Mutations in the Histone Methyltransferase Gene, *KMT2B* Cause Early Onset Dystonia

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ABSTRACT

Histone lysine methylation mediated by mixed-lineage leukemia (MLL) proteins, has emerged as critical in the regulation of gene expression, genomic stability, cell cycle and nuclear architecture. Although postulated to be essential for normal development, little is known about the specific functions of the different MLL lysine methyltransferases. Here, we report heterozygous mutations in \textit{KMT2B} (\textit{MLL4}) in 27 unrelated individuals with a complex progressive childhood-onset dystonia, often associated with a typical facial appearance and characteristic findings on brain magnetic resonance imaging. Over time the majority developed prominent cervical, cranial and laryngeal dystonia. Marked clinical benefit was observed following deep brain stimulation (DBS), leading to even restoration of independent ambulation in some cases. Decreased gene expression of \textit{THAP1} and \textit{TOR1A} was evident in cultured skin fibroblasts from subjects with \textit{KMT2B} mutations, with reduced THAP1 protein levels on immunoblotting. Analysis of cerebrospinal fluid from \textit{KMT2B} mutation-positive patients revealed markedly reduced levels of dopamine 2 receptor protein, with increased tyrosine hydroxylase levels. Our findings highlight a major new, clinically recognizable, and potentially treatable form of genetic dystonia, demonstrating the crucial role of KMT2B in the physiological control of voluntary movement.
INTRODUCTION

The control of voluntary movement is governed by interactive neural networks within the brain, involving the basal ganglia, sensorimotor cortex, cerebellum and thalamus\(^1\). Disruption of such pathways can lead to the development of a variety of motor disorders. Dystonia is one such movement disorder characterized by sustained or intermittent muscle contractions, causing abnormal, often repetitive movements and postures affecting the limbs, trunk, neck and face. Dystonic movements are typically patterned, twisting, and may be tremulous, often initiated or worsened by voluntary action and associated with overflow muscle activation\(^2\).

Dystonia is the 3\(^{rd}\) most commonly reported movement disorder worldwide\(^1\). It is described in a broad spectrum of genetic and acquired disorders, either in isolation or combined with other neurological and systemic features\(^2\). The precise pathophysiological processes remain yet to be fully elucidated, but defective dopaminergic signaling is thought to play an important role in many forms of isolated and complex dystonia\(^1,3-5\).

Despite genetic advances, the underlying cause remains elusive for a significant proportion of individuals with childhood-onset dystonia, hindering future prognostication and treatment strategies\(^6\). Here we report 27 individuals with an early-onset, complex, combined progressive dystonia associated with mono-allelic mutations in \textit{KMT2B} (\textit{MLL4}, OMIM *606834). \textit{KMT2B} encodes a lysine histone methyltransferase, involved in H3K4 methylation, an important epigenetic modification associated with active gene transcription.
RESULTS

Chromosomal microdeletions and intragenic KMT2B mutations in early-onset dystonia

We identified a cohort of 34 patients with undiagnosed childhood-onset dystonia for further molecular genetic investigation ([Online Methods, Supplementary Table 1, Supplementary Fig. 1]). On routine diagnostic testing, one case (Patient 1) was found to have a microdeletion at 19q13.12 of undetermined significance. Diagnostic chromosomal microarray was therefore undertaken in as many patients as logistically possible from this cohort (n=20) and overlapping microdeletions were detected in 5 more children (Supplementary Table 1, Patients 2-6). Using established networks (Online Methods, Supplementary Fig. 1), 4 more cases (Patients 7-10) with microdeletions were identified. In total, 10 patients (Patients 1-10) were found to have overlapping heterozygous interstitial microdeletions at 19q13.11-19q13.12 (Table 1a, Fig.1). Deletions detected on diagnostic microarray studies were confirmed by standard established laboratory protocols and confirmed de novo where parental testing was possible (Supplementary Table 2a). The smallest region of overlap extended from 36,191,100-36,229,548bp (GRCh37/Hg19), and contained two HUGO Gene Nomenclature Committee curated genes, ZBTB32 (zinc finger and BTB domain containing 32) and KMT2B (MLL4) (Fig. 1).

Of the remaining 28 patients from the original cohort, we undertook research exome (n=6) and genome sequencing (n=9) in 15 patients (Online Methods). Heterozygous variants of KMT2B were identified in 6/15 cases (Patients 13, 14, 17,
Subsequent Sanger sequencing of *KMT2B* in the other 13/28 individuals from the original cohort detected one more mutation-positive case (Patient 16). A further 10 cases (Patients 11, 12, 15, 18, 19, 20, 23, 24, 25, 26a) were ascertained through both national and international collaborators (Online Methods, Supplementary Fig. 1). In total, 17 patients with intragenic heterozygous *KMT2B* variants were identified, harboring frameshift insertions (n=1), frameshift deletions (n=6), splice site (n=1), stop-gain (n=2) and missense (n=7) mutations (Fig.1). All *KMT2B* mutations were confirmed on Sanger sequencing and parental segregation studies completed where DNA was available (Table 1a, Fig. 1, Supplementary Table 2a, Supplementary Fig. 2). No pathogenic variants in either *ZBTB32* or other known disease-associated genes (including genes causing clinically similar forms of dystonia) were otherwise identified in patients who had whole exome or genome sequencing. In the remaining patients, where further genetic testing was possible, mutations in *TOR1A, THAP1* and *GNAL* were excluded by diagnostic single gene testing, multiple gene panel testing or research Sanger sequencing (Supplementary Table 3).

**Phenotypic characterization of patients with KMT2B mutations**

Overall, we identified 27 patients (current age 6-40 years, 14 female, 13 male) with childhood-onset progressive dystonia (Table 1a, Table 1b, Supplementary Table 4, Supplementary Videos 1-7). Individuals presented in early childhood (1-9 years, median age 4 years) with either limb or cranio-cervical dystonia. Clinical presentation for those with microdeletions, frameshift, splice-site and stop-gain variants (mean age 4.1 years) occurred significantly earlier than for those with intragenic missense mutations (mean age 6.4 years) (p-value 0.0223)
Most patients (21/27) had lower limb symptoms at disease onset, leading to foot posturing, toe-walking and gait disturbance (Fig. 2a). 4/27 patients presented initially with upper limb symptoms associated with abnormal postures (Fig. 2b,c) and dystonic tremor, leading to reduced dexterity and handwriting difficulties (Supplementary Fig. 4a,b). With increasing age, cervical symptoms (torticollis, retrocollis) (Fig. 2d,e) and cranial involvement (facial dystonia, oromandibular involvement with dysarthria/anarthria and difficulties in chewing/swallowing) became prominent features in the majority of patients. In many patients, progressively severe dysphonia was suggestive of laryngeal involvement. None of the patients had airway compromise and videostroboscopy was not undertaken. Over time, the majority of patients (24/27) developed progressive generalized dystonia, 2-11 years after initial presentation (Fig. 2f). The dystonia was persistent in nature, absent in sleep, worsened by voluntary action and associated with overflow muscle activation. Some patients had dystonic tremor. Sudden, brief, involuntary muscle jerks, clinically consistent with myoclonus was evident in 2 cases (Patients 14 and 27). For a few subjects, dystonia was exacerbated when systemically unwell. Stepwise deterioration following intercurrent illness was particularly evident in Patient 14, and status dystonicus, triggered by a urinary tract infection, was reported in Patient 3.

Many patients with KMT2B mutations had further clinical findings. Additional neurological symptoms and signs were evident in some patients, including microcephaly, seizures, spasticity and eye movement abnormalities (strabismus, saccade initiation failure and oculomotor apraxia) (Table 1b). Dysmorphic features and characteristic facial appearance (elongated face and bulbous nasal tip) (Fig. 2g, Table 1b) were commonly reported. Delay in neurodevelopmental milestones,
intellectual disability, systemic (dermatological, renal, respiratory) features and psychiatric symptoms were also present in some individuals (Table 1b, Supplementary Table 4, Supplementary Fig. 4c). Malignancies were not reported in any patients. Cerebrospinal fluid (CSF) neurotransmitter analysis was undertaken in 13 patients revealing no major derangement of monoamine metabolites (Supplementary Table 5a). Magnetic resonance (MR) imaging revealed a characteristic signature in 17/22 patients who had imaging sequences suitable for assessment (Supplementary Table 5b). Subtle symmetrical hypointensity of the globus pallidi (with a hypointense streak of bilateral globus pallidus externa) was evident on MR images known to be sensitive to the magnetic resonance phenomenon of susceptibility (T2*-susceptibility- and echo-planar imaging b0-diffusion-imaging datasets) (Fig. 3). The mean age at neuroimaging was significantly lower for patients with MR abnormalities (11.7 years) than for those with normal brain scans (19.0 years) (p-value 0.0167) (Supplementary Fig. 3b).

Single positron emission tomography using 123I (DaTSCAN™) and 18 FDG-PET-CT glucose uptake studies, each undertaken in 3 patients, were normal (Supplementary Table 5b, Supplementary Fig. 4d).

Deep brain stimulation: clinical benefit in KMT2B-dystonia

Overall, medical therapies were not of clinical benefit in this patient cohort. None of the patients had a sustained response to levodopa treatment, nor other commonly used anti-dystonic agents (Table 1a). Due to the medically intractable, progressive nature of disease, 10 patients had symptomatic treatment with bilateral globus pallidus interna-deep brain stimulation (GPI-DBS) (Table 1a). All showed clinical benefit with DBS (which was particularly striking in some of the younger patients).
with overall amelioration of dystonia, improved oromandibular symptoms, better upper and lower limb function and even restoration of independent ambulation in some patients. Patient 6 showed significant improvement of torticollis and retrocollis, as well as in overall function and gait after DBS. Patient 8 showed a sustained clinical response 6 years after DBS insertion, with improvement of dystonia, even more evident after replacement of a faulty right DBS lead. Patient 9 had generalized dystonia and could not walk independently pre-DBS. Two weeks post-DBS insertion he dramatically regained independent ambulation with marked improvement of dystonic symptoms (Supplementary Video 8). Patient 17 and 21 were predominantly wheelchair-dependent pre-DBS insertion, but both patients showed restoration of independent walking and improvement of dystonia after DBS (Supplementary Video 9,10). Patient 19 had improvement in oromandibular symptoms with DBS. Patient 20 had DBS insertion at age 32 years and although most benefits were only transient, sustained improvement of foot posture was reported. Patient 23 had significant amelioration of dystonia symptoms after DBS insertion. Patient 22, now 9 months post-DBS (Supplementary Video 11) and Patient 25, 4 months post-DBS have both shown significant gains in hand function and independent walking with reduction of dystonia. Five patients in the cohort are now over three years post-surgery, and the observed reduction of dystonia, restoration of function and prevention of progressive disability is evidence of sustained clinical benefit.

**KMT2B is constrained for missense and predicted protein truncating variants**

Four individuals (Patient 13, 14, 17 and 21) had whole genome sequencing as part of the NIHR-funded BioResource-Rare Disease project. Enrichment analysis was undertaken in this cohort in order to determine whether predicted protein truncating
variants (PPTVs) in KMT2B are observed more frequently in patients than would be expected by chance. Given the size and sequence context of KMT2B, $5.73 \times 10^{-03}$ de novo PPTVs are expected to occur by chance in KMT2B in the subset of the NIHR BioResource- Rare Diseases cohort who have pediatric onset neurological disease, but 3 PPTVs are observed. This represents a significant enrichment (p-value $3.12 \times 10^{-08}$). Furthermore in ExAC, KMT2B is also highly constrained for PPTVs. In the ExAC database of 60,706 individuals (Exome Aggregation Consortium (ExAC), Cambridge, MA (URL: http://exac.broadinstitute.org, accessed July 2016)$^8$, there are only 5 PPTVs that are not flagged as having dubious variant annotation. All are extremely rare (4 are found in a single individual and one occurs in 2 individuals). Given the size and sequence context of the gene, the presence of so few PPTVs in a cohort of 60,706 individuals reveals KMT2B to be highly constrained for such variation, providing supportive evidence of its pathogenicity. Regarding variants in the ExAC database, there are 712 reported non-synonymous changes. Most of these are rare, as expected for a cohort of this size, and the median CADD score$^9$ for these variants is 22.9. The median CADD score for missense mutations identified in our KMT2B-dystonia cohort is significantly higher at 29.1 (p-value $0.0001364$; Supplementary Table 2b). Furthermore, given the size and sequence context of KMT2B, 956 missense variants are predicted to occur by chance, suggesting that KMT2B may also be constrained for missense variation (z=4.06)$^8$.

**KMT2B variants are predicted to destabilize protein structure**

*In silico* homology modelling studies were undertaken to generate hypotheses regarding the predicted effects of mutations on KMT2B structure-function properties (Supplementary Results). Based on Pfam domain assignments, KMT2B has a
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CXXC zinc finger domain, multiple PHD domains, an F/Y rich N-terminus (FYRN), FYRC (F/Y rich N-terminus) domain and a C-terminal SET domain (Fig. 4a). The modelled mutations occurred in residues within the PHD-like, FYRN, SET and FYRC-SET linking domains (Fig. 4b-d). Evaluation of a number of mutations using MAESTRO\textsuperscript{10} and DUET\textsuperscript{11} suggests change in free energy, with a predicted structure destabilizing effect (Supplementary Results).

Mutations Phe1662Leu and Gly1652Asp occur within a PHD-like domain (residues 1574-1688), predicted to facilitate interaction with DNA, protein-protein interaction and recognition of methylated/unmethylated lysines\textsuperscript{12-14}. Extensive hydrophobic interactions hold the globular structure of this region, which is important for its function\textsuperscript{12}. Phe1662 is fully buried at the core, stabilizing the structure of this PHD-like domain while Gly1652 is partially buried (Fig. 4b,e,f). Phe1662 is involved in multiple hydrophobic contacts at the core of the PHD domain, and mutation to leucine is predicted to cause loss of contacts at the core (Fig. 4g). Gly1652 is located on a loop (Fig. 4e) and mutation to aspartic acid is predicted to alter surface charge, with possible effect on the interaction network in the vicinity, involving a positively charged Arg1635, part of the helix $\alpha$3 implicated in DNA binding\textsuperscript{12}. Arg1762 and Leu1781 occur in a FYRN domain. FYRN and FYRC regions, particularly common in MLL histone methyltransferases, interact to form a compact structural unit (Fig. 4c,h), important in maintaining the active structure\textsuperscript{15,16}. Arg1762 forms hydrogen bonds with the backbone carboxyls of Arg2463 and Leu2464 of FYRC domain. Substitution of Arg1762 by cysteine is predicted to abolish these contacts and hence contribute to destabilization of FYRC-FYRN association. Leu1781, at the interface between FYRN and FYRC (Fig. 4h,i) is surface exposed and involved in backbone hydrogen bonds stabilizing the beta sheet formed
together by the two domains. Mutation to proline is predicted to disrupt the backbone hydrogen bond at this position, because it lacks one hydrogen bond donor and its backbone torsion angles are not compatible with that of a beta sheet, with a predicted destabilizing effect on sheet structure, potentially affecting the normal association of FYRN and FYRC domains. Arg2517 resides in the region linking FYRC and SET domains, known to bind WDR5, an effector required for trimethylation of histone H3\textsuperscript{17}, presenting methylated histone H3 substrates to the MLL complex for further methylation\textsuperscript{18}. Arg2517 is thought to be involved in a salt-bridge interaction with Asp172 of WDR5 (Fig. 4j) and Arg2517Trp is predicted to lead to loss of this interaction. Ile2674, Tyr2688 and Ile2694 occur in the catalytic methyltransferase SET domain common to histone lysine methyltransferases. Ile2674 is buried in the hydrophobic core, adjacent to the catalytic site (Fig. 4d,k). Mutation to threonine is predicted to lead to loss of contacts at the core of the domain (due to the shorter side chain) and also introduces a buried polar group (Fig. 4k,l). Tyr2688Thr occurs at the core of SET domain involving extensive hydrophobic interactions and a hydrogen bond interaction with Ser2661 (Fig 4m). The frameshift mutation Tyr2688Thrfs*50 with insertion of 50 additional residues, is predicted to destabilise the core and affect contacts due to the substitution with a shorter non-aromatic side-chain. Ile2694 is involved in the extensive hydrophobic contacts stabilizing the core of this domain. \textit{In silico} analysis predicts that the frameshift mutation Ile2694Serfs*44 will disrupt the domain fold and affect methytransferase activity.

\textit{KMT2B is ubiquitously expressed with reduced expression in KMT2B-dystonia}
We confirmed widespread *KMT2B* expression in a variety of control fetal and adult human tissues ([Fig. 5a](#)). Moreover, *KMT2B* is ubiquitously expressed in the brain, with higher expression in the cerebellum than any other region ([Fig. 5b](#)). We ascertained fibroblasts from 4 patients (Patient 2, 13, 14, 16, with either microdeletions or PPTVs in *KMT2B*) and detected a statistically significant decrease in fibroblast *KMT2B* expression on quantitative RT-PCR when compared to control fibroblasts ([Fig. 5c](#)).

**Histone H3K4 methylation is not globally reduced in KMT2B-dystonia**

To determine the effect of *KMT2B* mutations on methylation of lysine 4 on histone H3 (H3K4 methylation), we assayed tri-methylated H3K4 (H3K4me3) and di-methylated H3K4 (H3K4me2). Immunoblotting of histones extracted from fibroblasts of Patient 14 and 16 showed no significant reduction in H3K4me3 or H3K4me2 relative to control samples ([Fig. 5d](#), Supplementary Fig. S5a). We used the model species *Dictyostelium discoideum* to test the effect of SET domain mutation Ile2647Thr on *in vivo* histone methyltransferase activity. The SET domain of KMT2B shares 56% sequence identity with the *Dictyostelium* orthologue DdSet1, and Ile2647 is conserved (corresponding amino acid in *Dictyostelium* is Ile1447) ([Fig. 1h](#)). DdSet1 is the only H3K4 methyltransferase in *Dictyostelium* and targeted knockout of *DdSet1* (set1) results in loss of all methylation at H3K4[19]. We constitutively expressed wild-type DdSet1 (WT-DdSet1) and mutant-DdSet1 (m-DdSet1), both with N-terminal GFP fusions, in set1- *Dictyostelium* cells and compared the resulting levels of H3K4 methylation. Expression of either GFP-WT-DdSet1 or GFP-mDdSet1 in set1- cells resulted in rescue of H3K4 tri-methylation to wild type levels ([Fig. 5e](#), Supplementary Fig. S5b, S5c).
Fibroblast THAP1 gene and protein expression is reduced in KMT2B-dystonia

In order to determine whether KMT2B-dystonia is associated with dysregulation of specific genes implicated in the control of movement, we investigated the expression profiles of TOR1A and THAP1. Fibroblasts derived from 4 patients (Patients 2, 13, 14, 16) showed significantly reduced transcript levels of THAP1 and TOR1A when compared to control fibroblasts (Fig. 5f). Fibroblast immunoblotting studies showed a statistically significant reduction in THAP1 protein expression in all 4 patients when compared to control samples (Fig. 5g). A statistically significant reduction in TOR1A protein level was evident in Patient 14, though not in the other patients (Fig. 5h).

Abnormal CSF levels of dopaminergic proteins in KMT2B-dystonia

CSF immunoblotting studies were undertaken in two patients for whom samples were available for research testing (Patient 2 and 16). Both patients had markedly reduced levels of dopamine 2 receptor (D2R), 56.9% and 59.8% of levels observed in control CSF (Controls = 1.09±0.21SD, KMT2B patients = 0.64±0.02SD). In contrast, an increase in tyrosine hydroxylase (TH) levels was seen in both mutation-positive patients (173.3% and 170.9% of levels seen in control CSF) (Controls = 0.52±0.08SD, KMT2B patients = 0.90±0.01SD) (Fig. 5i).

DISCUSSION

We report 27 individuals with heterozygous mutations in the lysine methyltransferase gene, KMT2B, and define a new genetic movement disorder that importantly, is amenable to treatment with DBS. Using the current classification system², KMT2B-dystonia is defined as an inherited autosomal dominant, complex,
combined dystonia usually of infantile or childhood-onset. In most patients, the
dystonia is persistent and progressive in nature. The majority of individuals develop
4-limb dystonia with particularly prominent cervical, laryngeal and oromandibular
symptoms. Whilst the majority of patients in this cohort seem to follow this disease
trajectory, we also report atypical cases with relatively little limb involvement and
either mainly oromanibular features (Patient 18) or paroxysmal cervical dystonia
(Patient 26a).

For many patients, KMT2B-dystonia is associated with a number of additional
clinical features, including other neurological symptoms, intellectual disability,
psychiatric co-morbidity, dysmorphia, skin lesions and other systemic signs. Given
the association with active gene expression, is possible that KMT2B could account
for these additional disease features. For Patients 1-10, is also possible that other
genes within their microdeletion could contribute to aspects of their clinical
phenotype. Indeed, cutis aplasia and ectodermal dysplasia have been reported in
patients with more proximal deletions of chromosome 19q13.11\textsuperscript{20}. KMT2B is
therefore a complex dystonia, and affected patients should have close surveillance
of development during childhood, regular neurology assessments, routine
dermatological review and formal neuropsychiatric testing.

In KMT2B-dystonia, the majority of patients had a characteristic pattern on MR
imaging, with very subtle low pallidal signal on T2\textsuperscript{*}-, diffusion- and susceptibility-
weighted sequences, particularly affecting the lateral aspect of the globus pallidus
externa (Fig. 3). Although genotype did not appear to influence whether MR
findings were evident, those with abnormal imaging had scans undertaken at a
significantly younger age than those with normal imaging. Indeed, MR abnormalities
could possibly be an age-dependent phenomenon, perhaps becoming less
apparent with increasing age, as was evident in Patient 22 (Supplementary Table 5b, Supplementary Fig. 3b,c,d). The overall significance of the identified neuroradiological abnormalities remains unclear. Such radiological findings are reminiscent of, but much more subtle and different to those reported in classical Neurodegeneration with Brain Iron Accumulation (NBIA) syndromes\textsuperscript{21,22}. Similar non-specific features of T2*-weighted hypointensity are increasingly recognized in a number of other neurological conditions, including Huntington’s disease, \textit{TUBB4A}-related disorders, GM1 gangliosidosis, alpha-fucosidosis and mitochondriocytopathies.

In the original UCL-ICH Dystonia cohort, \textit{KMT2B} mutations were identified in 13/34 (38\%) individuals with a relatively homogenous phenotype of early onset progressive dystonia. In other screened cohorts, mutation detection rates varied from 1.3-30\%, with more cases identified from cohorts that were tightly phenotyped for dystonia (Supplementary Fig. 1). In screened cases where \textit{KMT2B} mutations were not detected, it is likely that these individuals have another underlying etiology accounting for their symptoms, although it is possible that (i) single/multiple exon \textit{KMT2B} deletions and duplications may have been missed on microarray, Sanger sequencing and whole exome/genome sequencing and (ii) promoter mutations and deeply intronic \textit{KMT2B} variants may not have been detected by whole exome and Sanger sequencing.

The majority of individuals with \textit{KMT2B} mutations (20/27, Patients 1-20) had either heterozygous interstitial microdeletions leading to \textit{KMT2B} haploinsufficiency, or variants predicted to cause protein truncation, protein elongation, splicing defects or nonsense-mediated mRNA decay. The remaining 7 patients (Patients 21-27) had previously unreported non-synonymous variants of \textit{KMT2B}, all affecting conserved
residues within key protein domains (Fig. 1c-h), and in silico studies predict destabilization of protein structure. Notably, initial disease presentation was significantly earlier in Patients 1-20 than in those with missense variants (Supplementary Fig. 3a). In KMT2B-dystonia, genotype did not however influence the rate of symptom evolution, disease severity or response to DBS.

For the majority of patients, KMT2B mutations were confirmed as de novo where parental testing could be undertaken. In our cohort, 3 patients had missense mutations that were all maternally inherited (Patient 22, 26a, 27). Given this observation of maternal inheritance, the possibility of imprinting at the disease locus was considered, but deemed unlikely, given (i) de novo microdeletions in Patients 2 and 10 occurred on paternally inherited alleles and (ii) there is evidence of bi-allelic expression of KMT2B single nucleotide polymorphisms in human tissues, including brain (Supplementary Fig. S6). Importantly, whole exome sequence analysis undertaken in Patients 22, 26a and 27 did not identify other rare or de novo variants to account for their disease. Interestingly, Patient 26a inherited p.Arg2517Trp from his symptomatic mother (26b) who also had (milder) disease symptoms. She reported gait abnormalities and a progressive inability to run, as well as periodic paroxysmal upper limb and neck dystonia. She also had a bulbous nasal tip, like her son (Fig. 2g). In contrast, both mothers of Patients 22 and 27 were clinically examined, and neither had evidence of a motor phenotype, intellectual disability, other neurological features, neuropsychiatric symptoms, facial dysmorphia, skin lesions or other systemic signs. The identification of both symptomatic and asymptomatic carriers suggests that there may be either 'apparent' incomplete penetrance, due to parental mosaicism, or true incomplete disease penetrance, a phenomenon commonly reported in a number of other autosomal dominant genetic
dystonias\textsuperscript{23,24}. Furthermore, other genetic, epigenetic and environmental modifiers may also influence disease penetrance and phenotypic presentation in KMT2B-dystonia.

\textit{KMT2B} encodes an ubiquitously expressed lysine methyltransferase specifically involved in H3K4 methylation\textsuperscript{25,26}, an important epigenetic modification associated with active transcription. H3K4me3 is enriched at promoters, marking transcription start sites of actively transcribed genes, whereas H3K4me1 is associated with active enhancer sequences\textsuperscript{27}. H3K4me2 is less specifically localized, but may be enriched at transcription factor binding sites\textsuperscript{28}. Members of the SET/MLL protein family, including KMT2B, are responsible for the generation of H3K4me1, H3K4me2, and H3K4me3, essential for gene activation in normal development\textsuperscript{29}.

Using patient-derived fibroblasts and a \textit{Dictyostelium discoideum} model, we demonstrated that \textit{KMT2B} mutations are not associated with widespread alterations in overall levels of H3K4 methylation. This is not surprising, given that haploinsufficiency of other MLL family members have not been convincingly shown to affect global H3K4 levels. The fundamental physiological role of MLL proteins is however further affirmed by the observation that loss-of-function heterozygous mutations in MLL-encoding genes are reported in human developmental disorders\textsuperscript{30}, namely Wiedemann Steiner (\textit{KMT2A, MLL1})\textsuperscript{31}, Kleefstra-like (\textit{KMT2C, MLL3})\textsuperscript{32}, Kabuki (\textit{KMT2D, MLL2})\textsuperscript{33} syndrome, and most recently \textit{SETD1A}-related disease (\textit{KMT2F})\textsuperscript{34}. The physiological functions of MLL proteins are yet to be fully characterized, however, the observation that mutations in different MLL genes cause phenotypically distinct syndromes (\textbf{Supplementary Table 6}) suggests that each MLL protein has a unique role regulating the expression of a specific set of genes\textsuperscript{35,36}. 
Amongst the 4 reported MLL-gene disorders, dystonia appears unique to KMT2B-related disease and is not described in other MLL syndromes (Supplementary Table 6), providing further evidence that different MLL proteins mediate the activation and transcription of a specific set of genes, with temporal and cellular context. We utilized fibroblasts and CSF derived from patients to investigate downstream effects of KMT2B mutations on specific gene expression. The rationale for investigating THAP1 and TOR1A in the first instance was based on a number of factors, namely that (i) loss-of-function mutations in both genes cause progressive generalized dystonia with cervical, oromandibular and laryngeal symptoms, similar to those seen in KMT2B-dystonia, (ii) both genes are expressed in fibroblasts, facilitating investigation in patient-derived tissue, (iii) analysis of methylation profiles using ENCODE demonstrates a sharp H3K4me3 peak at the 5' region of both THAP1 and TOR1A in a wide range of cell types, including brain cells (Supplementary Fig. 7), (iv) on human brain expression profiles, THAP1 and KMT2B similarly display highest expression in the cerebellum (Fig 5b, Supplementary Fig. 8). We detected statistically significant reduced levels of THAP1 and TOR1A gene expression and THAP1 protein expression in patient fibroblasts. The mechanisms causing such alterations in KMT2B-dystonia remain yet to be elucidated. Whilst H3K4 methylation is clearly associated with the process of active transcription, several studies have shown that H3K4 methylation is required, not for absolute transcriptional output, but rather for transcription stability or consistency. Recent studies have suggested that H3K4Me is required to minimize transcriptional variability between cells in a population, rather than absolute expression, so the effects of KMT2B haploinsufficiency on THAP1/TOR1A levels could conceivably operate via an intermediary sensitive to
stochastic fluctuations. Whilst our study focuses on two genes, it is highly likely that
dysregulation of other genes and proteins are also involved in the disease
pathophysiology of KMT2B-dystonia. Further studies will determine whether
expression profiles of other genes are affected in KMT2B-dystonia and contributory
to the phenotype.

CSF analysis is increasingly recognized as a highly useful tool for studying synaptic
proteins and dysregulation of the dopaminergic system\textsuperscript{43}. In our study, CSF
immunoblotting studies revealed significant reduction of D2R protein and increase
in TH levels in patients with KMT2B-dystonia when compared to control CSF
samples. Downregulation of D2R could conceivably impair post-synaptic activation
of coupled G-proteins with subsequent downstream effects, as seen in other
inherited dystonias. Reduced D2R striatal availability is reported in patients with
DYT1 and DYT6 dystonia\textsuperscript{44,45}. Furthermore, D2R dysfunction is described in murine
DYT1 models, where aberrant D2R-mediated responses are associated with
reduced D2R protein levels and impaired G-protein activation\textsuperscript{46}. The observed rise
in CSF TH levels could be secondary to reduced pre-synaptic D2 autoreceptors\textsuperscript{47},
which could conceivably impact dopamine synthesis and bioavailability.
Interestingly patients with KMT2B-dystonia had normal CSF levels of the stable
dopamine metabolite, homovanillic acid (HVA), indicating normal dopamine
turnover (conversion of dopamine to HVA by monoamine oxidase and catechol-o-
methyl transferase). Whilst CSF HVA levels accurately reflect dopamine turnover,
they are not always a true indicator of dopamine synthesis and bioavailability. Low
CSF HVA levels often correlate with low dopamine levels in patients with inherited
disorders of dopamine deficiency such as TH deficiency, aromatic L-amino acid
decarboxylase deficiency, and many inherited pterin defects\textsuperscript{3}. However HVA levels
may also be normal (autosomal dominant GTP cyclohydrolase deficiency) or elevated (Dopamine Transporter Deficiency Syndrome) in diseases known to be associated with dopamine deficiency\textsuperscript{3,48}. Normal HVA levels in KMT2B-dystonia may therefore not reflect true dopamine levels. Indeed, the effect of \textit{KMT2B} mutations on dopamine synthesis and bioavailability remain yet to be fully elucidated.

In conclusion, we report \textit{KMT2B} mutations in 27 patients with a clinically recognizable, distinct form of dystonia. To date, the underlying etiology is only genetically resolved in a minority of childhood-onset cases of dystonia, which precludes confirmatory diagnosis, accurate disease prognostication and selection of appropriate treatment strategies. We have shown that many patients with molecular confirmation of KMT2B-dystonia have significant, sustained clinical improvement with DBS and referral for DBS assessment should thus be considered for this group of individuals. Identification of additional cases will allow further characterization of the full phenotypic disease spectrum. Our report highlights mutations in \textit{KMT2B} as a new and important cause of early-onset dystonia, emphasizing the crucial role of KMT2B in the control of normal voluntary movement.
References


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Meyer et al 2016


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Disclosures

HP has unrestricted support for Educational Activity from Medtronic.
Figure 1
Figure 1:

**Molecular Genetics Findings in Patients with KMT2B Mutations**

(a) Top panel: Schematic representation of chromosome 19. Middle panel: Ten microdeletions on 19q13.11-19q13.12 are displayed as horizontal red bars (GRCh37/Hg19). Lower panel: The smallest overlapping region comprises two genes, **ZBTB32** and **KMT2B**. (b) Schematic of exon-intron structure of **KMT2B** (based on NCBI Reference Sequence: NM_014727.2) is shown indicating the location of the 7 frameshift insertions and deletions (blue, above gene), 2 stop-gain mutations (dark red, above gene), 1 splice site variant (purple, below gene) and 7 missense changes (in green, below gene).

(c) Alignment of KMT2B amino acid sequences from seven different species, the human paralog KMT2A (another member of the MLL protein family) and the human PHF6 protein used to model the PHD-like domain. Gly1652 (in red) is highly conserved in all listed amino acid sequences, while the Phe1662 residue (in red) is either conserved or tolerates replacement by the similar amino acid, Tyr without predicted functional effect (**Supplementary Fig. 10**). (d-g) Alignment of KMT2B amino acid sequences from seven different species and the human paralog KMT2A. (d) Arg1705 (in red) is conserved to zebrafish and in human KMT2A. (e) Arg1762 is fully conserved throughout species. (f) Leu1781 (in red) is conserved in all listed mammalian homologs of KMT2B. (g) Arg2517 (in red) is conserved to zebrafish and in human KMT2A. (h) Alignment of KMT2B amino acid sequences from seven different species, the human paralog KMT2A and SET domain containing proteins (human SETD1A/SETD1B, yeast Set1, *Dictyostelium discoideum* Set1. Ile2674 (in red) is highly conserved in all listed amino acid sequences. Residues matching human KMT2B (grey), not matching (white), amino acids conserved in all representative sequences (dark grey). * positions of fully conserved residues; :
conservation between groups of strongly similar properties and conservation between groups of weakly similar properties.

Figure 2
Figure 2:

Clinical Features of Patients with KMT2B Mutations

(a) Patient 17, age 13 years, evidence of gait disturbance with dystonic posturing of the four limbs. (b) Patient 27, age 19 years and (c) Patient 14, age 18 years both showing bilateral upper limb dystonic posturing. (d, e) Patient 23, age 8 years with retrocollis. (f) Patient 12, age 6 years, generalized dystonia, with jaw-opening dystonia and 4-limb posturing. (g) Montage of patient faces: Top row (left to right) Patients 1, 2, 3, 4, 8, 9; Middle row (left to right) Patients 11, 12, 13, 14, 16, 17 and Bottom row (left to right) Patients 21, 23, 25, 26a, 26b, 27. Facial elongation, broad nasal base and bulbous nasal tip, particular evident in Patients 1, 2, 4, 9, 11, 12, 14, 17, 23, 25, 26a, 26b, 27.
Figure 3: Radiological Features in KMT2B-patients

Magnetic resonance imaging (MRI) with T2*-weighted sequences (a-d), diffusion-imaging datasets with b-value of zero (e-h) and susceptibility weighted sequences (i-l). Abnormal findings indicated by yellow arrows. (a,e,i) Representative MRI from control subjects for T2*-weighted sequences (a: age 10y2m), diffusion-weighted sequences (e: age 10y4m) and susceptibility weighted sequences (i: age 10y8m) indicating normal appearances of basal ganglia on all three sequences. (b,f,j) Patient 1, age 9y5m, (c,g,k) Patient 13, age 11y3m, (d) Patient 9, age 15y1m, (h) Patient 22 age 13y1m and (l) Patient 25, age 16y.
all show evidence of bilateral subtle hypointensity of the globus pallidus with hypointense lateral streak of globus pallidus externa.
Figure 4
Figure 4:

Homology Modelling of KMT2B Protein Structure

(a) Schematic of domain architecture of KMT2B. (b-d) The degree of amino conservation is displayed in the structural models for the different domains. Red to blue indicates increasing conservation. (b) Model of PHD-like domain shows the mutation sites Gly1652 and Phe1662 (c) Model of the FYRN domain presents the position and conservation of Arg1762 and Leu1781. (d) Model of the SET methyltransferase domain indicates the position and conservation of Ile2674, Tyr2688 and Ile2694. (e) Location of Gly1652 in the PHD-like domain model and the hydrogen bond network in the vicinity are shown. Helix α3 is also indicated. (f) Hydrophobic packing involving Phe1662 (green) is displayed. (g) Change to leucine (green) at position 1662 is predicted to cause loss of contact within the hydrophobic core. Residue side chains are presented as spheres highlighting van der Waals contacts. (h) Interactions involving Arg1762 (green) from FYRN with Arg2463 and Leu2464 of FYRC. The hydrogen bond interactions are highlighted. (i) Leu1781 shown at the interface of FYRN (pink)/FYRC (blue) domains. The backbone hydrogen bonds stabilizing the sheet structure are highlighted. (j) Interactions involving Arg2517 (green) and WDR5 (brown). The salt bridge interaction between Arg2517 of KMT2B and Asp172 of WDR5 is highlighted. (k) Location and contacts involving Ile2674 (green) in the hydrophobic core of the SET domain are exhibits. SAH is displayed in light brown. (l) Conversion to threonine (green) at position 2674 is predicted to result in loss of contacts in the core. (m) Interactions involving Tyr2688 (light green) and Ile2694 (dark green) in the core of the SET domain. The hydrogen bond between Tyr2688 and Ser2661 is highlighted.
**Figure 5**

- **Panel a:** KMT2B expression
- **Panel b:** THAP1/TOR1A expression
- **Panel c:** THAP1 protein expression
- **Panel d:** TOR1A protein expression
- **Panel e:** Dopaminergic Markers in CSF
**Figure 5:**

**Functional Investigation of the Downstream Effects of Mutations in** *KMT2B*

(a) PCR analysis of human fetal and adult cDNA for expression of *KMT2B*. *KMT2B* is widely expressed in a range of human tissues, including fibroblasts, brain tissue and midbrain dopaminergic neurons. (b) Box plots of *KMT2B* mRNA expression levels in 10 adult brain regions (source: BRAINEAC; http://www.braineac.org/). The expression levels are based on exon array experiments as previously described and are plotted on a log2 scale (y axis). This plot shows that *KMT2B* is ubiquitously expressed across all 10 brain regions analyzed, with expression higher in the cerebellum than in any other region. Putamen (PUTM), frontal cortex (FCTX), temporal cortex (TCTX), occipital cortex (OCTX), hippocampus (HIPP), substantia nigra (SNIG), medulla (specifically inferior olivary nucleus, MEDU), intralobular white matter (WHMT), thalamus (THAL), and cerebellar cortex (CRBL). “N” indicates the number of brain samples analyzed to generate the results for each brain region. Whiskers extend from the box to 1.53 the interquartile range. (c) Quantitative RT-PCR indicates that patients with *KMT2B* mutations (n=4) have significantly decreased fibroblast mRNA levels of *KMT2B* when compared to controls (Controls = 1.01±0.16SD, n=3 technical replicates of 2 biological samples; Patients = 0.57±0.12SD, n=3 technical replicates of 4 biological samples; two-tailed unpaired t-test, p-value 0.0182). (d) Quantification of immunoblotting of tri-methyl H3K4 (left) and di-methyl H3K4 (right) in histones extracted from patient-derived fibroblasts (Patient 14 and 16), and two control fibroblast cell lines. Methylation values are normalized to pan-histone H3 levels. Individual data-points are plotted with center bar showing mean and error bars showing standard deviation. Differences between control and patient-derived samples are not significant (H3K4me3: Controls = 96.63±19.98SD; Patient 16 = 104.1±40.31SD; Patient 14 = 94.75±38.36SD; p=0.62; H3K4me2: Controls = 94.33±19.25SD; Patient 16 = 127.8±20.79SD; Patient 14 = 80.23±31.09SD; p=0.07). n=3 fibroblast samples (technical
replicates). (e) Quantification of immunoblotting of tri-methyl H3K4 in *Dictyostelium* cell lysates. Tri-methyl H3K4 intensity values are normalized against levels of total histone H3. H3K4 tri-methylation is impaired in set1- cells compared to wild type. Expression of GFP-DdSet1 or GFP-DdSet1(Ile1447Thr) in set1- cells rescues levels of H3K4Me3. Individual data-points are plotted with center bar showing mean and error bars showing standard deviation (Wild type = 115±48.25SD; set1- = 5.94±9.37SD; set1- GFP-DdSet1(I1447T) 1 = 133.7±38.11SD; set1- GFP-DdSet1(I1447T) 2 = 129.8±42.34SD; set1- GFP-DdSet1(I1447T) 3 = 96.07±31.82SD; set1- GFP-DdSet1 = 110.5±12.02SD). n=3 samples (technical replicates). (f) Quantitative RT-PCR of *THAP1* and *TOR1A*, indicates that patients have a reduction of *THAP1*, and to a lesser extent of *TOR1A* transcripts in comparison to controls (*THAP1*: Controls = 1.04±0.40SD, n=3 technical replicates of 2 biological samples; Patients = 0.55±0.05SD, n=3 technical replicates of 4 biological samples; two-tailed unpaired t-test, p-value 0.0498; *TOR1A*: Controls = 1.00±0.05SD, n=3 technical replicates of 2 biological samples; Patients = 0.79±0.06SD, n=3 technical replicates of 4 biological samples; two-tailed unpaired t-test, p-value 0.0140). (g) Immunoblotting studies in fibroblasts indicate a significant reduction in THAP1 for Patient 2, 13, 14 and 16 when compared to controls (Control 1 = 1.34±0.02SD; Control 2 = 1.80±0.11SD; Patient 2 = 0.83±0.06SD; Patient 13 = 0.77±0.17SD; Patient 14 = 0.53±0.04SD; Patient 16 = 0.54±0.06SD; Kruskal-Wallis test, p-value 0.0078). n=3 fibroblast protein samples (technical replicates). (h) Immunoblotting studies in fibroblasts indicate a statistically reduced level of TOR1A in Patient 14 when compared to controls (Control 1 = 1.08±0.02SD; Control 2 = 1.13±0.22SD; Patient 14 = 0.64±0.05SD; two-tailed unpaired t-test, p-value 0.0196), but not for Patient 2, 13 and 16 (Patient 2 = 1.09±0.27SD; Patient 13 = 1.39±0.18SD; Patient 16 = 1.13±0.29SD; Kruskal-Wallis test, p-value 0.0812). n=3 fibroblast protein samples (technical replicates). (i) CSF immunoblotting studies on Patient
2 and 16 show markedly reduced levels of D2R and increased levels of TH when compared to control CSF (D2R: Controls = 1.09±0.21SD, n=4 control CSF samples (biological replicates); Patients = 0.64±0.02SD, n=2 patient CSF samples (biological replicates); two-tailed unpaired t-test, p-value 0.0471; TH: Controls = 0.52±0.08SD, n=4 control CSF samples (biological replicates); Patients = 0.90±0.01SD, n=2 patient CSF samples (biological replicates); two-tailed unpaired t-test, p-value 0.0036).
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<tr>
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<td>Age (y)</td>
<td>Sex</td>
<td>KMT2B mutation</td>
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<td>Onset of dystonia (y)</td>
<td>Bilateral LL involvement (y)</td>
<td>Bilateral UL involvement (y)</td>
<td>Onset of cranial, cervical, laryngeal dystonia (y)</td>
<td>Symptoms of cranial, cervical, laryngeal dystonia</td>
<td>Trial of medication and clinical response</td>
<td>Deep brain stimulation (DBS)</td>
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<td>16</td>
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<td>Bilateral LL increasing falls Gait disturbance</td>
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<td>5</td>
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<td>17</td>
<td>17</td>
<td>M</td>
<td>c.6515_6518delins</td>
<td>Gait disturbance</td>
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<td>12</td>
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<td>L-dopa trial – no benefit TBZ – no benefit BLF and THP – mild benefit</td>
<td>Inserted age 16y Very good clinical response 4m post-DBS with restoration of independent ambulation</td>
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<tr>
<td>18</td>
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<td>c.8061del</td>
<td>Clumsy movements Difficulties with speech articulation</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>Infancy</td>
<td>Dysarthria Dysphonia Swallowing and chewing difficulties</td>
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<td>28</td>
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<td>Bilateral LL Toe walking Severe speech delay</td>
<td>2</td>
<td>3</td>
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<td>Anarthria Jaw opening dystonia Tongue protrusion Swallowing difficulties PEG 8y L torticollis R laterocollis</td>
<td>L-dopa trial – no benefit THP and TBZ – reduced tongue protrusion</td>
<td>Inserted age 27y Improvement of jaw opening dystonia and tongue protrusion</td>
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<td>c.3528+2T&gt;A</td>
<td>LLL Gait disturbance L foot dragging Clumsiness</td>
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<td>8</td>
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<td>Severe dysarthria Dysphonia L Torticollis</td>
<td>L-dopa trial – no benefit TBZ, THP, SUL – no benefit</td>
<td>Inserted age 32y – no benefit. Electrode replaced in 2009 with sustained improvement in foot posture but only transient benefit to cervical, UL and LL dystonia.</td>
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<tr>
<td>21</td>
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<td>12</td>
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<td>Dysarthria Dysphonia Swallowing difficulties</td>
<td>L-dopa trial – no benefit THP – not tolerated</td>
<td>Inserted age 15y Sustained clinical benefit 3y post-DBS, improved dystonia and independent walking</td>
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<td>22</td>
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<td>Right foot posturing Abnormal gait</td>
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<td>8</td>
<td>13</td>
<td>5-6</td>
<td>Dysarthria Dysphonia Swallowing difficulties Torticollis</td>
<td>L-dopa trial – no benefit BLF – no benefit THP – low dose, mild benefit BTX neck – reduction in pain but no functional</td>
<td>Inserted age 20y Very good clinical response 9m post DBS with improved dystonia and independent walking</td>
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<td>Bilateral LL involvement (y)</td>
<td>Bilateral UL involvement (y)</td>
<td>Onset of cranial, cervical, laryngeal dystonia (y)</td>
<td>Symptoms of cranial, cervical, laryngeal dystonia</td>
<td>Trial of medication and clinical response</td>
<td>Deep brain stimulation (DBS)</td>
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<tr>
<td>23</td>
<td>8</td>
<td>M</td>
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<td>3</td>
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<td>6</td>
<td>6.5</td>
<td>Dysarthria, Torticollis</td>
<td>L-dopa trial – no benefit CLZ, THP, IT BLF – some benefit</td>
<td>Inserted age 7y with considerable benefit</td>
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<tr>
<td>24</td>
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<td>c.5284C&gt;T p.Arg1762Cys</td>
<td>LLL Tiptoe walking and in-turning of L foot</td>
<td>6</td>
<td>6</td>
<td>7</td>
<td>7</td>
<td>Dysarthria, Anarthria from 14-15y Reduced tongue movements Swallowing preserved</td>
<td>L-dopa trial – no benefit THP- no benefit</td>
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<td>25</td>
<td>19</td>
<td>F</td>
<td>c.5342T&gt;C p.Leu1781Pro</td>
<td>RLL Right foot posturing Gait disturbance</td>
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<td>12</td>
<td>13</td>
<td>10</td>
<td>Dysarthria, Dysphonia Swallowing difficulties Torticollis</td>
<td>L-dopa trial – no benefit LVT – mild benefit</td>
<td>Inserted age 19y Very good clinical response 4m post-DBS with improved dystonia(3)</td>
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<tr>
<td>26a</td>
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<td>c.7549C&gt;T p.Arg2517Trp</td>
<td>Delayed speech Delayed motor development</td>
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<td>-</td>
<td>-</td>
<td>8</td>
<td>Severe paroxysmal retrocollis and jaw dystonia</td>
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<td>Bilateral UL UL posturing Torticollis Inability to walk long distances and run</td>
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<td>27</td>
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<td>F</td>
<td>c.8021T&gt;C p.Ile2674Thr</td>
<td>RUL Posturing, tremor Difficulty handwriting Myoclonic jerks</td>
<td>9</td>
<td>11-13</td>
<td>10</td>
<td>9-10</td>
<td>Dysphonia</td>
<td>L-dopa trial – no benefit THP – no benefit LVT – no benefit CBZ – initial benefit, not sustained CLZ – not tolerated</td>
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BLF: baclofen; BTX: botulinum toxin; CLZ: clonazepam; GBP: gabapentin; IT: intrathecal; LL: lower limbs; LLL: left lower limb; LVT: levetiracetam; m: months; NGF: nasogastric feeding; Pat: patient; PEG: percutaneous endoscopic gastrostomy; RLL: right lower limb; RUL: right upper limb; SUL: sulpiride; UL: upper limbs; TBZ: tetrabenazine; THP: trihexyphenidyl; y: years

(1) based on NCBI Reference Sequence: NM_014727.2
(2) onset shortly after being fitted with orthodontic braces
(3) had undergone 2 posterior cranial fossa explorations and palatal surgery before DBS
<table>
<thead>
<tr>
<th>Patient</th>
<th>KMT2B mutation</th>
<th>Number of genes in microdeletion</th>
<th>Intellectual disability</th>
<th>Dysmorphic features</th>
<th>Additional neurological features</th>
<th>Psychiatric features</th>
<th>Abnormal skin features</th>
<th>Other systemic manifestations</th>
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<td>Deletion: Chr19: 35,608,666-36,233,508</td>
<td>38</td>
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<tr>
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<td>Cutis aplasia&lt;sup&gt;(1)&lt;/sup&gt;</td>
<td>Retinal dystrophy</td>
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<td>Deletion: Chr19: 36,191,100-36,376,860</td>
<td>14</td>
<td>V mild - subtle memory problems</td>
<td>Elongated face Broad nasal bridge Bulbous nasal tip</td>
<td>Not reported</td>
<td>Prone to anxiety&lt;sup&gt;(2)&lt;/sup&gt;</td>
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<td>Not reported</td>
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<tr>
<td>5</td>
<td>Deletion: Chr19: 31,725,360-36,229,548</td>
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<td>Moderate</td>
<td>Sparse hair Blepharophimosis Absent eyelashes of lower eyelids Low set, posteriorly rotated ears Epicantathic folds Narrow nasal bridge, ridge and point Largely bifid tongue Microgathia Teeth overcrowding Finger contractures 5&lt;sup&gt;th&lt;/sup&gt; finger clinodactyly Toe over-riding Dysplastic toenails</td>
<td>Microcephaly</td>
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<td>Absence seizures</td>
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<td>Not reported</td>
<td>Absent right testis</td>
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<td>Mild</td>
<td>5&lt;sup&gt;th&lt;/sup&gt; finger clinodactyly</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Ectodermal dysplasia</td>
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<tr>
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<td>Not reported</td>
<td>Cleft palate</td>
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<td>Strabismus</td>
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<td>Short stature Bronchiectasis</td>
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<td>Dysmorphic features</td>
<td>Additional neurological features</td>
<td>Psychiatric features</td>
<td>Abnormal skin features</td>
<td>Other systemic manifestations</td>
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<td>11</td>
<td>c.399_400insT</td>
<td>p.Pro134Serfs*24</td>
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<td>c.1690C&gt;T</td>
<td>p.Arg564*</td>
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<td>Bulbous nasal tip, short nasal root, hypertelorism, large mouth with full lower lip</td>
<td>Epilepsy</td>
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<td>p.Glu1009Glyfs*9</td>
<td>V mild - difficulties with attention</td>
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<td>Not reported</td>
<td>Not reported</td>
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<td>p.Ile2694Serfs*44</td>
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<td>Delay in saccade initiation and hypometric vertical saccades</td>
<td>ADHD(+) with no response to Ritalin</td>
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<td>Spasticity in lower limbs from 6y</td>
<td>Not reported</td>
<td>Ichthyotic skin lesions with criss-cross pattern under the feet and at knees, broad scarring after operation</td>
<td>Episodic vomiting</td>
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</table>

(1) Supplementary Figure 3c  
(2) Identified on formal psychology review  
(3) Diagnosed by psychiatrist and under regular psychiatry review  
(4) Under regular review with psychiatrist (ICD-10-CM F06.30; ICD-10-CM F42)  
ADHD: attention deficit hyperactivity disorder; GFR: glomerular filtration rate; V: very; y: years