Nodo-Paranodopathy: Beyond the Demyelinating and Axonal Antibody-Mediated Neuropathies

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Abstract

In the last decade, antibodies targeting the node and paranode of myelinated peripheral nerves have been increasingly identified in patients with acquired immune-mediated neuropathies, commonly termed ‘nodo paranodopathies’. These patients present with clinical features not usually seen with the most common immune mediated neuropathies, Guillain-Barré syndrome and chronic inflammatory demyelinating polyneuropathy. They respond poorly to conventionally used immunomodulatory therapies. Emerging evidence that these are pathologically distinct diseases has further prompted the use of more targeted treatment, such as the B cell depleting monoclonal antibody rituximab, which has been reported to significantly improve functional outcomes in this subset of patients. It is now apparent that antibodies directed against several region-specific cell adhesion molecules, including neurofascin, contactin and contactin-associated protein, can be linked to phenotypically distinct peripheral neuropathies. Importantly, the immunological characteristics of these antibodies facilitate the prediction of treatment. Electro-clinico-pathological diagnostic criteria of these acquired immune mediated neuropathies thus need to be reclassified in the light of this disease entity of ‘nodo-paranodopathies’. In this review the readership is directed to explore the presence of these specific antibodies that cause this “nodo-paranodopathy” immune phenotype that has a different therapy paradigm and prognosis compared to the conventional acute and chronic neuropathy syndromes that we know of so very well.

Keywords: Guillain-barre syndrome, paranode node, CIDP; Node of Ranvier, Neurofascin, Contactin contactin-associated protein 1

Introduction

Traditionally inflammatory polyneuropathies, be it acute -like Guillain-Barre syndrome or chronic like CIDP have been diagnosed based on clinical, electrophysiological and pathological criteria1. However, clinical experience indicates that some of these neuropathies have responded differentially to immunotherapy and the temporal profile was different in a small subsection of both acute and chronic neuropathies of inflammatory background2. The pathogenesis of the immune neuropathies was explained by a combination of the cell-mediated, autoantibody induced and complement mediated immune injury mechanisms as well as disruption of the blood nerve barrier3. This then led to the identification of ganglioside autoantibodies in both the acute and chronic immune inflammatory neuropathies and their specific attachment to nerve proteins4. In specific situations autoantibodies against paranodal proteins have led to the identification of paranodopathies that are being increasingly recognised in the last few years. It is now apparent that antibodies directed against several region-specific cell adhesion molecules including neurofascin, contactin and contactin associated protein can be linked to phenotypically distinct peripheral neuropathies5. The immunological characteristics of these antibodies facilitate the production of treatment responsiveness. In what was initially identified to be refractory CIDP patients who responded to Rituximab we now have this specific diagnosis of “nodo-paranodopathy” to explain the disorder.

History

The node of Ranvier was originally described in 1871 to demonstrate that the nodes were places where ionic currents generated action potentials for saltatory conduction. Only in the last two decades this same nodal region has been recognised as a possible site of specific autoimmune attack in peripheral neuropathies6. Further delineation of the nodal region has led to the identification of the node, the paranode, the juxtaparanode and the internode regions with specific and characteristic distribution of the ion channels (predominantly sodium and potassium channels) and various proteins that are densely arranged to facilitate adhesion of myelin to axon and help maintain the correct localisation of these voltage-gated sodium and potassium channels facilitating effective saltatory conduction.
Of the spectrum of antibodies, immunoglobulin G4 (IgG4) specifically targeting nodal neurofascin 186 (NF186), the para nodal neurofascin 155 (NF155), contactin-1 (CNTN1) and contactin–associated protein –1 (Caspr1) have been identified in phenotyping distinct groups of patients among the paranodopathies.

**Pathogenesis**

The clues to the currently identified paranodopathies goes back to the early 1990s when the connection between acute motor primarily axonal subtype of the Guillain-Barre syndrome with antecedent Campylobacter jejuni infection and antiganglioside antibodies was established. Pathology studies of patients with this acute motor axonal neuropathy (AMAN) showing Wallerian degeneration with no demyelination and inflammation and with the earliest changes showing as lengthening of the node of Ranvier. IgG and complement deposition at the nodes of Ranvier preceded the development of Wallerian like degeneration. This led to the hypothesis of antibody binding resulting in paralysis with little structural changes and with the potential for rapid recovery. Despite this evidence, the simple explanation of axonal degeneration and the electrodiagnostic features lacking in demyelination overlooked the role of the para nodal protein dysfunction through antibody mediated conduction failure. Further identification of anti-GM1 and GD1a antibodies showing conduction block and or conduction slowing that then promptly resolved without the development of excessive temporal dispersion of the compound muscle action potentials characteristic of demyelination were reported. This was termed reversible conduction failure (RCF) due to temporary failure of conduction at the nodes of Ranvier due to loss of sodium voltage-gated channels. Next came the demonstration of nerve conduction velocity decreasing by blockage of these voltage-gated sodium channels by intravenous infusion of lignocaine or tetrodotoxin. These observations and resolution of understanding of the nerve conduction and the role of the nodes of Ranvier led to the concept of autoimmune nodo-paranodopathy. The reversible conduction failure (RCF) phenomena were not only detected in motor fibres but also in sensory nerves with lengthening of the nodes in the dorsal root fibres and in patients with the pharyngeal – cervical -brachial subtypes of Guillain-Barre syndrome. Pure involvement of the sensory fibres by RCF is seen in acute sensory ataxia neuropathy and in the Miller Fisher syndrome (MFS). Subsequently anti GQ1b antibodies revealed elongated nodes, myelin splitting at the para node and macrophage invasion of the internodal axon without any features of segmental demyelination quite similar to the pathology in acute motor axonal neuropathy (AMAN).

Likewise, in CIDP research in 2012 demonstrated that CIDP patients carried IgG to CNTN1 or CNTN1 and Caspr1. These Spanish patients had a common phenotype of advanced age, aggressive onset, severe axonal damage, motor predominance and poor response to intravenous immunoglobulin. The Spanish group also had patients with antibodies to NF155 with disabling tremors, a severe distal sensorimotor neuropathy and poor response to intravenous immunoglobulin. A further patient was identified as having Caspr1 antibodies with CIDP and prominent neuropathic pain. Thus 10% of all patients classified as CIDP had antibodies against proteins of para nodal junctions. This further subclassification among the AIDP and CIDP led to the paranodopathy diagnosis.

Pathology: (FIGURE ONE and TWO) the axonal cell adhesion molecules CNTN1 and Caspr1, and the glial NF155 represent an essential complex in the formation and stability of the para nodal junction and responsible through the transverse bands of addition of terminal myelin loops to axolemma. The paranodal junctions also act as a fence for the juxta para nodal segregation of voltage-gated potassium channels and overall contribution to the saltatory conduction of myelinated fibres.
Again, antibodies to axo-glial proteins are predominantly of the IgG 4 isotype. Of the five antibody isotypes IgG is the most abundant and is further subdivided into subclasses 1 to 4. The subclasses have different constant regions and vary in their ability to induce specific humoral or cellular effector functions. IgG1 and IgG3 are more potent activators of complement than IgG2 and IgG4. IgG4 antibodies are most frequently detected in paranodal antibody positive (PNAbs+) patients and appeared to be highly specific for the presence of an immune mediated neuropathy phenotypically distinct to seronegative CIDP. Patients with only non-IgG4 subclass PNAbs have been reported to be clinically indistinguishable from seronegative patients and may have a more favourable response to IVIG. The interested readership is advised to refer to the references on this intricate and emerging interesting immuno phenotyping and immune pathogenesis of the paranodal neuropathies.

Clinical feature: (TABLE ONE)

Various findings do increase the likelihood of detecting a paranodal/nodal antibody. These are mostly adult patients presenting with a severe and symmetrical motor predominant and distal predominant polyneuropathy with a more rapidly progressive disease than those with seronegative CIDP and the peak clinical severity was also significantly greater. Hence the initial impression of Guillain-Barre syndrome was significantly more frequent in this group. In the subsequently chronically progressive or relapsing remitting disease with characteristic electrophysiology, patients met the criteria for definitive CIDP. However, patients had additional features of sensory ataxia and tremor. This tremor and cerebellar/sensory ataxia commonly occurred in anti NF155 seropositive patients while an aggressive onset neuropathy associated with cranial neuropathies, autonomic dysfunction and respiratory insufficiency occurred in patients with antibodies cross reacting with both neurofascin isoforms (NF155 and NF186–pan NF). Nephrotic syndrome was associated in those with CNTN1 antibodies. Underlying haematological disorders (Hodgkin's lymphoma, chronic lymphocytic leukaemia and myeloma) was found in pan NF patients characterised by the presence of an IgG Lambda para protein. This group was characteristically incompletely responsive to first-line therapies in particular IVIG and had a more sustained improvement from alternative immunomodulatory therapy such as Rituximab.

1. Diagnosis: Electrophysiology: (TABLE TWO) no definitive electrophysiology features unequivocally diagnostic of PNAbs seropositivity is present. Axonal conduction block without temporal dispersion of compound muscle action potentials characteristic of paranodal/nodal pathology can also develop; this may either rapidly resolve (reversible conduction failure) or progress to axonal degeneration. Even in such serious cases clinical recovery can be complete contrary to the general perception of an axonopathy. With the emergence of the nodo-paranodopathies conventional electrophysiological criteria for demyelinating and axonal neuropathies would be in need for revision.

2. Testing for antibodies: (TABLE THREE) Venous blood taken before treatment to avoid contamination with an external source of IgG (from intravenous immunoglobulin) or dilution/removal or suppression of the antibody by plasma exchange or immunosuppression is the source of antibodies. Recommendations are to test for antibodies against CNTN1, Caspr1, NF155 and NF186 in parallel. The value of CSF examination is not determined. In the testing laboratories a live cell-based assay is used for initial screening. Some laboratories use Western blot or fluorescence activated cell sorting.
TABLE: 1

<table>
<thead>
<tr>
<th>Localisation</th>
<th>Antibodies</th>
<th>Associated neuropathy</th>
<th>Notable phenotypic features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Node</td>
<td>Gangliosides</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>Glycolipids</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>NGF 116</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Paraneuronal</td>
<td>Gangliosides</td>
<td>Anti-GQ1b Abs suggest good treatment response/prognosis</td>
<td>NTF18s and glycolipid Abs are associated with an animal model of ADSc, and are found in patients with MMN</td>
</tr>
<tr>
<td></td>
<td>NF 155</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>CNTN1</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Corporate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Juxtaparanodal</td>
<td>CNTN2/TAG1</td>
<td></td>
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<td></td>
<td>Corporate2</td>
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</tbody>
</table>

Characteristics suggesting a paranodal/nodal antibody-positive (PNAb+) neuropathy

**Clinical**

<table>
<thead>
<tr>
<th>Symptoms/signs</th>
<th>Electrophysiological [38, 39]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute/subacute onset</td>
<td>Reversible conduction failure</td>
</tr>
<tr>
<td>Severe nadiq disability</td>
<td>Reduced motor conduction velocity</td>
</tr>
<tr>
<td>Predominant distal motor weakness</td>
<td>Prolonged distal motor latencies</td>
</tr>
<tr>
<td>Severe sensory ataxia</td>
<td>Conduction blocks</td>
</tr>
<tr>
<td>Tremor</td>
<td>Decreased CMAP amplitudes</td>
</tr>
<tr>
<td>Neuropathic pain</td>
<td>Early axonal loss</td>
</tr>
<tr>
<td>Cranial neuropathies</td>
<td></td>
</tr>
<tr>
<td>Autonomic dysfunction</td>
<td></td>
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<tr>
<td>Respiratory insufficiency</td>
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</tr>
</tbody>
</table>

**Associated disorders**

Concurrent nephrotic syndrome

IgG paraprotein lymphoproliferative disorders

**Treatment response**

No/transient response to intravenous immunoglobulin or plasma exchange

Good response to rituximab and/or corticosteroids

**Histological**

Nerve root/plexus enhancement

Axonal degeneration enlargement

Nodal widening

Paranodal myelin loop detachment

No overt inflammation or segmental demyelination

CMAP, compound muscle action potential; IgG, immunoglobulin G


TABLE: 2

Nodo-Paranodopathy: Beyond the Demyelinating and Axonal Antibody-Mediated Neuropathies

**Table: 3**

<table>
<thead>
<tr>
<th>Ganglioside antibodies</th>
</tr>
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<tbody>
<tr>
<td>IgG class GQ1b antibodies help identify complex ataxic/brainstem syndromes as inflammatory, providing diagnostic reassurance and support for the use of immunomodulatory therapy.</td>
</tr>
<tr>
<td>Testing for other antiganglioside antibodies in patients with GBS helps differentiate pathological subtypes but does not add to prognostication, and, as yet, does not support the use of non-standard therapies.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Paranodal protein antibodies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antibodies against the nodal proteins NF155, CNTN1 and Caspr aid diagnosis of inflammatory neuropathies with atypical features.</td>
</tr>
<tr>
<td>Consider testing for;</td>
</tr>
<tr>
<td>nNF155 antibodies in the context of neuropathies with young age of onset, motor predominance, ataxia, tremor or additional central nervous system (CNS) involvement.</td>
</tr>
<tr>
<td>CNTN1 antibodies in neuropathies with aggressive onset, motor predominance.</td>
</tr>
<tr>
<td>Caspr antibodies in patients with rapidly progressive, motor predominant neuropathies and/or prominent neuropathic pain.</td>
</tr>
<tr>
<td>In patients with any IgG4 subclass paranodal protein antibody, IVlg is often ineffective and rituximab should be considered.</td>
</tr>
</tbody>
</table>


**Treatment**

“Seropositive patients” appear less responsive to first-line treatments for inflammatory neuropathies, and it would seem appropriate to consider initiating Rituximab early in the disease course even if there was a transient response to initial IVIG therapy likely indicating an ongoing and potentially treatment responsive immune process. The reason for transient effectiveness of IVIG is through the suppression of complement activity whereas IgG4 subclass antibodies do not fix complement and exert their pathogenic effects by other mechanisms. The effect of plasma exchange is more difficult to judge. Since Rituximab is an anti-CD20 monoclonal antibody it targets immature CD20+ B cells rather than mature plasma cells. The immature CD20+ B cells are responsible for generating short lived antibody secreting cells/plasma blasts that may be primarily responsible for nodal/paranodal antibody production early on.

**Prognosis**

The disability at clinical nadir is high in paranodal antibody positive patients. However, they have the potential to achieve long lasting clinical remission. This is particularly seen in patients with fulminating neuropathy requiring intensive care with presence of pan NF antibodies who go on to have a complete recovery. Falling antibody titres are linked to clinical and electrophysiology remission facilitating more aggressive escalation of therapy. It is more difficult to prognosticate the long-term fate of patients in whom seropositivity persists.

In summary, treatment of chronic immune mediated sensorimotor neuropathies (CIN) is challenging. Up to 20% of patients do not adequately respond to first-line therapy with steroids (STE), immunoglobulins (IVIg), and plasmapheresis (PE) and require further immunotherapy. Studies on these treatments are limited. Two drugs used commonly in patients refractory to first-line treatment or with severe disease course are rituximab (RTX) and cyclophosphamide (CYP). RTX is a monoclonal anti-CD20 antibody selectively depleting premature B-cells. A total of 60 patients with CIDP with a response rate of 78% to RTX were reported in different case series. CYP has also been reported to be a sufficient immunosuppressant therapy. In a case series of 51 CIDP patients, 35 (69%) benefited from CYP therapy. A third novel treatment in refractory CIDP is bortezomib (BTZ) – a proteasome inhibitor. In a previous case series, our group described the efficacy of BTZ in 10 CIDP patients. Further reports about the effectiveness of these therapies in CIN are necessary. The aim of this study was to retrospectively analyze clinical response to treatment with CYP, RTX, and BTZ in a cohort of 200 CIN patients.
Guidelines

European Academy of Neurology/Peripheral Nerve Society guideline on diagnosis and treatment of chronic inflammatory demyelinating polyradiculoneuropathy/Paranodopathies:

Antibody testing (PICO 5) Good Practice Points • The TF suggested to consider testing for nodal and paranodal antibodies in all patients with clinical suspicion of CIDP: when nodal and paranodal (anti-NF155, anti-CNTN1, anti-Caspr1) and possibly anti-NF140/186 antibody testing is available and meeting quality standards. Testing of nodal and paranodal antibodies is advised in CIDP patients with the following features: • resistance to standard therapy with IV Ig and corticosteroids. • acute or subacute aggressive onset, previous diagnosis of GBS or A-CIDP. • low-frequency tremor, ataxia disproportionate to the sensory involvement or other cerebellar features or predominantly distal weakness. • respiratory failure and cranial nerve involvement. • associated nephrotic syndrome. • very high CSF protein levels. • The TF advised using for nodal and paranodal autoantibody testing: a cell-based assay using mammalian expression vectors encoding human NF155, CNTN1, NF186/NF140, and Caspr1. Expression vectors should avoid the use any protein tag at the N-terminal site, any protein tag at the C-terminal site for CNTN1 and avoid the use, in general, of GFP-tagged expression vectors. a confirmatory test with ELISA (using human recombinant proteins) or teased-nerve immunohistochemistry. The order of assays can be interchanged. The application of additional confirmatory tests to the protocol is strongly recommended for low titre sera or dubious staining on the cell-based assay to avoid false positives. • The TF advised anti-MAG antibody testing in all patients with an IgM paraprotein fulfilling CIDP diagnostic criteria (especially distal CIDP) because a high titre of anti-MAG antibodies (>7000 Bühmann Titre Units, BTU) 153 would strongly imply a different diagnosis than CIDP. • The TF advised for anti-MAG antibody testing: Bühmann test ELISA, or Locally validated ELISA, Western blot or immunohistochemistry assays. Considerations supporting the Good Practice Points (supporting information) Evidence summary: Data from 16 cohort studies assessing the presence of nodal-paranodal and anti-MAG antibodies were extracted and analysed. Diagnostic utility seems strong for anti-NF155 and anti-CNTN1 IgG,67,69,72 and anti-Caspr1 IgG,73–75 More evidence is needed for anti-NF155 IgM,154 anti-nodal NF140/186 IgG,76,77 and anti-MAG without an apparent paraprotein.155 For autoantibodies against CNTN1 and NF155, replication studies and a systematic review156 are available with clear associations to clinically relevant features and a high diagnostic specificity. For autoantibodies against Caspr1, nodal NF, and MAG, only small case series or anecdotal cases have been reported. Evidence that autoantibody detection may inform treatment remains anecdotal. Several case reports and case series associate the detection of nodal-paranodal antibodies, especially anti-NF155 and anti-CNTN1 with poorer responses to conventional therapies. There is anecdotal evidence that these patients may respond well to rituximab. Although the evidence is weak due to the low numbers of patients, the response to rituximab has been replicated in independent cohorts and the magnitude of the effect is, at least for a subset of patients, very significant. Rationale: Nodal-paranodal or MAG antibody testing should be considered in patients who fulfill criteria for CIDP, when they present with particular characteristics (Flowchart 1) and when they do not respond well to proven effective treatments for CIDP. Anti-MAG antibodies are relevant, if associated with a distal CIDP phenotype and an IgM paraprotein. The antibody testing has a low cost and positive results have significant implications for diagnosis and treatment. Access to antibody testing requires specialized laboratory procedures that are not available worldwide and standardization of the assays through interlaboratory validation needs to be performed. Patient burden is negligible.

Concluding points

⇒ Patients with paranodal/nodal antibodies (PNaBs) have a distinct clinical phenotype, with different electrophysiological and histological characteristics compared with seronegative chronic inflammatory demyelinating polyneuropathy (CIDP).

⇒ PNaB+ patients appear less responsive to standard first-line treatments for CIDP/Guillain-Barré syndrome (GBS) (in particular intravenous immunoglobulin), and are likely to respond to rituximab clinically and serologically, although this is currently based on low-quality evidence.

⇒ Patients with an acute, subacute or ‘atypical’ (including treatment resistant) presentation of suspected GBS or CIDP should be prioritised for PNaB testing, and we advocate early testing of all patients with suspected immune-mediated neuropathies.

⇒ Prolonged suppression of antibody titres may be associated with sustained clinical remission, but the reverse is not necessarily true; we need more data to establish the prognostic implications and clinical utility of serial antibody measurements.

References


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