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Research

BRF1 mutations alter RNA polymerase III–dependent transcription and cause neurodevelopmental anomalies

Guntram Borck,^{1,20} Friederike Hög,^{2,20} Maria Lisa Dentici,^{3,20} Perciliz L. Tan,^{4,20} Nadine Sowada,¹ Ana Medeira,⁵ Lucie Gueneau,⁶ Holger Thiele,⁷ Maria Kousi,⁴ Francesca Lepri,³ Larissa Wenzek,² Ian Blumenthal,⁸ Antonio Radicioni,⁹ Tito Livio Schwarzenberg,¹⁰ Barbara Mandriani,^{11,12} Rita Fischetto,¹³ Deborah J. Morris-Rosendahl,¹⁴ Janine Altmüller,^{7,15} Alexandre Reymond,⁶ Peter Nürnberg,^{7,16,17} Giuseppe Merla,¹¹ Bruno Dallapiccola,^{3,21} Nicholas Katsanis,^{4,21} Patrick Cramer,^{18,21} and Christian Kubisch^{1,19,21}

¹Institute of Human Genetics, University of Ulm, 89081 Ulm, Germany; ²Gene Center Munich and Department of Biochemistry, Center for Integrated Protein Science CIPSM, Ludwig-Maximilians-Universität München, 81377 Munich, Germany; ³Bambino Gesù Children's Hospital, IRCCS, 00165 Rome, Italy; ⁴Center for Human Disease Modeling, Duke University, Durham, North Carolina 27710, USA; ⁵Serviço de Genética, Departamento de Pediatria, Hospital S. Maria, CHLN, 1649-035 Lisboa, Portugal; ⁶Center for Integrative Genomics, University of Lausanne, 1015 Lausanne, Switzerland; ⁷Cologne Center for Genomics (CCG), University of Cologne, 50931 Cologne, Germany; ⁸Molecular Neurogenetics Unit and Psychiatric and Neurodevelopmental Genetics Unit, Center for Human Genetic Research, Massachusetts General Hospital, Boston, Massachusetts 02114, USA; ⁹Department of Experimental Medicine, Sapienza University, 00161 Rome, Italy; ¹⁰Department of Neonatology, Sapienza University, 00161 Rome, Italy; ¹¹IRCCS Casa Sollievo Della Sofferenza, Medical Genetics Unit, 71013 San Giovanni Rotondo, Italy; ¹²PhD Program, Molecular Genetics applied to Medical Sciences, University of Brescia, 25121 Brescia, Italy; ¹³U.O. Malattie Metaboliche PO Giovanni XXIII, AOU Policlinico Consorziale, 70120 Bari, Italy; ¹⁴Genomic Medicine, National Heart and Lung Institute, Imperial College, London SW3 6LY, United Kingdom; ¹⁵Institute for Human Genetics, University of Cologne, 50931 Cologne, Germany; ¹⁶Cologne Excellence Cluster on Cellular Stress Responses in Aging-Associated Diseases (CECAD), University of Cologne, 50674 Cologne, Germany; ¹⁷Center for Molecular Medicine Cologne (CMMC), University of Cologne, 50931 Cologne, Germany; ¹⁸Max Planck Institute for Biophysical Chemistry, Department of Molecular Biology, 37077 Göttingen, Germany; ¹⁹Institute of Human Genetics, University Medical Center Hamburg-Eppendorf, 20246 Hamburg, Germany

RNA polymerase III (Pol III) synthesizes tRNAs and other small noncoding RNAs to regulate protein synthesis. Dysregulation of Pol III transcription has been linked to cancer, and germline mutations in genes encoding Pol III subunits or tRNA processing factors cause neurogenetic disorders in humans, such as hypomyelinating leukodystrophies and pontocerebellar hypoplasia. Here we describe an autosomal recessive disorder characterized by cerebellar hypoplasia and intellectual disability, as well as facial dysmorphic features, short stature, microcephaly, and dental anomalies. Whole-exome sequencing revealed biallelic missense alterations of *BRF1* in three families. In support of the pathogenic potential of the discovered alleles, suppression or CRISPR-mediated deletion of *brf1* in zebrafish embryos recapitulated key neurodevelopmental phenotypes; in vivo complementation showed all four candidate mutations to be pathogenic in an apparent isoform-specific context. *BRF1* associates with BDPI and TBP to form the transcription factor IIIB (TFIIIB), which recruits Pol III to target genes. We show that disease-causing mutations reduce *Brf1* occupancy at tRNA target genes in *Saccharomyces cerevisiae* and impair cell growth. Moreover, *BRF1* mutations reduce Pol III–related transcription activity in vitro. Taken together, our data show that *BRF1* mutations that reduce protein activity cause neurodevelopmental anomalies, suggesting that *BRF1*-mediated Pol III transcription is required for normal cerebellar and cognitive development.

[Supplemental material is available for this article.]

Three RNA polymerases synthesize the different classes of RNAs in eukaryotic cells. Pol I synthesizes most ribosomal RNAs; Pol II mRNAs and miRNAs; and Pol III in a variety of noncoding RNAs

with a structural and catalytic function, e.g., tRNAs, 7SK RNA, 5S rRNA, and U6 snRNA (Schramm and Hernandez 2002; White 2011; Dieci et al. 2013). Pol II–dependent transcription has attracted most

²⁰These authors contributed equally to this work.

²¹These authors jointly supervised the work.

Corresponding author: guntram.borck@uni-ulm.de

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attention because its mRNA products are protein coding, but recent years have seen an increasing interest in Pol III transcription, not least because of its implication in human disorders. Increased Pol III-dependent transcription has been linked to cell transformation and cancer (Marshall and White 2008; Cabarcas and Schramm 2011), and germline mutations in components of the Pol III or tRNA processing machinery have been associated with neurogenetic disorders, such as pontocerebellar hypoplasia (PCH), a heterogeneous group of severe, often lethal, progressive neurodegenerative conditions characterized by cerebellar hypoplasia, progressive microcephaly, seizures, and profound developmental impairment. At least 10 types of PCH are currently recognized, and mutations causing the most prevalent subtypes, PCH2 and PCH4, are predicted to lead to loss of function of tRNA splicing endonuclease (TSEN); other PCH types display anomalies of tRNA processing as well (Namavar et al. 2011; Akizu et al. 2013; Rudnik-Schöneborn et al. 2014; Schaffer et al. 2014). However, the pathomechanism leading to PCH is poorly understood in many subtypes.

More recently, mutations in the catalytic Pol III subunits POLR3A and POLR3B have been identified in syndromic hypomyelinating leukodystrophies, such as hypomyelination, hypodontia, and hypogonadotropic hypogonadism (4H syndrome) (Bernard et al. 2011; Saitu et al. 2011; Tetreault et al. 2011). *POLR3A* mutations lead to a decrease in POLR3A levels and are predicted to interfere with the interaction with other Pol III subunits, which in turn would perturb Pol III-mediated transcription; a similar mechanism has been postulated for POLR3B-associated leukodystrophies. Although the molecular pathophysiology of these diseases is not fully understood, these findings suggest that an intact Pol III apparatus is essential for cognitive development and the structural and functional integrity of the cerebellum and white matter.

The basal Pol III transcription machinery requires the orchestrated interaction of Pol III, a multiprotein enzyme consisting of 17 polypeptide subunits, TFIIB (transcription factor IIB), and TFIIC (White 2011; Vannini and Cramer 2012). These factors, as well as the underlying mechanisms of transcription initiation, are conserved in eukaryotes. TFIIC recognizes and binds to specific sequence blocks in internal promoters (type 1 and 2 promoters) of target genes, such as tRNA genes. TFIIC then recruits TFIIB, which is usually composed of the B-related factor BRF1, B double prime 1 BDP1, and the TATA box-binding protein TBP, each of which is required for TFIIB function *in vitro* (White 2011). TFIIB recruits Pol III for transcription initiation. For the transcription of human RNA genes with external (type 3) promoters, such as U6 snRNA, BRF1 is replaced by BRF2 in TFIIB, and TFIIC is not required (Schramm and Hernandez 2002). Notably, *TBP* is a candidate gene for cognitive disorders in humans (Rooms et al. 2006), and TBP polyglutamine expansions cause spinocerebellar ataxia type 17 (Nakamura et al. 2001).

Here, we describe an autosomal recessive human syndrome characterized by intellectual disability and cerebellar, dental, and skeletal anomalies. We used whole-exome sequencing (WES) (Bamshad et al. 2011) and *in vivo* and *in vitro* assays to provide evidence that partial loss-of-function *BRF1* mutations cause this recognizable disease by impairing Pol III-mediated transcription.

Results

Clinical delineation of a cerebellar-facial-dental syndrome

By comparing the clinical features of three pairs of siblings (three female and three male) (Fig. 1A–C) from three unrelated families,

we observed a striking clinical overlap pointing to a previously undescribed disorder characterized by cerebellar hypoplasia, intellectual disability, characteristic facial dysmorphisms, and dental anomalies (Table 1; for the complete version of Table 1, see Supplemental Material). In each of the three families, two affected children were born to unaffected parents (Fig. 1C), suggesting autosomal recessive inheritance. In family 3 with first-cousin parents, Leber congenital amaurosis (LCA) segregated independently from cerebellar hypoplasia in two siblings (Fig. 1C). The six children affected by the cerebellar-facial-dental syndrome showed similar dysmorphic features that included sparse eyebrows, wave-shaped palpebral fissures, apparently low-set ears, malocclusion, and prominent upper incisors (Fig. 1A). They had proportionate short stature and microcephaly of prenatal onset, mild-to-moderate intellectual disability with speech delay, scoliosis, and sparse hair (Table 1). Three children had laryngeal stridor or laryngomalacia. Cranial magnetic resonance imaging revealed a pattern that was reminiscent of TSEN-related PCH, with a very thin corpus callosum, a flattened brainstem, and cerebellar vermis hypoplasia; however, the pons was relatively well formed (Fig. 1B). On skull and orthopantomaxillary x-rays, we saw bialveolar protrusion with prominent alveolar processes as well as taurodontism, a specific malformation of the pulp of molar teeth. Array-CGH did not detect causative copy number variants. We refer to this syndrome as cerebellar-facial-dental syndrome.

Identification of *BRF1* mutations

We performed WES in one or two affected individuals from each of the three families. Under the simplest assumption of genetic homogeneity for this rare autosomal recessive phenotypic constellation, we filtered for variants in the same gene with a predicted impact on protein function or RNA splicing that are rare in control populations of European descent. Because of parental consanguinity in family 3 and the lack of consanguinity in families 1 and 2, we searched for a gene harboring a rare homozygous variant in the affected individuals from family 3 and rare compound heterozygous (or homozygous) variants in the affected children in families 1 and 2. Subsequent to WES analysis and filtering, there was a single gene that contained biallelic mutations in all affected children: *BRF1*. Each affected child had two variants that predicted missense alterations, for a total of four distinct mutations. We confirmed all four *BRF1* alleles, as well as cosegregation with the cerebellar-facial-dental syndrome, in each family by Sanger sequencing (Fig. 1C; Supplemental Fig. 1). In family 1, the affected siblings were compound heterozygous for a maternally inherited c.677C > T mutation and a paternally inherited c.776C > T mutation, predicting p.Ser226Leu (S226L) and p.Thr259Met (T259M), respectively. In family 2, the affected sisters had inherited a c.667C > T mutation from their mother and carried a c.875C > A mutation that was likely transmitted from their father, from whom no DNA was available (Fig. 1C). These mutations predict p.Arg223Trp (R223W) and p.Pro292His (P292H) missense alterations, respectively. The two affected boys in family 3 were homozygous for the c.677C > T (p.Ser226Leu) mutation that had been identified in family 1. This mutation cosegregated with the cerebellar-facial-dental syndrome in family 3, while LCA was caused by a homozygous mutation in *RDH12* (c.784dupG; p.Ala262Glyfs*11), a gene previously shown to be mutated in LCA (Janecke et al. 2004). The four *BRF1* mutations were absent from ethnically matched controls as shown by sequencing and were not present or very rare in public SNP databases (Table 2).

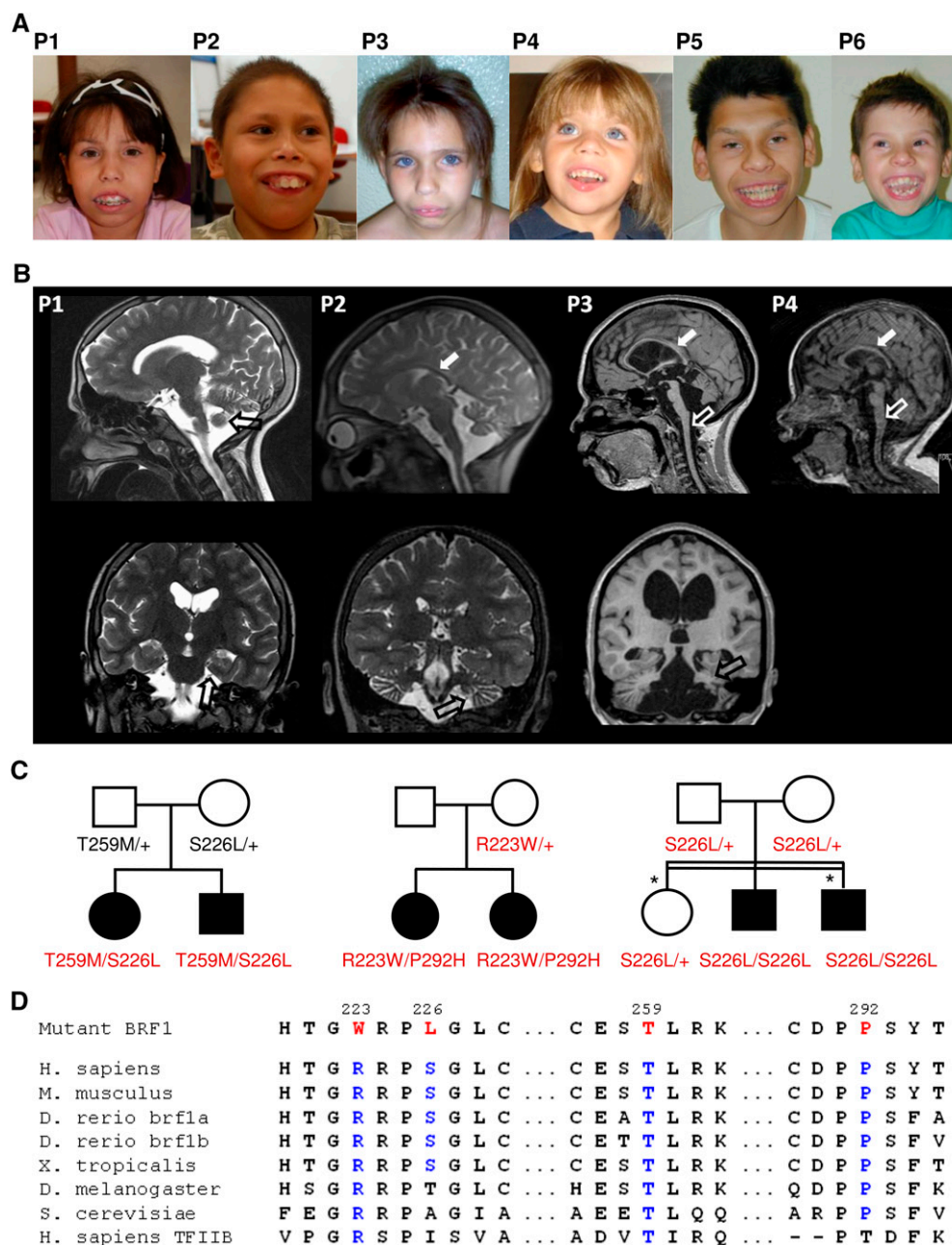


Figure 1. *BRF1* mutations cause a cerebellar-dental-skeletal syndrome. (A) Patients 1 and 2 (family 1) at the ages of 12 and 10 yr, patients 3 and 4 (family 2) at the ages of 10 and 4 yr, and patients 5 and 6 (family 3). Note characteristic facial dysmorphism and dental anomalies. (B) Brain MRI (*top*, sagittal scans; *bottom*, coronal scans) of patients 1–4 (P1–P4) at the ages of 14 yr, 9 yr, 12 yr, and 18 mo, respectively, showing a thin corpus callosum (white filled arrows), flattened brainstem (white unfilled arrows), and cerebellar hypoplasia (black unfilled arrows). (C) Pedigrees of family 1 (*left*), family 2 (*middle*), and family 3 (*right*) with genotypes for *BRF1* missense alterations. In family 2, the p.Pro292His mutation was likely transmitted by the unaffected father who did not participate in the study. In family 3, the two individuals denoted by an asterisk had Leber congenital amaurosis caused by a homozygous *RDH12* mutation. (D) Multiple sequence alignment of *BRF1* orthologs and human TFIIIB showing evolutionary conservation of mutant *BRF1* amino acid residues.

The longest *BRF1* isoform is a 677-amino-acid protein that, together with TBP and BDP1, forms the transcription factor IIIB (TFIIIB), which recruits Pol III to its templates and is involved in promoter opening (White 2011). The N-terminal region of *BRF1* has a role in DNA binding and Pol III recruitment (Kassavetis et al. 1998), is homologous to the Pol II transcription factor TFIIIB, and contains an N-terminal zinc ribbon domain and two cyclin domains (Khoo et al. 2014). The *BRF1* gene on human chromosome 14q32.33 en-

codes several functional isoforms, some of which lack the zinc ribbon domain (McCulloch et al. 2000). The four identified missense alterations invariably affect amino acids that are evolutionarily conserved, including in orthologs from mouse and zebrafish (Fig. 1D), but are not conserved in *BRF2*. In *Saccharomyces cerevisiae*, *Brf1* (Colbert and Hahn 1992), Arg223, Thr259, and Pro292 are conserved, whereas Ser226 is replaced conservatively by an alanine; for clarity, we will use the human amino acid numbering henceforth.

Table 1. Clinical features of individuals with cerebellar-dental-skeletal syndrome

Sex	Family 1 (Italy)		Family 2 (Portugal)		Family 3 (Italy)	
	Patient 1 F	Patient 2 M	Patient 3 F	Patient 4 F	Patient 5 M	Patient 6 M
Neonatal period	2420 g (39)	2640 g (39)	2530 g (40)	2820 g (40)	3050 g (40)	2900 g (40)
Weight at birth (weeks of gestation)	31 cm (-3 SD) 12 yr	33 cm (-2 SD) 9 yr 3 mo	31 cm (-3 SD) 12 yr	32 cm (-2 SD) 18 mo	n.a. 21 yr	n.a. 15 yr
OFC at birth	122 cm (<3rd)	109 cm (<3rd)	124 cm (<3rd)	71 cm (<3rd)	157 cm (<3rd)	151 cm (<3rd)
Last growth parameters (age)	51 cm (<3rd)	48 cm (<3rd)	48 cm (<3rd)	42 cm (<3rd)	47 cm (<3rd)	48 cm (<3rd)
Height (centile)	Mild Dyslalia	Mild Short sentences	Mild-moderate Short sentences	Mild Speech delay	Severe Few words	Severe Few words
Cognitive development	+	+	+	+	+	+
Intellectual disability	+	+	+	+	+	+
Speech	+	+	+	+	+	+
Craniofacial anomalies	+	+	+	+	+	+
Characteristic facial dysmorphism	+	+	+	+	+	+
Prominent upper incisors and malocclusion	+	+	+	+	+	+
Skeletal and limb anomalies	+	+	+	+	+	+
Scoliosis	+	+	+	+	+	+
Delayed bone age	+	+	+	+	+	+
Other clinical features	+	+	+	+	+	+
Laryngeal anomalies	-	Laryngomalacia	Laryngeal stridor	Laryngeal stridor	-	-
Ocular anomalies	Strabismus	-	Bilateral cataract	-	-	LCA due to <i>RDH12</i> mutation
Cardiac anomalies	Mitral valve prolapse; flat T-waves (ECG)	Mild dilatation of the ascending aorta; negative T-waves (ECG)	-	Ventricular septal defect	Systolic murmur	-
Thin hair and sparse eyebrows	+	+	+	+	+	+
Brain anomalies on MRI	+	+	+	+	+	+
Cerebellar hypoplasia	Enlarged lateral ventricles	Enlarged cisterna magna	Thin corpus callosum; enlarged lateral ventricles; enlarged cisterna magna	Thin corpus callosum	Enlarged ventricles; Thin corpus callosum; Hypoplastic pons and mesencephalon; enlarged cisterna magna	Enlarged ventricles; thin corpus callosum; hypoplastic pons and mesencephalon; enlarged cisterna magna
Other						
Skull and orthopantom x-rays						
Bialveolar protrusion	+	+	+	+	+	-
Prominent alveolar processes	+	+	+	+	+	-
Taurodontism	+	+	+	+	+	+
<i>BRF1</i> mutations						
<i>BRF1</i> mutation maternal allele		c.677C > T p.(Ser226Leu)	c.667C > T p.(Arg223Trp)			c.677C > T p.(Ser226Leu)
<i>BRF1</i> mutation paternal allele		c.776C > T p.(Thr259Met)	c.875C > A p.(Pro292His)			c.677C > T p.(Ser226Leu)

(LCA) Leber congenital amaurosis; (ECG) electrocardiogram; (n.a.) not available.

Table 2. *BRF1* mutations

<i>BRF1</i> mutation (DNA)	<i>BRF1</i> alteration (protein)	Affected protein domain	Frequency in control chromosomes (sequencing)	Frequency in control chromosomes (ESP-EVS)	PolyPhen-2 prediction	SIFT prediction
c.677C > T	p.(Ser226Leu)	Cyclin 2	0/200	0/12,994	Probably damaging	Not tolerated
c.776C > T	p.(Thr259Met)	Cyclin 2	0/200	1/13,006	Probably damaging	Not tolerated
rs373957300						
c.667C > T	p.(Arg223Trp)	Cyclin 2	0/100	2/12,996 (heterozygous)	Probably damaging	Not tolerated
rs370270828						
c.875C > A	p.(Pro292His)	–	0/100	0/13,006	Probably damaging	Not tolerated

(ESP-EVS) National Heart, Lung and Blood Institute (NHLBI) Exome Sequencing Project Exome Variant Server.

Functional analysis of *brf1* in zebrafish

As a first test of the functional candidacy of *BRF1*, we sought an in vivo model with a credible phenotypic surrogate. We noted that all probands in our study have cerebellar hypoplasia and microcephaly, both of which have been modeled previously in zebrafish embryos for a variety of neurodevelopmental disorders (Golzio et al. 2012; Dauber et al. 2013; Margolin et al. 2013; Bernier et al. 2014). We therefore turned to developing zebrafish embryos and asked whether either suppression or overexpression of *brf1* has an effect on neurogenesis, head size, and/or cerebellar morphology.

Given that *brf1* has two copies in the zebrafish genome due to the teleost-specific genome duplication, we generated splice blocking morpholinos (sbMOs) for both copies (*brf1a* and *brf1b*), and we scored for neurogenesis by measuring the area of the optic tectum, the area between the eyes as a surrogate for brain size, and the cerebellar integrity. We note that while *brf1a* is known to be expressed at multiple stages of development as well as in multiple organs, little is known about *brf1b* in zebrafish. Our in-house RNA-seq data that interrogate the transcripts expressed specifically in the head of 5 d post fertilization (d.p.f) zebrafish embryos indicated that *brf1b* has significantly increased expression when compared to *brf1a*. Indeed, *brf1b* was expressed, on average, 11.59 CPM (counts per million mapped reads), while *brf1a* expression was 4.32 CPM. In triplicate experiments scored by two investigators masked to the injection cocktails, suppression of *brf1a* gave no appreciable phenotypes (data not shown). In contrast, suppression of *brf1b* with a MO against the donor site of exon 8 (MO#1) (Supplemental Fig. 2) resulted in a significant reduction of both the size of the optic tectum (Fig. 2B',D,F',H) and the total area between the eyes (Fig. 2B,C,E,G). Furthermore, we observed cerebellar hypoplasia as determined by the lack of axons in the midline of the morphant zebrafish cerebellum (Fig. 2B',F',I). We did not observe further overt morphological abnormalities or developmental delay in the injected embryos as judged by the shape and position of the heart, the absence of edema, the number and shape of somites, and the presence of a normal swim bladder. The observed phenotypes were specific; coinjection of MO and wild-type (WT) human mRNA was able to rescue the phenotypes (Fig. 2B–D,F–I). To assess the functionality of the alleles, we coinjected MO with human mRNA bearing one of the variants of interest (P292H, R223W, S226L, and T259M) at a time. In the context of *BRF1* isoform 2 (Fig. 2E), P292H and R223W scored as functionally nulls as they failed to rescue the reduction in head size and optic tecta as well as the cerebellar hypoplasia (Fig. 2F–I), while S226L and T259M scored as hypomorphic: The mutants were significantly more affected than MO + WT but not as severe as MO alone. Strikingly, testing of three of the alleles (R223W, S226L, T259M) in

isoform 1 showed them all to be benign (rescue was indistinguishable from WT) (Fig. 2A–D), suggesting that the effect of the mutations is isoform specific. Overexpression of WT human *BRF1* and human P292H, R223W, S226L, or T259M *BRF1* showed no difference in head size, optic tecta, or cerebellar hypoplasia (Fig. 2G',H',I').

To confirm the specificity of our assay, we designed a second sbMO against the donor site of *brf1b* exon 6 (MO#2). Injections of MO#2 recapitulated the results obtained with MO#1, showing a reduction of head size and optic tecta as well as cerebellar disorganization phenotypes in 3 d.p.f. embryos; again, the phenotypes could be at least partially rescued by coinjection of human WT *BRF1* mRNA (Supplemental Fig. 3). Further supplementing these findings, we utilized the CRISPR system and induced aberrations in *brf1b* by *CAS9/gRNA*. In the F0 *brf1b*^{Cas9/gRNA}, we observed phenotypes that are fully concordant with the *brf1b* morphant phenotypes produced by both sbMOs (Supplemental Fig. 3). We conclude that suppression of *brf1* leads to phenotypes that recapitulate key symptoms of the human pathology, arguing in favor of the candidacy of this gene for disease pathogenesis.

BRF1 mutations affect cell growth

We next turned to the question of the precise effect of the discovered missense alleles. Consistent with their predicted pathogenic nature, yeast lacking the chromosomal *BRF1* gene and expressing *BRF1* with mutations corresponding to R223W or P292H did not grow in spot dilution tests (Fig. 3A), consistent with loss of Brf1 function and similar to our zebrafish functional assay in which R223W and P292H were classified as nulls. Yeast cells transformed with plasmids containing *BRF1* substitutions T259M or S226L (as found in families 1 and 3) or A226S (reflecting the human reference sequence) grew normally. We observed similar results in liquid cultures (Supplemental Fig. 4). To mimic the compound heterozygosity present in the families, we transformed yeast with two distinct plasmids expressing the *BRF1* variants R223W/P292H or S226L/T259M. Whereas S226L/T259M yeast showed essentially normal growth, the R223W/P292H combination was lethal (Fig. 3A). With respect to the mutations identified in family 1, these results further supported the candidacy of human biallelic *BRF1* missense alterations causing the cerebellar-facial-dental syndrome.

BRF1 mutations impair Pol III–dependent transcription

To investigate whether the altered *BRF1* residues may be important for DNA binding, we mapped the mutated *BRF1* residues on the known crystal structure of the homologous human TFIIB-TBP-DNA complex (Tsai and Sigler 2000). Residues Arg223 and Thr259 are conserved in human TFIIB (Fig. 1D) and are predicted to contact

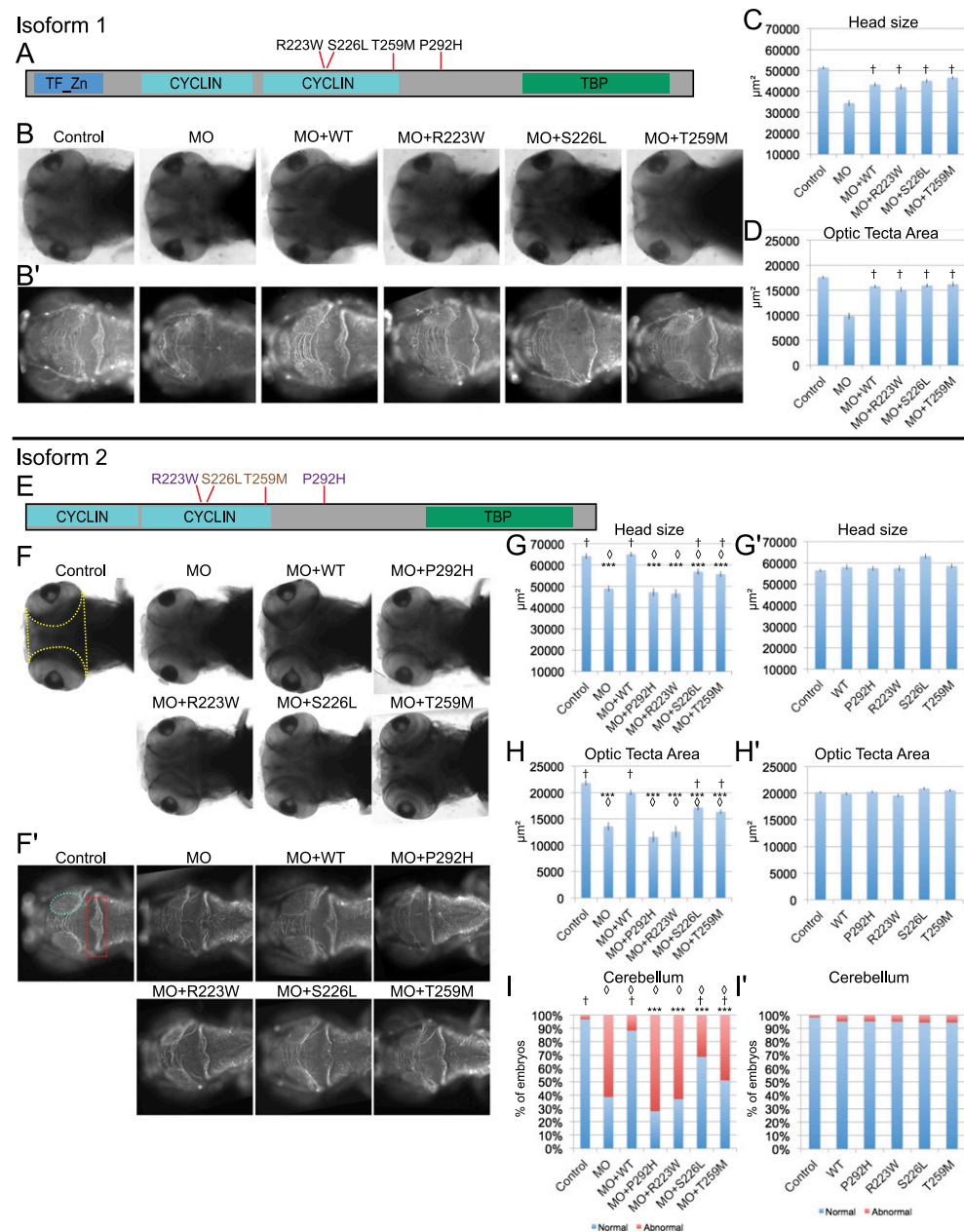


Figure 2. Functional annotation of variants in isoforms 1 and 2 of *BRF1* and its effects on the head size, optic tectum size, and cerebellar formation in zebrafish embryos. (A,E) Schematic representation of the location of *BRF1* variants examined in zebrafish within each of the two isoforms evaluated: isoform 1 (NP_001229717.1; 650 aa) and isoform 2 (NP_001229715.1; 584 aa). In purple are alleles that when tested are shown to be null, and in brown are alleles that score as hypomorphs based on zebrafish assays. (B,B') Dorsal views of control zebrafish embryos and embryos injected with *brf1b* MO, *brf1b* MO + WT human *BRF1*, and *brf1b* MO + variant (R223W, S226L, or T259M) human *BRF1* RNA in the context of isoform 1, respectively, at 3 d.p.f., stained with anti- α acetylated tubulin. (B) Head size measurements were taken using brightfield images (highlighted with a yellow outline in panel F). (B') The area of the optic tecta was measured in the fluorescent images (highlighted with a cyan oval in panel F'). (C,D) Bar graphs showing the relative head size (C) and the optic tecta area (D). Data are presented as mean \pm SE. Two-tailed *t*-tests were performed to assess statistical significance. The embryos injected with the MO and each of the variants were statistically different from MO alone but not statistically different when compared to embryos injected with MO + WT human *BRF1*, therefore scoring as benign. (F-I') Functional assessment of the *BRF1* missense variants in the context of isoform 2. Three d.p.f. embryos injected with *brf1b* MO, *brf1b* MO + WT human *BRF1*, *brf1b* MO + variant human *BRF1* RNA (P292H, R223W, S226L, or T259M), or variant human *BRF1* RNA alone in the context of isoform 2 were observed following staining with anti- α acetylated tubulin for head size (F) and for optic tecta area (F'), as well as for cerebellar defects (F'; illustrated with a red dashed box where maximum disorganization is observed). (G-I') Bar graphs showing average head size, optic tecta area, as well as the percentage of embryos with cerebellar defects evaluated among each condition. To assess statistical significance among the evaluated conditions, two-tailed *t*-tests were performed for head size and optic tecta, and χ^2 tests were performed for cerebellar disorganization to evaluate statistical significance across the conditions. In the context of isoform 2, functional analysis using zebrafish show that P292H and R223W are null, while S226L and T259M are hypomorphic alleles, showing an isoform-specific effect in which head size and optic tecta size are reduced and cerebellar disorganization occurs. No significant effects were observed for any of the phenotypes when the variants themselves were overexpressed. Each experiment was done at minimum in triplicate with at least 50 embryos per condition per replicate. (***) *P*-value \leq 0.001 relative to MO + WT; (<) *P*-value \leq 0.001 relative to controls; (†) *P*-value \leq 0.01 relative to MO.

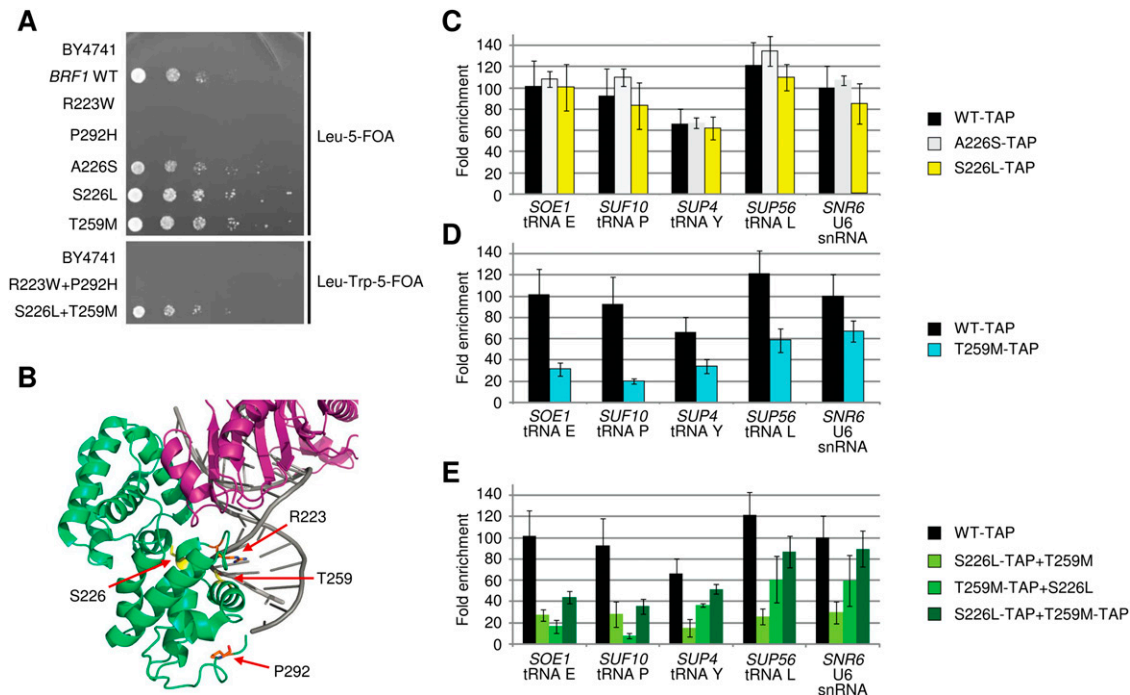


Figure 3. *BRF1* mutations cause growth defects and reduced target promoter occupancy. (A) *BRF1* mutations affect cell growth. Spot dilutions of the variants introduced into a *BRF1* knockout strain grown at 30°C. Wild-type (WT) and variant Brf1 were encoded on plasmids. For the combination of two mutations, two plasmids were used, each harboring one mutation and a distinct marker. (5-FOA)5-fluoroorotic acid. (B) Three-dimensional modeling of human *BRF1* missense alterations. The four identified amino acid substitutions were mapped to the structure of human TFIIB (green) in a complex with TBP (purple) and DNA (pdb code 1C9B) (Tsai and Sigler 2000). Amino acids affected by mutations in families 1 and 2 are shown in yellow and orange, respectively. (C–E) *BRF1* variants show a decreased occupancy of tRNA promoters in yeast. Fold enrichments of ChIP experiments performed with tandem affinity purification (TAP)—tagged *BRF1* variants in yeast. Data are presented as mean \pm SD. (C) Mutations A226S to serine or S226L show no effect on fold enrichment compared to WT. (D) Mutation T259M leads to decreased fold enrichment on tRNA genes. (E) Combination of the two variants results in a much lower occupancy of the S226L and T259M variants. Summing up both signals results in less occupancy of the two mutated *BRF1* than the WT *BRF1*.

DNA (Fig. 3B). Residues Ser226 and Pro292 are not conserved in human TFIIB and are predicted to be in the proximity of the DNA, but not in direct contact (Fig. 3B). These modeling results suggested that the *BRF1* mutations might affect *BRF1* binding to DNA, impair TFIIB-mediated Pol III recruitment, and reduce Pol III-mediated transcription.

To test whether the *BRF1* mutations influence Pol III recruitment to gene promoters *in vivo*, we performed chromatin immunoprecipitation (ChIP) with tandem affinity purification (TAP)-tagged versions of *BRF1* variants on Pol III target genes in yeast (Fig. 3C–E). Consistent with our predictions, the *BRF1* variant T259M showed strongly decreased promoter occupancy at four tested tRNA loci and a significant decrease in occupancy of the U6 snRNA promoter (Fig. 3D). The ChIP signals for TAP-tagged A226S and S226L variants were not different from wild-type Brf1 (Fig. 3C). Although the R223W and P292H variants could not be tested due to their lethal effect, these results indicate that at least one *BRF1* alteration can affect occupancy of different Pol III promoters *in vivo*.

Finally, to test whether the *BRF1* mutations cause a transcriptional defect, we used an established *in vitro* assay (Hahn and Roberts 2000). This assay is based on a nuclear extract from a temperature-sensitive yeast strain carrying the *BRF1* substitution p.Trp107Arg (W107R). Extracts from this strain are transcriptionally inactive, but addition of recombinant Brf1 protein restores activity on a template encoding the native *SUP4* promoter. When we used purified recombinant Brf1 variants in this assay, we observed significantly less transcription for the *BRF1* variant R223W

in comparison with the same concentration of WT Brf1 (Fig. 4). Milder, but both significant and reproducible defects were observed for T259M, P292H, A226S, and S226L. *BRF1* variants combining two mutations also showed transcriptional defects in this assay (Fig. 4). Taken together, these results show that *BRF1* mutations identified in patients can impair transcription in a well-controlled, *in vitro* transcription assay.

Discussion

Whereas intellectual disability can be due to mutations in synaptic genes (Pawlowsky et al. 2012; Zoghbi and Bear 2012), recent studies have highlighted impairments of basal cellular functions and pathways such as transcription and translation in the etiology of cognitive disorders and neurogenetic syndromes (Najmabadi et al. 2011). Some examples include pontocerebellar hypoplasias, Pol III-related leukodystrophies, X-linked intellectual disability, and leukoencephalopathy with vanishing white matter (Bugiani et al. 2010; Namavar et al. 2011; Borck et al. 2012; Daoud et al. 2013). Complete inactivation of processes such as translation initiation or tRNA transcription is not compatible with life, suggesting that most disease-causing mutations are hypomorphs.

Our identification and characterization of *BRF1* mutations causing the syndromic cerebellar-facial-dental phenotype is in line with these observations. No mutant *Brf1* mouse has been reported to our knowledge; in yeast, deletion of *BRF1* is lethal (Colbert and Hahn 1992). Accordingly, one of the missense variants we have

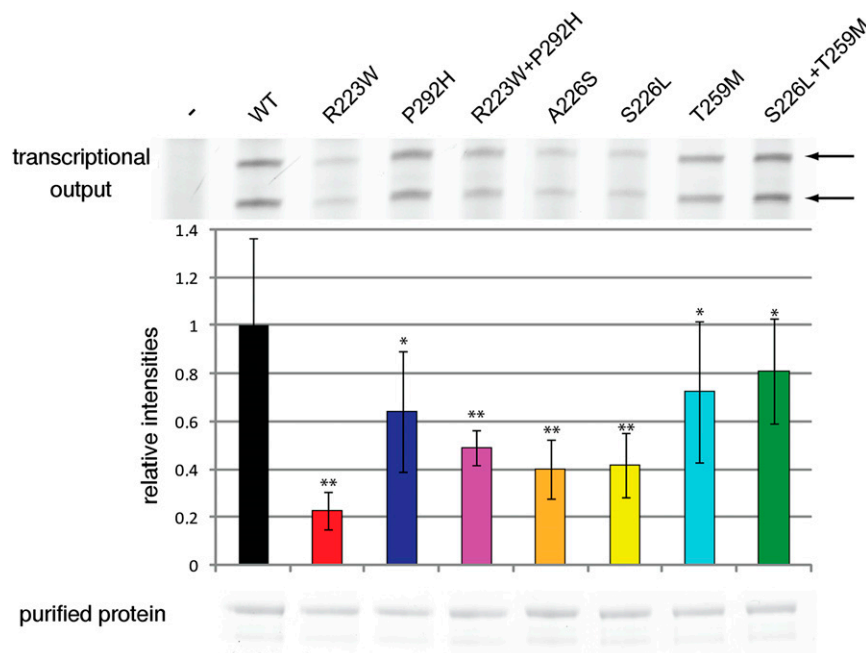


Figure 4. In vitro transcription defects caused by *BRF1* mutations in yeast. (Top) A representative gel of the in vitro transcription reaction using a nuclear extract harboring a deficient Brf1 protein (p.Trp107Arg, W107R) that is impaired in Pol III–dependent transcription. Addition of WT Brf1 rescues activity whereas the different Brf1 variants and the combinations present in the affected children show a defect in transcription. Data are presented as mean \pm SD. A quantification is shown in the middle panel. (*) $P < 0.01$; (**) $P < 0.0001$. The lower panel shows a twelve times excess of the Brf1 protein amount used for this assay separated on an SDS gel.

identified is recruited to a lesser extent than wild type to target genes. Several other mutants have reduced transcriptional activity, suggesting that in the context of this assay they behave as hypomorphs. *BRF1* mutations likely confer a specific phenotype; we did not identify *BRF1* mutations in three individuals with clinically overlapping but distinct malformation syndromes or in seven individuals with genetically unresolved forms of PCH (data not shown). However, there is compelling evidence for the pathogenicity of the identified variants in the cerebellar-facial-dental syndrome: *BRF1* was the only gene exome-wide that harbored very rare or unique biallelic mutations in the three affected sibships. These missense variants affect conserved amino acids that lie within a predicted functional domain, and two of the affected amino acids likely interact directly with DNA. In vivo experiments in zebrafish embryos showed that suppression of *brf1b* phenocopied some of the key pathognomonic features of the syndrome. Of note, while we cannot exclude a possible effect of overexpression on the outcome of our zebrafish assays, the effects of the variants seemed to be restricted to isoform 2 of *BRF1*. Thus our results possibly illustrate an isoform-specific effect underlying the phenotype and offer a plausible hypothesis about how mutations in a ubiquitous, key component of the cellular machinery can induce specific, albeit severe, phenotypes. Similar splice isoform-specific pathogenic effects have been found in other disorders (Sarparanta et al. 2012; Schulte et al. 2014), suggesting that this mechanism might be a significant regulator of pleiotropy. Previous studies have shown that full-length human *BRF1* is the canonical isoform in pol III transcription, and whether other isoforms, including isoforms lacking the evolutionary conserved Zn ribbon domain such as isoform 2, are biologically active remains unknown. Biochemical data indicate that overexpression of iso-

forms lacking the Zn ribbon domain can induce *BRF1*–TBP interactions in vitro and reconstitute transcription in crude systems (McCulloch et al. 2000; Schramm and Hernandez 2002). While the in vivo role of different *BRF1* isoforms remains to be elucidated, our zebrafish data provide further evidence of the pathogenic nature of the four missense variants.

Further studies showed each of the missense variants to affect a variety of relevant activities in yeast, including target gene promoter occupancy, in vitro transcription, and cell growth. We do not know the precise mechanism by which *BRF1* contributes to the phenotype in humans. We note, however, that mouse *Brf1* is expressed in anatomical structures that are relevant to the observed human pathology: In situ hybridization of post-natal day 56 mouse brain had previously revealed prominent *Brf1* expression in the olfactory bulb, hippocampus, and the Purkinje and granule cell layers of the cerebellum (Allen Brain Atlas) (Sunkin et al. 2013), underscoring the important role of *BRF1* in brain development.

We speculate that one plausible pathomechanism underlying the cerebellar-facial-dental syndrome is a reduced steady-state level of tRNAs that would be predicted to impair translation elongation; this may explain the growth defect in yeast and contribute to short stature in the affected individuals. Whether the major drivers of the phenotype are tRNAs or other Pol III–dependent transcripts remains to be determined. Finally, it is also unknown why partial disruption of Pol III transcription leads to organ- and tissue-specific phenotypes. There are ~ 450 nuclear encoded tRNA genes in the human genome for only 61 anticodons. It has been shown previously that up to 26% of tRNA genes are active in one cell type versus another (Barski et al. 2010) and that tRNA expression varies by as much as tenfold among human tissues (Dittmar et al. 2006), suggesting the existence of active cell and tissue-specific regulation of Pol III transcription (Oler et al. 2010). Although the genome-wide architecture of Pol III–mediated transcription in neurons or specific brain regions is unknown, a pilot study using a microarray containing 42 probes for nuclear encoded tRNAs and assaying their relative expression among eight human tissues revealed a high overall tRNA expression level in brain (Dittmar et al. 2006). This observation may partially explain the vulnerability of brain regions toward impaired tRNA transcription or processing. Whether other affected tissues, such as the teeth, also show specific tRNA expression patterns remains to be investigated. We note that dental anomalies such as delayed dentition, hypodontia, oligodontia, and abnormally placed or shaped teeth are part of the Pol III–related leukodystrophy syndromes 4H; ataxia, delayed dentition, and hypomyelination (ADDH); and leukodystrophy with oligodontia (LO) (Daoud et al. 2013). An alternative explanation for the tissue-restricted phenotypes engendered by *BRF1* dysfunction might be a partial functional redundancy with *BRF2*. It has been shown, however, that *BRF1* and *BRF2* are almost completely mutually exclusive as components of TFIIB under physiological

conditions (Moqtaderi et al. 2010), and whether BRF2 can partially compensate for defective BRF1 is unknown.

In conclusion, our study defines a syndrome with cerebellar hypoplasia, intellectual disability, and growth retardation that is caused by partial deficiency of the conserved Pol III transcription factor BRF1. These results add an example to the short list of genetic diseases caused by dysregulation of the Pol III machinery, most of which predominantly affect the central nervous system and suggest that improved understanding of the potential target specificity provided by BRF1 might offer pathomechanistic clues.

Methods

Subjects

The study was performed with the approval of the University of Ulm Ethics Committee. Affected individuals were evaluated at the genetics outpatient clinics of the Bambino Gesù Children's Hospital, Rome (family 1); the Hospital S. Maria, Lisboa (family 2); and IRCCS Casa Sollievo Della Sofferenza Hospital, San Giovanni Rotondo and Giovanni XIII Hospital, Bari, Italy (family 3). Patients were enrolled with written parental consent for participation in the study. The clinical evaluation included medical history interviews, a physical examination, and review of medical records. Clinical information of affected individuals from the first two families had been independently submitted to the web-based Dysmorphology Diagnostic System (DDS) of the DYSCERNE network (a European network of centers of expertise for dysmorphology) (Douzgou et al. 2014) by the respective attending geneticists (M.L.D. and B.D.; A.M.). After review of the clinical information and pictures, the DDS Expert Panel (including B.D.) concluded that the four affected individuals were likely affected by a previously unreported autosomal recessive disorder. Blood samples were obtained from each participating individual, and genomic DNA was extracted by standard procedures.

Whole exome and BRF1 sequencing

Families 1 and 2

Genomic DNA was enriched for exonic and adjacent splice site sequences with the SeqCap EZ Human Exome Library v2.0 kit, and libraries were run on an Illumina HiSeq 2000 Sequencer via a paired-end 100-bp protocol (Hussain et al. 2013). For data analysis, the Cologne Center for Genomics Varbank pipeline v2.6 and user interface was used. Primary data were filtered according to signal purity by the Illumina Realtime Analysis (RTA) software v1.8. Subsequently, the reads were mapped to the human genome reference build GRCh37/hg19 (<http://www.genome.ucsc.edu/>) using the BWA-SW alignment algorithm (Li and Durbin 2010). GATK v1.6 (McKenna et al. 2010) was used to mark duplicated reads, perform a local realignment around short insertion and deletions (indels), recalibrate the base quality scores, and call SNPs and short indels. Mean coverage was $>100\times$ in both exomes and $\sim 90\%$ of target bases were covered more than $30\times$. In the target regions, we detected 25,894 SNVs and 2004 short indels in the family 1 individual and 35,642 SNVs and 2795 short indels in the family 2 individual. Scripts developed in-house at the Cologne Center for Genomics were applied to detect protein changes, affected donor and acceptor splice sites, and overlaps with known variants. Splice site variants were analyzed with a maximum entropy model (Yeo and Burge 2004). We filtered for high-quality rare variants (MAF $< 0.1\%$; based on the 1000 Genomes database, build 20110521 [The 1000 Genomes Project Consortium 2012] and the NHLBI Exome Sequencing Project Exome Variant Server [EVS],

build ESP5400 [Tennessen et al. 2012]) with a predicted impact on protein sequence or splicing. We also filtered against an in-house database containing variants from 511 exomes from individuals with epilepsy to exclude pipeline-related artifacts.

Family 3

WES and data analysis were performed as previously described (Alfaiz et al. 2014). Briefly, exomes were captured using the Agilent SureSelect human all exon V4 enrichment kit and sequenced on an Illumina HiSeq platform. Variants were filtered based on adherence to an autosomal recessive inheritance pattern, prediction by SIFT (<http://sift.jcvi.org/>) (Kumar et al. 2009) and PolyPhen-2 (<http://genetics.bwh.harvard.edu/pph2/>) (Adzhubei et al. 2010), and absence from dbSNP129 (<http://www.ncbi.nlm.nih.gov/projects/SNP/>).

PCR and Sanger sequencing were performed according to standard protocols for the validation of variants of interest and for cosegregation studies. BRF1 mutation nomenclature is based on transcript NM_001519.3 and RDH12 mutation nomenclature on transcript NM_152443.2.

In vivo modeling in zebrafish

Reciprocal BLAST of the human BRF1 protein sequence against the zebrafish genome identified two orthologs, suggesting that this gene is duplicated in the zebrafish genome: *brf1a* (NP_956192.1, 60% identity, 72% similarity) and *brf1b* (NP_956183.1, 60% identity, 72% similarity). We targeted each of these transcripts with a sbMO designed and obtained from Gene Tools. The MOs target the donor site of exon 6 for *brf1a* sbMO (TATTGTGAAGCTTACC GATAGCTGGT), the donor site of exon 8 for *brf1b* sbMO (AGCTGTGACATTCACCTTTTCTCA), and the donor site of exon 6 for *brf1b* sbMO#2 (GTCATTACAGGGAGACTTACCGACA). For exon experiments, human WT mRNAs of isoform 1 (NP_001229717.1) and isoform 2 (NP_001229715.1) of BRF1 were cloned into pCS2+ vector and transcribed in vitro using SP6 mMessage machine kit (Ambion). All variants were introduced with Phusion high-fidelity DNA polymerase (New England Biolabs) and custom designed mutagenesis primers and Sanger sequence confirmed on a 3730 ABI sequencer. We injected 1 nL of solution (12 ng MO and/or 200 pg of human mRNA, (WT, P292H, R223W, S226L, or T259M) into one- to four-cell-stage WT zebrafish embryos (Niederiter et al. 2013) and collected them at 3 d.p.f., fixed in Dent's fixative overnight at 4°C. Standard immunohistochemistry was then performed using control embryos and embryos injected with *brf1b* MO, MO + WT human BRF1, and MO + variant human BRF1 (P292H, R223W, S226L, or T259M) stained with anti- α acetylated tubulin (T7451, mouse, Sigma-Aldrich, dilution 1:1000) as the primary antibody and Alexa Fluor (goat anti-mouse IgG (A21207, Invitrogen, dilution 1:1000) as a secondary antibody. Stained zebrafish were observed using the Nikon AZ100 Microscope with a DS-Qi1MC digital camera head. All experiments were performed in triplicate (~ 200 embryos), and significance was determined with a Student's *t*-test.

CRISPR target sequences (*brf1b*: CATGCATGAATCCGGC GCA) were identified using crispr.mit.edu and cloned into pT7-gRNA (Addgene) following the protocol described at <http://www.addgene.org/crispr/Chen/>. gRNAs and mRNAs were synthesized by a MEGAscript T7 kit (Ambion), and Cas9 mRNAs were synthesized by mMESAGE mMACHINE SP6 (Ambion). One hundred two pictograms gRNA + 150 pg Cas9 mRNA were injected into one-cell stage zebrafish and collected at 3 d.p.f in Dent's fixative overnight at 4°C and stained as mentioned above.

For RNA sequencing (RNA-seq), total RNA from 5 d.p.f. zebrafish heads was extracted using a TRIzol/chloroform protocol. Libraries were constructed using a customized version of strand-specific

dUTP and run on an Illumina HiSeq 2000 platform (Sugathan et al. 2014).

Yeast strains and growth assays

To introduce *BRF1* mutations in yeast, the shuffle vector system developed by Sikorski and Hieter was used (Sikorski and Hieter 1989). *BRF1* mutations were introduced in the vector pRS315, which contains yeast *BRF1* expressed under the control of its own promoter. Mutations were introduced using site-directed mutagenesis and confirmed by DNA sequencing. The identified mutations from humans (h) were transferred to yeast (y). The corresponding amino acid positions are as follows: hR223 = yR218, hS226 = yA221, hT259 = yT254, and hP292 = yP288. For combinations of two mutations, *BRF1* variants T259M and S226L were cloned into pRS314 vector and cotransformed with the S226L and T259M mutations in the pRS315 vector, respectively. Plasmids were transformed into the yeast strain SHY285 (gift from S. Hahn, *MAT α* , *ade2 Δ ::hisG*, *his3 Δ 200*, *leu2 Δ 0*, *lys2 Δ 0*, *met15 Δ 0*, *trp1 Δ 63*, *ura3 Δ 0*, *brf1 Δ ::HIS3/pSH524*), and WT *BRF1* was replaced with the mutant gene by plasmid shuffle at 30°C. The rescue plasmid pSH524 contains a URA3 marker (Ars Cen URA3, *BRF1*) (Hahn and Roberts 2000). For the spotting assays, strains were grown overnight in selective media, normalized to equal densities (10^7 cells/200 μ L), serially diluted (1:5), and spotted on YPD media or selective media (Leu-, Leu/Trp-) supplemented with 5-fluoroorotic acid (5-FOA; Carl Roth). Plates were incubated for 3 d at 30°C. For the liquid assays, cultures were inoculated in selective media, grown overnight and normalized to equal densities (OD_{600nm} = 0.33). We used 3.5 μ L of these dilutions to inoculate a 96-well plate with 200 μ L of liquid media (YPD and selective media supplemented with 5-FOA). Plates were incubated for 3 d at 30°C. As a control strain we used WT BY4741 yeast.

Protein purification

Protein purification was performed essentially as previously described (Alexander et al. 2004), with minor modifications. Proteins were expressed in *Escherichia coli* BL21-Codon plus (DE3) RIL cells in LB medium. Protein expression was induced in 2 l cultures with 1 mM IPTG at an OD₆₀₀ of 0.7. Cells were grown for 3 h at 37°C and harvested (20 min, 4000 rpm, 4°C). The pellet was washed with 40 mL LB medium and flash-frozen in liquid nitrogen. The cells were thawed on ice and resuspended in 50 mL lysis buffer (25 mM Tris/HCl at pH 7.5 and 25°C; 200 mM KCl; 12.5 mM MgCl₂; 10% [v/v] glycerol; 1 \times PI). Subsequently, 0.8 mg lysozyme (Roth) per 1 mL lysate was added and incubated rotating in the cold room. After 10 min, 0.2% IGEPAL CA-630 (MP Biomedicals) was added for further 5 min. For cell disruption, lysate was sonicated for 15 min, at 25% duty cycle and 40 output value. Brf1 inclusion bodies was washed twice in H.35 buffer (20 mM HEPES at pH 8.0 and 25°C; 350 mM KCl; 2 mM MgCl₂; 1% NaOH; 20% [v/v] glycerol; 0.1% [v/v] IGEPAL CA-630; 1 \times PI). Inclusion bodies were extracted with 5 mL G6 buffer (100 mM Tris/HCl at pH 7.5 and 4°C; 6 M GdmCl; 2 mM β -mercaptoethanol) and rotation for 1 h at 4°C. The solution was bound in batch to 2.5 mL Ni-NTA bead volume in G6 buffer overnight at 4°C. Beads were washed with 12.5 mL G6 buffer and protein eluted in batch with three times 2.5 mL Ni-NTA elution buffer (100 mM Tris/HCl at pH 7.5 and 4°C; 6 M GdmCl; 10 mM β -mercaptoethanol; 500 mM imidazole; 1 \times PI). To the combined eluate, ZnSO₄ and DTT were added to final concentrations of 10 μ M and 5 mM, respectively. Finally, the denatured Brf1 was refolded via rapid dilution and dialysis. For dilution, 100 μ L of the eluate was placed in the lid of a 1.5-mL Eppendorf tube containing 1 mL dialysis buffer (20 mM Tris ace-

tate 80% cation at pH 7.5 and 4°C; 200 mM KCl; 2 mM MgCl₂; 20% [v/v] glycerol; 0.1% [v/v], IGEPAL CA-630; 1 \times PI), and the lid was closed quickly. For every tube this could be repeated twice. After flash dilution the protein was dialyzed in dialysis buffer for 2 h, frozen in liquid nitrogen, and stored at -80°C. Usually the purification resulted in 30 mL of 0.4–0.8 mg/mL protein solution.

ChIP assay

For the ChIP assay, TAP-tagged *BRF1* and its mutant variants were introduced in the pRS315 vector, which contains *BRF1* expressed under the control of its own promoter. For combinations of the mutations, S226L and T259M were cloned into pRS314 vector and cotransformed with the TAP-tagged *BRF1* variants T259M-TAP and S226L-TAP in the pRS315 vector, respectively. ChIP experiments were performed as previously described (Aparicio et al. 2005).

In vitro transcription assay

The Pol III-dependent in vitro transcription assay was performed as described by Hahn and Roberts (2000), with minor modifications. The nuclear extract was prepared as described by Seizl et al. (2011). The template used contained the *SUP4* gene from [-159] – [+244] with respect to the transcription start site of the tRNA. *SUP4* encodes a tyrosine tRNA. The 25 μ L transcription reaction contained 100 mM KOAc (pH 7.6), 20 mM HEPES (pH 7.6), 1 mM EDTA, 5 mM MgOAc, 2.5 mM DTT, 150 ng template, 192 μ g phosphocreatine, 0.2 μ g creatine phosphokinase, 10 U RiboLock RNase inhibitor (Fermentas), 50 μ g temperature-sensitive nuclear extract harboring a temperature-sensitive W107R mutation in *BRF1*, 40 μ M α -amanitin, and 100 ng of recombinant protein where indicated. Proteins were added to the transcription reaction mix for 5 min before addition of 0.1 mM NTPs. The transcription reaction was carried out for 60 min at 30°C. The RNA was isolated, and for primer extension, 0.125 pmol primer was annealed for 45 min at 60°C in 20 μ L 5 mM Tris (pH 8.3), 75 mM KCl, 1 mM EDTA (pH 8.0). The sequence of the 5'-Cy5 labeled primer was TCTCCCGGGGGCGAGTCGAACGCC. Two micrograms of Actinomycin D was added before 0.25 units MuLV reverse transcriptase in 60 μ L 5 mM Tris (pH 8.3), 75 mM KCl, 4.5 mM MgCl₂, 15 mM DTT, and 0.1 mM dNTPs extended the primer. The resulting cDNA was isolated by ethanol precipitation and resuspended in 4 μ L 0.04 mg/mL RNase A and 4 μ L formamide buffer (80% formamide, 25 mM EDTA, 1.5% bromphenolblue). Before loading, samples were boiled for 2 min at 95°C and put directly on ice. Samples were run on a 7 M urea, 8% polyacrylamide (35:1) gel in 1 \times TBE for 45 min at 180 V. The gel was pre-run for 5–10 min, and the pockets were rinsed before loading. Gels were analyzed and quantified with a typhoon scanner FLA9400 and ImageQuant Software (GE Healthcare).

Data access

The zebrafish RNA-seq data have been submitted to the NCBI Gene Expression Omnibus (GEO; <http://www.ncbi.nlm.nih.gov/geo/>) under accession number GSE63191. *BRF1* mutation data have been submitted to the NCBI ClinVar database (<http://www.ncbi.nlm.nih.gov/clinvar/>) under accession numbers SCV000192004, SCV000192005, SCV000192006, and SCV000195766.

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