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Autosomal Recessive Cerebellar Ataxias With Elevated Alpha-Fetoprotein: Uncommon Diseases, Common Biomarker

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ABSTRACT: alpha-Fetoprotein (AFP) is a biomarker of several autosomal recessive cerebellar ataxias (ARCAs), especially ataxia telangiectasia (AT) and ataxia with oculomotor apraxia (AOA) type 2 (AOA2). More recently, slightly elevated AFP has been reported in AOA1 and AOA4. Interestingly, AOA1, AOA2, AOA4, and AT are overlapping ARCAs characterized by oculomotor apraxia, with oculocephalic dissociation, choreo-dystonia, and/or axonal sensorimotor neuropathy, in addition to cerebellar ataxia with cerebellar atrophy. The genetic backgrounds in these disorders play central roles in nuclear maintenance through DNA repair [*ATM* (AT), *APTX* (AOA1), or *PNKP* (AOA4)] or RNA termination [*SETX* (AOA2)]. Partially discriminating thresholds of AFP have been

proposed as a way to distinguish between ARCAs with elevated AFP. In these entities, elevated AFP may be an epiphenomenon as a result of liver transcriptional dysregulation. AFP is a simple and reliable biomarker for the diagnosis of ARCA in performance and interpretation of next-generation sequencing. Here, we evaluated clinical, laboratory, imaging, and molecular data of the group of ARCAs that share elevated AFP serum levels that have been described in the past two decades. © 2020 International Parkinson and Movement Disorder Society

Key Words: recessive ataxia; alpha-fetoprotein; oculomotor apraxia; DNA/RNA repair

Autosomal recessive cerebellar ataxias (ARCAs) are a heterogeneous group of inherited neurodegenerative disorders that affect the cerebellum. Cerebellar ataxia (CA) with onset before age 30 years in a context of consanguinity and/or similar cases in siblings with healthy parents is

suggestive of ARCA. Some biomarkers, such as alpha-fetoprotein (AFP), are crucial for the diagnosis of ARCAs.¹

Recently, a group of ARCAs caused by mutations in genes implicated in DNA repair and/or transcriptional deficiencies has emerged. They are characterized by the combination of CA with oculomotor disturbances, such as oculomotor apraxia (OMA; see Supporting Information Video S1 and Fig. 1)^{2,3} or oculocephalic dissociation (OCD; see Supporting Information Video S6 and Fig. 1), with dystonia, chorea, sensorimotor axonal neuropathy, and elevated AFP serum level.¹ OMA is identified by constant horizontal saccades with increased latency sometimes associated with hypometric staircase saccades during head thrust and/or OCD that may also provide clues. OMA also includes difficulty in making voluntary saccades and may be responsible for OCD in some patients. OMA is easier to diagnose with

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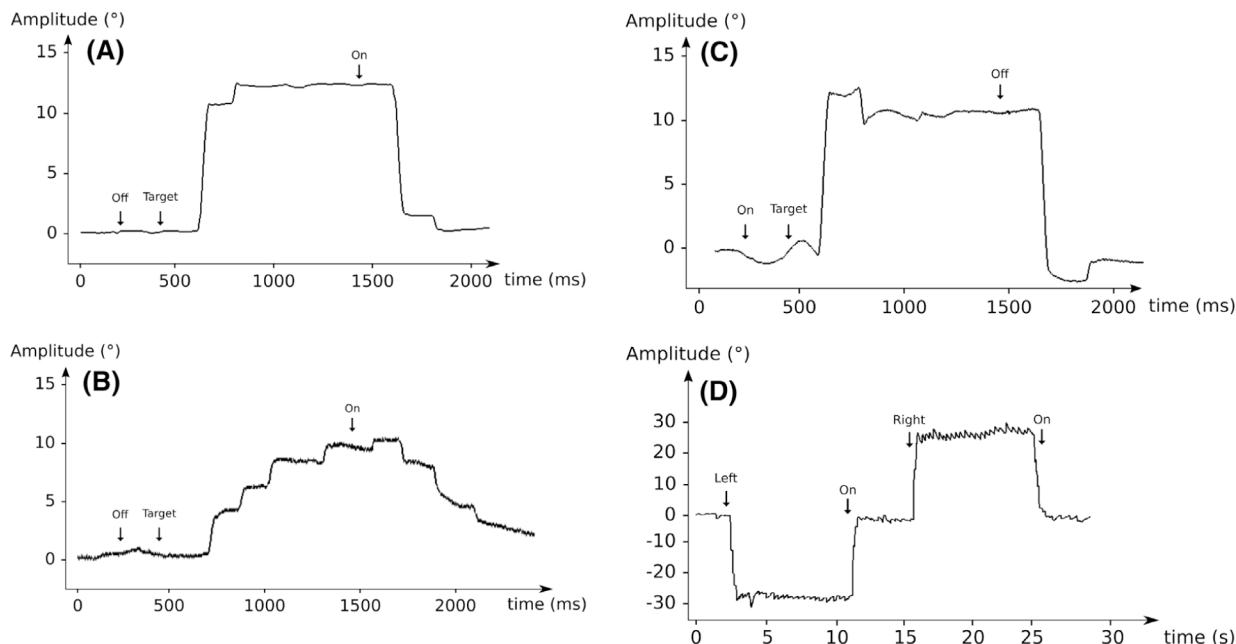


FIG. 1. Eye movement recordings. **(A)** Normal rightward saccade of control subject with a normal latency. **(B)** Hypometric rightward saccade with markedly increased latency in a patient with ataxia with oculomotor apraxia type 1 (AOA1). The latency increases in the patient with a saccade, starting later than the control. **(C)** Hypermetric centrifugal rightward saccade and hypermetric centripetal saccade in a patient with ataxia telangiectasia. **(D)** Gaze-evoked nystagmus in a patient with AOA2. Left, target on the left; off, center off; on, center on; right, target on the right; target, right target appears.

oculographic recording and may not be present in some patients with ataxia with oculomotor apraxia type 1 (AOA1) or AOA2, which can be a misleading factor. OCD dissociation is a delay of eye movement during a task of head rotation where the head reaches the target before the eyes. OCD may be caused by oculomotor apraxia or more frequently by very hypometric horizontal saccades. OCD is easier to diagnose clinically than OMA.

This new group includes ataxia telangiectasia [AT, *ataxia telangiectasia mutated (ATM)*],⁴ ataxia with AOA1 (*APT*X),⁵ AOA2 [*SET*X (*senataxin*)],⁶ and

AOA4 [*polynucleotide kinase 3'-phosphatase (PNKP)*].⁷ Here we review the group of ARCAs associated with an elevated AFP serum level (AT, AOA1, AOA2, and AOA4) (Table 1), which are rare, similar diseases that must be recognized and distinguished from each other for the best care of these patients.

AT, ATM Gene (OMIM: 208900)

AT is the second most frequent ARCA after Friedreich's disease. It is characterized by neurodegeneration, immunodeficiency, and predisposition to malignancy; with progressive CA, OCD and/or OMA, dystonia, sensorimotor axonal neuropathy, and elevated AFP serum level.^{4,8,9} Disease onset occurs most frequently around age 2 years and worsens progressively, leading to the loss of locomotion by 10 years, followed by death within the next decade.¹⁰ Dystonia is frequently accompanied by chorea, myoclonus, tremor, or parkinsonism.^{11,12} According to Levy and Lang,¹² the prevalence of movement disorders during the course of AT includes cerebellar symptoms (96%), myoclonus (92%), dystonia (89%), choreoathetosis (89%), tremor (74%), and parkinsonism (41%). Brain magnetic resonance imaging (MRI) reveals cerebellar atrophy. With later onset, a milder phenotype and consequently a longer life span are possible. Patients with the milder expression of AT typically have neurological deficits in childhood with slower age-related neurodegeneration. The predominant neurological symptoms at disease

TABLE 1. The ARCAs with elevated AFP

Mechanism	Gene	Protein	Disease
DNA double-strand break repair	<i>ATM</i>	ATM	AT
DNA single-strand break repair	<i>APT</i> X <i>PNKP</i>	Aprataxin PNKP	AOA1/EOAH AOA4 (MCSZ)
RNA termination/maturation	<i>SET</i> X	Senataxin	AOA2 (ALS4)
Channel	<i>ANO10</i>	ANO10/ TMEM16K	ARCA3

Abbreviations: ARCA, autosomal recessive ataxia; AFP, alpha-fetoprotein; ATM, ataxia telangiectasia mutated; AT, ataxia telangiectasia; AOA1, ataxia with oculomotor apraxia type 1; EOAH, early-onset ataxia with oculomotor apraxia and hypoalbuminemia; PNKP, polynucleotide kinase 3'-phosphatase; AOA4, ataxia with oculomotor apraxia type 4; MCSZ, microcephaly, seizures, and developmental delay; AOA2, ataxia with oculomotor apraxia type 2 (AOA2); ALS4, amyotrophic lateral sclerosis 4 (juvenile); TMEM16K, transmembrane protein 16K; ARCA3, autosomal recessive ataxia type 3 (ARCA3).

TABLE 2. Clinical and biological characteristics of ARCAs with elevated AFP

	AT	ATLD	AOA1/EOAH	AOA2	AOA4	ARCA3
Age of onset (years)	Infancy/2–5 (possible late onset)	Childhood	Infancy/childhood 2–10	Teens 10–15	Infancy/childhood 1–9	Adulthood/ 10–40
Age of onset, extreme range (years)	1–41	3–47	2–49	7–30	1–10	6–43
Progression/Severity	Rapid/Severe	Intermediate/Moderate	Rapid/Moderate	Rapid/Moderate	Rapid/Moderate	Slow/Moderate
Neurological signs	Dystonia Chorea Neuropathy OCD/OMA	Dystonia Chorea Neuropathy OMA	Intellectual disability Dystonia Chorea Neuropathy OCD Hypometric saccades Optional OMA	Dystonia Chorea Neuropathy Optional OCD Optional OMA	Dystonia Cognitive impairment Neuropathy OMA	Extensor plantar reflexes Spasticity Cognitive decline Downbeat nystagmus Lack of neuropathy
Extraneurological signs	Telangiectasias Immunodeficiency and recurrent infections Photosensitivity Risk for cancers (lymphoid malignancies)	Lack of telangiectasias Possible risk for cancers	None	None	None	None
AFP elevation (levels)	Marked: mostly 200 µg/L increase with disease duration	Normal level: <5 µg/L	Slightly increased in 40% of AOA1: 5–20 µg/L increase with disease duration	Always increased: 15–65 µg/L stable	Variably increased: 10–30 µg/L	Rarely increased 5–15 µg/L
AFP, extreme range (µg/L)	5–2800	0–7	1–17	5–185	6–30	1–17
Others biomarkers	Mixed immune deficiency (CD4/CD8 deficiency, IgA/IgG deficiency) Chromosomal rearrangements (7:14 translocations) on the karyotype		Optional hypoalbuminemia and hypercholesterolemia	Elevated CK	Elevated CK Optional hypoalbuminemia	

Abbreviations: ARCA, autosomal recessive ataxia; AFP, alpha-fetoprotein; AT, ataxia telangiectasia; ATLD, ataxia telangiectasia-like disorder; AOA1, ataxia with oculomotor apraxia type 1; EOAH, early-onset ataxia with oculomotor apraxia and hypoalbuminemia; AOA2, ataxia with oculomotor apraxia type 2 (AOA2); AOA4, ataxia with oculomotor apraxia type 4; ARCA3, autosomal recessive ataxia type 3 (ARCA3); OCD, oculocephalic dissociation; OMA, oculomotor apraxia; IgA, immunoglobulin A; IgG, immunoglobulin G; CK, creatine kinase.

onset and during the early stages may be dystonia, myoclonus, or tremor, with ataxia appearing subsequently. AFP should therefore be considered part of the workup in early-onset, unexplained dystonia. OMA may also appear later or not at all⁹ (Tables 2 and 3; Supporting Information Videos S1, S2, and S3).

Extraneurological findings are conjunctival, auricular, or buccal telangiectasia; immunodeficiency with recurrent infections (sinusitis and lung infections caused by *Hemophilus influenzae* or *Streptococcus pneumoniae*), increased cancer risk (especially hematological, such as lymphomas and leukemias, but also solid tumors)^{9,13}; and sensitivity to ionizing radiation. Follow-up of patients with AT is recommended to prevent or treat infections (intravenous immunoglobulin, antibiotics) or malignancies. Early detection may allow a better prospective evaluation of AT and thus assist in more

effective treatment for both the neurological and immune abnormalities.^{14,15} Van Os et al.¹⁶ have proposed guidelines for response to the medical problems encountered during the course of AT with regard to neurological, immunological, infectious, oncological, endocrinological, and nutritional disorders, as well as perioperative risk. In this particular form of AT, immunodeficiency may occur but is less common, whereas malignancies tend to appear later in life and include a higher proportion of nonhematopoietic cancers.⁹

The AFP serum level is markedly increased in 90% of patients, often greater than 100 µg/L (normal < 5 µg/L). A few patients with AT did not experience AFP elevation or a delayed increase of AFP during the clinical course of the disease.¹⁷ CD4 and CD8 immunodeficiencies, IgA or IgG immunodeficiency, and chromosomal translocations (especially 7:14 on the

TABLE 3. Comparison of the clinical features in ARCAs with OCD/OMA

Neurological features	AT	ATLD	AOA1	AOA2	AOA4
Intellectual disability	+/-	-	++	-	++
Learning difficulties					
Cerebellar atrophy	++	++	++	++	++
Sensorimotor polyneuropathy	++	+	+++	++	++
Dystonia	+++	++	++	++	++
	May be the predominant sign				
Chorea	++	+	++	+	?
			Could improve with disease progression		
Myoclonus	++	?	+	+	?
OCD	++	+	++	+	++
OMA	Genuine OMA		Pseudo-OMA (hypometric saccades) > OMA		Not severe
					Early and severe

Abbreviations: ARCA, autosomal recessive ataxia; OCD, oculocephalic dissociation; OMA, oculomotor apraxia; AT, ataxia telangiectasia; ATLD, ataxia telangiectasia-like disorder; AOA1, ataxia with oculomotor apraxia type 1; AOA2, ataxia with oculomotor apraxia type 2; AOA4, ataxia with oculomotor apraxia type 4.

karyotype⁸) are frequent.¹⁸ The variability of AT phenotypes and the large size of *ATM* (66 exons) may render diagnosis difficult; in support, laboratory testing proves very helpful.

The *ATM* gene¹⁹ encodes a phosphatidylinositol 3-kinase involved in cell-cycle progression, cellular response to DNA alterations, and maintenance of genome stability⁴ (Table 1).

Two rare forms of ataxia telangiectasia-like disorder (ATLD) have been described with a similar clinical picture and the same pathway, yet without AFP elevation: ATLD [*meiotic recombination 11 (MRE11)* gene] and ATLD2 [*proliferating cell nuclear antigen (PCNA)* gene].^{20–22} The *MRE11* gene encodes a protein (Mre11) with nuclease and DNA-binding activity. Together with Rad50 (DNA-repair protein 50) and Nbs1 (Nijmegen breakage syndrome protein 1 or Nibrin), Mre11 forms the MRN complex, which is a target of ATM kinase and is also involved in the signaling network of cellular response to DNA damage (Fig. 2B).

AOA1, *APTX* Gene (OMIM: 208920)

AOA1 has been described as a CA of early onset, most frequently before age 10 years.²³ Patients suffer from intellectual deficiency (50%) accompanied by cerebellar atrophy, OCD and/or OMA, choreo-dystonia, and severe peripheral neuropathy.^{5,24} OCD and/or OMA may not be present at disease onset but are expressed a few years later (Tables 2 and 3; Supporting Information Videos S6, S7, and S8).

Recently, we reported that OMA correlates with diseases of greater severity; the p.Trp279* *APTX* mutation was the most frequent among the white population, whereas the *APTX* missense mutations were associated with a milder phenotype.²⁵ In Japan, AOA1

is the most frequent ARCA, named early-onset AOA and hypoalbuminemia.

During the course of AOA1, decreased serum albumin levels and elevated total cholesterol levels may be observed, suggesting hypoalbuminemia and hypercholesterolemia, found in 63% of patients, as the most characteristic biochemical findings in AOA1, even if they are optional.^{25,48} AFP serum level was reported to be elevated in 40% of the 80 patients with AOA1 studied.^{25,26} This slight increase of AFP, ranging from 15 to 20 µg/L, may be significant.

Whether patients with AOA1 have coenzyme Q10 (CoQ10) deficiency is not yet clear.^{26,27}

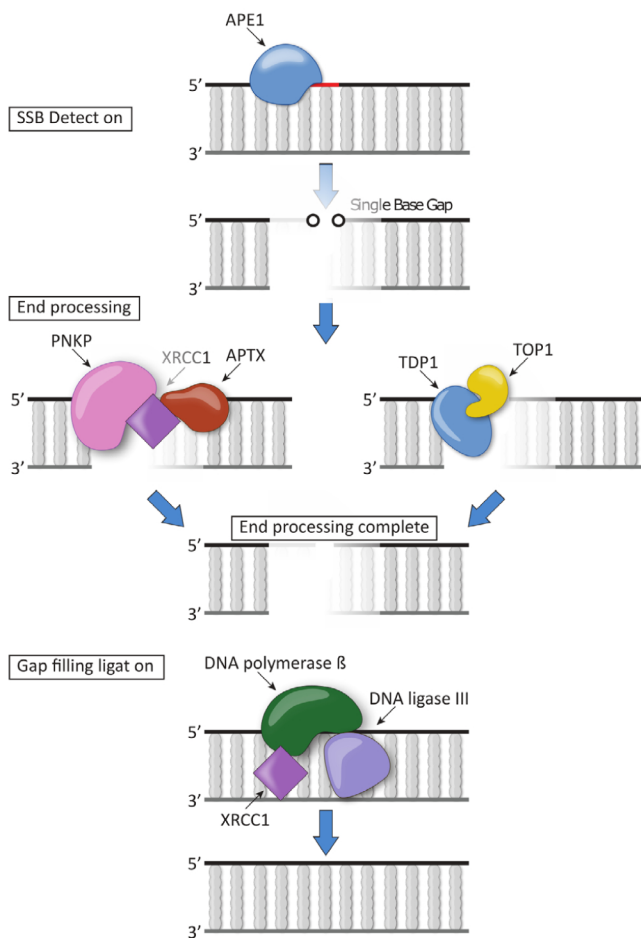
AOA2, *SETX* Gene (OMIM: 606002)

Disease onset in AOA2 is usually between ages 12 and 20 years, which is later than in AOA1 and AT.^{8,24,28} Patients with AOA2 experience progressive CA, along with sensorimotor neuropathy, occasional OCD and/or OMA, strabismus, chorea, dystonia, and elevated serum AFP levels. Brain MRI reveals marked cerebellar atrophy (Tables 2 and 3; Supporting Information Videos S9, S10, S11, and S12). Recently, three cases of disruption of spermatogenesis and infertility were reported in AOA2.²⁹

AFP serum levels are between 15 and 65 µg/L, averaging around 50 µg/L.³⁰ Healthy subjects who carry a heterozygous *SETX* mutation may have a mildly elevated AFP serum level, while laboratory examination may reveal elevated creatine kinase serum levels^{31,32} (Table 2).

SETX encodes senataxin, a protein containing a DNA/RNA helicase domain, which is involved in RNA processing, RNA termination, and coordination of transcription events, and may also act as a genome guardian^{6,28,32} (Table 1 and Fig. 3). Missense mutations

(A) DNA single-strand break repair (SSBR)



(B) DNA double-strand break repair (DSBR)

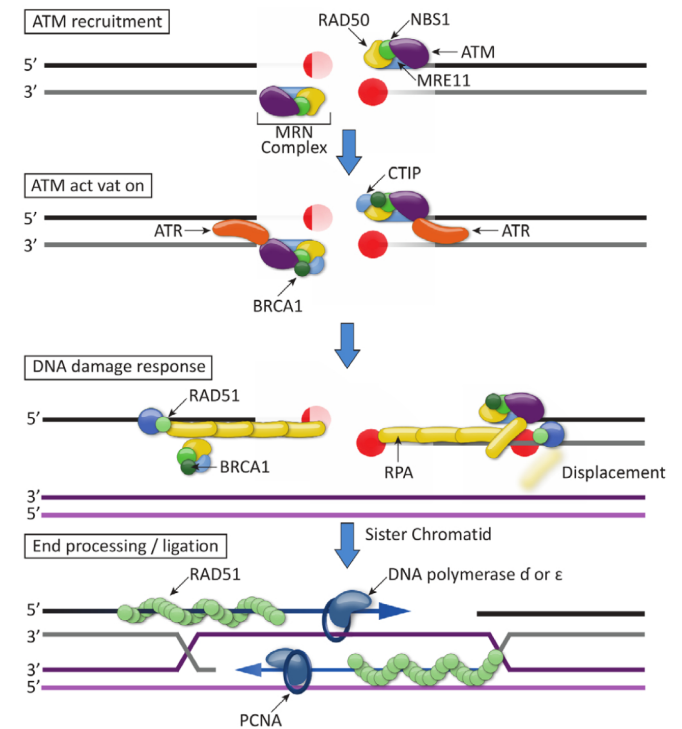


FIG. 2. DNA repair in autosomal recessive cerebellar ataxias (ARCA). Two distinct groups of ARCA are caused by defective DNA repair. **(A)** DNA single-strand break repair (SSBR). The first group includes ataxia with oculomotor apraxia type 1 (AOA1) caused by mutations in *APT*X AOA4 (*PNKP*), spinocerebellar ataxia with axonal neuropathy (SCAN1, *TDP1*), and cerebellar ataxia and peripheral neuropathy caused by *XRCC1*. The proposed role of *APT*X, *PNKP*, *TDP1*, and *XRCC1* genes in repair and maintenance of single-strand break (SSB) is shown. First, the damaged base is recognized by a DNA glycosylase (not indicated). An endonuclease (*APE1*, for example, in *A*) cleaves the phosphoribose backbone, leaving a single-base gap in the DNA. A specific end-processing enzyme such as *PNKP* (AOA4), *APT*X (AOA1), or *XRCC1* modifies the DNA ends to facilitate further repair. Alternatively, *TDP1* (SCAN1) and *TOP1* can process the ends of the DNA breaks. DNA polymerase β removes the phosphoribose left by *APE1* and fills the gap with a new base. The resulting nick is sealed by DNA ligase I or III. **(B)** DNA double-strand break repair (DSBR). The second group includes ataxia telangiectasia (AT, *ATM*), ataxia telangiectasia-like disorder (ATLD, *MRE11*), and recessive ataxia caused by abnormal cellular responses to DNA double-strand break (DSB). The role of ATM in the cellular response to DNA damage mediated by the BRCA-associated genome complex is shown. This homologous recombination is a highly complex pathway that uses a sister chromatid as a template to repair a double-strand DNA break. ATM is intimately associated with the MRE11-RAD50-NBS1 (MRN) complex. Following MRN complex binding, ATM is activated and autophosphorylates its inactive dimer at serine 1981. Monomeric, active ATM is then recruited to the damage site, where it phosphorylates downstream targets, including SMC1, Nbs1, Chk2, BRCA1, and histone H2AX. Furthermore, ATM phosphorylates p53 at serine 15. The DSB is recognized by the MRN complex by activation of ATM. The MRN complex and CTIP determine the DNA break point and resect the DNA to yield 3' protruding ends stimulated by BRCA1. The single-stranded DNA is coated by the RPA complex. RPA is recognized by ATR, which becoming activated together with ATM promotes cell-cycle arrest. BRCA1 recruits RAD51 and initiates invasion of the DNA to the homologous region on the sister chromatid. DNA is then synthesized by DNA polymerase δ or with the assistance of PCNA by using the homologue DNA as a template. *APE1*, apurinic-apyrimidinic endonuclease 1; *APT*X, aprataxin; *ATM*, ataxia telangiectasia mutated; *ATR*, ataxia telangiectasia and RAD3-related; *BRCA1*, breast cancer 1; *CTIP*, C-terminal binding protein interacting protein; *MRE11*, meiotic recombination 11; *NBS1*, Nijmegen breakage syndrome protein 1 or Nibrin; *PCNA*, proliferating cell nuclear antigen; *PNKP*, polynucleotide kinase 3'-phosphatase; *RAD50*, DNA-repair protein 50; *RPA* complex, Replication protein A complex; *TDP1*, tyrosyl-DNA phosphodiesterase 1; *TOP1*, DNA topoisomerase 1; *XRCC1*, X-ray repair cross-complementing protein 1. [Color figure can be viewed at wileyonlinelibrary.com]

located in the helicase domain were more frequently associated with dystonia.²³

Heterozygous mutations of *SETX* have been reported in patients affected with juvenile-onset amyotrophic lateral sclerosis (ALS4),³³ although this is a subject of debate.³⁴

AOA4, *PNKP* Gene (OMIM: 616267)

AOA4 has been described as a combination of CA, OMA, and/or OCD, and sensorimotor peripheral neuropathy with onset between ages 1 and 9 years because of mutations in the *PNKP* gene.⁷ Some patients

experience cognitive impairment, and brain imaging reveals cerebellar atrophy^{7,35,36} (Tables 2 and 3). In AOA4, AFP may be increased from 1.5 to 4 times the normal level. Some cases express low albumin levels and elevated cholesterol as in AOA1.^{36,37}

Recently, a patient with very late onset (50 years) AOA4 had been described who had ataxia, polyneuropathy, OMA, and cognitive impairment.³⁸

AOA4 may be the second most frequent recessive ataxia in Portugal, after Friedreich's ataxia.⁷ Elsewhere, *PNKP* mutations underlie another neurodevelopmental disorder named microcephaly with seizures.³⁹ The *PNKP* gene encodes PNKP that has a crucial role in DNA damage repair⁴⁰ (Fig. 3), including both single-strand breaks (SSBs) (Fig. 2A) and double-strand breaks.⁴¹

Elevated AFP in ARCAs (AT, AOA1, AOA2, AOA4, and ARCA3)

AFP is a 70-kDa glycoprotein produced by the liver that is involved in the transport of different ligands, chemotaxis, oxygen free radical scavenging, and lipid peroxidation.^{42,43} During physiological development, an inverse correlation exists between serum concentrations of AFP and albumin.⁴⁴

Among ARCAs, elevated AFP is typical of patients with AT or AOA2⁴² and may be encountered in AOA4, AOA1, and to some extent ARCA3.^{7,25,28,45} ARCA3, which is due to mutations of ANO10, a calcium-activated chloride channel,⁴⁶ causes progressive CA with cerebellar atrophy, optional extensor plantar reflexes, and spasticity, although without peripheral neuropathy, OCD, OMA, or movement disorders. Almost all patients with AOA2 and AT have increased AFP serum levels over the course of the disease, higher in fact than in patients with AOA1, whose AFP levels remain above controls.²⁵ Mariani et al.²⁸ have reported that application of discriminatory thresholds of AFP could be helpful in distinguishing AOA1, AOA2, and AT with high specificity and predictive values. The levels include: AT: AFP > 65 µg/L; AOA2: 15 µg/L < AFP < 65 µg/L; AOA4: 10 µg/L < AFP < 30 µg/L; AOA1 and ARCA3: 5 µg < AFP < 20 µg/L (Table 2). The same study reported that AFP levels of 7 to 15 µg/L had a specificity of 93%, positive predictive value (PPV) of 78%, and negative predictive value (NPV) of 86% for patients with AOA1 relative to AOA2 and AT; AFP levels of 15 to 65 µg/L had a specificity of 86%, PPV of 64%, and NPV of 89% for patients with AOA2 relative to AOA1 and AT; AFP levels greater than 65 µg/L had a specificity of 90%, PPV of 83%, and NPV of 73% for patients with AT relative to AOA1 and AOA2.²⁸

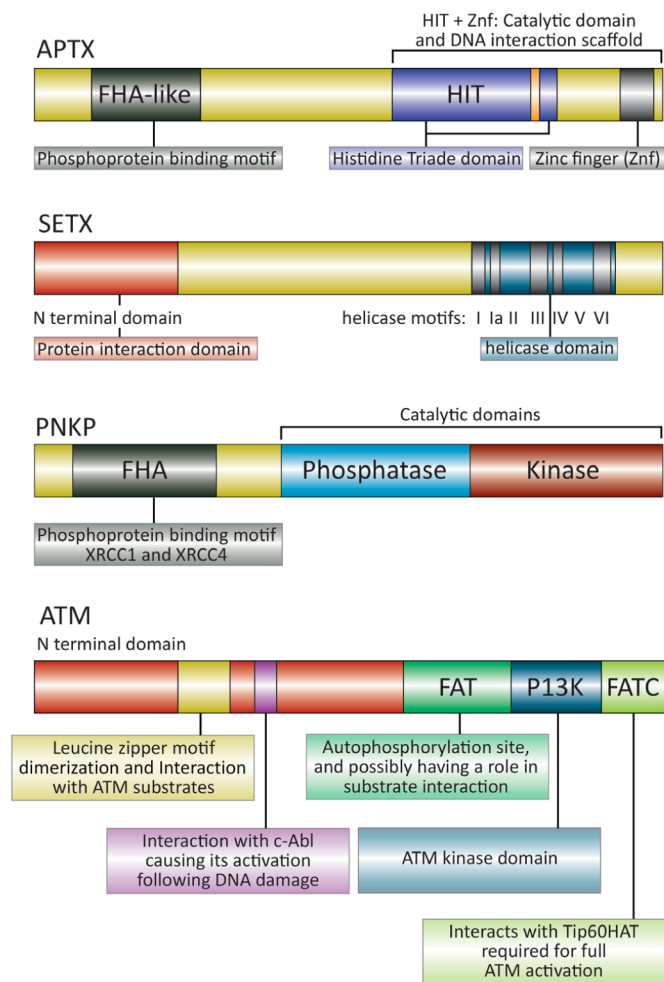


FIG. 3. Schematic representation of the domains of the APTX, SETX, PNKP, and ATM proteins (AOA1, AOA2, AOA4, and AT). APTX and PNKP: APTX interacts with several proteins involved in excision repair. FHA represents the forkhead-associated domains in APTX and PNKP, while HIT and Zinc finger represent the histidine triad and Zinc finger domains in APTX. The HIT motif (HxHxH) is highlighted in orange. For PNKP, we show only the pathogenic variants for AOA4, because interestingly, they are located on the inside of the kinase domain for AOA4, but not MCSZ. SETX: the Senataxin protein diagram shows two key domains, including the amino-terminal protein interaction domain and the helicase domain (with the seven regions of highest conservation indicated as helicase motifs). ATM: the ATM protein is a nuclear serine/threonine protein kinase activated by autophosphorylation after DNA damage. We show the domain architecture of ATM and the associated functions. The ATM protein has a focal adhesion targeting domain, a kinase domain, and focal adhesion targeting C-terminal domain at the extreme C terminus. The C-terminal kinase domain of ATM classifies it in the PIKK superfamily of protein kinases. AFP, alpha-fetoprotein; AOA1, ataxia with oculomotor apraxia type 1; AOA2, ataxia with oculomotor apraxia type 2; AOA4, ataxia with oculomotor apraxia type 4; APTX, aprataxin; AT, ataxia telangiectasia; ATM, ataxia telangiectasia mutated; FAT, focal adhesion targeting; FATC, FAT C-terminus; FHA, Forkhead-associated domain; HIT, histidine triad; MCSZ, microcephaly with seizures; PI3K, phosphatidylinositol 3-kinase; PNKP, polynucleotide kinase 3'-phosphatase; PIKK, phosphatidylinositol 3-kinase-related kinase; SETX, senataxin; XRCC1, X-ray repair cross-complementing protein 1. [Color figure can be viewed at wileyonlinelibrary.com]

Only in very rare cases is AFP normal in AT, while serum AFP increases continuously with age in patients with classical AT disease.¹⁷ Thus, even if initially

normal, AFP serum level assessments should be repeated during the evolution of CA, dystonia, and/or chorea of unknown etiology (Table 2).

Elevated AFP level in these ARCAs could be an epiphenomenon not directly linked to DNA repair disruption but to liver transcriptional dysregulation.^{1,47} Indeed, ATM kinase has many effects, not only on DNA repair, explaining the predisposition to cancers, but also on transcription, which could induce both neurological signs and hepatic effects leading to increased AFP.

In AOA1, elevated AFP and hypoalbuminemia are two consequences of transcriptional impairment in the liver, where AFP and albumin genes have transcriptional regulatory mechanisms of opposite character.^{43,44} We may hypothesize that the same mechanism exists in AOA4, because hypoalbuminemia has also been described in this recently identified entity.³⁶

In ARCA3, the AFP serum level has been found to increase mildly in two cases [16.6 and 11.6 µg/L (normal < 5 µg/L)] in a previous series of nine patients.⁴⁵ Whether AFP could be a biomarker for ARCA3 remains to be elucidated by further analysis of future cases.

DNA Repair Pathways (AT, AOA1, AOA2, and AOA4) and Susceptibility to Malignancies (AT)

Patients with AT have a highly increased incidence of cancers (approximately 38% lifetime risk despite the short life span), not only lymphomas and leukemias, but also lymphoid and solid tumors (breast cancer, pancreatic cancer, and melanoma). ATM is known as the central regulator of the DNA damage response (Fig. 2B). During induction of DNA double-strand breaks, the MRN complex, containing Mre11, Rad50, and Nbs1, binds to the damaged site.^{49,50} Loss of Nbs1, which is responsible for Nijmegen breakage syndrome 1, leads to a different phenotype, characterized by microcephaly, short stature, and recurrent infections, but not AFP elevation.⁴⁹ Rad50 deficiency, named Nijmegen breakage syndrome-like disorder, includes microcephaly, intellectual deficiency, short stature, and lymphoid malignancy.⁵¹ Activation and repression of ATM's downstream targets leads to senescence, genome repair, or apoptosis⁵² (Figs. 2B and 3). It has not yet been elucidated whether ATLD confers a risk for malignancies, but two brothers with ATLD died of lung cancer at ages 9 and 16 years, respectively, which is unusual.⁵³ Susceptibility to cancer has not been described for AOA1, AOA2, or AOA4.

The presence of a common single-strand break repair pathway links AOA1, AOA2, and AOA4. Tyrosyl DNA-phosphodiesterase 1 (*TDP1*), aprataxin (*APTX*),

and polynucleotide kinase/phosphatase (*PNKP*) each act on a discrete set of modified strand-break termini⁵⁴ (Fig. 2A). Spinocerebellar ataxia with axonal neuropathy (*SCAN1*) caused by mutated *TDP1* is another ARCA due to defective SSB repair, especially arising from abortive topoisomerase-1 activity.⁵⁵ Similar to AT, AOA1, and AOA4, patients with *SCAN1* have ataxia with CA, although the onset of the disease is delayed, and OCD, OMA, or elevated AFP were not reported.⁵⁶ However, the biological features of *SCAN1* include hypoalbuminemia and hypercholesterolemia, as in AOA1.

Many DNA repair syndromes are known to be associated with transcriptional defects, and disruptions in transcription could contribute to clinical variations, as happens in subgroups of xeroderma pigmentosum (XP) and related syndromes characterized by skin damage. Recently, two patients were described with CA and chorea as a result of mutations in *ERCC4* encoding XPF (XP type F) protein.⁵⁷ XPF is part, with *ERCC1*, of a two-subunit nuclease, which participates in nucleotide excision repair of DNA and in chromosome stability maintenance. Here, too, neurodegeneration may be related to transcriptional disruptions rather than to a DNA repair defect, because the dual role of XP factors in both nucleotide excision repair and initiation of transcription is well established.⁵⁸ Whereas DNA repair is significantly altered among the ARCAs with elevated AFP, and because DNA repair defects account for susceptibility to malignancies, we hypothesize that transcriptional deficiency could account for the neurodegeneration in all ARCAs with elevated AFP.

Clinical and Biological Overlap Between ARCAs With OMA and/or OCD and Elevated AFP

In clinical practice, the association of early-onset progressive CA with OMA and/or OCD and AFP is very suggestive of a limited number of ARCAs: AT, AOA1, AOA2, and AOA4. OMA is an interesting oculomotor finding, but its definition remains controversial. OMA (see earlier) was initially described under congenital conditions as the inability to turn the eyes voluntarily in a direction that aligns with a compensatory head movement.^{2,59} However, because apraxia is defined as the inability to move a certain part of the body in accordance with a proposed purpose, the motility of this part being otherwise preserved,⁵⁹ the word "apraxia" is not really appropriate to describe oculomotor disorders in this group of ARCAs, given that the motility of the eyes is not preserved. Whereas repetitive, hypometric saccades or a saccade of increased latency may be difficult to distinguish without oculographic recording, it may be much easier to recognize head

thrust and OCD than genuine OMA (Supporting Information Videos S1, S6, S9, and S10). In contrast with AT, genuine OMA may not be present in AOA1 (where very hypometric saccades are seen frequently) or AOA2 (Tables 2 and 3). Thus, we may consider that OCD, head thrust, and to some extent hypometric staircase saccades should be used where appropriate, rather than the misleading OMA, to describe the oculomotor abnormalities (Supporting Information Videos S1, S6, S8, S9, and S10). OMA has not often been found in other ARCAs; therefore, when present in the context of ataxia, it is very suggestive of either AOA1, AOA2, AOA4, AT, or ATLD, although its presence does not help distinguish between these entities. At present, video-oculography cannot differentiate AT, AOA1, and AOA2, and is not mandatory in the diagnostic workup of patients suspected to have ARCA; an appropriate oculomotor examination remains crucial.²⁸ Complex oculomotor impairment, including gaze-evoked nystagmus, hypermetric saccades, hypometric saccades, saccades with increased latencies, and antisaccade task disturbances, reflects diffuse cerebellar (including flocculus/paraflocculus, fastigial nuclei, and vermis), as well as pons, midbrain, and parietofrontal cortex, involvement in these entities.^{2,24,28,60}

Age at onset and disease progression may be helpful in distinguishing ARCAs with elevated AFP.^{1,54} The mean age at onset in AT, AOA1, and AOA4 has been reported to be lower than in AOA2. The course of AOA2 appears to be less severe than in AT or AOA1. Patients with AOA2 become wheelchair bound following a mean disease duration of 15 to 20 years, whereas most patients with AT and AOA1 are wheelchair bound after a mean disease duration of 10 and 11 years, respectively^{24,61} (Tables 2 and 3). Forms of AT in adults have been described either with early onset but a slow disease progression and sustained survival, or with a later onset (such as AOA1, or even later, such as AOA2). Thus, when assessed in adulthood, the severity and picture of AT, AOA1, and AOA2 may be surprisingly similar, including complex oculomotor disturbances, movement disorders, and elevated AFP.^{30,62}

A combination of several types of movement disorder are commonly found in AT, ATLD, AOA1, AOA2, and AOA4. Dystonia may be encountered in almost all patients with AT, AOA1, AOA2, and AOA4⁶³ (Supporting Information Videos S2, S5, S8, and S12). Chorea is also found in all of these entities, but more frequently in AOA1.⁶³ Interestingly, chorea and/or dystonia may improve during the course of AOA1.²⁴ Hyperkinesias were reported to improve spontaneously over time in one patient with AOA4.³⁶ Myoclonus is more frequent in AT than in AOA1 and AOA2. Parkinsonism seems to be more specific in AT, but it is found in only a third of patients with AT.³⁰ Dopaminergic nigrostriatal neuron loss has been reported in some

patients with AOA1 in the absence of obvious parkinsonism.⁶⁴ Occasionally, rather than CA, the hyperkinetic movement disorder may be the initial and/or most prominent sign of the disease. It should be kept in mind that cases of isolated dystonia, myoclonus dystonia, or chorea also occur, particularly in AT^{65,66} (Tables 2 and 3). Furthermore, prominent camptocormia has been reported in AT.⁶⁷

In 2019, Ronsin et al.⁶⁸ reported a new MRI abnormality in AOA. Hypointensity caused by iron deposition normally observed on susceptibility weighted imaging in the dentate nuclei is not observed in patients with AOA. Thus, loss of low signal in the dentate nuclei on fluid-attenuated inversion recovery (FLAIR) and susceptibility weighted imaging could be a promising marker of AOA.

Diagnostic Strategies for ARCAs, Therapeutic Options, and Management in Clinical Practice

Currently, the number of genes that when mutated lead to ataxia increases every month thanks to next-generation sequencing (NGS).⁶⁹ Our team created and validated the Recessive Ataxia Ranking Differential Diagnosis Algorithm (RADIAL), designed to improve the differential diagnostic approach toward ARCA by using patient features to predict the underlying gene responsible.⁷⁰ AFP assessment may guide the reverse phenotyping in case of first genotyping (ie, laboratory tests are performed based on the results of NGS, but not prior to the technique), and it may be helpful to interpret the multiple variants of unknown significance provided by premature NGS (ie, NGS performed before suitable phenotyping).

Today, the tendency is not to perform direct sequencing except for particular cases (very high AFP and/or telangiectasia for AT), or if an additional case has been diagnosed in the same family. Targeted-panel NGS is an interesting approach, because it provides a reliable clinical application.⁷¹ Whole-exome sequencing (WES) is an alternative genetic testing approach, particularly in patients for whom targeted-panel methodologies have not identified causative sequence variations or the clinical presentation is very complex (with multiple neurological and/or extraneurological signs), especially when the DNA of the parents is available. For patients in whom genetic disease is strongly suspected and targeted-panel sequencing and WES have both failed, whole-genome sequencing should be considered.⁷² WES or even whole-genome sequencing will probably become the first-line genetic test in the near future. In all cases, though, we recommend proper clinical examination, brain MRI, electromyography, and some biomarkers, such as AFP and vitamin E, before genetic testing to define the phenotype as precisely as possible.

Symptomatic, multidisciplinary management involving physiotherapy, speech therapy, management of orthopedic complications, and spasticity is necessary for each patient, as well as treatment of pain, anxiety, or depression when present. Intensive home-based speech therapy can improve dysarthria in ARCAs.⁷³ Continuous coordinative training leads to long-term improvements that translate to real-world function.⁷⁴

These measures improve the quality of life by trying to minimize the consequences of symptoms in everyday life. The establishment of a molecular diagnosis is crucial to provide suitable genetic counseling for patients and their relatives, diagnose treatable ARCAs,^{1,77} and include willing patients in clinical trials. Confirmation of the diagnosis of AT at an early stage can have important implications because of the risk for cancers (in patients and parents), recurrent infections, and avoidance of ionizing radiation.

Oxidative stress has been identified recurrently in different progressive cerebellar diseases and represents a widely investigated target for treatment.⁷⁵ In 2015, Romano et al.⁷⁶ described a novel and potentially important application of riluzole in patients with SCA or Friedreich's ataxia. The investigators noted that riluzole significantly improved ataxia at 3 and 12 months, and supported the idea that it could be a treatment for CA. Disease-specific trials, some already in progress in ataxia, are needed to confirm whether these findings can be applied in clinical practice. An ongoing French clinical trial assesses the efficacy of CoQ10 versus placebo in AOA1 patients, because CoQ10 deficiency has been reported in some patients with AOA1.

Gene-related therapy increasing genetic expression is a promising approach, because many ARCAs are caused by mutations leading to loss of function. Gene-related therapies for ARCAs have been developed furthest in Friedreich's ataxia; recent strategies to restore frataxin have revealed a potential for slowing or even stopping disease progression. Several of the treatment principles might well be applicable by analogy to other ARCAs in the future.⁷⁷

Hoch et al.⁷⁸ have identified PARP1 as a possible target for treating CAs associated with an unrepaired SSB. The development of selective inhibitors of PARP1 that prevent DNA binding by this enzyme could have therapeutic potential.

Future research in cerebellar dysfunction should include analysis of specific and potential candidate drugs for both symptomatic and disease-modifying studies, with sensitive clinical measures and reliable biomarkers.^{77,79} Novel therapies that target deleterious pathways, such as therapies acting on RNA, are also under development.⁸⁰

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