ORIGINAL ARTICLE



Exome sequencing in patient-parent trios suggests new candidate genes for early-onset primary sclerosing cholangitis

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Abstract

BACKGROUND & AIMS: Primary sclerosing cholangitis (PSC) is a rare bile duct disease strongly associated with inflammatory bowel disease (IBD). Whole-exome sequencing (WES) has contributed to understanding the molecular basis of very early-onset IBD, but rare protein-altering genetic variants have not been identified for early-onset PSC. We performed WES in patients diagnosed with PSC ≤ 12 years to investigate the contribution of rare genetic variants to early-onset PSC.

METHODS: In this multicentre study, WES was performed on 87 DNA samples from 29 patient-parent trios with early-onset PSC. We selected rare (minor allele frequency < 2%) coding and splice-site variants that matched recessive (homozygous and compound heterozygous variants) and dominant (de novo) inheritance in the index patients. Variant

Abbreviations: GWAS, genome-wide association studies; HLA, human leukocyte antigen; IBD, inflammatory bowel disease; MHC, major histocompatibility complex; PSC, primary sclerosing cholangitis: WES, whole-exome sequencing.

Patrick F van Rheenen, Cleo C van Diemen are shared last authors.

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pathogenicity was predicted by an in-house developed algorithm (GAVIN), and PSC-relevant variants were selected using gene expression data and gene function.

RESULTS: In 22 of 29 trios we identified at least 1 possibly pathogenic variant. We prioritized 36 genes, harbouring a total of 54 variants with predicted pathogenic effects. In 18 genes, we identified 36 compound heterozygous variants, whereas in the other 18 genes we identified 18 de novo variants. Twelve of 36 candidate risk genes are known to play a role in transmembrane transport, adaptive and innate immunity, and epithelial barrier function.

CONCLUSIONS: The 36 candidate genes for early-onset PSC need further verification in other patient cohorts and evaluation of gene function before a causal role can be attributed to its variants.

KEYWORDS

genetic, inflammatory bowel disease, sclerosing cholangitis

Key points

It is rare to diagnose a child with primary sclerosing cholangitis (PSC) before its 13th birthday. We screened the portion of the DNA that codes for proteins in 29 young PSC children and their biological parents and identified 54 rare genetic variants in 36 genes with an assumed deleterious effect on protein function. Whether these variants play a part in the aetiology of PSC will require further verification.

1 | INTRODUCTION

Primary sclerosing cholangitis (PSC) is a rare chronic cholestatic disease characterized by progressive inflammation and obliterative fibrosis of the intra- and extrahepatic bile ducts. Disease progression is inevitable in the majority of PSC patients, with the development of biliary cirrhosis and portal hypertension requiring repeated endoscopic procedures. Liver transplantation is the only curative treatment option, but the disease recurs in 20%-25% of transplanted patients. Cholangiocarcinoma and colorectal cancer are feared complications in PSC and the most common causes of death.

There is a strong relation between PSC and inflammatory bowel disease (IBD). Patients who initially present with isolated PSC may go on to develop IBD years later. In adult-onset disease, approximately two thirds of patients with PSC have concurrent IBD. The co-occurrence of PSC and IBD is higher in children than in adults, varying from 76% to 97%. The pathogenesis of PSC is largely unknown, but there is a strong genetic component. Genome-wide association studies (GWAS) in adult-onset PSC, carried out by the International PSC Study Group, identified 23 risk loci and 9 suggestive findings. Not surprisingly, considering the large clinical overlap, the human leukocyte antigen (HLA) locus is by far the strongest signal in GWAS in both PSC and IBD. 9,10

Next to the common variants found by GWAS that so far explain <10% of PSC susceptibility, rare variants with large monogenic effect size may play a role in the onset of PSC. These variants are so rare in

allele frequency (many of them private variants) that their genetic signals are hard to detect by GWAS. In contrast, whole-exome sequencing (WES) in patients with extreme phenotypes, such as early-onset IBD, has led to the identification of a growing number of rare monogenic disorders presenting with IBD-like intestinal inflammation. 11,12 Additionally, a rare monogenic variant (a loss of function mutation in doublecortin domain containing protein 2 (DCDC2) has recently been identified in a PSC-like disorder called neonatal sclerosing cholangitis. 13 Although neonatal sclerosing cholangitis is a different entity than PSC, histological similarities of the cholangiocytes of these patients lacking primary cilia suggest that some of the underlying pathogenic mechanisms could be shared between the two diseases. Likewise, we expect that a monogenic or oligogenic inheritance pattern may play a role in a subset of patients with early-onset PSC as well. We therefore performed WES in a Dutch cohort of patients with early-onset PSC and their parents to identify rare, but possibly causative genetic variants.

2 | METHODS

2.1 | Study design, participants and setting

In this multicentre parent-offspring study, we collected DNA from PSC patients with disease-onset prior to their 13th birthday and from their biological parents. PSC diagnosis was confirmed by cholangiography (presence of multifocal strictures, focal dilatation or beading of the

biliary tree) or liver histology (presence of bile duct damage, onionskinned peri-ductal fibrosis, inflammation, portal oedema or fibrosis, ductopenia, ductular proliferation or cholestasis), or both. Patients with sclerosing cholangitis as a result of secondary causes such as surgery, trauma, cancer or infection were excluded from participation.

Patients were recruited in five tertiary care hospitals in the Netherlands-University Medical Center Groningen (UMCG, a referral paediatric liver transplant centre), Erasmus University Medical Center-Sophia Children's Hospital, VU University Medical Center, Amsterdam University Medical Center-Emma Children's Hospital, University Medical Center Utrecht-Wilhelmina Children's Hospitaland one large general teaching hospital, the Isala Hospital. Eligible patients were those regularly attending the (paediatric) gastroenterology and hepatology clinics as part of standard care. After informed consent was given, the following information was obtained from the local patient records and entered in an online clinical registry using Castor Electronic Data Capture (Amsterdam, the Netherlands): age at PSC diagnosis, findings on cholangiography and/or histology, and follow-up data on medication use and appearance of biliary cirrhosis, portal hypertension or liver transplantation. Between January 2017 and July 2017, blood was collected from patients and volunteering parents for genomic DNA extraction according to standard protocols.

2.2 | Ethical considerations

The Medical Ethical Committee of the UMCG approved the study protocol (METC 2016/289), and secondary approval was obtained from all participating centres. All participating parents and teenagers 12-19 years old gave informed consent prior study inclusion.

2.3 | Whole-exome sequencing

Libraries were prepared using the Illumina Nextera prep kit and hybrid capture (Illumina Rapid Capture Enrichment – 37 Mb target), and sequencing was performed using the Illumina HiSeq 2500 at the Broad Institute of MIT and Harvard. All raw data underwent quality control steps (https://hub.docker.com/r/broadinstitute/gatk/) without any noticeable negative features to achieve 86.06 million high-quality reads per sample with 98.85% of reads aligned, on average, resulting in a coverage of 81% of the target region with a read depth of >30X. Sequence reads were aligned to the human reference genome using Novoalign (http://www.novocraft.com). Next, the Genome Analysis Toolkit of the Broad Institute¹⁴ was used for calling single-nucleotide variant and insertions/deletions.

2.3.1 | Variant annotation

Variants were annotated with SNPEff¹⁵ using publicly available data from Ensembl and Refseq, and with GAVIN, an annotation tool with an algorithm that scores the likely pathogenicity of the variants.¹⁶

Additional annotations at the variant, exon and gene level were obtained from the 1000 Genomes Project (http://www.1000genomes.org); National Heart, Lung and Blood Institute GO Exome Sequencing Project Exome Variant Server (http://evs.gs.washington.edu/EVS); PolyPhen2¹⁷ and the Exome Aggregation Consortium (ExAC, http://exac.broadinstitute.org). To facilitate further discoveries in this fairly narrow field of research, the VCF files of our study are now publicly available in The Groningen Data Catalogue (catalogus.helpdesk@umcg.nl) at (https://groningendatacatalogus.nl/menu/groningendatacatalogue/dataexplorer/details/umcg_collections/aaaac526ldngr6qwh2xc53aaae).

2.3.2 | Variant filtering

We used variants with a sequence coverage of ten or greater. We used a Genomics Data Management System (Alissa Interpret –Agilent technologies) to create a filtering tree specifically designed for this study (see Figure 1).

We performed patient-parent trio analyses. On the variant-level, we selected variants matching recessive (homozygous and compound heterozygous variants) and dominant (de novo) inheritance in the index patients. HLA variants were excluded from this analysis as it is unfeasible to distinguish functional polymorphism from random variation in a patient cohort of modest size. Minor allele frequency (MAF) cut-offs from the GnomAD database (http://gnomad.broad institute.org/) were <2% for recessive variants and <0.01% for dominant (de novo) variants. Variants were selected when they fulfilled the following two criteria: (a) deemed to be coding (missense- and nonsense mutations, frameshift insertions and deletions) or to have

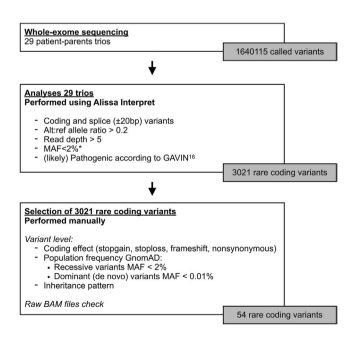


FIGURE 1 Variant selection. * Population databases used: ExAC, GnoMAD, 1000 Genomes project. Abbreviation: MAF, Minor allele frequency

an effect on splicing; and (b) predicted to be (likely) pathogenic according to ${\sf GAVIN.}^{16}$

2.3.3 | Variant categorization

We then searched in Genecards (www.genecards.com), Reactome (www.reactome.org) and OMIM (Online Mendelian Inheritance in Man; (www.omim.org) databases for information about gene function and their involvement in diseases. We prioritized the genes into three categories (Box 1).

2.3.4 | Variant verification and validation

De novo variants were manually checked for coverage and allelic balance in the BAM files. If there was doubt about the validity of the variant, confirmatory Sanger sequencing was performed.

3 | RESULTS

3.1 | Patient and disease characteristics

A total of 29 patient-parent trios were enrolled in this study (see Figure 1). Table 1 shows the characteristics of the affected persons, who were diagnosed with PSC at a median age of 10 years (range: 2-12) and were predominantly male (72%). Twenty-two of 29 patients (76%) had concurrent IBD, with ulcerative colitis significantly more prevalent than Crohn's disease (73% vs 27%). Other autoimmune disorders included celiac disease (n = 1), idiopathic thrombocytopenic purpura (n = 1) and vitiligo (n = 1). None of the parents was known to have liver disease, but three had IBD. The median time (range) between PSC diagnosis and study enrolment was 4 (0-30) years. Biliary complications, including cholangitis or bile duct obstruction, had occurred in one patient (3%), and cirrhosis with portal hypertension had developed in nine (31%). Two patients underwent a liver transplantation after a disease duration of 10 and 11 years, respectively,

BOX 1 Prior	itization of genes
Category 1	Well-known gene function connected with PSC or a similar phenotype (i.e., immunological, inflammatory or bile salt homeostasis).
Category 2	Unknown gene function and not (or rarely) reported in literature, and therefore cannot be excluded from having a potential role in the disease pathogenesis,
Category 3	Well-known gene function not directly associated with the disease phenotype:

and two other patients are currently listed for liver transplantation. One of the cirrhotic patients had experienced bleeding of oesophageal varices and required a transjugular intrahepatic portosystemic shunt procedure.

3.2 | Patient-parent trio analyses of WES data

Figure 1 provides an overview of WES variant selection and prioritization. In 22 of 29 trios, we identified at least 1 candidate pathogenic variant. We identified a total of 54 candidate variants with predicted pathogenic effect, annotated to 36 genes (see Table 2). In 15 trios we identified 36 unique compound heterozygous variants, and in 12 trios we identified 18 unique de novo variants. Twelve of 36 candidate risk genes were assigned to prioritization category 1, 11 to category 2 and 13 to category 3.

The category 1 genes are depicted in Figure 2. The genes *SLC9B1* and *ABCB6* encode for membrane transporter proteins and relate to the '*Transport of glucose and other sugars, bile salts and organic acids, metal ions and amine compounds*' pathway (www.pathcards.genec ards.org). The gene *NOTCH2NLA* regulates the '*Notch Signaling*' pathway (www.pathcards.genecards.org) and *NOTCH2* variants are related to syndromes with cholestatic phenotypes (www.omim.org/entry/600275). The genes *MARCH1*, *PTX4*, *TIRAP*, *ADAM32*, *DDX47*, *USP17L2*, *EHD1 and EDC4* are associated with defects of the immune system. The *CDHR2* gene is involved in the epithelial barrier function.

4 | DISCUSSION

4.1 | Key results

In this study we examined the exomes of patients with early-onset PSC and their biological parents. With this trio-analysis approach we identified 54 rare variants with predicted large effects on protein function in 36 genes. We prioritized 12 candidate risk genes that are most likely to contribute to the development of PSC in patients with early-onset disease based on their presumed role in (auto) immunity pathways, membrane transport (including bile salt homeostasis) and epithelial barrier functioning.

4.2 | Prioritized genes and their possible role in PSC pathogenesis

Previous GWAS studies on the genetics of PSC suggested the autoimmune origin of the disease, with strong associations with the genes encoding for the HLA complex on chromosome 6, along with several susceptibility genes that are critically involved in T-cell function. ^{1,8} Liver tissue from biopsied patients with PSC showed mainly T cells and, to a lesser degree, macrophages and neutrophils in the infiltrates. ¹ We found a de novo stop-gain variant located at the

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	Candidate gene identified [Prioritized genes printed in bold]		,	ABCB6	ADAM32 , MARCH1, PTX4 , PLXDC1		EDC4	PHC2, TNRC18	,	HMCN2	CCN4	ı	DNAH6, EHD1, LAMA2	DNAH7	KRTAP5-1, MCM8	CASKIN2	PPFIA4	CDHR2, DDX47	CALCRL, USP17L2	ANKRD36, DNAH11, NOTCH2NLA	1	JMJD1C, DACT1	TSPYL5	SLC9B1	TRDN	FAM234B	SMCHD1, TIRAP	•	CHST11	KIF4A, ZNF30
Complicated disease course?	Biliary complications [B]; Cirrhosis with portal hypertension [C]; liver transplantation [LT]		,	1			,	C, LT		1	O	O	1	O	1	1	,	ı	,		1	U	C, LT	1	C, LT		В		S	O
	1st degree relative with IBD or PSC?	N _o	IBD (father)	No	°N N	IBD (mother)	N _o	°N	°N °N	oN N	o _N	No	o _N	oN N	°N °N	oN N	o _N	No	N _o	No	o _N	No	°N N	N _o	o _N	°N	No	IBD (father)	No	°N ON
IBD present?	[No, UC or CD]	8	nc	nc	° Z	nc	No	No	oN	No	СО	No	СО	No	OC	CD	nc	nc	СО	nc	nc	nc	nc	nc	nc	nc	nc	UC	СО	nc
	Liver histology											\boxtimes		\boxtimes		\boxtimes								\boxtimes						
PSC confirmed by	ERCP											\boxtimes		\boxtimes									\boxtimes							
PSC conf	MRCP						\boxtimes	\boxtimes		\boxtimes														\boxtimes		\boxtimes			\boxtimes	
	Age at PSC diagnosis (in years)	11	10	7	œ	11	12	œ	11	12	10	7	6	7	12	12	10	5	2	11	10	11	11	10	5	œ	7	12	12	9
	Patient identification number	1	2	ო	4	2	9	7	80	6	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29

 TABLE 2
 Complete list of 36 candidate risk genes for early-onset PSC from 22 patient-parent trios

(Continues)

Protein function	Binds heme and porphyrins and functions in their ATP-dependent uptake into the	mitochondria. ^{20,2,1} Mutations in this gene underlie familial pseudohyperkalemia (OMIM 609 153) and dyschromatosis universalis hereditarian (OMIM 615 402).	This gene encodes a member of the disintegrin family of membrane-anchored proteins that	play a role in diverse biological processes such as brain development, fertilization, tumour development and inflammation. ²⁸	Downregulates surface expression of major histocompatibility complex (MHC) class Il molecules and other glycoproteins by directing them to the late endosomal/lysosomal compartment. ^{18,19}	Pentraxins are part of the humoral arm of innate immunity and behave as functional ancestors of antibodies by mediating agglutination, complement activation and opsonization. PTX4 is a new unique member of the pentraxin superfamily, conserved in evolution. Further studies are needed to define its function. ²⁹	Plays a critical role in endothelial cell capillary morphogenesis. ³⁰	Enhancer Of MRNA Decapping. ³¹ Diseases associated with EDC4 include Human	Granulocytic Anaplasmosis and Anteroseptal Myocardial Infarction.	Component of a Polycomb group (PcG) multiprotein PRC1-like complex, a complex	class required to maintain the transcriptionally repressive state of many genes, including Hox genes, throughout development. ³²
Literature category P	1 B		Τ .		<u> </u>	4	2 P	Ε		2 C	
CADD- score	35.0	32.0	12.93	29.8	38.0	12.5	23.8	10.93	27.7	23.2	18.51
Amino Acid change	p.R723Q	p.G588S	p.A314T	p.G634R	p.W4*	p.V63 M	p.A351V	p.S275G	p.A1230V	p.I753 M	p.D471N
GnomAD allele count Population frequency	217 = 0.00077	1236 = 0.0043	30 = 0	192 = 0.001	Unobserved	3 = 0.000012	Unobserved	447 = 0.002	21=0	41 = 0	40 = 0
Candidate risk Inheritance mode gene (parental allele)	Compound heterozygous (mother)	Compound heterozygous (father)	Compound heterozygous (father)	Compound heterozygous (mother)	De поvо	De поvо	De novo	Compound heterozygous (mother)	Compound heterozygous (father)	Compound heterozygous (father)	Compound heterozygous (mother)
Candidate risk gene	ABCB6	ABCB6	ADAM32	ADAM32	MARCH1	PTX4	PLXDC1	EDC4	EDC4	PHC2	РНС2
rs number	rs148211042	rs145526996	rs745952927	rs150114293	Unknown	rs775407157	Unknown	rs111231628	rs563149577	rs376869490	rs142759750
Chr: position:alleles	2:220075521:C/T	2:220078006:C/T	8:39044452:G/A	8:39103683:A/G	4:164775272:C/T	16:1537911:C/T	17:37234300:G/A	16:67911677:G/A	16:67916920:C/T	1:33794634:G/C	1:33820146:T/C
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Trio	Chr: position:alleles	rs number	Candidate risk gene	Candidate risk Inheritance mode gene (parental allele)	GnomAD allele count Population frequency	Amino Acid change	CADD- score	Literature	Protein function	
	7:5427971:C/A	Unknown	TNRC18	De поvо	0	p.G495V	14.88	2	Protein Coding gene. Diseases associated with TNRC18 include Atrial Septal Defect and Seckel Syndrome. ³³ Lead CpGs at TNRC18 map to active enhancers in kidney cortex and are associated with renal fibrosis ³⁴	INTERNATIONAL
6	9:133047587:C/G	Unknown	HMCN2	Compound heterozygous (father)	8 = 0	p.Q161E	22.8	2	Protein Coding gene. Diseases associated with HMCN2 include Posterior Myocardial Infarction. 35	CT SEE
	7:153503673:A/G	rs559/3/4101	HIMCINZ	Compound neterozygous (mother)		p.G4273E	3.1/1			E-MACIAN .
10	8:134225273:G/A	Unknown	CCN4	De novo	Unobserved	р.С79Ү	27.2	ю	Mediates diverse developmental processes, such as control of cell proliferation, adhesion, cell polarity and establishment of cell fates. ³⁶	
12	2:84816032:T/A	Unknown	DNAH6	De novo	Unobserved	р.L855Н	22.8	м	Force generating protein of respiratory cilia. Produces force towards the minus ends of microtubules. Diseases associated include Primary Ciliary Dyskinesia and Anomalous Left Coronary Artery From The Pulmonary Artery. ³⁷ :	
	11:64645648:G/A	rs747258453	EHD1	Compound heterozygous (mother)	Unobserved	p.F97L	27.8	1	Important motif in proteins involved in protein- protein interactions and in intracellular	
	11:64645841:C/T	Unknown	ЕНD1	Compound heterozygous (father)	2 = 0	ت. ا	15.68		sorting. The protein encoded by this gene is thought to play a role in the endocytosis of IGF1 receptors. ³⁸	
	6:129621952:A/T	Unknown	LAMA2	Compound heterozygous (father)	Unobserved	p.11037F	18.85	ო	It is thought to mediate the attachment, migration and organization of cells into	
	6:129824345:T/C	rs151334775	LAMA2	Compound heterozygous (mother)	22 = 0	p.P2823S	25.3		tissues during embryonic development by interacting with other extracellular matrix components. Diseases associated with LAMA2 include Muscular Dystrophy. ³⁹	
13	2:196720589:C/A	rs749776504	DNAH7	Compound heterozygous (father)	0 = 9	p.R2847S	12.06	ო	Force generating protein of respiratory cilia. Produces force towards the	
	2:196889160:A/G	rs182086316	DNAH7	Compound heterozygous (mother)	439 = 0.002	p.R246C	29.2		minus ends of microtubules."*Uiseases associated with DNAH7 include Situs Inversus and Dextrocardia With Situs Inversus.	
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Protein function	In the hair cortex, hair keratin intermediate filaments are embedded in an interfilamentous matrix, consisting of hair keratin-associated protein (KRTAP), which are essential for the formation of a rigid and resistant hair shaft through their extensive disulphide bond crosslinking with abundant cysteine residues of hair keratins. ⁴¹ :	The protein encoded by this gene is one of the highly conserved mini-chromosome maintenance proteins (MCM) that are essential for the initiation of eukaryotic genome replication. Diseases associated with MCM8 include Premature Ovarian Failure 10 and Amenorrhea. ⁴²	This gene encodes a large protein that contains six ankyrin repeats, as well as a Src homology 3 (SH3) domain and two sterile alpha motif	(SAM) domains, which may be involved in protein-protein interactions. ⁴³	May regulate the disassembly of focal adhesions. May localize receptor-like tyrosine	phosphatases type 2A at specific sites on the plasma membrane, possibly regulating their interaction with the extracellular environment and their association with substrates. ⁴⁴	Intermicrovillar adhesion molecule that controls the packing of microvilli at the apical membrane of epithelial cells. ^{23,24}	Involved in apoptosis. May have a role in rRNA processing and mRNA splicing. Associates with pre-rRNA precursors. ⁴⁵ :	Receptor for calcitonin gene-related peptide (CGRP) and adrenomedullin.46
Literature	м	м	2		2		\leftarrow	₩	2
CADD- score	,	23.1	18.14		15.66	33	16.9	24.5	29.6
Amino Acid change	p.S105Wfs*115	p.E94G	p.R623Q p.P454L	!	1	p.R1041L	p.=SPLICE_SITE DONOR	p.P90S	p.P209L
GnomAD allele count Population frequency	Unobserved	14 = 0	150 = 0.001 $83 = 0$,	43 = 0	0 = 6	5 = 0.00003	2 = 0	Unobserved
Candidate risk Inheritance mode gene (parental allele)	De novo	De novo	Compound heterozygous (mother) Compound heterozygous	(father)	Compound heterozygous (mother)	Compound heterozygous (father)	De novo	De novo	De novo
Candidate risk gene	KRTAP5-1	MCM8	CASKIN2 CASKIN2		PPFIA4	PPFIA4	CDHR2	DDX47	CALCRL
rs number	rs761147271	rs377435486	rs200947487		rs61756414	rs528573275	rs780769740	rs780873695	Unknown
Chr: position:alleles	11:1606144: ACAAGAGCCACAG CCCCCTTGG/.	20:5935281:A/G	17:73499287:T/C 17:73500515:G/A		1:203018043:G/C	1:203036894:G/T	5:176002840:G/A	12:12974228:C/T	2:188228104:G/A
Trio	41		15		16		17		18

(Continues)

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Protein function	Has deubiquitinating activity. Also regulates cell proliferation and apoptosis through	deubiquitination of SUDS3 a regulator of histone deacetylation. ⁴⁷	Protein Coding gene. Diseases associated with ANKRD36 include Giant Axonal	Neuropathy. ⁴⁸ .	Force generating protein of respiratory cilia. Produces force towards the minus ends of	microtubules. ⁴⁹ Diseases associated with DNAH11 include Ciliary Dyskinesia, Primary and Primary Ciliary Dyskinesia.	Human-specific protein that promotes neural progenitor proliferation and evolutionary expansion of the brain neocortex by regulating the Notch signalling pathway via direct interaction with NOTCH2.50 NOTCH2 variants are associated with Alagille syndrome, including cholestasis phenotypes (www.omim.org/entry/600275).	A candidate histone demethylase thought to be a co-activator for key transcription factors.	Plays a role in the DNA-damage response pathway. ⁵¹	Interacts with, and positively regulates, dishevelled-mediated signalling pathways	during development. ⁵² Associated with Townes-Brocks syndrome-2 (OMIM 617 466).	Involved in modulation of cell growth and cellular response to gamma radiation probably	via regulation of the Akt signalling pathway. Involved in regulation of p53/TP53. ⁵³ :
Literature	1		2		ო		П	ო		ო		ო	
CADD- score	ı	0.034	11.81	18.47	35	19.07	24.1	26.5	26.2	23.6	25.5	17.38	23.3
Amino Acid change	p.L82l	p.D50N	p.A240T	p.T428 M	p.R2997Q	p.K3779N	p.T50 M	p.E2531K	р.D374Н	p.T8 M	p.W679R	p.S279G	p.A21S
GnomAD allele count Population frequency	51=0	76 = 0	16 = 0	12 = 0	3087 = 0.011	Unobserved	4 = 0	1162 = 0.0041	132 = 0.00047	5 = 0	166 = 0.001	1053 = 0.004	485 = 0.003
Candidate risk Inheritance mode gene (parental allele)	Compound heterozygous (mother)	Compound heterozygous (father)	Compound heterozygous (mother)	Compound heterozygous (father)	Compound heterozygous (mother)	Compound heterozygous (father)	De novo	Compound heterozygous (father)	Compound heterozygous (mother)	Compound heterozygous (mother)	Compound heterozygous (father)	Compound heterozygous (father)	Compound heterozygous (mother)
Candidate risk gene	USP17L2	USP17L2	ANKRD36	ANKRD36	DNAH11	DNAH11	NOTCH2NLA	JMJD1C	<i>JMJ</i> D1С	DACT1	DACT1	TSPYL5	TSPYL5
rs number	rs201734663	rs199985479	rs770051110	rs567234399	rs35865357	rs751035617	rs782819394	rs71508957	rs200016210	Unknown	rs200977826	rs151015596	rs79679520
Chr: position:alleles	8:11996026:T/G	8:11996122:C/T	2:97790321:A/G	2:97823866:C/T	7:21805095:A/G	7:21901605:G/C	1:145273295:C/T	10:64927837:C/T	10:64974807:C/G	14:59104943:C/T	14:59113376:T/C	8:98289238:T/C	8:98290012:A/C
Trio			19					21				22	

(Continues)

	NAME OF	1
/er	CAL PR	-Wiley

and JNK, and resulting in cytokine secretion and the inflammatory response 58	Catalyses the transfer of sulphate in chondroitin. ⁵⁹ Diseases associated with CHST11 include Mucinoses and Costello Syndrome (OMIM 618 167).	Iron-sulphur (Fe-S) cluster binding motor protein that has a role in chromosome segregation during mitosis. ⁶⁰	May be involved in transcriptional regulation. ⁶¹ Diseases associated with ZNF30	include Chromosome 19Q13.11 Deletion Syndrome and Brugada Syndrome.
	m	ო	2	
	32.	10.11	23.2	25.8
	p.G213S	p.N907S	p.C182R	p.R589W
	Unobserved	Unobserved	43 = 0	542 = 0.002
	De novo	De novo	Compound heterozygous (mother)	Compound heterozygous (father)
	CHST11	KIF4A	ZNF30	ZNF30
	Unknown	Unknown	rs367651957	rs140215760
	12:105151159:G/A Unknown	X:69623814:A/G	19:35434411:C/T	19:35435632:C/T
	28	29		

essential role in X chromosome inactivation.⁵⁷

Involved in DNA management and plays an

က

31

p.D849N

Unobserved

Syndrome and Autosomal Dominant Non-

 $Syndromic\ Intellectual\ Disability.^{56};$

Protein Coding gene. Diseases associated ventricular tachycardia (OMIM 615 441)

7

35

p.R295W

23 = 0

Compound heterozygous

FAM234B

rs367547952

12:13219604:T/C

25

(mother)

25.6

p.F4441

150 = 0.001

Compound heterozygous

FAM234B

rs140271825

12:13221607:T/A

(father)

De novo

SMCHD1

Unknown

18:2722603:G/A

26

with FAM234B include Temtamy

Adapter involved in TLR2 and TLR4 signalling

┛

23.6

p.R215H

49 = 0

De novo

TIRAP

rs185114125

11:126162948:G/A

pathways in the innate immune response.

activation of NF-kappa-B, MAPK1, MAPK3 Acts via IRAK2 and TRAF-6, leading to the

Ca2 + release via the sarcoplasmic reticulum

Contributes to regulation of luminal

7

n.a.

p.S297Ffs*32

Unobserved

De novo

rs201431159 TRDN

6:123786033:./A

24

36.

2 = 0.000011

Sodium/hydrogen exchanger and transmembrane protein.⁵⁴

Protein function

Literature category

CADDscore

Amino Acid

count Population **GnomAD allele**

Candidate risk Inheritance mode

(parental allele)

gene

Chr: position:alleles

Trio

TABLE 2 (Continued)

De novo

SLC9B1

rs75599926 rs number

4:103832611:G/A

23

change p.R305*

frequency

calcium release channels. 55 Associated with

Abbreviations: CADD-score, Combined Annotation-Dependent Depletion scoreChr, Chromosome.

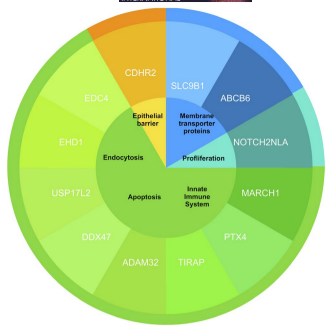


FIGURE 2 Function of the 12 prioritized candidate risk genes for early-onset PSC. The core indicates the protein functions and the inner ring shows the 12 candidate risk genes. Information on genetic functions comes from multiple databases including Genecards (www.genecards.com), Reactome (www.reactome.org) and OMIM (Online Mendelian Inheritance in Man; www.omim.org)

very beginning of the MARCH1-gene (transcript position 4) in patient #4, a boy of 8 with PSC-autoimmune hepatitis overlap syndrome, also called autoimmune sclerosing cholangitis. Such an early stop-gain results in loss of the corresponding protein from this allele and is predicted to be highly pathogenic. As the inheritance mode of this gene is still unknown, we cannot predict the biological effect of this variant. Functional studies of MARCH1 suggested that this gene mediates the immunosuppressive effect of the anti-inflammatory cytokine interleukin 10 (IL10) on antigen presentation in monocytes. ^{18,19} Knockdown of MARCH1 strongly inhibited IL-10-dependent down-regulation of cell surface HLA-DR. ¹⁹ The exact contribution of the MARCH1 gene regulation to immunopathology remains to be explored.

We identified variants in genes *ABCB6* and *SLC9B1*, which are (according to www.pathcards.genecards.org) related to the 'transport of glucose and other sugars, bile salts and organic acids, metal ions and amine compounds' pathway. Patient #3, a girl with both PSC and IBD diagnosed at age 7, had compound heterozygous *ABCB6* variants. The gene encodes a member of the ATP-binding cassette (ABC) transporter superfamily and is known to bind heme and porphyrins and function in their ATP-dependent uptake in the mitochondria.^{20,21} Both variants are predicted to be damaging and to disrupt the highly conserved ABC transport and ABC transmembrane regions of the protein, respectively. However, whether this genetic variant is truly involved in alteration of bile salt homeostasis has to be investigated further.

Another possible cause of development of PSC is the epithelial cell lining of the bile ducts. Bile salts are toxic in high concentrations.²² Damage of the epithelial border may result in leakage of bile and could be an important driver of toxicity. In patient #17 we identified one de novo intronic variant positioned exactly at a splice-donor consensus sequence site in *CDHR2*, predicted to disrupt splicing of the transcript and causing loss of function. *CDHR2* plays a central role in the integrity of epithelial tissues such as the bile duct epithelium.^{23,24}

Uncovering the functional consequences of the newly discovered candidate variants, in particular for the genes without a known function, and the pathways involved in the onset of PSC will require detailed functional experiments involving different functional readouts, given the broad nature of the identified genes, and further verification of our findings in independent cohorts.

4.3 | From theory to definitive proof

This is the first explorative study of high-impact rare coding variants in young patients with PSC which could help to identify causative genes. To prove that our set of candidate genes contains a causative gene, replication in an independent patient cohort is a first step. One method to replicate low-frequency and rare variants in complex immune diseases is by targeted genotyping using the Illumina exome chip array (HumanExome BeadChips, Illumina, Inc. San Diego, CA). This next-generation genotyping array includes several regions on the genome that are known to play a role in immune mediated disease based on the results from existing re-sequencing datasets. In a first attempt to further study the contribution of rare variants in PSC, our research group is in the process of replication genotyping of identified variants in a large international exome array case-control study and identified several genetic loci containing rare variants that are associated with PSC. Adding our new candidate risk genes and variants to such arrays may contribute to efficient replication.

Furthermore, replication in other early-onset PSC cohorts is needed. In 2013, the BROAD Institute partnered with researchers worldwide to develop a collaborative exome sequencing network in IBD, and this initiative is currently ongoing. A similar project is now up and running in PSC with the aim to meta-analyse the exomes of more than 1000 patients of European ancestry. This will enable modelling of the combined contribution of polygenic and oligogenic variants to the inheritance of PSC.

Convincing evidence to prove that a gene is causal in a disease is identifying a similar genetic variant in the same gene in another patient. In monogenic diseases, a freely accessible Web-based tool called GeneMatcher (www.genematcher.org) is used to identify additional individuals with rare phenotypes who have variants in the same candidate gene.²⁵ A similar tool does not yet exist for complex genetic diseases but data sharing such as GeneMatcher might be useful here as well.

4.4 | Implications for clinical practice

At this moment, there is no curative therapy available for PSC. Quality-of-life undermining complications may eventually justify a liver transplantation. The ultimate goal of investigating the genetic basis of this disease is to help reveal mechanisms of disease pathology and guide the selection of new targets for drug discovery. Each genetic risk locus can be seen as a potential drug target and the starting point of new treatment opportunities. This has successfully been demonstrated in the field of IBD, in which small-molecule inhibitors were used to recapitulate the anti-inflammatory function of CARD9 variants associated with protection from IBD.²⁶ Scientists now recognize that genes with evidence for causality in disease are more promising for identification of new drug targets, and this has led to an increased interest in disease-associated genes with variants that reduce gene function, such as nonsense, frameshift or essential splice-site variants.²⁷ Our study is one of the first steps on this road to drug discovery for patients with PSC.

5 | CONCLUSION

We identified 54 rare protein-altering genetic variants in 36 genes, of which 12 were prioritized as candidate risk genes. The functional consequences of these variants and their causality to PSC will need to be studied in replication cohorts and with functional testing.

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CONFLICTS OF INTEREST

None.

AUTHOR CONTRIBUTIONS

SMH, RKW, PFvR and CCvD contributed to study concept and design, interpretation of data and drafting of the manuscript. BAEdK, MEJ, BGPK, TdM, VMW and ON contributed to the collection of patient data. SMH obtained the data. Whole-exome sequencing and quality control were performed by MJD, CS, RJX and MAR. SMH and CCvD analysed the data. HJV, RB, DBHJ, NF and MV contributed important intellectual content. PFvR and CCvD had full responsibility for the study.

ETHICS APPROVAL STATEMENT

The Medical Ethical Committee of the UMCG approved the study protocol (METC 2016/289), and secondary approval was obtained from all participating centres.

PATIENT CONSENT STATEMENT

All participating parents and teenagers 12-19 years old gave informed consent prior study inclusion.

CONFERENCE PRESENTATION

This study was selected for a lecture presentation during the *Digestive Disease Week*® in May 2019 in San Diego.

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