

Perturbations of genes essential for Müllerian duct and Wölfian duct development in Mayer-Rokitansky-Küster-Hauser syndrome

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Summary

Mayer-Rokitansky-Küster-Hauser syndrome (MRKHS) is associated with congenital absence of the uterus, cervix, and the upper part of the vagina; it is a sex-limited trait. Disrupted development of the Müllerian ducts (MD)/Wölfian ducts (WD) through multifactorial mechanisms has been proposed to underlie MRKHS. In this study, exome sequencing (ES) was performed on a Chinese discovery cohort (442 affected subjects and 941 female control subjects) and a replication MRKHS cohort (150 affected subjects of mixed ethnicity from North America, South America, and Europe). Phenotypic follow-up of the female reproductive system was performed on an additional cohort of *PAX8*-associated congenital hypothyroidism (CH) ($n = 5$, Chinese). By analyzing 19 candidate genes essential for MD/WD development, we identified 12 likely gene-disrupting (LGD) variants in 7 genes: *PAX8* ($n = 4$), *BMP4* ($n = 2$), *BMP7* ($n = 2$), *TBX6* ($n = 1$), *HOXA10* ($n = 1$), *EMX2* ($n = 1$), and *WNT9B* ($n = 1$), while LGD variants in these genes were not detected in control samples ($p = 1.27E-06$). Interestingly, a sex-limited penetrance with paternal inheritance was observed in multiple families. One additional *PAX8* LGD variant from the replication cohort and two missense variants from both cohorts were revealed to cause loss-of-function of the protein. From the *PAX8*-associated CH cohort, we identified one individual presenting a syndromic condition characterized by CH and MRKHS (CH-MRKHS). Our study demonstrates the comprehensive utilization of knowledge from developmental biology toward elucidating genetic perturbations, i.e., rare pathogenic alleles involving the same loci, contributing to human birth defects.

Mayer-Rokitansky-Küster-Hauser syndrome (MRKHS [MIM: 277000]), also referred to as Müllerian aplasia, is characterized by congenital absence of the uterus, cervix, and upper part of the vagina in females with a normal karyotype (46, XX).¹ With an incidence of 1 in 4,500–5,000 newborn females, MRKHS is the second most common

cause of primary amenorrhea after gonadal dysgenesis.² MRKHS is further divided into MRKHS type I (isolated) and MRKHS type II (syndromic) according to the presence of multi-organ involvement.³

Formation and morphogenesis of the Müllerian ducts take place during weeks 5–6 of human embryogenesis

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and are induced by the Wölfian ducts (mesonephric ducts).⁴ Knock-out of genes expressed in the Müllerian duct (MD) or the Wölfian duct (WD) such as *Lhx1*,⁵ *Pax8*,⁶ *Wnt9b*,⁷ and *Tbx6*⁸ lead to MRKHS-like phenotypes in mice. In addition, knock-out of abdominal B homeobox genes (*Hoxa9*, *Hoxa10*, *Hoxa11*, *Hoxa13*), which are expressed along the anterior-posterior axis in a segmental pattern, can also disrupt the development of the corresponding part of the Müllerian duct, Wölfian duct, and the spine.⁹

Despite numerous candidate genes and pathways, only *WNT4* (MIM: 603490) has been well established to be associated with a clinically distinct subtype of MRKHS characterized by Müllerian aplasia with hyperandrogenism (MIM: 158330).¹⁰ To fully explore the molecular pathogenesis and genetic architecture of MRKHS, we took advantage of the comprehensive knowledge base of genitourinary developmental biology and performed a mutational burden analysis in a cohort of Chinese individuals with MRKHS. We also studied a global multi-center replication cohort consisting of North American, South American, and European individuals with MRKHS, and a Chinese phenotypic follow-up cohort of individuals with congenital hypothyroidism, to further investigate findings from the discovery cohort.

In the discovery cohort, we recruited 442 Chinese individuals of whom 105 derived from trios in which parents were unaffected and 941 female control subjects. The affected individuals consisted of 330 (74.7%) individuals with MRKHS type I and 112 (25.3%) with MRKHS type II. The replication cohort recruited 150 multi-ethnic individuals with MRKHS, including 78 singletons, 66 affected subjects with unaffected familial samples, and 3 families that each have 2 affected members (supplemental methods). This cohort contained 84 (56.0%) with MRKHS type I, 61 (40.7%) with MRKHS type II, and 5 (3.3%) with MRKHS unspecified (Table 1, Figure 1B). Proband and available familial samples from both cohorts underwent exome sequencing (ES, details provided in supplemental methods). The case-control dataset then underwent harmonization processing, where an individual coding sequence site was excluded from the analysis if the abso-

lute difference in percentages of cases compared to controls with at least 10× coverage differed by greater than 8%.¹¹ Afterward, the mutational burden of likely gene-disrupting (LGD) variants and deleterious missense (D-mis) variants in a set of 19 MD/WD development-associated genes (Table S1) was evaluated. Specifically, LGD variants were defined as variants predicted to cause loss-of-function (LoF) of the gene, and included nonsense, frameshift, and canonical splice site variants. The LGD + D-mis model additionally included missense variants predicted to be damaging by *in-silico* tools (SIFT score¹² < 0.05, PolyPhen-2 score¹³ > 0.95, CADD score¹⁴ > 15).

As a result, we identified 12 LGD variants in 7 candidate genes, including *PAX8* (MIM: 167415), *BMP4* (MIM: 112262), *BMP7* (MIM: 112267), *TBX6* (MIM: 602427), *HOXA10* (MIM: 142957), *EMX2* (MIM: 600035), and *WNT9B* (MIM: 602864) (Table 2, Figure 2A), while LGD variants were not detected in any of the candidate genes from the exomes of 941 female control samples ($p = 1.2E-06$, SKAT-O). Of the 12 LGD variants, 8 were protein-truncating and 4 affect the canonical splice sites. All of the truncating variants except one in *WNT9B* and one in *BMP4* were predicted to result in unstable mutant RNA susceptible to nonsense-mediated mRNA decay by NMDescPredictor¹⁵ (Figure S1). A significant burden of LGD + D-mis variants was also observed from cases (35/442) versus controls (38/941) (odds ratio = 1.98, $p = 3.8E-03$) in the discovery cohort (Table 2, Figure 2A).

With the identification of four LGD variants (c.156_157dupCG [p.Val53AlafsTer24], c.25+1G>T [splice donor], c.195delC [p.Tyr66ThrfsTer10], and c.322C>T [p.Arg108Ter], GenBank: NM_003466.3), *PAX8* represents the most significant disease-associated gene (false discovery rate [FDR] adjusted $p = 0.01$) (Tables 2, 3, and S2). Notably, *Pax8*^{-/-} mice recapitulate the human MRKHS phenotype: normal ovaries/oviducts and loss of the uterus.⁶ In humans, haploinsufficiency of *PAX8* is associated with congenital hypothyroidism (CH) caused by thyroid dysgenesis.¹⁶ However, all four individuals carrying *PAX8* heterozygous LGD variants presented with isolated MRKHS type I without a clinically observable CH phenotype from the initial visit. Clinical follow up confirmed

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Table 1. Cohort demographic information and phenotypes

Information	Discovery cohort (n = 442)	Replication cohort (n = 150) ^a
Age of visit, y, mean ± SD	24.0 ± 5.3	17.2 ± 3.8
Classification, n (%)		
MRKHS type I	330 (74.7%)	84 (56.0%)
MRKHS type II	112 (25.3%)	61 (40.7%)
Unspecified	–	5 (3.3%)
Complications, n (%)		
Skeletal anomaly	90 (20.4%)	21 (22%)
Kidney anomaly	46 (10.4%)	32 (33%)
Other deformities	3 (0.7%)	8 (8%)

^aIn the replication cohort of 150 samples, data regarding MRKHS classification were available for all the individuals; data regarding age of visit and complications were available for 96 individuals. The percentages in this column were calculated based on individuals with available information. Abbreviations: MRKHS, Mayer-Rokitansky-Küster-Hauser syndrome.

that the thyroid hormone levels of MRK49 (c.25+1G>T) and MRK467 (c.322C>T [p.Arg108Ter]) were within normal range (Table S3). Intriguingly, sequencing of the parental DNA from these two individuals revealed that their *PAX8* LGD variants were both paternally inherited (Figure 2B), consistent with a dominant disease trait with sex-dependent penetrance.

In addition to LGD variants, we also identified three D-mis *PAX8* variants (c.542C>T [p.Ser181Phe], c.266T>C [p.Val89Ala], c.236C>G [p.Ser79Cys]) from three individuals in the discovery cohort (Table 3). We experimentally explored the potential effect on the DNA-binding ability of these three D-mis variants using a luciferase reporter assay and found that one of the three variants (c.236C>G [p.Ser79Cys]) caused decreased transactivation potential of *PAX8* protein on its consensus binding sequence (Figure 2C), thus experimentally supporting its characterization as a LoF allele. We followed up this individual with the c.236C>G (p.Ser79Cys) variant (MRK330), and intriguingly, this variant was also found to be inherited from her father (Figure 2B). Moreover, MRK330 and her father both presented with subclinical hypothyroidism (Table S3), supporting the genetic pleiotropy of *PAX8*.

From the replication MRKHS cohort, we identified one nonsense variant in *PAX8* (c.619C>T [p.Arg207Ter]) and two heterozygous D-mis variants in *PAX8* (c.136G>A [p.Asp46Asn] and c.727C>G [p.Gln243Glu]) in three subjects affected with MRKHS type I (Table 3). The c.136G>A (p.Asp46Asn) variant was revealed to cause LoF of *PAX8* by luciferase assay (Figure 2C), and this variant was also confirmed to be paternally inherited, consistent with an autosomal-dominant mode of inheritance, with sex-dependent phenotypic expression of the disease trait (Figure 2B). Notably, LGD and D-mis variants in *PAX8* were enriched in the DNA-binding PAX domain

of the protein (Figure 2D), implicating an important role of the DNA-binding function of *PAX8* in the pathogenesis of MRKHS.

In order to further investigate the genetic pleiotropy of *PAX8*, we enrolled five female individuals ascertained as having pathogenic *PAX8* variants from an independent Chinese cohort with CH.¹⁷ Interestingly, ultrasound and MRI examination of the pelvis during clinical follow-up revealed that one of the individuals (CH123) had previously unobserved aplasia of the uterus, suggesting a clinical diagnosis of MRKHS (Table S4, Figure S2). The *PAX8* variant in this individual was c.68G>T (p.Gly23Val) (Table S3) and was validated to be a LoF allele *in vivo* (Figure 2C). The phenotype of this individual is consistent with a syndromic condition, which we define as congenital hypothyroidism-Mayer-Rokitansky-Küster-Hauser syndrome (CH-MRKHS).

In addition to *PAX8*, two genes encoding bone morphogenetic proteins, namely *BMP4* and *BMP7*, showed a significant mutational burden in the discovery cohort (FDR adjusted p = 0.03 for both *BMP4* and *BMP7*), implicating the contribution of BMP signaling perturbations to MRKHS. *BMP4* and *BMP7* are mainly expressed in the Wölfian duct epithelium and regulate the early formation of the Müllerian duct¹⁸ (Figure 2A). To date, genetic studies have been focused on the function of these genes in the development of the urinary system but not the reproductive system.^{19–21}

A heterozygous stop-gain variant c.367G>T (p.Glu123-Ter) and a heterozygous splice acceptor variant c.–132–1G>A in *BMP4* (GenBank: NM_001202.3) were identified in MRK644 and MRK166, respectively (Table S2). MRK166 presented with MRKHS type I, and MRK644 had a more complex clinical presentation with idiopathic scoliosis. Interestingly, MRK644 inherited her *BMP4* allele from a mother mosaic for the c.367G>T variant. The *BMP4* variant c.367G>T (p.Glu123Ter) in the proband was detected in 26/55 (47%) variant/total reads by ES, indicating a putative heterozygous variant. In contrast, her mother's blood DNA had Vr/Tr = 13/60 (22%) by ES, consistent with a mosaic variant allele. The mosaicism was further supported by digital droplet PCR (ddPCR), results of which suggested a mutational ratio of around 50% in the proband and 10% in the mother (Figure S3). In the second individual (MRKH166), the splice acceptor variant was paternally inherited, presenting a similar sex-limited inheritance pattern as *PAX8* (Figure S3). ES studies in the replication cohort identified another LGD variant in *BMP4* (c.766C>T [p.Arg256Ter]) from an individual affected with MRKHS type I, further suggesting the association of *BMP4* variants with MRKHS (Table S2). In *BMP7* (GenBank: NM_001719.2), a splice acceptor variant c.1036–2A>G from MRK444 and a frameshift variant c.275_287dupTGGAGGAGGGCGG (p.Pro98GlyfsTer31) from MRK342 were identified (Table S2). Both individuals presented with MRKHS type I, with unknown parental origin.

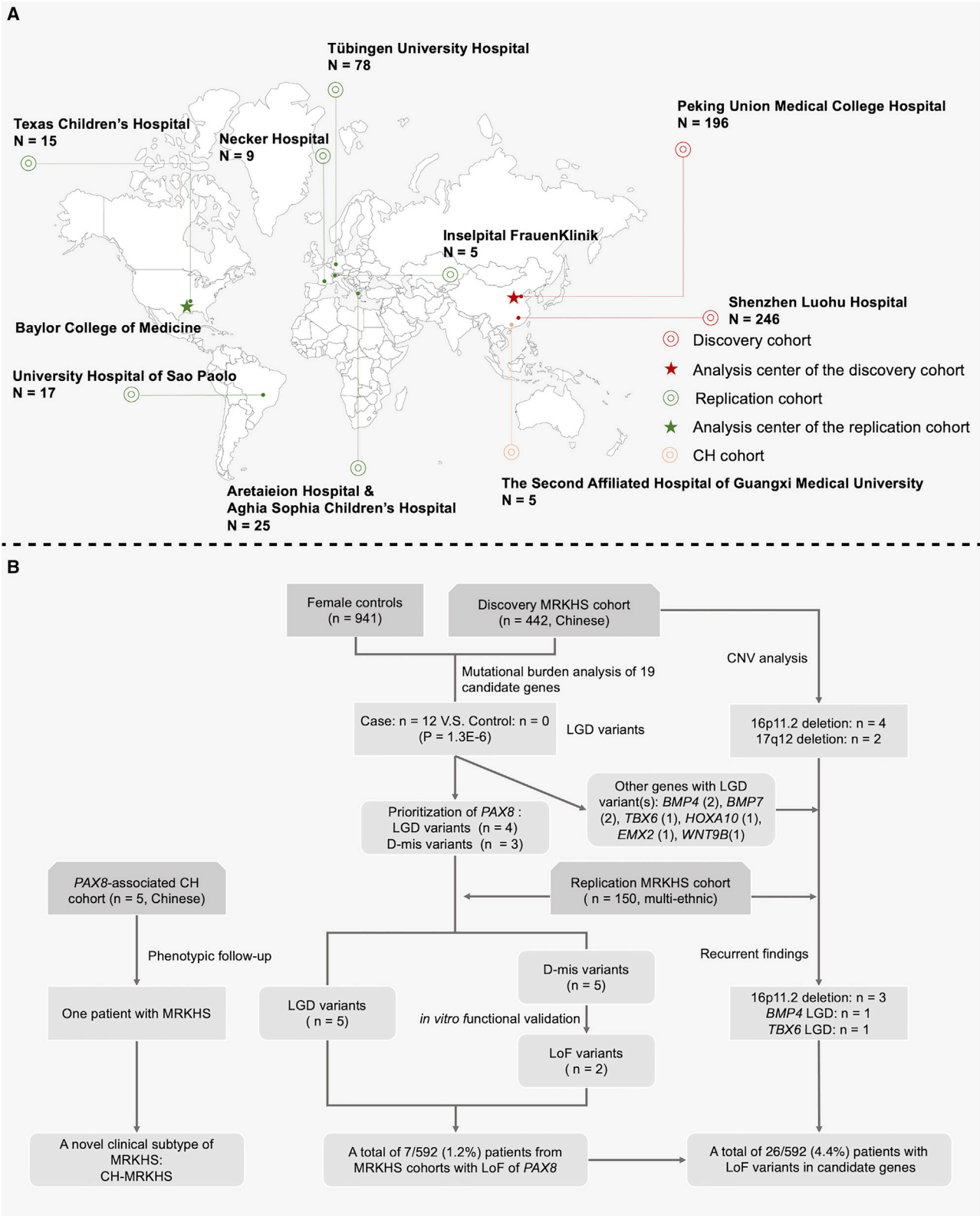


Figure 1. Study design and workflow of the study

(A) Centers of recruitment and analysis for MRKHS- and CH-affected individuals.

(B) Workflow and main findings of this study. Abbreviations: MRKHS, Mayer-Rokitansky-Küster-Hauser syndrome; CH, congenital hypothyroidism; LGD, likely gene-disrupting; LoF, loss-of-function; D-mis, deleterious missense.

Table 2. Mutational burden analysis of 19 candidate genes among 442 affected subjects and 941 female control subjects

Gene symbol	pLI	LGD variants			LGD + D-mis variants			
		Case (n = 442)	Control (n = 941)	FDR adjusted p value	Case (n = 442)	Control (n = 941)	OR	FDR adjusted p value
<i>PAX8</i>	0.90	4	0	0.01	7	4	3.77	0.16
<i>BMP4</i>	0.97	2	0	0.03	5	1	10.77	0.05
<i>BMP7</i>	0.98	2	0	0.03	4	1	8.62	0.13
<i>TBX6</i>	0.03	1	0	0.08	6	4	3.23	0.27
<i>HOXA10</i>	0.68	1	0	0.08	3	2	3.23	0.35
<i>EMX2</i>	0.94	1	0	0.08	1	2	1.08	0.78
<i>WNT9B</i>	0.05	1	0	0.08	3	5	1.29	0.59
<i>HNF1B</i>	1.00	0	0	–	2	1	4.31	0.35
<i>HOXA9</i>	0.00	0	0	–	1	1	2.15	0.59
<i>GATA3</i>	0.88	0	0	–	1	2	1.08	0.78
<i>LHX1</i>	0.29	0	0	–	0	0	–	–
<i>WNT7A</i>	0.44	0	0	–	0	0	–	–
<i>HOXA11</i>	0.86	0	0	–	1	3	0.72	0.81
<i>WT1</i>	–	0	0	–	1	3	0.72	0.81
<i>HOXA13</i>	–	0	0	–	0	2	0.00	0.59
<i>PAX2</i>	0.12	0	0	–	0	2	0.00	0.59
<i>PBX1</i>	0.91	0	0	–	0	2	0.00	0.59
<i>WNT4</i>	0.15	0	0	–	0	1	0.00	0.81
<i>WNT5A</i>	0.97	0	0	–	0	2	0.00	0.76
Total	–	12	0	1.2E–06	35	38	1.98	3.8E–03

Rare variants were analyzed using SNP-set (Sequence) Kernel Association Test-Optimized (SKAT-O) test to determine association of mutational burden in the 19 genes. D-mis variants are defined by missense variants with CADD > 15 and predicted to be deleterious by both SIFT and PolyPhen. Abbreviations: pLI, probability of loss-of-function intolerance from ExAC database; D-mis, damaging missense; FDR, false discovery rate; OR, odds ratio.

We also identified LGD variants in genes previously related to MRKHS, including *TBX6* and *WNT9B*.^{22–24} A *TBX6* (GenBank: NM_004608.3) splice donor variant, c.621+1G>A, was identified in individual MRK639 with MRKHS type II. Besides Müllerian aplasia, MRK639 was affected with congenital scoliosis caused by a hemivertebra at L1. Sanger sequencing revealed that this individual carried the hypomorphic T-C-A haplotype (composed of three SNPs, rs2289292-rs3809624-rs3809627²⁵) in *trans* with the c.621+1G>A splice variant, which is consistent with the compound inheritance gene dosage model of *TBX6*-associated congenital scoliosis (TACS).^{25,26} From the replication cohort, we identified another frameshift variant (c.856_859delAATG [p.His286CysfsTer28]) in a non-canonical transcript of *TBX6* (ENST00000553607) in Mul23, an individual diagnosed with MRKHS type II (Table S2). *WNT9B* is a key signaling molecule which directs early formation of the Müllerian ducts.⁴ Missense variants in *WNT9B* were first described in a Chinese cohort of 42 MRKHS-affected individuals.²⁷ In the present study, we identified a heterozygous nonsense variant c.976C>T (p.Gln326Ter) (GenBank:

NM_001320458.2) in MRK57 with isolated MRKHS type I (Tables 2 and S2).

By screening for CNVs that encompass one or more of the 19 candidate genes in the discovery cohort, we identified four deletions of 16p11.2 (encompassing *TBX6*) and two deletions of 17q12 (encompassing *LHX1* [MIM: 601999] and *HNF1B* [MIM: 189907]) (Table S5, Figure S4). Both regions have been linked to MRKHS in previous studies.²⁸ All four individuals with 16p11.2 deletion were classified as MRKHS type II: one presented with idiopathic scoliosis, and the other three presented with vertebral malformations. Haplotyping of the 16p11.2 locus revealed that the individual with idiopathic scoliosis carried the C-T-C haplotype in the remaining allele, and the other three carried the T-C-A haplotype (Table S5), consistent with the compound inheritance, gene dosage model of TACS.^{25,26,29} One of the two individuals with 17q12 deletion presented with MRKHS type I, and the other had a clinical picture complicated by idiopathic scoliosis (Table S5). From the replication cohort, three 16p11.2 deletion CNVs were identified in three individuals (Figure S5), including two individuals with MRKHS

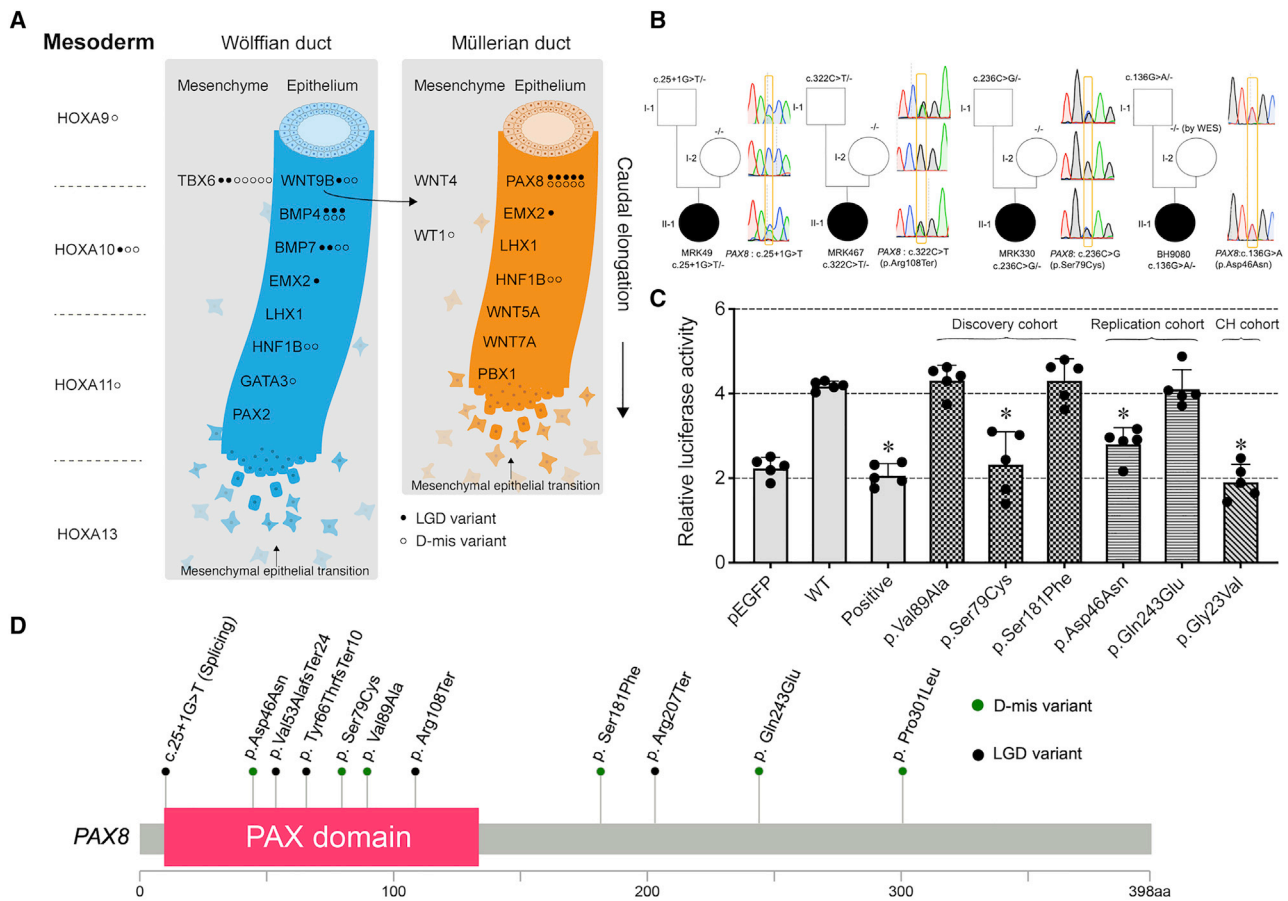


Figure 2. Identification of variants in genes essential for Müllerian duct (MD) and Wölffian duct (WD) development

(A) The elongation process of Müllerian duct/Wölffian duct is depicted. Protein located in the corresponding regions and with prioritized variants in them are also presented. Abbreviations: LGD, likely gene-disrupting; D-mis, damaging missense.

(B) Pedigree and Sanger sequencing results of three families with *PAX8* variants from the discovery cohort and one family from the replication cohort.

(C) Luciferase assay on *PAX8* missense variants from the discovery cohort and the replication cohorts. WT, wild type; R31C, a known deleterious variant, was used as the positive control; CH, congenital hypothyroidism. The data are the mean of $n = 5$ independent experiments. Error bars show one standard deviation (* $p < 0.05$, Dunnett's multiple comparison test).

(D) The mutational spectrum of LGD variants and damaging missense variants in *PAX8*.

type I and one individual with hip dysplasia and thus MRKHS type II. No 17q12 deletion CNVs were found in the replication cohort.

In summary, candidate variants in genes associated with MD/WD development were identified in MRKHS. A sex-limited penetrance with paternal inheritance was observed in multiple families. Of them, *PAX8* represents the most significant gene underlying the etiology of 7/592 (1.2%) individuals and is associated with a syndromic condition characterized by CH and MRKHS (CH-MRKHS). Other candidate genes such as *BMP4* and *BMP7* still warrant further genetic and functional studies. Our study demonstrates the comprehensive utilization of knowledge from developmental biology toward elucidating genetic perturbations, i.e., rare variant pathogenic alleles involving the same loci, contributing to human birth defects.

Data and Code Availability

The datasets supporting the current study have not been deposited in a public repository due to institutional ethics restrictions but are available from the corresponding author on request.

Supplemental Information

Supplemental Information can be found online at <https://doi.org/10.1016/j.ajhg.2020.12.014>.

Consortia

The members of the DISCO consortium are Guixing Qiu, Zhihong Wu, Terry Jianguo Zhang, Nan Wu, Shengru Wang, Jiaqi Liu, Sen Liu, Yuzhi Zuo, Gang Liu, Chenxi Yu, Lian Liu, Jiashen Shao,

Table 3. Summary of PAX8 variants identified from the discovery cohort and the replication cohort

Sample ID	MRKHS classification	Ethnicity	Zygoty	CHR	POS	vR	tR	Mutation type	Gene name	cDNA change	Protein change	gnomAD-AF	ExAC-AF	Inheritance	CADD score	Functional validation
Discovery cohort																
MRK51	type I	Chinese	het	2	114004364	22	46	frameshift	PAX8	c.156_157dupCG	p.Val53AlafsTer24	0	0	unknown	–	–
MRK49	type I	Chinese	het	2	114035946	105	265	splice_donor	PAX8	c.25+1G>T	–	0	0	paternal	–	–
MRK442	type I	Chinese	het	2	114002198	53	101	frameshift	PAX8	c.195delC	p.Tyr66ThrfsTer10	0	0	paternal	–	–
MRK467	type I	Chinese	het	2	114002071	49	89	stop_gained	PAX8	c.322C>T	p.Arg108Ter	0	0	unknown	–	–
MRK236	type I	Chinese	het	2	113999644	33	76	missense	PAX8	c.542C>T	p.Ser181Phe	0	0	unknown	28.8	normal
MRK283	type I	Chinese	het	2	114002127	58	99	missense	PAX8	c.266T>C	p.Val89Ala	0	0	unknown	26.7	normal
MRK330	type I	Chinese	het	2	114002157	56	112	missense	PAX8	c.236C>G	p.Ser79Cys	0	0	paternal	26.5	LoF
Replication cohort																
SEA13832	type I	European	het	2	113999286	38	91	stop_gained	PAX8	c.619C>T	p.Arg207*	0	0	unknown	–	–
BH9080	type I	European	het	2	114004386	84	179	missense	PAX8	c.136G>A	p.Asp46Asn	0	0	paternal	29.3	LoF
SEA13831	type I	European	het	2	113999178	59	224	missense	PAX8	c.727C>G	p.Gln243Glu	0	0	unknown	22.4	normal

GenBank: NM_003466.3 was used for annotations of cDNA and protein changes. Abbreviations: Chr, chromosome; gnomAD, genome aggregation database; ExAC, Exome Aggregation Consortium; AF, allele frequency; Ref, reference; VR, variant reads; TR, total reads; VAF, variant allele frequency; –, not applicable; LoF, loss-of-function; CADD, combined annotation dependent depletion.

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Declaration of interests

J.R.L. has stock ownership in 23andMe, is a paid consultant for Regeneron Pharmaceuticals and Novartis, and is a co-inventor on multiple United States and European patents related to molecular diagnostics for inherited neuropathies, eye diseases, and bacterial genomic fingerprinting. The Department of Molecular and Human Genetics at Baylor College of Medicine derives revenue from the chromosomal microarray analysis and clinical genomic sequencing offered in the Baylor Genetics Laboratory (<http://baylorgenetics.com>).

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Web Resources

Combined annotation dependent depletion (CADD), <https://cadd.gs.washington.edu/snv>
ExAC database, <http://exac.broadinstitute.org/>
GenBank, <https://www.ncbi.nlm.nih.gov/genbank/>
Genome aggregation database (gnomAD), <https://gnomad.broadinstitute.org/>
Online Mendelian Inheritance in Man (OMIM), <https://www.omim.org/>

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