

Adult-Onset Alexander Disease: New Causal Sequence Variant in the GFAP Gene

Tsepo Goertler, Dr med, Letizia Zanetti, MSc, Maria Regoni, MSc, Karl Egger, Dr med, Elias Kellner, Dr med, Cornelius Deuschl, Dr med, Christoph Kleinschnitz, Prof Dr med, Jenny Sassone, PhD, and Stephan Klebe, Prof Dr med

Correspondence

Dr. Klebe
stephan.klebe@uk-essen.de

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Abstract

Objectives

Alexander disease (AD) is a rare disorder of the CNS. Diagnosis is based on clinical symptoms, typical MRI findings, and mutations in the glial fibrillary acid protein (GFAP) gene. In this case study, we describe a new mutation (p.L58P) in *GFAP* that caused a phenotype of adult-onset AD (AOAD).

Methods

In our outpatient clinic, a patient presented with cerebellar and bulbar symptoms after brain concussion. We used MRI and performed next-generation exome sequencing (NGS) to find mutations in *GFAP* to diagnose AD. The mutation was then transfected into HeLa cell lines to prove its pathogenicity.

Results

The brain MRI finding showed typical AD alterations. The NGS found a heterozygous variant of unknown significance in *GFAP* (c.173T>C; p.L58P). After transfecting HeLa cell lines with this mutation, we showed that GFAP-L58P formed pathogenic clusters of cytoplasmic aggregates.

Discussion

We have found a new mutation that causes AOAD. We recommend that AOAD is included in the diagnostic workup in adult patients with gait ataxia and cerebellar and bulbar symptoms in association with a traumatic head injury.

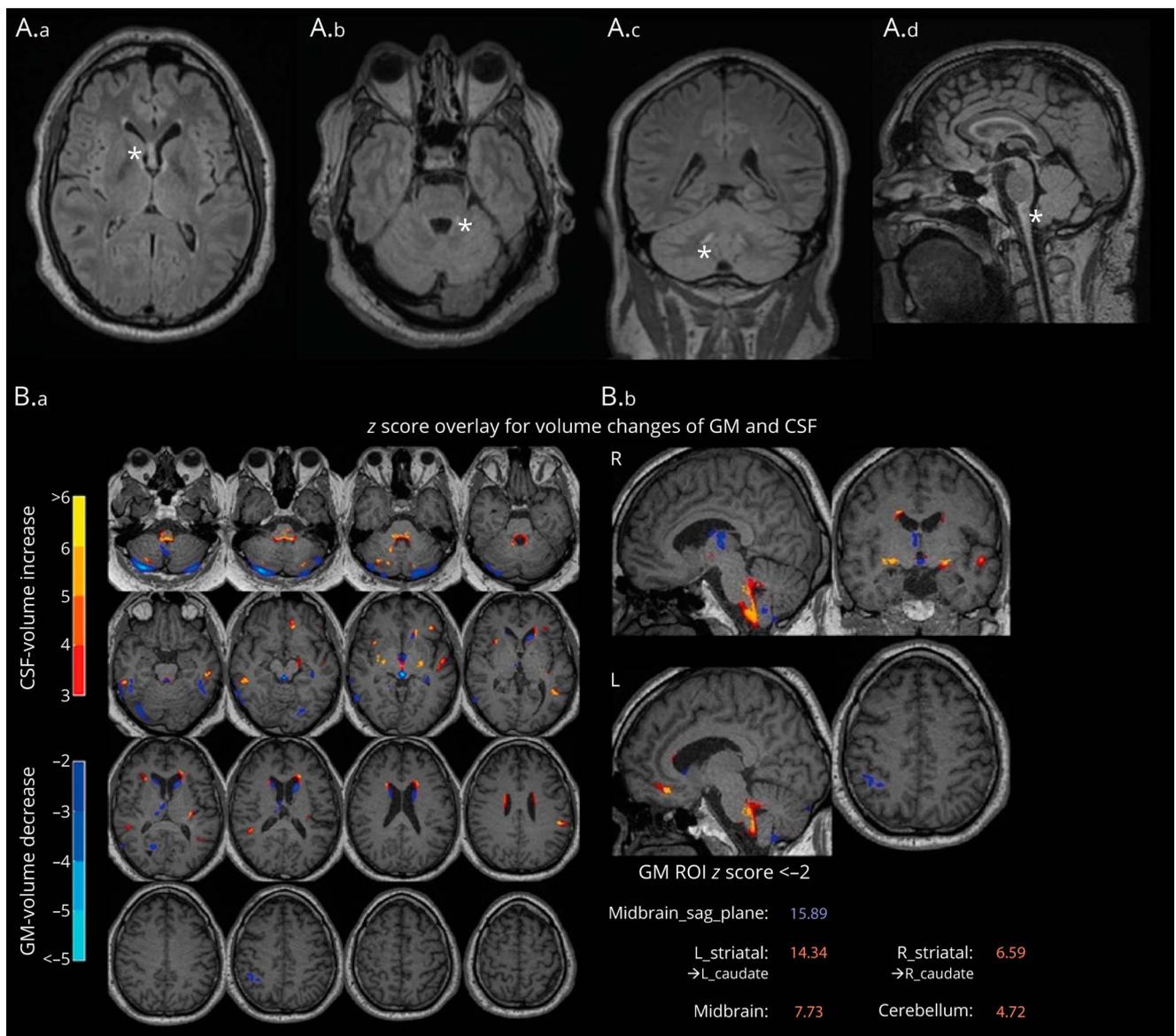
From the Department of Neurology (T.G., C.K., S.K.), Essen University Hospital, Germany; Division of Neuroscience (L.Z., M.R., J.S.), San Raffaele Scientific Institute; Vita-Salute San Raffaele University (L.Z., M.R., J.S.), Milan, Italy; Department of Radiology (K.E.), Tauernklinikum Zell am See, Academic Teaching Hospital of the Paracelsus University Salzburg, and Medical University of Vienna, Austria; Department of MR Physics (E.K.), Medical Center, University of Freiburg, Faculty of Medicine, University of Freiburg, Germany; and Institute of Diagnostic and Interventional Radiology and Neuroradiology (C.D.), University Hospital Essen, Germany.

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Figure 2 MRI and Automated Brain Volumetry Analysis of the Patient



(A.a–d) Brain MRI T2-weighted fluid-attenuated inversion recovery sequences in the axial plane (A.a, A.b), coronal plane (A.c) and sagittal plane (A.d) showing hyperintensities periventricular (*) (A.a), in the left cerebellar peduncle (*) (A.b), and dentate nuclei (*) (A.c). The typical tadpole sign signaling spinal cord atrophy is shown in (*) (A.d). (B.a–b) Automated brain volumetry analysis using the software VEOmorph (VEObrian GmbH). In the results of the automated, combined voxel and region (CVR)-based whole-brain volumetry using 3-dimensional T1-weighted MR images, regional volume increase in CSF and regional volume decrease in gray matter (GM) are superimposed in red to yellow and in light blue to dark blue (depending on the according z score) onto the patient's individual brain MRI in the transverse, sagittal, and coronal orientation (B.a). Besides the visually detectable atrophy of the medulla oblongata, additional volume abnormality is shown in the midbrain, cerebellum, and striatal regions of both hemispheres (B.b). Abnormal areas are defined based on a volume change of at least 2 SDs (z score 2) in comparison with an age-matched and sex-matched healthy control group. Egger K. *Neuroimage Clin.* 2018; doi: 10.1016/j.nicl.2018.09.013.

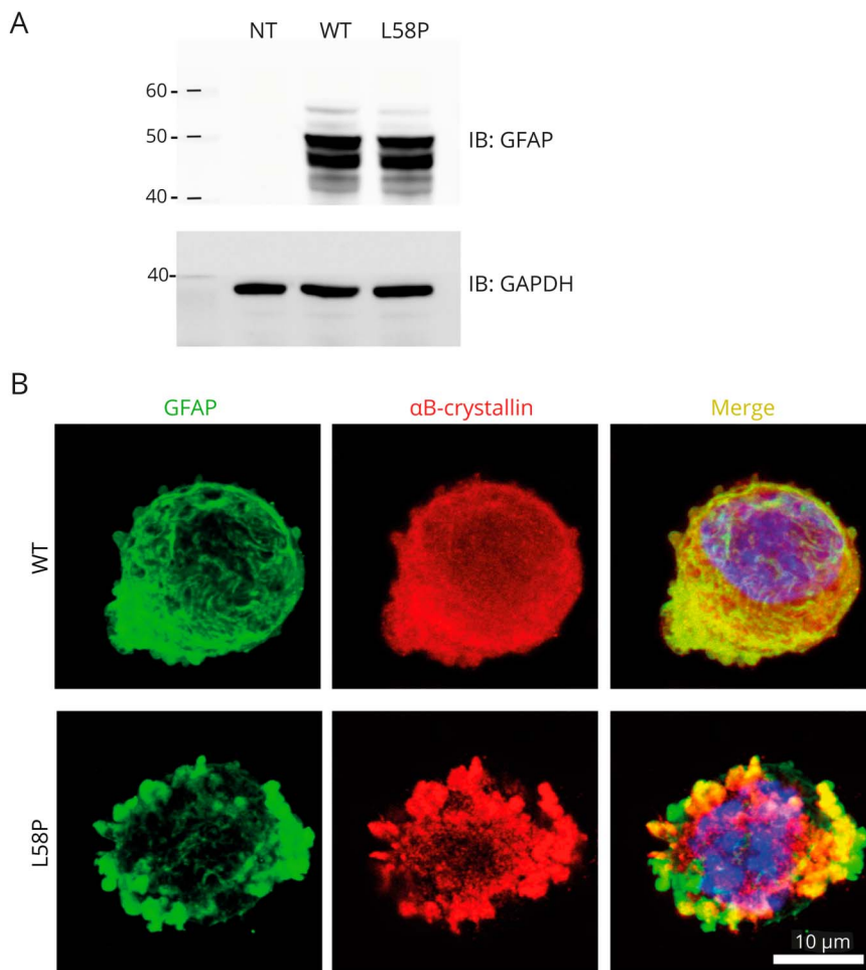
or *GFAP-L58P*. Forty-eight hours after transfection, cell lysates were analyzed by Western blot to demonstrate the plasmid expression (Figure 3A), and cell monolayers were immunostained with an antibody against GFAP. Fluorescence images showed that wild-type GFAP assembled into bundled filaments that extended throughout the cytoplasm, whereas *GFAP-L58P* formed clusters of cytoplasmic aggregates (Figure 3B). Because cytoplasmic inclusions within astrocytes of patients with AD also contain the chaperones α B-crystallin, we costained the cells with an antibody against α B-crystallin. The GFAP containing aggregates in the cells

transfected with *GFAP-L58P* gene were positive for α B-crystallin. Written informed consent was obtained from the patient.

Discussion

Owing to the primarily nonspecific clinical symptoms of cerebellar ataxia, bulbar symptoms, and positive pyramidal signs, only the brain MRI led to the suspected diagnosis of AOAD. Using NGS, the heterozygote variant (c.173T>C;

Figure 3 Western Blot and Immunocytochemistry of the Transfected HeLa Cell Lines



(A) Representative Western blot performed in lysates from HeLa cells transfected with plasmids encoding human GFAP-WT or mutant GFAP-L58P (GFAP: mAb #3670 cell signaling 1:1000, GAPDH: sc-25778 Santa Cruz, 1:1000). (B) Representative confocal images showing HeLa cells transiently transfected with plasmids encoding human wild-type GFAP or mutant GFAP-L58P and labeled with GFAP antibody (mAb #3670 cell signaling 1:300, green fluorescence). The image shows that wild-type GFAP assembled in filament networks, whereas mutant L58P formed dot-like aggregates. Cells were costained with α β -crystallin antibody (sc-137129 Santa Cruz 1:100, red fluorescence) in cells transfected with mutant GFAP-L58P, and α β -crystallin formed dot-like aggregates that colocalized with the GFAP signal. Images are representative of 50 analyzed cells from 3 independent experiments. DAPI is indicated in blue in the merge images on the right.

p.L58P) in *GFAP* was found and categorized as a variant of unknown significance. In vitro experiments demonstrated that this variant represented a novel mutation that affected the formation of the intermediate filament network and confirmed the diagnosis of AOAD.

Our patient stated that he experienced gait ataxia, clumsiness, and vertigo after minor brain concussion due to an accident. A correlation between AOAD and traumatic head injuries has been described before, with a latency between trauma and symptom onset of up to 10 years.⁷⁻⁹ Considering that severe symptoms may appear many years later, these incidences might be underrated. Similar to dystonia, we hypothesize a second-hit theory in the emergence of AOAD.

AD must be considered as a differential diagnosis in adult patients with new ataxia, bulbar symptoms, and leukodystrophy and the tadpole sign in brain MRI. Furthermore, anamnestic hints for traumatic head injuries exposing the disease onset must be taken into account. The second-hit theory is an interesting concept in the emergence of AOAD that should be considered in upcoming research.

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Disclosure

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Appendix Authors

Name	Location	Contribution
Tsepo Goerttler, Dr med	Department of Neurology, Essen University Hospital, Germany	Drafting/revision of the article for content, including medical writing for content; major role in the acquisition of data; study concept or design; and analysis or interpretation of data
Letizia Zanetti, MSc	Division of Neuroscience, San Raffaele Scientific Institute; Vita-Salute San Raffaele University, Milan, Italy	Analysis or interpretation of data
Maria Regoni, MSc	Division of Neuroscience, San Raffaele Scientific Institute; Vita-Salute San Raffaele University, Milan, Italy	Analysis or interpretation of data
Karl Egger, Dr med	Department of Radiology, Tauernklinikum Zell am See, Academic Teaching Hospital of the Paracelsus University Salzburg, and Medical University of Vienna, Austria	Analysis or interpretation of data
Elias Kellner, Dr med	Department of MR Physics, Medical Center, University of Freiburg, Faculty of Medicine, University of Freiburg, Germany	Analysis or interpretation of data
Cornelius Deuschl, Dr med	Institute of Diagnostic and Interventional Radiology and Neuroradiology, University Hospital Essen, Germany	Drafting/revision of the article for content, including medical writing for content; and analysis or interpretation of data
Christoph Kleinschnitz, Prof Dr med	Department of Neurology, Essen University Hospital, Germany	Drafting/revision of the article for content, including medical writing for content

Appendix (continued)

Name	Location	Contribution
Jenny Sassone, PhD	Division of Neuroscience, San Raffaele Scientific Institute; Vita-Salute San Raffaele University, Milan, Italy	Drafting/revision of the article for content, including medical writing for content; analysis or interpretation of data
Stephan Klebe, Prof Dr med	Department of Neurology, Essen University Hospital, Germany	Drafting/revision of the article for content, including medical writing for content; major role in the acquisition of data; study concept or design; and analysis or interpretation of data

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