Genotypes and phenotypes of patients with Lafora disease living in Germany

David Brenner1,*, Tobias Baumgartner2, Sarah von Spiczak3, Jan Lewerenz1, Roger Weis4, Anja Grimm5, Petra Gaspiro6, Claudia D. Wurster1, Wolfram S. Kunz1,7, Jan Wagner1, Berge A. Minassian8, Christian E. Elger2, Albert C. Ludolph1, Saskia Biskup9, Dennis Döcker3
1Department of Neurology, University of Ulm, Ulm, Germany.
2Department of Epileptology, University of Bonn Hospital, Bonn, Germany.
3DRK-Northern German Epilepsy Center for Children and Adolescents, Schwentinental, Raisdorf, Germany.
4Neuropediatric Center, Rheinhessen-Fachklinik Mainz, Mainz, Germany.
5Epilepsie-Zentrum Berlin-Brandenburg, Berlin, Germany.
6Klinikum St. Marien Amberg, Amberg, Germany.
7Institute of Experimental Epileptology and Cognition Research, University Bonn, Bonn, Germany.
8Department of Pediatrics, University of Texas Southwestern, Dallas, USA.

This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.

*Correspondence: david.brenner@uni-ulm.de.

Authors’ contributions
DB designed the study, examined patients and documented medical histories, analysed the data and drafted the manuscript. TB, SvS, JL, PG, JF, BAM, CEE, ACL and DD examined patients, analysed the data, documented medical histories and drafted the manuscript. RW, AG and CW examined patients and documented medical histories. WSK and SB documented medical histories and analysed the data. All authors read and approved the final manuscript.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Additional file
Additional file 1: Table S1. Anticonvulsive pharmacotherapy of LD patients living in Germany. (DOCX 16 kb)

Availability of data and materials
The datasets used and analysed during the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
This is a case series with informed consent by the patients or their parents, respectively. A separate ethics approval has not been considered necessary by the ethics committee of the University of Ulm.

Consent for publication
All patients or parents, respectively, gave their consent for publication.

Competing interests
B.A. Minassian holds patents for diagnostic testing of the following genes: EPM2A, EPM2B, MECP2, and VMA2I; and has received license fee payments/royalty payments from patents for diagnostic testing of the following genes: EPM2A, EPM2B, MECP2, and VMA2I.

The other authors declare that they have no competing interests.
Background: Lafora progressive myoclonus epilepsy (Lafora disease) is a rare, usually childhood-onset, fatal neurodegenerative disease caused by biallelic mutations in *EPM2A* (Laforin) or *EPM2B* (*NHLRC1*, Malin). The epidemiology of Lafora disease in Germany is largely unknown. The objective of this retrospective case series is to characterize the genotypes and phenotypes of patients with Lafora disease living in Germany.

Methods: The patients described in this case series initially had the suspected clinical diagnosis of Lafora disease, or unclassified progressive myoclonus epilepsy. Molecular genetic diagnostics including next generation sequencing-based diagnostic panel analysis or whole exome sequencing was performed.

Results: The parents of four out of the 11 patients are nonconsanguineous and of German origin while the other patients had consanguineous parents. Various variants were found in *EPM2A* (six patients) and in *EPM2B* (five patients). Eight variants have not been reported in the literature so far. The patients bearing novel variants had typical disease onset during adolescence and show classical disease courses.

Conclusions: This is the first larger case series of Lafora patients in Germany. Our data enable an approximation of the prevalence of manifest Lafora disease in Germany to 1.69 per 10 million people. Broader application of gene panel or whole-exome diagnostics helps clarifying unclassified progressive myoclonus epilepsy and establish an early diagnosis, which will be even more important as causal therapy approaches have been developed and are soon to be tested in a phase I study.

Introduction

Lafora disease (LD; EPM2A/B; OMIM #254780) is a severe form of progressive myoclonus epilepsy inherited in an autosomal recessive mode and caused by biallelic mutations in *EPM2A* (Laforin) or *EPM2B* (*NHLRC1*, Malin) [1, 2]. Pathogenic homozygous or compound heterozygous mutations in both genes cause cytoplasmatic precipitation, aggregation and accumulation of neurotoxic poorly branched and insoluble glycogen forming polyglucosan inclusions (so-called Lafora bodies) leading to progressive neurodegeneration. In most cases, the disease starts with epileptic seizures in late childhood or adolescence. Apart from multiple types of seizures (tonic-clonic, myoclonic, absence, atonic, or visual), LD patients develop progressive cerebellar ataxia, dysarthria, dementia and neuropsychiatric symptoms. LD usually leads to death within 10 years after symptom onset. The prevalence is below 1–9 in 10^6, depending on the population and the frequency of consanguinity [3]. The epidemiology as well as the genotypic and phenotypic spectrum of LD in Germany is largely unknown. At present, therapy is merely symptomatic and palliative. Recent preclinical studies have shown that downregulation of glycogen synthesis prevents LD in mice [4]. Antisense oligonucleotides (ASO) targeting the glycogen synthase mRNA significantly reduce Lafora body load in LD mice (unpublished observation). Consequently, an ASO targeting the human glycogen synthase mRNA is being developed.
towards an upcoming clinical trial. A two-year natural history study prerequisite for this therapy trial is currently ongoing (NCT03876522).

Here, we describe the genotypes and phenotypes of 11 index patients with LD living in Germany.

Methods

DNA analysis

Molecular diagnostics was performed as part of the clinical follow-up in all cases. DNA was extracted from EDTA blood samples according to standard procedures. Next generation sequencing (NGS)-based diagnostic panel analysis or whole exome sequencing was performed in 11 index patients with suspected clinical diagnosis of Lafora disease, or unclassified progressive myoclonus epilepsy. Annotation of reported variants is based on human reference genome *Homo sapiens* GRCh37 (hg19). All patients or parents, respectively, gave their consent to publication of their cases.

Results

Four out of the 11 patients are of German origin. Variants in *EPM2A* or *EPM2B* were detected by NGS in all 11 index patients with suspected LD or unclassified progressive myoclonus epilepsy. Variants of various types were detected in *EPM2A* in six patients (three of them of German origin) and in *EPM2B* in five patients (one of them of German origin; see Table 1). Five *EPM2A* variants (c.259A > G, c.290 T > G, c.759delinsCATGCA, c.836G > T, c.917A > T) and three *EPM2B* variants (c.385C > T, c.583del, and c.730delG) have not been reported in the LD literature or in the Lafora gene mutation database (http://projects.tcag.ca/lafora/) before [5]. The mean age of disease onset is 13.7 ± 0.6 years (mean ± SEM) in the patients bearing *EPM2A* variants and 14.6 ± 2.3 years in those carrying *EPM2B* variants. The cumulative mean age of onset of this cohort is 14.1 ± 1.0 years (range: 8–21 years).

In the following paragraphs, we shortly outline the disease courses of the patients bearing novel variants (also see Table 2). The anticonvulsants valproate (11/12), perampanel (8/12), levetiracetam/brivaracetam (7/12), and clobazam (6/12) were the most used ones in our cohort indicating a good of effectiveness of these drugs in LD (see Additional file 1: Table S1).

Case vignettes of index patients with novel variants (Table 2)

**Patient nr. 1** is a homozygous carrier of the *EPM2A* variant c.259A > G. Her parents are of Turkish origin and are consanguineous. Retrospectively, the first symptom were recurrent episodes of explosive headache at the age of 12 years. Next, myoclonic and atonic seizures with consecutive falls developed. Aged 15 years, she developed tonic-clonic, visual, gelastic and absence seizures and (negative) myoclonus aggravated. Shortly later, she also developed severe ataxia and progressive cognitive decline. At the age of 19 years, the patient is wheelchair-bound and exhibits severe dysarthria as well as dysphagia requiring percutaneous endoscopic gastrostomy (PEG).
Patient nr. 2 is compound heterozygous for the known variant c.269_275del and the novel variant c.917A > T in EPM2A. Her parents are German and nonconsanguineous. She developed myoclonic seizures as first disease symptom at the age of 15 years. However, an EEG at the age of 6 years, performed due to headaches showed posterior slow waves, which were interpreted as a variant of the basic rhythm at that time. Since the beginning of the epilepsy, the young woman lost major cognitive and motor functions and suffers from ataxia. She developed polyneuropathy of the lower extremities. About 3 years later, she can walk only few steps with lots of support and needs help with all activities of daily living.

Patient nr. 3 is a homozygous carrier of the EPM2A variant c.290 T > G. She is of German origin and her parents are considered nonconsanguineous based on family history. She had onset of tonic-clonic seizures at the age of 14 years. Shortly later, she developed myoclonic and absence seizures, ataxia and cognitive decline. She exhibits a slow progression and is still able to walk and speak 12 years after onset of symptoms.

Patient nr. 5 is homozygous for the EPM2A variant c.759delinsCATGCA. His parents are Russian and are consanguineous. He developed tonic-clonic seizures at the age of 15 years. The patient showed striking social difficulties and increased aggressiveness. After the age of 17 years, psychomotor skills reduced, and the patient developed tremor, ataxia and progressive cognitive decline. At the age of 20, he has lost many everyday skills and suffers from frequent seizures, while refusing medication. Due to severe dysphagia a PEG was implanted. At the age of 23 years he is wheelchair-bound and shows severe dysarthria.

Patient nr. 6 carries the homozygous c.836G > T variant in EPM2A. He is of German origin and his parents are considered nonconsanguineous based on family history. Myoclonic seizures started at the age of 16. About the same time, visual auras occurred followed by tonic-clonic and absence seizures about 1 year later. Thereafter, he developed mild ataxia and a cognitive decline. At the age of 24, daily drop attacks and myoclonic seizures were the main problem of the patient. His sister carried the same homozygous mutation in EPM2A, but her course of the disease was more severe and she died at the age of 24.

Patient nr. 7 carries the homozygous c.385C > T variant in EPM2B. She is of Syrian origin and her parents are consanguineous. She suffered the first and second tonic-clonic seizure at the age of 11 and 13 years. Thereafter, the frequency of seizures increased and since the age of 15, she has been suffering from intermittent myoclonic seizures and progressive dysarthria, ataxia, optic hallucinations and dementia. At the current age of 19 years, the patient is wheelchair-bound and suffers from severe dysphagia requiring PEG.

Patient nr. 10 is compound heterozygous for the known variant c.436G > A and the novel variant c.730delG in EPM2B. She is German and her parents are nonconsanguineous. She first developed tonic-clonic and absence seizures at the age of 16. About 1 year later, myoclonic seizures started. A juvenile myoclonic epilepsy was suspected. After initiating a therapy with Valproate, tonic-clonic seizures were well under control for about 4 years. Meanwhile, myoclonic seizures accelerated under different medications and the EEG worsened. At the age of 23, the patient’s cognitive skills are still preserved, and he does not suffer from ataxia.
**Patient nr. 11** is homozygous for the c.583del variant in *EPM2B*. He is Iraqi and his parents are consanguineous. His epilepsy started at the age of 9 years with generalized tonic-clonic seizures. Later, he developed myoclonic seizures (especially negative myoclonus), psychogenic seizures and cognitive decline. His motor abilities are relatively well preserved (now aged 14).

**Discussion**

This is the first larger case series of Lafora patients in Germany. Consistent with consanguinity being uncommon in Germany and the very rare prevalence of this recessive disease, only four out of the 11 patients are of German origin. In our cohort of LD patients, variants in *EPM2A* and *EPM2B* show a similar frequency. We report eight novel variants in *EPM2A* and *EPM2B*. Six novel variants are located maximally three nucleotides away from aforesaided pathogenic variants [5]. While the parents of the seven Non-German patients were consanguineous, those of the four German patients were nonconsanguineous. Although not verifiable by family history, the fact that two out of the four German patients bear homozygous variants should indicate that there may be a common ancestor. The patients bearing novel variants in *EPM2A* or *EPM2B* had typical disease onset during adolescence and developed characteristic symptoms including multiple types of seizures, ataxia, dysarthria and dysphagia, as well as cognitive decline during the disease course. The cumulative mean age of onset of this LD patient cohort of 14 ± 1 years coincides with the reported peak age of onset [3]. Consistent with previous reports [6], the three patients bearing the known *EPM2B* c.436G > A variant (patients nr. 8–10) had a late onset of symptoms and show a comparatively slow disease progression. Remarkably, the 35-year old sister of patient nr. 8 who is also homozygous for the c.436G > A variant (the genetic result was confirmed twice by independent laboratories) is asymptomatic to date. Noteworthy, patient nr. 8 bearing the *EPM2B* c.436G > A variant developed severe diabetes mellitus with diabetic coma and cataract, a finding that recently has also been described in a LD patient with an *EPM2B* c.386C > A variant [9]. Further, patient nr. 2 bearing the c.[269_275del]; [917A > T] variants in *EPM2A* developed a polynuropathy of the lower limbs. Although a causal link remains speculative, polyglucosan formation in axons indeed causes severe axonopathy in patients with adult polyglucosan body disease [10]. Based on the data of this report and the knowledge about two additional LD patients not included here the estimated prevalence of manifest LD in Germany is 14 per 83,019,000 [11] or 1.69 per 10 million people, respectively. Including the pre-symptomatic mutation carriers known to us the prevalence raises to 2.05 per 10 million people.

**Conclusions**

This is the first larger case series of Lafora patients in Germany. Our data enable an approximation of the prevalence of manifest Lafora disease in Germany to 1.69 per 10 million people. Early genetic testing is central when suspecting this condition given that causal therapy approaches have been developed and are soon to be tested in a phase I study. Broader application of gene panel or whole-exome diagnostics helps clarifying unclassified PME and establish an early diagnosis.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements

The authors thank the families and patients for their participation to this study.

Funding

Dr. Brenner’s contribution was supported by the B. Braun Foundation, Germany, under grant number BBST-D-18-00064. Dr. Minassian’s contribution was supported by the National Institute of Neurological Disorders and Stroke of the NIH under award number P01 NS097197.

Abbreviations

| Abbreviation | Description                        |
|--------------|------------------------------------|
| ASO          | Antisense oligonucleotide          |
| LD           | Lafora disease                     |
| NGS          | Next generation sequencing         |
| PEG          | Percutaneous endoscopic gastrostomy|
| PME          | Progressive myoclonus epilepsy     |

References

1. Minassian BA, Lee JR, Herbrick J-A, et al. (1998). Mutations in a gene encoding a novel protein tyrosine phosphatase cause progressive myoclonus epilepsy. Nature Genetics, 20, 171–174. [PubMed: 9771710]
2. Chan EM, Young EJ, lanzano L, et al. (2003). Mutations in NHLRC1 cause progressive myoclonus epilepsy. Nature Genetics, 35, 125–127. [PubMed: 12958597]
3. Turnbull J, Tiberia E, Striano P, et al. (2016). Lafora disease. Epileptic Disorders, 18, 38–62. [PubMed: 27702709]
4. Pederson BA, Turnbull J, Epp JR, et al. (2013). Inhibiting glycogen synthesis prevents Lafora disease in a mouse model. Annals of Neurology, 74, 297–300. [PubMed: 23913475]
5. lanzano L, Zhang J, Chan EM, et al. (2005). Lafora progressive myoclonus epilepsy mutation database-EPM2A and NHLRC1 (EMP2B) genes Human Mutation, 26, 397–397. Wiley.
6. Gómez-Garre P, Sanz Y, Rodríguez De Córdoba SR, & Serratosa JM (2000). Mutational spectrum of the EPM2A gene in progressive myoclonus epilepsy of Lafora: High degree of allelic heterogeneity and prevalence of deletions. European Journal of Human Genetics, 8, 946–954. [PubMed: 11175283]
7. Lanoiselée H-M, Genton P, Lesca G, Brault F, & De Toffol B (2014). Are c. 436G>A mutations less severe forms of Lafora disease? A case report Epilepsy & Behavior Case Reports, 2, 19–21. Elsevier. [PubMed: 25667860]
8. Ferlazzo E, Canafoglia L, Michelucci R, et al. (2014). Mild Lafora disease: Clinical, neurophysiologic, and genetic findings. Epilepsia, 55, e129–e133. [PubMed: 25270369]
9. Nicolescu RC, Al-Khawaga S, Minassian BA, & Hussain K (2019). Diabetes mellitus in a patient with Lafora disease: Possible links with pancreatic β-cell dysfunction and insulin resistance. Frontiers in Pediatrics, 6, 424. [PubMed: 30701169]
10. Robitaille Y, Carpenter S, Karpati G, & DiMauro SD (1980). A distinct form of adult polyglucosan body disease with massive involvement of central and peripheral neuronal processes and astrocytes: A report of four cases and a review of the occurrence of polyglucosan bodies in other conditions such as Lafora’s disease and normal ageing. Brain, 103, 315–336. [PubMed: 6249438]

Neur Res Pract. Author manuscript; available in PMC 2020 June 25.
11. Bevölkerungsstand 12/2018 - Statistisches Bundesamt. https://www.destatis.de/DE/Themen/Gesellschaft-Umwelt/Bevoelkerung/Bevoelkerungsstand/Tabellen/zensus-geschlecht-staatsangehoerigkeit-2018.html. Accessed 19 July 2019.
## Table 1

### Genotypes of LD patients living in Germany

| P. nr. | Variant | Alleles | Ref. |
|--------|---------|---------|------|
| **EPM2A** | NC_000006.11; NM_005670.3; | | |
| 1<sup>a</sup> | g.146056376T > C | c.259A > G | p.(Lys87Glu) | Homo. |
| 2<sup>a</sup> | g.146056360_146056366del; [145,948,631 T > A] | c.[269_275del];[917A > T] | p.[(Lys90Ser fs*35);(Asp306Val)] | Comp. het. | [6] |
| 3<sup>a</sup> | g.146056345A > C | c.290 T > G | p.(Leu97Arg) | Homo. |
| 4 | g.146007412G > A | c.322C > T | p.(Arg108Cys) | Homo. | [1] |
| 5<sup>a</sup> | g.145948789delinsTGCATG | c.759delinsCATGCA | p.(Ala258Met/fs*33) | Homo. |
| 6<sup>a</sup> | g.145948712C > A | c.836G > T | p.(Gly279Val) | Homo. |
| **EPM2B** | NC_000006.11; NM_198586.2; | | |
| 7<sup>a</sup> | g.18122453G > A | c.385C > T | p.(Pro129Ser) | Homo. |
| 8, 9 | g.18122402C > T | c.436G > A | p.Asp146Asn | Homo. | [2, 7] |
| 10<sup>a</sup> | g.18122402C > T; [18122108delC] | c.[436G > A];[730delG] | p.[(Asp146Asn);(Val244Ser fs*51)] | Comp. het. | [2, 7] |
| 11<sup>a</sup> | g.18122255del | c.583del | p.(Asp195Ile/fs*37) | Homo. |

Patients with novel variants are indicated by a superscripted letter (<sup>a</sup>). Annotation based on human reference genome *Homo sapiens* GRCh37 (hg19)
| Patient number | Nationality | Gender | Consanguinity | Variant | Age of onset | Current age | Current disease stage | Special features |
|----------------|-------------|--------|---------------|---------|--------------|-------------|-----------------------|-----------------|
| 1<sup>a</sup>   | Tur         | F      | yes           | EMP2A   | c.259A > G (homo.) | 14          | 15                    | Gelastic seizures |
| 2<sup>a</sup>   | Ger         | F      | no            |         | c.[269_275del]; [917A > T] | 17          | 15                    | Congenital hypothyroidism, polyneuropathy |
| 3<sup>a</sup>   | Ger         | F      | no            | EMP2A   | c.290T > G (homo.)   | 14          | 15                    | Psychogenic seizures |
| 4<sup>a</sup>   | Leb         | F      | yes           |         | c.322C > T (homo.)   | 12          | n/a                   | Vision loss     |
| 4<sup>b</sup>   | Leb         | M      | yes           |         | c.322C > T (homo.)   | 11          | n/a                   | Vision loss     |
| 5<sup>a</sup>   | Rus         | M      | yes           | EMP2B   | c.759delCATGCA      | 14          | 14                    | Aggressiveness  |
| 6<sup>a</sup>   | Ger         | M      | no            | EMP2B   | c.836G > T (homo.)   | 18          | 16                    | Vision loss     |
| 7<sup>a</sup>   | Syr         | F      | yes           |         | c.385C > T (homo.)   | 11          | 15                    | Optic hallucinations, hypothyroidism, hepatomegaly |
| 8               | Tur         | M      | yes           |         | c.436G > A (homo.)   | 17          | 17                    | Diabetes mellitus with coma and cataract, hypothyroidism, arterial hypertension |
| 9               | Tur         | F      | yes           |         | c.436G > A (homo.)   | 23          | 21                    | Vision loss     |
| 10<sup>a</sup>  | Ger         | F      | no            |         | c.[436G > A]; [730delG] | 16          | 17                    | Vision loss     |
| 11<sup>a</sup>  | Ira         | M      | yes           |         | c.583del (homo.)     | 8           | 13                    | Psychogenic seizures |

Patients with novel variants are indicated by a superscripted letter (<sup>a</sup>). Patients 4 <sup>a</sup> and <sup>b</sup> are siblings. *Abbreviations: Ger German, Ira Iraqi, Leb Lebanese, Rus Russian, Syr Syrian, Tur Turkish.* Disease stage score was assessed according to Ferlazzo et al. [8]: (1) mild cognitive and motor impairment, preserved daily living activities, and social interaction; (2) moderate mental decline, limitations in motor activities, and limited social interaction; (3) severe mental and motor impairment, needing help in walking and regular assistance in daily living activity, and poor social interaction; (4) patient wheelchair-bound or bedridden, and no significant daily living activities or social interaction. Consanguinity or non-consanguinity status was based on family history.