

Genetic and clinical characterization of Pakistani families with Bardet-Biedl syndrome extends the genetic and phenotypic spectrum

Maleeha Maria^{1,2,#}, Ideke J.C. Lamers^{2,3,#}, Miriam Schmidts^{2,3,4,5}, Muhammad Ajmal¹, Sulman Jaffar⁶, Ehsan Ullah^{7,8}, Bilal Mustafa¹, Shakeel Ahmad¹, Katia Nazmutdinova⁴, Bethan Hoskins⁹, Erwin van Wijk^{10,11}, Linda Koster-Kamphuis¹², Muhammad Imran Khan², Phil L. Beales^{4,13}, Frans P.M. Cremers^{2,11}, Ronald Roepman^{2,3}, Maleeha Azam^{1,*}, Heleen H. Arts^{2,3,14,*}, Raheel Qamar^{1,15,16,*}

Supplementary Table S1: The results of biochemical tests with abnormal values.

Family ID	Individual ID	Test	Result (Normal)	
F01	IV:2	RFT		
		Urea	198 mg/dL (13-43 mg/dL)	
			Creatinine	21.26 mg/dL (0.7-1.3 mg/dL)
	IV:3	LFT		
		ALT	45 U/L (<40 U/L)	
		ALP	321 U/L (98-279 U/L)	
		LPT		
serum triglycerides		167 mg/dL (<160 mg/dL)		
		HDL cholesterol	28 mg/dL (35-40 mg/dL)	
F02	IV:1	RFT		
		Creatinine	0.7 mg/dL (0.7-1.3 mg/dL)	
		LFT		
		ALT	86 U/l (<40 U/L)	
		ALP	283 U/L (98-279 U/L)	
		AST	63 U/L (<38 U/L)	
		LPT		
		Serum triglycerides	350 mg/dL (<160 mg/dL)	
		Serum cholesterol	204 mg/dL (<200 mg/dL)	
		HDL cholesterol	30 mg/dL (35-40 mg/dL)	
	GTPT			
	LH	0.58 mIU/mL (1.14-8.75 mIU/mL)		
	Testosterone	0.39 ng/mL (1.66-8.77 ng/mL)		
	Lactate			
	Plasma Lactic acid	22.5 mg/dL (4.5-19.8 mg/dL)		
	IV:2	GTPT		
		Testosterone	1.27 ng/mL (1.66-8.77 ng/mL)	
F03	IV:2	RFT		
		Creatinine	0.7 mg/dL (0.7-1.3 mg/dL)	
		LFT		
		ALT	50 U/l (<40 U/L)	
		ALP	368 U/L (98-279 U/L)	
		LPT		
		Serum triglycerides	253 mg/dL (<160 mg/dL)	
		Serum cholesterol	243 mg/dL (<200 mg/dL)	
		LDL cholesterol	155 mg/dL (100-130 mg/dL)	
GTPT				
		Progesterone	0.1 ng/mL (1.2-15.9 ng/mL)	

F04	IV:2	LFT	
		ALT	40 U/l (<40 U/L)
		ALP	462 U/L (98-279 U/L)
		LPT	
		LDL cholesterol	90 mg/dL (100-130 mg/dL)
		HDL cholesterol	27 mg/dL (100-130 mg/dL)
		GTPT	
		Prolactin	32.63 ng/mL (3.46-19.40 ng/mL)
		Lactate	
		LDH	329 U/L (125-243 U/L)
Plasma Lactic acid	20.1 mg/dL (4.5-19.8 mg/dL)		
F05	V:2	RFT	
		Creatinine	1.3 mg/dL (0.7-1.3 mg/dL)
		LFT	
		ALT	55 U/l (<40 U/L)
		ALP	340 U/L (98-279 U/L)
		LPT	
		Serum cholesterol	90 mg/dL (100-130 mg/dL)
		Serum triglycerides	180 mg/dL (100-130 mg/dL)
		HDL cholesterol	34 mg/dL (100-130 mg/dL)
		GTPT	
Testosterone	0.0029 ng/mL (1.66-8.77 ng/mL)		
	V:4	GTPT	
		Testosterone	0.0008 ng/mL (1.66-8.77 ng/mL)

Abbreviations: ALP: Alkaline Phosphatase, ALT: Alanine Transaminase, AST: Aspartate aminotransferase, GTPT: Gonadotropin profile Tests, HDL: High density lipoprotein, ID: Identity, LDH: Lactate Dehydrogenase, LDL: Low Density Lipoprotein, LFT: Liver Function Tests, LH: Luteinizing Hormone, LPT: Lipid Profile Tests, RFT: Renal Function Tests

Supplementary Table S2: Targeted mutations prescreened in five probands from affected families. These mutations were reported from Pakistani BBS population (Khan et al.²⁹).

Gene	RefSeq Id	Nucleotide variant	Protein variant	Exons	References
<i>BBS1</i>	NM_02464.9.4	c.47+1G>T	p.(?)	Ex1	1
<i>BBS1</i>	NM_02464.9.4	c.442G>A	p.(D148N)	Ex5	2
<i>BBS10</i>	NM_024685.3	c.271dup	p.(C91Lfs*5)	Ex2	3
<i>BBS10</i>	NM_024685.3	c.1091del	p.(N364Tfs*5)	Ex2	3
<i>BBS10</i>	NM_024685.3	c.1958_1967del	p.S653Ifs*4	Ex2	4
<i>BBS10</i>	NM_024685.3	c.2121dup	p.(K708*)	Ex2	3
<i>BBS12</i>	NM_152618.2	c.1589T>C	p.(L530P)	Ex3	5
<i>BBS12</i>	NM_152618.2	c.2102C>A	p.(S701*)	Ex3	6
<i>BBS2</i>	NM_031885.3	c.1237C>T	p.(R413*)	Ex11	5
<i>ARL6</i>	NM_032146.3	c.123+1119del	p.(?)	Intron 4	7
<i>BBS5</i>	NM_152384.2	c.2T>A	p.(M1K)	Ex1	5
<i>TTC8</i>	NM_144596.2	c.1049+2_1049+4del	p.(?)	Ex11	8

Abbreviations: RefSeq Id: Reference sequence identity

References

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2. Beales, P.L. *et al.* Genetic interaction of BBS1 mutations with alleles at other BBS loci can result in non-Mendelian Bardet-Biedl syndrome. *Am J Hum Genet* **72**, 1187-99 (2003).
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Supplementary Table S3: Transcripts and exons of 21 BBS genes sequenced in targeted next generation sequencing panel.

Chr#: Position	Gene	Transcript	Exons
chr3:97486937-97516899	<i>ARL6</i>	NM_032146	2, 3, 4, 5, 6, 7, 8
chr3:97486937-97516899		NM_177976	2, 3, 4, 5, 6, 7, 8
chr10:112660235-112677929	<i>BBIP1</i>	NM_001195304	0, 1, 2, 3
chr10:112660111-112677929		NM_001195305	0, 1, 2
chr10:112660111-112677929		NM_001195306	0, 1, 2
chr10:112660111-112677929		NM_001195307	0, 1
chr12:76739586-76742152	<i>BBS10</i>	NM_024685	0, 1
chr4:123663033-123665186	<i>BBS12</i>	NM_001178007	2
chr4:123663033-123665186		NM_152618	1
chr11:66278116-66278183	<i>BBS1</i>	NM_024649	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16
chr16:56518666-56553788	<i>BBS2</i>	NM_031885	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16
chr15:73016911-73029934	<i>BBS4</i>	NM_001252678	6, 7, 8, 9, 10, 11, 12, 13, 14
chr15:72978554-73029934		NM_033028	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15
chr2:170336049-170361098	<i>BBS5</i>	NM_152384	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11
chr4:122749289-122791482	<i>BBS7</i>	NM_018190	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17

chr4:122747008-122791482		NM_176824	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18
chr7:33185850-33644844	<i>BBS9</i>	NM_001033604	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20, 21
chr7:33185850-33644844		NM_001033605	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20, 21
chr7:33185850-33644844		NM_014451	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20
chr7:33185850-33644844		NM_198428	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20, 21, 22
chr1:32667522-32670855	<i>CCDC28B</i>	NM_024296	1, 2, 3, 4, 5
chr12:88442954-88535098	<i>CEP290</i>	NM_025114	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52
chr22:37154348-37171765	<i>IFT27</i>	NM_001177701	0, 1, 2, 3, 4, 5, 6
chr22:37154348-37171765	<i>IFT27</i>	NM_006860	0, 1, 2, 3, 4, 5, 6
chr3:45867799-45883497	<i>LZTFLI</i>	NM_020347	0, 1, 2, 3, 4, 5, 6, 7, 8, 9
chr20:10385888-10394176	<i>MKKS</i>	NM_018848	0, 1, 2, 3
chr20:10385888-10394176		NM_170784	0, 1, 2, 3
chr17:56283433-56296886	<i>MKSI</i>	NM_001165927	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17
chr17:56283433-56296605		NM_017777	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17
chr2:63349134-63665031	<i>WDPCP</i>	NM_001042692	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11
chr3:45867799-45879509	<i>LZTFLI</i>	NM_001276378	0, 1, 2, 3, 4, 5, 6, 7, 8

chr3:45867812-45954652		NM_001276379	0, 1, 2, 3, 4, 5, 6, 7, 8
chr3:97486937-97516899	<i>ARL6</i>	NM_001278293	1, 2, 3, 4, 5, 6, 7
chr14:89291037-89343760	<i>TTC8</i>	NM_001288781	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15
chr14:89307857-89343760		NM_001288782	5, 6, 7, 8, 9, 10, 11, 12, 13
chr14:89323526-89343760		NM_001288783	8, 9, 10, 11, 12, 13, 14
chr1:32667522-32670405	<i>CCDC28B</i>	NM_001301011	1, 2, 3, 4
chr1:243419461-243663093	<i>SDCCAG8</i>	NM_006642	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17
chr8:94770761-94828686	<i>TMEM67</i>	NM_001142301	2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28
chr8:94767128-94828686		NM_153704	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 19, 20, 21, 22, 23, 24, 25, 26, 27
chr9:119460007-119461989	<i>TRIM32</i>	NM_001099679	1
chr9:119460007-119461989	<i>TRIM32</i>	NM_012210	1
chr14:89291037-89343760	<i>TTC8</i>	NM_144596	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15
chr14:89291037-89343760		NM_198309	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14
chr14:89291037-89343760		NM_198310	1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13
chr2:63349134-63815419	<i>WDPCP</i>	NM_015910	0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17

Abbreviation: Chr #: Chromosome number

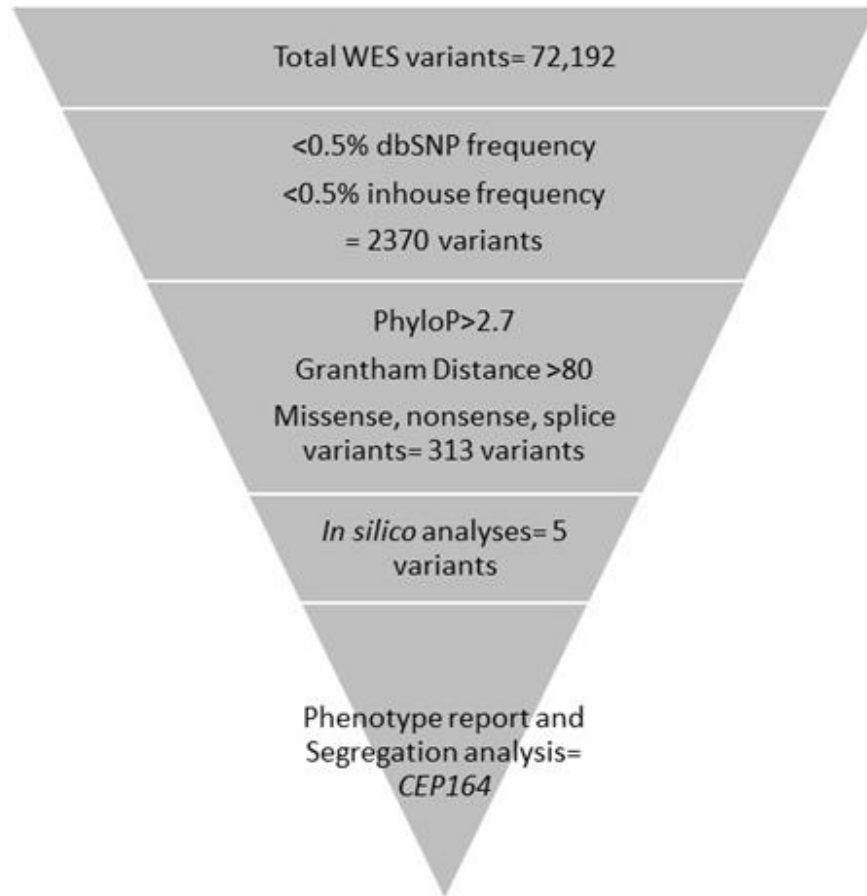
Supplementary Table S4: Primers used for the Sanger sequencing of RT-PCR products in minigene splicing assay.

Primer	Primer Sequence 5'-3'
pCI-Neo-Rho-Insert Fw	cggaggtaacaacgagtct
pCI-Neo-Rho-Insert Rev	aggtgtaggggatgggagac

Supplementary Table S5: Homozygosity mapping on WES data of the proband from family F05 using GRCh37/hg19 reference assembly. The cutoff selected for significant homozygous regions is 1Mb.

Rank	Chr	From Mb-to Mb	Size Mb	Retnet genes	Genes from WES
1	8	38,027,483-81,892,766	43.9	<i>ADAM9, RP1, TTP1, PXMP3</i>	
2	14	34,247,631-59,730,423	25.5		
3	12	12,243,828-29,936,501	17.7	<i>PDE6H</i>	
4	11	104,871,322-120,823,608	16.0	<i>C1QTNF5, MFRP</i>	<i>CEP164</i>
5	21	35,276,440-48,063,476	12.8		
6	6	112,671,813-124,759,146	12.1		
7	1	154,728,279-164,853,647	10.1	<i>SEMA4A</i>	
8	13	96,705,739-106,124,923	9.4		
9	4	62,861,964-71,232,388	8.4		
10	11	28,232,879-34,668,143	6.4		
11	14	89,656,709-94,908,876	5.3	<i>FBLN5</i>	
12	1	177,250,670-181,479,906	4.2		
13	17	41,466,121-45,234,407	3.8		
14	13	111,368,316-114,838,992	3.5	<i>GRK1</i>	
15	17	1,490,409-4,859,123	3.4	<i>PRPF8</i>	
16	11	46,907,827-48,346,681	1.4		
17	19	43,877,081-45,117,163	1.2		
18	11	1,630,342-2,790,019	1.2		

Abbreviations: Chr: Chromosome, Mb: Megabases, WES: Whole exome sequencing

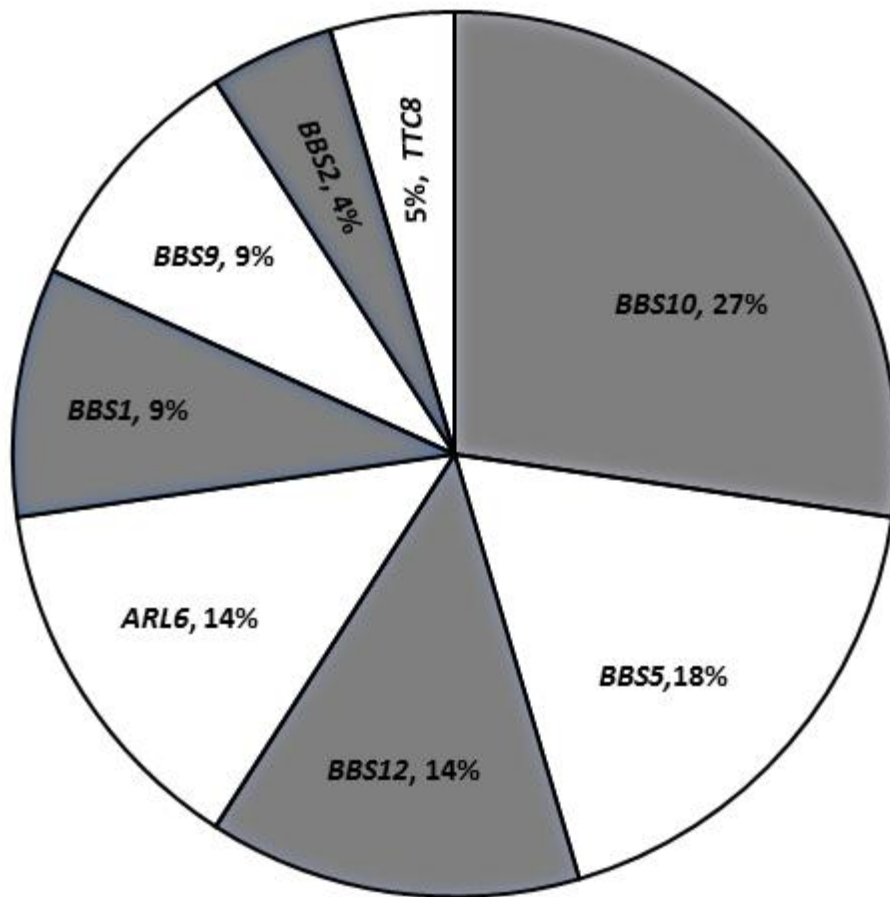


Supplementary Fig. S1: WES data filter summary for family F05.

ARL6, c.534A>G; p.=



Supplementary Fig. S2: *In silico* analysis of novel synonymous mutation in *ARL6* (c.534A>G; p.=) identified causative factor in family F01. The changes in scores between upper panel (reference sequence) and lower panel (mutated sequence) indicate an effect on splicing of exon 8. Source: Alamut visual Version 2.7.1 from Interactive Biosoftware (<http://www.interactive-biosoftware.com>).



Supplementary Fig. S3: Genetic frequency of eight BBS-associated genes from 22 Pakistani BBS families. Eighteen families were previously reported while 4 BBS families are part of this study, where most of the Pakistani families carry mutations in *BBS10*.