Antiparanodal antibodies and IgG subclasses in acute autoimmune neuropathy

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Abstract

Objective

To determine whether IgG subclasses of antiparanodal autoantibodies are related to disease course and treatment response in acute- to subacute-onset neuropathies, we retrospectively screened 161 baseline serum/CSF samples and 66 follow-up serum/CSF samples.

Methods

We used ELISA and immunofluorescence assays to detect antiparanodal IgG and their subclasses and titers in serum/CSF of patients with Guillain-Barré syndrome (GBS), recurrent GBS (R-GBS), Miller-Fisher syndrome, and acute- to subacute-onset chronic inflammatory demyelinating polyradiculoneuropathy (A-CIDP). We evaluated clinical data retrospectively.

Results

We detected antiparanodal autoantibodies with a prevalence of 4.3% (7/161), more often in A-CIDP (4/23, 17.4%) compared with GBS (3/114, 2.6%). Longitudinal subclass analysis in the patients with GBS revealed IgG2/3 autoantibodies against Caspr-1 and against anti–contactin-1/Caspr-1, which disappeared at remission. At disease onset, patients with A-CIDP had IgG2/3 anti–Caspr-1 and anti–contactin-1/Caspr-1 or IgG4 anti–contactin-1 antibodies, IgG3 being associated with good response to IV immunoglobulins (IVIg). In the chronic phase of disease, IgG subclass of one patient with A-CIDP switched from IgG3 to IgG4.

Conclusion

Our data (1) confirm and extend previous observations that antiparanodal IgG2/3 but not IgG4 antibodies can occur in acute-onset neuropathies manifesting as monophasic GBS, (2) suggest association of IgG3 to a favorable response to IVIg, and (3) lend support to the hypothesis that in some patients, an IgG subclass switch from IgG3 to IgG4 may be the correlate of a secondary progressive or relapsing course following a GBS-like onset.

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Glossary

A-CIDP = acute-onset chronic inflammatory demyelinating polyradiculoneuropathy; Caspr-1 = contactin-1-associated protein-1; CBA = cell-based assay; CIDP = chronic inflammatory demyelinating polyradiculoneuropathy; GBS = Guillain-Barré syndrome; HEK = human embryonic kidney; HLA = human leukocyte antigen; ICU = intensive care unit; IVIg = IV immunoglobulin; MFS = Miller-Fisher syndrome; NF = neurofascin; OD = optical density; R-GBS = recurrent GBS.

Autoantibodies against the paranodal antigens contactin-1, contactin-associated protein-1 (Caspr-1), and neurofascin-155 (NF155) have been described as biomarkers for a new entity of inflammatory neuropathies classified as paranodopathies.¹⁻³ In the chronic phase of disease, IgG4-seropositive patients do not respond to IV immunoglobulins (IVIg), but to rituximab.4-7 Noninflammatory IgG4 autoantibodies are pathogenic, possibly by inhibition of the interaction between contactin-1/Caspr-1 and NF155 and by NF155 depletion.⁸⁻¹¹ Autoantibodies of the IgG3 subclass have been described (1) in monophasic disease, (2) at the subacute onset, (3) in patients with anti-panneurofascin autoantibodies and severe course of disease, and (4) most recently in chronic inflammatory demyelinating polyradiculoneuropathy (CIDP) with clinical features indistinguishable from seronegative patients but with a good response to IVIg.^{6,7,11–13} Proinflammatory IgG3 antibodies lead to complement deposition in vitro¹⁴ and in vivo, resulting in reversible conduction failure in Lewis rats intraneurally injected with anti-contactin-1 IgG3¹⁵ and may therefore play a role in the acute onset of paranodopathies. However, data on antiparanodal autoantibodies in Guillain-Barré syndrome (GBS) or the acute onset of CIDP are scarce because previous studies mostly focused on CIDP and patients were recruited during the chronic phase of disease. IgG subclass distribution and the associated clinical phenotype have never been investigated longitudinally. We therefore aimed at determining the prevalence and IgG subclass of paranodal autoantibodies in a cohort of patients with acute to subacute inflammatory neuropathies including followup of seropositive patients. We hypothesize that IgG subclass and titer are related to the course of disease and therapeutic response in acute-onset paranodopathy.

Methods

Patients and controls

One hundred sixty-one patients with suspected GBS and subacute inflammatory neuropathy (peak of symptoms \leq 90 days) who had undergone diagnostic lumbar puncture at the University Hospitals of Kiel and Magdeburg between 2001 and 2016 were included into the study. We assessed clinical data retrospectively by analysis of discharge letters and documented laboratory, electrophysiologic, and MRI examinations. Table e-1, links.lww.com/NXI/A274, summarizes demographic data. The diagnosis of GBS was confirmed in n = 114 patients with diagnostic certainty according to the Brighton criteria^{16,17}: level 1 in 73 patients, level 2 in 29 patients, level 3 in 6 patients, and level 4 in 6 patients. We classified 6 patients as recurrent GBS (R-GBS) according to the criteria adapted from Kuitwaard

et al.¹⁸ and 18 patients as Miller-Fisher syndrome (MFS). In 23 patients, the initial diagnosis was GBS, but was later reverted to CIDP because of a disease progression >2 months (diagnostic certainty according to the EFNS criteria¹⁹: definite CIDP in 10 patients, probable CIDP in 3 patients, possible CIDP in 1 patient, and EFNS electrodiagnostic criteria not fulfilled in 9/23 patients). Eighteen of 23 patients with CIDP fulfilled the criteria for acute-onset CIDP (A-CIDP) proposed in previous publications, $^{20-22}$ and 5/23 patients with CIDP showed a subacute-onset (peak \leq 90 days). In the following, all patients with acute- to subacute-onset CIDP are referred to as part of the A-CIDP cohort. Follow-up sera and CSF samples were available in 66 patients, including 3 seropositive patients. We included sera of 40 healthy controls recruited in former studies.⁶ Sera of all patients had already been tested for anti-NF155 autoantibodies in a previous study.¹²

Binding assays on murine teased fibers

We used binding assays on murine teased fibers at a serum dilution of 1:100 and 1:500 to screen all patients' and controls' sera for antiparanodal autoantibodies as previously described.²³ Titers of seropositive samples were measured with a dilution series using Cy3TM-conjugated goat anti-human IgG 1:300 (Jackson, West Grove). In positive sera, we performed double immunofluorescence with sera diluted 1:50 and 1:100 and rabbit anti-Caspr-1 diluted 1:1,000 (Abcam, Cambridge, United Kingdom), using MFP488-conjugated goat anti-human IgG diluted 1:500 (MoBiTec, Göttingen, Germany) and Cy3TM-conjugated donkey anti-rabbit IgG diluted 1:300 (Jackson) as secondary antibodies. For subclass analysis, we used FITC-conjugated secondary antibodies diluted 1:100, namely mouse anti-human IgG1, IgG4 (Abcam), mouse anti-human IgG2 (Merck, Darmstadt, Germany), and sheep anti-human IgG3 (Rockland Immunochemicals, Inc., Pottstown, PA). A healthy control serum and serum of patients with anti-Caspr-1 IgG3 and IgG4 antibodies from a previous study⁷ stained on separate slides served as negative and positive controls for double immunofluorescence and subclass analysis. We tested CSF of seropositive patients of the first assessment and follow-up on teased fibers in a dilution of 1:20 and 1:50 as described above. CSF of five patients who were seronegative served as negative controls.

ELISA

We performed ELISA for detection of anti–contactin-1 in duplets with sera of all patients and 40 healthy controls as previously described.^{6,12,23,24} For anti–Caspr-1 ELISA, we coated Chinese hamster ovary cell line–derived recombinant human Caspr-1 protein (Research and Diagnostic Systems, Inc., Minneapolis, MN) at a dilution of 2 μ g/mL on MaxiSorb 96-well

plates (Thermo Scientific Fisher, Minneapolis, MN) and tested baseline/follow-up sera of seropositive patients and 40 healthy controls or rabbit anti-Caspr-1 (Lifespan Biosciences, Seattle, WA) in duplets, using corresponding horseradish peroxidaseconjugated secondary antibodies⁶ and conjugated anti-rabbit IgG (DakoCytomation, Glostrup, Denmark). For each sample, we subtracted corresponding values of uncoated duplet wells to reduce background signals. We set the threshold for both assays at 5 SDs above the mean of healthy controls. Antibody titers were determined by serial dilutions. Subclass-specific secondary antibodies served for subclass analysis as previously described.¹² We tested CSF of seropositive patients in a dilution of 1:20 in anti-contactin-1 ELISA and anti-Caspr-1 ELISA (exception: patient 4 baseline CSF in both ELISAs and patient 3 in anti-Caspr-1 ELISA due to lack of material) as described above, with CSF of 5 seronegative patients as negative controls.

Preabsorption experiments

To determine the specific paranodal target of the autoantibodies, we serially preincubated seropositive sera and serum from a healthy control with contactin-1 (*CNTN1*)– and Caspr-1 (*CNTNAP1*)–transfected human embryonic kidney (HEK) 293 cells as previously described⁷ and afterward performed binding assays on murine teased fibers as described above.

Cell-based assay

For specification, we tested sera of patients positive in teased fibers assay or ELISA and sera of five controls on HEK293 cells transfected with plasmids of human Caspr-1 (*CNTNAP1*) and rat contactin-1 (*CNTN1*) as previously described^{7,23} and used polyclonal rabbit anti–contactin-1 IgG (Abcam) and monoclonal mouse anti–Caspr-1 IgG1 (Santa Cruz Biotechnology, Dallas, TX) for double immunofluorescence in positive patients with corresponding Cy3TM-conjugated donkey anti-rabbit IgG (Abcam), donkey anti-mouse IgG, and MFP488-conjugated goat anti-human IgG (MoBiTec) secondary antibodies. Sera of patients with anti–Caspr-1 and anti–contactin-1 antibodies from previous studies^{6,7} served as positive controls and sera of healthy controls as negative controls in the double immunofluorescence assays.

Statistical analysis

We performed statistical analysis (descriptive statistics, calculation of OR, and 95% CI) with SPSS 25.0 (IBM, Armonk, NY).

Standard protocol approvals, registrations, and patient consents

The Ethics Committees of the University of Würzburg and the University of Kiel approved to conduct this study. Written informed consent was obtained from all participants in the study.

Data availability

Data not published within this article are available at the University Hospital of Würzburg or will be shared in an anonymized manner on request from any qualified investigator for purposes of replicating procedures and results.

Results

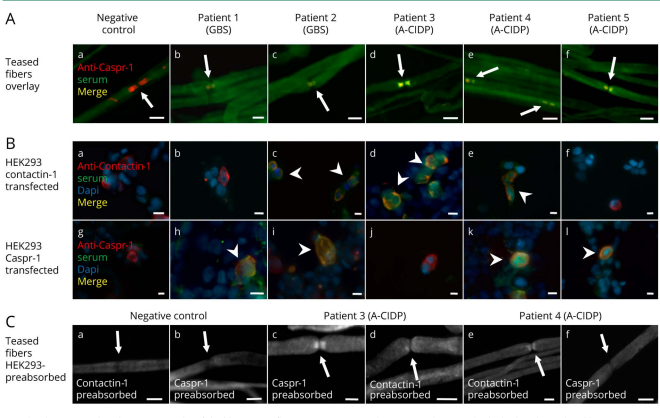
Antiparanodal autoantibodies of different IgG subclasses in 4.3% of the patients

We applied immunofluorescence binding assays on murine teased fibers as a screening tool for detection of antiparanodal autoantibodies. We observed serum binding to the paranodal junctional region colocalizing with the commercial anti-Caspr-1 antibody (figure 1A) in seven patients of the total cohort (7/161; prevalence 4.3%), in none of the patients of the R-GBS (0/6) and MFS (0/18) subcohort, nor in the healthy controls (0/40), but in 3/114 (2.6%) of the GBS subcohort and 4/23 (17,4%) of the A-CIDP subcohort. Thus, paranodal autoantibodies occurred more frequently in A-CIDP compared with GBS (OR = 7.79, 95% CI 1.61-37.95). Two of the sera with paranodal binding (1 GBS and 1 A-CIDP) had been identified as anti-NF155 seropositive in a recent study¹²; therefore, experimental findings and clinical associations of these patients are not reported here. Target antigens of the other five seropositive patients were determined using cell-based assay (CBA; figure 1B) and ELISA (see below) and confirmed by preabsorption experiments (figure 1C). In the GBS subcohort, the two seropositive patients showed anti-Caspr-1 (patient 1) and anti-contactin-1/Caspr-1 (patient 2) antibodies. Seropositive A-CIDP patients carried antibodies against either contactin-1 (patient 3), contactin-1/Caspr-1 (patient 4) or anti-Caspr-1 (patient 5). At the onset of disease, the two patients with GBS showed IgG2 and IgG3 antibodies (table 1). During the acute phase at onset, sera of the three patients later classified as A-CIDP carried IgG2/4 (patient 3) and IgG2/3 (patients 4 and 5), as detected by binding assays and ELISA (figure 2, A and B). Autoantibody titers are shown in table 1. The results of the binding assays on teased fibers, ELISA, and CBA showed concordance regarding the specific target antigen of the antibodies and the predominating IgG subclass. There were only minor differences of the antibody titer and coexisting subclasses in ELISA and teased fibers assay. In case of double positivity for anti-contactin-1/ Caspr-1 (patients 2 and 4), preabsorption assays detected only the dominating antibody (table 1).

Subclass switch from IgG3 to IgG4 at follow-up in one patient

In 66/161 patients, follow-up serum and CSF samples were available, including three seropositive patients (1 GBS and 2 A-CIDP). None of the follow-up sera of initially seronegative patients or controls showed positive results in the teased fibers assay (data not shown) nor in the contactin-1 ELISA (figure 3A). Serum of the patient with GBS acquired in remission 55 months after the first assessment, when the patient did not show any remaining neurologic deficits, was seronegative in all experimental assays (figure 3, B and C).

Figure 1 Detection of antiparanodal autoantibodies by binding assays on teased fibers, CBA, and preabsorption assay



(A) Colocalization overlay photomicrographs of double immunofluorescence assays with commercial autoantibody displayed in red and human serum in green. Colocalization at the paranodes (arrows) appears yellow and indicates antiparanodal autoantibodies in patients 1–5 (A.b.–A.f), but not in the negative control serum (A.a). (B) The specific target of the autoantibodies is illustrated by colocalization overlay photomicrographs on transfected HEK293 cells (arrowheads) with commercial antibody in red, serum in green, and nucleus staining in blue. Colocalization appears yellow (merge) and confirms anti-contactin-1 in patients 2 (B.c), 3 (B.d), and 4 (B.e), but not in patients 1 (B.b), 5 (B.f), and the negative control (B.a). Sera of patients 1, 2, 4, and 5 (B.h, B.i, B.k, and B.l) colocalize on Caspr-1 (*CNTNAP1*)-transfected HEK293 cells as proof of anti–Caspr-1 autoantibodies, whereas sera of the negative control (B.g) and patient 3 (B.j) do not show anti–Caspr-1 positivity. (C) Single immunofluorescence on murine teased fibers after HEK293 contactin-1 and Caspr-1 preabsorption shows that antiparanodal autoantibodies disappear after contactin-1 preabsorption in patient 3 (C.d) and after Caspr-1 preabsorption in patient 4 (C.f), proving specificity to the respective target antigen. Scale bar = 10 µm. CBA = cell-based assay; HEK = human embryonic kidney.

Follow-up serum of one patient with A-CIDP two months after onset, i.e., still in the acute phase of disease (patient 5), showed IgG3 autoantibodies against Caspr-1, with no difference in antibody titer and subclass compared with onset (figure 3, B and C). A follow-up-sample during the chronic stage (28 months after the onset) revealed a reduction of the titer (1:50 in teased fibers, negative in ELISA, table 1) of anti-Caspr-1 IgG autoantibodies with subclasses not measurable due to the low titer (data not shown). The patient now presented with only very mild sensorimotor impairment and ameliorated nerve conduction studies. Follow-up serum and CSF of the other patient with A-CIDP (patient 4) who was initially IgG3-anti-contactin-1/Caspr-1 double positive were acquired 120 months after the acute onset, when the patient had developed relapsing-remitting sensorimotor CIDP. At follow-up, we could no longer detect anti-contactin-1, but detected and confirmed anti-Caspr-1 autoantibodies via anti-Caspr-1 ELISA, CBA, and preabsorption assay (table 1 and figure 3, B and C). Subclass analysis now revealed reactivity against IgG2/4, but not IgG3 subclass anymore, giving evidence of a switch of IgG

subclasses between the acute and chronic stage of the disease (figure 2C).

Intrathecal antiparanodal autoantibodies in patients with high antibody titer and bloodbrain barrier dysfunction

ELISA with CSF samples of the seropositive patients and five disease controls showed elevated optical density (OD) at a dilution of 1:20 only in anti-contactin-1 patient 3 with a very high serum titer (see appendix e-1, links.lww.com/ NXI/A275) and anti-Caspr-1 patient 5 with signs of severely disrupted blood-brain barrier (high CSF protein, MRI cauda equina enhancement, see appendix e-1, links.lww. com/NXI/A275), but equally low OD values for the other seronegative and seropositive patients (see figure e-1, links. lww.com/NXI/A273). Binding assays on murine teased fibers showed weak binding of CSF to the paranodes in these two patients, but not in the other seropositive patients or controls. None of the seropositive patients showed signs of intrathecal autoantibody production, with CSF IgG index and antiparanodal autoantibody specific ASI (antibody

	CSF			Serum		
		Teased fibers and ELISA		Preabsorption	СВА	
	Paranodal	binding/OD	Titer	lgG subclass		
Patient 1 (baseline, no follow-up)						
Teased fibers	Negative	Positive	1:100	lgG3		
Contactin-1	0.202	0.465	n.d.	n.d.	Negative	Negative
Caspr-1	0.072	2.961	1:200	n.m.	Positive	Positive
Patient 2 (baseline)						
Teased fibers	Negative	Positive	1:200	n.d.		
Contactin-1	0.247	3.538	1:200	lgG2	Negative	Positive
Caspr-1	0.129	3.965	1:1,000	lgG2	Positive	Positive
Patient 2 (follow-up)						
Teased fibers	n.d.	Negative	n.d.	n.d.		
Contactin-1	n.d.	0.039	n.d.	n.d.	n.d.	Negative
Caspr-1	n.d.	0.025	n.d.	n.d.	n.d.	Negative
Patient 3 (baseline, no follow-up)						
Teased fibers	Negative	Positive	1:30,000	lgG4		
Contactin-1	0.516	4.492	1:40,000	lgG4/2	Positive	Positive
Caspr-1	n.d.	0.073	n.d.	n.d.	Negative	Negative
Patient 4 (baseline)						
Teased fibers	Negative	Positive	1:500	lgG3		
Contactin-1	0.267	2.294	1:200	lgG3	Negative	Positive
Caspr-1	n.d.*	3.011	1:500	lgG3/2	Positive	Positive
Patient 4 (follow-up)						
Teased fibers	Negative	Positive	1:500	lgG4		
Contactin-1	0.227	0.206	n.d.	n.d.	Negative	Negative
Caspr-1	0.138	3.618	1:500	lgG2/4	Positive	Positive

 Table 1
 Summary of the autoantibody status of seropositive patients 1–5 at baseline and follow-up

Continued

	CSF	Serum				
		Teased fibers and ELISA			Preabsorption	СВА
	Paranodal	binding/OD	Titer	IgG subclass		
Patient 5 (baseline)						
Teased fibers	Positive	Positive	1:100	lgG3		
Contactin-1	0.219	0.260	n.d.	n.d.	Negative	Negative
Caspr-1	0.136	n.d.*	n.d.*	n.d.*	Positive	Positive
Patient 5 (first follow-up)						
Teased fibers	Positive	Positive	1:100	lgG3		
Contactin-1	0.236	0.243	n.d.	n.d.	Negative	Negative
Caspr-1	0.364	3.802	1:200	lgG3/2	Positive	Positive
Patient 5 (second follow-up)						
Teased fibers	n.d.	Positive	1:50	n.m.		
Contactin-1	n.d.	0.168	n.d.	n.d.	n.d.	Negative
Caspr-1	n.d.	0.013	n.d.	n.d.	n.d.	Negative

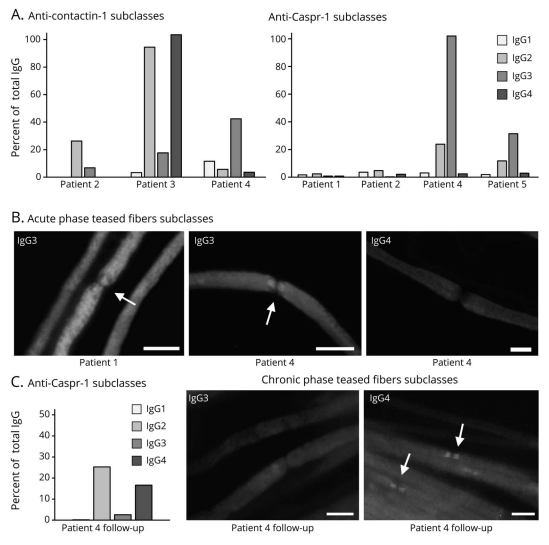
Table 1 Summary of the autoantibody status of seropositive patients 1–5 at baseline and follow-up (continued)

Abbreviations: A-CIDP = acute-onset chronic inflammatory demyelinating polyradiculoneuropathy; Caspr-1 = contactin-1-associated protein-1; CBA = cell-based assay; GBS = Guillain-Barré syndrome; n.d. = not done; n.d.* =

not done due to lack of material; n.m. = not measurable; OD = optical density. Results of teased fibers assays are displayed in the first row for each patient; contactin-1 and Caspr-1-specific results are displayed in lines 2 and 3. ODs above the threshold and results considered positive are highlighted in bold letters. There is a 100% concordance regarding target specificity in the teased fibers assay, ELISA, and CBA, but minor differences in IgG titer and coexisting subclass between the teased fibers assay and ELISA. In case of double positivity, preabsorption assays only detect the target antigen with the higher antibody titer (see ELISA titer).

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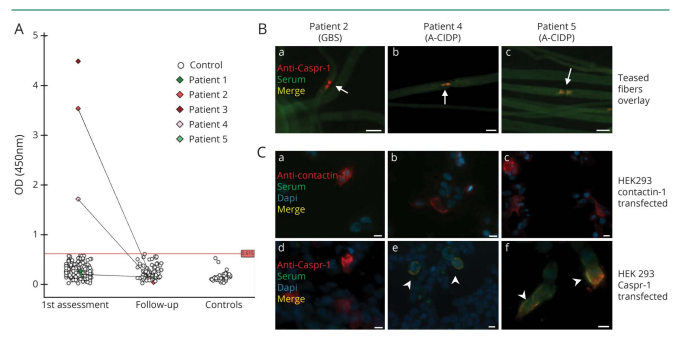
(A) Results of anti-contactin-1 and anti-Caspr-1 ELISA with baseline sera and subclass-specific IgG1-4 autoantibodies (grayscale) are shown as % of the OD of the total IgG (using anti-human IgG autoantibody) on y-axis. In patient 1, ELISA IgG subclasses were not measurable. ELISA revealed mainly IgG2 subclass in patient 2 (GB5), IgG2/IgG4 in patient 3 (A-CIDP), and IgG2/3 in patients 4 and 5 (A-CIDP). (B) Acute phase teased fibers subclass analysis with baseline sera showed binding to the paranodal regions (arrows) only when using IgG3-specific secondary antibody in patients 1 (GBS) and 4 (A-CIDP), visualized by photomicrographs of single immunofluorescence staining on teased fibers. (C) Anti-Caspr-1 IgG subclass ELISA with follow-up serum of patient 4 now revealed IgG 2/4 subclass. Photomicrographs show immunofluorescence staining of murine teased fibers with subclass-specific IgG3 and IgG4 autoantibodies at follow-up. Paranodal binding disappeared using IgG3 antibody, but occurred using IgG4 subclass-specific antibody, proving subclass switch in patient 4. Scale bar = 10 µm. A-CIDP = acute-onset chronic inflammatory demyelinating polyradiculoneuropathy; GBS = Guillain-Barré syndrome.

specificity index) being in a normal range (see appendix e-1, links.lww.com/NXI/A275).

Clinical features of seropositive patients

Figure 4 schematically summarizes the clinical course related to the experimental results. Appendix e-1, links.lww.com/ NXI/A275, summarizes clinical data of seropositive patients 1–5 and provides a detailed description of the clinical disease course in individuals. The two seropositive patients with GBS were both middle-aged women presenting with severe motor, but only mild sensory involvement. Cranial nerve involvement, autonomic symptoms, and neuropathic pain were present in 1/2 patients with GBS only. Both patients required treatment on the intensive care unit (ICU). The three patients with A-CIDP presented with moderate to severe paresis and multimodal sensory impairment at disease onset. During the course of disease, all three patients with A-CIDP reported neuropathic pain requiring pregabalin/gabapentin or even opioid treatment. Two of three patients with A-CIDP presented sensory ataxia. Nerve conduction studies showed demyelinating features in all seropositive patients with conduction block in 3/5 patients. Sural nerve biopsy was performed in one patient (patient 5) six months after the onset and showed axonal loss without signs of demyelination. CSF analysis revealed cytoalbuminologic dissociation and blood-brain barrier dysfunction in all seropositive patients. The patient with subacute-onset CIDP, highly elevated CSF protein, and detection of anti–Caspr-1 in both serum and

Figure 3 Antiparanodal IgG autoantibodies at follow-up assessment



(A) Contactin-1 IgG ELISA of all patients at first assessment, of 66/161 patients at follow-up, and of 40 healthy controls. Optical density (OD) at 450 nm is displayed on the y-axis. The threshold for positive results is set at 5 SDs above the mean of controls (0.615). Patients 2–4 show positive results at first assessment. The other patients and controls show values below the threshold. Sera of patients 2, 4, and 5 are negative in anti-contactin-1 IgG ELISA at follow-up, as well as 63 other follow-up sera. Patients 1 and 3 were lost to follow-up. (B) Overlay photomicrographs of teased fibers double immunofluorescence and (C) contactin-1 and Caspr-1 (*CNTNAP1*)-transfected HEK293 cells show loss of anti-contactin-1/Caspr-1 autoantibodies in patient 2 (GBS) at follow-up (B.a, C.a, C.d), but colocalization of commercial anti-Caspr-1 antibody and sera at paranodes in patients 4 and 5 with A-CIDP (B.b, B.c, arrows). Both sera bind to caspr-1 (*CNTNAP1*)-transfected HEK293 cells (C.e, C.f, arrowheads), but not to contactin-1-transfected cells (C.b, C.c). Scale bar = 10 µm. A-CIDP = acute-onset chronic inflammatory demyelinating polyradiculoneuropathy; GBS = Guillain-Barré syndrome; HEK = human embryonic kidney.

CSF (patient 5) developed action tremor during the course of disease.

Antibody status is concordant with disease activity and therapy response

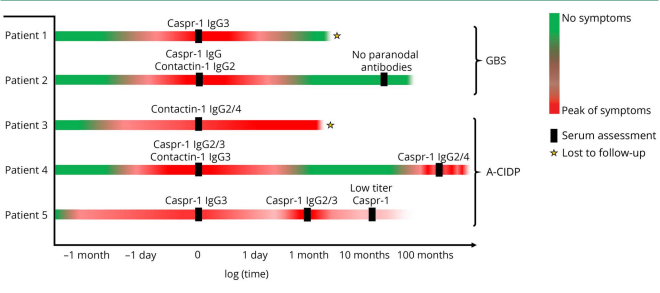
One of two patients with GBS (patient 2) received IVIg, which did not lead to amelioration. This patient was IgG2 > IgG3 seropositive (patient 2, figure 2A). Both patients with GBS responded well to plasma exchange. The two patients with A-CIDP and IgG3 autoantibodies (patients 4 and 5) showed good response to IVIg treatment in the acute phase of disease. Ongoing IVIg treatment led to further improvement of symptoms and was stopped after 25 cycles in patient 5 because of a remission of symptoms. In the other patient with A-CIDP and anti-contactin-1/Caspr-1 IgG3 at disease onset (patient 4), IVIg showed a good therapeutic effect in the acute phase and led to remission not requiring further therapy. In the chronic phase, IVIg was initiated again and stopped after 26 cycles because of a loss of the therapeutic effect. At that stage, the autoantibody subclass had already switched from IgG3 to IgG4 (for detailed treatment response in the course of disease, see appendix e-1, links.lww.com/NXI/A275). The patient with A-CIDP and high-titer anti-contactin-1 IgG4 autoantibodies (patient 3) had to be treated on ICU because of the severity of symptoms, developed renal dysfunction, did not respond to IVIg treatment, and did not ameliorate during the course of disease.

Discussion

We detected antiparanodal autoantibodies in 4.3% of patients with acute to subacute immune-mediated neuropathy, more frequently in CIDP with acute to subacute onset than in GBS. We found IgG2/3 in monophasic GBS, which supports the notion of previous studies that IgG3 is the predominant subclass in monophasic disease.⁷ By analyzing sera from different time points, we could show that the antibody status was concordant with the disease severity. Therefore, our data support pathogenicity of autoantibodies and suggest that autoantibody status and titer are valid indicators for disease activity, as proposed in previous studies on paranodopathy and further IgG4-related neurologic disease.^{25–27}

In one patient, we could demonstrate a subclass switch from IgG3 in acute disease to IgG4 in chronic disease, a finding that we had hypothesized in a previous study.⁶ In other autoimmune diseases, e.g., in membranous glomerulonephritis, subclass switch of anti-PLA2R autoantibodies from IgG1 to IgG4 also associates with progression to chronic disease.²⁸ According to a linear model of autoantibody response, unspecific IgM response is followed by IgG isotype, produced by antigen-experienced B cells. These IgG are of the IgG3 subclass and sequentially switch to IgG1, 2 and 4.^{29,30} Concurrently, autoantibodies strongly gain antigen affinity, with IgG4 showing the highest affinity to its target.^{31,32} Disease progression in paranodopathy might therefore depend on IgG subclass properties

Figure 4 Scheme of the autoantibody status in the course of disease in seropositive patients



The course of disease is shown on the x-axis by pseudo-logarithmic display of the time course. Time point 0 is set at baseline assessment of serum/CSF. The color code indicates severity of symptoms. Results of serum assessment are displayed in black. Patients 1 and 3 were lost to follow-up.

and affinities. Gain of specificity and affinity at switch from IgG3 to IgG4 might explain different binding characteristics of antiparanodal autoantibodies in the patient whose IgG3 autoantibodies reacted both against contactin-1/Caspr-1 epitopes, but whose IgG4 autoantibodies were specific to Caspr-1. Here, we report IgG2/4 (1) in patient 4 in the chronic phase after subclass switch and (2) in another patient at disease onset, showing nonameliorating disease with high disability similar to previous reports on anti–contactin-1 IgG4-related disease.^{11,13,33} So far, studies have only reported antiparanodal IgG4 autoantibodies in patients with chronic disease.^{7,11,34,35} We therefore suggest that high-affinity IgG4 autoantibody binding leads to chronic disease, with functional impairment possibly caused by disturbance of the antigen interaction, paranodal architecture, and paranodal complex formation as demonstrated previously.^{6,8–11}

Patients of our study with predominant IgG3 subclass showed good response to IVIg, whereas patients with IgG2 and IgG4 initially did not respond to IVIg or lost responsiveness in the disease course. Several studies described good responses to IVIg in IgG3-associated paranodopathy, but poor responses in IgG4-associated disease.^{5,6,11,13,34} IgG3 can initiate complement activation, opsonization, antigen cross-linking, and internalization, whereas IgG2 shows little C1q binding and IgG4 completely lacks these Fc-mediated effector functions.³¹ In an in vivo study on anti-contactin-1 pathogenicity, we suggested complement-mediated functional impairment at the paranodes as the pathophysiologic correlate in IgG3-mediated paranodopathy earlier and could show in vitro that IVIg inhibits complement deposition and activation dose-dependently.^{14,15} Our data therefore support the hypothesis that therapy responsiveness might depend on IgG subclass and subclassrelated pathomechanism of antiparanodal autoantibodies. If validated in larger studies, these findings might have a direct impact on the treatment regime in affected patients. Testing of autoantibody subclass during the course of disease may be useful as an indicator of a positive response to IVIg in case of IgG3, whereas IgG4-related paranodopathy shows very good response to rituximab.^{4,6,7,11}

The trigger of the autoantibody production and the pathomechanism behind possible IgG switching are still unknown. Three of five seropositive patients in this study had diabetes mellitus type 2 as a comorbidity diagnosed years before the onset of disease. Previous studies report disruption of the bloodbrain barrier and the nodal architecture in patients with diabetes mellitus, thereby possibly exposing paranodal antigens to the immune response.^{36,37} A recent study has identified human leukocyte antigen (HLA)-DRB15 as a risk factor for CIDP associated with anti-NF155 antibodies, and another study reported different HLA antibody patterns with subclass switch after renal transplantation.^{38,39} Whether interindividual differences in antibody production and switching during the course of disease depend on (1) HLA alleles, (2) antigen exposure, or (3)further immunologic mechanisms has to be investigated in further studies.

Regarding characteristic clinical features of paranodopathy, we report neuropathic pain in anti–Caspr-1–related disease as previously described in patients with higher autoantibody titers.⁷ Different titers might explain discrepancies between our results and a recent study that did not report pain in anti–Caspr-1–positive patients.^{7,11} In our patients, further features of IgG4-related chronic paranodopathy such as tremor, sensory ataxia, and nephropathy^{5,6,13,40} were only present in the course of disease, suggesting that they evolve due to chronic autoantibody

exposure. The pathomechanistic correlate of action tremor, which recently has also been described in anti–Caspr-1 IgG3related paranodopathy,¹¹ is still under research, but studies suggest cerebellar origin.^{5,6,12} In this study, we detected intrathecal autoantibodies in a patient with tremor, as previously reported in a case of anti–pan-neurofascin–positive paranodopathy.¹² The antiparanodal autoantibodies most probably diffuse into the intrathecal compartment due to a blood-brain barrier dysfunction, as indicated by normal IgG and ASI indices in our patients. Our CSF data suggest that detection of antiparanodal antibodies in CSF shows little sensitivity and depends on either high titer or strong leakage of the blood-brain barrier. We therefore encourage further studies on intrathecal antiparanodal autoantibodies targeting sensitivity, pathogenicity, and association to clinical symptoms.

Low prevalence of autoantibodies against paranodal proteins in acute inflammatory neuropathy, retrospective assessment, and the low number of seropositive patients available for follow-up limit our study. Clinical characteristics at acute onset and subclassrelated seropositivity as a possible risk factor for chronic progression when detected at the onset should be investigated in large multicentric studies, considering low overall prevalence of GBS and A-CIDP as well as further possible confounders in analysis of paranodopathy-specific clinical symptoms, e.g., coinciding painful diabetic neuropathy. The question of sequential subclass switching in paranodopathy needs to be addressed in larger prospective studies. Nevertheless, our data (1) propose autoantibody subclass screening to be considered in patients with GBS and A-CIDP, as results might have direct implications on diagnosis and therapeutic regime during the course of disease, and (2) pave the way for further clinical and pathomechanistic studies on antiparanodal autoantibodies in acute to subacute immune-mediated neuropathy.

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Disclosure

L. Appeltshauser, K-P. Wandinger, R. Junker, C. Sommer, F. Leypoldt, and K. Doppler work for an academic institution offering commercial antibody diagnostics. A.M. Brunder, A. Heinius, P. Körtvelyéssy, and C. Villmann report no disclosures relevant to the manuscript. C. Sommer has served on scientific advisory boards for Algiax, Alnylam, Air Liquide, Akcea, Astellas, Bayer, Grifols, Takeda, and UCB. She reports being a member of speakers' bureau and receiving speaker honoraria from Akcea, Alnylam, Novartis, Pfizer, Sanofi-Aventis, and Teva. C. Sommer served as a journal editor, associate editor, or editorial advisory board member for the European Journal of Neurology, PLoS ONE, and PAIN Reports. F. Leypoldt reports speaker honoraria from Bayer, Roche, Novartis, and Fresenius, travel funding from Merck, Grifols, and Bayer, and serving on advisory boards for Roche, Biogen, and Alexion. Go to Neurology.org/NN for full disclosures.

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Appendix Authors

Name	Location	Contribution	
Luise Appeltshauser, MD	University Hospital of Würzburg	Major role in the acquisition, analysis, and interpretation of the data and drafting and revising the manuscript for intellectual content	
Anna-Michelle Brunder	University Hospital of Würzburg	Major role in the acquisition and analysis of the data	
Annika Heinius	University Hospital of Schleswig-Holstein, Campus Kiel	Major role in the acquisition and analysis of the data	
Peter Körtvélyessy, MD	University Hospital of Magdeburg	Revising the manuscript fo intellectual content and contribution of vital reagents/tools/patients	
Klaus-Peter Wandinger, MD	University Hospital of Schleswig-Holstein, Campus Kiel	Revising the manuscript for intellectual content and contribution of vital reagents/tools/patients	
Ralf Junker, MD	University Hospital of Schleswig-Holstein, Campus Kiel	Revising the manuscript for intellectual content and contribution of vital reagents/tools/patients	
Carmen Villmann, PhD	University Hospital of Würzburg	Revising the manuscript for intellectual content and contribution of vital reagents/tools/patients	
Claudia Sommer, MD	University Hospital of Würzburg	Conceptualization of the study; revising the manuscript for intellectual content; and major role in interpretation of data	
Frank Leypoldt, MD	University Hospital of Schleswig-Holstein, Campus Kiel	Drafting and revising the manuscript for intellectual content; design and conceptualization of the study; and major role in interpretation of data	

Appendix	(continued)
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Name	Location	Contribution
Kathrin Doppler, MD	University Hospital of Würzburg	Drafting and revising the manuscript for intellectual content; design and conceptualization of the study; major role in analysis and interpretation of data; and study supervision

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