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# Genetic, Phenotypic, and Interferon Biomarker Status in ADAR1-**Related Neurological Disease**

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# Abstract

We investigated the genetic, phenotypic, and interferon status of 46 patients from 37 families with neurological disease due to mutations in ADAR1. The clinicoradiological phenotype encompassed a spectrum of Aicardi-Goutières syndrome, isolated bilateral striatal necrosis, spastic paraparesis with normal neuroimaging, a progressive spastic dystonic motor disorder, and adult-onset psychological difficulties with intracranial calcification. Homozygous missense mutations were recorded in five families. We observed a p.Pro193Ala variant in the heterozygous state in 22 of 23 families with compound heterozygous mutations. We also ascertained 11 cases from nine families with a p.Gly1007Arg dominant-negative mutation, which occurred de novo in four patients, and was inherited in three families in association with marked phenotypic variability. In 50 of 52 samples from 34 patients, we identified a marked upregulation of type I interferon-stimulated gene transcripts in peripheral blood, with a median interferon score of 16.99 (interquartile range [IQR]: 10.64–25.71) compared with controls (median: 0.93, IQR: 0.57–1.30). Thus, mutations in ADAR1 are associated with a variety of clinically distinct neurological phenotypes presenting from early infancy to adulthood, inherited either as an autosomal recessive or dominant trait. Testing for an interferon signature in blood represents a useful biomarker in this context.

# Keywords

Aicardi-Goutières syndrome; bilateral striatal necrosis; spastic paraparesis; dystonia; idiopathic basal ganglia calcification

## Introduction

Adenosine deaminases acting on RNA (ADARs) catalyze the hydrolytic deamination of adenosine to inosine in double-stranded RNA, and thereby potentially alter the information content and structure of cellular RNAs. ADAR1 is encoded by a single-copy gene that

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#### **Authors' Contributions**

J.H.L. and Y.J.C. collated and reviewed all clinical and radiological data. G.I.R. performed quantitative PCR analysis, with assistance from N.K., M.B., T.A.B., A.C.E.B., M.L.C., A. M.C., C.C.,R.C.D., F.R.D.,N.D., B.De A., V.De G., C.G.E.L. De G.,I. D., C De L., A.E., M.C.F., P.F., A.F., E.F., M.P.G., N.R.G., M.H., M.A. K., N.L., J.-P.S.-M.L., M.A.L., S.S.M., R.M., L.M.-S., G.M., M.M., V. N., S.O., J.D.O.-E., B.P.-D., F.P., K.M.R., M.R., F.R., P.R.-P., A.R., T.I. S., M.B.T., A.T., F.U., N.U., A.V., and A.W. provided clinical samples and critically reviewed clinical and immunological patient data. Y.J.C. conceived the study and wrote the initial draft with the assistance of G.I.R. All authors critically reviewed the article and agreed to its publication.

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maps to human chromosome 1q21 and is present in two main isoforms in mammalian cells. In mice, a loss of ADAR1 activity leads to a dramatic upregulation of interferon-stimulated gene (ISG) expression, which is dependent on the editing activity of ADAR1 and specific to the interferon-inducible full-length p150 isoform of the protein.<sup>2–4</sup>

In 2012, we reported mutations in *ADAR1* to cause a phenotype consistent with the infantile encephalopathy Aicardi–Goutières syndrome (AGS), and demonstrated that, similar to the *ADAR1* null mouse, the mutant genotype was associated with an upregulation of type I interferon signaling.<sup>5</sup> Further to this, in 2014, we described both bilateral striatal necrosis (BSN), sometimes occurring after a trivial childhood infection, and otherwise nonsyndromic, slowly progressive spastic paraparesis associated with normal intellect occur due to ADAR1 dysfunction, again in association with the enhanced expression of type I interferon-induced gene transcripts.<sup>6–8</sup> These data indicate that neurological disease can occur through inappropriate induction of the innate immune system by self-derived nucleic acids.

Here, we present an update of our experience of screening for *ADAR1* mutations, describing the clinical, radiological, molecular, and interferon biomarker characteristics of a cohort of 46 patients from 37 families with neurological dysfunction due to mutations in *ADAR1*.

## **Materials and Methods**

#### **Patients and Methods**

We ascertained clinical and molecular data through direct contact and/or via collaborating physicians. The study was approved by the Leeds (East) Research Ethics Committee (reference number 10/H1307/132), and the Comité de Protection des Personnes (ID-RCB/EUDRACT: 2014-A01017-40).

A diagnosis of AGS was suggested by characteristic clinical and neuroimaging features including cerebral atrophy, white matter disease, and intracranial calcification. BSN was diagnosed in the context of an acute or subacute onset of a dystonic/rigid motor disorder associated with magnetic resonance imaging features of bilateral striatal signal change with or without swelling. Spastic paraparesis/tetraparesis and spastic dystonia were diagnosed according to clinical signs, in the presence of either normal neuroimaging or mild nonspecific changes sometimes including calcification of the basal ganglia. Assessment of the motor and communication status of patients over the age of 1 year was made using the Gross Motor Function Classification System (GMFCS), 10 the Manual Ability Classification System (MACS), 11 and the Communication Function Classification System (CFCS). 12

#### **Mutational Analysis**

Primers were designed to amplify the coding exons of *ADAR1* (Supplementary Table S1, online-only). Purified polymerase chain reaction (PCR) amplification products were sequenced using BigDye terminator chemistry and an ABI 3130 DNA sequencer. Mutation description is based on the reference cDNA sequence NM\_001111.4, with nucleotide numbering beginning from the first A in the initiating ATG codon. Variants were assessed using the in silico programs SIFT (http://sift.jcvi.org) and Polyphen2 (http://

genetics.bwh.harvard.edu/pph2/), and population allele frequencies obtained from the ExAC (http://exac.broadinstitute.org) and gnomAD (http://gnomad.broadinstitute.org) databases.

#### Interferon Score

## Results

#### **Molecular Data**

We collected data on 46 patients from 37 families of pan-ethnic origin with either biallelic mutations in *ADAR1* (28 families) or the single known dominant-negative mutation p. Gly1007Arg (nine families) (Table 1; Fig. 1). In four families, the p.Gly1007A mutation was considered to have occurred de novo, while in three families, inheritance was confirmed or inferred (two paternal half-siblings born to an unaffected father unavailable for testing), with somatic mosaicism recorded in one case. In two families, inheritance could not be determined because DNA from both parents was not available. We observed three distinct homozygous mutations in five families (two families each sharing the same mutation), in four of which the parents were knowingly related. All of these mutations were missense. Of 23 families with compound heterozygous mutations, 22 carried the p. Pro193Ala mutation on one allele. In 13 of 22 families segregating this p.Pro193Ala substitution, the second molecular lesion was a null or splicing variant.

#### **Clinical Radiological Phenotype**

Clinical radiological characteristics of all patients are summarized in Table 2, and characteristic radiological appearances are summarized in Fig. 2. Median age of disease onset was 14 months (range: birth–30 years). We observed 21 and 25 affected females and males, respectively. Although spasticity and dystonia were common features present in the majority of patients, clinically and radiologically distinct phenotypes could be defined, including classical AGS (15 patients), BSN (16 patients), apparently isolated spastic paraparesis (1 patient)/tetraparesis (2 patients), and a progressive spastic dystonic motor disorder (7 patients). In two of these latter cases, the initial presentation was of isolated

lower limb spasticity, with a dystonic component and involvement of the upper limbs only becoming evident several years later. Four patients demonstrated radiological features of both AGS and BSN. The mother of a child with an AGS presentation was diagnosed at the age of 30 years with subtle psychological features and marked intracranial calcification. We identified three patients with significant neurological disease (a spastic/dystonic phenotype) in the absence of changes on brain imaging at presentation.

A total of 25 patients were considered to have demonstrated normal development prior to disease onset, in 18 of whom there was a history of either vaccination (4 patients) or a notable infectious episode (14 patients) in the period shortly preceding the development of clinical signs (Fig. 3A). Several patients experienced a rapid onset of dystonia/spasticity and loss of skills, with two patients being admitted to intensive care due to severe dystonic crisis. Others exhibited a more slowly progressive onset over weeks or months. Definite clinical progression beyond the initial presentation was recorded in 16 cases. Nine patients are deceased, between the ages of 10 months and 19 years, six of whom had early-onset disease consistent with AGS.

An assessment of gross motor function, manual ability, and communication status at last contact was made in 45 patients, of whom 27 were recorded to have none of any purposeful gross motor, hand and communication function (score of 5 on all three scales) (Fig. 3B). Five patients were able to walk with no or some support (GMFCS I–III). Eleven patients were capable of effective sender and receiver communication (CFCS I–III). Although formal testing was not undertaken, seven patients were considered to have normal intellectual function.

Five patients were reported to demonstrate hypo/hyper-pigmentation consistent with dyschromatosis symmetrica hereditaria (DSH) 1, and two patients were described with chilblain-like vasculitic lesions. Four patients were documented with autoimmune hemolytic anemia. Glaucoma was not recorded in any patient.

# **Interferon Status**

We derived 52 interferon scores from 34 patients, 50 of which were abnormal, with a median interferon score across the group of 16.99 (interquartile range [IQR]: 10.64–25.71) compared with controls (median: 0.93, IQR: 0.57–1.30) (Fig. 4). Positive scores were observed up to 25 years after disease onset. We also tested 20 interferon scores from 16 parental carriers of a recessive mutation in *ADAR1*. Two samples from seven parents heterozygous for the recurrent p.Pro193Ala mutation demonstrated a positive interferon score, versus six samples from nine parents carrying a different mutation (Supplementary Fig. S1, online-only).

# **Discussion**

In 2012, *ADAR1* mutations were described in the context of the early-onset encephalopathy AGS, associated with the presence of intracranial calcification, white matter disease, and severe developmental delay.<sup>5</sup> Subsequently, in 2014, mutations in *ADAR1* were also shown to underlie cases of apparently nonsyndromic BSN, and of isolated spastic para-paresis with

normal neuroimaging.<sup>6,7</sup> Here, we confirm these associations, thus emphasizing the need to consider ADAR1-related disease in several distinct clinical scenarios triggering different investigative algorithms. Furthermore, we now describe a patient with a dominant-negative mutation in *ADAR1* demonstrating an adult-onset phenotype evocative of "idiopathic" basal ganglia calcification characterized by intracranial calcification and subtle psychological disturbance. Our clinical and radiological findings highlight the propensity of ADAR1-related disease to incur basal ganglia dysfunction, and the value of basal ganglia calcification, frequently only appreciated on computed tomography, as a diagnostic indicator. In general, mutations in *ADAR1* should be considered in the context of a motor disorder characterized by spasticity and dystonia. The onset of disease can occur after a period of normal development, sometimes associated with a rapid loss of skills, or a much slower progression over many years. Assessments using the GMFCS, MACS, and CFCS rating scales indicate that disease outcome in the cases that we have ascertained is frequently severe. It is of note that we observed cases with completely preserved intellect +/- normal neuroimaging in the face of significant motor disability.

Our own research focus is biased toward the ascertainment of pediatric disease. However, Tojo et al described a female patient with the dominant-negative p.Gly1007Arg mutation, presenting at the age of 17 years with gait disturbance and dystonic posturing of the legs, who experienced intellectual deterioration from 21 years of age, and became wheelchair bound a year later. Together with our observation of an adult female whose clinical phenotype only became evident at the age of 30 years, it is clear that later onset disease can occur due to ADAR1 deficiency. This latter case also illustrates the significant intrafamilial variability which can be seen in association with ADAR1 dysfunction, the mother presenting in adulthood with subtle psychological disturbance, while her son experienced a devastating early-onset encephalopathy.

ADAR1-related neurological disease can be inherited as either an autosomal recessive or autosomal dominant trait. We observed homozygosity for a missense mutation in five of 28 families segregating recessive disease. As previously suggested, the absence of patients with homozygous null mutations indicates that, as for the ADAR1 null mouse, complete loss of ADAR1 protein activity is likely embryonic lethal.<sup>5</sup> Our molecular data reveal a remarkably high frequency of the p.Pro193Ala substitution, seen in 22 of 23 families with compound heterozygous molecular lesions in ADAR1. This mutation, which is recorded on 602 of 282,636 alleles in the gnomAD database, was not observed in the homozygous state in our cohort. That this variant was seen in combination with a null mutation in 13 families suggests that homozygosity for the p.Pro193Ala allele leads to a milder, later onset, or distinct phenotype not ascertained here, or may not be associated with disease. Perhaps of note, the gnomAD database includes one individual homozygous for this mutation. Finally, our molecular data highlight the dominant-negative p.Gly1007Arg mutation, which can occur de novo, or be inherited with variable expression and/or nonpenetrance at least into mid-adult life. The proximity of Gly1007 to the backbone of its RNA ligand, and the possibility for an arginine residue to make polyvalent interactions there suggests a mechanism whereby Arg1007 might bind more tightly to RNA and thus act as a competitive inhibitor of wild-type protein, while being itself catalytically inactive. <sup>14</sup> In keeping with this model, we previously demonstrated that a plasmid expressing Gly1007Arg showed stronger

inhibition of wild-type ADAR1 than equivalent amounts of a plasmid expressing catalytic inactive ADAR1.<sup>5</sup>

More than 130 different *ADAR1* mutations have been documented in patients with DSH, an autosomal-dominant disorder characterized by the childhood onset of hypopigmented and hyperpigmented macules on the face and dorsal aspects of the extremities. <sup>15</sup> DSH has only very rarely been reported outside of Japan and China, and even within identified families, a marked variability in expression is well recognized. In our series, five patients were noted to demonstrate pigmentary lesions consistent with DSH. The frequent observation of stop and frameshift variants in DSH indicates haploinsufficiency as the likely molecular pathology, consistent with the recent confirmation of our previous suggestion that two individuals with DSH would be at one in four risk of a pregnancy with ADAR1-related neurological disease.

Loss-of-function mutations in *ADAR1* have been classified within the so-called type I interferonopathy grouping, a novel set of inborn errors of immunity where it is proposed that an upregulation of type I interferon signaling is central to disease pathogenesis. <sup>17,18</sup> The AGS phenotype can arise due to mutations in any one of seven genotypes within this grouping (*AGS1–7: TREX1, RNASEH2A, RNASEH2B, RNASEH2C, SAMHD1, ADAR1*, and *IFIH1*), and apparently isolated spastic paraparesis has been reported in patients mutated in three of these genes (*RNASEH2B, ADAR1*, and *IFIH1*). In contrast, in an overview of 374 patients from 299 families with mutations in *AGS1–7*, BSN, the most frequently ascertained phenotype in the current series, was only recorded in the context of ADAR1-related disease, suggesting discrete factors relevant to gene/protein expression and disease mechanism consequent upon ADAR1 dysfunction. <sup>19</sup> Also possibly reflective of this apparent specificity, in comparison to other genotypes, is the frequency of clinical progression, and the low risk of developing glaucoma and chilblain-like lesions (since we recorded no examples of the former and only two cases of the latter in our cohort).

The consistent finding of a positive interferon signature in peripheral blood in the series of patients reported here indicates the potential utility of this biomarker as a screening test for ADAR1-related disease, for the interpretation of ADAR1 genetic sequence variants of uncertain significance, and in the possible monitoring of treatment efficacy as anti-interferon therapies is developed.<sup>20,21</sup> We emphasize that the interferon signature remains elevated many years after disease onset, providing evidence of ongoing pathology. ADAR1 is expressed throughout the brain including the basal ganglia (http://www.brain-map.org), and it has been shown that a loss of ADAR1 renders cells more susceptible to apoptosis following stress, including infection.<sup>22</sup> We cannot rule out the possibility that the occurrence of fevers prior to frank neurological regression represents a prodrome in some cases. However, a history of vaccination or an apparently discrete infectious episode in several patients considered to be completely developmentally normal prior to disease onset, of whom 12 demonstrated BSN on neuroimaging, raises the possibility that the acute degeneration of striatal tissue seen in many patients with ADAR1 mutations might relate to a rapid induction of apoptosis triggered by viral infection/metabolic stress. Beyond this possibility, there is strong evidence that interferon is a neurotoxin, <sup>23–27</sup> and we consider it likely that inappropriate and chronic exposure to type I interferons may be directly relevant

to the ADAR1-related neurological phenotypes described here, perhaps induced by dsRNA species which are normally edited by ADAR1, thereby rendering them as immunology inert/marking them as self. 1,3,4,28 These observations highlight the potential utility of treatments for ADAR1-related disease, which recent data suggest might be usefully targeted at antagonism of type I interferon signaling. 29

# **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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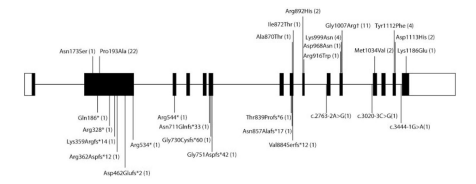
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**Fig. 1.**Schematic of *ADAR1* gene showing mutations (according to protein nomenclature) ascertained in the present study. Missense and nonsense mutations are annotated above and below, respectively. Numbers in brackets indicate the number of families in which each mutation was observed. †Indicates mutation acting as a dominant negative.

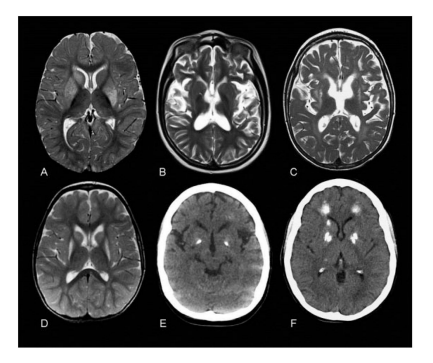
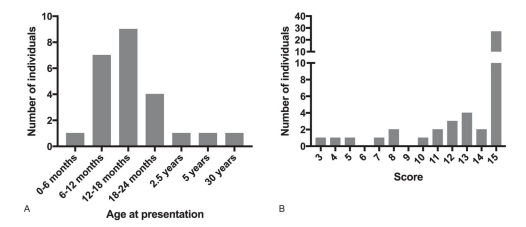
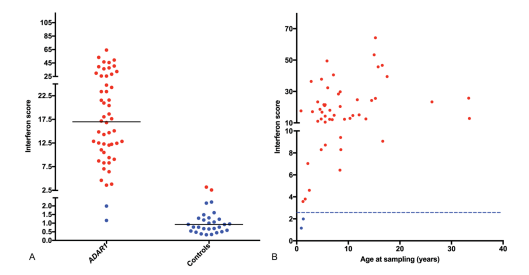


Fig. 2.
Characteristic neuroradiological features of ADAR1-related disease. Images (**A**) and (**D**) are axial T2 images of AGS251, presenting at 9 months of age with bilateral striatal necrosis following varicella zoster infection, showing characteristic high signal and swelling of head of caudate and putamen (**A**). (**D**) Follow-up at 35 months shows persisting signal change and shrinkage of caudate and putamen. Images (**B**) and (**E**) are from AGS150, a 10-year-old child presenting with an Aicardi–Goutières syndrome phenotype. (**B**) T2 axial MR shows cerebral atrophy with mildly increased signal in white matter. (**E**) CT shows dense bilateral globus pallidus calcification. Image (**C**) is of a patient presenting with an Aicardi–Goutières syndrome phenotype (AGS810\_P1). (**C**) T2 axial MR at 5 years shows marked cerebral atrophy, white matter high signal, and signal change and shrinkage of the putamen. (**F**) CT scan of his mother (AGS810\_P2) aged 34 years shows dense calcification of globus pallidus, head of caudate, and deep frontal white matter. Her MR (not shown) was normal. CT, computed tomography; MR, magnetic resonance.



**Fig. 3.**Age at presentation and associated disability. (**A**) Age at presentation in patients developing disease after a period of clearly normal development. (**B**) Assessment of gross motor function, manual ability, and communication status in living patients with mutations in *ADAR1* over 1 year of age.



**Fig. 4.** Interferon score data in *ADAR1*-mutated patients and controls. Summary of interferon score data (**A**) in *ADAR1*-mutated patients and controls and (**B**) in *ADAR1*-mutated patients by age. Circles indicate results above +2 SD of the mean of 29 controls (= 2.466, considered "positive"). Solid horizontal lines indicate median value of *ADAR1*-mutated and control groups. Dotted line indicates positive/negative boundary (2.466) of interferon score.

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Table 1

Family structure, ethnicity, and molecular data of ascertained ADARI mutation-positive cases

gnomAD frequency	602/282636 1 hom	1/252010	602/282636 1 hom	Novel	Novel	Novel	Novel	Novel	602/282636 1 hom	1/252270	602/282636 1 hom	Novel	602/282636 1 hom	1/252010	Novel	Novel	602/282636 1 hom	1/30224	144/282658 1 hom	Novel	602/282636 1 hom	Novel	602/282636 1 hom	2/252366	602/282636 1 hom	Novel
ExAc frequency	260/ 121402	Novel	260/ 121402	Novel	Novel	Novel	Novel	Novel	260/ 121402	1/121342	260/ 121402	Novel	260/ 121402	Novel	Novel	Novel	260/ 121402	Novel	34/121366	Novel	260/ 121402	Novel	260/ 121402	Novel	260/ 121402	Mossol
CADD Phred	23.9	35	23.9	34	33	34	33	34	23.9	26.9	23.9	Frameshift	23.9	35	34	34	23.9	Frameshift	24.3	Frameshift	23.9	Frameshift	23.9	Stop	23.9	
Polyphen2	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Frameshift	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Probably damaging 1.000	Frameshift	Probably damaging 0.999	Frameshift	Probably damaging 1.000	Frameshift	Probably damaging 1.000	Stop	Probably damaging 1.000						
SIFT	Deleterious 0	Deleterious 0.01	Deleterious 0	Deleterious 0	Deleterious 0.02	Deleterious 0	Tolerated 0.17	Deleterious 0.03	Deleterious 0	Deleterious 0.01	Deleterious 0	Frameshift	Deleterious 0	Deleterious 0.01	Deleterious 0	Deleterious 0	Deleterious 0	Frameshift	N/A	Frameshift	Deleterious 0	Frameshift	Deleterious 0	Stop	Deleterious 0	
Inheritance	Maternally inherited	Paternally inherited	Paternally inherited	Maternally inherited	Both parents het	De novo (paternity confirmed)	Not tested	Both parents het	Maternally inherited	Paternally inherited	Maternally inherited	Paternally inherited	Maternally inherited	Paternally inherited	De novo (paternity confirmed)	Presumed inherited from asymptomatic Father	Paternally inherited	Maternally inherited	Paternally inherited	Maternally inherited	Not known	Not known	Paternally inherited	Maternally inherited	Not known	
Allelic status	Het	Het	Het	Het	Hom	Het	Hom	Hom	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	Het	
Protein	p.Pro193Ala	p.Arg892His	p.Pro193Ala	p.Ala870Thr	p.Asp1113His	p.Gly1007Arg	p.Tyr1112Phe	p.Lys999Asn	p.Pro193Ala	p.Ile872Thr	p.Pro193Ala	p.Lys359Argfs*14	p.Pro193Ala	p.Arg892His	p.Gly1007Arg	p.Gly1007Arg	p.Pro193Ala	p.Asn857Alafs*17	p.Asn173Ser	p.Thr839Profs*6	p.Pro193Ala	p.Val884Serfs*12	p.Pro193Ala	p.Arg544*	p.Pro193Ala	I 1 19001
cDNA	c.577C>G	c.2675G>A	c.577C>G	c.2608G>A	c.3337G>C	c.3019G>A	c.3335A>T	c.2997G>T	c.577C>G	c.2615T>C	c.577C>G	c.1076_1080del	c.577C>G	c.2675G>A	c.3019G>A	c.3019G>A	c.577C>G	c.2565_2568del	c.518A>G	c.2515del	c.577C>G	c.2647_2648dup	c.577C>G	c.1630C>T	c.577C>G	
Ethnicity	White European		Italian		Pakistani	Brazilian	Pakistani	Indian	White European		Italian/ Hispanic		Spanish		White European	White European	White European		Greek/ Lebanese		White European		White European		White European	
Consanguinity	No		No		Yes	No	Yes	No	No		No		No		No	No	No		No		No		oN		No	
Individuals tested	3A, M, F		1A, M, F		2A, M, F	1A, M, F	1A	1A, M, F	1A, M, F		1A, M, F		2A, M, F		1A, M, F	2A, M	1A, M, F		1A, M, F		1A		2A, M, F		1A	
AGS number	AGS081		AGS093		AGS107	AGS150	AGS219	AGS228	AGS251		AGS327		AGS430		AGS474	AGS530	AGS550		AGS567		AGS582		AGS663		AGS679	

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AGS	Individuals tested	Consanguinity	Ethnicity	cDNA	Protein	Allelic status	Inheritance	SIFT	Polyphen2	CADD Phred	ExAc frequency	gnomAD frequency
AGS699	1A, M, F	No	White European	c.3019G>A	p.Gly1007Arg	Het	De novo (genotyping not undertaken)	Deleterious 0	Probably damaging 1.000	34	Novel	Novel
AGS703	1A	No	Asian/ White European	c.577C>G	p.Pro193Ala	Het	Not known	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.3100A>G	p.Met1034Val	Het	Not known	Deleterious 0.03	Possibly damaging 0.760	25.8	Novel	Novel
AGS720	1A, M, F	No	White European	c.577C>G	p.Pro193Ala	Het	Maternally inherited	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.2250del	p.Gly751Aspfs*42	Het	De novo (genotyping not undertake n)	Frameshift	Frameshift	Frameshift	Novel	Novel
AGS759	1A, M, F	No	White European	c.577C>G	p.Pro193Ala	Het	Paternally inherited	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.2902G>A	p.Asp968Asn	Het	Maternally inherited	Tolerated 0.06	Probably damaging 1.000	34	Novel	Novel
AGS765	1A	No	White European	c.577C>G	p.Pro193Ala	Het	Not known	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.556C>T	p.Gln186*	Het	Not known	Stop	Stop	Stop	Novel	Novel
AGS788	1A, M, F	No	White European	c.577C>G	p.Pro193Ala	Het	Maternally inherited	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.1386_1390del	p.Asp462Glufs*2	Het	De novo (paternity confirmed)	Frameshift	Frameshift	Frameshift	Novel	Novel
AGS810	1A, MA, F	No	White European	c.3019G>A	p.Gly1007Arg	Het	Inherited from symptomatic mother	Deleterious 0	Probably damaging 1.000	34	Novel	Novel
AGS943	1A, M, F	No	North African	c.3019G>A	p.Gly1007Arg	Het	De novo (genotyping not undertaken)	Deleterious 0	Probably damaging 1.000	34	Novel	Novel
AGS1115	1A, M, F	Yes	Persian	c.2997G>T	p.Lys999Asn	Hom	Both parents het	Deleterious 0.03	Probably damaging 1.000	34	Novel	Novel
AGS1170	1A	No	Asian	c.577C>G	p.Pro193Ala	Het	Not known	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.3100A>G	p.Met1034Val	Het	Not known	Deleterious 0.03	Possibly damaging 0.760	25.8	Novel	Novel
AGS1315	2A, M, F (mosaic)	No	White European	c.3019G>A	p.Gly1007Arg	Het	Father mosaic	Deleterious 0	Probably damaging 1.000	34	Novel	Novel
AGS1456	1A	No	White European	c.577C>G	p.Pro193Ala	Het	Not known	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.3020–3C>G	Splicing	Het	Not known	Splicing	Splicing	Splicing	Novel	Novel
AGS1507	1A, M, F	No	Asian/ White European	c.577C>G	p.Pro193Ala	Het	Maternally inherited	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.2763–2A>G	Splicing	Het	Paternally inherited	Splicing	Splicing	Splicing	Novel	Novel
AGS1537	1A	No	White European	c.3019G>A	p.Gly1007Arg	Het	Not known	Deleterious 0	Probably damaging 1.000	34	Novel	Novel
AGS1542	2A, M, F	Yes	Asian	c.3335A>T	p.Tyr1112Phe	Hom	Both parents het	Tolerated 0.17	Probably damaging 1.000	33	Novel	Novel
AGS1824	1A	No	White European	c.577C>G	p.Pro193Ala	Het	Paternally inherited	Deleterious 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom
				c.1084_1085del	p.Arg362Aspfs*12	Het	Maternally inherited	Frameshift	Frameshift	Frameshift	Novel	Novel
AGS1980	1A	No	White European	c.577C>G	p.Pro193Ala	Het	Not known	Deleterions 0	Probably damaging 1.000	23.9	260/ 121402	602/282636 1 hom

Individuals tested	Consanguinity	Ethnicity	cDNA	Protein	Allelic status	Inheritance	SIFT	Polyphen2	CADD Phred	ExAc frequency	gnomAD frequency
			c.2130dupC	p.Asn711Glnfs*33	Het	Not known	Frameshift	Frameshift	Frameshift	Novel	Novel
	No	South American	c.577C>G	p.Pro193Ala	Het	Paternally inherited	Deleterious 0	Probably damaging 1.000 23.9	23.9	260/ 121402	260/ 121402 602/282636 1 hom
			c.2187_2198delinsGT p.Gly730Cysfs*60	p.Gly730Cysfs*60	Het	Maternally inherited	Frameshift	Frameshift	Frameshift	Novel	Novel
	No	White European	c.577C>G	p.Pro193Ala	Het	Not known	Deleterious 0	Probably damaging 1.000 23.9	23.9	260/ 121402	602/282636 1 hom
1			c.982C>T	p.Arg328*	Het	Not known	Stop	Stop	Stop	Novel	1/252210
1	No	White European	c.577C>G	p.Pro193Ala	Het	Paternally inherited	Deleterious 0	Probably damaging 1.000 23.9	23.9	260/ 121402	260/ 121402   602/282636 1 hom
			c.2746C>T	p.Arg916Trp	Het	Maternally inherited	Deleterious 0	Probably damaging 1.000 35	35	Novel	Novel
	No	Hispanic	c.3019G>A	p.Gly1007Arg	Het	M WT, F not tested	Deleterious 0	Probably damaging 1.000 34	34	Novel	Novel

Abbreviations: A, affected; F, father; Het, heterozygous; Hom, homozygous; M, mother; MA, mother affected; WT, wild type.

Note: Nucleotide numbering based on transcript ADARINM\_001111.4. ExAc browser Beta version accessed on October 28, 2016 (http://exac.broadinstitute.org), gnomAD browser \(\beta\) version accessed on October 28, 2016 (http://gnomad.broadinstitute.org).

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Table 2

Clinical and radiological data relating to ascertained ADARI mutation-positive cases

Summary	AGS	AGS	AGS	AGS/BSN	AGS	AGS	AGS	AGS	AGS	BSN	AGS/BSN	AGS	AGS	AGS	BSN	BSN
CFCS	Λ	Λ	^	>	^	>	>	>	< 1 year	IV	>	Λ	^	Λ	>	IV
MACS	Λ	Λ	>	^	Λ	^	>	^	< 1 year	Λ	>	Λ	Λ	>	^	IV
GMFCS	Λ	Λ	Λ	^	Λ	^	Λ	^	< 1 year	Λ	>	Λ	Λ	>	^	^
Interferon scores (age, deci- malized years)	24.267 (14.53); 53.356 (15.01); 45.676 (15.78)	NT	37.822 (4.82); 21.590 (5.28)	25.608 (15.26); 46.665 (16.59)	64.22 (15.26)	IN	14.69 (10.88)	NT	IN	28.367 (8.1); 12.301 (9.27)	23.382 (4.07); 9.402 (8.52)	8.296 (4.75); 21.538 (5.53)	12.444 (4.75); 14.306 (5.53)	20.961 (5.42); 32.319 (5.88); 49.463 (6.02)	12.502 (13.41)	23.385 (26.21)
Neuroima- ging	Characteristic of AGS	Characteristic of AGS	Characteristic of AGS	Characteristic of AGS and BSN	Characteristic of AGS	Characteristic of AGS	Some white matter disease and calcification of GP	Characteristic of AGS	Characteristic of AGS	BSN	AGS with features of BSN	Characteristic of AGS	Characteristic of AGS	Characteristic of AGS	BSN	BSN
Status at last con- tact	SDT with severe ID	SDT with severe ID	SDT with severe ID	SDT with severe ID	SDT with severe ID; AIHA	SDT with severe ID; AIHA	SDT with some ID	SDT with severe ID; AIHA	SDT with severe ID	SDT with some ID; CB	SDT with severe ID; DSH	SDT with severe ID	SDT with severe ID	SDT with severe ID	SDT with some understanding	SDT with some understanding
Progres- sive course	Yes	Yes	Not obvious	Not obvious	Not obvious	Not obvious	No	Not obvious	Not obvious	Not obvious	Not obvious	Uncertain	Not obvious	Yes, with worsening respiratory function and overall neurological deterioration	Yes	Yes
Current age/ age at death (cause were known)	Died aged 17 y	Died aged 23 mo	9 y	20 y	Died aged 19 y	14 y	15 y	Died aged 6 y	Died aged 10 mo	12 y	8 y	Died aged 6 y	9 y	8 y	17 y	29 y
Features at presenta- tion	DD, dystonia, irritability	DD, dystonia, irritability, microcephaly	Raised CSF IFN at birth with transient thrombocytopenia and petechiae	DD, irritability, sleep and feeding disturbance	DD, dystonia, irritability, microcephaly	DD, dystonia, irritability, microcephaly	Loss of head control, sitting and speech	DD, poor head control	IUGR, thrombocytopenia, HSM	Loss of skills over a few weeks	DD, encephalopathy, irritability	DD, dystonia, irritability, microcephaly	DD, dystonia, irritability, microcephaly	Nystagmus, gross and fine motor delay	Subacute loss of skills becoming rigid over a few months	Subacute loss of skills becoming rigid over a few months
Age at in- itial ascertain- ment	5 mo	5 mo	Neonatal	1 mo	< 7 mo	Neonatal	18 mo	om 9 >	Prenatal	9 mo	8 mo	< 2 mo	< 2 mo	4 mo	5 y	1 y
Possible trigger	No	No	No	No	No	No	No	No	No	Varicella infection	Possible viral infection (otitis media)	No	No	Vaccination	No	No
Develop- mental status prior to onset	Delayed	Delayed	Diagnosed at birth	Delayed	Delayed	Delayed	Mild delay	Delayed	Delayed	Normal	Delayed	Delayed	Delayed	Normal	Normal	Normal
Sex	F	F	M	M	F	н	Н	M	н	F	M	F	н	×	ц	F
Individual	PI	P2	P3	PI	P1	P2	P1	PI	PI	PI	PI	P1	P2	PI	PI	P2
AGS number	AGS081		1	AGS093	AGS107		AGS150	AGS219	AGS228	AGS251	AGS327	AGS430		AGS474	AGS530	

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Summary	BSN	BSN	BSN	BSN	BSN	BSN	SP	SDT	BSN	SDT	SDT	BSN	AGS/BSN	ICC with psychiatric features	SP becoming SDT with preserved intellect	AGS	AGS/BSN
CFCS	>	^	>	Ш	V	Ħ	I	Ш	Ш	Ш	Ŋ	>	>	1	I	>	>
MACS	V	Λ	>	Ш	^	Ш	I	IV	>	П	>	>	^	I	Ш	Λ	>
GMFCS	>	Λ	^	IV	Λ	п	п	>	^	Ш	VI	>	>	I	I	^	>
Interferon scores (age, deci- malized years)	6.429 (8.39)	36.387 (2.81)	NT	IN	38.13 (17.53)	3.802 (1.66)	16.833 (4.91)	20.427 (8.44); 29.817 (8.44)	12.057 (6.90)	11.048 (4.09); 18.633 (4.53)	NT	1.99 (1.29); 4.59 (2.46)	40.571 (7.13); 14.851 (7.27)	25.743 (33.34); 12.836 (33.48)	24.753 (11.75); 15.074 (12.11)	NT	17.627 (0.84); 1.158 (0.90); 3.578 (1.23)
Neuroima- ging	BSN	BSN	BSN	BSN	BSN	BSN	Normal	Initially structurally normal MRI; BG calcification noted 2 years later	BSN	Calcification of caudate and putamen	BG signal changes and atrophy with subcortical hypomyelination	BSN	Characteristic of AGS	Normal except for BG, WM, and Cb calcification	Some cortical atrophy with BG and WM calcification	Characteristic of AGS	Initial bilateral high signal and swelling of BG progressing to extensive WM and
Status at last con- tact	SDT with some understanding	SDT with moderate ID; DSH	SDT with moderate ID	SDT moderate ID	SDT	Dystonic gait and clumsy hand finger movements; intellectually normal	Major LL spasticity; intellectually normal	SDT with severe ID	SDT; intellectually normal; DSH	SDT; intellectually normal	SDT with some ID	SDT with severe ID	SDT with severe ID	Normal clinical examination; subtle psychological difficulties	SDT; intellectually normal	SDT with severe ID	SDT with severe ID
Progres- sive course	Yes	Yes	Yes	No	Not obvious	Yes, then some recovery	Yes	Yes	No	No	No	Not obvious	Not obvious	Possibly	Yes, developing asymmetric dystonia of upper limbs 7 y after initial presentation	Not obvious	Not obvious
Current age/ age at death (cause were known)	Died aged 9 y (pneumonia)	6 у	Died aged 10 y	12 y	Died age 18 years	4 y	8 y	11 y	9 y	6 y	7 y	3 y	y 9	35 y	13 y	2 y	2 y
Features at presenta- tion	Sudden-onset motor regression	Sudden onset motor regression	Loss of skills	Sudden onset motor regression	Sudden onset motor regression	Sudden onset motor regression	Falling	Loss of skills over a few weeks	Rapid loss of skills	Motor regression and speech arrest	Rapid loss of skills	Acute regression, dystonia, extra- pyramidal movements, orofacial dyskinesia	Rapid psychomotor regression, axial hypotonia, spastic dystonic tetraparesis	Pain, fatigue, anxiety, sleep problems	SP	Hypotonia and dystonia	Acute regression with dystonia necessitating ICU admission
Age at in- itial ascertain- ment	16 то	om 6	14 mo	11 mo	11 months	18 mo	2 y	2 y	18 mo	14 mo	11 mo	15 mo	12 mo	30 y	22 mo	4 mo	9 mo
Possible trigger	D & V	Bronchiolitis	No	URTI	URTI	Unspecified viral infection	No	N <sub>o</sub>	Unspecified viral infection	URTI	URTI	URTI/meningitis C vaccination	URTI	No	Vaccination	No	URTI
Develop- mental status prior to onset	Normal	Mild delay	Normal	Normal	Normal	Normal	Normal	Mild delay	Normal	Normal	Normal	Normal	Mild delay	Normal	Normal	Unknown	Normal
Sex	M	М	M	M	М	压	М	M	F	F	ц	Ħ	M	F	M	M	F
Individual	PI	PI	P1	P1	P2	PI	PI	ΡΙ	P1	P1	PI	PI	P1 (son to P2)	P2 (mother to P1)	P1	P1	PI
AGS number	AGS550	AGS567	AGS582	AGS663		AGS679	AGS699	AGS703	AGS720	AGS759	AGS765	AGS788	AGS810	<u> </u>	AGS943	AGS1115	AGS1170

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	ST	ST	Normal	SDT	BSN	SDT	SP becoming SDT	Clinically AGS- like (but no imaging)	BSN	BSN	BSN	AGS	BSN
	I	IV	I	Λ	>	Ш	IV	>	V	Ш	۸	Λ	IV
	п	IV	I	Λ	>	IV	Ш	>	Λ	IV	Λ	Λ	IV
	IV	V	I	>	>	^	VI	>	Λ	VI	Λ	Λ	Ŋ
	10.506 (5.53)	17.147 (3.06)	2.692 (30.18)	9.063 (16.68)	8.293 (8.56)	12.865 (10.33)	12.24 (6.38); 18.051 (6.43)	7.031 (2.17)	8.713 (5.55)	NT	NT	NT	ŢN
cortical atrophy and severely atrophied putamina (no CT)	Normal MRI (at 2 years) and CT (at 4 years)	MRI normal, BG and PV calcification on CT	No imaging	Mild hyper-intensity of the BG (no CT)	BSN	BG calcification (CT); normal MRI at age 10 years	Normal (no CT)	No imaging	BSN with BG calcification	BSN (no CT)	BSN with BG calcification	Characteristic of AGS	BSN with BG calcification
	ST; intellectually normal;	ST with severe ID	Normal	SDT with severe ID; DSH	SDT with some ID; DSH	SDT with some ID; AIHA	SDT with some ID	SDT with severe ID	SDT with severe ID; CB	SDT with some ID	SDT with severe ID	SDT with severe ID	SDT with severe ID
	Fluctuations	Yes, age 2.5 y episode of definite regression	No	Yes, with intermittent flares of encephalopathy and slowly progressive dystonia	Yes, episode of definite regression at age 4 years	No	Yes, with progressive involvement of UL and spastic dystonia	No	No	Yes, from uni- to bi- lateral; however, some skills (e.g., crawling, pulling to stand) subsequently reacquired	No	No	No
	6 y	4 y	31 y	17 y	y 9	11 y	٧ ٢	19 то	5 y	2 y	4 y	3 у	6 y
	ST with normal intellect	ST and speech delay	Always normal	Lethargy, dystonia, global regression	Developmental arrest with onset of generalized dystonia	Motor delay with spastic tetraparesis	Rapidly progressive SP	Onset of dystonia and loss of skills	Acute regression with dystonia necessitating ICU admission	Left hemiparesis with loss of ambulation	Tremor and rapid loss of skills	Developmental regression with loss of crawling and other skills	Developmental regression with loss of skills
	2.5 y	DD obvious by 1 y	Always normal	15 mo	13 mo	15 mo	21 mo	14 mo	11 mo	14 mo	12 mo	15 mo	13 mo
	No	No	NR	Otitis media	N <sub>o</sub>	No	°Z	No	Unspecified viral infection	Febrile illness	Otitis media	Febrile respiratory illness	MMR and varicella vaccination
	Normal	Delayed	Always normal	Normal	Moderate delay	Delayed	Normal	Likely delayed	Normal	Normal	Normal	Possible mild delay	Normal
	M	M	M	M	M	日	M	M	M	M	M	M	н
	P1 (brother to P2, son of P3)	P2 (brother to P1, son of P3)	P3 (Father to P1 and P2; mosaic)	P1	P1	PI	PI	P2	P1	PI	P1	P1	P1
	AGS1315	<u>I</u>		AGS1456	AGS1507	AGS1537	AGS1542	!	AGS1824	AGS1980	AGS1989	AGS2007	AGS2009
	cortical atrophy and severely atrophied putamina (no CT)	P1 (cortical atrophy and severety atrophied severet	P1 (brother M Normal No Dabvious by ST and speech delay	Hother Mornal Mornal No Delayed Mornal Mornal Mays normal Mays nor	Harring Line Line Line Line Line Line Line Line	Pi Corolle M Normal Normal and learning and the correction of the control and correctly according and correctly according and correctly according and a correctly according to the corr	Holistic Management of the control o	Pictorial arroy of secretary arroy of the control arrow of the control a	Fig. 10   Fig. 10   Fig. 10   Fig. 11   Fig.	Fig. 10   Fig. 20   Fig.	Principal Materials   Principal Materials	Part   Part	Property   Property

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GMFCS MACS CFCS Summary
Interferon scores (age, deci- malized years)
Neuroima- ging
Status at last con- tact
Progressive course
Current age/ age at death (cause were known)
Features at presenta- tion
Age at in- itial ascertain- ment
Possible trigger
Sex
AGS number Individual Sex Developmental status status prior prior to onset
AGS number

disability; IFN, interferon; IUGR, intrauterine growth retardation; LL, lower limb; MACS, Manual Ability Classification System; MRI, magnetic resonance imaging; NR, not relevant; NT, not tested; PV, periventricular; SD, spastic dystonia; SDT, spastic dystonic tetraparesis; SP, developmental delay; DSH, dyschromatosis symmetrica hereditaria; D & V, diarrhea and vomiting; GMFCS, Gross Motor Function Classification System; GP, globus pallidus; HSM, hepatosplenomegaly; ICC, intracranial calcification; ICU, intensive care unit; ID, intellectual Abbreviations: AGS, Aicardi-Goutières syndrome; AIHA, autoimmune hemolytic anemia; BG, basal ganglia; BSN, bilateral striatal necrosis; CB, chilblains; CFCS, Communication Classification System; CSF, cerebrospinal fluid; CT, computed tomography; DD, spastic paraparesis, ST, spastic tetraparesis, UL, upper limb; URTI, upper respiratory tract infection; WM, white matter.

Note: AGS1315\_P3 (different shading) is not included in the patient data analysis because of mosaic status; disability scales were not calculated for AGS228 because of age < 1 year at last contact.