Anti–pan-neurofascin IgG3 as a marker of fulminant autoimmune neuropathy

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Abstract

Objective
To identify and characterize patients with autoantibodies against different neurofascin (NF) isoforms.

Methods
Screening of a large cohort of patient sera for anti-NF autoantibodies by ELISA and further characterization by cell-based assays, epitope mapping, and complement binding assays.

Results
Two different clinical phenotypes became apparent in this study: The well-known clinical picture of subacute-onset severe sensorimotor neuropathy with tremor that is known to be associated with IgG4 autoantibodies against the paranodal isoform NF-155 was found in 2 patients. The second phenotype with a dramatic course of disease with tetraplegia and almost locked-in syndrome was associated with IgG3 autoantibodies against nodal and paranodal isoforms of NF in 3 patients. The epitope against which these autoantibodies were directed in this second phenotype was the common Ig domain found in all 3 NF isoforms. In contrast, anti–NF-155 IgG4 were directed against the NF-155–specific Fn3Fn4 domain. The description of a second phenotype of anti–NF-associated neuropathy is in line with some case reports of similar patients that were published in the last year.

Conclusions
Our results indicate that anti–pan-NF-associated neuropathy differs from anti–NF-155–associated neuropathy, and epitope and subclass play a major role in the pathogenesis and severity of anti–NF-associated neuropathy and should be determined to correctly classify patients, also in respect to possible differences in therapeutic response.

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The node of Ranvier has long been considered a potential site of attack in inflammatory neuropathies. With respect to the concept of complement-mediated reversible conduction block induced by antiganglioside autoantibodies, the term “paranodopathy/nodopathy” was introduced by Uncini et al.1,2 In the past few years, non–complement-fixing IgG4 autoantibodies against paranodal proteins were described in patients with immune neuropathies and were also comprised under that term.3,4 Patients with autoantibodies against paranodal proteins present with a distinct clinical phenotype of subacute-onset severe sensorimotor neuropathy with poor response to intravenous immunoglobulins but excellent response to rituximab.5–10 Autoantibodies are mostly of the IgG4 subtype, but IgG3 and IgG2 autoantibodies have also been described, possibly associated with early disease or monophasic course.9,10 In these cases, complement deposition could be demonstrated.11 Anti-neurofascin (NF)-155 appears to be the most prevalent paranodal autoantibody and is strongly associated with tremor.6,12,13 NF-155 is located at the paranodes, whereas NF-186 and -140 are nodal isoforms, the latter mainly expressed during embryonic development.14,15 Although all 3 isoforms share the Ig domain and some parts of the fibronectin domain,16 most autoantibodies described so far were selectively directed against NF-155.6,8 However, 2 recent studies could also detect anti-NF-186/140 autoantibodies, mostly of the IgG4 subclass, in patients with chronic inflammatory demyelinating polyradiculoneuropathy (CIDP).17,18 Single cases of patients with autoantibodies against all isoforms and severe neuropathy have been described.17,19 In contrast to anti–NF-155-associated neuropathy, not much is known about neuropathy with anti–pan-NF autoantibodies.

In the present study, we aimed to identify and characterize patients with autoantibodies against NF to potentially establish an anti–pan-NF-associated neuropathy as a distinct clinical phenotype.

Methods

Patients

Three different subcohorts of patients were included: sera of 182 patients with clinically suspected inflammatory neuropathy that were sent to our department for testing of paranodal autoantibodies, further referred to as “comprehensive diagnostic testing cohort,” sera of 306 patients with a differential diagnosis of inflammatory neuropathy who attended our department for diagnostic workup (32 finally diagnosed as Guillain-Barré syndrome (GBS), 81 as CIDP fulfilling the European Federation of Neurological Societies criteria,20 and 193 as noninflammatory neuropathy or suspected inflammatory neuropathy not fulfilling the EFNS criteria; “Würzburg cohort”), and 181 sera of a patient cohort from the University Hospitals of Kiel and Magdeburg that were collected between 2004 and 2016 (146 GBS, 21 CIDP, and 15 Miller-Fisher syndrome/Bickerstaff encephalitis; “Kiel/Magdeburg cohort”). Diagnosis of GBS was assessed by the Brighton criteria.21 CSF was obtained from diagnostic lumbar puncture. The study was approved by the Ethics Committees of the Universities of Würzburg and Kiel. Demographic data are summarized in table 1.

Enzyme-linked immunosorbent assay

All patients were screened for anti-NF-155 autoantibodies by ELISA, and 252 patients of the Würzburg cohort and all anti–NF-155-positive patients were also screened for anti-NF-186. All anti–NF-155-positive patients were tested for anti-NF-140. CSF was available from 3 seropositive patients and was tested at a dilution of 1:20. Five patients without a clinical history of neuropathy who had received lumbar puncture for other diagnostic purposes served as controls. NF-155/-186 ELISA was performed as previously described using the extracellular domains of human NF-155 and -186.8 For anti-NF-140 ELISA, wells were coated with recombinant human NF-140 protein (1 μg/mL, Sino Biological Inc, Beijing, China). Normal controls were run in each assay. The normal value was set at 5 SDs above the mean of all healthy controls that were tested in previous studies.22 Titters were determined by measuring serial dilutions of anti–NF-positive sera. Subclasses were detected using subclass-specific horseradish peroxidase-conjugated secondary antibodies (anti-IgG1: 1:1,000, anti-IgG2: 1:4,000, anti-IgG3: 1:3,000, and anti-IgG4: 1:2,000; Invitrogen, Carlsbad, CA).

Binding assays with murine teased fibers, sympathetic ganglia, cerebellar granular neurons, and rat cerebellum sections

All sera were screened for paranodal and nodal autoantibodies using binding assays with murine teased fibers as previously described.9 The preparations were incubated with patients’ sera diluted 1:100 and 1:500 overnight at 4°C following incubation with appropriate anti–human Cy3-conjugated secondary antibodies (Dianova, Hamburg, Germany, 1:100). Cervical sympathetic ganglia were dissected from adult Lewis rats, and 10-μm cryosections were cut. Binding assays with patients’ and controls’ sera were performed with a dilution of 1:100 and 1:500 by incubating overnight. Cy3-conjugated anti-human secondary antibodies were added for 2 hours. All slides were assessed using a fluorescence microscope (Zeiss Ax10, Zeiss, Oberkochen, Germany). Cerebellar granular neurons were prepared from P5 mice: the cerebellum was dissected, and cells were dissociated and suspended using

Glossary

CIDP = chronic inflammatory demyelinating polyradiculoneuropathy; GBS = Guillain-Barré syndrome; HEK = human embryonic kidney; MFI = mean fluorescence intensity; MFS = Miller-Fisher syndrome; NF = neurofascin.
a papain dissociation system (Worthington Biochemical Corporation, Lakewood, NJ). Cells were plated on cover slips at a density of 200,000 cells per well and fixed with 4% paraformaldehyde after 3 days, and binding assays with sera of patients and controls were performed at a dilution of 1:500 using an anti-human Cy3-conjugated secondary antibody (Dianova, 1:300). Anti–pan-NF served as a positive control.

Binding assays on rat cerebellum sections were performed overnight at a serum dilution of 1:200 or CSF dilution of 1:4 on 6-μm sagittal adult rat brain cryosections optimized for detection of surface antigens as has been previously described.23

**Binding assays with NF-transfected human embryonic kidney (HEK) 293 cells**

Binding assays with HEK293 cells transiently transfected with plasmids of human NF-155 and -186 were performed as previously described.22 Sera were used at a dilution of 1:200 (range) and controls. After incubation with transfected HEK293 cells, sera were used for binding assays on teased fibers as described above.

**Preabsorption experiments**

Preabsorption was performed by serial incubation of patients’ sera (1:500) 1 hour each in 3 wells of a 24-well plate with NF-155/-186-transfected HEK293 cells as described before.9 Sera incubated in wells with contactin-1–transfected cells served as controls. After incubation with transfected HEK293 cells, sera were used for binding assays on teased fibers as described above.

**Epitope characterization**

TE671 human rhabdomyosarcoma cells stably transfected with NF-155, NF-186, or 6 truncated forms of NF-155 and NF-186 were previously described.8 Cells were incubated with patients’ and controls’ sera (1:100). Detection of binding was performed as described,8 with some modifications: Serum IgG binding was detected using Biotin-SP-conjugated AffiniPure F(ab’)2 antibodies (Jackson ImmunoResearch) and AF647-conjugated streptavidin (Jackson ImmunoResearch). Mean fluorescence intensity (MFI) of the AF647 channel was calculated after gating on enhanced green fluorescent protein (EGFP) high cells for NF truncations and empty EGFP vector–transfected cells, and delta MFI was calculated from these values. Patient 5 was only tested for Fn3F4, as the sample was obtained at a later time point and NF-155 isoform specificity had already been shown by ELISA.

**Complement binding assay**

ELISA plates were coated, blocked, and incubated with patients’ sera as described above. The next day, the wells were incubated with C1q (10 μg/mL, Sigma-Aldrich, St. Louis, MO) for 2 hours at room temperature, following incubation with horseradish peroxidase–conjugated anti-C1q (LifeSpan Biosciences, Seattle, WA, 1:200) for 30 minutes. Measurement of optical density was made as described above.

**Data availability statement**

Any data not published within the article will be shared by request from any qualified investigator in anonymized form.

**Results**

**Detection of anti-NF autoantibodies**

Anti-NF-155 autoantibodies were detectable by ELISA in sera of 6 patients (figure 1). Titters ranged from 1:300 to 1:6,000 (table 2). Three of the 6 anti–NF-155-positive patients were also positively tested for anti-NF-186 and -140 (patients 1–3, table 2). Sera of 2 patients showed an optical density slightly above the cutoff value in the anti-NF-155 ELISA, but a second serum sample of these patients obtained during the course of disease was clearly negative. Of note, in none of the 252 neuropathy cases tested for anti-NF-186 independent of NF-155 positivity, did we observe isolated anti-NF-186 positivity. Anti-NF-155 and/or -186 autoantibodies were confirmed by binding assays with NF-155– and NF-186–transfected HEK293 cells in 5 of these patients (patients 1–5, figure 1). Binding assays on murine teased fibers revealed a distinct

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**Table 1 Demographic data of patients of all subcohorts**

| Würzburg cohort (n = 306) | Kiel/Magdeburg cohort (n = 181) | Comprehensive diagnostic testing cohort (n = 182) |
|---------------------------|---------------------------------|-----------------------------------------------|
| Median age (range)        | 62 (8–84)                       | 59 (18–87)                                    | 59 (18–84)                                   |
| Sex                       | 218 male, 88 female             | 91 male, 90 female                            | 136 male, 46 female                          |
| Diagnosis                 |                                 |                                               |                                               |
| GBS                       | 32 (22 level 1, 6 level 2, 3 level 3, and 1 level 4) | 146 (75 level 1, 41 level 2, 6 level 3, and 24 level 4) | NA                                           |
| CIDP                      | 81                              | 21                                            | NA                                           |
| Others                    | 193                             | 14 MFS, 1 Bickerstaff                         | NA                                           |

**Abbreviations:** CIDP = chronic inflammatory demyelinating polyradiculoneuropathy; GBS = Guillain-Barré syndrome; MFS = Miller-Fisher syndrome.
paranodal staining in the patients with anti-NF-155 auto-
antibodies only (patients 4–6; figure 2, F–H) and a nodal
and paranodal staining in the 3 patients with anti–pan-NF
autoantibodies (patients 1–3; figure 2, A, C, and D). In
patient 1, paranodal staining was stronger than nodal
staining, and in patients 2 and 3, nodal staining was
stronger (Figure 2A, C–E). In patients 3 and 6, the fluo-
rescence signal was weaker than in the other patients, most
probably due to the low titer (Figure 2E, H). In patients 1
and 2, paranodal binding became stronger during the
course of disease (Figure 2B, D).

Preincubation assays were performed to confirm paranodal/
nodal binding as anti-NF specific. In the 2 patients who were
only positive for anti-NF-155 (patients 4 and 5), paranodal
binding was lost after preincubation with NF-155–transfected
cells, not after incubation with NF-186–transfected cells. In 2
of the patients with autoantibodies against both isoforms
(patients 1 and 2), paranodal and nodal binding were lost after
incubation with NF-155-transfected cells, but weaker para-
nodal binding persisted after incubation with NF-186–transfected
cells, suggesting the presence of at least 2 auto-
antibodies with different epitopes (supplemental figure 1,
links.lww.com/NXI/A132, and figure 3).

CSF was available from patients 2, 3, and 6. No anti-NF
autoantibodies were detectable in CSF by ELISA in patients 3
and 6. In patient 2, no anti-NF autoantibodies were detectable
in CSF by ELISA at the onset of disease (July 2014) or after 2
months (September 2014). However, 6 months after the
onset of disease, anti-NF autoantibodies were detectable,
most probably due to disruption of the blood-brain barrier
(CSF/serum albumin index 37.2 × 10⁻³). There was no evi-
dence of intrathecal IgG production at this time point, as no
oligoclonal bands were detectable and the CSF IgG index was
within the normal range (0.59).

Comparison of clinical phenotypes in patients
with pan-NF or NF-155–specific antibodies
All 6 anti–NF-positive patients had a clinical diagnosis of
CIDP except for patient 3 who had a monophasic course and
was classified as GBS. Yet detailed analysis of clinical pictures
showed that anti–NF-positive patients segregated into 2 dis-
tinct phenotypes.

Patients 1–3 who were seropositive for anti-NF-155, -NF-186,
and -NF-140 had a fulminant course of disease. After GBS-
like onset, patients 1–3 quickly developed tetraplegia and
severe cranial nerve involvement. No antiganglioside

Figure 1 Binding assays with NF-155– and NF-186–transfected HEK293 cells (A–G) and ELISA (H)

Patients 1, 2, and 3 show a clear binding to NF-
155– and NF-186–transfected cells (A–C),
whereas patients 4 and 5 only bind to NF-155–
transfected and not to NF-186–transfected cells
(F and G). Healthy control serum does not bind
to the cells (E), and binding of commercial
anti–pan-NF autoantibodies to NF-155– and NF-
186–transfected cells is shown in (D). NF =
neurofascin; OD = optical density.
| Patient | 1       | 2       | 3       | 4       | 5       | 6       |
|---------|---------|---------|---------|---------|---------|---------|
| ELISA   | NF-155, -186, -140 positive | NF-155, -186, -140 positive | NF-155, -186, -140 positive | Only NF-155 positive | Only NF-155 positive | Only NF-155 positive |
| ELISA titer (onset) | 1:4,000 (NF-155), 1:2,000 (NF-186) | 1:1,000 (NF-155), 1:500 (NF-186) | 1:300 (NF-155), 1:200 (NF-186) | 1:6,000 (NF-155) | 1:2,500 (NF-155) | 1:500 (N-155) |
| IgG subclass (anti-NF-155 ELISA) | IgG3 | IgG3 (course of disease: weak IgG4) | Weak IgG3 | IgG4>IgG2 | IgG4 | IgG3 |
| Cell-based assay | NF-155, -186 positive | NF-155, -186 positive | NF-155, -186 positive | NF-155 positive | NF-155 positive | Negative |
| Truncated NF (flow cytometry) | 6Ig domain > Fn3 | 6Ig domain >> Fn3 | 6Ig domain | Fn3Fn4 | Fn3Fn4 | NA |
| IgG subclass (flow cytometry) | IgG3 | IgG3 | Weak IgG3 | IgG4 | IgG4 | NA |
| Teased fiber binding | Paranodal >> nodal | Nodal > paranodal | Weak nodal and paranodal | Paranodal | Paranodal | Weak paranodal |

**Summary of autoantibody finding**

| Cohort | Compl. diag. testing | Compl. diag. testing | Würzburg | Compl. diag. testing |
|--------|----------------------|----------------------|----------|----------------------|
| Diagnosis | CIDP | CIDP | GBS | CIDP | CIDP | CIDP |
| Age/sex | 71, male | 63, male | 74, female | 28, male | 18, male | 49, male |
| CSF (cell count/protein) | <1/μL/907 mg/L | 2/μL/621 mg/L | <1/μL/298 mg/L | 4/μL/2011 mg/L | 3/μL/4,400 mg/L | 3/μL/616 mg/L |
| Autonomic symptoms | Cardiac resuscitation | Severe blood pressure instability, paralytic ileus | No | Orthostatic hypotension, pollakiuria, hyperhidrosis | No | No |
| Cranial nerve involvement | Almost complete, locked-in | Almost complete, locked-in | Dysphagia | No | No | No |
| Respiratory insufficiency | Ventilated | Ventilated | No | No | No | No |
| Motor symptoms | Tetraplegia | Tetraplegia | Almost tetraplegia | Distal > proximal paresis, tremor | Mild distal paresis, tremor | Severe distal > proximal tetraparesis |
| Sensory symptoms | Pallanesthesia, distal hypesthesia | Hyperesthesia | Hypesthesia | Neuropathic pain | Mild distal hypesthesia | Hyperesthesia |
| Concomitant diseases | Cervical myelopathy | Cervical myelopathy, bullous pemphigoid | Chronic lymphatic leukemia | No | No | No |
| Response to treatment | None | Glucocorticoids? | Plasma exchange | Mild improvement on IVIG, PE, and rituximab | Response to PE and rituximab | Mild response to IVIG |
| Outcome | Died due to sepsis 10 mo after onset | Complete recovery of cranial nerves, residual paresis of legs and arms (October 2018) | Complete recovery | Only minor improvement | Improvement of tremor | Wheelchair user, distal paresis, and hypesthesia (February 2009) |

Abbreviations: CIDP = chronic inflammatory demyelinating polyradiculoneuropathy; GBS = Guillain-Barré syndrome; IVIG = intravenous immunoglobulins; NA = not available; NF = neurofascin; PE = plasma exchange.
antibodies were detectable. Patient 3 quickly recovered within a few months. Patients 1 and 2 needed mechanical ventilation for several months with complete tetraplegia and were almost locked-in. Patient 1 died due to sepsis, and patient 2 had a very protracted course with almost 2 years in tetraplegia and still recovering almost 5 years after onset of symptoms.

Patients 4 and 5 were young men with anti-NF-155 autoantibodies and presented with motor more than sensory neuropathy associated with tremor, as described in multiple reports.6,13 Remarkably, autonomic involvement was a major problem in patient 4. In patient 5, action and postural tremor and gait ataxia were the dominant symptoms with only minor neuropathic complaints and no autonomic symptoms.

In patient 6 anti-NF-155 autoantibodies could not be confirmed by cell-based assay. The patient had severe sensorimotor neuropathy without tremor.

For detailed clinical data, course of disease, and response to treatment, see Table 2 and Supplements (supplement course of disease, links.lww.com/NXI/A137).

**Different patterns of reactivity of autoantibodies against truncations of NF**

To find out whether the autoantibodies in patients 1–3 were directed against different epitopes or target a common epitope of all NF isoforms and were thus responsible for the positive reactivity to all 3 NF, binding assays with truncated NF mutants were performed. Patients 1–3 showed immunoreactivity against the Ig domains of NF that are part of all isoforms. In addition, patients 1 and 2 were found to be weakly immunoreactive to Fn3, a component of the NF-155 isoform. Immunoreactivity to Fn3 as a second NF-155–specific epitope is in line with the results of the preincubation assays, explaining persistence of weak paranodal binding after preincubation with NF-186–transfected HEK293 cells. Additional immunoreactivity to Fn3 as a NF-155–specific epitope also accounts for higher titers of anti-NF-155 compared with anti-NF-186. Patients 4 and 5 were immunoreactive against Fn3Fn4, a component of the fibronectin domain of NF-155 that is isoform specific (figure 3).

**IgG subclasses and C1q binding**

ELISA with subclass-specific secondary antibodies detected IgG4 and IgG2 as the predominant subclasses in patient 4, IgG4 in patient 5, and IgG3 in patients 1, 2, and 6. In patient 1, IgG2 was detectable at the onset of disease. No specific subclass was detectable in patient 3, most probably due to the low titer and lower sensitivity of the subclass-specific ELISA (figure 4). In patient 2, a minor titer of IgG4 was transiently detectable during the course of disease at a time when IgG3 titers peaked at extraordinarily high titers (supplemental table, links.lww.com/NXI/A133). Results of subclasses were confirmed by flow cytometry. Complement deposition induced by autoantibody binding was found in patients 1, 2, and 4 but not in patients 3 and 6 (both with low autoantibody titers and patient 6 without confirmation of anti-NF-155 by cell-based assays) and patient 5 (with IgG4 only) (figure 4). In patients 1 and 2, sera of different time points of disease with different autoantibody titers were measured, and in total, C1q deposition correlated with IgG3 immunoreactivity (p = 0.009, r = 0.72).

**Autonomic and cranial nerve involvement in anti-NF-positive patients and binding to the cerebellum**

Autonomic involvement was present in 3 of the patients. Patients 1 and 2 developed severe autonomic dysfunction during intensive care treatment. Patient 1 was resuscitated after cardiac arrest. Patient 4 had disabling orthostatic dysfunction, increased transpiration, and polyuria. As there is no obvious link between NF and the autonomic nervous
system and postganglionic autonomic fibers are unmyelinated, binding assays with patients’ sera on sympathetic cervical ganglia were performed, but we did not detect any specific binding of patients’ sera to autonomic neurons (data not shown). To assess binding of anti–NF-positive sera to preganglionic B-fibers, we performed binding assays with anterior root teased fibers of rats that contain preganglionic sympathetic B-fibers. B-fibers are distinguishable from A-alpha- and A-gamma-fibers by their diameter. Binding of patients’ sera to ventral root fibers of all diameters (supplemental figure 3, links.lww.com/NXI/A135) and also to A-beta fibers of the dorsal root was detected. As former studies have proposed binding of anti-NF sera to the cerebellum as a possible correlate of tremor,6,13 we also performed binding assays with cerebellum sections that showed clear binding to the molecular and granular layer in patients 1–4 and to the molecular layer only in patient 5 and no cerebellar binding in patient 6. Of note, strong cerebellar binding was observed in patient 2 using CSF, which was not evident in patients 3 and 6 (figure 5). Positive binding to cerebellar granular neurons was also detectable in these patients (supplemental figure 3, links.lww.com/NXI/A135).

Cranial nerve involvement was detectable in the 3 patients with pan-NF autoantibodies. Patients 1 and 2 were almost locked-in with complete facial and oculomotor palsy and dysphagia. Patient 3 had dysphagia requiring a percutaneous endoscopic gastrostomy tube. As concomitant glomerulonephritis has been described in patients with paranodal autoantibodies,9,18,24 patients’ records were checked for renal symptoms. There was no evidence of glomerulonephritis in

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**Figure 3** Epitope mapping of anti-NF autoantibodies by binding assays to truncated NF mutants and flow cytometry

The NF isoforms consist of a common Ig6 domain and different isoform-specific fibronectin domains (A, right section). Reactivity of patient sera to NF-155 and NF-186 (A) and truncations of NF (B) by flow cytometry: patients 1, 2, and 3 show reactivity to both NF-155 (blue line) and NF-186 (orange line) compared with control (empty RSV vector, black, filled), whereas patients 4 and 5 only react to NF-155 (A). Epitope analysis using NF truncations (B) demonstrated that reactivity was against the 6Ig domain in 3 of the patients (patients 1–3) who were positive for both NF isoforms (green line). Patients 1 and 2 also reacted to the Fn3 domain (red line). The Fn3Fn4 domain (yellow line) was recognized by the sera of patients 4 and 5, who reacted only to NF-155. Other truncated NF variants (Fn1Fn2 = dark blue; Fn4-Mucin-Fn5 = brown; Mucin-Fn5 = purple; GFP-only control = gray filled) were not recognized by any of the sera tested (sera 1–4 and 6 tested for all mutants, serum 5 for Fn3Fn4 only). NF = neurofascin.
patients 2–6, but patient 1 had acute-on-chronic renal failure with mild proteinuria and hematuria since the onset of disease, so concomitant glomerulonephritis is possible, although it was not formally diagnosed.

**Discussion**

In the present study, we identified 5 patients with definite anti–NF-associated neuropathy, 3 of them presenting with a distinct clinical phenotype of tetraplegia and cranial nerve involvement. Our data suggest that the clinical picture is associated with the IgG subclass and epitope—IgG4 autoantibodies against the Fn3Fn4 domain of NF-155 being associated with the well-known phenotype of severe sensorimotor neuropathy with tremor and IgG3 autoantibodies against the Ig domain of all isoforms leading to tetraplegia and cranial nerve involvement.

We describe multiple clinically uniform cases of this more severe form of anti–pan-NF-associated neuropathy, and the distinct clinical phenotype is strengthened by identification of a common pattern of reactivity to the 6Ig domain that is common to all NF isoforms. The notion of anti–pan-NF-associated neuropathy as a distinct subtype of anti-NF neuropathy is supported by reports of single cases similar to our 3 patients in the last years: In a cohort by Burnor et al., 1 patient with autoantibodies against NF-155, -186, and -140 clinically very much resembled our patients but had IgG4 subclass autoantibodies. Only recently, another patient with tetraplegia, respiratory failure, almost locked-in syndrome, and IgG3 autoantibodies against NF-186/-140 and also NF-155 was reported, and ultrastructural analysis of nodal architecture revealed occlusion of the nodal gap by myelin layers.

In a recent study, 5 patients with anti-NF-186/-140 autoantibodies were compared with patients with anti-NF-155 IgG autoantibodies. Anti–NF-186/-140-positive patients were found to be clinically different from anti-NF-155 patients, and 2 anti–NF-186/-140-positive patients presented with cranial nerve involvement and respiratory failure, one of them with predominance of the IgG3 subclass. In contrast to our study, patients with pan-NF antibodies of the IgG4 subclass were also described in these studies, so possibly the epitope appears to be the major determinant of the clinical picture and not the IgG subclass. However, these subclasses may still contribute to the acuteness or other clinical aspects of the disease. This suggests that the unique clinical picture of our patients 1–3 may be mostly explained by the unique epitope—a common domain of nodal NF-186/-140 and paranodal NF-155—but may also be associated with the IgG3

**Figure 4** ELISA of total IgG and IgG subclasses (bars) and C1q deposition in the complement binding assay (line)

IgG3 can be identified as the dominant subclass in patients 1, 2, and 6, IgG 2 and 4 in patient 4, and IgG4 in patient 5. Complement deposition can be measured in patients 1 and 2 with IgG3 predominance and in patient 4 with IgG2 and IgG4 autoantibodies. No complement deposition was detectable in patient 5 with predominance of IgG4 and patients 3 and 6 with low titers of total IgG. IgG = immunoglobulin G.
subclass. In contrast to IgG4, IgG3 autoantibodies activate complement and are considered to have a strong proinflammatory effect. Indeed, C1q binding induced by anti–pan-NF autoantibodies could be shown in our patients by complement binding assays. Remarkably, complement deposition was also detectable in one of the patients with the typical anti-NF-155 IgG4 phenotype, most probably due to additional IgG2 autoantibodies that were detectable in this case. Complement deposition may contribute to the more severe acute course of disease in this patient compared with the other anti-NF-155 IgG patient in our cohort. Complement-mediated conduction block may either be reversible, as demonstrated by the case of patient 3 with GBS, or result in secondary axonal damage or nodal alterations (as demonstrated in an ultrastructural case description by Vallat et al.19) with a prolonged phase of recovery or lack of recovery as in patients 1 and 2. Pathophysiologic studies including passive transfer need to be conducted to further unravel the connection between clinical phenotype and autoantibodies in this subgroup of autoimmune neuropathy. CNS axonal damage and deterioration of clinical severity of experimental allergic neuritis induced by autoantibodies against the common domain of NF-155 and NF-186 were demonstrated by injection of pan-NF antibodies to rats indicating a pathogenic effect of autoantibodies against the common domain of NF.8,26

Patient 6 who was anti–NF-155 positive (IgG3) by ELISA did not present with the typical clinical phenotype of anti–NF-155-associated neuropathy. Lack of binding in the cell-based assay, cerebellar binding studies, and of complement deposition and only weak nodal/paranodal binding in teased fibers may reflect a low affinity of autoantibodies to the in vivo conformation of NF or even a protein associated with NF in this patient and may explain the mild phenotype.

Whereas all previous case reports of patients with pan-NF autoantibodies only described nodal but not paranodal binding on teased fibers,18,19 clear paranodal binding was found in all our anti–pan-NF patients. Stronger paranodal binding may be explained by additional binding to Fn3, part of the NF-155–specific fibronectin domain, in 2 of our patients. Remarkably, these 2 patients showed the most severe clinical phenotype and a very long course of disease. Thus, multiple epitopes may be a predictor of disease severity as also described in other autoimmune diseases.27,28 The underlying mechanism of multiple epitopes remains unclear: intramolecular epitope spreading could be an explanation and has also been described in other autoimmune diseases.27 Indeed, in patient 2, the Fn3 epitope was only detectable during the course of disease but not at disease onset. Paranodal binding in teased fiber assays increased respectively.

Tetraplegia was the dominant symptom in all of our patients with anti–pan-NF autoantibodies and in single cases described in the literature.17,29 We did not find any tremor in these patients, but tremor may be masked by tetraplegia. Tremor in anti-NF-155 neuropathy has been attributed to autoantibody binding to cerebellar neurons,6 and we could show cerebellar binding of anti-NF-155 and also anti–pan-NF...
serum. Our study is the first to detect paranodal autoantibodies in CSF: Anti-pan-NF autoantibodies were detectable in the CSF of patient 2 as a potential correlate of cerebellar involvement. In comparison to the 5 patients with anti-NF-186/-140 autoantibodies described by Delmont et al.,18 that also bound to the Ig domain, the course of disease was more severe in our patients and refractory to treatment. Lack of tremor, cranial nerve involvement, and older age compared with anti-NF-155 patients are features that were also found in the patients by Delmont et al.18 Differences may be explained by IgG3 subclass in our patients and mostly IgG4 in Delmont’s patients.

Remarkably, 3 of our patients including a patient with only anti-NF-155 autoantibodies had severe autonomic involvement, a feature that has not yet been investigated in anti–NF-associated neuropathy. However, autonomic symptoms were also described in the single anti–pan-NF-positive patient in the study by Burnor et al.17 As postganglionic autonomic fibers are unmyelinated and we could not show any binding to sympathetic ganglia, this can most probably be explained by radiculitis involving preganglionic autonomic fibers.

The role of IgG subclasses in the pathogenicity of anti-NF autoantibodies and particularly association with therapeutic response needs to be further studied. Detection of complement binding in our study and by others argues for complement-mediated inflammation, which extends the concept of anti–NF-associated neuropathy as a pure IgG4-mediated disease. Our data indicate that testing of IgG subclasses should be performed when detecting autoantibodies against paranodal proteins, and testing of anti-NF autoantibodies should not be restricted to the NF-155 isoform.

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|-----------------------|-----------------------------------------------|-------------------------------------------------------------------------------|
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| Atay Vural, MD, PhD   | Biomedical Center and University Hospital     | Data acquisition; drafting/revising the manuscript; and analysis or interpretation of data |
| Anna-Michelle Brunder | University Hospital Würzburg                  | Data acquisition                                                               |
| Annika Heinius         | Universitätsklinikum Schleswig-Holstein Campus Kiel, | Data acquisition and drafting/revising the manuscript                          |
| Luise Appeltshauser, MD| University Hospital Würzburg                  | Analysis or interpretation of data and acquisition of data                   |
| Bianca Fiebig          | University Hospital Würzburg                  | Analysis or interpretation of data and acquisition of data                   |
| Florian Giese, MD     | University Hospital Halle                     | Drafting/revising the manuscript                                              |
| Christian Dresel, MD  | University Hospital Mainz                     | Acquisition of data and drafting/revising the manuscript                     |
| Alkaterini Papagianni, MD| University Hospital Würzburg               | Drafting/revising the manuscript and acquisition of data                   |
| Frank Birklein, MD, PhD| University Hospital Mainz                     | Data acquisition and drafting/revising the manuscript                          |
| Joachim Weis, MD      | RWTH Aachen University                        | Data acquisition; drafting/revising the manuscript; analysis or interpretation of data; and contribution of vital reagents/tools/patients |
| Tessa Huchtemann, MD  | University Hospital Magdeburg                 | Data acquisition                                                               |
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| Peter Körtvelyessy, MD| University Hospital Magdeburg                 | Data acquisition                                                               |
| Carmen Villmann, PhD  | University Hospital Würzburg                  | Drafting/revising the manuscript; study concept or design; and contribution of vital reagents/tools/patients |
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