprascapular nerve entrapment with isolated paralysis of the infraspinatus muscle is uncommon, but has been reported in volleyball players.4,6 A larger than normal range of mobility of the shoulder joint with greater traction injury during the game has been postulated as the probable cause.10

Shoulder pain is a common symptom in patients with suprascapular nerve entrapment and is characterised as a dull ache, exacerbated by overhead activities.7 The pain is articular in origin based on the sensory supply of the acromioclavicular joint and surrounding structures by the suprascapular nerve.

The interesting aspect of the cases reported here is the almost identical symptoms in two siblings, brother and sister, both volleyball players of repute. Both played in state teams for six or seven years before they developed shoulder pain and then muscle wasting. Thus the pattern of their disorder suggested a common anatomical peculiarity for the entrapment rather than coincidence. (There were six other siblings, three brothers and three sisters, who were not athletic and did not have this condition.)

There was clear evidence of maximal involvement of the right infraspinatus muscle in both, but the supraspinatus muscle was also involved clinically and electrophysiologically. In addition, there was slight winging of the scapula with demonstrable weakness of the rhomboid muscles. Serratus anterior muscle weakness produces prominent scapular winging, especially on pushing forward, with difficulty in lifting the arm beyond 90°. Weakness of the rhomboids may also cause mild winging of the scapula without interference with arm abduction and elevation.10 The patients reported here had no difficulty or weakness in raising the arm fully at the right shoulder with no further increase in scapular winging during such movements. The possibility of weakness of the rhomboid muscles should also be considered in the differential diagnosis as that too may produce winged scapula, without interfering with arm elevation. This problem was thus highlighted in both patients reported here. The dorsal scapular nerve (C5 nerve to the rhomboid muscles) also arises from the brachial plexus. Therefore there may be injury or entrapment of the C5 root somewhere in its course in these patients. However, this seemed unlikely, with the deltoid and other C5 root muscles being completely spared. Hence, the possibility of entrapment not only of the suprascapular nerve but also of the nerve to the rhomboids, with the suprascapular nerve dominating the clinical picture, arises. No previous report has shown involvement of the rhomboid muscles. Another possibility is an aberrant supply of the rhomboid muscles by the suprascapular nerve, but this
Saphenous nerve injury after fasciotomy for compartment syndrome

D Pyne, A S M Jawad, N Padhia

A case is reported of chronic exertional compartment syndrome treated by fasciotomy. The decompression procedure was complicated by injury to the saphenous nerve. The importance of accurate placement of the postero-medial incision line to avoid saphenous nerve injury is highlighted.

Compartment syndrome refers to muscle ischaemia following small vessel occlusion resulting from an increase in local tissue pressure within a closed fascial compartment (fig 1). The condition may be acute (usually secondary to trauma) or less commonly chronic, secondary to exertion. The definitive treatment is surgical decompression of the compartment by fasciotomy. We describe a case of chronic exertional compartment syndrome (CECS) treated by fasciotomy. The decompression procedure was complicated by injury to the saphenous nerve.

CASE HISTORY
A 24 year old kickboxer, with no previous medical history, presented with pain in both shins and posteromedial aspect of his calf muscles, after an increase in his training level. The pain would slowly resolve over hours on cessation of training but would recur when training was restarted. Physical examination failed to show any abnormal clinical signs. Radiographs of tibiae and fibulae were normal. An isotope bone scan excluded any stress fractures. Intracompartmental pressure measurements were made at rest and during a variety of sprinting exercises. The resting measurements were 25 mm Hg in both deep posterior compartments rising to 35 mm Hg in the left posterior compartment during exercise. The resting pressures were 35 mm Hg in both anterior compartments rising to 45 mm Hg in both compartments during exercise. This confirmed the diagnosis of CECS in both anterior and left deep compartments.1 The patient subsequently underwent bilateral compartmental decompression. His original symptoms resolved, but two weeks after surgery he developed a diffuse burning pain extending from below the knee at the end of the anterior incision, down the medial shin to the region of the arch of the left foot. The pain was persistent. The ankle and knee jerks were both present, and there was no evidence of any weakness. A repeat of the intracompartmental pressure measurements and an isotope bone scan showed no abnormality. Nerve conduction studies found no activity in the left saphenous nerve, indicating that the nerve had been severed. Tinel’s sign could not be elicited. The stump of the nerve was identified by nerve conduction studies, but identification was also aided by the presence of a small neuroma which was hypersensitive to the touch. This

References
stump was blocked with bupivacaine with immediate symptomatic relief.

DISCUSSION

The saphenous nerve is the cutaneous branch of the femoral nerve. Dysfunction of the saphenous nerve causes loss of sensation extending from below the knee down the medial shin to the region of the arch of the foot. Occasionally the infrapatellar branch of the saphenous nerve is damaged as the result of mild trauma or knee operations, producing the syndrome of gonyalgia paraesthetica, which includes sensory loss in a patch of several centimetres diameter below the knee, with intermittent lancinating pains.

Fasciotomy is generally a safe and effective treatment for CECS, with success rates greater than 90%. 2 Complications include wound infection, haematoma, ulceration, deep venous thrombosis, and muscle herniation. 2 Injury to the superficial peroneal nerve is recognised, with fasciotomy for anterior and lateral compartments where the incision line lies halfway between the fibula shaft and the tibial crest, directly over the superficial peroneal nerve.

For superficial and deep compartment decompression, a posteromedial incision 5–6 cm long is used 2 cm posterior to the tibial margin. An incision made anteriorly risks damage to the underlying saphenous nerve (fig 2).

The saphenous nerve is in particular danger during operations on varicose veins. 3 Removal of the saphenous vein for use as an arterial graft poses the same risk. 4 5 Saphenous nerve damage occurred in 3% of 421 patients in one operative study. 6 Cannulation of the saphenous vein at the ankle may injure the nerve and produce permanent painful paraesthesia. 7

Many patients with a saphenous neuropathy have relatively trivial numbness in the distribution of this nerve, but severe neuropathic pain can occur. There is often sensory loss and a tender neuroma may be present. The course of the nerve should be palpated carefully for areas of tenderness and a Tinel’s sign.

To our knowledge this complication has not been previously reported after fasciotomy for compartment syndrome. This case report highlights the importance of accurate placement of the posteromedial incision line to avoid saphenous nerve injury, which should be added to the list of potential complications of decompression fasciotomy.

Authors’ affiliations

D Pyne, A S M Jawad, N Padhiar, Department of Rheumatology, The Royal London Hospital (Mile End), Bancroft Road, London E1 4DG, UK

Correspondence to: Dr Pyne; dpyne@supanet.com

Accepted 5 November 2002

REFERENCES


Figure 2 Diagram showing the relation of the fasciotomy line to the saphenous nerve.