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Bioinformatics-Based Identification of Expanded Repeats: A Non-reference Intronic Pentamer Expansion in RFC1 Causes CANVAS

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Genomic technologies such as next-generation sequencing (NGS) are revolutionizing molecular diagnostics and clinical medicine. However, these approaches have proven incomplete in identifying pathogenic repeat expansions. Here, we apply a collection of bioinformatics tools that can be utilized to identify either known or novel expanded repeat sequences in NGS data. We performed genetic studies of a cohort of 35 individuals from 22 families with a clinical diagnosis of cerebellar ataxia with neuropathy and bilateral vestibular areflexia syndrome (CANVAS). Analysis of whole-genome sequence (WGS) data with five independent algorithms identified a recessively inherited intronic repeat expansion [(AAGGG)<sub>exp</sub>] in the gene encoding Replication Factor C1 (RFC1). This motif, not reported in the reference sequence, localized to an Alu element and replaced the reference (AAAAAG)<sub>11</sub> short tandem repeat. Genetic analyses confirmed the pathogenic expansion in 18 of 22 CANVAS-affected families and identified a core ancestral haplotype, estimated to have arisen in Europe more than twenty-five thousand years ago. WGS of the four RFC1-negative CANVAS-affected families identified plausible variants in three, with genomic re-diagnosis of SCA3, spastic ataxia of the Charlevoix-Saguenay type, and SCA45. This study identified the genetic basis of CANVAS and demonstrated that these improved bioinformatics tools increase the diagnostic utility of WGS to determine the genetic basis of a heterogeneous group of clinically overlapping neurogenetic disorders.

Introduction

Repetitive DNA sequences constitute approximately one third of the genome and are thought to contribute to diversity within and between species.1 Microsatellites or short tandem repeats (STRs) are mini-repeats of DNA, typically two to five base-pairs in length, which are usually present in a concatamer of between 5 and 50 repeated elements. There are thousands of STRs scattered through the human genome and recent studies have suggested important roles for STRs in the regulation of gene expression.2,3 STRs display considerable variability in length between individuals, which is presumed to have no detrimental consequences for humans,4,5 unless the repeat number is expanded beyond a gene-specific threshold.6,7 Pathogenic repeat expansions (REs) have been shown to underlie at

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least 30 inherited human diseases, the majority being disorders of the nervous system. These disorders, which vary from autosomal-dominant, autosomal-recessive, and X-linked inheritance, have an overall prevalence of ~1:20,000. They display a broad onset age and are characterized by progressive cerebellar ataxia with dysarthria, oculomotor abnormalities, cognitive dysfunction, and other symptoms (see GeneReviews in Web Resources). Additional novel pathogenic REs likely remain to be identified. For example, putative spinocerebellar ataxia (SCA) loci, including SCA25 (MIM: 608703) and SCA30 (MIM: 613371), remain to be identified, and unsolved hereditary ataxias such as cerebellar ataxia with neuropathy and bilateral vestibular areflexia syndrome (CANVAS [MIM: 614575]) display extensive clinical similarities with known RE disorders.

CANVAS is a cerebellar ataxia with combined cerebellar, vestibular, and somatosensory dysfunction. Historically, individuals with CANVAS have been assigned the diagnosis of idiopathic late-onset cerebellar ataxia. More recently, CANVAS is clinically recognized and has been incorporated into the contemporary research and teaching of both cerebellar and vestibular diseases. Unifying the otocerebellar and neuropathology, CANVAS is a neuronopathy (ganglionopathy) affecting the vestibular and dorsal root ganglia. The progression of these clinical features can be measured longitudinally using a specific neurophysiological protocol. A characteristic radiological pattern of cerebellar atrophy has also been described and verified on post-mortem pathology. The characteristic oculomotor abnormality seen in combined cerebellar and vestibular impairment is the visually enhanced vestibulo-ocular reflex (VVOR), and this can now be evaluated using a commercially available instrumented assessment tool. Altogether, these advances have allowed the formulation of diagnostic criteria to aid identification of CANVAS, contributing both research and clinical benefits including improved prognostication and targeted management.

The majority of individuals and families with CANVAS have been identified in individuals of European ancestry, although CANVAS has recently been reported in two individuals of Japanese ethnicity, a 68-year-old male and a 76-year-old female. A genetic cause of CANVAS is highly plausible given the observation of 13 affected siblings and families with multiple affected individuals over several generations. The pattern of inheritance suggests an autosomal-recessive trait, although autosomal-dominant inheritance with incomplete penetrance cannot be excluded. CANVAS symptoms overlap considerably with SCA3 (also known as Machado-Joseph disease [MIM: 109150]) and Friedreich ataxia (MIM: 229300), both genetic forms of ataxia caused by the inheritance of a pathogenic RE. These observations are consistent with the hypothesis that a novel pathogenic STR expansion may underlie CANVAS.

Historically, the detection of REs has been time consuming and expensive. Indeed, it is only in recent years that computational methods have been developed to screen for RE in short-read whole-exome sequence (WES) and WGS data, leading to the discovery of previously undescribed, disease-causing REs. For example, a pentanucleotide RE was identified to underlie autosomal-dominant spinocerebellar ataxia 37 (SCA37 [MIM: 615945]). Moreover, pathogenic REs of intronic pentamers (TTTCA)n and (TTTAA)n were identified as the cause of benign adult familial myoclonus epilepsy locus 1, 6, and 7 (BAFME1 [MIM: 618073], BAFME6 [MIM: 618074], BAFME1 [MIM: 618075]).

A number of bioinformatics tools now exist that allow screening of short-read sequencing data for expanded STRs. Initially, STR detection tools, such as lobSTR and hipSTR, were limited to short STRs that were encompassed by a single sequencing read. However, in the last two years, multiple methods have been released that can screen WES and WGS datasets for REs without being limited by read length. These include ExpansionHunter (EH), exSTRa, TREDPARSE, STRetch, and GangSTR. These are all reference based methods, i.e., they rely on a catalog of STR loci and motifs and are therefore limited to detecting expansion of previously defined STRs, such as those cataloged in the UCSC track. Moreover, the normal variability in STR length and repeat composition remains poorly described, particularly for rare STRs or those larger than ~100 bp. Therefore, there is a need for bioinformatics tools that are unbiased to the limited catalogs of STR loci available. Ideally, these tools will be able to search genome-wide for expanded repeat sequences in NGS data, independent of prior knowledge of either the location or composition of the RE. Here, we utilized a STR reference-free method called Expansion Hunter De Novo (EHdn), in combination with multiple reference-based tools, to show that CANVAS is caused by the homozygous inheritance of a previously undescribed expanded intronic pentamer (AAGGG)exp in the gene encoding Replication Factor C Subunit 1 (RFC1). An independent study, published while this work was under review, similarly identified this causal pentamer in RFC1. Cortese and colleagues defined a small linkage region from ten families with CANVAS and the causative RE was identified by WGS and visual inspection of the aligned read pairs inside the linkage region.

### Material and Methods

#### Recruitment, Linkage, and Next-Generation Sequence Data
The Royal Children’s Hospital Human Research Ethics Committee approved the study (HREC 28097). Informed consent was obtained from all participants and clinical details were collected from clinical assessments and review of medical records. Genomic DNA was isolated from peripheral blood. Single-nucleotide polymorphism (SNP) genotype data were generated for two affected......
siblings from three families (CANVAS1, 2, and 3) and all six siblings from family CANVAS4 using the Illumina Infinium HumanOmniExpress BeadChip genotyping array. SNP genotypes for individuals from CANVAS9 were extracted from WES data. Parametric multipoint linkage analysis was subsequently performed using LINKDATAGEN and MERLIN, specifying a rare recessive disease model with complete penetrance, and overlapping linkage signals were detected using BEDtools. WES was performed on individuals from CANVAS9 using Agilent SureSelect XT Human All exon V5 + UTR on the Illumina HiSeq2000 platform at 50× mean coverage. WES was performed on an additional 23 individuals in 15 families in collaboration with the Johns Hopkins Center for Inherited Disease Research (CIDR) as part of the Baylor-Hopkins Center for Mendelian Genomics (BHCMG). WGS was performed in two stages. Libraries for the first round of samples, including two affected individuals from CANVAS1 and CANVAS9 and 31 individuals lacking a clinical diagnosis of CANVAS (subsequently referred to as control subjects although some have a diagnosis other than CANVAS), were prepared using the TruSeq nano PCR-based Library Preparation Kit and sequenced on the Illumina HiSeq X platform. Libraries for the second round of WGS, including affected individuals with evidence of an alternate RE motif (CANVAS2 and 8) or lacking the pathogenic RE in RFC1 (CANVAS11, 13, 17, and 19), were prepared using the TruSeq PCR-free DNA HT Library Preparation Kit and sequenced on the Illumina NovaSeq 6000 platform. PCR-free WGS data from 69 unrelated Coriell control subjects28 was obtained from Illumina. GTEx samples (SRA files, 133 WGS with matching cerebellar RNA-seq) were downloaded from the dbGAP (phs000424.v7.p2).

Alignment and Variant Calling
Alignment and haplotype calling were performed based on the GATK best practice pipeline. All WES and WGS datasets were aligned to the hg19 reference genome using BWA-mem, then duplicate marking, local realignment, and recalibration were performed with GATK. Merged VCF files were annotated using vcfanno and ANNOVAR. Candidate variant filtering was performed using CAVALIER, an R package for variant interpretation in NGS data (Web Resources). Standard variant calling was performed on WGS data for CANVAS samples negative for the pathogenic RE in RFC1. Candidate variants were defined as (1) occurring in known ataxia genes, as defined by OMIM, (2) exonic, with a minor allele frequency of less than 0.0001 in gnomAD (both genome and exome data), and (3) predicted pathogenic by both SIFT and PolyPhen2. RNA-seq data was aligned to the hg19 reference genome (ENSEMBL, Homo_sapiens.GRCh37.75) using STAR. 39

STR Analysis
Genome-wide screening for putative REs was performed using Expansion Hunter Denovo (EHDn) v.0.6.2, an open-source method that is being developed by Illumina, the Walter Eliza Hall Institute, and others. EHDn operates by performing a genome-wide search for read pairs where one mate has confident alignment (anchor) and the second mate consists of repetition of a repeat motif (in-repeat read). The program reports the counts of in-repeat reads with anchor mates stratified by the repeat motif and genomic position of their anchor mate. For this analysis we defined a confidently aligned read as one aligned with MAPQ of 50 or above. The counts of in-repeat reads with anchor mates were subsequently compared for each region in case subjects (CANVAS) and control subjects using a permutation test (10,000 permutations). The resulting p values were used to rank candidate sites with higher counts in individuals with CANVAS than in the control subjects for further computational validation. These candidates were subsequently annotated with ANNOVAR.

Computational validation was performed using five independent STR detection tools for short-read NGS after updating the STR catalog reference files to incorporate the identified motifs. The RE candidates were screened in the two individuals with CANVAS and the 31 non-CANVAS control subjects using exSTRA and EH, then the top candidate [(AAGGG)exp STR in RFC1] was further validated with TREDPARSE, GangSTR, and STRetch. All tools were used with default parameters, with the following additional parameters for EH: read-depth of 30 and min-anchor-mapq of 20. All five tools were also used to screen for the (AAGGG)exp RFC1 STR in the 69 Coriell control WGS datasets. A short-list of (AAGGG)exp carriers was generated based on consensus calling from at least four of the five tools.

Individuals diagnosed with CANVAS lacking the (AAGGG)exp RFC1 RE were further screened with EHDn for novel STRs and for known pathogenic STRs using exSTRA and EH. The WES datasets could not be analyzed for the (AAGGG)exp RFC1 RE as the intrinsic locus (chr4:39350045–39350095, hg19) was not captured during library preparation. However, the region was visualized in the alignment using the Integrative Genomics Viewer (IGV) to identify potential off-target reads, which could provide supportive evidence for the presence of the (AAGGG)exp motif. Only samples with at least one read mapping at the STR in the RFC1 locus were considered.

Haplotyping and Mutation Dating
Haplotyping was performed on the WES data. Variants were filtered based on read depth (≥30), including both exonic and non-exonic variants. A core haplotype was defined based on sharing among a majority of affected individuals. A method based on haplotype sharing42 was used to determine the most recent common ancestor (MRCA) from whom the core haplotype was inherited, as well as dating additional sub-haplotypes shared by clusters of individuals, which are likely to be individuals with a MRCA who is more recent than that for the whole group (see Mutation dating in Web Resources).

Molecular Genetic Studies
We designed a PCR assay to test for presence of additional inserted sequence, not present in the reference database, at the RFC1 STR. The primers (Table S1) flank the STR and are predicted to amplify a 253 bp fragment using standard PCR conditions with a 30 s extension cycle. Presence of the pathogenic RFC1 RE was tested by repeat-primed PCR utilizing three primers: TPP_CANVAS_FAM_2F, 5R_TPP_M13R_CANVAS_RE_R, and TPP_M13R (Table S1). The FAM labeled forward primer is locus specific, while the repeat-specific primer (5R_TPP) includes a tag M13R sequence. PCR was performed in a 20 μL reaction with 20 ng genomic DNA, 0.8 μM of both the FAM labeled forward primer and TPP_M13R, and 0.2 μM 5R_TPP using GoTaq Long PCR Polymerase (Promega). A standard 60TD55 protocol was utilized (94°C denaturation for 30 s, 60TD55°C anneal for 30 s, and 72°C extension for 2 min), products were detected on an ABL3730xl DNA Analyzer and visualized using PeakScanner 2 (Applied Biosystems).
Results

Case Recruitment

The workflow for this study is summarized in Figure 1. Individuals with a clinical diagnosis of CANVAS were recruited following neurological assessment and investigation in accordance with published guidelines.43 While variable between case subjects, data leading to the clinical diagnosis included evidence of combined cerebellar and bilateral vestibular impairment, cerebellar atrophy on MRI, neurophysiological evidence of impaired sensory nerve function, and negative genetic testing for pathogenic RE at common SCA loci (typically SCA1, 2, 3, 6, and 7) and FRDA (Friedreich ataxia, FRDA). In total, the cohort consisted of 35 individuals with a clinical diagnosis of CANVAS (Table 1). The individuals came from 11 families with a single affected individual, 7 families with affected sib pairs and 4 larger/multigenerational families (Figure S1). A full clinical description of the cohort will be reported in a forthcoming manuscript.

Linkage Analysis

CANVAS typically presents in families with one or multiple affected individuals in a single generation, consistent with a recessive inheritance. For example, in the second-degree consanguineous family CANVAS9, four siblings were diagnosed with CANVAS and two were classified as unaffected at the time phenotyping was performed (Figure 2A). Parametric multipoint linkage analysis was performed on five CANVAS-affected families (CANVAS1, 2, 3, 4, and 9; Figure S1) specifying a rare recessive disease model with complete penetrance. This identified linkage regions with logarithm of odds (LOD) scores ranging from 0.6 for smaller pedigrees (two affected siblings) to a statistically significant linkage region on chromosome 4 in CANVAS9 (LOD = 3.25, Figure 2B). Intersection of the linkage regions from the five families identified a single region on chromosome 4 (chr4:38941465–40390306, combined LOD = 7.04) common to all families (Figure 2C). CNV analysis utilizing PennCNV did not identify any potential copy number variants in the minimal linkage region.44 The 1.5 Mb shared region contains 42 genes, of which 14 are protein coding, none with any association with ataxia in OMIM or the published literature (Table S2).

Large-Scale WES Analysis Did Not Identify Candidate Pathogenic Variants

WES was used to screen 27 affected individuals with CANVAS from 15 families for potentially pathogenic rare variants (MAF < 0.001) shared across multiple pedigrees.
in a homozygous or compound heterozygous inheritance pattern. No candidate mutations were detected, either within the chromosome 4 linkage region or elsewhere in the genome.

Identification of an Undescribed (AAGGG)exp RE in the Linkage Region

The lack of candidate variants identified from the WES data suggested the possibility of (1) intronic or intergenic mutations or (2) that CANVAS might be caused by a non-standard mutation, such as a pathogenic RE of an STR. Therefore, WGS was performed on two individuals from different pedigrees (CANVAS1 and 9) who share the chr4 linkage region. EHdn was used to perform a genome-wide screen for STRs in the two individuals with CANVAS compared to WGS data from 31 unrelated control subjects. This identified 19 regions with a p value < 0.005 (Table S3), although genome-wide significance could not be achieved after adjustment for multiple testing due to the skewed ratio of the number of case subjects to control subjects (2 versus 31). These candidate STRs were visualized with the Integrative Genomics Viewer (IGV) tool, which suggested that the (AAGGG)exp STR within intron 2 of the gene encoding Replication Factor C1 (RFC1) was likely real and present in both alleles in the affected individuals, consistent with the recessive inheritance pattern hypothesized for CANVAS (Figure S2). In both individuals with CANVAS, the (AAAAG)11 motif is substituted by the (AAGGG) exp motif, with potential interruptions to the Alu element (Figure 2D).

<table>
<thead>
<tr>
<th>Family</th>
<th>Participants (Sex)</th>
<th>SNP Array</th>
<th>WES</th>
<th>WGS</th>
<th>RFC1 STR in WES</th>
<th>PCR Wild-Type Allele</th>
<th>Repeat-Primed PCR</th>
<th>Genetic Diagnosis</th>
<th>Haplotype</th>
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<td>Y</td>
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Abbreviations: M, male; F, female; U, deidentified; NA, not applicable; ND, not detected; Other*, a different haplotype OR shortened A haplotype. The gene reference sequences utilized were GenBank: NC_000004.11 and NM_002913.4 (RFC1).
Confirmation of (AAGGG)\textsubscript{exp} STR in Off-Target WES Reads

While WGS was only performed in two individuals with CANVAS, the majority of the cohort (n = 27) was analyzed by WES. The putative pathogenic CANVAS RE is located in intron 2 of \textit{RFC1}, 2,863 bp downstream of exon 2 and 2,952 bp upstream of exon 3. Therefore WES data are \textit{a priori} assumed to be uninformative for this RE as it is not targeted during DNA capture. However, given that WES data includes off-target reads, we hypothesized that some reads might map to the \textit{RFC1} RE locus. Visual assessment of the WES data in IGV identified 14 individuals with informative reads; the maximum off-target read coverage at this locus was two, with a median of one read. While 3 individuals only had reads that correspond to the reference genome STR sequence (AAAAG), 11 affected individuals from 9 families had reads containing (AAGGG) repeats (Table 1). Furthermore, single affected individuals from families CANVAS2 and CANVAS8 had single, independent reads identifying (AAGGG) and (AAAGG) motifs at the RFC1 STR locus. This observation raised the possibility that CANVAS might result from pathogenic expansions of different pentanucleotide motifs.

Computational Validation with Existing STR Detection Tools

Multiple tools have been developed in recent years that test for the presence of REs at pre-defined STRs. Therefore, we inserted the \textit{RFC1} STR motifs into the STR reference files and used exSTRa, EH, TREDPARSE, STRetch, and GangSTR to estimate the size of the STR and/or detect REs in the WGS data from the two original CANVAS samples (CANVAS1 and 9) and seven additional individuals with CANVAS. The seven additional CANVAS samples selected for WGS were those with WES evidence for an alternate...
The library preparation of this second round of WGS was PCR free as PCR amplification has previously been shown to affect RE detection.8 Using exSTRa, we confirmed the homozygous inheritance of the (AAGGG)exp RE in three individuals (CANVAS1, 2, and 9; Figure 3). The empirical cumulative distribution function (ECDF) pattern for CANVAS2 is consistent with the presence of one shorter and one longer (AAGGG)exp RE, while CANVAS8 appears to only have a single (AAGGG)exp allele. Screening of all datasets for the (AAAGG)exp motif at the chr4 RFC1 locus using exSTRa identified an expansion of this STR only in CANVAS8, suggesting that this individual is a compound heterozygote, with an (AAGGG)exp RE on one allele and an (AAAGG)exp RE on the other. Visualization in IGV confirmed the presence of the (AAAGG)n motif embedded within the reference STR: (AAAAG)6-(AAAGG)n-(AAAAG)6 (Figure S3). This observation raises the possibility that an expanded (AAAGG)exp motif might also be associated with CANVAS.

EH, TREDPARSE, and GangSTR were used to estimate the length of the AAGGG motif on each allele (Figure 3). The results were highly variable depending on the tool used. EH reported minimum and maximum allele sizes ranging from 30 to 68 in individuals with the RE. The allele size ranges estimated by GangSTR and TREPARSE were 2 to 27 and 7 to 14, respectively. Furthermore, all three tools inferred the presence of two alleles, even in individuals who carry a single allele, and hence do not appear to be distinguishing read contributions between the alleles, also contributing to unreliable size estimates. Reads comprised of the (AAGGG)exp motif in particular also showed evidence of high read sequencing error. Based on these results, we can infer that while the (AAAGG) motif (families CANVAS2 and 8), and those who did not appear to have a RE at the RFC1 locus based on the PCR/RP-PCR studies described below (CANVAS11, 13, 17, and 19 families). The library preparation of this second round of WGS was PCR free as PCR amplification has previously been shown to affect RE detection.8 Using exSTRa, we confirmed the homozygous inheritance of the (AAGGG)exp RE in three individuals (CANVAS1, 2, and 9; Figure 3). The

Figure 3. Computational Validation of the (AAAGG)exp RE

The (AAAGG)exp RE at the coordinates chr4:39350045–39350095 was added to the reference databases of the tools exSTRa, EH, GangSTR, TREDPARSE, and STReach and WGS data from four unrelated individuals with CANVAS was analyzed (CANVAS11, orange; CANVAS2, blue; CANVAS8, red; and CANVAS9, green). The non-CANVAS control subjects are presented in gray. Plots have been divided into PCR-based and PCR-free WGS (left and right columns, respectively). The Y and X axes for ExpansionHunter, GangSTR, and TREDPARSE refer to the number of repeat units on the longer and shorter allele per individual, respectively. The Y axis for the STReach plot refers to the number of individuals.
CANSV samples were all correctly identified as having homozygous RE at RFC1; estimates of expansion size are inconsistent and appear likely to significantly underestimate the actual repeat size.

The consensus of the different tools was that CANSV11, 13, 17, and 19 families did not encode a pathogenic RE [either (AAGGG)exp or (AAAGG)exp] at the RFC1 locus, which we confirmed by PCR analyses (see below). However, the RFC1 (AAGGG)exp RE was present in three of the control WGS datasets (two heterozygous, one homozygous, allele frequency ~0.06 [4/62]; Figure 3). No control individuals were identified to carry the (AAAGG)exp motif. As with the CANSV samples, the STR sizing estimates using the different tools was inconsistent, so no conclusions could be drawn from this in silico analysis regarding the relative size of the (AAGGG)exp RE in control subjects compared to individuals with CANSV.

We then analyzed a larger in-house collection of unrelated control Coriell WGS samples (n = 69) and again failed to identify the (AAAGG)exp motif. However, we identified six individuals heterozygous for the (AAAGG)exp RE, representing a frequency estimate of ~0.04 (6/138; Figure S4). Using the NGS QC software tool peddy, we found evidence that two of these heterozygous individuals are of European ancestry and that two further individuals are of admixed Native American ancestry. Finally, we accessed WGS from GTEx for 133 individuals who have matching brain (cerebellum) RNA-seq. Our analysis identified 11 heterozygous individuals who have matching brain (cerebellum) RNA-seq. (AAAGG)exp RE in control subjects compared to individuals with CANSV.

Validation of the (AAGGG)n RE as the Causal Variant for CANSV

We developed a PCR assay that amplifies across the repeat tract to rapidly screen for the presence of a non-expanded allele at the RFC1 STR. Although the screen does not distinguish between the (AAAGG)11 reference STR, the (AAGGG) STR, or any other potential motif, amplification of an ~250 bp fragment indicates at least one allele is not expanded. Moreover, the presence of two distinct non-expanded products is indicative of a heterozygous non-mutant state. Conversely, the complete absence of the PCR product provides indirect evidence of a RE affecting both alleles of the RFC1 locus. Analysis of all available DNA samples from individuals with CANSV suggested that the reference STR at the RFC1 locus was not present in 30 clinically diagnosed individuals from 18 (of 22) CANSV-affected families (Table 1, Figure 4). Notably, unaffected individuals from RFC1-positive families carried at least one non-expanded RFC1 allele (Figure S5). To directly confirm expansion of the (AAGGG) motif in RFC1, we developed a locus-specific repeat-primed PCR assay, using a primer located adjacent to the RFC1 repeat and an AAGGG-specific primer. Consistent with the PCR assay, affected individuals from the 18 families demonstrated a saw-toothed “ladder” when the repeat-primed PCR products were analyzed by capillary array (Table 1, Figure 4).

These results suggest that a homozygous RE underlies CANSV in these 18 families, and at least one pathogenic allele encodes the (AAGGG)exp RE. Molecular analysis of the DNA for the three in-house control individuals with the in silico predicted (AAGGG)exp motif (Figure 3) demonstrated a ~250 bp product in both heterozygous samples but no product in the homozygous individual. The repeat-primed assay demonstrated a saw-toothed ladder in all three samples (Figure S6). Collectively, these analyses suggested all three control individuals have at least one copy of the pathogenic (AAGGG)exp RE at RFC1, although the size of the RE cannot be determined by these analyses.

In four families (CANSV11, 13, 17, and 19), the presence of the expected reference PCR amplicon and lack of a repeat-primed PCR product suggested that the pathogenic RFC1 RE was not present on either allele. This implied that these individuals have a different CANSV-causing mutation in RFC1, or there is locus heterogeneity. A third possibility is that they do not have CANSV but instead a related ataxia. Therefore, we performed WGS on these individuals and initially screened for known REs associated with ataxias using exSTra and EH. A CAG trinucleotide expansion in ATXN3, associated with spinocerebellar ataxia type 3 (SCA3 [MIM: 109150], also known as Machado-Josephs disease) was identified in CANSV13 (Figure S7) and confirmed by diagnostic testing. The WGS was then screened for unreported or rare SNPs and indels in genes known to cause ataxia. No de novo or rare variants were identified in RFC1, but a potential genomic re-diagnosis was achieved in two additional families. In CANSV17, two variants (GenBank: NM_001278055; c.12398delT [p.Phe4133Serfs*28] and GenBank: NM_001278055; c.5306T>A [p.Val1769Asp]) were identified in the gene encoding saccin (SACS) and segregation analysis confirmed they were in trans. Biallelic mutations in SACS cause spastic ataxia of the Charlevoix-Saguenay type (MIM: 270550). In CANSV19, a heterozygous variant in the gene encoding FAT tumor suppressor homolog 2 (FAT2; GenBank: NM_001447.2; c.4370T>C [p.Val1457Ala]) was identified. Heterozygous mutations in FAT2 have recently been associated with SCA45 (MIM: 604269). No potentially pathogenic variants were identified in CANSV11, but a variant of unknown significance was identified in the gene encoding Ataxin 7 (ATXN7; GenBank: NM_001177387.1; c.2827C>G [p.Arg943Gly]). CANSV11 was also screened genome-wide with EHDn for potentially pathogenic unreported RE, but no additional candidate REs were identified.

A Single Founder Event for the (AAGGG)n RE in RFC1

We performed haplotype analysis to determine whether the (AAGGG)exp RE arose more than once in human history. Analysis of haplotypes inferred from the WES data identified a core ancestral haplotype, comprised of 27 SNPs (Figure 5A), that was shared by most individuals except CANSV14 (Table 1, Table S4). The core haplotype
spans four genes (TMEM156, KLHL5, WDR19, and RFC1) and is 0.36 Mb in size (chr4:38995374–39353137, hg19). Inspection of this region in the UCSC browser suggested that the core haplotype overlaps with a region of strong linkage disequilibrium in European and Asian populations (Han Chinese and Japanese from Tokyo), but not the Yoruba population (an ethnic group from West Africa, Figure 5B). Using a DNA recombination and haplotype-based mutation dating technique,\(^4\) we estimate that the most recent common ancestor (MRCA) of the CANVAS cohort lived approximately 25,880 (CI: 14,080–48,020) years ago (Figure 5C). This age estimate corresponds to the size of the haplotype and LD block and is roughly equivalent to the origin of modern Europeans as represented by the HAPMAP CEU cohort. Further investigation of the haplotypes allowed us to infer a simple phylogeny based on identified clusters of shared haplotypes extending beyond the core haplotype, suggesting that some individuals have common ancestors more recent than that of the MRCA for the whole group. This approach identified four subgroups. Group A had a MRCA dating back 5,600 (CI: 2,120–15,520) years and group B (further divided into groups B1 and B2) have a MRCA dating back 4,180 years (CI: 2,240–7,940). Furthermore, one individual shared part of their haplotype with both groups A and B, suggesting that group B is a distant branch of the MRCA of group A. Another subgroup, C, has a MRCA that lived 1,860 (CI: 560–7,020) years ago. The final group, labeled

Figure 4. Genetic Validation of the (AAGGG)\(^{\text{exp}}\) RE

(A) PCR analysis of the RFC1 STR failed to produce the control ~253 bp reference product in 18 of 22 CANVAS-affected families. (B and C) Representative images of the repeat-primed PCR for the (AAGGG)\(^{\text{exp}}\) RE demonstrating a saw-toothed product with 5 base pair repeat unit size, amplified from gDNA of individuals from CANVAS1 (B) and CANVAS9 (C). (D and E) No product was observed for the unaffected control (D) and no gDNA template negative control (E).
Next, we compared the haplotype of the nine control samples (three in-house control subjects and six from the Coriell collection) that carry the (AAGGG)exp RE to the core haplotype defined in the individuals with CANVAS. All control subjects shared at least part of the core haplotype, again suggesting that the (AAGGG)exp RE arose once in history. Finally, we determined that 9 of the 11 individuals from GTEx heterozygous for the (AAGGG)exp RE also shared the same core haplotype identified in individuals with CANVAS. The haplotype-specific SNP rs2066782 (exon 18, chr4:39303925, A>G) enabled us to analyze the expression of the (AAGGG)exp RFC1 allele in the cerebellum RNA-seq data and confirm that the STR did not inhibit the expression of RFC1 compared to the reference (AAAAG)11 allele. The remaining two carriers do not appear to share the core haplotype. As they do not have any additional sharing beyond the core haplotype.

Figure 5. The Majority of Individuals with CANVAS Encode an Ancestral Haplotype
(A) Analysis of WES data identified an ancestral haplotype surrounding RFC1 in all affected individuals confirmed to carry the (AAGGG)exp RE.
(B) The core haplotype (blue highlight) was intersected with the linkage disequilibrium (LD) track in the UCSC browser (converted to hg18 coordinates). The three LD tracks represent the Yoruba population (top track), Europeans (middle), and Han Chinese and Japanese from Tokyo (bottom). Red areas indicate strong linkage disequilibrium. The core CANVAS haplotype spans a large LD block in Europeans, which is broken up into two LD blocks in Japanese and Chinese, suggesting an ancient origin for the CANVAS repeat expansion allele.
(C) Haplotype sharing between individuals with CANVAS was used to determine the age of the most recent common ancestor (MRCA) of the cohort.
not have heterozygous SNPs in their exons, allele-specific expression could not be determined.

**Discussion**

Since the first description of the syndrome of cerebellar ataxia with bilateral vestibulopathy in 2004 and proposal of CANVAS as a distinct clinical entity in 2011, there has been little progress made in delineating the etiology of the disorder. While most affected individuals are described as idiopathic, reports of multiple affected sib pairs and a disorder. While most affected individuals are described as being likely. The genetic basis of CANVAS has now been identified and validated in two independent studies, one recently published by Cortese et al. and this study. Both studies utilized a similar study design, with linkage analysis to reduce the genomic search space to a modest interval (<2 Mb), but no plausible causal variant(s) could be identified in WES data. WGS was then performed on multiple individuals and Cortese et al. successfully identified the RE by visual inspection of the aligned read pairs inside the linkage region using the Integrative Genomics Viewer. In contrast, we utilized a bioinformatics approach and performed genome-wide analysis of WGS data to identify potential RE and then prioritized the RE located within the linkage interval. While both approaches were successful, the bioinformatics approach to RE detection, as described in this study, is likely more sensitive and practical and can be applied even in the absence of a small, or indeed any, linkage region. Furthermore, using a bioinformatics approach allows simultaneous testing of other potentially causal RE due to differential diagnoses. For example, we quickly re-diagnosed an affected individual with a pathogenic SCA3 RE.

Previously, the only variant associated with CANVAS was a heterozygous missense variant in the gene encoding E74 Like ETS Transcription Factor 2 (ELF2), which segregated with the disorder in three individuals in a single family. It is now apparent that the majority of individuals with CANVAS result from the homozygous inheritance of an expanded intronic pentamer in RFC1. We found the (AAGGG)exp in 30 of 31 individuals with a RE at this locus. In only a single individual did we observe a different, presumably pathogenic motif; CANVAS8 had one allele with the (AAGGG)exp whereas the second allele appeared to consist of an (AAAAG)11 allele. Interestingly, Cortese et al. were unable to determine a mechanism of action. The (AAGGG)exp RE did not appear to alter expression levels of RFC1 or surrounding genes as determined by bulk RNA-seq and qRT-PCR. Similarly, RFC1 expression and protein levels appeared unchanged in peripheral or brain tissue derived from individuals with CANVAS, and no AAGGG RNA foci deposits were observed. While RFC1 has not been previously associated with any disorder, it appears extremely intolerant to LoF (pLI = 0.97; observed/expected = 0.18, CI 0.12–0.31). In addition, siblings in the families studied carried the pathogenic RE in a heterozygous state but did not manifest any signs of the disorder. This observation is analogous to Friedreich ataxia, a recessive genetic ataxia caused by loss of function (LoF) of FRDA due to a pathogenic intronic RE.

RFC1 encodes a subunit of replication factor C, a five-subunit protein complex required for DNA replication and repair. Analysis of the Genotype-Tissue Expression (GTEx) database demonstrated significant expression of RFC1 in brain tissue, particularly the cerebellum. Replication factor C catalyzes opening the protein ring of proliferating cell nuclear antigen (PCNA), allowing it to encircle the (AAGGG)exp allele did not inhibit expression of RFC1 compared to the reference (AAAAG)11 allele. Interestingly, Cortese et al. were unable to determine a mechanism of action. The (AAGGG)exp RE did not appear to alter expression levels of RFC1 or surrounding genes as determined by bulk RNA-seq and qRT-PCR. Similarly, RFC1 expression and protein levels appeared unchanged in peripheral or brain tissue derived from individuals with CANVAS, and no AAGGG RNA foci deposits were observed. While RFC1 has not been previously associated with any disorder, it appears extremely intolerant to LoF (pLI = 0.97; observed/expected = 0.18, CI 0.12–0.31). In addition, siblings in the families studied carried the pathogenic RE in a heterozygous state but did not manifest any signs of the disorder. This observation is analogous to Friedreich ataxia, a recessive genetic ataxia caused by loss of function (LoF) of FRDA due to a pathogenic intronic RE.
the DNA and function as a scaffold to recruit proteins involved in DNA replication, repair, and remodeling. Mutations in multiple DNA replication and repair genes such as TCD1, PKNP, XRCC1, and APTX result in ataxia, highlighting the central role of this pathway in these overlapping disorders. One of the best-known examples is the severe and early-onset autosomal-recessive disorder ataxia telangectasia, which is caused by mutations in the gene encoding ATM serine/threonine kinase (ATM), which is important for the repair of DNA double-strand breaks.

The minimum pathogenic length and fine structure of the RFC1 RE is currently unclear. While Cortese et al. reported a pathogenic range of ~400–2,000, the individual repeat composition and a more precise repeat length was not determined. The short-read NGS technologies utilized in this study were unable to extend more than ~100 bp into the repeat sequence and efforts to amplify across the region using long-range PCR were unsuccessful. While the repeat-primed PCR assay indicates the presence of the (AAGGG)exp motif, it does not extend beyond ~250 bp (50 repeat units). The application of long-read sequencing technologies currently being developed for RE disorders will be required to accurately elucidate both the length of the pathogenic allele and the repeat composition. Both of these parameters provide important clinical information regarding onset, progression, and pathogenicity in other genetic ataxias such as SCA1 and Friedreich ataxia. Additional studies will also be required to elucidate the nature of the RFC1 STR in control individuals. Cortese et al. demonstrated considerable variability is present in the size and composition of the STR, but details regarding the size and composition of both normal and pathogenic alleles are yet to be fully determined. We show that the (AAGGG)exp RE occurs within the 3’ end of the Alu element, AluSx3. Alu elements typically have A-rich tails and in the reference sequence the RFC1 Alu has an A-rich tail containing an (AAAAG)11 STR. There is some evidence that motifs that follow the pattern A_mG_n especially (AAGG)_n and (AAAGG)_n display strong base-stacking interactions and are more likely to expand through replication slippage. This suggests an inherent mitotic instability of A- and G-rich motifs, consistent with what we observe in CANVAS. Notably, a number of pathogenic RE located with Alu have previously been described, including SCA10, SCA31, SCA37, and Friedreich ataxia.

Genomic Re-diagnosis in CANVAS

Of 22 families enrolled in this study with a clinical diagnosis of CANVAS, 4 did not harbor the RE or any other potentially pathogenic variants in the RFC1 locus. CANVAS13 was re-diagnosed with SCA3 after the WGS data was analyzed using our computational pipeline for detecting known pathogenic REs. In addition to cerebellar ataxia, individuals with SCA3 not uncommonly manifests a somatosensory impairment and vestibular involvement may be variably present, resulting in a phenotype indistinguishable from CANVAS. This molecular re-diagnosis highlights the power of modern STR detection techniques to diagnose RE ataxias. In addition, NGS data provide the opportunity to simultaneously identify non-RE-mediated causes of ataxia. In CANVAS17 we identified biallelic variants in SACS as the likely cause of disease. While individuals with spastic ataxia of the Charlevoix-Saguenay type may present with the combination of cerebellar ataxia and a peripheral neuropathy, as seen in CANVAS, to our knowledge vestibular involvement has not previously been described, and so this potentially constitutes a previously undescribed manifestation of the disease. In addition, a very plausible heterozygous variant was identified in EAT2 in CANVAS19. While classified as a VUS using ACMG guidelines, the variant is observed only once in gnomAD and was predicted pathogenic by multiple in silico algorithms. Very recently, heterozygous point mutations affecting the last cadherin domain (p.Lys3586Asn) or the linker region (p.Arg3649Gln) of EAT2 have been associated with SCA45, adding weight to classifying the variant (p.Val1457Ala) in the 13th cadherin domain, as likely pathogenic. While the published clinical phenotype and mutation spectrum in SCA45 is limited, in common with CANVAS, it is a late-onset and slowly progressive cerebellar ataxia.

Strengths and Limitations of Current STR Detection Tools

In this study, we implemented multiple computational tools to identify and validate the presence of an (AAGGG)exp RE in the majority of individuals with a clinical diagnosis of CANVAS. In particular, the use of EHdn, with its non-reference based RE discovery framework, was crucial in identifying a putative candidate, with the reference-based STR detection tools facilitating the follow-up analysis. Although all tools gave highly variable estimated repeat sizes, which are likely to be significantly less than the actual repeat size, they provided consistent evidence that the (AAGGG) motif was expanded. This level of evidence is helpful before embarking on the potentially complex process of molecular validation. In our analysis, only a single tool (GangSTR) failed to detect the alternate (AAGGG)exp RE. It is not clear why this was the case, although it could be related to the more complicated [(AAAGG)_n(AAAGG)exp(AAAAG)g]_n repeat structure. Notably, EHdn was able to identify the (AAGGG)exp RE in CANVAS8-8, but genome-wide significance was not achieved and the motif was less highly ranked based on p value than the initial discovery of AAGGG (Table S3). This is likely due to the fact that power was reduced as there was only a single case subject and a small number of control subjects. EHdn appears most effective as a discovery tool when used with multiple case subjects or larger numbers of control subjects. The results of this study highlight the importance of utilizing multiple tools to provide
redundancy in the data analysis pipeline. We have now updated the exSTRA package (see Web Resources below) to include CANVAS and other recently described pathogenic RE, providing additional utility to the research community to rapidly identify these RE in their cohorts. An additional issue we encountered, which potentially limited all tools, was the poor sequencing quality in reads containing the (AAGGG) motif compared to other STRs.

In conclusion, in this study we show that a recessively inherited, ancient RE located in intron 2 of RFC1 is the predominant cause of CANVAS. Recently developed RE discovery tools facilitated the identification and verification of this previously undescribed RE, in addition to identifying other genetic causes of disease in the cohort. Despite the RE being located in an intron, we demonstrate that previously generated WES data with low-coverage genome-wide off target reads were helpful in providing increased statistical confidence in RE identification. Therefore, reanalysis of previously generated WES datasets potentially offers a cost-effective approach to facilitate identification of intronic RE, not currently described in the genome reference sequence, in discovery projects. Finally, we anticipate that implementation of these tools into routine diagnostic pipelines has the potential to significantly increase the current diagnostic rates of <40%, recorded for clinical exome and targeted panel analyses of individuals with ataxia.61,62

Data and Code Availability

The ClinVar details for the RFC1 variants reported in this paper are SCV000898492.1 and SCV000898493.1.

Supplemental Data

Supplemental Data can be found online at https://doi.org/10.1016/j.ajhg.2019.05.016.

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Declaration of Interests

The authors declare no conflicts of interest.

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Web Resources

Cavalier, https://github.com/bahlolab/cavalier
European Genome-phenome Archive (EGA), https://www.ebi.ac.uk/ega
exSTRA, https://github.com/bahlolab/exSTRA
gnomAD Browser, https://gnomad.broadinstitute.org/
GTEx Portal, https://gtexportal.org/home/
OMIM, https://www.omim.org/
UCSC Genome Browser, https://genome.ucsc.edu
Varsome, https://varsome.com

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