Congenital Diarrhea and Cholestatic Liver Disease: Phenotypic Spectrum Associated with MYO5B Mutations

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Abstract: Myosin Vb (MYO5B) is a motor protein that facilitates protein trafficking and recycling in polarized cells by RAB11- and RAB8-dependent mechanisms. Biallelic MYO5B mutations are identified in the majority of patients with microvillus inclusion disease (MVID). MVID is an intractable diarrhea of infantile onset with characteristic histopathologic findings that requires life-long parenteral nutrition or intestinal transplantation. A large number of such patients eventually develop cholestatic liver disease. Bi-allelic MYO5B mutations are also identified in a subset of patients with predominant early-onset cholestatic liver disease. We present here the compilation of 114 patients with disease-causing MYO5B genotypes, including 44 novel patients as well as 35 novel MYO5B mutations, and an analysis of MYO5B mutations with regard to functional consequences. Our data support the concept that (1) a complete lack of MYO5B protein or early MYO5B truncation causes predominant intestinal disease (MYO5B-MVID), (2) the expression of full-length mutant MYO5B proteins with residual function causes predominant cholestatic liver disease (MYO5B-PFIC), and (3) the expression of mutant MYO5B proteins without residual function causes both intestinal and hepatic disease (MYO5B-MIXED). Genotype-phenotype data are deposited in the existing open MYO5B database in order to improve disease diagnosis, prognosis, and genetic counseling.

Keywords: congenital diarrheal diseases; enteropathy; microvillus inclusion disease; MYO5B; myosin Vb; progressive familial intrahepatic cholestasis; PFIC; genotype–phenotype correlation; lack of protein; tail domain

1. Introduction

Microvillus inclusion disease (MVID) is a severe congenital enteropathy with intractable watery diarrhea, most often starting in the first days after birth but at times starting within the first months of life. Histopathology demonstrates hypoplastic villous atrophy; characteristic cytoplasmic inclusions of brush border microvilli and "secretory granules", a subapical accumulation of aberrant vesicular compartments, are detected
by electron microscopy [1]. Both immunohistochemistry and electron microscopy reveal the mislocalization of brush border transporter proteins and rarefication of microvilli ultimately leading to osmotic and secretory diarrhea, malabsorption, and failure to thrive. The prognosis is generally poor and continuous total parenteral nutrition (TPN) and bowel transplantation are therapeutic options. A number of patients die at a young age as a consequence of septicemia [2–4]. MVID is autosomal recessively inherited with locus heterogeneity, where the majority of patients harbors biallelic mutations in the myosin 5b gene (MYO5B), and defects in syntaxin 3 (STX3) and syntaxin binding protein 2 (STXBP2) account for additional cases of MVID [1,5–8].

Up to 54% of MVID patients develop persistent cholestatic liver disease, which appears unrelated to treatment with TPN and is clinically indistinguishable from progressive familial intrahepatic cholestasis (PFIC) types I, II, IV, and V (PFIC-1, PFIC-2, PFIC-4, PFIC-5) [9–11]; there is persistent cholestasis with normal serum gamma-glutamyl transferase (GGT) concentrations, reduced concentrations of primary bile acids in bile, and progressive liver damage that frequently requires liver transplantation in childhood [12–14]. PFIC-1, PFIC-2, PFIC-4, and PFIC-5 are caused by biallelic mutations in genes encoding the canalicular membrane transporter proteins ATP8B1, the aminophospholipid flippase FIC-1 and ABCB11, the bile salt export pump (BSEP), the tight junction protein-2 (TJP2), and the transcription factor NR1H4. Recently, bi-allelic MYO5B mutations have also been identified in a small number of patients with a low-GGT PFIC phenotype in the absence of congenital diarrhea [9–11]; MYO5B is therefore considered both the main disease gene for MVID and to represent one of the increasing number of disease genes for low-GGT type PFIC, and the delineation PFIC6 has been suggested for this disorder [15].

Biallelic MYO5B mutations are thus identified in patients with a spectrum of clinical manifestations, ranging from intestinal disease (MYO5B-MVID) to intestinal disease combined with cholestatic liver disease in the same patients (MYO5B-MIXED), where MVID and the liver disease can be of equal or unequal medical concern, to predominant cholestatic liver disease, clinically indistinguishable from low-GGT PFIC (MYO5B-PFIC). Very recently, molecular mechanisms were proposed that can explain the occurrences of predominant intestinal, predominant hepatic and mixed phenotypes [15]. We compiled the clinical outcomes of 114 novel and previously reported patients with mostly biallelic MYO5B mutations and probed the dataset for genotype–phenotype correlation with respect to proposed disease mechanisms. Our study includes 44 novel patients with biallelic MYO5B mutations, of which 35 MYO5B mutations are novel. Novel data will be entered into the international online registry for patients with MYO5B mutations and their phenotype information (http://www.MVID-central.org) [16]; this patient registry already contains incomplete phenotype or genotype information on a small number of patients reported in our study; our study enables straightforward complementation of these entries.

2. Materials and Methods

The Department of Pediatrics I at the Medical University of Innsbruck participates in a pediatric liver transplantation program in Austria, and there is both a diagnostic and research focus on pediatric liver diseases as well as a research focus on the delineation of congenital and inherited diarrheas. Patients and patient samples are referred continuously for genetic testing to determine the etiology of the apparent hepatic or intestinal disorder. Written informed consent for molecular research investigations was obtained from the patients or from minor patients’ parents, and the studies were approved by the local ethics committee (vote No. AN2016-0029 359/4.5).

In this study, we retrospectively identified all cases referred to our institution between 2007 and 2020 with chronic diarrhea, with cholestatic liver disease, and with both chronic diarrhea and chronic liver disease; all cases involved patients in whom we identified biallelic pathogenic or likely pathogenic MYO5B variants. We then re-contacted the referring physicians to provide a standardized, questionnaire-based follow-up on their referrals. The
following data were abstracted from the medical records of each patient: family history, sex, ethnicity, parental consanguinity, age at onset of diarrhea, failure to thrive, dependence on parenteral nutrition, single- or multi-organ transplantation, characteristic histological findings of duodenal biopsies, outcome at last patient contact. Routine hepatic function criteria were determined in patients with chronic diarrhea and with chronic liver involvement. Genomic DNA was isolated from patients’ peripheral blood leukocytes by standard procedures. DNA samples from patients’ healthy parents were obtained and tested for MYO5B variant segregation with disease.

Targeted Sanger sequencing of the MYO5B gene was conducted in 42 patients as reported previously [17,18]; MYO5B gene analysis was performed in whole-exome-sequencing (WES) data in the remaining 27 patients; WES was conducted and data processed as reported [19]; variant calling was restricted to the MYO5B region (chr18:47349156-47721542 in the hg19 reference sequence) in patients with a clinical diagnosis of MVID. In five instances, MYO5B mutation analysis had been conducted initially in a collaborating diagnostic laboratory. In patients referred with a clinically indeterminate congenital diarrhea, the genomic regions of genes known to cause forms of congenital diarrhea and immune deficiency syndromes were evaluated (gene panel list available from the authors on request). Patients with cholestatic liver disease, in whom mutations in PFIC1-3 genes had been excluded, were evaluated for MYO5B variants. Categorization and interpretation of sequence variants were based on variant allele frequencies <0.005 in public domain population databases, and predicted effects on protein function by in-silico evaluation using Polyphen2 and CADD programs [20]. The nomenclature of identified variants and patients’ genotype follows the Human Genome Variation Society guidelines (HGVS v 2.0, www.hgvs.org/mutnomen). The MYO5B variant designation is based on the NCBI reference sequence for transcript NM_001080467.3 and the genomic reference sequence NG_012925.2. The exon numbering is based on NG_033082.1. Nucleotide numbering uses +1 as the A of the ATG translation initiation codon in the reference sequence, with the initiation codon as codon 1.

We evaluated the MYO5B mutation-related data from our center together with published mutation data that were retrieved by a PubMed search with search terms “MYO5B mutations, microvillous atrophy, microvillus atrophy, microvillus inclusion disease” and in which associated phenotype data were reported. Compiled data (Figure 1) were analyzed with respect to genotype–phenotype correlations and previously proposed cellular disease mechanisms [11,15,21].

The statistical significance of genotype–phenotype correlations was determined using Fisher’s exact test with \( p < 0.05 \).

3. Results

3.1. Study Population

In this study, we present a cohort of 114 patients with MYO5B mutations. Since the first observation of MYO5B gene mutations in 9 MVID patients [17], we have identified pathogenic and likely pathogenic MYO5B variants in a total of 67 patients at our center within the last decade. We previously reported findings in 23 of these patients [1,17,18,22]. The remaining 44 patients’ corresponding genetic and clinical data are presented in this study for the first time. From our literature search, we extracted genotype and phenotypic data of 47 additional patients with mostly biallelic MYO5B mutations [9–11,23–33] (Figure 1).

Patients were of diverse ethnicity, and parental consanguinity was known in one third of all cases. The median age at last patient contact was 5 years (IQ-range 9 years). For the 108 patients of the cohort whose gender was known, there was no bias (62 males and 46 females, \( p = 0.24 \), chi-squared test).

3.2. Clinical Characteristics

Final diagnoses of 114 patients are shown in Table 1. Age at disease onset was known in 107 out of 114 patients; disease manifested in the first week of life in 63.6% of patients
and after one year of age in 6 patients. Thirty-three out of 114 patients had died at a median age of 1 year (IQ-range 2.5 years). For 19 of 114 patients, outcome information was missing. Cause of death was mainly septicemia. Necrotizing enterocolitis, liver failure, macrophage activation syndrome, or drug resistant epilepsy with respiratory distress accounted for deaths in single cases. Twenty-two deceased patients had been diagnosed with MYO5B-MVID, and 11 patients with MYO5B-MIXED.

25 out of 114 patients (21.9%) received single or multi-organ transplants; information regarding transplantation was not available for 8 of 114 patients. Median age at 1st transplant surgery was 3.5 years (IQ-range 1 year), median age at 2nd transplant surgery was 9.25 years (IQ-range 6.88 years); first transplants were 17 x small bowel, 2 x small and large bowel, 5 x small bowel and liver, 1 x no data. Non-transplanted MVID patients were dependent on TPN, with the exception of 4 patients who required partial PN, and four patients could be weaned completely from PN at ages of 3–7 years but developed liver disease during childhood. We obtained no information on the current treatment of 8 patients.

Forty-one out of 114 patients had additional medical concerns, of which polyhydramnios, premature birth, failure to thrive, and renal Fanconi syndrome were seen in more than one patient. Congenital lung hypoplasia, necrotizing enterocolitis, bowel dissection, hematuria, proteinuria, calcuria, exocrine pancreatic insufficiency, low IgG level, hypothyroidism, gastroesophageal reflux disease, eczema, juvenile idiopathic arthritis, language development delay, pyramidal syndrome, epilepsy, short stature, microcephaly, and polydactyly were present in single cases.
Detailed information on all 114 individual patients is provided in Supplementary Table S1.

3.3. Molecular Findings in the Study Cohort

The 114 reported patients implicate 114 distinct MYO5B alleles in causing disease (Figure 2), of which we report 35 novel mutations here. These mutations include 4 intragenic deletions of one or several exons, 16 noncoding/splicing mutations, and 45 frameshifts/nonsense/initiation codon mutations, which are all predicted to lead to MYO5B truncation, as well as 49 missense mutations and one in-frame insertion.

Figure 2. Schematic overview of myosin Vb (MYO5B) protein and annotated mutations. (A) Full-length MYO5B with domain-positions in amino acids. Protein length and domain annotation are shown as in UniProt (http://www.uniprot.org/uniprot/Q9ULV0). The Motor (B), IQ 1–6 (C), coiled-coil (D) and globular tail domain (E) are shown separately. (B–E) Missense mutations are annotated on the left of the respective domain and truncating mutations on the right. Mutations on opaque background are associated with liver disease and novel mutations first described in this study are in bold letters. Mutations in italic letters are in-silico predicted to result in truncated or abnormal protein. * denotes a premature stop codon.
An analysis of 67 previously reported patients along with the 44 newly identified patients with MYO5B mutations is consistent with the reported autosomal recessive inheritance of disease (Supporting Information Table S2). Forty-seven of 114 patients (41%) were homozygous for MYO5B mutations, and all but 7 of the remainder were compound-heterozygotes. Seven patients displayed characteristic phenotypes, but only one mutated MYO5B allele was found with the current techniques that address only exonic and flanking intronic gene sequences. The large proportion of homozygous individuals is consistent with the high incidence of MYO5B patients amongst consanguineous marriages.

Missense mutations cluster in the head domain of MYO5B (Table 2), while truncating mutations are more uniformly distributed over the whole MYO5B protein. Missense mutations are found in all subdomains of the myosin head domain. With the exception of the arginine residue at position 401, which is mutated to cysteine in MYO5B-PFIC, and is mutated to histidine in MYO5B-MVID, the MYO5B-PFIC-associated and MYO5B-MIXED-associated missense mutations affect distinct amino acid residues (Figure 2).

Table 2. MYO5B missense mutation per domain distribution.

| Domain       | Head | IQ | Coiled Coil | Tail | Total |
|--------------|------|----|-------------|------|-------|
| Mutation count (%) | 37 (80.4) | 4 (8.7) | 2 (4.4) | 3 (6.5) | 46 (100) |
| Amino acid residues count (%) | 692 (41.9) | 130 (7.9) | 554 (33.5) | 277 (16.7) | 1653 (100) |

3.4. Genotype–Phenotype Correlation and Mutations of Particular Interest

Out of the cohort of all 114 patients, 28 were excluded from our genotype–phenotype correlation due to incomplete phenotyping (Figure 1). The remaining 86 patients were allocated to 3 different groups according to their phenotypes: MYO5B-PFIC, MYO5B-MVID, and MYO5B-MIXED. For 78 out of these 86 cases, the biallelic mutational genotypes were known (Table 3). A detailed list of all patients’ genotypes is provided in Table 4.

The MYO5B mutational and genotypic spectrum in patients with the predominant hepatic phenotype, i.e., MYO5B-PFIC, is different from the spectrum in patients with the predominant intestinal phenotype, MYO5B-MVID (Table 4). Biallelic truncating genotypes are significantly more frequent in the MYO5B-MVID group than in the MYO5B-PFIC group (Fisher’s exact test, \( p < 0.001 \)).

It is noteworthy that each MYO5B genotype was only observed within one phenotype category, implying a strong genotype–phenotype correlation. In addition, only four out of 114 different MYO5B mutations were observed in different phenotype categories, including p.R1016 * and p.M392T. Interestingly, arginine-401 was mutated to cysteine (p.R401C) in one patient with cholestatic liver disease [11], while being mutated to histidine (p.R401H) in two patients with MVID [18].

Table 3. Biallelic MYO5B genotypes (n = 78) observed within 3 phenotype categories.

| Genotype       | Biallelic Missense | Biallelic Truncating | Missense-Truncating | Total |
|----------------|--------------------|----------------------|---------------------|-------|
| Phenotype      | count (%)          | count (%)            | count (%)           | total |
| MYO5B-MVID     | 9 (31.0)           | 15 (51.7)            | 5 (17.3)            | 29 (100) |
| MYO5B-PFIC     | 11 (57.9)          | 0 (0)                | 8 (42.1)            | 19 (100) |
| MYO5B-MIXED    | 12 (40.0)          | 11 (36.7)            | 7 (23.3)            | 30 (100) |
| total          | 32                 | 26                   | 20                  | 78     |
Table 4. MYO5B genotypes \((n = 86)\) associated with MYO5B-PFIC, MYO5B-MVID, MYO5B-MIXED.

| MYO5B-PFIC | MYO5B-MVID | MYO5B-MIXED |
|------------|------------|-------------|
| **Study ID** | **cDNA** | **Protein** | **Class** | **Study ID** | **cDNA** | **Protein** | **Class** | **Study ID** | **cDNA** | **Protein** | **Class** |
| this study | 10D2367 | c.1669G > T | p.V557L | Mis Mis this study 18D1383 | c.2014A > T | p.K672 * | Tru Tru this study 18D4596 | c.1247T > A | Mis Mis |
| this study | 11D1388 | c.1669G > T | p.V557L | Mis Mis this study 10D0098 | c.3190C > A | p.K672 * | Tru Tru this study 10D0028 | c.1475A > G | Mis Mis |
| [26] | pat 1 | c.274C > T | p.R92C | Mis Mis this study 16D2984 | c.4399C > T | p.Q1467 * | Tru Tru this study 11D2081 | c.1966C > T | Mis Mis |
| [26] | pat 3 | c.2470C > T | p.R824C | Mis Mis this study 15D1631 | c.1322-2A > G | Tru Tru this study 12D1383 | c.1247T > A | Mis Mis |
| [11] | pat 4 | c.3237G > C | p.Q1079H | Mis Mis this study 09D0802 | p.R1016 * | Tru Tru | Mis Mis |
| [11] | pat 5 | c.7976C > T | p.C266R | Mis Mis this study 10D0875 | p.R799W | Tru Tru | Mis Mis |
| [11] | pat 6 | c.1748G > A | p.S83N | Mis Mis this study 11D0567 | c.502G > A | Tru Tru | Mis Mis |
| [11] | pat 10 | c.2470C > T | p.R824C | Mis Mis this study 11D1811 | c.483 + 1G > A | Tru Tru | Mis Mis |
| [10] | pat 1 | c.274C > T | p.R292C | Mis Mis this study 12D2180 | c.881C > T | p.M17 * | Tru Tru | Mis Mis |
| [10] | pat 2 | c.1499T > C | p.I500T | Mis Mis this study 17D2697 | c.1582C > T | Tru Tru | Mis Mis |
| [10] | pat 3 | c.356A > G | p.Y119C | Mis Mis this study 18D0872 | p.R1016 * | Tru Tru | Mis Mis |
| this study | 14D0856 | c.274C > T | p.R292C | Mis Tru this study GOSH 1 | c.1087C > T | Tru Tru | Mis Mis |
| this study | 14D0857 | c.437C > T | p.Q1375 | Mis Tru this study GOSH 2 | c.502G > A | Tru Tru | Mis Mis |
| [11] | pat 8 | c.3046C > T | p.R1016 * | Mis Tru | 114 | c.3544_3545del | Tru Tru | Mis Mis |
| [11] | pat 9 | c.3046C > T | p.R1016 * | Mis Tru | 115 | c.1110_1111del | Tru Tru | Mis Mis |
| [26] | pat 2 | c.274C > T | p.R92C | Mis Mis this study 14D0669 | c.1582C > T | Tru Tru | Mis Mis |
| [10] | pat 4 | c.2395C > T | p.R284C | Mis Mis this study 18D4787 | c.1087C > T | Tru Tru | Mis Mis |
| [10] | pat 5 | c.2395C > T | p.R92C | Mis Mis this study 10D1884 | c.1582C > T | Tru Tru | Mis Mis |
### Table 4. Cont.

| Study          | ID          | cDNA          | Protein Class | Study          | ID          | cDNA          | Protein Class | Study          | ID          | cDNA          | Protein Class | Study          | ID          | cDNA          | Protein Class |
|----------------|-------------|---------------|---------------|----------------|-------------|---------------|---------------|----------------|-------------|---------------|---------------|----------------|-------------|---------------|---------------|
| [11] pat 3     | [11] pat 1  | c.1021C > T   | p.R401C       | [11] pat 2     | [11] pat 7  | c.3538-1G > A | p.R697Cfs * 74 | [11] pat 3     | [11] pat 7  | c.2090delC   | p.E82K        | [11] pat 3     | [11] pat 7  | c.1201C > T   | p.R401C       |
|                |             | p.c.2414 + 5G > T |               |                |              |               |               |                |              | c.4852 + 11A > G | p.E82K        |                |              | c.1021C > T   |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              | c.1021C > T   |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              | p.Q341*       |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              |               |               |
|                |             |               |               | [11] pat 1     | [11] pat 1   | c.656G > A    | p.M392T        | [11] pat 4     | [11] pat 7  | c.2259-2262dup | p.T686M       | [11] pat 4     | [11] pat 7  | c.1708dup     | p.T686M       |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              | p.K783       |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              | p.E82K        |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              | p.E82K        |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              | p.E82K        |               |
|                |             |               |               |                |              |               |               |                |              |               |               |                |              |               |               |
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§ Deletes the intron-26 acceptor splice-site; predicts exclusion of exon 27 and a protein with a 22-amino-acid frameshift. & Weakens the motif of the exon 19-intron 19 donor splice-site; usage of an upstream splice-site is predicted, with the consequence of exclusion of the terminal 27 bp of exon 19 and a 9-amino-acid frameshift deletion of the protein. $ No in-silico splicing effect predicted; phase of the genotype was not determined, both identified variants might reside on the same allele with an unknown variant on the other allele that might allow for protein expression. Red font: Mutations that appear in more than one group of phenotypes. Mis Mis, biallelic missense mutations; Mis Tru, compound-heterozygous for missense and truncating mutation, Tru Tru, biallelic truncating mutations; ?, mutation on one allele not identified or consequence ambiguous.
MYO5B-PFIC was reported in 27 patients, of whom 75% developed the disorder within the first year of life. Twenty-eight patients were diagnosed with MYO5B-MIXED. However, exact differentiation between TPN-associated liver disease and MYO5B-PFIC cannot be clearly made in every case.

Few mutations were seen in more than two families. The p.P660L mutation represents a known founder mutation among Navajo Indian patients with MVID and cholestasis, i.e., MYO5B-MIXED [25], and was counted only once in this study. The p.R1016 * mutation was the most common mutation in our study and was present in 10 disease alleles; it was identified in patients of Han Chinese, Anglo-American, and Hispanic ancestry. The corresponding DNA point mutation affects a CpG island, which might be prone to independent mutational events. This mutation was present in homozygous state in a single patient and compound-heterozygous with an exon 2 deletion in another patient, both times associated with MYO5B-MVID. Taking p.R1016 * mutation as an example of the large number of early premature stop codons observed in MYO5B, this mutation is expected to trigger nonsense-mediated mRNA decay and to abolish protein production.

It is of interest that the p.R1016 * mutation was seen in compound-heterozygous state with a p.M392T mutation in two unrelated patients with a notable phenotype: clinical and histopathological characteristics of MVID were documented in the first years of life in both patients, but MVID resolved completely during childhood, whereas MYO5B-PFIC manifested subsequently at ages of 7 and 15 years in these patients. The former patient received a liver transplant for cholestatic liver disease at age 11 years, the latter received an endogenous internal diversion. Both compound-heterozygotes were on a normal diet at last contact at 11 and 24 years of age, respectively; the older patient has been reported twice [26,32], and the genetic testing was performed in our laboratory. This transient-MVID and late-MYO5B-PFIC phenotype was encountered in two other patients with distinguishable genotypes, p.E82K/p.E82K and p.H81/p.Q1600 *. Regarding genotype p.E82K/p.E82K, two siblings of 9 and 5 years were also diagnosed with MVID postnatally, and MVID resolved completely; so far, no hepatic disease occurred.

Mutation p.R656C, predicted to be functionally damaging by all in-silico tools, is a candidate founder mutation in Turkish and Arab patients. In 3 patients from the United Arab Emirates, the MYO5B p.R656C allele harbored an additional p.P1615L mutation on the same allele. Three homozygotes for p. R656C presented with MYO5B-MIXED and were alive at ages 8, 12, and 17 years. The youngest p.R656C homozygote died at age 3.5 years, and chronic cholestatic liver involvement was not recorded but cannot be excluded. Two compound-heterozygotes for p.R656C and an early truncating mutation, p.K672*, had presented with MYO5B-MVID until their deaths at 7 and 9 months of age.

4. Discussion

Biallelic MYO5B mutations have emerged as the major cause of MVID since 2008 [17,18, 23–25]. Currently, MVID represents one disease entity among approximately 50 monogenic disorders that are termed congenital diarrhea and enteropathies (CODEs). CODEs are typically associated with persistent and severe diarrhea presenting in the first weeks of life and with feeding intolerance and malabsorption [34]. CODEs require significant dietary and therapeutic interventions, including specialized formulas or PN to sustain appropriate growth, electrolyte, and nutrient balance. MVID was recognized as an entity on clinical and histopathological grounds in 1978 [2]. Marked but variably severe hepatic involvement was recognized in a significant percentage of MVID patients thereafter, which is considered at least in part to represent a consequence of TPN and of a disrupted enterohepatic circulation in young children [4,35]. Hepatic involvement most often manifested in the first year of life and was similar to low-GGT PFIC. Recently, four studies reported a small series of patients with PFIC who were negative for mutations in the known PFIC genes and harbored biallelic MYO5B mutations, or pathogenic monoallelic MYO5B mutations, where a mutation on the second allele had not been detected with the current diagnostic techniques [9–11,26].
Since our first reporting of MYO5B mutations in MVID, we identified MYO5B mutations in 67 patients. In this study, we compile genetic and clinical findings from 44 new patients with MYO5B mutations and from 47 patients with clinical and genetical findings from the literature. Our review of a total of 114 patients demonstrates that there are indeed 3 distinctive phenotypes associated with MYO5B genotypes: MYO5B-PFIC, pure MYO5B-MVID, and MYO5B-MIXED. Our study shows that there is a nearly complete genotype–phenotype correlation, i.e., each MYO5B genotype belongs to one of the above classes, as was suggested in previous studies with fewer patients with known genotypes and in functional studies [7,11,15].

How do different biallelic MYO5B genotypes cause isolated hepatic or intestinal disease, or concomitant hepatic and intestinal disease? An evaluation of our comprehensive cohort of patients with known MYO5B genotypes and their phenotypes is consistent with the following hypotheses [11,15,21]:

1. Biallelic mutations that predict nonsense-mediated mRNA decay are always associated with MYO5B-MVID. This indicates that the MVID phenotype is caused by a loss-of-function of MYO5B, which causes clinical symptoms via disrupted enterocyte polarization [1,17,22,36]. The small number of missense mutations associated with MYO5B-MVID might all lead to misfolding and MYO5B degradation, also resulting in lack of MYO5B protein. The loss of MYO5B motor function alone does not cause liver disease.

2. Contrastingly, a distinct set of missense mutations is identified in patients with MYO5B-PFIC, and these mutant MYO5B proteins are expressed, as has been shown for two of these missense mutations, p.C266R, and p.S158F [11,15,37]. Apparently, intestinal MYO5B function of these mutant proteins is preserved, at least to the degree in that such biallelic missense mutations allow sufficient degree of enterocytic function to maintain intestinal autonomy. However, displacement of bile canicular transporters to the cytoplasm of hepatocytes was shown in liver biopsies of patients with MVID presenting with cholestasis and homozygous missense mutations p. P660L [37] as well as in patients with MYO5B-PFIC with the homozygous missense mutations p. C266R [11] and p.S158F mutation [15]. Mutagenesis experiments showed that the disrupting effect of PFIC-associated MYO5B motor domain mutants on the localization of canalicular proteins was critically dependent on their preserved ability to interact with active RAB11a, as loss of MYO5B did not affect RAB11-dependent vesicle trafficking [15].

3. The conclusion from recent in vitro and in vivo studies is indirectly supported by our genotype–phenotype correlation, which shows that MYO5B is not required for the correct localization of hepatic canalicular proteins. It was shown that PFIC-associated MYO5B mutants require active RAB11a for their disruptive effect on canalicular protein localization, and this previous study indicated a direct and simple explanation for the observed genotype–phenotype correlation in patients with MYO5B mutations.

4. Patients with MYO5B-MIXED most often present with a missense or late-truncating mutation in trans with a loss-of-function mutation in MYO5B. In these instances, the presence of a single mutant expressed protein cannot prevent MVID development in enterocytes and causes cholestatic liver disease by putatively interfering with RAB11- or RAB8-dependent [38] processes.

The correlation between loss-of-function and expressed mutant proteins is displayed schematically in Figure 3.
Correct localization of hepatic canalicular proteins. It was shown that PFIC-associated MYO5B mutants require active RAB11a for their disruptive effect on canalicular protein localization, and this previous study indicated a direct and simple explanation for the observed genotype–phenotype correlation in patients with MYO5B mutations.

Patients with MYO5B-MIXED most often present with a missense or late-truncating mutation in trans with a loss-of-function mutation in MYO5B. In these instances, the presence of a single mutant expressed protein cannot prevent MVID development in enterocytes and causes cholestatic liver disease by putatively interfering with RAB11- or RAB8-dependent processes.

The correlation between loss-of-function and expressed mutant proteins is displayed schematically in Figure 3.

Very rarely do patients present with transient MVID in the first years of life and develop liver disease in childhood. Interestingly, this appears to occur only with certain MYO5B genotypes. It might be speculated that MVID is present at times where nutritional load per surface area and intestinal growth are highest in the first years of life. Liver disease might ensue with increasing load of the enterohepatic circulation. Along similar lines, the intestinal microbiome is a relevant modulator of bile acids and the enterohepatic circulation, via bile acid modification, and only fully matures during the first years of life [39]. This then might lead to the misbalance of a fragile hepatocytic homeostasis and result in MYO5B-PFIC. It is known that therapeutic intervention can change the MYO5B-related phenotype. For example, MYO5B-MVID patients develop MYO5B-PFIC following a bowel transplantation, supposedly due to an altered enterohepatic circulation [7].

The majority of MYO5B mutations were observed in individual patients or families, highlighting the need for a complete MYO5B sequence analysis on suspicion of MYO5B-MVID, MYO5B-PFIC, and MYO5B-MIXED. Such an analysis needs to incorporate testing for copy-number-variants (CNVs) as intragenic exon deletions constitute 4 of 114 distinct disease variants and represent 5 out of 230 disease alleles. CNV calling can be included in any next-generation-sequencing pipeline suited to analyzing gene panels for congenital diarrheas and for pediatric cholestatic disorders.

5. Conclusions

Our data support the concept that (1) MVID results from the loss of MYO5B function in enterocytes, (2) that a complete lack of protein and greatly truncated MYO5B proteins do not cause primary liver disease, (3) that solitary primary cholestatic liver disease

Figure 3. Schematic overview of enterocytic and hepatocytic phenotypes observed upon mutations in myosin Vb (MYO5B). (A) healthy enterocytes and hepatocytes establish proper epithelial differentiation. Apical proteins (e.g., transmembrane transporters and enzymes; red) localize at the apical brush border in enterocytes or bile canalicular membrane in hepatocytes. (B) lack of MYO5B results in MYO5B-MVID: disrupted apical brush border, subapical “secretory” granules (blue), mislocalization of apical proteins (red) and microvillus inclusions. Hepatocyte function is unaffected. (C) Expressed mutated MYO5B results in varying degree of disease-severity and enterocytic phenotypes but leads to mislocalisation of apical bile transporters and cholestasis in hepatocytes.
results from the expression of mutant MYO5B proteins that cause aberrant protein–protein interactions in hepatocytes, (4) and that expressed non-functional MYO5B protein causes both intestinal and hepatic disease.

The existence of an experimental mouse model for MYO5B disease will provide a useful platform for testing mechanistic hypotheses [22,40,41].

Supplementary Materials: The following are available online at https://www.mdpi.com/2077-0383/10/3/481/s1, Table S1: Detailed information on all 114 individual patients.

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References

1. Vogel, G.F.; Janecke, A.R.; Krainer, I.M.; Gutleben, K.; Witting, B.; Mitton, S.G.; Mansour, S.; Ballauf, A.; Roland, J.T.; Engevik, A.C.; et al. Abnormal Rab11-Rab8-vesicles cluster in enterocytes of patients with microvillus inclusion disease. Traffic 2017, 18, 453–464. [CrossRef]
2. Davidson, G.P.; Cutz, E.; Hamilton, J.R.; Gall, D.G. Familial enteropathy: A syndrome of protracted diarrhea from birth, failure to thrive, and hypoplastic villus atrophy. Gastroenterology 1978, 75, 783–790. [CrossRef]
3. Cutz, E.; Rhoads, J.M.; Drumm, B.; Sherman, P.M.; Durie, P.R.; Forstner, G.G. Microvillus inclusion disease: An inherited defect of brush-border assembly and differentiation. N. Engl. J. Med. 1989, 320, 646–651. [CrossRef]
4. Ruemmele, F.M.; Schmitz, J.; Goulet, O. Microvillus inclusion disease (microvilli atrophy). Orphanet J. Rare Dis. 2006, 1, 22. [CrossRef]
5. Wiegerinck, C.L.; Janecke, A.R.; Schneeberger, K.; Vogel, G.F.; van Haafsen-Visser, D.Y.; Escher, J.C.; Adam, R.; Thoni, C.E.; Pfaller, K.; Jordan, A.J.; et al. Loss of syntaxin 3 causes variant microvillus inclusion disease. Gastroenterology 2014, 147, 65–68. [CrossRef]
6. Stepensky, P.; Bartram, J.; Barth, T.F.; Lehmburg, K.; Walther, P.; Amann, K.; Philips, A.D.; Beringer, O.; Zur Stadt, U.; Schulz, A.; et al. Persistent defective membrane trafficking in epithelial cells of patients with familial hemophagocytic lymphohistiocytosis type 5 due to STXBP2/MUNC18-2 mutations. Pediatr. Blood Cancer 2013, 60, 1215–1222. [CrossRef] [PubMed]
7. Dheke, H.S.; Pylypenko, O.; Overeem, A.W.; Ferreira, R.J.; van der Velde, K.J.; Rings, E.; Posovszky, C.; Swertz, M.A.; Houdusse, A.; van IJzendoom, S.C.D. MYO5B, STX3, and STXBP2 mutations reveal a common disease mechanism that unifies a subset of congenital diarrheal disorders: A mutation update. Hum. Mutat. 2018, 39, 333–344. [CrossRef] [PubMed]
8. Vogel, G.F.; van Rijn, J.M.; Krainer, I.M.; Janecke, A.R.; Posovszky, C.; Cohen, M.; Searle, C.; Jantchou, P.; Escher, J.C.; Patey, N.; et al. Disrupted apical exocytosis of cargo vesicles causes enteropathy in FHL5 patients with Munc18-2 mutations. JCI Insight 2017, 2. [CrossRef] [PubMed]
9. Girard, M.; Lacaille, F.; Verkarre, V.; Mategot, R.; Feldmann, G.; Grodet, A.; Sauvat, F.; Irtan, S.; Davit-Spraul, A.; Jacquemin, E.; et al. MYO5B and bile salt export pump contribute to cholestatic liver disorder in microvillus inclusion disease. Hepatology 2014, 60, 301–310. [CrossRef] [PubMed]
10. Gonzales, E.; Taylor, S.A.; Davit-Spraul, A.; Thebaut, A.; Thomassinn, N.; Guettier, C.; Whittington, P.F.; Jacquemin, E. MYO5B mutations cause cholestasis with normal serum gamma-glutamyl transferase activity in children without microvillus inclusion disease. *Hepatology* 2017, 65, 164–173. [CrossRef] [PubMed]

11. Qiu, Y.L.; Gong, J.Y.; Feng, J.Y.; Wang, R.X.; Han, J.; Liu, T.; Lu, Y.; Li, L.T.; Zhang, M.H.; Sheps, J.A.; et al. Defects in myosin VB are associated with a spectrum of previously undiagnosed low gamma-glutamyltransferase cholestasis. *Hepatology* 2017, 65, 1655–1669. [CrossRef] [PubMed]

12. Bull, L.N.; Carlton, V.E.; Stricker, N.L.; Baharloo, S.; DeYoung, J.A.; Freimer, N.B.; Magid, M.S.; Kahn, E.; Markowitz, J.; DiCarlo, F.J.; et al. Genetic and morphological findings in progressive familial intrahepatic cholestasis (Byler disease [PFIC-1] and Byler syndrome): Evidence for heterogeneity. *Hepatology* 1997, 26, 155–164. [CrossRef] [PubMed]

13. Sambrota, M.; Straunzieks, S.; Papoulis, E.; Rushton, P.; Clark, B.E.; Parry, D.A.; Logan, C.V.; Newbury, L.J.; Kamath, B.M.; Ling, S.; et al. Mutations in TJP2 cause progressive cholestatic liver disease. *Nat. Genet.* 2014, 46, 326–328. [CrossRef] [PubMed]

14. Droge, C.; Bonus, M.; Baumann, U.; Klindt, C.; Lainka, E.; Kathemann, S.; Brinkert, F.; Grabhorn, E.; Pfister, E.D.; Wenning, D.; et al. Sequencing of FIC1, BSEP and MDR3 in a large cohort of patients with cholestatics revealed a high number of different genetic variants. *J. Hepatol.* 2017, 67, 1253–1264. [CrossRef]

15. Overeem, A.W.; Li, Q.; Qiu, Y.L.; Carton-Garcia, F.; Leng, C.; Klappe, K.; Drunker, J.; Hsiao, N.H.; Wang, J.S.; Arango, D.; et al. A Molecular Mechanism Underlying Genotype-Specific Intrahepatic Cholestasis Resulting From MYO5B Mutations. *Hepatology* 2020, 72, 213–229. [CrossRef]

16. van der Velde, K.J.; Dhekne, H.S.; Swertz, M.A.; Sirigu, S.; Ropars, V.; Vinke, P.C.; Rengaw, T.; van den Akker, P.C.; Rings, E.H.; Houdusse, A.; et al. An overview and online registry of microvillus inclusion disease patients and their MYO5B mutations. *Hum. Mutat.* 2013, 34, 1997–1605. [CrossRef]

17. Muller, T.; Hess, M.W.; Schiefermeier, N.; Pfalder, K.; Ebner, H.L.; Heinz-Erian, P.; Pomstingl, H.; Partsch, J.; Rollinghoff, B.; Kohler, H.; et al. MYO5B mutations cause microvillus inclusion disease and disrupt epithelial cell polarity. *Nat. Genet.* 2008, 40, 1163–1165. [CrossRef]

18. Ruemmele, F.M.; Muller, T.; Schiefermeier, N.; Ebner, H.L.; Lechner, S.; Pfalder, K.; Thoni, C.E.; Goulet, O.; Lacaille, F.; Schmitz, J.; et al. Loss-of-function of MYO5B is the main cause of microvillus inclusion disease: 15 novel mutations and a CaCo-2 RNAi cell model. *Hum. Mutat.* 2010, 31, 544–552. [CrossRef]

19. Klee, K.M.C.; Janecke, A.R.; Civan, H.A.; Rosipal, S.; Heinz-Erian, P.; Huber, L.A.; Muller, T.; Vogel, G.F. AP1S1 missense mutations cause a congenital enteropathy via an epithelial barrier defect. *Hum. Genet.* 2020, 139, 1247–1259. [CrossRef]

20. Ghosh, R.; Oak, N.; Pion, S.E. Evaluation of in silico algorithms for use with ACMG/AMP clinical variant interpretation guidelines. *Genome Biol.* 2017, 18, 225. [CrossRef]

21. van Ijzendoorn, S.C.D.; Li, Q.; Qiu, Y.L.; Wang, J.S.; Overeem, A.W. Unequal effects of MYO5B mutations in liver and intestine determine the clinical presentation of low-GGT cholestasis. *Hepatology* 2020. [CrossRef]

22. Schneebberger, K.; Vogel, G.F.; Teunissen, H.; van Ommen, D.D.; Begthel, H.; El Bouazzaoui, L.; van Vugt, A.H.; Beeekman, J.M.; Klumperman, J.; Muller, T.; et al. An inducible mouse model for microvillus inclusion disease reveals a role for myosin Vb in apical and basolateral trafficking. *Proc. Natl. Acad. Sci. USA* 2015, 112, 12408–12413. [CrossRef] [PubMed]

23. Szper, A.M.; Golachowska, M.R.; Bruinenberg, M.; Prekeris, R.; Thumnissen, A.M.; Karrenbeld, A.; Dijkstra, G.; Hoekstra, D.; Mercer, D.; Ksiazyk, J.; et al. Functional characterization of mutations in the myosin Vb gene associated with microvillus inclusion disease. *J. Pediatr. Gastroenterol. Nutr.* 2011, 52, 307–313. [CrossRef] [PubMed]

24. Golachowska, M.R.; van Dael, C.M.; Keuning, H.; Karrenbeld, A.; Gijbers, C.F.; Benninga, M.A.; Rings, E.H.; van Ijzendoorn, S.C.D. MYO5B mutations in patients with microvillus inclusion disease presenting with transient renal Fancconi syndrome. *J. Pediatr. Gastroenterol. Nutr.* 2012, 54, 491–498. [CrossRef] [PubMed]

25. Erickson, R.P.; Larson-Thome, K.; Valenzuela, R.K.; Whitaker, S.E.; Shub, M.D. Navajo microvillus inclusion disease is due to a mutation in MYO5B. *Am. J. Med. Genet.* 2008, 146A, 3117–3119. [CrossRef]

26. Cockar, I.; Foskett, P.; Strauntzieks, S.; Clinch, Y.; Fustok, J.; Rahman, O.; Sutton, H.; Mtegha, M.; Fessato, S.; Kontaki, E.; et al. Mutations in Myosin 5B in Children With Early-onset Cholestasis. *J. Pediatr. Gastroenterol. Nutr.* 2020, 71, 184–188. [CrossRef]

27. Comegna, M.; Amato, F.; Liguori, R.; Berni Canani, R.; Spagnuolo, M.I.; Morroni, M.; Guarino, A.; Castaldo, G. Two cases of microvillus inclusion disease caused by novel mutations in MYO5B gene. *Clin. Case Rep.* 2018, 6, 2451–2456. [CrossRef]

28. Perry, A.; Bensallah, H.; Martinez-Vinson, C.; Berrebi, D.; Arbeille, B.; Salomon, J.; Goulet, O.; Marinier, E.; Drunat, S.; Samson-Bouma, M.E.; et al. Microvillus atrophy: Atypical presentations. *J. Pediatr. Gastroenterol. Nutr.* 2014, 59, 779–785. [CrossRef]

29. Chen, C.P.; Chiang, M.C.; Wang, T.H.; Hsieh, C.; Chang, S.D.; Tsai, F.J.; Wang, C.N.; Chern, S.R.; Wang, W. Microvillus inclusion disease: Prenatal ultrasound findings, molecular diagnosis and genetic counseling of congenital diarrhea. *Taiwan. J. Obstet. Gynecol.* 2010, 49, 487–494. [CrossRef]

30. Mao, M.; Guo, L.; Zhang, Z.; Wang, B.; Huang, S.; Song, Y.; Chen, F.; Wen, W. Phenotypic and genetic analysis of a family affected with microvillus inclusion disease. *Zhonghua Yi Xue Yi Chuan Xue Za Zhi* 2016, 33, 792–796. [CrossRef]

31. Fernandez Caamano, B.; Quiles Blanco, M.J.; Fernandez Tome, L.; Burgos Lizaldez, E.; Sarria Oses, J.; Molina Arias, M.; Prieto Bozano, G. Intestinal failure and transplantation in microvillus inclusion disease. *An. Pediatr. (Barc.)* 2015, 83, 160–165. [CrossRef] [PubMed]

32. Croft, N.M.; Howatson, A.G.; Ling, S.C.; Nairn, L.; Evans, T.J.; Weaver, L.T. Microvillus inclusion disease: An evolving condition. *J. Pediatr. Gastroenterol. Nutr.* 2000, 30, 185–189. [CrossRef] [PubMed]
33. Sadiq, M.; Choudry, O.; Kashyap, A.K.; Velazquez, D.M. Congenital diarrhea in a newborn infant: A case report. World J. Clin. Pediatr. 2019, 8, 43–48. [CrossRef]
34. Thiagarajah, J.R.; Kamin, D.S.; Acra, S.; Goldsmith, J.D.; Roland, J.T.; Lencer, W.I.; Muise, A.M.; Goldenring, J.R.; Avitzur, Y.; Martin, M.G.; et al. Advances in Evaluation of Chronic Diarrhea in Infants. Gastroenterology 2018, 154, 2045–2059.e6. [CrossRef]
35. Goulet, O.; Ruemmele, F. Causes and management of intestinal failure in children. Gastroenterology 2006, 130, S16–S28. [CrossRef] [PubMed]
36. Vogel, G.F.; Klee, K.M.; Janecke, A.R.; Muller, T.; Hess, M.W.; Huber, L.A. Cargo-selective apical exocytosis in epithelial cells is conducted by Myo5B, Slp4a, Vamp7, and Syntaxin 3. J. Cell Biol. 2015, 211, 587–604. [CrossRef]
37. Schlegel, C.; Weis, V.G.; Knowles, B.C.; Lapiere, L.A.; Martin, M.G.; Dickman, P.; Goldenring, J.R.; Shub, M.D. Apical Membrane Alterations in Non-intestinal Organs in Microvillus Inclusion Disease. Dig. Dis. Sci. 2018, 63, 356–365. [CrossRef] [PubMed]
38. Roland, J.T.; Bryant, D.M.; Datta, A.; Itzen, A.; Mostov, K.E.; Goldenring, J.R. Rab GTPase-Myo5B complexes control membrane recycling and epithelial polarization. Proc. Natl. Acad. Sci. USA 2011, 108, 2789–2794. [CrossRef]
39. Ridlon, J.M.; Kang, D.J.; Hylemon, P.B.; Bajaj, J.S. Bile acids and the gut microbiome. Curr. Opin. Gastroenterol. 2014, 30, 332–338. [CrossRef]
40. Carton-Garcia, F.; Overeem, A.W.; Nieto, R.; Bazzocco, S.; Dopeso, H.; Macaya, I.; Bilic, J.; Landolfi, S.; Hernandez-Losa, J.; Schwartz, S., Jr.; et al. Myo5b knockout mice as a model of microvillus inclusion disease. Sci. Rep. 2015, 5, 12312. [CrossRef]
41. Weis, V.G.; Knowles, B.C.; Choi, E.; Goldstein, A.E.; Williams, J.A.; Manning, E.H.; Roland, J.T.; Lapiere, L.A.; Goldenring, J.R. Loss of MYO5B in mice recapitulates Microvillus Inclusion Disease and reveals an apical trafficking pathway distinct to neonatal duodenum. Cell. Mol. Gastroenterol. Hepatol. 2016, 2, 131–157. [CrossRef]