

**Figure 2.**

A 3.7-Mb microdeletion including *SCN2A* and *SCN1A* in patient 409. **(A)** Relative depth of coverage ratio for patient 409 indicates a microdeletion encompassing *SCN2A* and *SCN1A*. Different colors distinguish the target genes. **(B)** The array profile clearly shows a 3.7-Mb microdeletion at 2q24.3 in this patient. Thirteen RefSeq genes, including *SCN2A* and *SCN1A*, lie within the microdeletion (bottom). **(C)** Breakpoint-specific PCR analysis of the patient's family. Primers flanking the deletion were able to amplify a 1,607-bp product from the patient only, indicating that the translocation occurred de novo. **(D)** Deletion junction sequence. The top, middle, and bottom strands show the proximal, deleted, and distal sequences, respectively. A single inserted nucleotide (colored in red) was identified at the breakpoint.

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the capture of target genes to the detection of mutations and CNVs—can be done within a week, our workflow provides a fast, sensitive, and comprehensive genetic testing method for patients with epilepsy.

Whole-exome sequencing will reveal novel mutations in unexpected genes in patients with EOEEs. For example, *KCNQ2* mutations, which cause benign familial neonatal seizures (Biervert et al., 1998; Charlier et al., 1998), were identified in patients with OS by whole exome sequencing (Saitou et al., 2012a). Similarly, screening known and potential candidate genes in patients with EOEEs will reveal novel mutations in unexpected genes, in addition to mutations in well-known genes.

In our target capture analysis, some exons of genes such as *ARX* and *FOXG1* were insufficiently sequenced because repeat sequences hampered the design of capture probes. Repeat sequences also interfere with appropriate mapping of

sequence reads, resulting in low coverage. For these exons, Sanger sequencing should be added for complete analysis.

In conclusion, a rapid and efficient system of target capture sequencing can be applied to the comprehensive genetic analysis of EOEEs. Point mutations, small indels, and CNVs are all detected by this method, confirming the potential of this approach for efficient genetic testing.

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DISCLOSURE

None of the authors has any conflicts of interest to disclose. We confirm that we have read the Journal's position on issues involved in ethical publication and affirm that this report is consistent with those guidelines.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Figure S1. Flow chart of our variant detection and copy number analysis scheme.

Figure S2. Insufficient coverage of reads in two genes rich in repetitive sequences.

Clinical Correlations of Mutations Affecting Six Components of the SWI/SNF Complex: Detailed Description of 21 Patients and a Review of the Literature

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Mutations in the components of the SWItch/sucrose nonfermentable (SWI/SNF)-like chromatin remodeling complex have recently been reported to cause Coffin–Siris syndrome (CSS), Nicolaides–Baraitser syndrome (NCBRS), and *ARID1B*-related intellectual disability (ID) syndrome. We detail here the genotype–phenotype correlations for 85 previously published and one additional patient with mutations in the SWI/SNF complex: four with *SMARCB1* mutations, seven with *SMARCA4* mutations, 37 with *SMARCA2* mutations, one with an *SMARCE1* mutation, three with *ARID1A* mutations, and 33 with *ARID1B* mutations. The mutations were associated with syndromic ID and speech impairment (severe/profound in *SMARCB1*, *SMARCE1*, and *ARID1A* mutations; variable in *SMARCA4*, *SMARCA2*, and *ARID1B* mutations), which was frequently accompanied by agenesis or hypoplasia of the corpus callosum. *SMARCB1* mutations caused “classical” CSS with typical facial “coarseness” and significant digital/nail hypoplasia. *SMARCA4* mutations caused CSS without typical facial coarseness and with significant digital/nail hypoplasia. *SMARCA2* mutations caused NCBRS, typically with short stature, sparse hair, a thin vermillion of the upper lip, an everted lower lip and prominent finger joints. A *SMARCE1* mutation caused CSS without typical facial coarseness and with significant digital/nail hypoplasia. *ARID1A* mutations caused the most severe CSS with severe physical complications. *ARID1B* mutations caused CSS without typical facial coarseness and with mild digital/nail hypoplasia, or caused syndromic ID. Because of the common underlying mechanism and overlapping clinical features, we propose that these conditions be referred to collectively as “SWI/SNF-related ID syndromes”. © 2013 Wiley Periodicals, Inc.

Key words: Coffin–Siris syndrome; SWI/SNF complex; *SMARCB1*; *SMARCA4*; *SMARCA2*; *SMARCE1*; *ARID1A*; *ARID1B*; Nicolaides–Baraitser syndrome; intellectual disability (ID)

INTRODUCTION

Coffin–Siris syndrome (CSS; OMIM 135900) was first described by Coffin and Siris [1970]. It is a rare congenital anomaly syndrome characterized by developmental delay or intellectual disability (ID), coarse facial appearance, feeding difficulties, frequent infections, and hypoplastic-to-absent fifth fingernails and fifth distal phalanges [Devy and Baraitser, 1991; Fleck et al., 2001; Schrier et al., 2012]. We recently reported on mutations in six genes encoding components of the SWItch/sucrose nonfermentable (SWI/SNF)-like chromatin remodeling complex in 20 of 23 patients clinically diagnosed with CSS: *SMARCB1* in four patients, *SMARCA4* in six, *SMARCA2* in one, *SMARCE1* in one, *ARID1A* in three, and *ARID1B* in five [Tsurusaki et al., 2012]. In the same journal issue, truncating

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mutations in *ARID1B* were reported in three patients with CSS and microdeletions encompassing *ARID1B* were reported in three patients with ID and remnants of CSS [Santen et al., 2012]. Furthermore, missense mutations of *SMARCA2* were reported in 36 patients with Nicolaides–Baraitser syndrome (NCBRS; OMIM#601358) [van Houdt et al., 2012]. NCBRS was first described by Nicolaides and Baraitser [1993] and it is a recently delineated condition characterized by severe ID with absent/limited speech, seizures, short stature, sparse hair, typical facial characteristics, brachydactyly, prominent finger joints, and broad distal phalanges; the main differential diagnosis is CSS [Sousa et al., 2009]. *ARID1B* has also been reported to be a cause of ID. Nagamani et al. [2009] reported four patients with interstitial deletion of 6q25.2–q25.3 including *ARID1B*, all of whom manifested microcephaly, developmental delay, facial characteristics, and hearing impairment, and two of whom had agenesis of the corpus callosum. Halgren et al. [2012] reported eight patients with haploinsufficiency of *ARID1B* (de novo chromosomal translocation involving *ARID1B* in one, intragenic deletions in three, and microdeletion including *ARID1B* in four), who manifested agenesis or hypoplasia of the corpus callosum, ID with speech impairment, and autism. Hoyer et al. [2012] very recently concluded that haploinsufficiency of *ARID1B* is a relatively frequent cause of moderate-to-severe ID from their findings that 0.9% (8/887) of patients with unexplained ID had truncating mutations in the gene. Michelson et al. [2012] also very recently reported a patient with an interstitial 1.19 Mb deletion of 6q25.2 including *ARID1B* and *ZDHHC14*, who manifested global developmental delay, facial characteristics, dysgenesis of the corpus callosum, limb anomalies, and genital hypoplasia.

To delineate the clinical consequences of mutations affecting components of the SWI/SNF complex (CSS, NBS, and *ARID1B*-related ID syndrome), we report the individual clinical information for 20 previously reported patients [Tsurusaki et al., 2012] as well as an additional patient with an *SMARCA4* mutation. Furthermore, we create a comprehensive list of all reported patients (including our series) with mutations affecting components of the SWI/SNF complex (Tables Ia–Ic), which will be helpful when discussing similarities and differences among these conditions.

CLINICAL REPORTS

SMARCB1 Mutations

SMARCB1-1 (Subject 4 [Tsurusaki et al., 2012]; Fig. 1a–i): She was born at 42 weeks of gestation after an uncomplicated prenatal period. Her birth weight was 3,008 g (−0.5 SD). She had: cleft palate with exudative otitis media; congenital dislocation of the right hip; pectus excavatum; sucking/feeding difficulty. She underwent surgical correction of cleft palate and insertion of ventilation tubes at age 2 6/12 years. Hearing aids were required for bilateral, severe, mixed hearing impairment with a threshold of 80–90 dB. At age 4 years, she developed tonic seizures, which were treated with carbamazepine. Scoliosis, found at age 2 years, progressed with a Cobb angle of $\approx 150^\circ$. She showed hypotonia and motor development was severely delayed: she raised her head at age 1 8/12 years, sat alone at 2 3/12 years, and walked independently at 7 years. From ages 12 to 18 years, she vomited frequently and had recurrent infections. She had multiple dental caries, treated under general anesthesia at age 16 years. At age 21 years, she weighs 30 kg (−3.4 SD), her height is 112.5 cm (−8.4 SD), and her occipito-frontal circumference (OFC) is 51.2 cm (−2.9 SD). She understands simple commands in daily life, expresses herself with gestures, and likes to play portable personal computer games, but speaks no words. She has serious behavioral problems such as impulsiveness, hyperactivity, and self-injurious behaviors (including skin picking). She becomes exhausted easily.

SMARCB1-2 (Subject 11 [Tsurusaki et al., 2012]): Her prenatal period was complicated by intrauterine growth retardation. She was born at 38 weeks of gestation. Her birth weight was 2,088 g (−1.8 SD), length was 42 cm (−2.9 SD), and OFC was 33 cm (0 SD). She had a small ventricular septal defect (VSD). Diaphragmatic hernia was corrected surgically at age 5 months. She had sucking/feeding difficulties. She showed hypotonia and motor development was severely retarded: she rolled over at age 3 years. Generalized seizures developed and were controlled with valproic acid. Magnetic resonance imaging (MRI) of the brain showed cerebellar hypoplasia and Dandy–Walker malformation. She had visual impairment that was corrected with spectacles; hearing impairment with a threshold of 60 dB in the right ear and 50 dB in the left ear was noted. At age 7 years, her weight is 12 kg (−3.1 SD) and height is 105 cm (−2.7 SD). She sits for several seconds with her hands, distinguishes her family members from others, and smiles when called by her name. She has visual and hearing impairment.

SMARCB1-3 (Subject 21 [Tsurusaki et al., 2012]; Fig. 1j–p): Her prenatal period was complicated by intrauterine growth retardation and oligohydramnios. She was born at 38 weeks of gestation, followed by resuscitation through endotracheal intubation. Her

birth weight was 1,746 g (−2.6 SD). Surfactant treatment was undertaken for pulmonary hemorrhage. She had micrognathia, exotropia, and a dark complexion. She suffered from complex partial seizures. She sucked poorly and then had feeding difficulty associated with gastroesophageal reflux (GER), which required gastrostomy. She showed hypotonia with severe delay in motor development. A hypoplastic corpus callosum was observed. At age 7 years, she weighs 12 kg (−3.0 SD), has a height of 97 cm (−4.5 SD), and an OFC of 44 cm (−5.1 SD). She is unable to sit alone, and moves by rolling over. She cannot communicate with others or speak any words, but smiles when she appears to be happy.

SMARCB1-4 (Subject 22 [Tsurusaki et al., 2012]): He was born at 37 weeks of gestation. His birth weight was 2,784 g (+0.2 SD). He was admitted to hospital as a newborn for treatment of transient tachypnea. He had pyloric stenosis that was corrected surgically. He sucked poorly and then had feeding difficulties associated with GER, requiring gavage-feeding and resulting in failure to thrive. He showed hypotonia and motor development was severely delayed. MRI of the brain showed hypoplasia of the corpus callosum. He suffered from recurrent respiratory tract infections. At age 2 years, he weighed 9.7 kg (−2.3 SD), his height was 83.4 cm (−2.2 SD), and had an OFC of 43 cm (−3.3 SD). At age 3 years, he rolls over, but cannot sit alone or speak words.

SMARCA4 Mutations

SMARCA4-1 (Subject 9 [Tsurusaki et al., 2012]; Fig. 2a–c): He was born at 39 weeks of gestation. His birth weight was 2,880 g. Sucking or feeding difficulties were not reported, but abdominal distension occurred in infancy and constipation was frequent in childhood. Possible seizures developed once at age 1 2/12 years with unconsciousness but without electrocardiographic abnormalities. The submucosal cleft palate was corrected surgically at age 1 6/12 years. Congenital torticollis with vestigial (right) and shortened (left) cleidomastoid muscles was corrected surgically. His right chest was funnel-shaped with a hypoplastic right pectoral major muscle. Exudative otitis media in the left ear was recurrent. He showed hypotonia in his infancy and motor development was mildly delayed: he raised his head at age 4 months, sat alone at 10 months, crawled at 11 months, and stood alone at 1 3/12 years. His hair has been bristly and gray-streaked since childhood. His upper teeth were misaligned. At age 18 years, his height is 159 cm (−1.8 SD) and OFC is 53 cm (−2.3 SD). He can walk independently, talk (albeit with a stutter), and understand almost everything necessary for daily life. He has myopia and mild astigmatism which are corrected with spectacles. He has nocturnal enuresis which he finds hard to control.

SMARCA4-2 (Subject 7 [Tsurusaki et al., 2012]; Fig. 2d–g): His prenatal period was complicated by intrauterine growth retardation. He was born at 40 weeks of gestation. His birth weight was 2,250 g (−2.2 SD). He showed respiratory insufficiency and had sucking/feeding difficulties associated with laryngomalacia. He had bilateral ptosis, myopia, lacrimal duct stenosis, bilateral sensorineural hearing loss, and ankyloglossia. He showed hypotonia and motor development was severely delayed: he raised his head in late infancy, sat alone at age 2 years, and walked independently at 6 years. At age 20 years, he weighs 60 kg (−0.2 SD), has a height of

TABLE IA. Clinical Features of Patients With Mutations in the Components of SWI/SNF Complex

Gene	SMARCB1				SMARCA4						
Patient (subject no. in the previous report§)	1 [4]	2 [11]	3 [21]	4 [22]	1 [9]	2 [7]	3 [5]	4 [16]	5 [25]	6 [17]	7
Age at publication (years of age)	21	?	?	2	18	20	9	11	16	4	8
Sex	F	F	F	M	M	M	M	M	F	M	F
Mutation	p.Lys364del	p.Arg377His	p.Lys364del	p.Lys364del	p.Lys546del	p.Thr859Met	p.Arg885Cys	p.Leu921Phe	p.Met1011Thr	p.Arg1157Gly	p.Arg885His
<i>Growth</i>											
Prenatal growth (birth weight/length)#	-0.5/?	-1.8/-2.9	-2.6/?	+0.2/?	-	-2.2/?	-1.2/-0.9	-1.7/-1.6	-1.0/-1.9	-1.1/-2.3	-2.6/-2.7
Postnatal growth (weight/height)†	-3.4/-8.4	-3.1/-2.7	-3.0/-4.5	-2.3/-2.2	?/-1.8	-0.2/-2.6	-1.5/-3.2	-1.8/-3.1	-1.9/-1.9	-3.0/-3.4	-1.9/-1.8
<i>Psychomotor</i>											
Developmental delay/intellectual disability‡	Severe	Severe	Severe	Severe	Mild	Severe	Severe	Severe	Severe	Severe	Moderate
Speech delay	NW	NW	NW	NW	Mild	SW	NW	NW	SC	NW	Mild
Seizures	+	+	-	-	+	-	-	+	-	-	-
Hypotonia	+	+	+	+	+	+	+	+	+	+	+
Autistic features/behavioral abnormalities	HA,Im,SH	-	-	-	-	HA, Im	ASD (HA, HS, Ob, SH)	-	RB	-	HA
Microcephaly (≤2 SD) [SD score]	+[-2.9]	-	+[-5.1]	+[-3.3]	+[-2.3]	+[-3.8]	+[-3.6]	+[-2.9]	+[-2.3]	+[-2.7]	+[-3.0]
Brain anomaly	-	CH,DW	HCC	HCC	-	-	HCC,HCV	-	-	HCC	-
<i>Craniofacial</i>											
Sparse hair	+	+	+	+	-	+	-	+	-	+	-
Thick eyebrows	+	+	+	+	+	+	+	+	+	+	+
Thick eyelashes	+	+	+	+	+	+	+	+	+	+	+
Ptosis	-	+	+	+	+	+	+	+	-	+	+
Abnormal ears	+	+	+	+	-	+	+	+	+	+	-
Nasal bridge	Broad	Broad	Broad	Broad	Narrow	Narrow	Normal	Flat	Flat	Flat	Flat
Thick, anteverted alae nasi	-	-	-	+	-	-	-	-	-	-	-
Wide mouth	+	+	+	+	-	-	+	-	+	-	+
Philtrum	Broad	-	Long	Long	Short	Short	Short	-	Short	-	Short
Upper lip vermilion feature	Thin	-	Thin	Thick	Everted	Everted	Everted	Thin	Everted	-	-
Thick lower lip vermilion	+	+	+	+	-	+	+	+	+	+	+
Palatal abnormality	C	C	H	H	SMCP	H	C	H	H	C	H
<i>Skeletal-limb</i>											
Hypoplastic/absent fifth finger/toe	Fi/T	Fi/T	Fi/T	Fi/T	Fi	Fi/T	T	Fi/T	Fi/T	Fi/T	T
Hypoplastic/absent nail (fifth finger/toe)	Fi/T	+	Fi/T	Fi/T	Fi	Fi/T	T	Fi/T	Fi/T	Fi/T	T
Hypoplastic/absent nail (other fingers/toes)	Fi/T	-	Fi/T	Fi/T	Fi	Fi/T	T	Fi/T	Fi/T	Fi/T	-
Prominent interphalangeal joints	+	-	-	-	-	+	-	-	+	-	-
Prominent distal phalanges	+	+	-	-	+	+	+	+	+	-	-
Scoliosis/spinal abnormalities	+	-	+	+	-	+	-	-	-	-	-
Joint laxity	-	-	+	+	-	+	+	-	+	-	+
<i>Others</i>											
Hirsutism	+	-	+	+	+	+	+	+	+	+	+
Congenital heart defects	-	VSD	-	+	-	-	-	VSD, PDA	-	MA, PA, SRV, AtSD, PDA	+
Genitourinary defects	-	-	-	+	-	-	Cr	-	-	Cr	-
Gastrointestinal abnormalities	-	-	GER	PS	-	GOO	Co	DU	Co	GER	GER
Inguinal (I)/umbilical (U) hernia	-	-	I	I	-	I	-	I	-	Om	I/U
Sucking difficulty	+	+	+	+	-	+	+	+	+	+	+
Feeding difficulty	+	+	+	+	-	+	+	+	+	+	+
Frequent vomiting	+	+	+	+	-	+	+	+	+	+	+
Hearing impairment	+	+	+	+	-	+	+	-	+	-	-
Visual impairment	+	+	+	+	+	+	+	+	+	-	-
Recurrent infections	+	-	+	+	-	+	-	+	+	+	+

+, present; -, absent; blank, data not available; §, Tsurusaki et al. [2013]; #, SD score; †, SD score; ‡, at latest assessment; ASD, autism spectrum disorder; AtSD, atrial septal defect; C, cleft palate; CH, cerebellar hypoplasia; Co, constipation; Cr, cryptorchidism; DU, duodenal ulcer; DW, Dandy-Walker malformation; F, female; Fi, finger; GER, gastroesophageal reflux; GOO, gastric outlet obstruction; H, high palate; HA, hyperactivity; HCC, hypoplastic corpus callosum; HCV, hypoplastic cerebellar vermis; HS, hypersensitivity; I, inguinal; Im, impulsiveness; M, male; MA, mitral atresia; NW, no words; Ob, obsession; Om, omphalocele; PA, pulmonary atresia; PDA, patent ductus arteriosus; PS, pyloric stenosis; RB, repetitive behavior; SC, simple conversation; SH, self-harming behavior; SMCP, submucosal cleft palate; SRV, single right ventricle; SW, several words; T, toe; VSD, ventricular septal defect

TABLE IB. Clinical Features of Patients With Mutations in the Components of SWI/SNF Complex

Gene	SMARCA2	SMARCE1	ARID1A	ARID1B						
Patient (subject no. in the previous report[s])	1 [19]	NCBRS (n = 36)	1 [24]	1 [3]	2 [6]	3 [8]	1 [1]	2 [15]	3 [23]	4 [10]
Age at publication (years of age)	8	2.3/12–32	14	2	1	10	13	7	10	11
Sex	M	21M,15F	F	M	M	M	M	M	F	M
Mutation	Partial deletion [van Houdt et al., 2012]	Missense	p.Tyr73Cys	p.Ser11Alafs*91	p.Gln920*	p.Arg1335*	p.Ile560Glyfs*89	p.Gln635*	p.Arg1102*	p.Asp1878Metfs*96
<i>Growth</i>										
Prenatal growth (birth weight/length)#	–2.6/–3.1	10/34	–2.3/?	–1.0/–1.4	–0.6/–1.1	–	–0.9/+0.4	–1.2/–1.2	+0.5/–1.7	–1.0/–1.1
Postnatal growth (weight/height)†	–2.1/–3.8	19/36	–4.4/–6.8	–6.2/–8.9	–5.5/–8.2	50th/3rd	–0.5/–1.4	–1.1/–1.1	+1.5/–1.3	–0.5/–2.4
<i>Psychomotor</i>										
Developmental delay/intellectual disability‡	Moderate	3Mild,9Moderate, 24Severe	Severe	Severe	Severe	Severe	Moderate	Severe	Mild	Mild
Speech delay	NW	13/33NW	NW	NW	NW	NW	NW	NW	Mild	Mild
Seizures	+	22/35	+	–	–	–	–	+	+	–
Hypotonia	+	–	+	+	+	–	+	+	+	+
Autistic features/behavioral abnormalities	–	–	–	–	–	HA	ASD (HA, Im, O)	ASD (Im)	–	–
Microcephaly (≤ 2 SD) (SD score)	+ [–3.3]	19/34	+ [–6.7]	+ [–3.7]	+ [–3.1]	– (50th)	– [+0.7]	– [–1.4]	– [+1.8]	– [–1.1]
Brain anomaly	–	–	–	ACC, CH, DW	ACC	HCC	–	–	ACC, Colp	–
<i>Craniofacial</i>										
Sparse hair	–	35/36	+	+	+	+	+	–	–	+
Thick eyebrows	+	–	+	+	+	+	+	+	+	+
Thick eyelashes	+	23/35	+	+	+	+	+	+	+	+
Ptosis	+	–	+	–	–	–	–	–	–	–
Abnormal ears	+	–	+	+	+	+	+	+	–	+
Nasal bridge	Broad	22/36Narrow	Narrow	Broad	Broad	–	Broad	Broad	Broad	Broad
Thick, anteverted alae nasi	+	32/36	–	–	–	–	–	+	–	–
Wide mouth	+	34/36	+	+	+	+	+	+	–	+
Philtrum	Broad	31/36Broad, 29/36Long	Long	Short	Broad	Long	Broad	Broad	Long	Long
Upper lip vermilion feature	–	27/36Thin	–	–	Everted	–	Thin	Thick	Thin	Thin
Thick lower lip vermilion	+	32/36	+	+	+	+	–	+	+	+
Palatal abnormality	–	–	C	C	C	H	H	H	H	H
<i>Skeletal–limb</i>										
Hypoplastic/absent fifth finger/toe	–	–	Fi/T	–	Fi/T	Fi/T	Fi/T	Fi/T	Fi	Fi/T
Hypoplastic/absent nail (fifth finger/toe)	–	–	Fi/T	+	Fi/T	Fi/T	Fi/T	T	Fi	Fi/T
Hypoplastic/absent nail (other fingers/toes)	–	–	Fi/T	Fi/T	–	–	–	–	–	Fi/T
Prominent interphalangeal joints	+	28/35	–	–	–	–	–	–	–	+
Prominent distal phalanges	+	21/35	–	–	–	+	+	+	–	+
Scoliosis/spinal abnormalities	+	10/34	–	+	–	–	+	–	–	+
Joint laxity	–	–	–	+	+	–	+	+	–	+
<i>Others</i>										
Hirsutism	+	–	+	+	+	+	+	+	+	+
Congenital heart defects	–	6/34	TR, MR, AS	CoA, AtSD, VSD	CoA, AS, VSD	AtSD	–	–	–	+
Genitourinary defects	Cr, HS	13/20Cr, 2/32VUR	–	–	HS, Cr	–	–	–	–	Cr
Gastrointestinal abnormalities	–	–	–	–	AA, RUF	IO, GER	–	–	–	–
Inguinal (I)/umbilical (U) hernia	I/U	14/35	–	–	–	I	–	–	–	–
Sucking difficulty	+	–	+	+	+	+	+	+	+	+
Feeding difficulty	+	–	+	+	+	+	+	+	+	+
Frequent vomiting	+	–	–	–	–	–	–	–	–	–
Hearing impairment	+	–	–	+	–	+	–	+	–	–
Visual impairment	+	–	–	–	–	–	+	–	+	–
Recurrent infections	–	+	+	+	+	+	+	+	+	+

+, present; –, absent; blank, data not available; §, Tsurusaki et al. [2013]; #, SD score; †, SD score; ‡, at latest assessment; AA, anal atresia; ACC, agenesis of corpus callosum; AS, aortic stenosis; ASD, autism spectrum disorder; AtSD, atrial septal defect; C, cleft palate; CH, cerebellar hypoplasia; CoA, coarctation of aorta; Colp, colpocephaly; Cr, cryptorchidism; DW, Dandy-Walker malformation; F, female; Fi, finger; GER, gastroesophageal reflux; H, high palate; HA, hyperactivity; HCC, hypoplastic corpus callosum; HS, hypospadias; Im, impulsiveness; IO, intestinal obstruction; M, male; MR, mitral regurgitation; NBS, Nicolaides-Baraitser syndrome; NW, no words; O, obsession; RUF, retrourethral fistula; T, toe; TR, tricuspid regurgitation; VSD, ventricular septal defect; VUR, vesicoureteral reflux; fractions in the previous publications show patient numbers "feature-positive/data available"

TABLE 1C. Clinical Features of Patients With Mutations in the Components of SWI/SNF Complex

Gene	SWI5		ARID1B		ARID1A	
Patient (subject no. in the previous report§)	5 [12]	n = 4	n = 8	n = 9	n = 6	n = 1
Age at publication (years of age)	19	11/12–4	3–46	3 3/12–20	2–40	2
Sex	M	2M,2F	2M,6F	4M,5F	1M,5F	M
Mutation	Del (9.2Mb)	Del (3.77–13.81Mb) [Nagamani et al., 2009]	?Del (0.2–14.5Mb),1Tra [Halgren et al., 2012]	1Del(2.5Mb),1Dup(exon 5/6),3Ns,4Fs [Hoyer et al., 2012]	3Del(0.73–2.72),2Ns,1Fs [Santen et al., 2012]	Del(1.19Mb) [Michelson et al., 2012]
Growth						
Prenatal growth (birth weight/length)#	–2.3/–3.7	3/4		0/9	0/6	–
Postnatal growth (weight/height)†	–2.3/–6.1	1/3	5/7	3/9	3/6	–
Psychomotor						
Developmental delay/mental retardation‡	Severe	4/4(2Moderate, 2Severe)	8/8	5Moderate,4Severe	2Moderate, 4Severe	Moderate
Speech delay	NW	4/4(3NW, 1SW)	8/8(3NW,5SW)	9/9(2NW, 2SW, 3Se)	6/6(2NW, 3Severe, 1Moderate)	NW
Seizures	+	1FC	3/6	3/9		
Hypotonia	–	2	7/8	7/9		+
Autistic features/behavioral abnormalities	HA		5/7	1	1	
Microcephaly (≤2 SD) (SD score)	+	4/4	1/6	2/9	0/6	–
Brain anomaly	–?	2/3(ACC + Colp)	4/5(1ACC, 3HCC, 1HCV)	0/6	4/4(3ACC, 1HCC, 2Colp)	HCC
Craniofacial						
Sparse hair	+		1	1	1	
Thick eyebrows	+		1		6/6	
Thick/long/prominent eyelashes	–					
Ptosis	–	1				
Abnormal ears	+	2	1	7/9		
Nasal bridge	Narrow	3Broad	3Broad			
Thick, anteverted alae nasi						Broad
Wide mouth	–		2	1		+
Philtrum	Long	2Long	2Long	1Long		
Upper lip vermilion feature		2Thin	3Thin,1Thick	6/9(6Thin)		Thin
Thick lower lip vermilion	–	2Thick	3			
Palatal abnormality	H	2H		2/9(2H)		
Skeletal-limb						
Hypoplastic/absent fifth finger/toe	Fi/T				2/6(2Fi)	
Hypoplastic/absent nail (fifth finger/toe)	+		1T	1T	2/6(2Fi)	
Hypoplastic/absent nail (other fingers/toes)				1?	1/6	
Prominent interphalangeal joints	+					
Prominent distal phalanges	–	1				
Scoliosis/spinal abnormalities	+		1	1		
Joint laxity	–		3/4		3/5	
Others						
Hirsutism	+		3/4	2		3/6
Congenital heart defects	MR	1AtSD	1AtSD	2/9(1AtSD)	1AtSD	
Genitourinary defects	–	1PSW	1Cr, 1DoUr, 2RL	2Cr, 1MU	1RL, 1DoUr	
Gastrointestinal abnormalities	GU, GER		1Co, 1AA	1AA		
Inguinal (I)/umbilical (U) hernia	–					
Sucking difficulty	–	3	4			+
Feeding difficulty	–		5	+		
Frequent vomiting	–					
Hearing impairment	–	4	1/2	1/9	+	
Visual impairment	–	1St, 1Am	2Hy, 2My, 2St, 1 Cat, 1Ny		3St/9,4My	2St
Recurrent infections	+	1	1			

+, present; –, absent; blank, data not available; §, Tsurusaki et al. [2013]; #, SD score; †, SD score; ‡, at latest assessment; AA, anal atresia; ACC, agenesis of corpus callosum; Am, amblyopia; AtSD, atrial septal defect; Cat, cataract; Co, constipation; Colp, colpocephaly; Cr, cryptorchidism; Del, deletion; Dup, duplication; DoUr, double ureter; F, female; Fi, finger; Fs, frameshift mutation; GER, gastrointestinal reflux; GU, gastric ulcer; H, high palate; HA, hyperactivity; HCC, hypoplastic corpus callosum; HCV, hypoplastic cerebellar vermis; Hy, hypermetropia; M, male; MR, mitral regurgitation; MU, megareuter; My, myopia; Ns, nonsense mutation; NW, no words; Ny, nystagmus; PSW, penoscrotal webbing; RL, renal lithiasis; Se, sentences; St, strabismus; SW, several words; T, toe; Tra, translocation; fractions in the previous publications show patient numbers "feature-positive/data available"



FIG. 1. Clinical photographs of patients with an *SMARCB1* mutation. **SMARCB1-1:** Craniofacial features at age 2 months [a], 1 year [b], 6 years [c], and 18 years [d]. Note a round face with thick and arched eyebrows, a short nose with a bulbous tip and anteverted nostrils, a long philtrum, a small mouth, and micro-retrognathia in the early childhood. Later, note a broad nasal bridge without anteverted nostrils, a broad philtrum, a large tongue, and a protruding jaw. Poor posture due to severe scoliosis [e] as well as hypoplastic fingers [f, g] and toes [h, i] with nail hypoplasia, prominent interphalangeal joints, and prominent distal phalanges are noted at age 21 years. **SMARCB1-3:** Craniofacial features in the neonatal period [j, k], at age 2 years [l, m], and 7 years [n]. Note a round face with thick and arched eyebrows, a short nose with anteverted nostrils, a long philtrum, a small mouth, and micro-retrognathia in early childhood. Later, note a broad nasal bridge and a protruding jaw. Feet at age 5 years [o, p]. Note hypoplasia of the bilateral fifth toes and hypoplasia of all toenails. [Figure 1 originally published in Tsurusaki et al. [2012], in *Nature Genetics*.]

156 cm (-2.6 SD), and an OFC of 51.2 cm (-3.8 SD). He suffers from constipation, nocturnal enuresis, and unstable body temperature. He understands simple commands and speaks several words. He is friendly but also hyperactive and impulsive.

SMARCA4-3 (Subject 5 [Tsurusaki et al., 2012]; Fig. 2h–o): Increased nuchal translucency thickness was shown by fetal ultra-

sonography. He was born at 41 weeks of gestation. His birth weight was 2,756 g (-1.2 SD), length was 48 cm (-0.9 SD), and OFC was 32.0 cm (-0.9 SD). He was gavage-fed due to sucking/feeding difficulties until 7 months of age. He had right cryptorchidism which was corrected surgically at age 1 year. He showed hypotonia and motor development was delayed: he raised his head at age