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Genetic landscape of pediatric acute liver failure of indeterminate origin

Dominic Lenz¹  | Lea D. Schlieben^{2,3}  | Masaru Shimura^{3,4} |
 Alyssa Bianzano¹ | Dmitrii Smirnov^{2,3}  | Robert Kopajtich^{2,3} |
 Riccardo Berutti^{2,3} | Rüdiger Adam⁵ | Denise Aldrian⁶ | Ivo Baric⁷ |
 Ulrich Baumann⁸ | Neslihan E. Bozbulut⁹ | Melanie Brugger² |
 Theresa Brunet² | Philip Bufler¹⁰ | Birutė Burnytė¹¹ | Pier L. Calvo¹² |
 Ellen Crushell¹³ | Buket Dalgıç⁹ | Anibh M. Das¹⁴ | Antal Dezsőfi¹⁵ |
 Felix Distelmaier¹⁶ | Alexander Fichtner¹ | Peter Freisinger¹⁷ |
 Sven F. Garbade¹ | Harald Gaspar¹⁸ | Louise Goujon¹⁹ | Nedim Hadzic²⁰ |
 Steffen Hartleif²¹ | Bianca Hegen²² | Maja Hempel^{23,24} | Stephan Henning¹⁰ |
 Andre Hoerning²⁵ | Roderick Houwen²⁶ | Joanne Hughes²⁷ | Raffaele Iorio²⁸ |
 Katarzyna Iwanicka-Pronicka²⁹ | Martin Jankofsky²² | Norman Junge⁸ |
 Ino Kanavaki³⁰ | Aydan Kansu³¹ | Sonja Kaspar²⁵ | Simone Kathemann³² |
 Deidre Kelly³³ | Ceyda T. Kırsaçlıoğlu³¹ | Birgit Knoppke³⁴ | Martina Kohl³⁵ |
 Heike Kölbel³⁶ | Stefan Kölker¹ | Vassiliki Konstantopoulou³⁷ |
 Tatiana Krylova³⁸ | Zarife Kuloğlu³¹ | Alice Kuster³⁹ | Martin W. Laass⁴⁰ |
 Elke Lainka³² | Eberhard Lurz⁴¹ | Hanna Mandel⁴² | Katharina Mayerhanser² |
 Johannes A. Mayr⁴³ | Patrick McKiernan⁴⁴ | Patricia McClean⁴⁵ |
 Valerie McLin⁴⁶ | Karine Mention⁴⁷ | Hanna Müller⁴⁸ | Laurent Pasquier¹⁹ |
 Martin Pavlov^{2,3} | Natalia Pechatnikova⁴⁹ | Bianca Peters¹ |
 Danijela Petković Ramadža⁷ | Dorota Piekutowska-Abramczuk²⁹ | Denisa Pilic³² |
 Sanjay Rajwal⁴⁶ | Nathalie Rock⁴⁶ | Agnès Roetig⁵⁰ | René Santer²² |
 Wilfried Schenk⁵¹ | Natalia Semenova³⁸ | Christiane Sokollik⁵² |
 Ekkehard Sturm²¹ | Robert W. Taylor⁵³ | Eva Tschiedel⁵⁴ | Vaidotas Urbonas¹¹ |
 Roser Urreizti⁵⁵ | Jan Vermehren³⁴ | Jerry Vockley⁴⁴ |
 Georg-Friedrich Vogel^{6,56} | Matias Wagner² | Wendy van der Woerd²⁶ |
 Saskia B. Wortmann⁴³ | Ekaterina Zakharova³⁸ | Georg F. Hoffmann¹ |
 Thomas Meitinger² | Kei Murayama⁴ | Christian Stauffer¹  | Holger Prokisch^{2,3} 

¹Heidelberg University, Medical Faculty, University Hospital Heidelberg, Center for Child and Adolescent Medicine, Department I, Division of Pediatric Neurology and Metabolic Medicine, Heidelberg, Germany

²School of Medicine, Institute of Human Genetics, Klinikum rechts der Isar, Technical University of Munich, Munich, Germany

³Institute of Neurogenomics, Computational Health Centre, Helmholtz Munich, Munich Germany

⁴Department of Metabolism, Chiba Children's Hospital, Centre for Medical Genetics, Chiba, Japan

- ⁵Department of Paediatric Gastroenterology, Hepatology and Nutrition, University Children's Hospital, Medical Faculty Mannheim, Heidelberg University, Mannheim, Germany
- ⁶Paediatrics I, Medical University of Innsbruck, Innsbruck, Austria
- ⁷Department of Paediatrics, University Hospital Centre Zagreb, University of Zagreb, School of Medicine, Zagreb, Croatia
- ⁸Department of Paediatric Kidney, Liver, and Metabolic Diseases, Division for Paediatric Gastroenterology and Hepatology, Hannover Medical School, Hannover, Germany
- ⁹Department of Paediatric Gastroenterology, Gazi University Faculty of Medicine, Ankara, Turkey
- ¹⁰Department of Paediatric Gastroenterology, Nephrology and Metabolic Diseases, Charité - Universitätsmedizin Berlin, Berlin, Germany
- ¹¹Institute of Biomedical Sciences, Faculty of Medicine, Vilnius University, Vilnius, Lithuania
- ¹²Regina Margherita Children's Hospital, Paediatric Gastroenterology Unit, Torino, Italy
- ¹³National Centre for Inherited Metabolic Disorders, Children's Health Ireland, Dublin, Ireland
- ¹⁴Hannover Medical School, Clinic for Paediatric Kidney, Liver, and Metabolic Diseases, Hannover, Germany
- ¹⁵First Department of Paediatrics, Semmelweis University, Budapest, Hungary
- ¹⁶Department of General Paediatrics, Neonatology and Paediatric Cardiology, University Children's Hospital, Heinrich-Heine-University Düsseldorf, Düsseldorf, Germany
- ¹⁷Department of Paediatrics, Hospital Reutlingen, Reutlingen, Germany
- ¹⁸Department of Human Genetics, Inselspital, Bern University Hospital, University of Bern, Bern, Switzerland
- ¹⁹CLAD Ouest CHU Hôpital Sud, CRMR Déficiences intellectuelles, Service de Génétique Médicale, Rennes, France
- ²⁰King's College Hospital, Paediatric Liver, GI & Nutrition Centre, London, United Kingdom
- ²¹Eberhard Karls University Tuebingen, Paediatric Gastroenterology and Hepatology, Tuebingen, Germany
- ²²Department of Paediatrics, University Medical Centre Hamburg-Eppendorf, Hamburg, Germany
- ²³Institute of Human Genetics, University Hospital Heidelberg, Heidelberg, Germany
- ²⁴University Medical Centre Hamburg-Eppendorf, Institute of Human Genetics, Hamburg
- ²⁵Department of Paediatrics, University Hospital Erlangen, Erlangen, Germany
- ²⁶Paediatric Gastroenterology, UMC Utrecht, Utrecht, The Netherlands
- ²⁷Children's Health Ireland, Temple Street Hospital, Dublin, Ireland
- ²⁸Department of Translational Medical Sciences, University of Naples Federico II, Naples, Italy
- ²⁹Department of Medical Genetics, Children's Memorial Health Institute, Warsaw, Poland
- ³⁰Department of Paediatric Gastroenterology, Hepatology and Nutrition, Third Department of Paediatrics, Attikon University General Hospital, National and Kapodistrian University of Athens, Athens, Greece
- ³¹Department of Paediatric Gastroenterology, Ankara University, School of Medicine, Ankara, Turkey
- ³²Department of Paediatrics II, Paediatric Gastroenterology, Hepatology and Liver Transplantation, University Hospital Essen, Essen, Germany
- ³³Birmingham Children's Hospital NHS Trust, Liver Unit, Birmingham, UK
- ³⁴University Hospital Regensburg, KUNO University Children's Hospital, Regensburg, Germany
- ³⁵Department of General Paediatrics, University Medical Centre Schleswig-Holstein, Kiel, Germany
- ³⁶Department of Paediatric Neurology, Centre for Neuromuscular Disorders, Centre for Translational Neuro and Behavioral Sciences, University Duisburg-Essen, Essen, Germany
- ³⁷Department of Paediatrics and Adolescent Medicine, Medical University of Vienna, Vienna, Austria
- ³⁸Research Centre for Medical Genetics, Moscow, Russian Federation
- ³⁹Department of Neurometabolism, University Hospital of Nantes, Nantes, France
- ⁴⁰Department of Pediatrics, Faculty of Medicine and University Hospital Carl Gustav Carus, Technische Universität Dresden, Dresden, Germany
- ⁴¹Department of Paediatrics, Dr. von Hauner Children's Hospital, University Hospital, LMU Munich, Munich, Germany
- ⁴²Department of Paediatrics, Rambam Medical Centre, Meyer Children's Hospital, Metabolic Unit, Haifa, Israel
- ⁴³University Children's Hospital, Paracelsus Medical University Salzburg, Salzburg, Austria
- ⁴⁴University of Pittsburgh and Children's Hospital of Pittsburgh of UPMC, Pittsburgh Liver Research Centre, Pittsburgh, Pennsylvania, USA
- ⁴⁵Children's Liver Unit, Leeds Children's Hospital, Leeds, UK
- ⁴⁶Department of Paediatrics, Gynecology, and Obstetrics, Division of Paediatric Subspecialities, Swiss Paediatric Liver Centre, Paediatric Gastroenterology, Hepatology and Nutrition Unit, University of Geneva, Geneva, Switzerland
- ⁴⁷Jeanne de Flandres Hospital, Reference Centre for Inherited Metabolic Diseases, Lille, France
- ⁴⁸Department of Paediatrics, Division of Neonatology and Paediatric Intensive Care, University Hospital Marburg, Marburg, Germany

⁴⁹Healthcare Department Morozov Children's City Clinical Hospital, Moscow City, Moscow

⁵⁰Laboratory of Genetics of Mitochondrial Diseases, Imagine Institute, University Paris Cité, INSERM UMR, Paris, France

⁵¹Department of Paediatrics, University Hospital Augsburg, Augsburg, Germany

⁵²Department of Paediatrics, Division of Paediatric Gastroenterology, Hepatology and Nutrition, Inselspital, Bern University Hospital, University of Bern, Bern, Switzerland

⁵³Wellcome Centre for Mitochondrial Research, Translational and Clinical Research Institute, Faculty of Medical Sciences, Newcastle University, Newcastle upon Tyne, UK

⁵⁴Department of Paediatrics I, University Hospital Essen, University of Duisburg-Essen, Essen, Germany

⁵⁵Clinical Biochemistry Department, Hospital Sant Joan de Déu, IRSJD, Esplugues de Llobregat, Barcelona, Spain and Centro de Investigación Biomédica en Red de Enfermedades Raras (CIBERER)- Instituto de Salud Carlos III, Spain

⁵⁶Institute of Cell Biology, Medical University of Innsbruck, Innsbruck, Austria

Correspondence

Christian Staufner, University Hospital Heidelberg, Center for Child and Adolescent Medicine, Department I, Division of Pediatric Neurology and Metabolic Medicine, Im Neuenheimer Feld 430, D-69120 Heidelberg, Germany.
Email: Christian.Staufner@med.uni-heidelberg.de

Holger Prokisch, Institute of Human Genetics, Klinikum rechts der Isar, Technical University Munich, Trogerstraße 32, 81675 München, Germany
Email: Holger.Prokisch@helmholtz-munich.de

Abstract

Background and Aims: Pediatric acute liver failure (PALF) is a life-threatening condition. In Europe, the main causes are viral infections (12%–16%) and inherited metabolic diseases (14%–28%). Yet, in up to 50% of cases the underlying etiology remains elusive, challenging clinical management, including liver transplantation. We systematically studied indeterminate PALF cases referred for genetic evaluation by whole-exome sequencing (WES), and analyzed phenotypic and biochemical markers, and the diagnostic yield of WES in this condition.

Approach and Results: With this international, multicenter observational study, patients (0–18 y) with indeterminate PALF were analyzed by WES. Data on the clinical and biochemical phenotype were retrieved and systematically analyzed.

Results: In total, 260 indeterminate PALF patients from 19 countries were recruited between 2011 and 2022, of whom 59 had recurrent PALF. WES established a genetic diagnosis in 37% of cases (97/260). Diagnostic yield was highest in children with PALF in the first year of life (41%), and in children with recurrent acute liver failure (64%). Thirty-six distinct disease genes were identified. Defects in *NBAS* ($n=20$), *MPV17* ($n=8$), and *DGUOK* ($n=7$) were the most frequent findings. When categorizing, the most frequent were mitochondrial diseases (45%), disorders of vesicular trafficking (28%), and cytosolic aminoacyl-tRNA synthetase deficiencies (10%). One-third of patients had a fatal outcome. Fifty-six patients received liver transplantation.

Conclusions: This study elucidates a large contribution of genetic causes in PALF of indeterminate origin with an increasing spectrum of disease entities.

Abbreviations: ALF, acute liver failure; ALT, alanine aminotransferase; AST, aspartate aminotransferase; HPO, human phenotype ontology; INR, international normalized ratio; PALF, pediatric acute liver failure; PALFES, pediatric acute liver failure exome sequencing; PALFSG, PALF study group; *POLG*, polymerase gamma; RALF, recurrent acute liver failure; SI, IU system; TUM, Technical University Munich; WES, whole-exome sequencing.

Dominic Lenz, Lea D. Schlieben and Masaru Shimura share first authorship and contributed equally to the paper. Kei Muramaya, Christian Staufner and Holger Prokisch are shared last authors and contributed equally to the paper.

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The high proportion of diagnosed cases and potential treatment implications argue for exome or in future rapid genome sequencing in PALF diagnostics.

INTRODUCTION

Pediatric acute liver failure (PALF) is a rare, life-threatening clinical condition mainly affecting children in their first year of life.^[1] In the United States, PALF is mainly caused by paracetamol intoxication (13%), metabolic disorders (10%), and viral infections (8%).^[2] In Europe, a systematic collection of data on PALF etiologies is lacking. Single-center report inherited metabolic diseases (14%–28%) and viral infections (12%–16%) as the main causes of PALF in Europe.^[3,4] In US and European cohorts, the underlying etiology remained unclear in about half of cases, hampering clinical management including disease-specific therapies, particularly decision-making regarding liver transplantation.^[1–4] This uncertainty is critical for survival with a high burden for physicians, affected individuals, and their families. Therefore, establishing a causal diagnosis is central to PALF. Although standardized approaches regarding biochemical testing and targeted Sanger sequencing in children with PALF contributed to a higher diagnostic rate and helped to elucidate some PALF etiologies, a large fraction of PALF cases remained unclassified.^[5] Access to next-generation sequencing techniques unraveled novel or uncovered genetic causes of hitherto unsolved cases of PALF. An important milestone boosting whole-exome sequencing (WES) in acute liver failure (ALF) was the identification of biallelic variants in *NBAS* as a cause of recurrent acute liver failure (RALF) with onset in infancy (MIM: #616483).^[5,6] More recently, numerous *NBAS* cases^[7] but also reports on other rare genetic diseases associated with PALF were published, such as infantile liver failure syndrome type 1 due to variants in *LARS1* (Mendelian Inheritance in Man (MIM): #615438),^[8,9] infantile liver failure syndrome type 3 due to variants in *RINT1* (MIM: #618641),^[10] and transient infantile liver failure syndrome due to variants in *TRMU* (MIM: #613070).^[11] In the context of the rapidly growing knowledge on genetic causes of PALF, a single-center retrospective analysis of 148 PALF cases (between 2001 and 2011) identified the underlying genetic cause of formerly indeterminate cases in 27% using custom next-generation sequencing panel, indicating the high proportion of genetic causes among unsolved cases before the era of next-generation sequencing.^[12] However, a systematic WES study has not been performed to date in individuals with PALF of unknown etiology.

Therefore, in the present international, multicenter study, we aimed to study the proportion, type, and

biochemical and clinical presentation of genetic diseases identified by WES in a cohort of individuals with PALF of indeterminate etiology.

METHODS

Study design and recruitment of patients

The study was carried out as an international, multicenter study including both prospective and retrospective cases. The inclusion criterion was the clinical diagnosis of PALF of unknown etiology in children aged 0–18 years. The clinical diagnosis of PALF was made according to local criteria, including patient phenotype and laboratory parameters such as plasma alanine aminotransferase (ALT) and aspartate aminotransferase (AST) activity, bilirubin and INR (international normalized ratio). Individuals were eligible if diagnostic workup did not establish the etiology of PALF; workup was broadly consistent within the collaborating expert centers for pediatric hepatology (for details see Supplemental Table S1, <http://links.lww.com/HEP/I114>). When there was more than one episode of PALF, cases were characterized as RALF. Data on country of origin, sex, clinical signs, and symptoms using the human phenotype ontology (HPO)^[13] as well as laboratory and histology data were retrieved by means of a specific case report form. All research was conducted in accordance with both the Declarations of Helsinki and Istanbul and the ethical standards of the responsible committee on human experimentation. Informed consent to participate in the study was obtained from all patients and/or from their parents in case of minor patients. The study was approved by the ethical committees of the Technical University Munich and the Medical Faculty Heidelberg (study number: S-035_2014).

WES and variant prioritization

WES of cases and unaffected parents was performed at the Helmholtz Centre Munich (Munich, Germany) for research cases and the Institute of Human Genetics, Klinikum rechts der Isar, Technical University Munich (TUM, Munich, Germany) for diagnostic cases. WES data analysis was performed using an in-house pipeline of TUM.^[14] WES was performed on genomic DNA extracted from blood.^[15] Sequencing reads were aligned to human genome-build GRCh37/hg19 (UCSC Genome Browser) using the Burrows-Wheeler Aligner

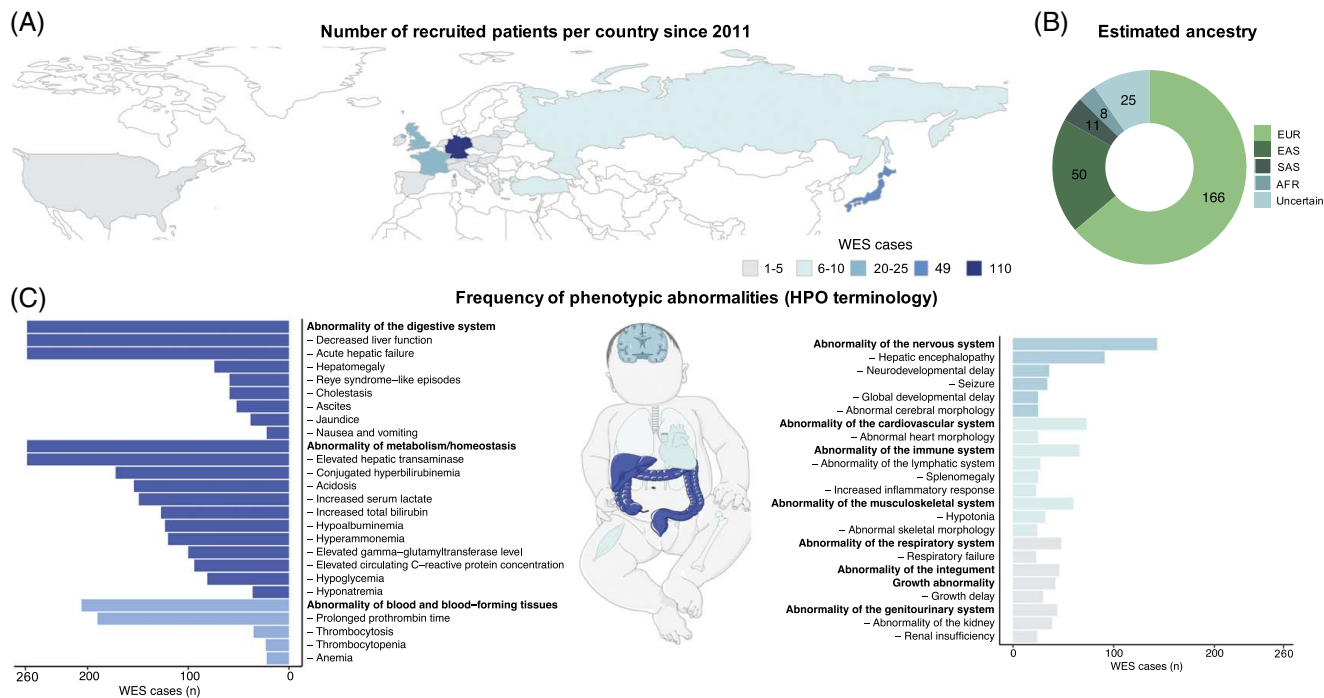


FIGURE 1 Geographical origin, clinical phenotypes, and affected organ systems. (A) Geographic origin: the number of patients recruited within the different countries is indicated by color. (B) Ancestry prediction based on WES data. (C) Frequency of phenotypic abnormalities within the whole cohort using the HPO. HPO terms with a frequency of 20 and higher are displayed. The exact INR values were not available for all cases, however, for all patients with available INR information, INR levels were abnormally elevated. All frequencies are shown in absolute numbers. Missing data may have led to low numbers in some of the items (eg, cholestasis, conjugated hyperbilirubinemia). Abbreviations: AFR, African; EAS, east Asian; EUR, European; HPO, Human Phenotype Ontology; SAS, south Asian; WES, whole-exome sequencing.

(v.0.7.5a).^[16] Single-nucleotide variants and small insertions and deletions were detected using the Genome Analysis Toolkit.^[17] Copy number variants were detected with ExomeDepth.^[18] Mitochondrial DNA (mtDNA) variants were assessed from exome data as described.^[19]

A special focus was given to genes previously reported to be associated with ALF. This gene list was manually created and extended through searching the OMIM database for all entries matching the keyword “liver failure.” This resulted in a set of 243 distinct candidate genes (Supplemental Table S2, <http://links.lww.com/HEP/I115>). Variants were classified according to the guidelines of the American College of Medical Genetics and Genomics^[20] using the Python package “InterVar” and the ClinVar annotation.^[21]

Phenotypic analysis

The following clinical variables were collected and analyzed: sex, weight, height, age at the time of the episode(s) of ALF/first symptoms of liver disease and at last assessment, patient survival, need for liver transplantation, and cause of death. Age ranges were defined as follows: neonatal period, age of onset at birth until day 28; infancy, age of onset before the second year of life; early childhood, age of onset between the second year of life and fifth year of life;

childhood, age of onset between the fifth and 12th year of life; adolescence, age of onset between the 12th and 18th year of life. Laboratory data during the episode of liver failure were collected using the IU system (SI). Additionally, the most prominent hepatic and extrahepatic clinical features were provided as HPO terms by the referring clinicians.^[13] Based on the provided HPO terms, ancestral HPO terms were derived from the ontology using the R package “OntologyX.”^[22]

Statistical analyses

Statistical analyses were performed using R version 4.0.4. For conducting survival analyses in R, the “Survival” and “Survminer” packages were used. The log-rank test was used to compare survival between different categorical variables.

Results

Study cohort

A total of 260 individuals were enrolled in this study between 2011 and 2022, with 111 individuals being prospectively enrolled and 149 patients enrolled

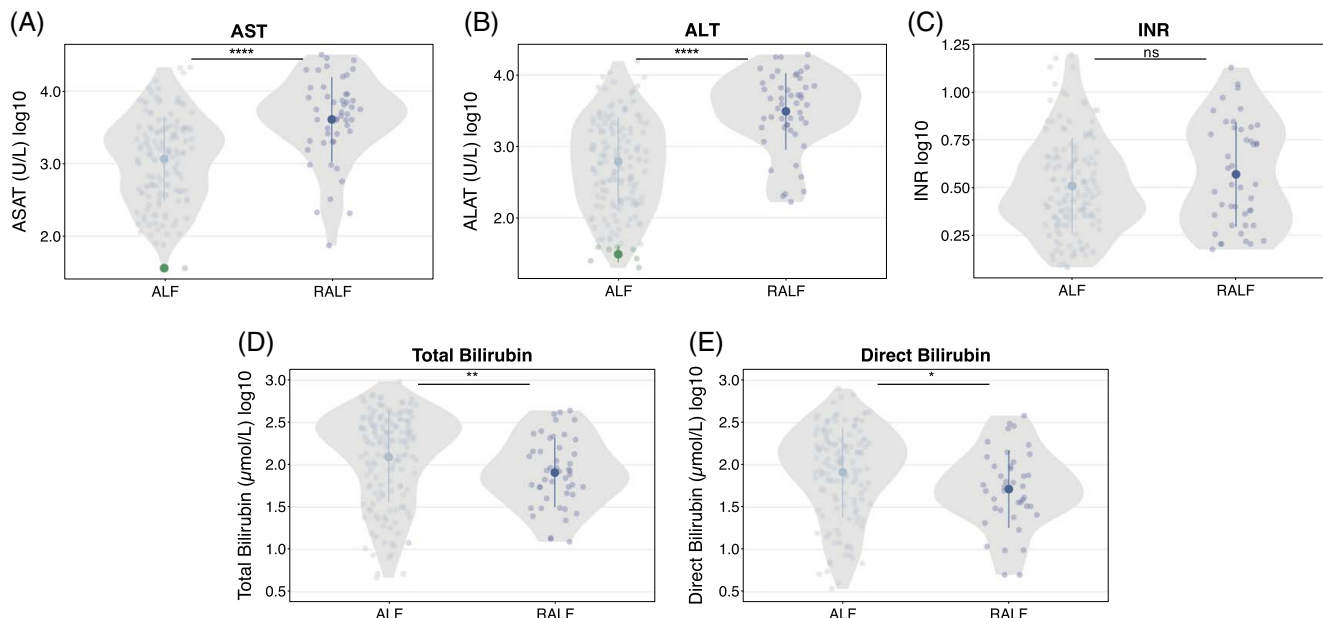


FIGURE 2 Biochemical characterization of the cohort. Laboratory characterization of ALF and RALF including aspartate aminotransferase, alanine aminotransferase, liver function (INR), total bilirubin as well as direct bilirubin using violin plots. Bold dots indicate the median, bars indicate the 25th to 75th percentile; green dots represent values within the reference range for normal values according to a consensus of Deutsche Gesellschaft für Klinische Chemie und Laboratoriumsmedizin and Verband der Diagnostika- und Diagnostikgeräte-Hersteller.^[29] * $p \leq 0.05$ ** $p \leq 0.01$ *** $p \leq 0.001$ **** $p \leq 0.0001$ by Wilcoxon test. Abbreviations: ALF, acute liver failure; ALT, alanine aminotransferase; AST, aspartate aminotransferase; RALF, recurrent acute liver failure.

retrospectively. Forty had been reported in publications on single disease gene discoveries or phenotypic spectrum studies.^[5–7,9–11,23–28] WES data were analyzed at the Technical University of Munich, of which 175 were examined as singletons and 85 by trio WES.

One hundred eighteen (118/260) patients were female (45%). Patients originated from centers in 19 countries in Europe, Asia, or North America (Figure 1A). The majority of patients (70%) were enrolled in Germany, Japan, and the United Kingdom, which is reflected in the estimated ethnic background based on the WES data (Figure 1B). Fifty-nine individuals (59/260) presented with RALF. On average, patients presented the second episode of ALF within 0.9 years (mean: 0.9 ± 0.9 y SD), whereas time intervals between ALF were significantly longer for further episodes (mean: 2.2 ± 4.8 y) (Supplemental Figure S1, <http://links.lww.com/HEP/I108>).

Three hundred seventy-eight (378) distinct, nonredundant HPO terms were reported, resulting in a median of 12 HPO terms per patient (range 2–29) spanning a median of 5 organ systems (Figure 1B, Supplemental Table S3, <http://links.lww.com/HEP/I116>).

With respect to clinical liver involvement, 74 patients presented with hepatomegaly, 59 had cholestasis, and 52 ascites. Jaundice was present in 38 individuals. Besides the liver, main phenotypic presentations included abnormalities of metabolism (100%), the blood and blood-forming tissues (79%), the nervous system (55%), the cardiovascular system (28%), and the immune system (25%) (Figure 1C).

All patients had abnormal blood homeostasis or abnormalities of metabolism (according to the HPO terminology), including elevated aminotransferases, hyperbilirubinemia, acidosis, increased serum lactate, hypoalbuminemia, hyperammonemia, and hypoglycemia. Concerning blood and blood-forming tissue, the most frequent clinical features besides prolonged prothrombin time were thrombocytosis, thrombocytopenia, and anemia. Frequently reported neurological abnormalities were HE, neurodevelopmental delay, and seizures. Organ involvement stratified by clinical phenotype revealed only few differences between ALF and RALF cases, but abnormalities of the head and neck, nervous and musculoskeletal system were more common in the RALF than in the ALF group (Fisher Exact test, nominal p -value < 0.05 ; Supplemental Figure S2, <http://links.lww.com/HEP/I108>).

Biochemical characterization of the whole cohort showed broad ranges for plasma ALT activity with a median of 907 U/L (range: 20–19,200 U/L) and AST activity with a median of 1939 (range: 36–31,800 U/L) (Figure 2A, B). When analyzed separately, patients with RALF had a significantly higher ALT activity with a median of 3,900 U/L (range: 167–19,200 U/L) (p -value 2.3×10^{-11} , Wilcoxon test) and AST activity with a median of 4804 (range: 74–31,800 U/L) (p -value 5.5×10^{-8} , Wilcoxon test) compared to the activity levels in the ALF group with ALT activity with a median of 582 U/L (range: 20–15,528 U/L) and AST activity with a median of 1395 (range: 36–21,227 U/L).

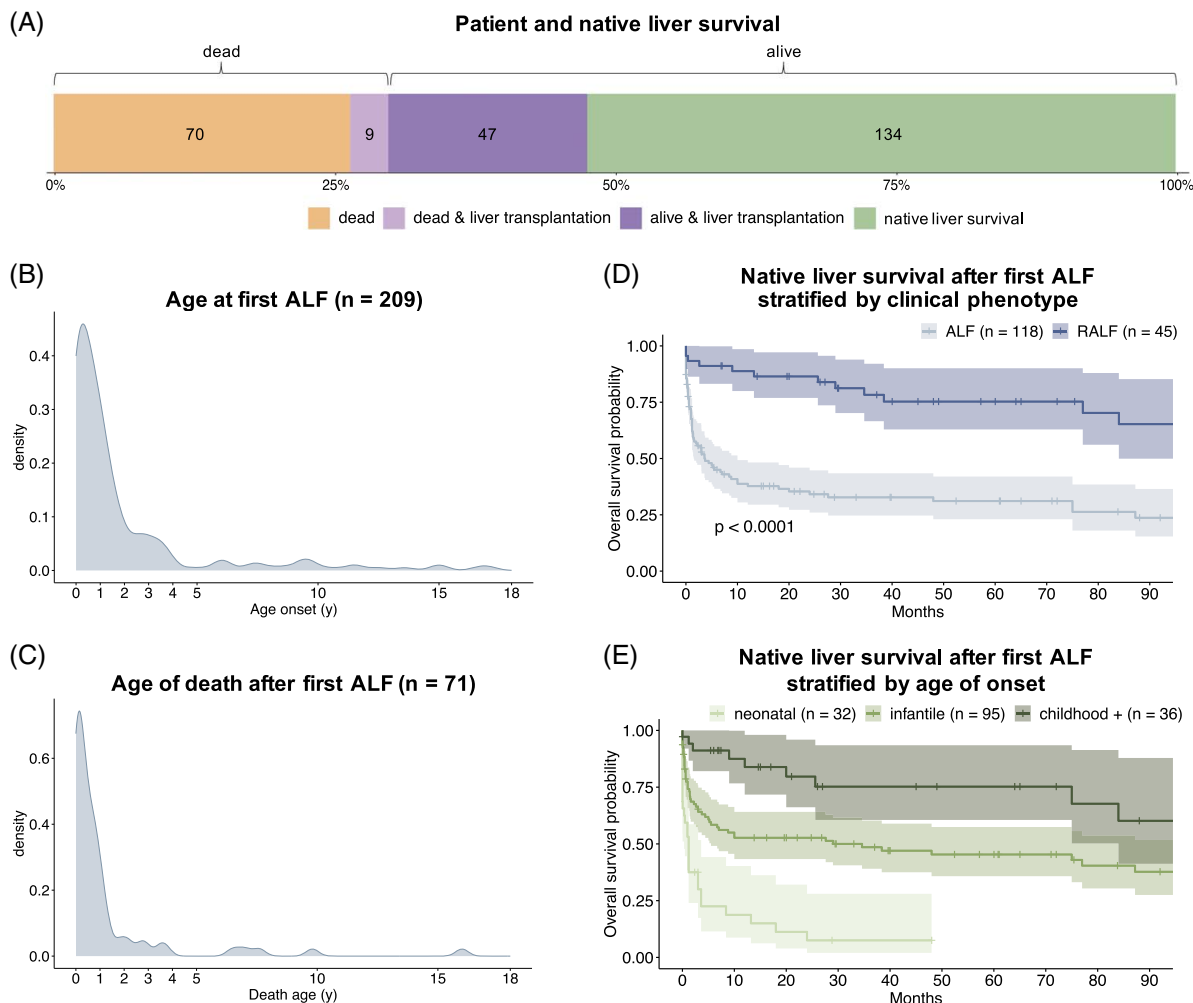


FIGURE 3 Outcome of patients with pediatric acute liver failure. (A) Patient and native liver survival. (B) Timeline for age at first ALF in years. (C) Timeline for age of death after the first ALF in years. (D) Kaplan-Meier plot for the native liver survival after the first ALF stratified by the clinical phenotype in months. p -value calculation by log-rank test. (E) Kaplan-Meier plot for the native liver survival after the first ALF stratified by the age of onset. Abbreviations: ALF, acute liver failure; RALF, recurrent acute liver failure.

Median maximal INR was 3 (range: 1.21–15.7), and median maximal total bilirubin was 140 $\mu\text{mol/L}$ (range: 4.62–1219.2 $\mu\text{mol/L}$). (Figure 2C–E). Sixteen patients in the ALF cohort showed a bilirubin level below 17.1 $\mu\text{mol/L}$ (1 mg/dL) and hence a low bilirubin phenotype. Five patients were reported to have a conjugated bilirubin below 5.1 $\mu\text{mol/L}$ (0.3 mg/dL). Bilirubin and INR levels did not differ significantly between patients with ALF and RALF.

Liver biopsy data were available for 104 patients, with steatosis (36/104) and fibrosis (34/104) being the most commonly reported histologic findings. Steatosis was found significantly more frequent in patients with RALF than in patients with ALF, whereas other histologic findings exhibited no differences between RALF and ALF (nominal p -value < 0.05 ; Supplemental Figure S3, <http://links.lww.com/HEP/1108>).

At the time point of data analysis, 181 patients were alive (134 with native liver survival, 47 with liver transplantation), while 79 children had died, 9 of those after liver transplantation (Figure 3A). Age of onset of ALF (first episode in patients with a RALF phenotype) ranged from neonatal to juvenile, with the majority of patients presenting within the first year of life (data available for 209 patients) (Figure 3B). The majority of deaths due to ALF (72%) occurred within the first year of life (Figure 3C). Comparison of the overall survival rate in the two clinical subgroups revealed a significantly higher native liver survival rate in the RALF subgroup as compared to the ALF subgroup (Figure 3D). When stratified by age of ALF onset, decreased native liver survival probability was associated with a younger age of onset (Figure 3E). A trigger for ALF was reported in 82 cases, with infections being the most prominent one (77 cases).

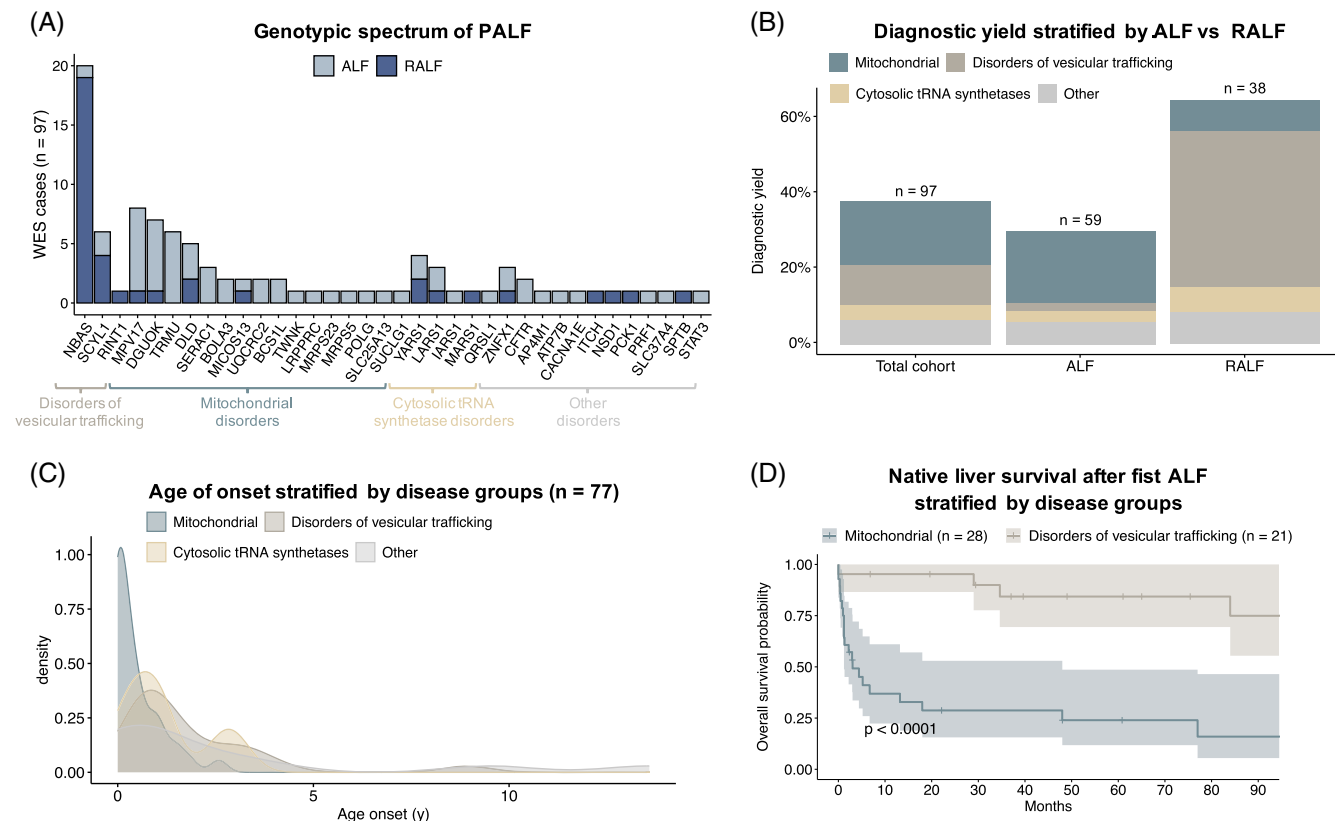


FIGURE 4 Molecular etiology of pediatric acute liver failure. (A) Genetic spectrum of pediatric acute liver failure; (B) diagnostic yield, stratified by ALF vs. RALF; (C) age of onset stratified by disease groups; and (D) native liver survival after first ALF stratified by the 2 most frequent disease groups. Abbreviations: ALF, acute liver failure; RALF, recurrent acute liver failure; WES, whole-exome sequencing.

Genetics

WES analysis established a genetic diagnosis in 97/260 of so far genetically unsolved cases (37%). Nineteen patients remained unsolved with variants of uncertain significance in OMIM disease genes or candidate disease genes. In the remaining 144/260 patients (55%), no variant could be prioritized. Pathogenic and likely pathogenic variants were detected in 36 distinct ALF disease-associated genes, of which more than half (21/36, 58%) were reported in single cases only (Figure 4A). Defects in *NBAS* (21%), *MPV17* (8%), and *DGUOK* (7%) occurred most frequently. In 92/97 cases, the causative genetic defect was inherited in an autosomal recessive fashion; among these, 53/92 were homozygous for disease-causing variants (Supplemental Table S4, <http://links.lww.com/HEP/I117>). In total, 108 distinct variants classified as “likely pathogenic” or “pathogenic” according to ACMG criteria within the 36 genes were found. Of the 108 variants, 56 (52%) were already reported as “likely pathogenic” or “pathogenic” in ClinVar at the time point of WES analysis. Missense variants formed the major proportion of all disease-causing variants. Out of the 36 cases in this study who had consanguineous parents, a genetic diagnosis was established in 22 (61%). Additionally, 20

out of 22 genetically diagnosed cases were found to be homozygous for the disease-causing variant.

When categorizing the disease genes into functional groups, most of the cases are associated with mitochondrial diseases (45%), followed by disorders of vesicular trafficking (28%) and cytosolic aminoacyl-tRNA synthetase deficiencies (10%) (Figure 4A). Diagnostic yield was higher in patients with ALF within the first 5 years of life (40%; n = 187) compared to children 6 years of age or above (14%; n = 21) (Supplemental Figure S4, <http://links.lww.com/HEP/I108>). Highest yield was achieved in infancy (41%; n = 123) and in children with RALF (64%; n = 59) (Figure 4B). The majority of patients with disorders of vesicular trafficking developed RALF. *NBAS* deficiency was the most frequent diagnosis in this group. In the majority of mitochondrial diseases (68%), ALF occurred within the first 6 months of life (median 2.5 months), whereas disorders of vesicular trafficking and cytosolic aminoacyl-tRNA synthetases were characterized by an age of onset of half a year and older (Figure 4C).

Patients with disorders of vesicular trafficking rarely died, while patients with mitochondrial disease showed a high mortality. For the other disease groups, patient numbers were too low for analyses of survival (Figure 4D).

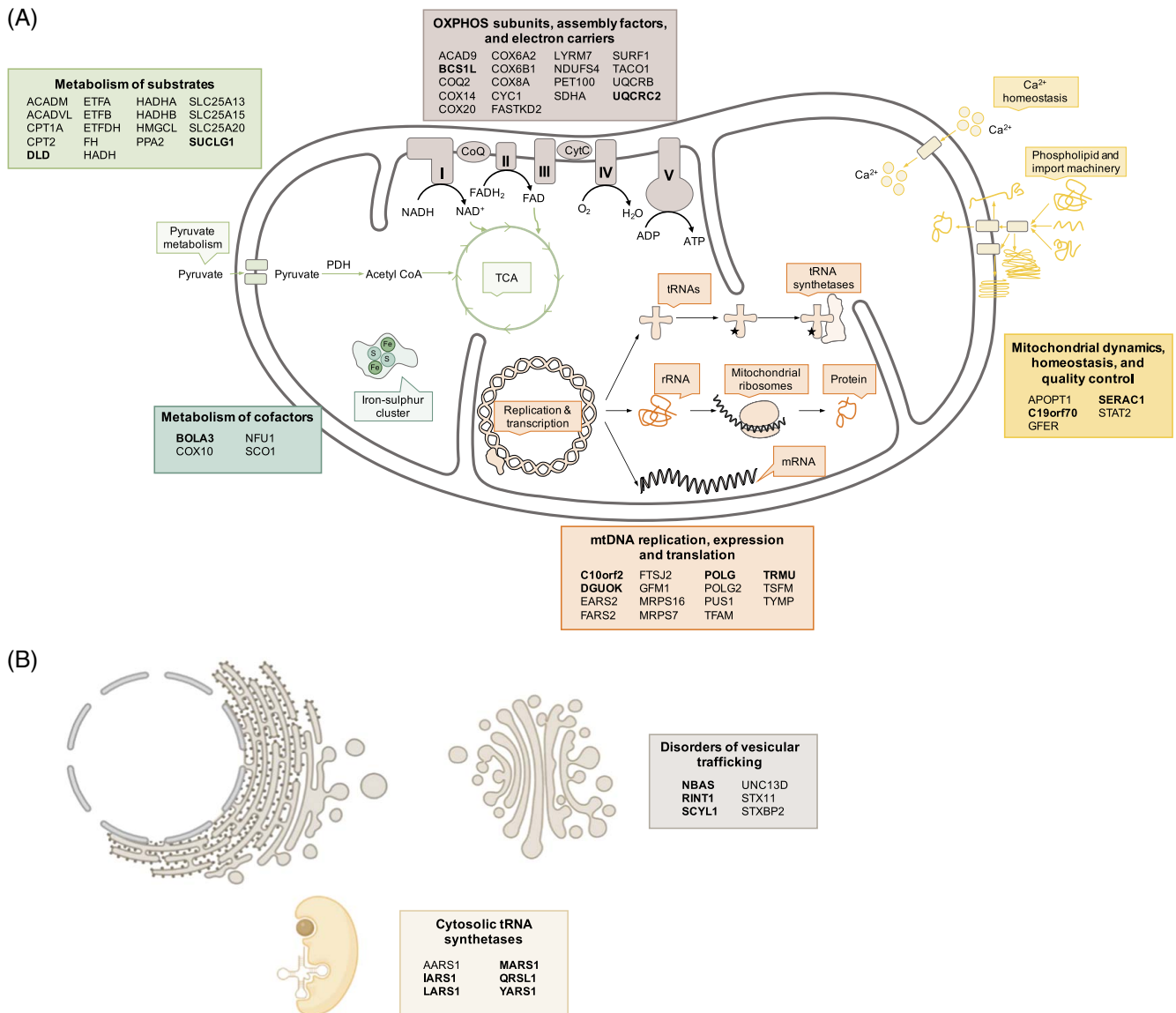


FIGURE 5 (A) Mitochondrial disease genes causing pediatric acute liver failure. (B) Disorders of vesicular trafficking and aminoacyl-tRNA synthetase deficiency (box) leading to pediatric acute liver failure. Genes with pathogenic variants that have been identified in cases within this study are shown in bold.

The 3 most frequent disease groups in this cohort presented a broad phenotypic spectrum with multiple organs involved (Supplemental Figure S5, <http://links.lww.com/HEP/I108>).

Liver biopsy data were available for 44 genetically solved cases (Supplemental Figure S6, <http://links.lww.com/HEP/I108>). While fibrosis was most common in patients with mitochondrial disorders, cirrhosis was found most frequently in patients with cytosolic tRNA synthetase deficiencies. Steatosis was the commonest finding in all three distinct disease groups, and necrosis was only rarely reported in any of the distinct disease groups. No significant differences in necrosis or inflammation incidence could be identified between the groups.

Subcohort fulfilling PALF study group inclusion criteria

The inclusion criteria of our cohort differed from those of the US PALF study group (PALFSG,^[1]). We investigated whether applying the more stringent PALFSG inclusion criteria to our cohort would yield different results. A total of 145/201 patients with ALF in our study fulfilled the PALFSG inclusion criteria. Comparing this subgroup with the overall cohort, there were no differences apart from INR (and the associated HPO umbrella term blood and blood-forming tissue), which was significantly higher in the subgroup fulfilling PALFSG criteria (for details see Supplemental Table S5, <http://links.lww.com/HEP/I118>).

DISCUSSION

PALF is a life-threatening event with very heterogeneous etiologies. Despite ongoing efforts of standardizing diagnostic approaches, in one-third to half of cases no causal diagnosis is achieved.^[30] In 2015, variants in *NBAS* were identified as a novel cause of PALF,^[5] the number of individuals diagnosed with *NBAS* deficiency rose rapidly^[6,7] and motivated to further explore the genetic landscape of PALF of unknown etiology. Hence, the pediatric acute liver failure exome sequencing (PALFES) study was started including up to now 260 individuals with PALF of unknown etiology from three different continents. This represents by far the largest cohort of patients with unresolved PALF studied by WES reported in the literature to date. A genetic diagnosis was established in 37% of patients with unresolved PALF; in patients with RALF the diagnostic yield was even higher (64%).

With the elucidation of about 40% of hitherto unexplained PALF cases, genetic diseases likely represent the largest group of PALF in terms of etiology. Nevertheless, it is noteworthy that 55% of patients in this series had no evidence of an underlying genetic disorder. Transcriptome or proteome analyses may further increase the diagnostic rate of genetic diseases by providing functional evidence on inconclusive candidate variants detected or by discovery of functional relevant variants missed by the genetic analysis of our cohort. Even disregarding this, a substantial part of indeterminate PALF is likely of nongenetic origin.^[31] According to Squires et al,^[32] the fraction of PALF cases with unknown etiology is particularly high in the first 3 years of life, which is the age range in which our study demonstrates the highest molecular diagnostic yield by WES.

Three main groups of genetic diseases underlying indeterminate PALF have been identified: mitochondrial disorders, disorders of vesicular trafficking, and cytosolic aminoacyl-tRNA synthetase deficiencies (Figure 5). Mitochondrial genetic diagnoses were noticeably more frequent within our study than had previously been reported.^[32] Still, in our cohort, one of the early reported genetic causes of PALF, mutations in mitochondrial polymerase gamma (*POLG*) associated with Alpers syndrome, was rarely observed. Because it is a known cause of PALF, *POLG*-suspected cases may have undergone targeted genetic testing and are therefore not included in this cohort. However, we cannot exclude the possibility of enrollment bias in the PALFES cohort, as some of the participating centers were studying a cohort of pediatric patients with mitochondrial diseases in parallel with the PALFES study, and thus mitochondrial-related ALF cases may have been assigned more frequently.^[33] In addition, the mere fact that a genetic study was conducted may have led to a higher inclusion rate of patients with suspected genetic causes.

Specifically, publications on new genetic causes of RALF during the study led to an increased sending of RALF cases. Therefore, the unbiased diagnostic yield should likely be lower, with a potentially lower rate of mitochondrial disease and genetic causes of RALF. This suggestion is supported by the high proportion of RALF cases in our cohort, for which a genetic cause is more likely to be expected.

A potential limitation of our study was the lack of an internationally standardized definition of PALF, which is also reflected in our cohort. Most of the individuals included in our cohort (201/260 including also RALF cases) met the stringent inclusion criteria of the longitudinal studies of PALF conducted by the "PALF Study Group" in the United States.^[1] Although the PALF Study Group inclusion criteria do not represent definition criteria of PALF, they are frequently used as such. Our analyses show that these more stringent criteria would not lead to different results, including diagnostic yield, age at onset, histology, survival, organ involvement, histology, and disease genes.

In general, PALF cases with a clear biochemical fingerprint (such as tyrosinemia type I, with elevated succinylacetone in urine or dried blood spots) are typically diagnosed based on metabolic investigations, and genetically confirmed in the further workup. The 3 disease groups identified in this study share the feature that no specific laboratory biomarker or metabolic fingerprint by itself can establish the molecular diagnosis or point specifically to a single disease gene. Accordingly, we could not identify a clear biochemical or clinical profile in the cases now diagnosed genetically by WES using already established investigations. More comprehensive metabolomic studies would be needed to extend the search for diagnostic biomarkers. Consequently, with our current knowledge, we advocate performing genetic analyses using WES or whole-genome sequencing in parallel to metabolic analyses as the timely decision on further appropriate treatment options such as transplantation and/or specific drug therapy or dietary management in case of metabolic diseases will depend on the underlying disorder.

Our study demonstrates that the lack of biomarker specificity also holds true for lactate in serum, as lactate concentration could not differentiate between mitochondrial disorders from other causes of PALF, indicating that lactic acidemia is also a secondary finding in severe liver dysfunction such as PALF. Extremely high levels of AST and ALT have been associated with *NBAS* deficiency,^[6] often presenting as RALF. This reflects the significantly higher AST/ALT levels in this group; however, diagnosis cannot be ascertained based on this finding. Age of onset can direct the clinical suspicion toward possible genetic disorders underlying PALF, with mitochondrial disorder most frequently occurring in the neonatal period. Another frequent cause of neonatal ALF is gestational alloimmune liver

disease presenting typically with low AST and ALT levels. Such cases have not been included in our study. In our cohort, only 5 out of 50 neonates showed ALT < 100 U/L. Of note, in 2 of the 5 cases with low ALT, pathogenic variants in *DGUOK* were found to be responsible for the phenotype. There is a substantial overlap among the different disease groups with respect to age of onset.

An unexpected histological finding in the PALFES cohort is the high percentage (30%) of progressive liver remodeling. This indicates that a subgroup has a preceding subclinical disease, with PALF as the first clinical presentation.

The most commonly affected extrahepatic organ system was the nervous system, with HE and neurodevelopmental delay reported most often. In analogy to biochemical parameters, also the clinical parameters in our study did not discriminate between cases with and without a genetic diagnosis. This again emphasizes the relevance and necessity of genome-wide genetic analyses in PALF to determine the underlying cause.

Establishing a (genetic) diagnosis is crucial for clinical management and outcome of children with PALF. While causative treatments are not available for most of the genetic disorders detected in our cohort, there are specific management approaches for several diseases. In patients with TRMU deficiency, it has been shown that cysteine supplementation improved survival significantly compared to patients without cysteine supplementation.^[11] Forced antipyretic management in ILFS1, ILFS2, and ILFS3 may help to avoid fever-triggered RALF.^[6,7,9,10] Additionally, the decision on transplantation is dependent on the underlying disorder and the expected outcome in relation to both transplant liver survival and extrahepatic features and finally, the risk of recurrence of PALF in another child of the family can only be predicted with a genetic diagnosis. Our study demonstrates that outcome differs depending on the underlying disease etiology; native liver survival is significantly higher in individuals with disorders of vesicular trafficking compared to mitochondrial diseases. The role of liver transplantation for individuals with PALF due to genetic diseases is discussed controversially in the literature especially for mitochondrial disorders, due to potentially unfavorable neurological, cardiac, or neuromuscular involvement and poor outcome, arguing against liver transplantation.^[34] However, there are mitochondrial disorders such as TRMU or hepatic DLD deficiency where extrahepatic involvement is scarce with a favorable neurological prognosis.^[11,35] Moreover, multisystemic mitochondrial disorders are not contradicting liver transplant in general, as there are reports of patients, for example, with *DGUOK* deficiency, with minor neurological involvement who received a liver transplantation and had a satisfactory post-transplant course.^[11,36,37]

In order to help decision-making, diagnosis of PALF and the identification of the etiology needs to be

established within a short period of time as the clinical situation typically is critical and decisions are time-sensitive. Turnaround time in our study was not assessed systematically, and individuals have also been enrolled retrospectively where turnaround time was not essential for clinical care. However, it is impractical to wait for a long time to receive WES results in critically ill patients. Hence, from a technical point of view, genome sequencing would be faster and more accurate than WES, with reported turnaround time as low as 24 hours, being a promising option to achieve genetic diagnoses especially in critically ill children.^[38,39] Higher investment costs and demanding computational power hampers availability of this technology in most hospitals, but will likely be available in the near future in many countries. Again, rapid establishment of a (genetic) diagnosis has the potential to both save lives and reduce costs.^[38,39]

In conclusion, this study demonstrates that a relevant number of indeterminate PALF cases can be solved by WES. This shifts PALF in the range of other rare disorders for which WES/whole-genome sequencing is now a well-established diagnostic procedure. Major identified groups are mitochondrial disorders, disorders of vesicular trafficking, and cytosolic aminoacyl-tRNA synthetase deficiencies. An ascertained diagnosis helps physicians make treatment decisions and can save lives and costs. For these reasons, we assert that WES, or in the near future whole-genome sequencing, should be within a first-line diagnostic approach for every child presenting with PALF.

AUTHOR CONTRIBUTIONS

Dominic Lenz, Lea D. Schlieben, Masaru Shimura, Kei Murayama, Christian Staufner, and Holger Prokisch were responsible for the conception and design of the study. Data acquisition was performed by Dominic Lenz, Lea D. Schlieben, Masaru Shimura, Alyssa Bianzano, Riccardo Berutti, Rüdiger Adam, Denise Aldrian, Ivo Baric, Ulrich Baumann, Neslihan E. Bozbulut, Philip Bufler, Birutė Burnytė, Pier L. Calvo, Ellen Crushell, Buket Dalgıç, Anibh M. Das, Antal Dezsőfi, Felix Distelmaier, Alexander Fichtner, Peter Freisinger, Harald Gaspar, Louise Goujon, Nedim Hadzic, Steffen Hartleif, Sven F. Garbade, Bianca Hegen, Maja Hempel, Stephan Henning, Andre Hoerning, Roderick Houwen, Joanne Hughes, Raffaele Iorio, Katarzyna Iwanicka-Pronicka, Martin Jankofsky, Norman Junge, Ino Kanavaki, Aydan Kansu, Sonja Kaspar, Simone Kathemann, Deidre Kelly, Ceyda T. Kırsaçlıoğlu, Birgit Knoppke, Martina Kohl, Heike Köbel, Vassiliki Konstantopoulou, Tatiana Krylova, Zarife Kuloğlu, Alice Kuster, Martin W. Laass, Elke Lainka, Eberhard Lurz, Hanna Mandel, Johannes A. Mayr, Patrick McKiernan, Patricia McClean, Valerie McLin, Karine Mention, Hanna Müller, Laurent Pasquier, Natalia Pechatnikova, Bianca Peters, Danijela

Petković Ramadža, Dorota Piekutowska-Abramczuk, Denisa Pilic, Sanjay Rajwal, Nathalie Rock, Agnès Roetig, René Santer, Wilfried Schenk, Natalia Semenov, Christiane Sokollik, Ekkehard Sturm, Robert W. Taylor, Eva Tschiedel, Vaidotas Urbonas, Roser Urreiziti, Jan Vermehren, Jerry Vockley, Georg-Friedrich Vogel, Wendy van der Woerd, Saskia B. Wortmann, and Ekaterina Zakharova.

Lea D. Schlieben, Dmitrii Smirnov, Robert Kopajtich, Theresa Brunet, Melanie Brugger, Matias Wagner, Katharina Mayerhanser, and Thomas Meitinger interpreted genetic results. Lea D. Schlieben was responsible for the statistical analysis and data visualization. Dominic Lenz and Lea D. Schlieben drafted the article. Masaru Shimura, Kei Murayama, Christian Staufner, Holger Prokisch, Georg F. Hoffmann, Thomas Meitinger, and Stefan Kölker critically revised the content. Christian Staufner and Holger Prokisch supervised the study.

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CONFLICTS OF INTEREST

Ulrich Baumann consults, advises, is on the speakers' bureau, and received grants from Albireo and Mirum. He advises and received grants from Alexion and Astellas. He consults for Vivet. Pier Luigi Calvo advises Albireo/Ipsen, Mirum, and Nestle. Felix Distelmaier received grants from Danone, DFG, Elterninitiative Kinderkrebsklinik E.V., Margarete-Breuer Stiftung, PTC Therapeutics, Santhera, and TANGO2 Research Foundation. Elke Lainka advises Albireo and Mirum. Eberhard Lurz consults, advises, and is on the speakers' bureau for Albireo, Mirum, Nutricia, and Takeda. The remaining authors have no conflicts to report.

ORCID

Dominic Lenz  <https://orcid.org/0000-0002-9561-8190>

Lea D. Schlieben  <https://orcid.org/0000-0001-6583-256X>

Dmitrii Smirnov  <https://orcid.org/0000-0002-5802-844X>

Christian Staufner  <https://orcid.org/0000-0002-1403-9075>

Holger Prokisch  <https://orcid.org/0000-0003-2379-6286>

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