



Repeat-associated ataxias in a German patient cohort analysed by targeted parallel long-read sequencing

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Hereditary adult-onset ataxias are a heterogeneous group of phenotypically overlapping conditions, often caused by pathogenic expansions of short tandem repeats. Currently, 18 repeat disorders with a core phenotype of adult-onset ataxia are known. Diagnosis typically relies on sequential PCR-based methods, which are labour-intensive and lack precision. Long-read sequencing (LRS) has the potential to overcome these limitations and is currently implemented and validated in clinical genetics. Using clinical nanopore Cas9-targeted sequencing (Clin-CATS) for parallel in-depth repeat analysis, we evaluated a diagnostic cohort of 513 adult-onset ataxia patients, determining frequencies of all known repeat-associated ataxias except Spinocerebellar ataxia 4 (SCA4), as well as the carrier frequencies for autosomal-recessive disorders, RFC1 spectrum disorder and Friedreich's ataxia (FRDA). Additionally, phenotypes of patients with established genetic diagnoses were characterized, especially those of patients living with RFC1 spectrum disorder and SCA27B.

Repeat-associated ataxias were confirmed in 33.3% of cases, including rare ataxias, such as SCA10, SCA36 and SCA37, alongside as the most prevalent conditions SCA27B and RFC1 spectrum disorder. Potentially pathogenic expansions in FGF14 were identified in an additional 4.7% of patients. Testing of another 347 patients for ZFH3 expansions linked to SCA4 did not identify any cases. Dual diagnoses were frequent, occurring in 6.4% of patients with repeat-associated ataxia. We confirmed a high RFC1 spectrum disorder carrier frequency (7.2%) and reclassified certain FXN expansions as likely non-pathogenic, resulting in a lower than estimated carrier frequency for FRDA of 0.8%. We also identified novel repeat configurations in several loci and illustrated the high heterogeneity of repeat expansions in RFC1, highlighting it as a potential source of false results when using PCR-based methods. This study underscores the diagnostic

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advantages of LRS for comprehensive repeat analysis and recommends its adoption as a standard in clinical genetics, replacing Southern blot and PCR-based approaches. Furthermore, based on our findings in a large patient cohort a re-evaluation of existing phenotype-genotype correlations is recommended as well as evaluating additional parameters alongside the repeat length to improve diagnostic precision of repeat analysis.

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Introduction

Adult-onset ataxias are a heterogeneous group of phenotypically overlapping conditions characterized by a progressive lack of coordination leading to stance and gait imbalance, ocular motor disturbance, dysarthria and dysidiadochokinesia. Additional signs and symptoms, such as neuropathy, basal ganglia dysfunction or cognitive and behavioural changes, may also occur.^{1,2} Based on their aetiology, these conditions are classified as acquired, sporadic-degenerative or hereditary.³

The most frequent genetic cause of ataxias is the pathogenic expansion of short tandem repeats. To date, 18 repeat disorders with

ataxia as a core phenotype have been identified, encompassing autosomal-dominant, autosomal-recessive and X-linked inheritance patterns (Supplementary Table 1).^{4–6} Due to the deep intronic location of their repeat expansions, two of these disorders, RFC1 spectrum disorder and spinocerebellar ataxia 27B (SCA27B), have only been described recently, despite their suspected high prevalence, likely accounting for a significant proportion of apparently sporadic degenerative ataxias.^{5,7–10} RFC1 spectrum disorder results from biallelic pathogenic repeat expansions in RFC1.^{7,8} Initially linked to the clinical triad of Cerebellar Ataxia, Neuropathy and Vestibular Areflexia Syndrome (CANVAS), it is now associated with a broader range of neurological conditions, in particular, incomplete

presentations of CANVAS and additional symptoms such as chronic cough, autonomic dysfunction and motoneuron involvement.^{11–14} SCA27B, an autosomal-dominant ataxia caused by intronic GAA expansions in *FGF14*, is often episodic, triggered by physical exertion and progresses slowly, complicating its diagnosis, despite the availability of symptomatic treatments, such as 4-aminopyridine.^{5,9,15–17}

Hereditary ataxias cannot be reliably diagnosed on clinical findings alone, making molecular genetic testing essential for accurate diagnosis.¹⁸ Standard diagnostic methods, including PCR-based methods and Southern blotting, focus on repeat length as the primary diagnostic parameter. These approaches are limited in their ability to detect complex repeat configurations, where pathogenicity depends on distinct motifs exceeding a specific length threshold, as seen in *RFC1* spectrum disorder, SCA27B, SCA31 and SCA37.^{19–21} Although repeat-primed PCR methods can detect the presence or absence of the expansion of a specific motif, they do not provide information on repeat length, nor do they capture all possible repeat motifs or complex repeat configurations, as is often the case for expansions in *RFC1*.^{19,22,23}

Long-read sequencing (LRS) has emerged as a transformative method for repeat analysis, offering parallel and detailed analysis of multiple repeat loci while overcoming current diagnostic limitations.^{24–26} We recently implemented clinical nanopore Cas9-targeted sequencing (Clin-CATS) for the comprehensive, in-depth repeat analysis of patients with adult-onset ataxia.²⁵ Initially covering the 10 most common repeat-associated ataxias known in 2020, we have since extended our assay to cover all known repeat-associated adult-onset ataxias. SCA4 was added after the completion of this study, following the recent identification of its molecular origin.^{6,27}

In this study, we applied Clin-CATS to a clinical cohort of 513 adult-onset ataxia patients in Germany. We determined the diagnostic yield, assessed the frequency of the different conditions and their genotypes, including configuration (repeat length plus repeat motif) of polymorphic repeat expansions in *RFC1*, and reassessed the carrier frequencies of the autosomal-recessive disorders, *RFC1* spectrum disorder and Friedreich's ataxia (FRDA). Based on our findings, we discuss challenges in clinically assessing repeat expansions and compare Clin-CATS to other LRS approaches.

Materials and methods

Patient DNA samples and study approval

In total, 513 individuals (>20 years old) with clinical diagnosis of cerebellar ataxia and without established genetic diagnosis were analysed with our updated Clin-CATS workflow. The cohort includes patients without prior genetic diagnostics as well as patients having had a negative result in a previous analysis. Only three families, with a total of seven individuals, were included in the study; the remaining 506 patients were unrelated. Patients were referred to genetic testing from general neurologists, as well as specialized centres for movement disorders from all over Germany likely resembling the general German population living with cerebellar ataxia. Genetic testing was requested only if there was no suspicion or confirmation of another, alternative aetiology of ataxia (e.g. infectious, autoimmune, paraneoplastic, toxic or structural) after thorough clinical characterization by board-certified neurologists and diagnostic procedures addressing aforementioned aetiologies. Clinical data were collected retrospectively from medical records. Another 347 patients were analysed with the most recent version of Clin-CATS including guides for *ZHFX3*. All patients were analysed as diagnostic samples within our institute between April 2023 and October 2024. Informed consent

was obtained from all patients. The study was approved by local institutions (Bayerische Landesärztekammer, 2019–210). All genetic analyses and investigations were performed in accordance with the guidelines of the Declaration of Helsinki.

Extraction of genomic DNA

Genomic DNA (gDNA) was obtained from total peripheral blood samples by extraction of white blood cells with a Biomek FX system (Beckman Coulter) using the NucleoMag® Blood 3 ml Kit (Machery-Nagel, 744502.1) as per the manufacturer's instructions. All DNA samples showed high purity as determined by optical density measurements ($A_{260}/A_{280} > 1.9$ and $A_{260}/A_{230} > 2.0$). The DNA Integrity Number (DIN) was measured on the TapeStation 4200 (Agilent Technologies, G2991BA) as per the manufacturer's instructions.

Repeat analysis by Oxford Nanopore Technologies LRS

All patients were analysed using Oxford Nanopore Technology (ONT) LRS-based Clin-CATS, a accredited diagnostic method in Germany.²⁵ Library preparation and flow cell loading were performed according to the ONT Cas9-targeted sequencing protocol using ONT's SQK-CS9109 kit and 5 µg of input gDNA. CRISPR RNAs (crRNAs) for the enrichment of 17 regions of interest were designed using CHOPCHOP 8 (Supplementary Table 2). Sequencing was performed with ONT FLO-MIN106D R9 flow cells on the GridION X5 sequencer. Base calling was performed using Guppy (v.5.0.16).^{28,29} The generated FASTQ files were aligned to the human reference genome (GRCh38/hg38) using Minimap2 (v.2.17) to identify the reads spanning the targets of interest.³⁰ Minimum criteria to evaluate samples in the diagnostic context was a 50× coverage across all target regions. For *RFC1* and *NOP56*, incomplete reads exceeding the current threshold of pathogenicity of the respective motif were considered for calculating the on-target coverage. For quality control of the aligned reads, NanoPlot (v.1.29.1) was used.³¹ The bioinformatics tool STRique (v.0.2.1) was used to determine the number of repeat units (RU) for all reads assigned to the regions of interest.³² Repeat length distributions obtained by STRique were visualized as violin plots and used to determine the repeat size for each allele by computing local maxima using FindPeaks (v.2.1.1) and visual assessment of the plots.³³ For extended repeat expansion (>100 RU) showing multiple local maxima, a size range was calculated starting with the lowest and ending with the highest significant peak as determined by FindPeaks. In all such cases, the median repeat length was calculated from the mapped reads, provided the repeat length exhibited an approximately Gaussian distribution, indicating sufficient coverage of the respective allele. Repeat configurations were analysed manually by visual inspection of all reads mapped to the respective regions in the Integrative Genomics Viewer (IGV). Repeat length distribution and median repeat length were extracted for each allele separately. For *ATXN8/ATXN8OS*, repeat lengths were determined as the sum of the non-pathogenic proximal (CTA • TAG)_{exp} repeat and the adjunct disease-associated (CTG • CAG)_{exp} repeat as established for PCR-based methods. Patients heterozygous for a pathogenic repeat expansion in *RFC1* or *FXN* were analysed for a pathogenic variant on the other allele by next-generation sequencing (NGS).

The validity of median repeat lengths for the repeat in *FGF14* was confirmed by comparing our results with those of the previously established sequencing of long-range PCR products using ONT LRS (LR-PCR amplicon sequencing; Supplementary Table 3 and

Supplementary Fig. 1).³⁴ Patients were diagnosed as positive for a repeat disorder when they carried a reduced-penetrance or full-penetrance repeat expansion. If more than one repeat expansion was detected, the primary diagnosis was based on the repeat expansion with full penetrance, while additional expansions with reduced penetrance were listed as secondary (and tertiary) diagnoses.

NGS

All patients being heterozygous carriers for RFC1 spectrum disorder and FRDA were analysed by NGS sequencing. Further patients were analysed, when requested by referring physicians based on non-standardized decision criteria. To identify pathogenic single nucleotide variants (SNVs) or copy number variations (CNVs) in known genes associated with ataxia (altogether 250 genes, [Supplementary Table 4](#)), gene-targeted enrichment was performed with the SureSelectXT gene panel custom kit (Agilent Biosciences) or the Twist Human Comprehensive Exome Kit (Twist Biosciences). Massively parallel sequencing was carried out on an Illumina NextSeq 500 or a Novaseq 6000 system (Illumina) as 150 base pairs paired-end runs using v.2.0 sequencing by synthesis chemistry (Illumina). Secondary and tertiary analysis was carried out using varvis@ 1.22.0. Pipeline versions fc9-063-00b and 8a2-0c8-080 were used for SNV and CNV analysis, respectively. Only SNVs and small insert and deletions (INDELs) in the coding and flanking intronic regions (± 50 bp) were evaluated. Variants were classified according to the ACMG/AMP (American College of Medical Genetics and Genomics and the Association for Molecular Pathology) guidelines.³⁵ In total, 185 patients were analysed, comprising 146 with a negative result (139 patients) or only an intermediate FGF14 allele (7 patients) and 39 with likely causative repeat expansions.

Results

Diagnosis of 513 ataxia patients by parallel LRS-based repeat analysis

We analysed a cohort of 513 patients with adult-onset ataxia (median age 61 years; range: 22–88; [Supplementary Table 5](#)) using our updated version of Clin-CATS.²⁵ This assay captured all known repeat-associated hereditary ataxias, except SCA4, whose molecular origin was only recently discovered ([Fig. 1](#) and [Supplementary Table 1](#)).^{6,27} Genetic analysis revealed pathogenic repeat expansions confirming hereditary ataxia in 171 patients (33.3%) ([Fig. 2](#) and [Supplementary Table 5](#)). An additional 24 patients (4.7%) harboured FGF14 expansions (180–249 RU), which, while potentially pathogenic with reduced penetrance for SCA27B, remain unconfirmed and are henceforth referred to as ‘intermediate FGF14 alleles’.^{16,34} The most common repeat expansions were compatible with RFC1 spectrum disorder (42 patients, 8.2%) and SCA27B [in total 83 patients (16.2%): 55 (10.7%) with (GAA)_{>300} RU expansions, 28 (5.4%) with (GAA)_{250–300} RU expansions]. Including intermediate FGF14 alleles, 107 patients (20.9%) had results consistent with SCA27B. Pathogenic repeat expansions compatible with a primary diagnosis of SCA8 were the third most common finding (11 patients, 2.1%), followed by SCA3 (7 patients, 1.4%), SCA6 (6 patients, 1.2%), Fragile X-associated-tremor/ataxia syndrome (FXTAS; 6 males, 1 female; 1.4%) and other rarer expansions: SCA1 (4 patients, 0.8%), FRDA (3 patients, 0.6%), SCA2 (2 patients, 0.4%), SCA10 (2 patients, 0.4%), dentatorubral-pallidolysian atrophy (DRPLA) (1 patient, 0.2%), SCA36 (1 patient, 0.2%) and SCA37 (1 patient, 0.2%). One patient exhibited a TBP expansion with reduced penetrance alongside a variant of uncertain significance

(VUS) in STUB1, indicating SCA17 and/or SCA48, with the intermediate allele in TBP acting as a disease modifier.³⁶ No cases of SCA7, SCA12 or SCA31 were detected in this cohort.

Testing of another 347 patients with adult-onset ataxia for SCA4

After completing the study of the 513 patients, another 347 patients were tested for SCA4 for an initial estimate of its frequency in patients with adult-onset ataxia. No pathogenic repeat expansions compatible with diagnosis of SCA4 were identified in this extended cohort.

Potential dual genetic diagnoses and additional repeat expansions

Among the 171 patients with confirmed hereditary ataxias, five (2.9%) carried reduced-penetrant repeat expansion(s) compatible with a potential dual diagnosis of two distinct late-onset ataxias ([Fig. 2B](#), dashed line, and [Supplementary Table 5](#)). An additional six patients (3.5%) carried an intermediate FGF14 allele as a second repeat expansion ([Fig. 2B](#)), while another six patients (3.5%) had biallelic expansions in loci associated with autosomal-dominant ataxia, including one patient each with SCA6 and SCA8, and four with SCA27B (exclusively intermediate alleles at the second allele). Furthermore, 10 patients (5.8%, not shown in [Fig. 2B](#)) were also carriers for RFC1 spectrum disorder (nine patients) or FRDA (one patient).

Family history

Among 119 patients with autosomal-dominant repeat disorders, information on family history was available for 73 patients ([Supplementary Table 5](#)). Of these, only 42% reported having a family member affected by ataxia. When restricting the analysis to the 79 patients with fully penetrant repeat expansion, this proportion increased slightly to 55%.

Genotypes and phenotypes of repeat-associated ataxias and reassessment of carrier frequencies

ATN1 (DRPLA)

One patient was diagnosed with DRPLA carrying a repeat expansion in the lower range of pathogenicity (50 RU) in line with late disease onset in the early seventies. The clinical presentation included gait ataxia and oculomotor disturbances. Notably, the patient did not exhibit other common DRPLA-associated features such as myoclonus, dystonia, chorea, epilepsy or additional movement disorders, but showed mild memory impairment.

ATXN1-3, CACNA1A (SCA1–3, 6)

Patients diagnosed as affected by SCA1–3 and 6 carried pathogenic repeat expansions with typical compositions. Notably, one patient with SCA6 was found to be homozygous for a 20-repeat expansion in CACNA1A, a finding consistent with previous reports of pathogenicity at this repeat size. Cumulative clinical features for all patients are summarized, with detailed individual data provided in [Supplementary Table 6](#). In addition to cerebellar symptoms, peripheral neuropathy was a consistent finding across all patients diagnosed with SCA1–3 and SCA6.

ATXN8/ATXN8OS (SCA8)

In addition to various cerebellar signs, 66.7% of patients with a pathogenic repeat expansion in ATXN8/ATXN8OS frequently

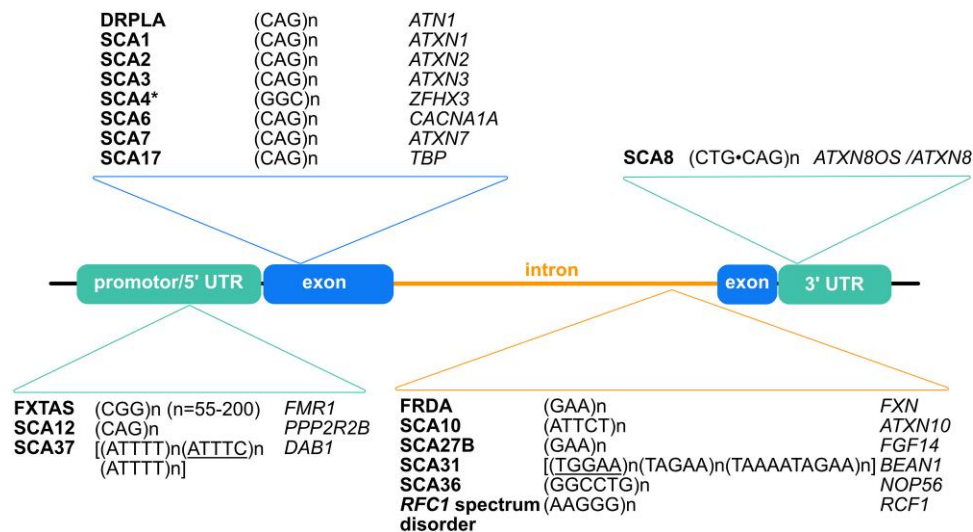


Figure 1 Overview of repeat-associated ataxias. Repeat-associated ataxias and their location in the genome, repeat motifs and associated genes. For all polymorphic repeat loci, the most frequent pathogenic repeat motif or repeat configuration is given. *Recently included within the last update of clinical nanopore Cas9-targeted sequencing (Clin-CATS).

exhibited additional non-cerebellar features. These included both hyperkinetic and hypokinetic movement disorders, as well as upper motor neuron signs (Supplementary Table 6). This reflects the known phenotypic variability associated with SCA8.

ATXN10 (SCA10)

In two related patients (father and daughter), we identified a complex repeat expansion in ATXN10 [(ATTCT)_{~130}(ATTCC)_{~1400}] compatible with SCA10. Unlike the pure expansions of the wildtype-motif (ATTCT)_{exp} typically associated with SCA10, these patients exhibited heterogeneous expansions of the wild-type motif (ATTCT)_{exp} and an alternative motif (ATTCC)_{exp}, previously suggested as a disease-modifier increasing risk for seizures.^{37,38} Clinically, neither patient exhibited seizures, and both primarily presented with gait disturbance and other cerebellar features. EEG showed a normal result in one patient, while abortive spike-wave complexes were present in the other patient.

DAB1 (SCA37)

A complex heterozygous [(ATTTT)₁₅₅(ATTTC)₁₀₀(ATTTT)₁₀] repeat expansion was identified in one female, compatible with SCA37. The repeat configuration differed slightly from previously reported patterns, which typically consist of [(ATTTT)₇₋₄₀₀(ATTTC)₃₁₋₇₉(ATTTT)₅₈₋₄₂₀].^{20,39} The pathogenic (ATTTC)_n insertion was unusually long and shifted towards the 3'-end of the ATTTT expansion, leading to an atypically short 3'-ATTTT repeat. Clinically, the patient presented with a pure cerebellar ataxia phenotype, including oculomotor disturbances, limb and gait ataxia, and dysarthria. The onset of symptoms was at the age of 43. No additional non-cerebellar features—such as motor neuron involvement or peripheral neuropathy—were observed.

FGF14 (SCA27B)

In addition to pure (GAA)_{>180 RU} expansions, 15 patients carried FGF14 expansions (>100 RU) involving an alternative repeat motif not yet associated with SCA27B. The most common variant, (GAAGGA)_{exp}, observed in 14 patients, was often flanked by short

stretches of (GAA)_{exp} as previously reported.³⁴ One patient exhibited a novel repeat configuration, (GGA)₇₅(GAA)₃₀₀, currently classified as of unknown significance. While the pathogenic (GAA)_{exp} expansion exceeds the established threshold of pathogenicity, its co-occurrence with another motif complicates interpretation.

Patients diagnosed as affected by SCA27B had a median age of onset of 62 years and most frequently presented with gait ataxia (95% of patients) and oculomotor signs (92%), while limb ataxia was only present in 66% of patients (Fig. 3A and Supplementary Table 6). Approximately half of the patients exhibited an episodic onset or course (52%) and downbeat nystagmus (54%), both considerable more specific features of SCA27B. Various forms of neuropathy were present in about 43% of patients. Notably, 25% of patients for whom respective clinical data were available fulfilled the clinical triad of CANVAS. No correlation between age of onset and repeat length could be observed ($r < 0.01$; Fig. 3C).

FMR1 (FXTAS)

Six male patients and one female patient were identified as carriers of an FMR1 premutation, potentially associated with FXTAS. The median age at onset of cerebellar ataxia was 66 years. Based on established diagnostic criteria for FXTAS, which include both clinical features and characteristic radiological findings, three male patients met the criteria for definite FXTAS, and two were classified as probable cases. Due to incomplete clinical and imaging data, the likelihood of FXTAS in the female carrier could not be assessed.⁴⁰

FXN (FRDA)

Among the three patients diagnosed with FRDA, two individuals—aged 38 (Patient 330) and 50 years (patient 263)—carried one short allele (100 and 160 RU, respectively) and one large expanded allele. This configuration is consistent with (very) late-onset FRDA, and both patients exhibited a milder phenotype being less specific for FRDA. Patient 330 presented with pronounced gait ataxia, dysarthria, spasticity, sensory neuropathy, and evidence of both pyramidal and dorsal column involvement based on evoked potential studies. Patient 263 showed cerebellar ataxia with marked gait ataxia,

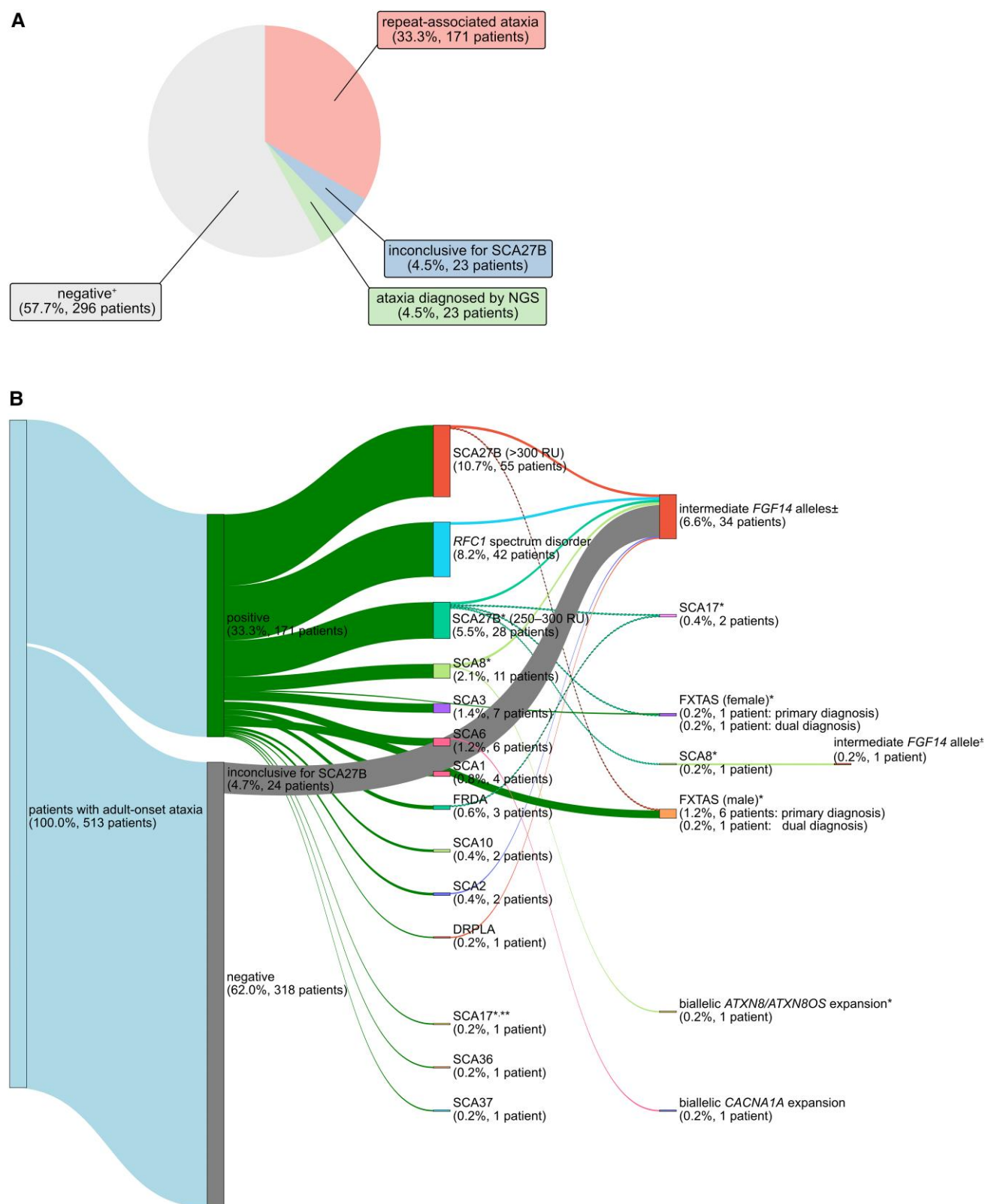


Figure 2 Diagnostic results within a cohort of 513 patients with adult-onset ataxia. (A) Classification of the 513 patients analysed by clinical nanopore Cas9-targeted sequencing (Clin-CATS, all patients) and next-generation sequencing (NGS, 185 patients). * NGS testing was performed in only 118 of these patients. (B) Diagnostic results based on Clin-CATS of 513 patients with adult-onset ataxia. Flow diagram visualizing diagnostic result (positive/negative), primary (dark green lines) and dual diagnoses. *Reduced penetrance; **Patient additionally carried a variant of uncertain significance (VUS) in *STUB1*; dashed lines indicate dual diagnoses; †Intermediate alleles in *FGF14* with unknown clinical significance. One patient with intermediate *FGF14* allele was diagnosed with SPG7 by NGS.

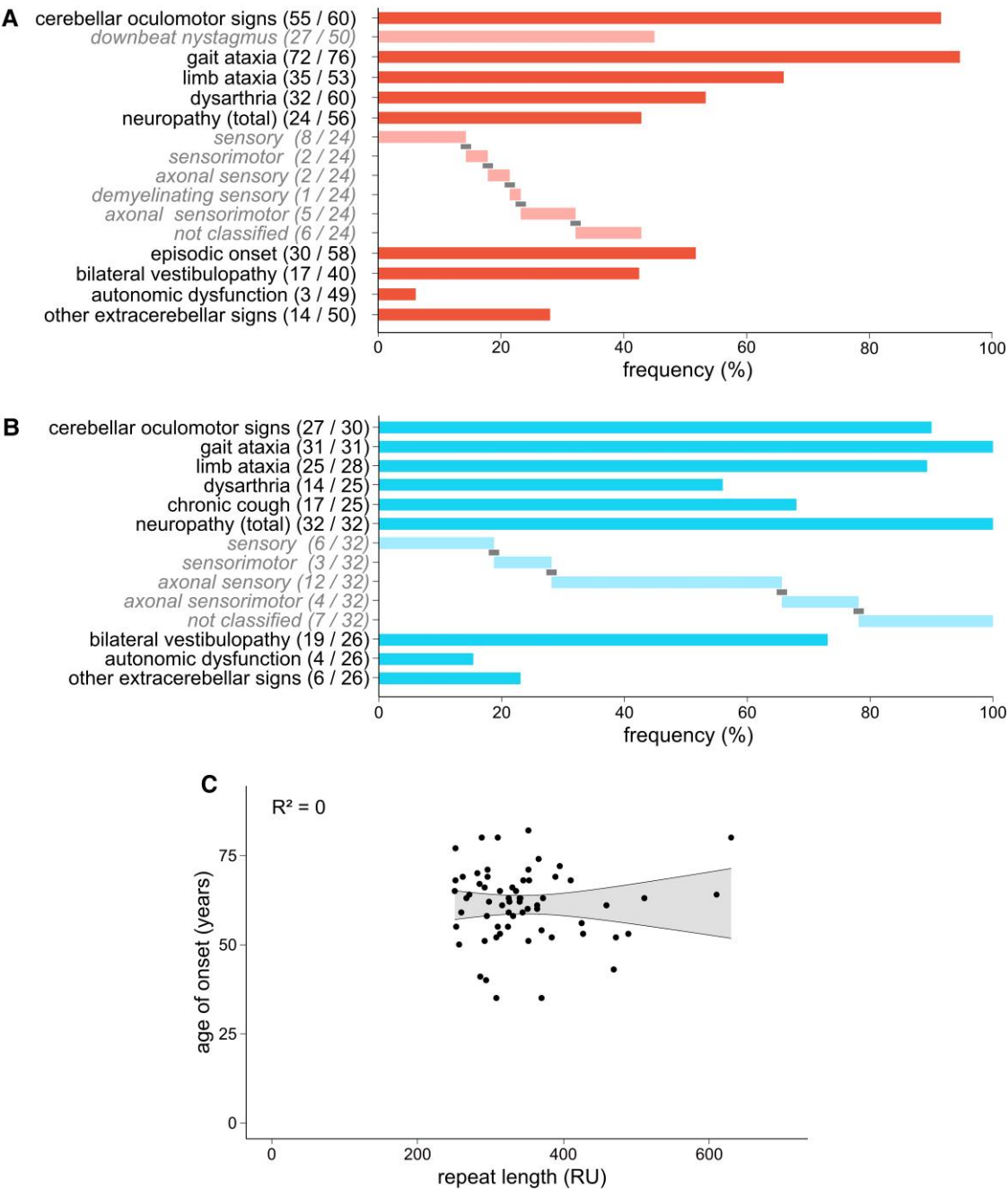


Figure 3 Clinical features in patients living with SCA27B and RFC1 spectrum disorder. Frequency of different symptoms in patients living with (A) SCA27B and (B) RFC1 spectrum disorder. (C) Correlation of age of onset and FGF14 repeat length.

dysarthria and spasticity, but retained deep tendon reflexes and had no evidence of dorsal column involvement or neuropathy based on sensory evoked potentials and electroneurography. This atypical presentation delayed diagnosis for over 9 years. The third patient (Patient 297), aged 24 years, carried large biallelic FXN repeat expansions along with an incompletely penetrant repeat expansion in TBP. According to the referring neurologist, this patient displayed a classical FRDA phenotype.

In addition, four individuals were identified as heterozygous carriers for FRDA. Two carried large expanded alleles (827 and 915 RU), typically associated with childhood- or adolescent-onset FRDA in biallelic cases. The other two carried short expansions

(91 and 93 repeat units), previously linked to (very) late-onset and atypical FRDA presentations. Truncating FXN variants in *trans* were excluded in all of these individuals.

Five additional individuals carried complex heterozygous repeat expansions with alternative motifs: four exhibited the known (GAAGGA)_{exp} configuration, and one harboured a novel (GAAGAG)_{exp} motif flanked by a (GAA)_{exp} segment. These were classified as variants of uncertain significance. While short (GAA)_{exp} expansions interrupted by alternative motifs have been associated with very late-onset FRDA, pure (GAAGGA)_{exp} expansions are currently considered likely non-pathogenic.^{12,41,42} Therefore, only the four individuals with a heterozygous pure (GAA)_{exp} expansion

were included as FRDA carriers. Based on the 510 individuals who tested negative for biallelic FXN expansions, the carrier frequency for FRDA in this cohort was estimated at 0.8% (1:125).

NOP56 (SCA36)

One patient was diagnosed with SCA36 based on a large GGCTG-expansions (>650 RU) in NOP56. The clinical presentation was consistent with a mild, pure cerebellar ataxia, including oculomotor abnormalities, discrete limb ataxia and mild gait disturbance. Notably, the patient did not exhibit dysarthria or hyper-reflexia—features commonly associated with SCA36. The disease onset was at the age of 45.⁴³ Atrophy and fasciculation of the tongue were not noticed as another common features of SCA36.⁴³

RFC1 (RFC1 spectrum disorder/CANVAS)

The majority of patients with RFC1 spectrum disorder (40 of 42 patients) carried biallelic pathogenic (AAGGG)_{>250 RU} repeat expansions, while the remaining two carried a heterozygous (AAAGG)_{>500 RU} expansion on one allele and a (AAGGG)_{>250 RU} expansion on the other. Characterization of RFC1 repeats revealed significant heterogeneity at this polymorphic locus, with 13 distinct repeat configurations identified (Fig. 4 and Supplementary Table 7). Notably, both interallelic and intraallelic heterogeneity were observed. The 'AAAGGG' motif was never observed as a pure expansion but was always part of a complex repeat structure containing 'AAAGG' and 'AAGGG' stretches [therefore denoted as (AAAGGG)_{complex}], similar to (ACGGG)_{complex} expansions. One allele exhibited a unique composite configuration, (AAAGG)₃₁₇/(AAGGG)₂₁₀, consisting of two pure repeats currently classified as of unknown significance. Sizing was possible for most RFC1 repeat expansions; however, 40 out of 108 (AAGGG)_{>250 RU} expansions could not be precisely measured due to broken reads. Additionally, 34 patients were identified as carriers for RFC1 spectrum disorder based on a pathogenic repeat expansion on one allele [(AAAGG)_{>500 RU} in 7 patients and (AAGGG)_{>250 RU} in 27 patients]. Of these, one patient with cerebellar ataxia and neuropathy carried a pathogenic (AAGGG)₅₅₆ expansion on one allele and an (AAAGG)₃₆₀ repeat expansion on the other, which fell below the current pathogenicity threshold. NGS analysis in all carriers excluded pathogenic variants in *trans*. Based on these 34 cases, we calculated a carrier frequency of 7.2% for RFC1 spectrum disorder among the 471 patients in whom RFC1 was not the primary cause of ataxia.

Patients diagnosed with RFC1 spectrum disorder had a median age of onset of 55 years (Supplementary Table 6). Chronic cough was a common symptom, reported in 68% of patients (Fig. 3B). Among those with available data, all were affected by peripheral neuropathy, which was classified as axonal based on electroneurography findings and in 18/25 patients with information available sensory and the other 7/25 patients sensorimotor. Of these, 26% presented only with neuropathy as an additional symptom to cerebellar ataxia, while the remaining 74% also exhibited bilateral vestibulopathy (BVP), thereby fulfilling the diagnostic criteria for the CANVAS triad.

TBP (SCA17)

One patient was identified carrying a reduced-penetrance allele in TBP along with a VUS in STUB1. The individual presented with a complex clinical phenotype, including cerebellar ataxia, seizures and myoclonus.

NGS analysis of patients with negative or inconclusive Clin-CATS results

Of the 342 patients without a likely causative repeat expansion, 141 underwent NGS to identify pathogenic SNVs or CNVs in ataxia-related genes (Supplementary Tables 5 and 8). A hereditary cause of ataxia was found in 23 patients (6.7%), revealing a spectrum of ataxia syndromes, including SCA11, 13, 14, 28, 42, 48, R8, episodic ataxia type 2, POLR3A-associated ataxia, ataxia teleangiectasia. Additional diagnoses included hereditary spastic paraplegias with overlapping ataxia phenotypes (SPG5, 7, 39, 79A) and syndromes with cerebellar involvement, such as Tay-Sachs disease, neuropathy-ataxia-retinitis pigmentosa (NARP) syndrome, and STXBP1-associated developmental and epileptic encephalopathy (as somatic mosaicism).

Discussion

Diagnostic findings and prevalence of repeat-associated ataxias

The field of hereditary repeat-associated ataxias has made remarkable advancements since 2019, e.g. marked by the discovery of two additional repeat disorders that likely account for a high proportion of ataxia previously classified as sporadic.^{5,7-9} These breakthroughs, driven by LRS, have not only facilitated novel disease discoveries but have also enabled long-pending revisions of clinical diagnostics of repeat disorders.^{24,25} However, the wide implementation of LRS in clinical genetics remains limited. In this study, we analysed 513 patients with adult-onset ataxia using LRS-based Clin-CATS. This is one of the largest cohorts of ataxia patients evaluated since the discovery of SCA27B, to date, and the only one analysed by parallel LRS-based repeat analysis.

The diagnostic yield in this clinical cohort was high, with 33% being diagnosed with repeat-associated ataxia, increasing to 38% when intermediate FGF14 alleles (180–249 RU) were considered. Even if causative SNVs and CNVs were not analysed in all patients, our study shows that repeat expansions are much more frequent than other genetic alterations causing ataxia (33% versus 7%). This emphasizes the importance of prioritizing repeat analysis as an initial genetic test in patients with adult-onset ataxia, either in parallel or after the exclusion of acquired forms of ataxia.

Only 55% of patients with fully penetrant autosomal-dominant repeat expansions reported a positive family history of ataxia. In the remaining 45%, the absence of a reported family history may be explained by subtle or subclinical symptoms in affected relatives—detectable only through targeted neurological examination—or by intergenerational repeat instability. In several repeat-associated disorders, such as SCA1–3, expansions are known to increase in size during transmission, leading to more severe phenotypes in subsequent generations, a phenomenon referred to as anticipation.⁴⁴⁻⁴⁶ Repeat expansions in FGF14 are especially unstable with substantial variations (increase and decrease) of repeat size across generations and, in some cases, shift between the pathogenic and non-pathogenic range within a single family.⁴⁷ As such, restricting diagnostic analysis to patients with a positive family history risks missing a considerable number of cases. Consistent with findings in other populations of European ancestry, SCA27B is the most frequent adult-onset ataxia in our cohort (16.2%), followed by RFC1 spectrum disorder (8.2%).⁴⁸⁻⁵¹ The prevalence of SCA27B may be even higher considering multiple lines of evidence indicating a lower threshold for pathogenicity (20.9% when considering intermediate FGF14

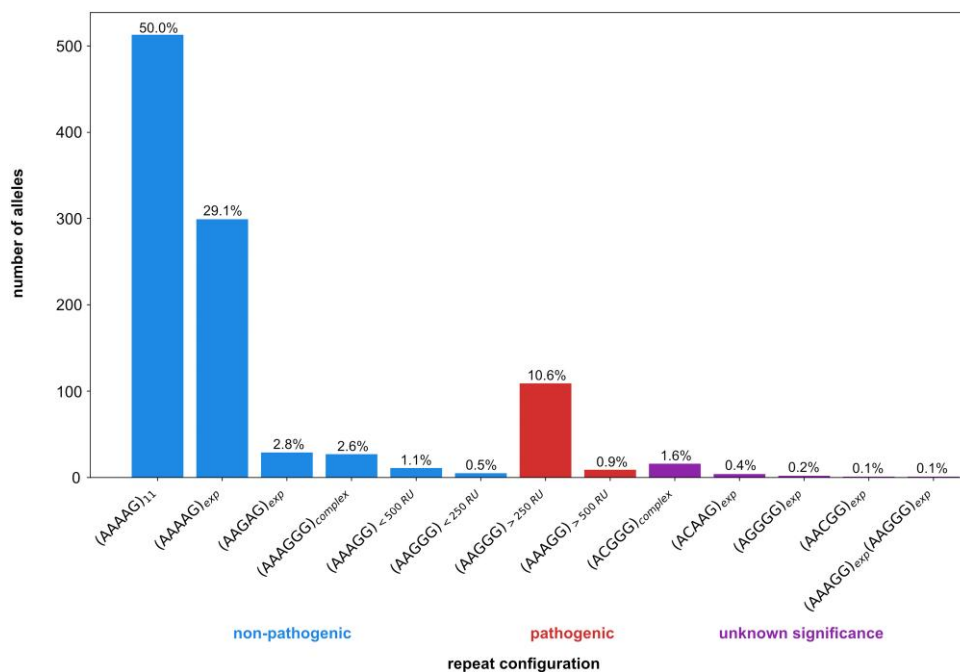


Figure 4 Repeat configurations of RFC1, their frequency and current clinical classification. Total number of alleles: 1026.

alleles).^{34,52,53} Many SCA27B cases may remain undiagnosed due to subtle, slowly progressive symptoms and age-related comorbidities.⁵⁴ Other repeat disorders were substantially less common, collectively accounting for as many cases of ataxia as SCA27B alone.

Parallel repeat analysis identified rare repeat-associated ataxias, including SCA10, 36 and 37, with only SCA37 previously reported in a German patient cohort.^{55–57} After updating our repeat assay by including the ZFH3 repeat, analysis of 347 additional ataxia patients did not reveal any pathogenic repeat expansion in this locus, indicating a low frequency of SCA4 in the population studied.

Determining the relative frequency of the individual repeat-associated ataxias is inherently challenging, regardless of the approach. The main bias in our study is the potential enrichment of patients who previously tested negative for established repeat disorders such as SCA1–3 and 6. A comparison with historical PCR-based repeat testing at our institute in the last two decades (Supplementary Fig. 2) suggests SCA1–3 and 6 may be 1.9 to 6.5 times underrepresented in this cohort. Nevertheless, SCA27B and RFC1 spectrum disease likely remain the most common repeat-associated ataxias, further supported by the high carrier frequency for RFC1 spectrum disorder.

Phenotypic overlap of repeat-associated ataxias and correlation of age of onset and repeat length in SCA27B

Clinical characterization in our cohort highlights the significant phenotypic overlap among different repeat-associated ataxia. Differentiation based on clinical features alone would have been challenging in many cases: patients diagnosed with SCA1–3, SCA6, SCA36 and SCA37 all exhibited non-specific signs of cerebellar ataxia, occasionally accompanied by peripheral neuropathy. Without genetic testing, accurate subtype classification would have been possible only to a limited extent. Among patients

diagnosed with SCA27B, 75% displayed either an episodic disease onset/course and/or downbeat nystagmus, features that have been suggested as relatively specific indicators for this condition.^{5,9,10,54,58} Similarly, 68% of patients with diagnosis of RFC1 spectrum reported a chronic cough, a known presyndromal hallmark of the disease.⁵⁹ However, the presence of the CANVAS triad (cerebellar ataxia, neuropathy and bilateral vestibulopathy) alone is insufficient to distinguish between these two entities. In our study, 25% of patients with SCA27B also fulfilled the CANVAS triad, consistent with previous reports, although the characteristics of neuropathy and vestibular involvement tend to differ between SCA27B and RFC1 spectrum disorder.^{16,60}

Current literature offers conflicting findings regarding an inverse relationship between repeat length and age of onset in SCA27B. In agreement with several previous studies, we did not observe a statistically relevant correlation in our cohort.^{15,34,50,58} Nevertheless, it remains plausible that a weak inverse correlation exists and may only become apparent in larger, more age-diverse cohorts.^{5,9,34}

It is important to note that clinical data in this study were collected retrospectively and not through standardized neurological assessments. The primary goal of evaluation was to determine the aetiology of cerebellar ataxia, and therefore the extent and type of clinical information varied across patients. As a result, phenotypic bias—particularly for conditions represented by a small numbers of cases—cannot be excluded.

Dual diagnoses and their phenotypic impact

Through parallel analysis, 17 of 171 patients with repeat-associated ataxia were found to carry an additional potentially pathogenic repeat expansion, either in the same or a different gene. The phenotypic impact of these additional repeat expansions is of particular interest. In this study, all additional expansions in other loci fell within the range of reduced penetrance or had unconfirmed

pathogenicity, raising the possibility that they do not contribute to the phenotype, as has been recently described for FGF14 expansions in patients with FRDA.⁶¹ Biallelic pathogenic expansions in autosomal-dominant ataxias were previously reported and were also observed in this cohort. In patients with SCA6 and SCA27B, the second expansion is likely to act as a disease modifier, increasing disease severity as previously reported.^{34,62,63} However, no such modifying effect could be confirmed for patients with SCA8.⁶⁴ Given the phenotypic overlap and variability in disease severity, longitudinal studies in larger cohorts, including family-based investigations, are needed to confirm the potential effects of dual diagnoses.

High carrier frequency of RFC1 spectrum disorder

The high RFC1 carrier frequency of 7.2% for spectrum disorder is striking, corresponding to an estimated prevalence of 1:772. In principle, this high carrier frequency in a cohort of ataxia patients could indicate monoallelic pathogenic expansions in RFC1 as a risk factor for sporadic ataxia or that a pathogenic variant on the second allele and, therefore, RFC1 spectrum disorder may be missed in some ataxia patients. However, the carrier frequency in this cohort aligns with that observed in a large cohort of 29 496 controls with European population background (8.0%) analysed by LRS.⁶⁵ This study can be assumed to be more representative than earlier studies, which observed lower heterozygote carrier frequencies of 0.7–6.5%.²³ As such, an enrichment of monoallelic pathogenic expansions in RFC1 in ataxia patients not affected by RFC1 spectrum disorder is unlikely. Compared with an estimated prevalence of sporadic adult-onset ataxias (2–12:100 000) and autosomal recessive hereditary ataxias (1–9:100 000), the estimated prevalence of RFC1 spectrum disorder exceeds that of both conditions together by at least 6-fold.^{66–68} This discrepancy may reflect underdiagnosis of RFC1 spectrum disorder due to mild or atypical presentation without ataxia (e.g. isolated neuropathy or vestibulopathy) or incomplete penetrance of biallelic pathogenic repeat expansions.⁶⁹ Longitudinal screening of older, unaffected individuals with biallelic repeat expansions could help to address this question.

Reassessing FRDA carrier frequency

For FRDA, we determined a carrier frequency of 0.8% (1:125) in our cohort, based on the assumption that short, heterogeneous alleles containing alternative motifs such as (GAAGGA)_{exp} are non-pathogenic. The observed carrier frequency for FRDA in our cohort is lower than reported estimates (1:60–1:100).⁷⁰

The most likely explanation for this discrepancy is statistical variation due to our cohort size. However, we also hypothesize that earlier studies may have overestimated carrier frequencies due to limitations of PCR-based genotyping methods, which cannot reliably distinguish pathogenic pure (GAA)_{exp} alleles from non-pathogenic expansions with alternative repeat motifs. Consequently, inclusion of such non-pathogenic alleles in previous analyses may have artificially inflated carrier frequency estimates.

To assess whether heterozygous carrier frequencies for FRDA can be directly translated into disease prevalence, future studies should evaluate the clinical relevance of biallelic short expansions (<100 repeat units). In our cohort, such short alleles were found among carriers at similar frequencies as large expansions. It remains to be determined whether these biallelic short expansions are associated with very late-onset FRDA in only a small subset of individuals, or whether they exhibit high penetrance more broadly.

Challenges in the assessment of repeat expansions

Advances in LRS-based repeat analysis highlight unresolved challenges in assessing repeat expansions, particularly in determining their pathogenicity, penetrance and the prediction of age of onset, severity and phenotypic presentation. These hurdles are pronounced for recently described conditions like RFC1 spectrum disorder and SCA27B, where a significant number of diagnoses remain inconclusive. For SCA27B, a major issue is the uncertain threshold of pathogenicity. While current studies set the threshold for pathogenicity with reduced penetrance at 250 RU, other studies suggest lower values (180 or 200 RU).^{15,16,34,52} Establishing a definitive threshold of pathogenicity is challenging due to the extensive mosaicism of repeat expansions in FGF14. Additionally, inherent biases in both PCR-based methods and amplification-free LRS techniques can lead to slight variations in the median repeat length (Supplementary Fig. 1). Consequently, studies aimed at determining length thresholds must carefully benchmark their methods to account for these potential discrepancies. In this study, 35 patients carried expansions in the 180–249 RU range, which require further validation regarding an SCA27B diagnosis. Additionally, novel FGF14 repeat configurations, such as (GGA)₇₅(GAA)₃₀₀, observed in this study, which carry large continuous stretches of the pathogenic motif, require further investigation.

In RFC1 spectrum disorder, up to 24 distinct repeat motifs have been identified (confirmed 12 motives in Supplementary Table 1), some extremely rare or highly heterogeneous, complicating pathogenicity assessment.^{19,22,24,25,71} In this study, 24 alleles (2.3%) carried repeat configurations of unknown significance, such as [(ACGGG)_{complex}, (AACGG)_{exp}, (ACAAG)_{exp} and (AGGGG)_{exp}]. Most patients carried a non-pathogenic repeat configuration on the second allele, largely excluding RFC1 spectrum disorder. However, one individual carried repeat configurations of unknown significance on both alleles [(ACGGG)_{complex}/(AACGG)_{exp}], hindering the exclusion of RFC1 spectrum disorder. Based on the assumption that GC content and repeat length determine pathogenicity, we speculate that the 80 RU 'AACGG' expansion is non-pathogenic, as it has the same GC content as the pathogenic 'AAGGG' motif but is shorter than the current pathogenicity threshold (250 RU).¹⁹ Similarly to FGF14, LRS deciphered composite expansions in RFC1, such as (AAAGG)₃₁₇(AAGGG)₂₁₀, which require an assessment of pathogenicity. As the formation of stable G-quadruplex structures has been associated with the pathogenic 'AAGGG' motif, analysing secondary RNA/DNA structures across all known motifs could contribute to the assessment of their clinical significance in conjunction with further elucidating the pathomechanism of RFC1 spectrum disorder.^{72–74} Adding to the complexity of RFC1 repeat expansion assessment, two motifs ('AAGGG' and 'AAAGG'), previously assumed as solely pathogenic or non-pathogenic, respectively, are now known to be pathogenic depending on their length.¹⁹ Current thresholds of pathogenicity require further confirmation and refinement, e.g. through large (meta)analyses.

A significant proportion of ataxia patients with likely causative repeat expansion, 44% (75 out of 171 patients) in this study, carry reduced-penetrance expansions in loci such as ATXN8/ATXN8OS, TBP, FGF14 and FMR1. Emerging data indicate that not only incomplete penetrance but also repeat expansions with currently assumed full penetrance are frequent in healthy individuals.^{65,75} Genetic findings must, therefore, be critically evaluated in the context of clinical findings, phenotype and differential diagnoses as penetrance might be lower than currently assumed, leading to a potentially unrelated coincidence of adult-onset ataxia and a repeat expansion associated with ataxia.

Given the phenotypic overlap in patients with adult-onset ataxia hindering clinical classification, molecular genetic diagnoses should be refined and may benefit from the inclusion of additional genetic parameters. Depending on the underlying pathomechanism, the prediction of penetrance and disease severity may be improved by analysing flanking regions, repeat interruptions, alternative repeat motifs, somatic mosaicism and methylation. For example, in the case of Huntington's disease, somatic instability contributes for variability in penetrance, age of onset and clinical severity, factors that extend beyond repeat length alone.^{76–78} The same might apply to certain SCAs, as indicated by a recent study.⁷⁹ The degree of somatic instability may be inferred from repeat length heterogeneity in blood, as accessible through Clin-CATS. Similarly, indirect markers, such as repeat interruptions and DNA repair gene variants, may further indicate repeat stability.^{80,81} For repeat disorders driven by loss-of-function mechanisms, such as FRDA and likely SCA27B, methylation patterns of regulatory regions could serve as markers of residual gene expression, potentially predicting penetrance and clinical severity.^{82,83} In addition to investigating additional parameters, pathogenic variants in other genes might impact or determine the phenotype, as described for SCA48, where intermediate alleles in *TBP* likely act as disease modifiers.³⁶

Targeted LRS as a superior tool for repeat analysis and remaining technological challenges

While NGS has fundamentally changed the detection of SNVs, methods for repeat analysis have stagnated for decades. The limitations of traditional PCR-based approaches are particularly evident in complex repeat disorders such as RFC1 spectrum disorder. Even combining several PCR-based analyses could not reliably determine disease and carrier status for all patients analysed in this study due to the inability to measure expansions larger than 150–200 RU and the restriction of the screening to a limited set of repeat motifs. Recent bioinformatic methods, such as ExpansionHunter, offer a valuable tool for screening repeat expansions in short-read genomes.⁸⁴ However, these methods cannot accurately determine the repeat length of larger expansions and repeat interruptions due to read length limitations. Moreover, while their sensitivity and specificity are high, they remain lower compared with PCR-based methods, resulting in a significant false-positive rate, which necessitates confirmation by complementary methods.^{85,86}

LRS has the potential to overcome persisting limitations while enabling the incorporation of additional genetic features, such as repeat interruptions and methylation. With decreasing costs and increasing sequencing accuracy, long-read genomes will likely become the gold standard in the diagnosis of hereditary ataxias, as repeat-associated ataxias can be detected in parallel to other genetic variants. However, current targeted approaches are required for cost-efficiency and precision. Current LRS platforms include ONT or Pacific Biosciences, with several targeted approaches including CRISPR/Cas9-based enrichment, and ONT bioinformatic-based enrichment with adaptive sampling.⁸⁷ However, a comprehensive benchmarking, comparison and clinical validation of all LRS approaches is currently lacking. Individual studies highlight specific limitations, such as repeat expansions in *FXN* or *RFC1* carrying specific motifs missed in PacBio genome sequencing or missed expansions in adaptive sampling due to low coverage.^{85,88}

In contrast, Clin-CATS, based on CRISPR/Cas9 target enrichment, ensures high coverage of all repeat loci and replaces at least 18 individual PCR analyses with a single, more comprehensive test. However, sizing very large expansions, such as the pathogenic

motif 'AAGGG' in *RFC1*, remains a challenge. While Southern blotting has detected expansions as large as 3885 RU, no 'AAGGG'-repeat expansion larger than 1395 RU was detected in our study.⁸⁹ This could be explained by the fragmentation of large repeat expansions composed of the pathogenic 'AAGGG' motif, potentially as a consequence of its tendency to form G-quadruplex DNA structures in combination with ONT's bias toward shorter fragments. As the presence of expansions above the threshold of pathogenicity is reliably detected also, in these cases, the diagnostic precision of Clin-CATS is not affected by this limitation. Further optimization could extend the upper limit of detectable expansions, but complementary methods such as Southern blotting or Optical Genome Mapping may still be required for extremely large expansions.⁹⁰ Finally, to maintain the robust performance of Clin-CATS in routine diagnostics, we adapted the workflow to be compatible with the new R10 flow cell chemistry following the discontinuation of the R9 flow cell in a recent work.⁹¹ Improvements to the library preparation protocol and bioinformatics pipeline have ensured the continuation of Clin-CATS with the latest technology.

Conclusion

This study found and confirmed the high prevalence of repeat disorders as the cause of adult-onset ataxia, with SCA27B and *RFC1* spectrum disorder as the most common conditions. We identified several novel genotypes and variations of the known repeat configurations, further illustrating the genetic heterogeneity of repeat-associated ataxias. Our study also highlights the methodological advantages of LRS-based repeat analysis with Clin-CATS, showing a stable and robust performance and cost-efficiency. Given the demonstrated superiority of parallel analysis with LRS, we strongly advocate for its adoption as first-tier analysis in patients with adult-onset ataxia before performing NGS exome or genome analysis.

Data availability

Anonymized data from this study are available from the corresponding author on reasonable request.

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The other authors report no competing interests.

Supplementary material

Supplementary material is available at [Brain](#) online.

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