Skin biopsy and small fibre neuropathies: facts and thoughts 30 years later

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ORIGINS
Between 1989 and 1990, two independent research groups of anatomists and histologists at the Karolinska Institute of Stockholm, Sweden, showed that antibodies against the cytoplasmic protein gene product 9.5 (PGP9.5) could visualise in bright-field immunohistochemistry the rich cutaneous innervation much better than neuropeptides and other antibodies previously did1,2 (figure 1). Their publications would soon widen the interests of neurology to the human skin.

This novel pathological technique was then used to understand the symptoms in a group of patients who had distal neuropathic limb pain, primarily in the feet, but in whom standard neurological examinations, aside from reduced pinprick sensations, standard nerve conduction studies3 and even nerve biopsy findings4 were normal. The findings of reduced or absent intraepidermal nerve fibres (IENF), either distally alone or in a length-dependent fashion, strongly suggested that the complaints of distal neuropathic limb pain and nearly normal examinations were due to involvement of the small nerve fibres in the skin, primarily those subserving nociception and thermal sensation. This fitted well with classic neurological education of the fundamental importance of clinical–physiological–pathological correlation in neurological disease.

The new protocol for the quantification of IENF density5 overcame the first and more complicated one used in the seminal work of 1959 by Arthur and Shilley6 (figure 2) and was then used to provide a measurement at different sites. The technique was initially available only at universities with interest in peripheral neuropathies. These academic centres developed, through international collaborations, normative reference values for the different techniques6–7 and published guidelines,8 providing sex and age-adjusted values of normal IENF density at the distal site of the leg for clinical use. This overcame the difficulties related to the development of single centre normative data, which requires the enrolment of very large numbers of healthy individuals. More recently, in the USA at least, commercial laboratories have sprung up claiming to provide the same service. This has led to an explosion of skin biopsies done to evaluate IENF density and a larger explosion of so-called ‘small fibre neuropathies’ that have gone far beyond the bounds set by classic neurologic clinical–physiological–pathological correlation. How are we to understand these developments?

FUNCTIONAL AND PATHOLOGICAL CORRELATION
IENF are the endings of small size dorsal root ganglion (DRG) neurons whose expression of the transient receptor potential vanilloid type 1 receptor9 reflects their nature as distal nociceptors. IENF lose the enwrapping of non-myelinated Schwann cells while crossing the dermal–epidermal barrier,10 like large myelinated sensory nerves do while twisting around dermal mechanoreceptors,11 to remain naked among antigen presenting cells (dendritic cells and melanocytes) and keratinocytes which are known to participate in the transduction of nociceptive signals, which explains why their loss is associated with a syndrome of pain hypersensitivity in vivo and in vitro.12,13

Figure 1 Immunofluorescence micrograph of section of human skin incubated with antibodies to PGP9.1 showing the innervation of the subepidermal area (A), a small artery (B), a sweat gland (C) (adapted from Dalsgaard et al, 1989) and the epidermis from the back (D) (adapted from Wang et al, 1990).

Figure 2 The pattern of cutaneous free-ending innervation. The epidermis is seen from above. Subepidermal nerves are drawn in blue and epidermal nerve in red (adapted from Arthur & Shilley, 1959).
Occasional essay

of sensory sensation.12 This makes the epidermis, rather than IENF alone, a huge polymodal receptor.

The density of IENF at the distal site of the leg in healthy individuals slightly declines over the decades, using either bright-field or immunofluorescence techniques, and is generally higher in women.6,7 However, the density does not differ between the right and left side, and, most importantly, it does not vary when reassessed after 3 weeks, which is the period of keratinocytes’ turnover, both in healthy individuals and in patients with neuropathy, strengthening the diagnostic reliability of the evaluation.13,14

Small size sensory neuron or axon damage can lead either to the loss of thermal and nociceptive sensations, to neuropathic pain, or a combination of both, which is common in several sensory length-dependent neuropathies and non-length dependent neuropathies. IENF loss seemed able to resolve the clinical–physiological–pathological correlation in almost any process selectively affecting DRG nociceptive neurons or their axons, whenever the related symptoms were appropriate. Indeed, this was reported for patients with congenital insensitivity to pain syndrome,14 for those with small fibre neuropathy (SFN) combining neuropathic pain and loss of pinprick sensation,15 and for those with painless diabetic neuropathy.16 Studies on sensory sensation and neuropathic pain after artificial destruction of these skin nerve fibres,17 their spontaneous regrowth in both experimental18 and disease-related loss,19 and changes in other painful20 or painless14 diseases confirmed such clinical–physiological–pathological correlation.

Noted early was the lack of correlation between IENF density and neuropathic pain,21 22 which in retrospect should not be a surprise. PGP9.5 staining simply detects those skin nerves intact enough to be identified, but it does not tell anything about their functions, which are driven by a precise molecular ontogenesis toward tissue targeting.23 Other molecular markers linked to different subtypes of skin nerves might provide a better clinical–physiological–pathological correlation, as suggested by the increased density of IENF expressing peptides in painful diabetic SFN24 and specific molecular signatures for regenerating nerves.25

WHAT IS SFN?

SFN is a disease of somatic, and commonly to a lesser extent autonomic, thin myelinated and unmyelinated nerve fibres. It is called ‘pure’ when those subtypes of sensory nerves alone are affected, but, more commonly, it occurs as part of a more diffuse impairment of peripheral nerves, of which SFN could be the earliest manifestation.

The classical presentation of SFN is that of a length-dependent neuropathy, a term that links the underlying pathophysiological mechanism of axonal dying-back to the clinical features and anticipates that the feet are first affected by sensory symptoms. Indeed, burning feet has been the eponym used both for familial and sporadic cases.26–28 Further clinical presentations have been proposed, from focal neuropathy underlying the cases of burning mouth syndrome29 30 or pain after inguinal hernia repair,31 in which the distribution of symptoms respected the neuroanatomical distribution, to a non-length dependent presentation reflecting the primary involvement of DRG neurons in paraneoplastic and non-malignant immune-mediated diseases and genetic syndromes.31 32

The intrinsic limitations of routine nerve conduction studies, psychophysical measurement of thermal thresholds, pain-related evoked potentials and autonomic function evaluation for achieving the diagnosis of SFN in individual patients have been overcome by skin biopsy at the ankle essentially for one reason: the possibility to provide an objective morphometric assessment of the target fibres in the body region corresponding to the symptoms.

Physicians understand that without context an abnormal laboratory value or imaging study has no intrinsic meaning. This perfectly applies also to IENF quantification. Indeed, for neurologists the interpretation of an abnormal IENF density should be just that, nothing more without context. One relevant example is that of sodium channelopathies. Sodium channels are key membrane proteins for the generation and propagation of action potentials and are expressed at low density throughout the length of small nerve fibres. Functional changes caused by gene mutations have been demonstrated in typically painful SFN, but not all patients showed a reduction of IENF.33–35 Similarly, patients with inherited erythromelalgia, a severe autosomal dominant syndrome with distal neuropathic limb pain and dysautonomia also caused by sodium channel mutations, can have normal IENF density.36 At the other end of the spectrum, patients with congenital insensitivity to pain have profound loss of IENF.37 These observations emphasise that epidermal nociceptors can be functionally and structurally altered in disorders causing loss of nociception without pain, and functionally abnormal but structurally normal in some painful syndromes. Similarly, specific sodium channel subunits could be involved in the generation of peculiar neuropathic pain symptoms caused by neurotoxic compounds without evidence of IENF degeneration.38

Therefore, the measurement of IENF density at the distal site of the leg is just one among other synergic clues needed to diagnose the classic, length-dependent, SFN.39 However, due to widespread use of skin biopsy, the commercial labs that promote it, and interpretative yet deceptive language used in some reports, the diagnosis of ‘SFN’ is frequently applied to any condition in which IENF at any site is found reduced. This has generated a blurred nosology, because SFN may be virtually (or erroneously) thought of as any disorder with reported loss of IENF, whatever the site of the biopsy, or the distribution or even the type of symptoms.

INTERPRETATION OF RESULTS IN THE CLINICAL CONTEXT

The loss of IENF is expected to lead to symptoms and clinical signs correlated to the physiological functions of small nerve fibres. Indeed, the old paradigm that SFN patients can complain of severe burning pain and yet have normal clinical examination has been forgotten. In most patients with SFN, the clinical picture is characterised by positive (eg, hyperalgesia) and/or negative (eg, loss of nociception) signs of small fibre dysfunction.33 Moreover, when looking at how symptoms correlate with clinical signs and the results of the diagnostic testing, a large study demonstrated that nearly 90% of patients with neuropathic-sounding symptoms but no clinical signs at examination had normal results for both quantitative sensory testing (QST) and IENF density, which remained normal after 18 months. This suggests that the diagnosis of SFN should not rely on symptoms alone, because they are not sufficiently specific.

Taking SFN as the benchmark, the same correlation should be expected also for other conditions in which there is loss of IENF, whatever is its distribution (eg, length-dependent or independent, focal). Conversely, this does not seem to be always the case.

One example is that of fibromyalgia, a clinical condition diagnosed by the presence of ‘multi-site’ pain, meaning that six out of nine sites are painful to the patient.39 Due to the lack of any clinical and electrophysiological evidence of large fibre sensory
nerve involvement, fibromyalgia patients have been investigated by skin biopsy and other methods for assessing small fibre functions, with conflicting results. Some have found diffuse IENF loss correlating with the disease severity. Others reported little contribution from peripheral small nerve fibres as compared with central pain processing impairment [s41], and yet others did not report clinically meaningful correlations between diagnosis of fibromyalgia and small nerve fibre pathway functioning [s42], thus challenging the definition of the small fibre pathology attributed to this disorder [s43].

This lack of correspondence between the distribution of painful symptoms and the site of skin biopsy also hinders the definition of the plausible neuroanatomical correlation required for diagnosing neuropathic pain [s44], contributing to such a blurred scenario. While the loss of IENF assessed in the correspondent clinical area allowed meeting this diagnostic requirement in focal painful syndromes like burning mouth [s45] and atypical pain of the sacral region [s46], even though no underlying pathological mechanism is increasingly filled with reports of SFN in several conditions where the a priori view would be that these distal small fibres are not involved [s47]. These include various conditions of central dysautonomia like postural orthostatic tachycardia syndrome [s50], Ehlers-Danlos syndrome [s51] or after vaccination [s52].

Recently, autism joined this list. Autism is a complex neurodevelopmental disorder in which a variety of molecular mechanisms likely underpin the clinical phenotype, which includes hypersensory and hyposensory responsiveness, and sensation-seeking [s53]. A recent paper reported the reduction of IENF at the distal site of the leg in about half of 32 adult autistic patients [s54]. Leaving aside the changes in psychophysical somatosensory thresholds and of contact heat evoked potentials amplitude, which might not be surprising within the complex multisystem sensory processing impairment of autism [s55], the simple association between reduced IENF density at one leg and tactile and autistic symptoms seems quite a large conceptual jump.

In some clinical situations, neurologists might expect a loss of IENF, those being mainly complex neurodegenerative diseases. While many of these affects one neuronal population primarily, affection to a minor degree of other neuronal populations might not be unexpected. Long ago, Professor P J Dyck showed that some patients with amyotrophic lateral sclerosis had large fibre neuropathy [s56], so it is not surprising that some also have reduction in IENF as indeed it was found [s57], and that insights on possible molecular mechanisms were described [s58]. The same logic would extend to Parkinson’s disease, in which neuropathy possibly attributed to L-dopa therapy has been first reported in few patients [s59], then found to involve unmyelinated axons from autopsy sural nerve biopsies [s60], and eventually confirmed by skin biopsies [s61], even though no underlying pathophysiological mechanisms has been proposed yet. It is unclear whether this could apply to other disorders based on one single anecdotal observation like fragile-X syndrome [s62] or when the diagnosis is made on screening questionnaires like for Pompe disease [s63].

**IMPLICATIONS ON TREATMENT**

The popularity of the idea that small nerve fibres could be affected in any disease, even lacking any clinical—physiological correlation, and being at the same time easily diagnosed, brings a number of concerns. Among these, the most important for many reasons is proposing disease-modifying treatments with unknown corresponding pathophysiological mechanisms nor evidence of efficacy from properly designed clinical trials. Intravenous immunoglobulin (IVIg) is in the top five drug categories in terms of annual spending in the USA and the yearly in-hospital cost per patient is about US$60 000 [s64]. In recent years, the hypothesis that patients with SFN associated with systemic immune-mediated diseases could benefit from IVIg originated from retrospective, anecdotal or small open-label studies [s65–67]. Following them, and with the same uncontrolled approach, this hypothesis of IVIg efficacy spread from fibromyalgia [s68] to ‘apparently’ autoimmune SFN of adults and children [s69, s70]. However, the largest of these studies showed a 1-point improvement of pain relief in a 0–10 scale [s70], which is hardly clinically meaningful as this is the same magnitude of change in the placebo group in placebo-controlled studies of pain [s71]. One answer arrived from the first randomised controlled trial designed to investigate if IVIg had any efficacy in providing SFN patients with pain relief. The administration of IVIg (2 g/kg body weight) or placebo, followed by infusions of IVIg (1 g/kg) or placebo at 3-week intervals in 60 skin biopsy-proven idiopathic SFN did not change the Pain Intensity Numerical Rating Scale score at 12 weeks compared with baseline [s72]. Even though the result of this trial cannot be extended to any disorder classified as SFN, it suggests that controlled trials are mandatory before suggesting the effectiveness of IVIg or any other treatment.

**WHAT ARE THE LESSONS?**

Reduced IENF in skin biopsy, like many tests, is a finding, not a diagnosis, which in medicine requires at least two clues converging within the clinical context. SFN should not be regarded as an exception. IENF density results need to be compared with a control population usually using published data, which are available for the two most commonly used techniques. Developing local controls without an interobserver quality control programme or controls for a specific condition is fraught with potential problems especially if a small sample size is used. The site where biopsy is taken is also important as control data do not exist for many sites now being biopsied such as the top of the foot. Moreover, data should be shown as both mean with SD/SE, mean and range/IQR, but also as percentage of truly abnormal by a predefined outcome measure. Some laboratories report values as ‘low normal’. Currently, this has no meaning and certainly no diagnostic significance.

In clinical studies either focusing on diagnosis or efficacy of treatments, blinding of skin biopsy reading is critical and should be specified in the protocol.

Adding skin biopsy to diagnostic criteria for a disease should be carefully considered. Rather than jumping to conclusions, understanding how the biopsy results might fit into the clinical context should be pursued. Some have used the biopsy alone, divorced from the clinical context, to diagnose SFN and to obtain authorisation for IVIg or other putative treatments. We have grave reservations about this line of thinking and see no
justification for IVIg in SFN aside from within well-designed controlled clinical trials, as recently demonstrated [327].

While skin biopsy for IENF density provides a pathological view of the state of small nociceptive and thermal fibres, the simple reduction of IENF, whatever its degree, when blind to the clinical context makes it impossible to predict symptoms, signs and aetiology of the underlying neuropathic process that could affect DRG nociceptors or their axons. Without denying the skin biopsy findings in some patients dissociated from the clinical picture, the hypothesis that assessing IENF at one lower limb may explain complex sensory symptoms or those localised in different body areas seems misguided. This is not trivial, as it has to do with the nosology of a disorder and the consequent diagnostic criteria applied in clinical practice and trials.

Thirty years after the appearance of skin biopsy, which has widened the diagnostic toolbox of neurologists providing structural information on nerves used to be considered clinically invisible, a new step toward the interpretation of the results should be undertaken to make it a measure for the improvement of the diagnostic reliability in the disorders potentially affecting small nerve fibres.

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