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Peripheral Nerve Entrapment and their Surgical Treatment

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Abstract

Nerves pass from one body area to another through channels made of connective tissue and/or bone. In these narrow passages, they can get trapped due to anatomic abnormalities, ganglion cysts, muscle or connective tissue hypertrophy, tumours, trauma or iatrogenic mishaps. Nearly all nerves can be affected. The clinical presentation is pain, paraesthesia, sensory and motor power loss. The specific clinical features will depend on the affected nerve and on the chronicity, severity, speed and mechanism of compression. Its incidence is higher under some occupations and is some systemic conditions: diabetes mellitus, hypothyroidism, acromegaly, alcoholism, oedema and inflammatory diseases. The diagnosis is suspected with the clinical presentation and provocative clinical test, being confirmed with electrodiagnostic and/or ultrasonographic studies. Magnetic Resonance Studies (MRI) rule out ganglion cysts or tumours. Conservative medical treatment is often sufficient. In refractory ones, surgical decompression should be performed before nerve damage and muscle atrophy are irreversible. The ‘double crash’ syndrome happens when a peripheral nerve is compressed at more than one point along its trajectory. In cases with marked muscle atrophy, a ‘supercharge end-to-side’ nerve transfer can be added to the decompression. After decompression in those few cases with refractory pain, a nerve neurostimulator can be applied.

Keywords: entrapment neuropathy, compression neuropathy, carpal tunnel syndrome, cubital tunnel syndrome, meralgia paraesthetica, cheiralgia paraesthetica, peroneal nerve entrapment, ulnar tunnel syndrome, radial tunnel syndrome, tarsal tunnel syndrome
1. Introduction

Nerves pass from one body area or cavity to another through holes and channels made of connective tissue and/or a bone channel, be it total (mental nerve) or partial (carpal tunnel). In these narrow passages, they can get trapped and/or injured due to congenital anatomical abnormalities, muscle or connective tissue hypertrophy, ganglion cysts, tumours, trauma or iatrogenic mishaps. Nearly all nerves can suffer an entrapment syndrome. Not all have the same incidence, some being very common (i.e. carpal tunnel) [1] and some exceedingly rare (i.e. tarsal tunnel) [2].

Entrapment neuropathies results in pain, paraesthesia and muscle power loss in the distribution of a peripheral nerve. With time muscle atrophy and skin trophic changes will appear. The clinical presentation will depend on the specific affected nerve, the chronicity, severity, speed and mechanism of compression [3, 4].

Nerve entrapment incidence is higher under some systemic conditions: diabetes mellitus, hypothyroidism, acromegaly, chronic alcoholism, extensive oedema and systemic inflammatory diseases [4]. Some occupations are associated with specific peripheral nerve entrapment syndromes. For example, occupations requiring repetitive wrist or finger movements or handling of vibrating tools have a higher incidence of carpal tunnel syndrome (CTS) [5–7].

Clinical presentation and provocative tests will suggest a diagnosis [3] confirmed or not with electrodiagnostic or ultrasonographic studies [4, 8]. Moreover, electrodiagnostic studies are also helpful to stage the severity and to rule out other confounding conditions (i.e. carpal tunnel and C7 radiculopathy, peroneal nerve compression vs. L5 radiculopathy) [3, 9] or generalized diseases (i.e. diabetic peripheral neuropathy) [9, 10]. MRI studies often show changes, ganglion cysts or tumours [11, 12] but the ultrasonography is less costly and more easily available [13].

Conservative treatment is sufficient in many cases (i.e. Saturday night palsy) but otherwise surgical decompression should be considered before irreversible peripheral nerve damage and muscle atrophy are established [3, 4, 8].

A nerve can be compressed at more than one single point, exacerbating the effects [14, 15]. This is called the ‘double crash’ syndrome and is common in some systemic diseases, particularly in diabetes mellitus [16].

In cases with advanced muscle atrophy, a ‘supercharge end-to-side’ nerve transfer is an option. After thorough decompression, a nearby healthy nerve is sectioned and sutured to the side of the previously compressed nerve, ideally distal to the entrapment point. The motor axons of the healthy nerve will grow inside of the damaged one much faster than the damaged axons of the damaged nerve. So, the healthy axons of the healthy nerve will keep the muscle alive while the axons of the damaged nerve recover [4]. This has been performed between the pronator quadratus (PQ) nerve branch and the motor fascicle
of the ulnar nerve (UN) at the forearm, between the flexor digitorum superficialis (FDS) and the anterior interosseous syndrome (AIN), between the triceps long head branch of the radial nerve (RN) and the axillary nerve, between the medial pectoral nerve and the axillary or the musculocutaneous nerve and between the spinal accessory nerve and the suprascapular nerve [4].

CTS is the most frequent entrapment syndrome, followed by meralgia paraesthetica and UN in the elbow. Decompression is always the treatment, removing the fibrous band, muscle or benign lesion causing the entrapment. After decompression, cases with refractory pain can undergo a nerve neurostimulator to block the pain transmission.

2. Upper extremity entrapment syndromes

2.1. Carpal tunnel syndrome (CTS)

This tunnel is formed by the ‘U’ of the carpal bones closed by the transverse carpal ligament. It is the most frequent entrapment neuropathy and one of the most common surgical conditions [17, 18]. Its estimated prevalence is 2% in men and 3% in women [17, 18], affecting a 3.72% of the USA population [1].

Idiopathic forms are due to a connective tissue proliferation of the flexor tendons synovium [19]. Some medical conditions predispose to its development: diabetes mellitus [20, 21], acromegaly [22], obesity [23], pregnancy [24], amyloidosis [25], hypothyroidism [22], rheumatoid arthritis [26], chronic kidney disease [27] and haemodialysis [28]. Its incidence is higher in occupations requiring repetitive finger and wrist movements [29], handling of vibrating tools [5–7] or repetitive blows with the palm of the hand (carpenters, sculptors) [29], but not with keyboard use [30]. It affects 30% of diabetics with polyneuropathy and 14% without it [31]. In pregnancy, it is most common in the third trimester [32].

Patients notice pain, numbness and tingling in the first three fingers of the hand. Initially symptoms are intermittent but become permanent with time, worsening with activity and at night [3]. Symptoms wake patients up at dawn, making them shake the affected hand to get rid of the symptoms (the so called flick sign). Paraesthesia may affect the whole hand. The pain may be an early symptom and radiate to the forearm or even to the whole arm up to the shoulder [33].

Sensory deficits affect the thumb, index and middle fingers and spare the thenar eminence [34], but 20% of clear-cut CTS show no sensory abnormalities [35]. Because the palmar cutaneous branch for the thenar eminence branches off the MN a few centimetres before the carpal tunnel the sensation of this area is normal in CTS. If this sensation is impaired pre-operatively it indicates proximal MN compression [3] while if damaged is only seen post-operatively it indicates iatrogenic injury.

Entrapment of this branch is possible but exceedingly rare [36].
Atrophy of the thenar muscles is a very late event in the progression of the disease [3] (Figure 1C), as are motor symptoms (Figure 1D) such as hand clumsiness, rigidity and loss of dexterity. Symptoms are usually bilateral but predominate in one hand.

The diagnosis is suspected by the symptoms and provocative manoeuvres (Phalen test (Figure 1A) and the Tinel and the carpal compression signs) [37]. The Phalen test indicates advanced disease [38], having a 75% sensitivity and a 47% specificity [39]. Electrodiagnostic studies confirm the diagnosis, rule out confounding conditions and stage the disease [40], with an 85% sensitivity and a 95% specificity [41]. Symptoms do not always correlate with electrodiagnostic findings. Ultrasonography is also useful [42]. Due to its higher costs, MRI is not used regularly [43].

Up to 20% of CTS cases improve with conservative treatments [44, 45]. Night-time wrist splints help 60% of patients but many eventually need an operation [46, 47]. Local corticosteroid injections can provide relief but often temporary [48]. Surgical decompression is the only proven long-term lasting relief [40, 49]. Any concomitant systemic disease predisposing to CTS should be treated at once although decompression is usually needed nonetheless [27]. The surgical procedure entails complete transverse carpal ligament section to decompress the MN. Local, regional or general anaesthesia are options, but local is faster and more cost effective [50, 51]. Open field (Figures 1F and G) or endoscopy has a similar time out of work, but the latter MN damage is more frequent [52–54]. Retinaculotomy decompression is similar to endoscopy but with less time and cost requirements [55] (Figures 1H and I). Re-operation is indicated in failure or recurrence. Incomplete decompression either at the distal carpal ligament or at the proximal antebrachial fascia is a frequent finding [56], but sometimes there is a thick scar tissue recreating the transverse carpal ligament and fixing the MN [4].

2.2. Pronator teres syndrome (PTS)

It is the MN compression as it passes through the pronator teres muscle (PTM) [57], the proximal arch of the FDS, the bicipital aponeurosis, the ligament of Struthers or an accessory head of the flexor pollicis longus muscle (FPL) (Gantzer’s muscle) [58]. There is proximal forearm deep pain with sensory and/or motor deficits in the distribution of distal MN [33, 57]. Repetitive pronation aggravates symptoms, and contrary to CTS, they appear at daytime and disappear during the night. Another difference is that in PTS there can be sensory loss in the thenar eminence [58]. The pronator compression test is a steady digital pressure on the proximal edge of the PTM 6 cm distal to the elbow crease and 4 cm lateral to the medial epicondyle for 30 s [57]. If positive, it should reproduce the symptoms. Forearm pain and hand MN paraesthesias can be induced by resistive pronation or by elbow extension with the forearm in pronation [57], elbow flexion with the arm supinated or flexion of the middle finger interphalangeal joint [58]. Electrodiagnostic studies can confirm the diagnosis [59]. Conservative treatment should be attempted, encouraging patients to avoid pronation, particularly against resistance. If symptoms persist, surgical decompression is recommended [60]. It is usually performed open field but some have reported an endoscopic approach [57]. The number of cases is limited, so no definitive conclusions on the best technique can be drawn yet.
2.3. Anterior interosseous syndrome

It is due to compression of this purely motor branch of the MN [61]. It induces a mild vague forearm pain accompanied by paresis or complete paralysis of the FPL and flexor digitorum profundus (FDP) of the second and at times part of the third finger [33]. The PQ is also affected but as the PTM remains intact the pronation is preserved. Patients notice lack of muscle power in the pinch between the thumb and index finger [62]. This induces difficulties on the writing hand [62], and the patient cannot make the OK sign [58]. There is no sensory deficit [61]. The causes are a tendinous origin of the PTM deep head or the third finger FDS, collateral ulnar vessel thrombosis, an accessory head of the FPL (Gantzer’s muscle), aberrant radial arteries or an enlarged bicipital bursa encroaching on the MN near the AIN site of origin [63]. Electrodiagnostic studies can confirm the diagnosis. If conservative treatment fails, surgical decompression is indicated [64]. The results are usually satisfactory provided that the nerve is freed on time.

2.4. Ulnar nerve compression at the elbow

Cubital tunnel syndrome (CubTS) is the entrapment of the UN at the elbow [65]. It is the most common site of UN entrapment and the second most common in the upper extremity nerve
[66]. Its estimated incidence is 25 new cases/100,000 inhabitants/year [67, 68], affecting males more often than females [68–72]. It is more common in jobs with constant leaning on the elbow (i.e. book keepers, drivers resting the elbow on the window frame) [73], gripping tools (gardeners, farmers, builders) [74], professional motorbike runners, cyclist [75, 76], repetitive elbow flexoextension [73, 74] and in floor cleaners [73, 77]. It is also more frequent in some systemic disorders like diabetes mellitus [16], acromegaly [78], rheumatoid arthritis [79] or amyloidosis [80]. CTS and CubTS in the same arm is not a rare finding [81–83].

Its clinical presentation consists of pain, sensory loss, paraesthesias, motor weakness and muscle atrophy at the forearm ulnar side and fourth and fifth fingers [33]. If untreated, it can lead to lack of sensation and muscle power, as well as pain and clumsiness in the affected hand [70]. Patients often complain of a dull pain at the elbow with shock-like sensations with any mild pressure or blow on this area. Some patients notice no sensory symptoms because of progressive weakness in the fourth and fifth fingers accompanied by muscle atrophy of the hand intrinsic muscles (Figures 2E and F) [84]. Symptoms get worse with activity and on flexing the elbow.

On clinical examination, the fifth finger remains in abduction due to weakness of the fourth palmar interosseous muscle (Wartenberg sign) (Figure 2A). This finger stays behind and out when the patient is attempting to put his/her hand inside the pocket [33]. The Froment sign is the flexion of the distal phalanx of the thumb when attempting to hold a piece of paper (Figure 2C). It is due to weakness of adductor pollicis, flexor pollicis brevis and first dorsal interosseous muscle, being substituted by the action of the FPL [33]. The weakness of the interossei and lumbrical muscles induces metacarpophalangeal joint hyperextension with flexion of the interphalangeal joint of the fourth and fifth fingers, creating the ‘claw hand’, ‘main en griffe’ or Duchenne sign (Figure 2B) [33]. Contrariwise to the hand of benediction seen with medial nerve damage at the forearm, the ulnar claw hand is due to the impossibility of the fourth and fifth fingers to extend. Meanwhile in the hand of benediction it is impossible to flex the thumb, index and middle digits when attempting to make a fist. In CubTS the weakness and atrophy of the first dorsal interosseous muscle (Figure 2D) is much more severe and earlier than the weakness and atrophy of the abductor digiti minimi (ADM) [33].

The most common site of UN entrapment is the retroepicondylar groove followed by the cubital tunnel 1.5–3 cm distal to the epicondyle [3]. About 40% of the cases are idiopathic [33]. The causes of compression are a bulky triceps muscle [85], the anconeus epitrochlearis muscle [86], fibrous bands bridging between the medial epicondyle and the olecranon [87], Osborne’s fascia [88] or the point where the UN crosses under the two heads of the flexor carpi ulnaris muscle [89]. It can occur after trauma or a protracted wrong position of the arm. It is the most common entrapment syndrome after anaesthesia for surgical procedures, particularly if they are long [90].

The diagnosis is based on the symptoms. Electrodiagnostic studies confirm the diagnosis and rule out other medical conditions (i.e. C8 radiculopathy) [3]. Some patients may improve with conservative measures like avoiding external elbow pressure, using a night time splint to keep the elbow extended or stopping any occupational activity that might be causing the disease. If that is not enough or the patient presents with muscle weakness and atrophy, a surgical decompression is indicated.
The techniques for UN decompression at the elbow are medial epicondylectomy, in situ decompression and transposition. The epicondylectomy is not popular anymore. Several clinical comparative studies [91–94], meta-analyses [95–97] and prospective randomized trials [98, 99] have shown that in situ CubTS decompression (Figures 3A–D) is just as effective as transpositions (Figures 4A–D) provided there is no UN subluxation on elbow flexoextension [97, 99]. Advantages of in situ decompression are smaller surgical incisions, less risk of medial antebrachial cutaneous nerve damage [100], no UN devascularisation [97], shorter operating time [101], smaller costs [99] and a faster recovery [102]. Transpositions need more surgical time, are more expensive and have more complications than in situ decompression [99], but can be used in the case of failure [102, 103]. It is imperative to avoid damaging the medial antebrachial cutaneous nerve or its branches regardless of the technique, as injury will induce post-operative neuropathic pain in the elbow area [104–106]. The in situ decompression can be performed open field or endoscopic. The latter shows a higher rate of post-operative surgical field haematoma [107]. The clinical results are equivalent for both procedures [100, 107, 108]. In recurrent cases, the most frequent finding is incomplete decompression either at the distal flexor-pronator muscle group or proximally at the intermuscular septum [109, 110].

Figure 2. Cubital tunnel syndrome. (A) Wartenberg sign; (B) claw hand; (C) Froment sign; (D) weakness of the first interosseous muscle; (E) atrophy thenar eminence; (F) atrophy thenar and hypothenar eminences; (G) surgical incision; (H) ulnar nerve (UN) compression by the (I) Osborne’s arcade (OS); (J) ulnar nerve fully decompressed with a compression mark at the retroepicondylar tunnel (depicted by the arrow).
Figure 3. In situ, ulnar nerve decompression. (A) patient position; (B) skin incision; (C) ulnar nerve (UN) decompression, the arrow points to the entrapment point at the retroepicondylar tunnel; (D) ulnar nerve fully decompressed.

Figure 4. Cubital tunnel syndrome. Subcutaneous ulnar nerve transposition. (A) patient position; (B) skin incision; (C) ulnar nerve (UN) decompressed; (D) ulnar nerve transposed subcutaneously.
Pre-operative and intra-operative electrophysiological inching studies have found that the compression point is at or immediately proximal to the cubital tunnel [87, 111–113], less often at the Osborne’s arcade but not proximally at the intermuscular septum [82]. Others with endoscopic assistance have reported no nerve constriction beyond 4 cm distally or proximally to the retroepicondylar tunnel [100]. So, extensive proximal decompression seems futile [100, 114]. Unsatisfactory results have been related to concomitant undiagnosed CTS or to weight gain [107].

2.5. Ulnar nerve compression at the hand

It is an uncommon site for UN entrapment (Figure 5A). Depending on the exact point of compression, it can be classified into five types [88, 115, 116]. In type I, the compression is proximal to Guyon’s canal with involvement of the superficial sensory, hypothenar motor, as well as deep motor branch. In type II, the compression is inside the canal and only the superficial sensory branch is affected. In type III, the compression is distal to the sensory branch with involvement of the hypothenar and deep motor branch proximal to the branch for the ADM. In type IV, the compression is distal to the superficial sensory and the hypothenar branch, so only the deep motor branch is affected. In type V, there is compression to the deep motor branch just proximal to the adductor pollicis and first dorsal interosseous muscles.

Usually there is the antecedent of an acute trauma [58] or chronic compression (cyclists) [76]. In other cases, there is a structural lesion in the area compressing the nerve, most commonly a ganglion cyst [116]. In cases of repetitive compression (i.e. cyclists), removal of the offending activity can be tried. If there is a lesion it has to be removed before irreversible UN damage develops (Figures 5B–E) [116].

2.6. Radial nerve (RN) entrapment syndromes

Its entrapment points are [3] at the spiral groove by the intermuscular septum between the triceps and brachialis (BaM) muscles, at the proximal forearm by the ligament of Frohse (posterior interosseous nerve or PIN), between the two heads of the supinator muscle (PIN) and by edge of the brachioradialis muscle (BRM) in the distal forearm (superficial cutaneous branch of the RN). It is the third most common upper limb entrapment [3].

The Saturday Night palsy or Honeymoon palsy is the most common compression of the RA occurring at the spiral groove [117]. It is usually due to local pressure on the posterior aspect of the arm under anaesthesia, drug intoxication (i.e. alcohol) or profound sleep with the arm over a hard surface or under the body of somebody else (Honeymoon palsy) [118]. In most cases, it improves spontaneously [117].

The radial tunnel syndrome is the entrapment of the deep branch of the RA [119]. This tunnel begins where the deep branch of the RN crosses over the radiohumeral joint, ending where this nerve becomes the PIN below the supinator muscle (SM) [33]. It has an average length of 5 cm. Its lateral wall is formed by the muscles BRM, extensor carpi radialis brevis (ECRB), BaM and extensor carpi radialis longus (ECRL). Its medial wall is created by the biceps tendon and the BaM. The floor of this tunnel is created by the radiocapitellar joint [120]. The compression can be due to the arcade of Frohse (tendinous border of the superficial layer of the SM),
the superomedial border of the ECRB, the inferior border of the superficial layer of the SM, fibrous bands at the radio-humeral joint and some radial vessels with a recurrent direction and a fibrous septum between the BRM and BaM muscles [119, 121, 122]. The treatment is conservative [123] avoiding the movements that induce pain but if it persists surgical exploration may be justified [124].

The PIN entrapment induces pain in the lateral aspect of the arm and forearm [125] and weakness in extension in all fingers (thumb included) and in thumb abduction (Figure 6A). There is no wrist drop because the ECRB is spared but on wrist extension the extensor carpi ulnaris weakness induces radial deviation [33]. There is no sensory deficit as the superficial RN is not involved. Symptoms exacerbate on hand or forearm repetitive movements [126]. Symptoms can be reproduced by direct pressure on the radial aspect of the forearm at 6 cm distal to the epicondyle. The provocative test consists in forceful hand supination with the shoulder in adduction and the elbow at 90° flexion or with extension of the middle finger [58]. It can be due to entrapment by the arcade of Frohse or on its way between both heads of the SM [126] (Figure 6C), ganglion cysts [125] or benign tumours (lipoma the most frequent) [127]. When symptomatic its treatment is surgical decompression [128] (Figures 6B–E).

Figure 5. Ulnar nerve entrapment at wrist. (A) Atrophy thenar eminence and claw hand; (B) skin incision; (C) ulnar nerve (UN) exposed in the distal third of the forearm; (D) ulnar nerve exposed in the wrist; (E) ulnar nerve deep (DB) and superficial branch (SB) liberated.
Cheiralgia paraesthetica or Wartenberg syndrome is the entrapment of the superficial cutaneous branch of the RN. There is numbness and pain in the dorsum of the hand, the thumb, index and middle fingers, accompanied by dysaesthesia, burning sensation and hyperesthesia [58]. The entrapment points are at the forearm between the tendons of the BRM and ECRB or at the wrist at its exit from beneath the fascia to the subcutaneous layer in the site where the fascia joins the tendons of the BRM and ECRL [129]. In the first case, it can be due to repetitive pronation and supination [58]. In this second case, it is usually due to external compression by hand-cuffs [130] or a wristwatch [131]. The best provocative test is to ask the patient to place their arm under maximum pronation with the wrist flexed to the ulnar side [132]. The Finkelstein test is positive, inducing confusion with a De Quervain syndrome [58]. The treatment is the removal of the cause. In case of persistence, surgical decompression of this nerve, particularly as it crosses the edge of the tendon of the BRM muscle, is indicated [131, 133] (Figures 7A–D).

2.7. Suprascapular nerve entrapment

It can be trapped at the suprascapular and spinoglenoid notches where the nerve is fixed by ligaments in a bony canal [134]. The first symptom is pain localized in the posterior aspect of the shoulder that gets worse with activity, when lying on the affected area, or by
shoulder adduction crossing the midline with the extended arm [33]. The weakness and atrophy of the supra and infraspinatus muscles induce paresis of shoulder abduction and external rotation.

On clinical examination, the affected shoulder is lower than the healthy one and the scapular muscles are atrophied (Figure 8A). The patient has difficulty raising the outstretched arm above the horizontal (Figures 8B and C).

Its treatment is surgical with section of the ligament that closes the suprascapular notch at the superior aspect of the scapula. It can be done open field [135] (Figures 8D and E) or endoscopically [136] with similar outcomes.

2.8. Thoracic outlet syndrome

There is pain in the inner aspect of the arm and forearm, sometimes reaching the fourth and fifth fingers [137]. This pain gets worse when lifting the arm above the horizontal [138]. Sometimes there is associated hand muscle atrophy [139]. Claw hand deformity can be
present in the long protracted cases [138]. The neurogenic type has an incidence of one case per million inhabitants [137]. It can be due to hypertrophy of some muscles at the root of the arm (typical of ceiling painters or swimmers) or to the existence of a cervical rib or fibrous ligament at the same point [137].

The clinical presentation is pins and needles with overhead activities (like painting a ceiling) [140], carrying heavy objects with the arms hanging down [138], combing hair and applying makeup [4].

In the Roos’ elevated arm stress test, the shoulder is abducted 90° and kept in external rotation with the elbow in 30° flexion. This provocative test is positive if opening and closing the hand for 1 min reproduces the symptoms [141].

In case of poor response to conservative treatment, surgical decompression is in order. The two possibilities are the transaxillary removal of the first rib [142] or supraclavicular scalenectomy [143] (Figures 9A–D). This depends on the causative mechanism and the surgeon’s preferences but the supraclavicular approach offers a better chance of solving any causative abnormality [143].

Figure 8. Suprascapular nerve entrapment. (A) Atrophy supra and infraspinatus muscles; (B) and (C) weakness of shoulder abduction; (D) suprascapular nerve (SSN) exposed with the ligament for the suprascapular notch (LSN) intact; (E) suprascapular nerve after removal of the ligament for the suprascapular notch.
3. Lower limb entrapment syndromes

3.1. Meralgia paraesthetica

The femoral cutaneous nerve is a purely sensory nerve which runs usually medial to the ASIS. The name meralgia paraesthetica comes from the Greek, *meros* meaning thigh and *algos* meaning pain. The clinical presentation is pain, paraesthesia, numbness and hypersensitivity in the anterolateral side of the thigh down to the knee (Figures 10A and B) [144]. There are no motor signs and if present a different medical condition should be suspected [145]. The pain worsens when standing or walking and improves on sitting [146]: it is aggravated on leg extension and improved on knee flexion [146]. Its estimated incidence is 36.2 cases/100,000 habitants/year in the USA [147] and 43 cases/100,000 persons/year in Europe [145]. Meralgia paraesthetica is more frequent in obese people, in the fourth–six decades of life [145, 147, 148] and in diabetics (seven times more than in the general population) [147, 148]. Although many cases are idiopathic oftentimes its cause is an external compression of the nerve as it passes underneath or through the anterior inguinal ligament at its origin on the ASIS [149]. This can be due to either internal causes like a bulging abdomen [150], obesity [147], pregnancy [151], ascites [152], external agents like tight clothes [153] or belts resting on the outer aspect of the thigh.

Figure 9. Thoracic outlet syndrome. (A) skin incision; (B) supraclavicular cutaneous nerve (SCCN), sternocleidomastoid muscle (SCM), clavicle (CL); (C) brachial plexus (BP) exposed; (D) first rib exposed, brachial plexus upper trunk (BPut).
repeatedly while standing (typical of hairdressers) or due to a prolonged position (lithotomy posture, cycling) [144]. It can also be induced by a pelvic fracture with psoas haematoma or by surgical procedures such as hip [154] or knee replacements [155] or an aortofemoral bypass [144, 147]. It has also been related to acute seat belt compression in car accidents [144].

The clinical presentation in the absence of motor signs helps to make the diagnosis. Electrodiagnostic studies can rule out confounding conditions [144, 156], but ultrasonography is very useful, particularly in obese patients [156, 157].

Initially, the treatment is to remove the compressing agent and/or lose weight. If insufficient, the area can be infiltrated with a local anaesthetic agent and corticosteroids [158]. The rebel cases require surgical treatment with nerve decompression (Figures 10C–E) or neurectomy [159].

3.2. Peroneal nerve entrapment

It is the most common lower limb entrapment neuropathy [162]. It is a mixed nerve that runs at the fibular head, reaching the anterior compartment of the leg distal to the knee [160]. At that level, it lies between the skin and bone. This makes it very sensitive to trauma or pressure.
particularly in bedridden lean patients [161]. It can also be due to mass lesions (i.e. ganglion cyst of the tibiofibular joint) or associated with systemic diseases (diabetes and vasculitis) [162]. It is more frequent in occupations requiring people to squat for long periods of time (strawberry pickers, farm workers and carpet layers) [163] or that sit crossing their legs [164].

The clinical presentation is pain at the fibular head and loss of strength in dorsiflexion (Figure 11A), which causes the foot to drag when walking. The patient notices foot slap with steppage gait and wearing the tip of the shoe as well as a sensory loss on the dorsal aspect of the foot between the first and second toe [165].

The treatment is surgical decompression (Figures 11B–G) [161, 166].

3.3. Anterior tarsal tunnel

It is the entrapment of the deep peroneal nerve. The clinical presentation is pain in the dorsum of the foot associated with sensory loss in the first foot web space [167]. The treatment is initially conservative, but surgical decompression with extensive fascial opening might be needed [168].

Figure 11. Peroneal nerve entrapment at the level of the fibular head. (A) foot dorsiflexion weakness; (B) skin incision and fibular head (FH) depicted; (C)–(G) steps in peroneal nerve (PN) decompression, subcutaneous fascia (SB). The arrow points at the marked impinged on the nerve by the compressing fascial band.
3.4. Tarsal tunnel syndrome

It is due to compression of the posterior tibial nerve at the tarsal tunnel behind the foot medial malleolus. It is very uncommon. Many cases are idiopathic (20–46%) [169]. Contributing factors are ankle sprain and fracture, tight-fitting foot wear and space occupying lesions [170]. The clinical presentation is pain, paraesthesia and numbness in the sole of the foot. This symptoms get worse on standing, walking and at night time [2]. The sensory loss affects the sole of the foot sparing the heel, supplied by the calcaneal branch [171]. The diagnosis is identified with the clinical presentation. Electrodiagnostic studies can be useful to rule out confounding medical conditions [172]. Ultrasonography [173] and MRI [174] can rule out associated space occupying lesions.

Its initial treatment is rest and anti-inflammatories, but if there is no improvement or relapse after an initial response, surgical decompression may be necessary. This can be done endoscopically [175], but for a good decompression, especially in the distal part, an open approach gives better results [176] (Figures 12A–D).

**Figure 12.** Tarsal tunnel syndrome. (A) skin incision; (B) exposure medial malleolus (MM); (C) section of roof tarsal tunnel, posterior tibial structures (PTS); (D) exposure posterior tibial structures.
3.5. Piriformis syndrome

It is a very rare disorder in which the sciatic nerve is compressed by the piriformis muscle. This is a flat, pyramid-shaped muscle located deep in the gluteal region, running between the femur and the iliac bone. It helps in hip external rotation [177]. Common sciatic nerve pain (i.e. lumbar disc hernia) is much more prevalent than this syndrome.

The conditions associated with this syndrome are sitting for extended periods of time, sitting with a large wallet in the rear pocket, repeated forward movements, running, bicycling, stiff sacroiliac joints, foot overpronation, Morton’s toe (the second toe is longer than the first one) and after a fall on the buttocks [177–179]. Approximately 50% of the cases are caused by trauma and the rest are spontaneous [178].

The symptoms are sciatica-like pain. Pain starts in the gluteal area and may travel through the back of the thigh and calf up to sole the foot. Patients might experience tingling, numbness, burning sensation and weakness. The sciatic pain aggravates with sitting or with activities that press the piriformis against the sciatica nerve, such as running, cycling or horse riding [178].

The diagnosis is usually made through physical examination. Certain tests may elicit sciatica nerve pain indicating the presence of the syndrome, especially internal rotation of the hip with the knee in full extension.

On MRI examination, it is possible to see the sciatic nerve with oedema when crossing under the piriformis muscle.

Conservative treatment is initially recommended. Alternate ice and heat treatment may provide relief. Ultrasound penetrates deep into the muscle alleviating the sciatica nerve pain. Stretching exercises to target the piriformis, hamstrings and hip muscles, will help increase the range of motion and decrease the sciatic nerve pain.

If all these treatments prove unsuccessful, injection of botulinum toxin in the piriformis muscle [180] under CT or MRI guidance can be attempted. In the case of failure, surgical decompression removing the piriformis muscle or the offending fibrous band could be indicated [179, 181] (Figures 13A–E). The results are inconsistent.

3.6. Pudendal nerve entrapment

It induces pain in the genital and sometimes gluteal areas [182]. The pain worsens with local pressure, sitting, defecating, and urinating and with sexual intercourse [183]. It is constant, intense and burning. The cause can be local pressure induced by repeated cycling [184] or by horse riding. The problem is that most patients are diagnosed late. Once suspected, it can be confirmed with electrodiagnostic studies [185]. When conservative treatments [186] fail, surgical decompression should be considered [187] (Figures 14A–E). The results are often poor, at times because patients are diagnosed much too late due to lack of awareness in the medical world.
Figure 13. Piriformis syndrome with sciatic nerve (SCN) entrapment. (A) Skin incision; (B) gluteus maximus muscle exposed; (C) piriformis muscle (PiM) isolated; (D) piriformis muscle sectioned exposing the sciatic nerve (SCN); (E) sciatic nerve free.

Figure 14. Pudendal nerve entrapment. (A) Skin incision; (B) gluteus maximus muscle dissection exposing the sacroischias ligament (SIL); (C) the pudendal nerve (PuN) is exposed after sectioning the sacroischial ligament; (D) the pudendal and its branches are freed (RBPuN, rectal branch pudendal nerve; PBPuN, perineal branch pudendal nerve); (E) post-operative scar.
4. Conclusion

Nerve entrapments syndromes are more frequent than currently thought. Their awareness is essential to diagnose and treat the patient on time. Although almost any nerve can suffer an entrapment syndrome, some are more common than others. The most frequent is CTS, followed by meralgia paraesthetica and ulnar nerve entrapment at the elbow. The clinical presentation is pain, paraesthesia and muscle power loss in the distribution of the affected nerve. Many cases are idiopathic, but others are induced by internal or external compressing mechanisms. Some systemic conditions are associated with an increased incidence of these syndromes. The clinical presentation together with the electrodiagnostic studies help in the diagnosis. The ultrasonography and the MRI are also helpful but not used so regularly.

In many cases, conservative medical treatment is sufficient. When it is not, surgical decompression has to be performed.

Open and endoscopic approaches are available. In each case, we will have to see which shows better outcomes. A few cases with persistent pain after surgical decompression might benefit from peripheral nerve neurostimulation.

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Appendices and nomenclatures

| Abbreviation | Full Form                                      |
|--------------|-----------------------------------------------|
| AIN          | Anterior interosseus syndrome                 |
| ASIS         | Anterior superior iliac spine                |
| BaM          | Brachialis muscle                             |
| BRM          | Brachioradialis muscle                        |
| CubTS        | Cubital tunnel syndrome                       |
| CTS          | Carpal tunnel syndrome                        |
| ECRB         | Extensor carpi radialis brevis                |
| ECRL         | Extensor carpi radialis longus                |
| FDS          | Flexor digitorum superficialis                |
| FDP          | Flexor digitorum profundus                    |
| FPL          | Flexor pollicis longus                        |
Peripheral Nerve Entrapment and their Surgical Treatment

MN  Median nerve
PIN  Posterior interosseous nerve (PIN)
PQ  Pronator quadratus muscle
PTM  Pronator teres muscle
RN  Radial nerve
SM  Supinator muscle
UN  Ulnar nerve

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