A Second Case With the V374A KCND3 Pathogenic Variant in an Italian Patient With Early-Onset Spinocerebellar Ataxia

Flavia Palombo, PhD, Chiara La Morgia, MD, PhD, Claudio Fiorini, PhD, Leonardo Caporali, PhD, Maria Lucia Valentino, MD, Vincenzo Donadio, MD, PhD, Rocco Liguori, MD, and Valerio Carelli, MD, PhD Correspondence Dr. Carelli valerio.carelli@unibo.it

Neurol Genet 2022;8:e200004. doi:10.1212/NXG.0000000000200004

Abstract

Background and Objectives

To date, approximately 20 heterozygous mainly loss-of-function variants in KCND3 have been associated with spinocerebellar ataxia (SCA) type 19 and 22, a clinically heterogeneous group of neurodegenerative disorders. We aimed at reporting the second patients with the V374A KCND3 mutation from an independent family, confirming its pathogenic role.

Methods

We describe the clinical history of a patient with SCA and conducted genetic investigations including mitochondrial DNA analysis and exome sequencing.

Results

This male patient was reported to have unstable gait with tremors at the lower limbs and dysarthric speech since childhood. A neurologic examination also showed dysarthria, nystagmus, action tremor, dysmetria, and weak deep tendon reflexes. He had marked cerebellar atrophy at brain MRI, more evident at vermis. Molecular analysis, including exome sequencing and an in silico panel analysis of genes associated with SCA, revealed the c.1121T>C [p.V374A] mutation in KCND3.

Discussion

This report consolidates the pathogenicity of the V374A KCND3 mutation and suggests that the ataxic paroxysmal exacerbations are not a key phenotypic feature of this mutation.

Go to Neurology.org/NG for full disclosures. Funding information is provided at the end of the article.

From the IRCCS Istituto delle Scienze Neurologiche di Bologna (F.P., C.L.M., C.F., L.C., M.L.V., V.C.), Programma di Neurogenetica, Italy; IRCCS Istituto delle Scienze Neurologiche di Bologna (C.L.M., V.D., R.L.), UOC Clinica Neurologica, Italy; and Department of Biomedical and NeuroMotor Sciences (DIBINEM) (M.L.V., R.L., V.C.), University of Bologna, Italy.

The Article Processing Charge was funded by Ricerca Corrente funding.

This is an open access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND), which permits downloading and sharing the work provided it is properly cited. The work cannot be changed in any way or used commercially without permission from the journal.

KCND3 gene encodes the voltage-dependent potassium channel Kv4.3, an alpha subunit of the Shal family of A-type K⁺ channels, essential in membrane repolarization in excitable cells. Heterozygous loss-of-function mutations in KCND3 have been associated with spinocerebellar ataxia type 19 and 22 (SCA19/22), a clinically heterogeneous group of neurodegenerative disorders characterized by variable degrees of cerebellar ataxia, parkinsonism, cognitive dysfunction, epilepsy, and extrapyramidal signs (MIM#607346).¹⁻³ Recently, it has been reported a case with paroxysmal ataxia exacerbations carrying the heterozygous pathogenic variant V374A in the *KCND3* gene.⁴ In this study, we present a second case carrying the same pathogenic variant, confirming this genotype-phenotype association.

Case Report

This male patient, now aged 50 years, with a negative family history, had unstable gait with tremors at the lower limbs and

dysarthric speech since childhood. Our first evaluation was at age 40 years. A neurologic examination showed dysarthria, nystagmus in all directions of gaze, action tremor, dysmetria, weak deep tendon reflexes, truncal ataxia, ataxic gait, and instability in upright position with a positive Romberg sign. He also had bilateral pes cavus. He had marked cerebellar atrophy at brain MRI, more evident at vermis (Figure, A–C). Somatosensory-evoked potential displayed slightly increased time in cortical responses. At the last cognitive evaluation in 2017, the Mini-Mental Status Exam score and the score at the Brief Test of Cognition (IQ at the lower range of the normal values) were normal. At the Brief Neuropsychological Examination test, the patient presented a mild intellectual disability with abnormalities in attentive/executive functions, unchanged compared with the cognitive evaluation in 2010. The abnormalities in executive/praxic functions are likely due to a deficit in planning strategies. EMG, EEG, and ECG findings and audiometry and ophthalmologic examination findings were normal. Polysomnography revealed the presence of moderate sleep apnea syndrome. The patient is overweight (weight: 125

Figure Clinical, Histologic, and Genetic Findings of the Patient



(A-C) Brain MRI shows severe cerebellar atrophy more evident at vermis. (D and E) Muscle biopsy (cytochrome c oxidase/succinic dehydrogenase staining) of the patient performed at the age of 39 years showed scattered cox-negative fibers (red arrows). (F) Chromatogram showing *KCND3* c.1121T>C segregation within the family in available members. Ages represent age at death (for those marked as deceased) or current age. Moreover, major clinical features are also indicated. MI = myocardial infarction.

V374A patients	Present case	II:1 ⁴	l:1 ⁴	
Ataxia	+	+	+	
Dysarthria	+	+	+	
Onset/current age (years)/ sex	6/50/M	25/35/M	46/62/F	
Deep tendon reflexes	Weak	Normal	Normal	
Cognitive impairment	Mild (verbal logic and constructive apraxia)	+	+	
Nystagmus	+	+	+	
Other features	Moderate obstructive sleep apnea syndrome, action tremor, dysmetria, bilateral pes cavus	Paroxysmal ataxia exacerbation; lower limb hypertonus	Mild hand posturing	
EEG	Normal	Slow activity in temporoparietal regions	Moderate diffuse slow activity	
EMG	Normal			
ECG	Normal	Normal	Normal	
Brain MRI	Severe cerebellar atrophy (vermis)	Severe cerebellar atrophy (vermis and superior part of the cerebellar hemispheres)	Mild atrophy in the cerebellum (vermis and superior cerebellar peduncles)	

Table 1 Clinical Features of Patients With KCND3 V374A Mutation

kg). Initially investigated for mitochondrial disease, lactic acid was normal, whereas 3 cytochrome c oxidase (COX)negative fibers were noted at muscle biopsy (Figure, D and E). A complete sequencing of mitochondrial DNA (mtDNA) extracted from skeletal muscle did not show any pathogenic variant (haplogroup U2e2a1c) and pathologic accumulation of macrodeletions. Finally, the mtDNA copy number assessment was normal. The only noticeable result was a relative increase of 7S DNA. Assessment of coenzyme Q in muscle biopsy was also normal.

Genetic investigation was then expanded to exome sequencing, and an in silico panel analysis of genes associated with spinocerebellar ataxia (SCA) revealed the presence of a heterozygous missense variant, c.1121T>C [p.V374A], in *KCND3* (NM_004980.5). This variant has not been reported in the gnomAD database and was predicted to be damaging with a 24.4 CADD-PHRED score. According to the American College of Medical Genetics classification, the c.1121T>C was classified as likely pathogenic with the PM1, PM2, PP2, and PP3 criteria. Segregation analysis was consistent with a possible de novo origin of the V374A variant in the patient because available relatives tested were negative and neurologic disturbances were never reported in the parents and siblings (Figure, F).

Data Availability

Anonymized data not published here will be made available by request from any qualified investigator.

Discussion

We reported an independent case of SCA associated with the V374A variant in *KCND3* gene. The cases originally

Neurology.org/NG

described were a mother and a son (index case) diagnosed with adult-onset slowly progressive cerebellar ataxia, bradyphrenia, and normal general intellectual ability despite low results in cognitive tests.⁴ Both presented cerebellar atrophy, more severe in the index case, with moderate-to-severe cerebellar hypometabolism (Table 1). Notably, the index case experienced paroxysmal ataxia exacerbations responsive to acetazolamide when exposed to accelerations/ decelerations. The V374A pathogenicity was confirmed in vitro: electrophysiology studies showed that the V374A variant was nonfunctional and caused a conductance reduction predicted to generate an increased Purkinje neuron firing frequency.⁴ This family presented an additional A671V variant in the *KCNC3* gene (SCA13), for which a potential synergistic effect was excluded in vitro.

Our patient, differently from those described by Paucar et al., was affected by early-onset cerebellar ataxia, which progressively worsened, apparently without paroxysmal exacerbations. Notably, mild abnormalities in cognitive testing were also observed. The finding of rare COX-negative fibers and increased 7S DNA may be envisaged as secondary reflection on mitochondrial metabolism due to dysfunctional energyconsuming ion channeling. To date, approximately 20 mutations SCA19/22 have been described in patients with heterogeneous clinical presentations, mainly including cerebellar ataxia, cognitive dysfunction, and movement disorders such as parkinsonism (Table 2).

In conclusion, our case consolidates the pathogenicity of this mutation, with a substantially overlapping phenotype except for the paroxysmal exacerbations, for which a synergistic effect of the A671V in *KCNC3* gene cannot be completely excluded, and the onset of the disease.

KCND3 variant	Clinical feature	Inheritance
p.K214R	Episodic gait disorder, vertigo, paraesthesia, pyramidal signs, abnormal ocular movement	AD with incomplete penetrance
p.F227 deletion	Slowly progressive cerebellar ataxia, onset from teenage to middle age; oculomotor abnormalities, pyramidal signs parkinsonism, epilepsy, or cognitive impairment have been reported in some cases	AD, recurrent mutation
p.R293_F295 duplication	Early-onset cerebellar ataxia, intellectual disability, oral apraxia, and epilepsy	De novo mutation
p.S301P	Early onset forms with neurodevelopmental disorder, epilepsy, parkinsonism-dystonia, and ataxia in adulthood	Apparently de novo mutation
p.C317Y	Cerebellar ataxia onset at teenage, developmental delay, intellectual disability, myoclonus, and dystonia	De novo mutation
p.V338E	Adult-onset cerebellar ataxia; cognitive dysfunction	AD
p.G345V	Adult-onset cerebellar ataxia; variable pyramidal signs and oculomotor abnormalities	AD with incomplete penetrance
p.S347W	Adult-onset slowly progressive cerebellar ataxia	Undetermined
р.Т352Р	Mild cerebellar ataxia, cognitive impairment; variable degree of oculomotor disturbance, neuropathy, tremor, and myoclonus	AD
p.I362M	Cerebellar ataxia	AD
p.M365T	Cerebellar ataxia	AD
p.M373L	Adult-onset pure cerebellar ataxia	AD
p.V374A	Progressive cerebellar ataxia and bradyphrenia, cognitive impairment, paroxysmal ataxia exacerbations Cerebellar ataxia, dysarthria, and mild cognitive impairment	AD Apparently de novo mutation
p.P375S	Teenage-onset or adult-onset cerebellar ataxia; cognitive dysfunction, dystonia, and bradykinesia	AD
p.T377M	Adolescent-onset or adult-onset cerebellar ataxia; cognitive impairment in some patients	Recurrent mutation
p.G384S	Cerebellar ataxia, intellectual disability, dystonia, and myoclonus	De novo mutation
p.S390N	Teenage-onset or adult-onset cerebellar ataxia; cognitive dysfunction in some patients	Recurrent mutation
p.V392I	Cerebellar ataxia, intellectual disability, epilepsy, early repolarization syndrome, and paroxysmal atrial fibrillation	Undetermined
p.R419H	Slowly progressive cerebellar ataxia, parkinsonism, and cognitive dysfunction	Sporadic case
p.R431C	Episodic ataxia	Sporadic case
p.L450F	Late-onset cerebellar ataxia and pyramidal signs	AD
p.P633S	Late-onset cerebellar ataxia, decreased reflexes, and vibration sense	Sporadic case

Table 2 Clinical Features and Inheritance of Patients With KCND3 Mutations

Adapted from Hsiao et al.³

Study Funding

Supported by the "Ricerca Corrente" funding (F.P., C.F., L.C. and V.C.), from the Italian Ministry of Health.

Disclosure

F. Palombo reports no disclosures. C. La Morgia reports Consultancies for Chiesi Farmaceutici, Regulatory Pharma Net, and Thenewway srl; speaker honoraria from Santhera Pharmaceuticals, Chiesi Farmaceutici, Regulatory Pharma Net, Thenewway srl, First Class srl, and Biologix; and PI/SI for clinical trials sponsored by GenSight Biologics and Santhera. C. Fiorini, L. Caporali, M.L. Valentino, and V. Donadio report no disclosures. R. Liguori acts as a scientific consultant in boards of Argenx BV, Alexion Pharma Italy s.r.l., and UCB Pharma S.p.A. and received speaker honoraria from Amicus Therapeutics s.r.l. and Editree s.r.l. V. Carelli acts as a scientific consultant in boards of GenSight Biologics, Stealth BioTherapeutics, Santhera Pharmaceuticals, and Chiesi and received speaker honoraria from Chiesi and an unrestricted research grant from Stealth BioTherapeutics. Go to Neurology.org/NG for full disclosure.

Publication History

Received by Neurology: Genetics January 24, 2022. Accepted in final form April 13, 2022. Submitted and externally peer reviewed. The handling editor was Stefan M. Pulst, MD, Dr med.

Appendix Authors

Name	Location	Contribution
Flavia Palombo, PhD	IRCCS Istituto delle Scienze Neurologiche di Bologna, Programma di Neurogenetica, Italy	Analyzed NGS data, performed molecular studies, and drafted the article
Chiara La Morgia, MD, PhD	IRCCS Istituto delle Scienze Neurologiche di Bologna, UOC Clinica Neurologica, Italy	Performed clinical assessment and drafted the article
Claudio Fiorini, PhD	IRCCS Istituto delle Scienze Neurologiche di Bologna, Programma di Neurogenetica, Italy	Performed wet phase of NGS
Leonardo Caporali, PhD	IRCCS Istituto delle Scienze Neurologiche di Bologna, Programma di Neurogenetica, Italy	Performed mitochondrial studies
Maria Lucia Valentino, MD	IRCCS Istituto delle Scienze Neurologiche di Bologna, UOC Clinica Neurologica, Italy Department of Biomedical and NeuroMotor Sciences (DIBINEM), University of Bologna, Italy	Performed and analyzed muscle biopsy and critical revision of the article
Vincenzo Donadio, MD, PhD	IRCCS Istituto delle Scienze Neurologiche di Bologna, UOC Clinica Neurologica, Italy	Performed clinical assessment and critical revision of the article

Appendix (continued) Contribution Name Location IRCCS Istituto delle Scienze Supervised the study and Rocco Neurologiche di Bologna, UOC Liguori, MD critical revision of the Clinica Neurologica, Italy article Department of Biomedical and NeuroMotor Sciences (DIBINEM), University of Bologna, Italy Valerio IRCCS Istituto delle Scienze Study design, supervised Carelli, MD, Neurologiche di Bologna, the study, and critical PhD Programma di Neurogenetica, revision of the article Italy Department of Biomedical and

NeuroMotor Sciences (DIBINEM), University of Bologna, Italy

References

- Lee YC, Durr A, Majczenko K, et al. Mutations in KCND3 cause spinocerebellar ataxia type 22. Ann Neurol 2012;72(6):859-869.
- Duarri A, Jezierska J, Fokkens M, et al. Mutations in potassium channel kcnd3 cause spinocerebellar ataxia type 19. Ann Neurol 2012;72(6):870-880.
- Hsiao CT, Tropea TF, Fu SJ, et al. Rare gain-of-function KCND3 variant associated with cerebellar ataxia, parkinsonism, cognitive dysfunction, and brain iron accumulation. Int J Mol Sci 2021;22(15):8247.
- Paucar M, Ågren R, Li T, et al. V374A KCND3 pathogenic variant associated with paroxysmal ataxia exacerbations. *Neurol Genet* 2021;7(1):e546. doi: 10.1212/NXG. 00000000000546.