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This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Rots, D., Choufani, S., Faundes, V., Dingemans, A.J.M., Joss, S., Foulds, N., et al. (2024). Pathogenic variants in KMT2C result in a neurodevelopmental disorder distinct from Kleefstra and Kabuki syndromes. *AMERICAN JOURNAL OF HUMAN GENETICS*, 111(8), 1626-1642 [10.1016/j.ajhg.2024.06.009].

Availability:

This version is available at: <https://hdl.handle.net/11585/981977> since: 2024-09-09

Published:

DOI: <http://doi.org/10.1016/j.ajhg.2024.06.009>

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Pathogenic variants in *KMT2C* result in a neurodevelopmental disorder distinct from Kleefstra and Kabuki syndromes

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30 **ABSTRACT**

31 Trithorax-related H3K4 methyltransferases, KMT2C and KMT2D, are critical epigenetic
32 modifiers. Haploinsufficiency of *KMT2C* was only recently recognised as a cause of
33 neurodevelopmental disorder (NDD), so the clinical and molecular spectrum of the *KMT2C*-
34 related (NDD) (now designated as Kleefstra syndrome 2) are largely unknown. We ascertained
35 98 individuals with rare *KMT2C* variants, including 75 with protein truncating variants (PTVs).
36 Notably, ~15% *KMT2C* PTVs were inherited. Although the most highly expressed *KMT2C*
37 transcript consists of only the last four exons, pathogenic PTVs were found in almost all the
38 exons of this large gene. We found that *KMT2C* variant interpretation can be challenging due to
39 segmental duplications and clonal hematopoiesis induced artefacts. Using samples from 27
40 affected individuals, divided into discovery and validation cohorts, we generated a moderate
41 strength disorder-specific *KMT2C* DNAm signature and demonstrate its utility in classifying
42 non-truncating variants. Based on 81 individuals with pathogenic/likely pathogenic variants, we
43 demonstrate that the *KMT2C*-related NDD is characterized by developmental delay, intellectual
44 disability, behavioral and psychiatric problems, hypotonia, seizures, short stature, and other
45 comorbidities. Using facial module of PhenoScore on photographs of 34 affected individuals,
46 we show that the *KMT2C*-related facial gestalt is significantly different from general
47 neurodevelopmental disorder population. Finally, using PhenoScore and DNAm signatures, we
48 demonstrate that the *KMT2C*-related NDD is clinically and epigenetically distinct from Kleefstra
49 and Kabuki syndromes. Overall, we define the clinical features, the molecular spectrum, and
50 the DNAm signature of the *KMT2C*-related NDD and prove them as distinct from Kleefstra and
51 Kabuki syndromes highlighting the need to rename this condition.

52 INTRODUCTION

53 Mendelian disorders of epigenetic machinery are amongst the most common forms of
54 neurodevelopmental disorders (NDDs)¹. Humans possess six histone-3 lysine-4 (H3K4)
55 methyltransferases that are divided into three sub-groups, including the trithorax-related
56 subgroup which consists of KMT2C and KMT2D. These proteins are important components of
57 the epigenetic machinery and are involved in spatiotemporal gene expression regulation^{2,3}.
58 KMT2C or KMT2D, together with WDR5, RBBP5, ASH2L, DPY30 (i.e., WRAD subunit),
59 KDM6A, and other proteins, form the COMPASS complex² that mono- (H3K4me1)⁴ and
60 trimethylates H3K4 (H3K4me3)⁵ at active chromatin sites of gene enhancers and promoters,
61 respectively.

62 Heterozygous loss-of-function *KMT2D* (MIM #602113) variants were identified to cause
63 Kabuki syndrome type 1 (MIM #147920) in 2010⁶. However, the consequences of germline
64 *KMT2C* (MIM #606833) variants in humans have been identified more recently. In a phenotypic-
65 led study we identified *de novo* loss-of-function *KMT2C* variants in individuals with clinical
66 characteristics overlapping Kleefstra syndrome^{3,7}. Kleefstra syndrome (MIM #610253) is
67 caused by haploinsufficiency of euchromatin histone methyltransferase 1 (*EHMT1*; MIM
68 #610253) and is characterized by intellectual disability (ID), autism spectrum disorder (ASD),
69 characteristic facial dysmorphisms, and other variable clinical features⁸. In an independent
70 genotype-led study, we studied variants in histone lysine methyltransferases (KMTs) and
71 demethylases (KDMs) in the Deciphering Developmental Disorders (DDD) cohort and
72 discovered that *KMT2C* loss-of-function variants result in a neurodevelopmental phenotype
73 with occasional physical anomalies⁹. However, the clinical and molecular spectrum of the
74 *KMT2C*-related disorder is largely unknown, as only a few such individuals have been reported.
75 Based on our experience, we hypothesized that the *KMT2C*-related NDD is a unique entity that
76 is clinically and molecularly different from Kleefstra and Kabuki syndromes.

77 In this study, we provide comprehensive clinical, molecular and DNA methylation (DNAm)
78 data for the *KMT2C*-related NDD based on a large previously unreported cohort of individuals,
79 as well as demonstrate that this disorder is different from the molecularly-related Kleefstra and
80 Kabuki type 1 syndromes.

81 **MATERIALS AND METHODS**

82 **Cohort recruitment**

83 This study was approved by the institutional review boards of the South Manchester NHS REC,
84 Radboudumc and the Hospital for Sick Children (Research Ethics Committee Approvals
85 11/H1003/3/AM02, 2011/188, and #1000038847, respectively). Individuals with rare
86 heterozygous pathogenic (P) or likely pathogenic (LP) variants or variants of uncertain
87 significance (VUSs) in *KMT2C* were identified in clinical diagnostic settings (using standard
88 chromosomal microarray, exome, or genome sequencing¹⁰) or from large NDD research cohorts
89 (the Deciphering Developmental Disorders (DDD) study¹¹, 100,000 Genomes project¹², Simons
90 Simplex collection (SSC)¹³, and MSSNG¹⁴). Individuals with rare reported single nucleotide
91 variants (SNVs), indels, and copy number variants (CNVs) in the *KMT2C* gene were included.
92 However, CNVs were limited only to those with deletions <1Mb which did not affect other
93 predicted haploinsufficient genes. Individuals with additional pathogenic variants in other genes
94 were also included in the study but were excluded from the clinical feature frequency analysis.
95 All included individuals or their caregivers/legal representatives consented to participate in this
96 research. Genetic and clinical data from individuals were collected via a customized proforma.

97
98 **Variant analysis**

99 The variants were annotated using the *KMT2C* MANE select¹⁵ transcript
100 (GenBank:NM_170606.3, GRCh37). All identified variants were re-classified according to the
101 ACMG guidelines 2015¹⁶ based on the clinical and molecular evidence obtained during this
102 study. The variants and their interpretations were submitted to the ClinVar database (ClinVar
103 accession numbers: SCV005044911-SCV005044983 and SCV005044991-SCV005045000).
104 Only individuals with LP/P *KMT2C* variants without known pathogenic variants in other genes
105 were included for further clinical feature analyses.

106
107 **DNA methylation analysis**

108 Sample processing, DNA methylation (DNAm) signature derivation and analysis were
109 performed similarly as described before¹⁷⁻¹⁹. Briefly, DNA samples underwent bisulfite
110 conversion using the EpiTect Bisulfite Kit (EpiTect PLUS Bisulfite Kit, QIAGEN) following
111 the manufacturer's protocol. The converted DNAs were then analyzed on the Illumina Infinium
112 Human Methylation EPIC V1 BeadChip (with ~850,000 CpG sites) at The Center for Applied
113 Genomics (TCAG), Hospital for Sick Children Research Institute, Toronto, Ontario, Canada.
114 The affected individuals' and controls' samples were randomly positioned on the arrays.

115 DNAm analysis was performed using the minfi R package²⁰. Briefly, the minfi package was
116 used for data preprocessing, quality control, normalization, transformation to β values and blood
117 cell composition estimation by Housman's method. Probes with a detection p-value >0.05 in
118 $>25\%$ of the samples (911 probes), probes located near common polymorphic variants with
119 minor allele frequencies $>1\%$ (166,596 probes), non-specific probes (35,613 probes), probes
120 with raw β value equal to 0 or 1 in $>25\%$ of samples (240 probes), non-CpG probes (2,377
121 probes), and chrX and chrY probes (17,485 probes) were removed from the analysis.
122 Additionally, 705 probes behaving like single nucleotide variants were removed using the
123 MethylToSNP package²¹. In total, 645,199 CpG sites remained for the differential methylation
124 analysis. All samples passed the quality control and were suitable for the analysis.

125 For the signature derivation, the remaining 645,199 CpG sites' β values were transformed to M
126 values, and differentially methylated CpGs were identified by using a linear regression with
127 monocyte count, batch and second principal components as covariates using the limma R
128 package²². Only differentially methylated CpGs, with a 10% methylation difference ($|\Delta\beta|>0.10$)
129 and false discovery rate (FDR)-adjusted p-values <0.05 were selected for the analysis. To
130 remove false positive sites, we further excluded 6 CpGs with methylation β values in cases and
131 controls that followed a SNP-like pattern. This resulted in a DNAm signature of 51 CpGs with
132 $|\Delta\beta|>0.10$. The results were visualized through principal component analysis (PCA) and
133 heatmap plots using the Qlucore Omics software.

134

135 **Machine learning models and classifications**

136 For the sample classification, we have developed a machine learning model – support vector
137 machine (SVM) with linear kernel using *caret* R package²³ as described before¹⁷. The model
138 was trained on the signature's CpG sites for the discovery cohorts (16 *KMT2C*-related NDD and
139 50 control samples). Because infants have noticeably different blood DNAm, 3 infant control
140 samples were added to the controls during the model training, to increase model's specificity
141 across different ages. Receiver operating characteristic analysis was used to select the optimal
142 model using the largest value. The SVM model was set to the “probability” mode, so the model
143 generated scores ranging between 0 and 1, where scores <0.25 were interpreted as “negative”,
144 >0.5 as “positive”, and 0.25-0.5 as “intermediate”.

145 The model's sensitivity and specificity were evaluated using the *KMT2C*-related NDD and
146 control validation samples, as well as Kabuki type 1 and Kleefstra syndromes samples. Finally,
147 the model was used to classify the 22 testing samples with *KMT2C* VUSs.

148 To evaluate the overlap between *KMT2C*-related NDD DNAm changes with DNAm changes

149 for other disorders of the epigenetic machinery, we analyzed all *KMT2C* samples on 17 other
150 available DNAm signatures deployed on EpigenCentral portal²⁴, as described before²⁵. To
151 evaluate specificity of the *KMT2C*-related NDD DNAm signature, we have utilized six
152 molecularly-confirmed Kabuki syndrome type 1 and six Kleefstra syndrome individuals, as well
153 as 165 healthy controls.

154

155 **Protein 3D structure analysis**

156 *KMT2C* is a large (4911 amino acids) multidomain protein, so there is no solved protein
157 structure of the whole *KMT2C* protein available. Therefore, for the analysis of possible
158 missense variant effects, we have used: 1) solved structures of HMG box (PDB:2YUK),
159 extended PHD6 (PDB:6MLC²⁶), and SET (PDB:6KIW²⁷, 5F6K, 5F59²⁸) domains; 2) homology
160 models for FYR (based on PDB:2WZO²⁹), and extended PHD2 (based on PDB:4NN2³⁰,
161 6U04³¹, 7MJU³², 7D8A³³) domains; 3) *ab initio* protein models for all missense positions. The
162 homology modeling, analysis, and visualization were performed using YASARA Structure
163 software³⁴. The *ab initio* protein models were generated by AlphaFold2³⁵ (multiple modeled
164 overlapping protein fragments of maximal size 1500 amino acids) and downloaded from the
165 AlphaFold Protein Structure Database³⁶. The predicted effect was assessed by at least two
166 different protein structures/models, where available (e.g., solved structure and *ab initio*; two
167 different *ab initio* structures).

168

169 **Clinical features analyses**

170 Consent acquisition, as well as clinical and molecular description of the recruited individuals
171 were provided by their physicians using a standardized proforma (**Table S1**). However, for
172 statistical analyses, only individuals with LP/P *KMT2C* variants and without confirmed
173 additional pathogenic variants in other genes were considered. For any feature, we excluded
174 individuals with “UNKNOWN” coding as previously described³⁷. WHO growth standards per
175 age and sex groups were used to evaluate the growth. Absolute and relative frequencies
176 (expressed as n[%]) were used for describing categorical variables, whereas medians (m) and
177 interquartile ranges (IQR) were used for describing continuous variables. Chi-square/Fisher’s
178 exact and Mann–Whitney tests were applied to study categorical and continuous variables,
179 respectively, and their associations with sex (females vs. males), variant group (protein-altering
180 [PAV] vs. protein-truncating variants [PTV]), and inheritance (inherited vs. *de novo* variants),
181 respectively. Two-tailed, Bonferroni-adjusted p-value <0.05 was considered significant for all
182 statistical analyses, which were carried out using the IBM SPSS Version 29 software.

183

184 **Facial photo analysis**

185 To identify whether *KMT2C*-related NDD has a facial gestalt, facial 2D photos from 29 out of
186 the 34 individuals with a LP/P *KMT2C* variant were compared against 29 sex-, age- and
187 ethnicity- matched individuals with NDDs as controls (sampling different controls 5 times) by
188 using PhenoScore as described before³⁸. Not all 34 individuals included in this study could be
189 included in the PhenoScore analyses, because a matched control was not found for five
190 individuals.

191 Briefly, the facial module of PhenoScore utilizes a state-of-the-art convolutional neural network
192 used in facial recognition (QMagFace³⁹) that recognizes facial features and allows for
193 objectively evaluating and statistically comparing different NDD facial gestalts.

194 Similarly, by using PhenoScore, the photos from these *KMT2C*-related NDD individuals were
195 compared against the sex, age, and ethnicity matched individuals with either *EHMT1* or *KMT2D*
196 pathogenic variants to objectively evaluate whether the *KMT2C*-related NDD facial gestalt
197 significantly differs from those with Kleefstra and Kabuki type 1 syndromes, respectively.

198 RESULTS

199 ***KMT2C*-related NDD cohort curation and identification of variant spectrum**

200 To systematically analyze the genetic and clinical spectrum of the *KMT2C*-related NDD, we
201 ascertained 98 individuals with rare reported *KMT2C* variants, irrespective of their phenotypes
202 through research databases, international collaborations, and previously published
203 individuals^{3,7,9}.

204 75 out of 98 individuals from 67 families had 61 distinct heterozygous *KMT2C* variants
205 predicted to be protein truncating (PTVs) (20 nonsense, 28 frameshift, 5 splice site and 8 large
206 deletions). These PTVs were classified as pathogenic (P) or likely pathogenic (LP) (**Figure 1**),
207 because loss of function is the currently accepted disease mechanism^{3,7,9} and the gene is
208 intolerant to loss-of-function variants in population (pLI=1; o/e = 0.08 (0.06 - 0.12))⁴⁰. Of note,
209 52 of these P/LP variants were *de novo*, 11 were inherited (6 maternal and 5 paternal), and in
210 12 individuals the inheritance was unknown. Importantly, *KMT2C* is a large gene consisting of
211 59 exons (NM_170606.3). Ensembl documents 70 transcripts for *KMT2C*, 31 of which are
212 annotated as protein coding: ranging in amino acid length from 4968 (ENST00000682283.1) to
213 83 (ENST00000684278.1). The MANE select transcript (canonical isoform
214 NM_170606.3/ENST00000262189.11) encodes a protein of 4911 amino acids with mean
215 expression across all adult tissues in the GTEx database of only 0.83 transcripts per million
216 (TPM), while the most highly expressed transcript (5.08 TPM) is ENST00000485655.2
217 consisting of only the last four exons of the gene⁴¹. This, however, might be related to the 3'
218 bias of the short-read GTEx data⁴². In our cohort, we found the PTVs are distributed throughout
219 the gene, with the most 5' and the most distal PTVs being located in the 3rd and 57th *KMT2C*
220 exons, suggesting that in spite of apparently relatively low expression of the canonical (MANE)
221 transcript in adult tissues, PTVs across the gene are likely to be pathogenic. We also did not
222 observe specific clustering of PTVs in the gnomAD database (**Figure S1**), but, based on the
223 skewed variant allele frequency, we noted that some of the observed PTVs are either artifacts
224 due to segmental duplications (e.g., c.2710C>T p.(Arg904Ter) and c.1173C>A p.(Cys391Ter)),
225 or somatic variants (e.g., c.6415C>T p.(Arg2139Ter)), as *KMT2C* is known clonal
226 hematopoiesis driver gene⁴³.

227 We observed a relatively high proportion of individuals with pathogenic CNVs in our cohort
228 (n=11, 14%; 5/11 are *de novo*). We, therefore, examined the genomic architecture and found
229 that the *KMT2C* contains several segmental duplications with high homology (>98%) to
230 sequences elsewhere in the genome (**Figure 1B**) making this region a “hot-spot” for structural
231 rearrangements⁴⁴. We examined the Database for Genomic Variants (DGV)⁴⁵ and gnomAD⁴⁰

232 for structural variants affecting this coding part of the *KMT2C* in the control population in the
233 UCSC genome browser⁴⁶ and identified in total 26 partial or intragenic gains and only a single
234 deletion. This suggests that the partial/intragenic gains are unlikely to be pathogenic (if not
235 leading to frameshift). We, however, cannot exclude frequent occurrence of artifacts due to the
236 limitations of current technologies in the regions affected by segmental duplications.
237 In contrast, non-PTVs have not been classified as pathogenic, so 23 out of 98 individuals from
238 22 families had 22 VUSs (18 missense and 2 splice variants and 2 large duplications). Of those,
239 10 were *de novo*, 6 were inherited, and 6 were of unknown inheritance.
240 Collectively, these data show that P/LP truncating *KMT2C* variants are spread across the gene,
241 affecting only MANE or multiple transcripts, and that they can be inherited in some cases.
242 Additionally, clinical classification of *KMT2C* non-truncating variants is currently challenging.

243 **Protein truncating *KMT2C* variants result in a DNAm signature in peripheral blood**

244 We and others have previously shown that pathogenic variants in genes encoding components
245 of epigenetic regulators are associated with genome-wide DNAm changes⁴⁷ from which disease-
246 associated DNAm signatures can be derived. Therefore, we performed genome-wide
247 methylation screening on peripheral blood-derived DNA from 16 *KMT2C*-related NDD
248 individuals (discovery cohort) with pathogenic *KMT2C* PTVs and 50 controls using Illumina
249 methylation Epic arrays. We identified 51 significant differentially methylated CpG sites at
250 $|\Delta\beta|>0.10$, and FDR-corrected $p<0.05$ representing the DNAm signature of the *KMT2C*-related
251 NDD (for simplicity, further called *KMT2C* DNAm signature) (**Table S3**). Most of the
252 signature's CpGs were hypomethylated (42/51, 82%). Interestingly, 4/51 (8%) of the signature's
253 CpGs sites mapped to two CpG islands of the WT1 gene (MIM #607102), which were mostly
254 hypomethylated (**Figure 2C**).

255 Based on the derived *KMT2C* DNAm signature, we were able to discriminate the discovery
256 *KMT2C* cases from healthy controls based on PCA and heatmap plots (**Figures 2A and 2B**,
257 respectively). Next, we trained an SVM model based on the discovery cohort and tested the
258 sensitivity and specificity of the *KMT2C* DNAm signature using 165 controls without known
259 developmental disorders, and 11 additional validation cases with P/LP PTV *KMT2C* variants.
260 On this SVM model, all controls were classified as negative (SVM values <0.25 , 100%
261 specificity), and 9/11 validation cases were classified as positive (SVM values >0.5 , 82%
262 sensitivity) (**Figure 2D**). Two validation cases were classified negatively: individual #48
263 presented with typical clinical features and has a *de novo* pathogenic frameshift variant
264 c.13107_13108dup p.(Thr4370Argfs*11); individual #52 is mildly affected with multiple
265 affected children and has a small likely pathogenic *KMT2C* exon 36 and 37 deletion resulting
266 in frameshift. However, we cannot exclude that these cases are high-level mosaics in blood.

267 Next, to test whether *KMT2C*-related NDD affected individuals share the DNAm signature with
268 other conditions, we analyzed all available samples with a LP/P *KMT2C* variant on 17 available
269 DNAm signatures deployed at EpigenCentral²⁴. All *KMT2C*-related NDD samples were
270 classified negatively on all signatures (**Figure 2E**).

271 These results show that significant changes in methylation patterns of DNA derived from the
272 peripheral blood of individuals with P/LP *KMT2C* variants exist, and that these changes can be
273 used to generate a moderate-effect DNAm signature that does not overlap with the other known
274 disorder-specific DNAm signatures.

275

276

277 **A DNAm signature can be used to re-interpret *KMT2C* VUS**

278 Next, we explored if DNAm signature could enable classification of the *KMT2C* VUSs by
279 testing DNA samples from 22/23 individuals (20 unrelated and one proband-father pair) with
280 21 distinct *KMT2C* VUSs on the *KMT2C* DNAm signature-derived SVM model (one individual
281 with VUS c.14501T>C p.(Val4834Ala) was not available for testing). The SVM model resulted
282 in classification of 5/21 VUS as positive for the *KMT2C* DNAm signature. One out of 21 VUS
283 was classified as intermediate (SVM score 0.25-0.5), and the rest were classified as negative on
284 the *KMT2C* DNAm signature. These results supported re-classification of 6/21 variants as LP
285 and 13/21 variants as benign (described below), while 2/21 variants remained in the VUS
286 category (one additional VUS was unavailable for testing).

287 Next, we examined the basis of pathogenicity of the PAVs in this cohort. The details for each
288 variant classification and the criteria applied utilizing various evidence are described in **Table**
289 **S4**. All six variants reclassified as LP/P were classified as positive (or intermediate) on the
290 DNAm signature. Out of the six VUSs reclassified as LP/P, two (*de novo*) variants were
291 (re)interpreted as PTVs: duplication of exons 3-38 which is predicted to result in frameshift, if
292 present in tandem, so based on the DNAm signature classification we concluded that the
293 duplication most likely is indeed in tandem; one variant was initially reported from trio exome
294 sequencing as *de novo* missense c.3499G>A p.(Asp1167Asn), located outside of known
295 domains, but the signature analysis aided the reanalysis of the variant which was later confirmed
296 by Sanger sequencing as c.3499+1G>T p.?. The remaining four LP PAVs (c.3233G>A
297 p.(Cys1078Tyr), c.13229A>G p.(Asp4410Gly), c.13783C>T p.(Arg4595Cys) and c.14335C>G
298 p.(Arg4779Gly)) are absent from gnomAD, predicted as pathogenic by *in silico* tools (REVEL⁴⁸
299 > 0.65 and AlphaMissense⁴⁹ > 0.56), and all are located in well-known functional domains of
300 the protein (PHD6, ePHD2, FYRN and SET domains, respectively). Moreover, these variants
301 are predicted to disrupt the 3D structure of the respective domains (**Figure S3**), except for the
302 c.13229A>G p.(Asp4410Gly) for which no reliable 3D structure or model was available.

303 In contrast, all 11/13 variants classified as benign are missense and are negative on the DNAm
304 signature, the majority are not classified as pathogenic by *in silico* tools and are not predicted to
305 affect protein 3D structure and/or located outside of the known functional domain's (largely in
306 disordered regions) (**Table S4**). Only three benign PAVs occurred *de novo*. Additionally, one
307 splicing variant c.1735+2dup was inherited from father, predicted to result in inframe skipping
308 of exon 12 that encodes a disordered region of the protein and was classified as negative by the
309 DNAm signature, so the variant was classified as benign. Similarly, one benign partial *KMT2C*
310 duplication (1-55 exons) with unknown inheritance was absent from population databases, but

311 similar *KMT2C* 5' partial duplications are common in the general population (**Figure 1B**), and
312 it is also classified as negative on the DNAm signature.

313 Finally, three missense VUSs remained classified as VUS due conflicting or insufficient
314 evidence. One of the missense variants (c.14501T>C p.(Val4834Ala)) is located in the SET
315 domain, is predicted as pathogenic by *in silico* tools (REVEL = 0.8 and AlphaMissense = 0.99)
316 and was predicted to disrupt the domain's 3D structure (**Table S2**), but the sample was not
317 available for DNAm testing. Two missense variants (c.9773A>C p.(His3258Pro) and
318 c.13298C>T p.(Ala4433Val)) were classified negatively on the DNAm signature but had
319 additional significant evidence for pathogenicity: both variants are *de novo*, absent in gnomAD
320 and predicted to be possibly disruptive by *in silico* tools and/or 3D protein analysis (**Table S2,**
321 **S4**).

322 These results show that the *KMT2C* DNAm signature can be used to reclassify most, but not all
323 VUSs in the gene and that PAVs disrupting functional domains are more likely to be P/LP.
324

325 ***KMT2C* loss-of-function variants result in a phenotypically heterogenous NDD syndrome**

326 In total, we identified 81 individuals from 73 families with P/LP *KMT2C* variants. We gathered
327 detailed clinical information on these individuals to understand the clinical spectrum of *KMT2C*-
328 related NDD. Out of 81 individuals, 6 had a second molecular diagnosis due to a pathogenic
329 variant in another NDD-associated gene. To understand the clinical consequences of P/LP
330 *KMT2C* variants, we analyzed the clinical features of 75 individuals with P/LP *KMT2C* variants
331 without any additional known P/LP variants in other NDD genes (**Table 1, S1**).

332 In this cohort, 45.3% (n=34) were females, and the median age at last examination was 11 years
333 (with IQR 5.17;20). The median weight at the last reported time was -1.01 SD (-1.94;-0.21), the
334 median height was -1.56 SD (-2.32;-1.19), and the median head circumference (HC) was -0.44
335 SD (-1.65;0.51). Importantly, short stature ($\leq -2SD$) was reported in ~54% of individuals, and 5
336 of them have received recombinant human growth hormone (rhGH) with an apparently good
337 response. Also, 9/53 (~17%) of individuals have displayed microcephaly ($\leq -2SD$) at least on
338 one available measurement. These findings suggest *KMT2C*-related NDD is mainly an
339 undergrowth condition.

340 Neurodevelopmental problems are the most common features in this cohort. Although gross
341 motor delay was described in 87% of individuals, all individuals older than 3.5 years of age had
342 achieved independent walking. Speech delay was reported in 80% of individuals, and ~15%
343 developed mutism at some point in their lifetime. Developmental regression was reported in 6
344 individuals. ID was reported in ~86% of individuals with half being classified as having
345 moderate or severe ID. Importantly, a proportion of individuals had normal cognitive abilities
346 (~14%; all with *de novo* variants) and some of them were tested due to non-NDD phenotypes
347 (e.g., short stature, seizures). However, we cannot exclude mosaicism in these cases.
348 Furthermore, behavioral, and psychiatric problems were reported in most individuals: features
349 or diagnosis of ASD or ADHD were reported in ~79% and 61% of individuals, respectively;
350 aggressive (both hetero- and self-) behavior was reported in 18% and obsessive/compulsive
351 behavior – in ~15%.

352 Abnormalities affecting the central nervous system (~37%; e.g., ventriculomegaly, white-matter
353 anomalies, syringomyelia) and the cardiovascular system (~24%; e.g., septal defects, valvular
354 anomalies) were frequent. Refractive errors (33%) and hearing loss (~29%) were the most
355 frequent sensory system problems. Recurrent infections (~26%), central/obstructive sleep apnea
356 (~17%), seizures (~15%), kyphosis and/or scoliosis (~16%), and constipation (~16%) were
357 some of the other most significant and frequently encountered medical issues (**Table 1**). Feeding

358 difficulty was the most frequent neonatal finding (49%), followed by neonatal hypotonia
359 (~29%).

360 Next, we compared the frequencies of clinical features between male and female individuals,
361 type of variants (PAVs vs. PTVs), and their inheritance (inherited vs. *de novo*). None of the
362 features was significant after correction for multiple testing ($p_{adj}>0.05$).

363 Craniofacial dysmorphisms were described in 90% of individuals with P/LP *KMT2C* variants
364 in our cohort. Photographs of 34 individuals were available in this cohort, with 13 consenting
365 for publishing (**Figure 3**). We used these photographs to analyze the facial features of
366 individuals with *KMT2C*-related NDD. The most frequent findings were frontal
367 bossing/prominent-broad forehead, thick/prominent eyebrows, synophrys, down-slanted
368 palpebral fissures, deep-set eyes, epicanthus, hypertelorism, midface retrusion, anteverted nares,
369 thin vermilion of the upper lip, micrognathia, low-set ears, thick ear helices and posteriorly
370 rotated ears. Although all individuals had craniofacial dysmorphism, overall, we did not identify
371 a clearly recognizable *KMT2C* facial gestalt.

372 Next, to objectively evaluate whether the *KMT2C*-related NDD has a specific facial gestalt, we
373 used PhenoScore and compared the facial photographs of individuals with the *KMT2C*-related
374 NDD with facial photographs of individuals in a general ID cohort. PhenoScore identified a
375 significantly different *KMT2C*-related facial gestalt (AUC = 0.91, $p<0.001$) (**Figure S2**),
376 indicating that the facial features the 29 individuals with a LP/P *KMT2C* variant for whom an
377 age-, sex- and ethnicity matched NDD control was available are distinguishable from the general
378 NDD population. The PhenoScore identified periorbital and nasal regions as the most different
379 (**Figure S2**), which fit the most common dysmorphic features described above.

380 Collectively, these observations show that *KMT2C*-related NDD is clinically a highly
381 heterogeneous disorder characterized by intellectual disability, short stature, congenital
382 anomalies, recurrent infections and craniofacial dysmorphism.

383

384 ***KMT2C*-related NDD is distinct from Kleefstra and Kabuki syndromes**

385 OMIM designates the *KMT2C*-related NDD as 'Kleefstra syndrome 2' (MIM #617768) which
386 may be interpreted that the *EHMT1* and *KMT2C* related disorders have significant clinical
387 overlap. However, this overlap has not been examined previously. Also, *KMT2C* and *KMT2D*
388 are part of the same gene family, but their overlap has not been examined previously. For this,
389 firstly, we compared the facial features of individuals with *KMT2C*-related NDD to the
390 photographs of 24 and 10 matched individuals with Kleefstra, and Kabuki type 1 syndromes,
391 respectively. PhenoScore revealed significant differences between the *KMT2C*-related NDD
392 facial gestalt and the facial gestalt of Kleefstra syndrome (AUC=1, $p<0.001$) and Kabuki
393 syndrome type 1 (AUC=0.8, $p=0.04$).

394 DNAm signatures can also be used to delineate epigenetically distinct disorders^{50,51}. Pathogenic
395 loss-of-function variants in the *EHMT1*⁵² or *KMT2D*⁶ result in unique DNAm signatures^{51,53}.
396 We analyzed DNA samples from 6 individuals with Kleefstra and 6 individuals with Kabuki 1
397 syndromes on the *KMT2C* DNAm signature. Samples from Kleefstra and Kabuki type 1
398 syndromes were classified negatively by the SVM (**Figure 2D; Table S5**) on the *KMT2C*
399 DNAm signature. Similarly, all tested *KMT2C*-related NDD individuals were classified
400 negatively on the Kleefstra and Kabuki syndrome signatures (**Figure 2E**), highlighting the
401 DNAm differences among the three conditions.

402 These findings confirm that the *KMT2C*-related NDD is clinically and epigenetically distinct
403 from both Kleefstra and Kabuki syndromes.

404 **DISCUSSION**

405 In this study, we have demonstrated that heterozygous loss-of-function *KMT2C* variants result
406 in a disorder that is characterized by variable combinations of developmental delay, intellectual
407 disability, short stature, congenital anomalies, craniofacial dysmorphism, recurrent infections,
408 and a moderate strength DNAm signature. We also demonstrate that the *KMT2C*-related NDD
409 is clinically and epigenetically distinct from (*EHMT1*-related) Kleefstra syndrome type 1 and
410 (*KMT2D*-related) Kabuki type 1 syndromes.

411 Our data show that despite the complexities of the transcripts encoded by *KMT2C*, the PTVs in
412 the gene have similar clinical or epigenetic consequences irrespective of their location in the
413 gene. This observation suggests four possible explanations: 1) the canonical isoform is
414 biologically and disease relevant in spite of its apparent low level of expression in tissues, 2)
415 the developmental expression profile of *KMT2C* may be different from its known adult
416 expression profile, 3) the *KMT2C* transcript expression pattern may not yet be fully known, or
417 4) the regulatory consequences of loss-of-function variants in the longer transcripts may have
418 complex consequences on the transcriptional network. Additionally, we cannot exclude that the
419 low expression of the canonical transcript in GTEx is a technical artifact of the short-read based
420 RNA sequencing, displaying 3' bias and short-isoform preferences⁴². Our analysis also shows
421 that *KMT2C* is located in a hot-spot for both benign and pathogenic structural rearrangements,
422 and, therefore, CNVs in this gene should be interpreted with caution. In our study, we also
423 showed that rare missense variants that affect conserved residues and disrupt the 3D structure
424 of the PHD Zn-fingers, the FYR domain, or the catalytic SET also cause the same disorder.
425 Interestingly, pathogenic Kabuki syndrome Type 1 causing missense variants in *KMT2D*, a
426 paralogue of *KMT2C*, cluster significantly in several similar functional domains⁵⁴. These
427 findings collectively indicate haploinsufficiency and functional haploinsufficiency as likely
428 mechanisms that underlie *KMT2C*-related NDD.

429 We also identify a moderate-effect DNAm signature for *KMT2C*-NDD with current sensitivity
430 of ~82%. To identify a reliable signature, we had to utilize a relatively large sample size cohort
431 (of different ages and sex) as initial signatures derived from smaller cohort sizes lacked
432 sensitivity and specificity. It is possible that variable expressivity is typical not only for the
433 clinical features, but also for the DNAm signature. It has been shown previously that the level
434 of DNAm changes correlate with the age of onset of dystonia in the *KMT2B*-related dystonia⁵⁵
435 (MIM #617284), and the Weaver-syndrome (MIM #277590) signature was found to be weaker
436 in mildly affected individuals with a pathogenic *EZH2* (MIM #601573) variant even within a
437 single family¹⁸. Therefore, it is possible that mildly affected individuals can be classified as

438 intermediate or negative on the DNAm signature despite the presence of pathogenic variants.
439 For example, one mildly affected individual with a pathogenic *KMT2C* variant (exons 36-37
440 deletions) classified as negative by the DNAm signature is the mother of three affected children.
441 This individual is mildly affected which might explain the DNAm classification results.
442 Unfortunately, DNA from the children was not available for testing. However, the second
443 individual with pathogenic variant c.13107_13108dup p.(Thr4370Argfs*11) was also classified
444 negatively despite having typical *KMT2C*-related NDD clinical features. However, we were not
445 able to be exclude mosaicism in either case, which may explain the negative classification
446 results. In the future, larger studies will be required to further elucidate the biological basis of
447 such correlations, which may have prognostic and management implications.

448 A large proportion of the identified *KMT2C* signature's CpG sites (8%) mapped to the *WT1*
449 gene's CpG islands, which are hypomethylated in *KMT2C*-related NDD individuals.
450 Hypomethylation of CpG islands, especially those located in the promoter of a gene, is usually
451 associated with upregulation of gene expression⁵⁶, which was previously shown for *WT1 in*
452 *vitro*⁵⁷. *WT1* is not a known interactor of *KMT2C*, so it is unclear whether the hypomethylation
453 at the *WT1* CpG islands is a primary effect of *KMT2C* disruption or secondary to disrupted
454 interactions of multiple genes. Loss-of-function or dominant-negative *WT1* variants are
455 associated with Wilms tumor (MIM #194070) or Denys-Drash syndrome (MIM #194080),
456 respectively, but to date there is no human phenotype associated with *WT1* overexpression. Until
457 now, individuals with *KMT2C*-related NDD have not displayed any tumors or anomalies of sex
458 development that are reported for the *WT1*-related disorders. Recently, a role for *WT1* in brain
459 and neuron development was also described⁵⁸⁻⁶⁰. Mariotinni et al.⁶¹, proved that *Wt1* is
460 important for synaptic plasticity and learning in mice, and that it functions as a long-term
461 memory suppressor. Additionally, they showed that *Wt1* overexpression causes a reduction of
462 memory retention⁶¹. In another study, Ji et al., showed that *Wt1* brain-specific loss in mice
463 resulted in depressive-like behavior, but the effects of overexpression were not investigated⁶⁰.
464 At this time, the relevance of *WT1* methylation changes to the phenotype of *KMT2C*-related
465 NDD is not known.

466 Several neurodevelopmental phenotypes were present in the majority of the individuals in the
467 study cohort, including developmental delay, intellectual disability (mostly mild, but ranging
468 from borderline to severe), ASD, and ADHD. Additionally, the individuals often have other
469 psychiatric issues, including hetero- or self-aggressive behavior, obsessive-compulsive
470 behavior, and selective mutism. In fact, pathogenic or *de novo* *KMT2C* variants have also been
471 found to be enriched in individuals with schizophrenia⁶², bipolar disorder⁶³ and other psychiatric

472 disorders⁶⁴. This suggests that *KMT2C*-related NDD is associated with multiple, overlapping
473 neurodevelopmental and psychiatric presentations.

474 Individuals with *KMT2C*-related NDD present with a broad spectrum of clinical features, but
475 the expressivity of the features is highly variable among individuals. For example, ~14% of the
476 individuals in the entire cohort did not have intellectual or learning disabilities. The variable
477 expressivity likely explains the relatively large proportion (11/75, ~15%) of inherited cases in
478 this cohort, as most of the parents with pathogenic variants were mildly affected.

479 Individuals with the *KMT2C*-related NDD had common dysmorphic features, including frontal
480 bossing or prominent forehead, downslanted palpebral fissures, deep-set eyes, and
481 hypertelorism, among others. These features are mild and non-specific, and therefore, the facial
482 gestalt was unrecognizable to clinicians, unlike Kleefstra and Kabuki 1 syndromes. However,
483 using PhenoScore³⁸, we were able to show that the *KMT2C*-related NDD had a typical facial
484 gestalt that is significantly different from Kleefstra and Kabuki 1 syndromes.

485 *KMT2C*-related NDD, Kleefstra, and Kabuki 1 syndromes are clearly different conditions
486 despite some clinical and molecular overlap, but it is currently impossible to objectively
487 compare the frequencies of specific features among these conditions head-to-head. Kleefstra
488 and Kabuki syndromes are clinically well-known and well-recognized, and most of the
489 published individuals are diagnosed through a phenotype-first approach^{8,65}. This results in
490 biased results with only the most typical affected individuals being initially described. However,
491 with the recent widespread application of exome or genome sequencing in clinical diagnostics,
492 it is possible to also identify non-specific and/or mildly affected individuals^{66,67}, as well as novel
493 morbid entities within the same gene⁵⁰, including *KMT2D*⁶⁸. *KMT2C*-related NDD is not
494 clinically recognizable, so all individuals were diagnosed through a genotype-first approach and
495 this study represents a clinical expansion and delineation of this condition. Therefore, studies
496 on the phenotype of Kleefstra and Kabuki 1 syndromes diagnosed through a genotype-first
497 approach would be necessary to better understand their clinical and molecular spectrum, as well
498 as the differences among the three conditions.

499 Although *KMT2C*-related NDD, Kleefstra and Kabuki 1 syndrome are caused by
500 haploinsufficiency of methyltransferases, where PTVs are the most frequent type of variant, the
501 proportion of inherited variants is considerably different among them. While Kleefstra and
502 Kabuki 1 syndrome mostly occur by *de novo* loss-of-function variants, with only a handful of
503 inherited cases described so far^{69,unpublished data}, ~15% of *KMT2C*-related NDD individuals
504 inherited their variants from similarly affected parents. Albeit this may be explained by a biased

505 approach to diagnosis, our study also showed that the expressivity of the *KMT2C*-related NDD
506 is highly variable, and individuals with few manifestations can be expected.

507 Several individuals in our study were primarily investigated due to short stature - a feature
508 present in more than a half of the *KMT2C*-related NDD individuals reported in this study. Short
509 stature is also common in Kleefstra and Kabuki 1 syndromes^{8,65}. The causes of short stature in
510 these syndromes are often unknown but likely complex⁶⁵. However, short stature in individuals
511 with monogenic NDDs is rarely investigated in-depth in routine clinical practice as it is usually
512 considered to be part of the syndrome. While it is known that growth hormone deficiency is
513 present in approximately a third of individuals with Kabuki 1 syndrome⁷⁰, this can partially
514 explain the presence of short stature. However, treatment with rhGH has proven to be highly
515 effective and safe⁷¹, and therefore, these individuals should be screened for this deficiency. In
516 this cohort, 40/75 individuals with endocrine anomalies had proven short stature, and 6/40 had
517 received rhGH with apparently good results. This should prompt further studies on the
518 prevalence of GH deficiency and the utility of rhGH therapy also for individuals with *KMT2C*-
519 related NDD and short stature.

520 While various psychiatric disorders are seen in *KMT2C*-related NDD, this is not frequent in
521 Kabuki 1 syndrome⁷². Also, although psychotic disorders such as schizophrenia⁷³ and
522 developmental regression⁷⁴⁻⁷⁶ have been frequently reported in Kleefstra syndrome, nearly all
523 individuals with Kleefstra syndrome manifest intellectual disability that ranges from moderate
524 to severe^{8,76}, which contrasts with our finding that ~15% of individuals with *KMT2C*-related
525 NDD demonstrate no cognitive impairment. Again, albeit this may be explained by a bias in our
526 diagnostic approach, our study shows that the penetrance and severity of this finding is
527 consistently reduced in the *KMT2C*-related NDD.

528 In summary, pathogenic variants in the *KMT2C* gene result in a distinct syndromic
529 neurodevelopmental disorder, which is different from Kleefstra and Kabuki 1 syndromes.
530 Therefore, the OMIM designation of *KMT2C*-related NDD as Kleefstra syndrome 2 should be
531 reconsidered, as it may be misleading to clinicians and patients' families.

532
533

534 **DECLARATION OF INTERESTS**

535 RW is a consultant (equity) for Alamy Health. The rest of the authors declare no competing
536 interests.

537

538 **ACKNOWLEDGEMENTS**

539 Described in the Supplemental information.

540

541 **WEB RESOURCES**

542 <https://epigen.ccm.sickkids.ca>

543

544

545 **DATA AND CODE AVAILABILITY**

546 Recruited individuals' with pathogenic *KMT2C* variants clinical details are provided in the
547 **Table S1**. Recruited individuals' with benign or VUS *KMT2C* variants details are provided in
548 the **Table S2**. The variants and their interpretations were submitted to the ClinVar database
549 (ClinVar accession numbers: SCV005044911-SCV005044983 and SCV005044991-
550 SCV005045000).

551 The *KMT2C* DNAm signature is available in supplementary material (**Table S3**). The raw
552 datasets supporting the current study have not been deposited in a public repository due to
553 institutional ethics restrictions. All software and R packages used in the study are publicly
554 available as described in the methods section. The *KMT2C* methylation classifier will be made
555 available through EpigenCentral at <https://epigen.ccm.sickkids.ca>.

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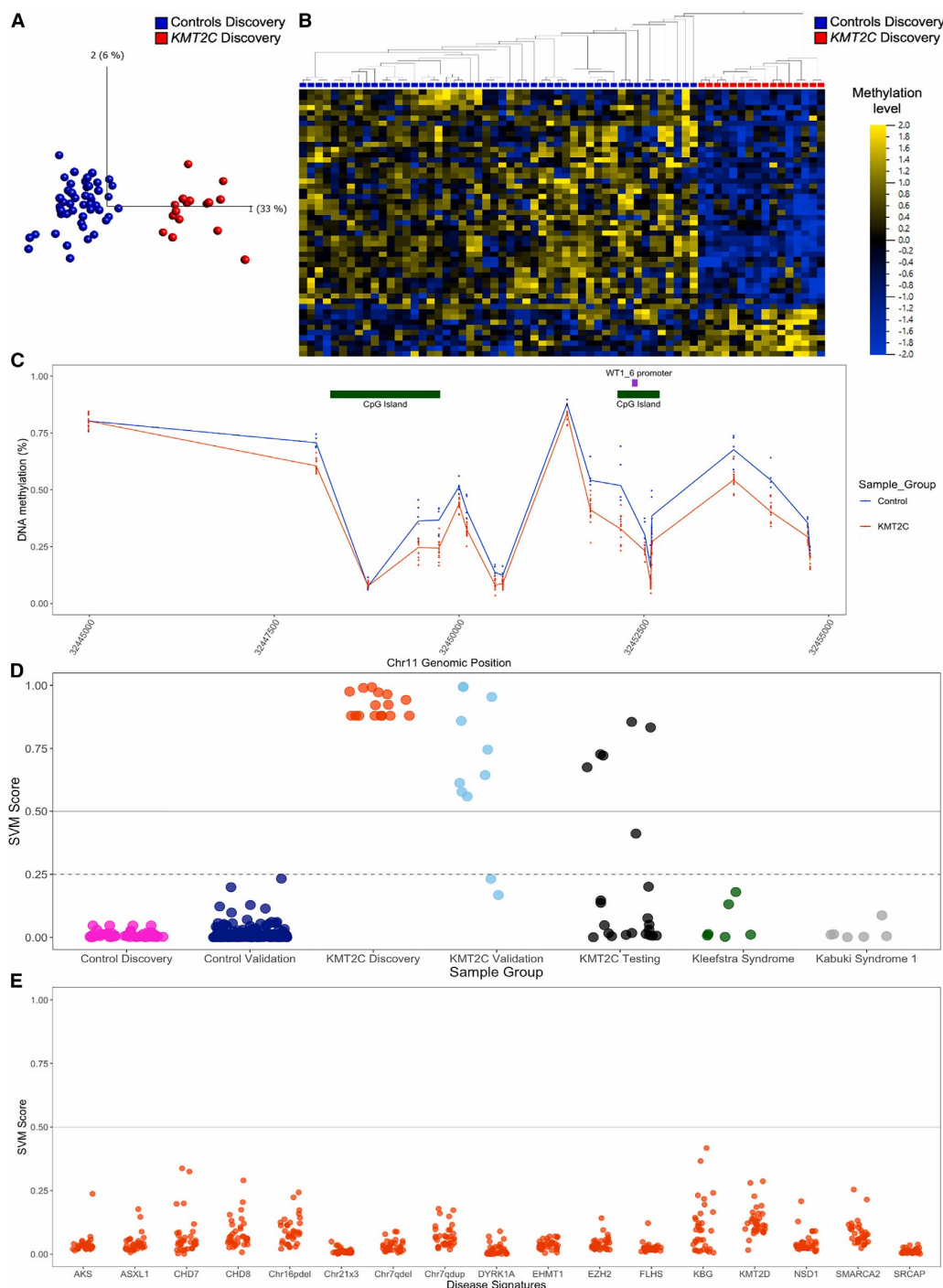
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854 comparison shown on the *KMT2C* genomic region with segmental duplications (grey), as well
 855 as populational deletions and duplications from DGV and gnomAD databases.
 856
 857

858 **Figure 2.** DNAm signature classification results.
 859



860
 861 **A)** PCA plot for the discovery cohort of 16 KMT2C-related NDD cases (red) and 50 controls
 862 (blue). **B)** Heatmap plot with hierarchical clustering for the discovery cohort of 16 KMT2C-
 863 related NDD cases (red) and 50 controls (blue), with hypermethylated sites shown in yellow
 864

865 and hypomethylated ones in blue. **C)** Differentially methylated region which maps to several
 866 *WT1* CpG islands (green) and one of the promoters (purple) (data obtained from the UCSC
 867 genome browser) where each dot shows methylation level at a CpG site in the region for each
 868 discovery cohort sample, and the line depicts the mean methylation for controls (blue) and cases
 869 (red). **D)** SVM model classification results for different groups: 16 discovery *KMT2C*-related
 870 NDD cases (red), 50 controls (magenta); validation *KMT2C*-related NDD cases (light blue)
 871 and their matched controls (dark blue); *KMT2C* VUSs (black); Kleefstra syndrome individuals
 872 (green) and Kabuki syndrome individuals (grey). **E)** *KMT2C*-related NDD classification results
 873 against 17 other DNAm signatures.

874

875 **Figure 3.** Photos of individuals with the *KMT2C*-related NDD.

876



877

878 *y.o.* = years old – the age of the individuals.

879 **Table 1.** Frequencies or distributions of clinical findings of *KMT2C*-related NDD.

Clinical finding ^a	Frequency or distribution
Respiratory anomalies (n [%]) (n/42)	14 (33.3)
Central/obstructive sleep apnea (n [%])	7 (16.7)
Asthma (n [%])	6 (14.3)
Other (n [%])	2 (4.8)
Palate anomalies (n [%]) (n/437)	12 (32.4)
High/narrow palate (n [%])	11 (29.7)
Cleft lip/palate (n [%])	1 (2.7)
Cardiovascular anomalies (n [%]) (n/54)	13 (24.1)
Septal defects (n [%])	6 (11.1)
Conduction disorder (n [%])	6 (11.1)
Valvular anomalies (n [%]) Other (n [%])	4 (7.4)
	3 (5.6)
Genitourinary anomalies (n [%]) (n/439) Uni/bilateral	8 (20.5)
inguinal hernia (n [%])	2 (5.1)
Other (n [%])	7 (17.9)
Dental anomalies (n [%]) (n/470)	13 (18.8)
Dental crowding (n [%])	6 (8.6)
Prominent upper incisors (n [%])	6 (8.6)
Other (n [%])	1 (1.4)

880
881 *a* The number of responders are detailed for every feature, and their frequencies/distributions
882 were calculated according to that number. Individuals with another NDD were excluded.* at
883 any moment documented <-2SD.
884 ADHD=Attention Deficit/Hyperactivity Disorder; ASD=Autism Spectrum Disorder;
885 CNS=Central nervous system; IQR=Interquartile range m=median; SD=Standard deviation
886
887

888 **SUPPLEMENTAL INFORMATION**

889 **Table S1.** Recruited individual with *KMT2C*-related NDD detailed clinical description with
890 variants.

891 **Table S2.** Recruited individual with benign/VUS *KMT2C* variants detailed clinical description
892 with variants and their classification.

893 **Table S3.** *KMT2C* DNAm signature CpG sites.

894 **Table S4.** *KMT2C* VUS classification.

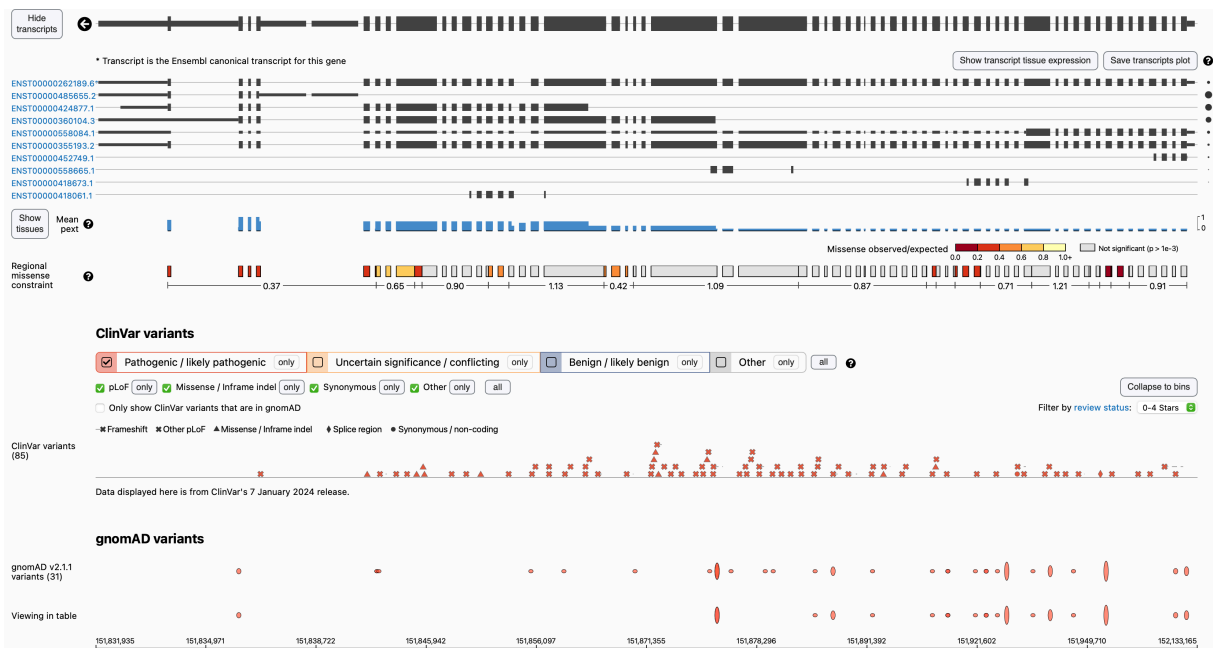
895 *Blue = evidence for pathogenicity; Green = benign evidence; Grey – unknown or uncertain*
896 *significance*

897

898 **Table S5.** Description of Kleefstra and Kabuki syndrome 1 samples used for the testing of the
899 *KMT2C* signature.

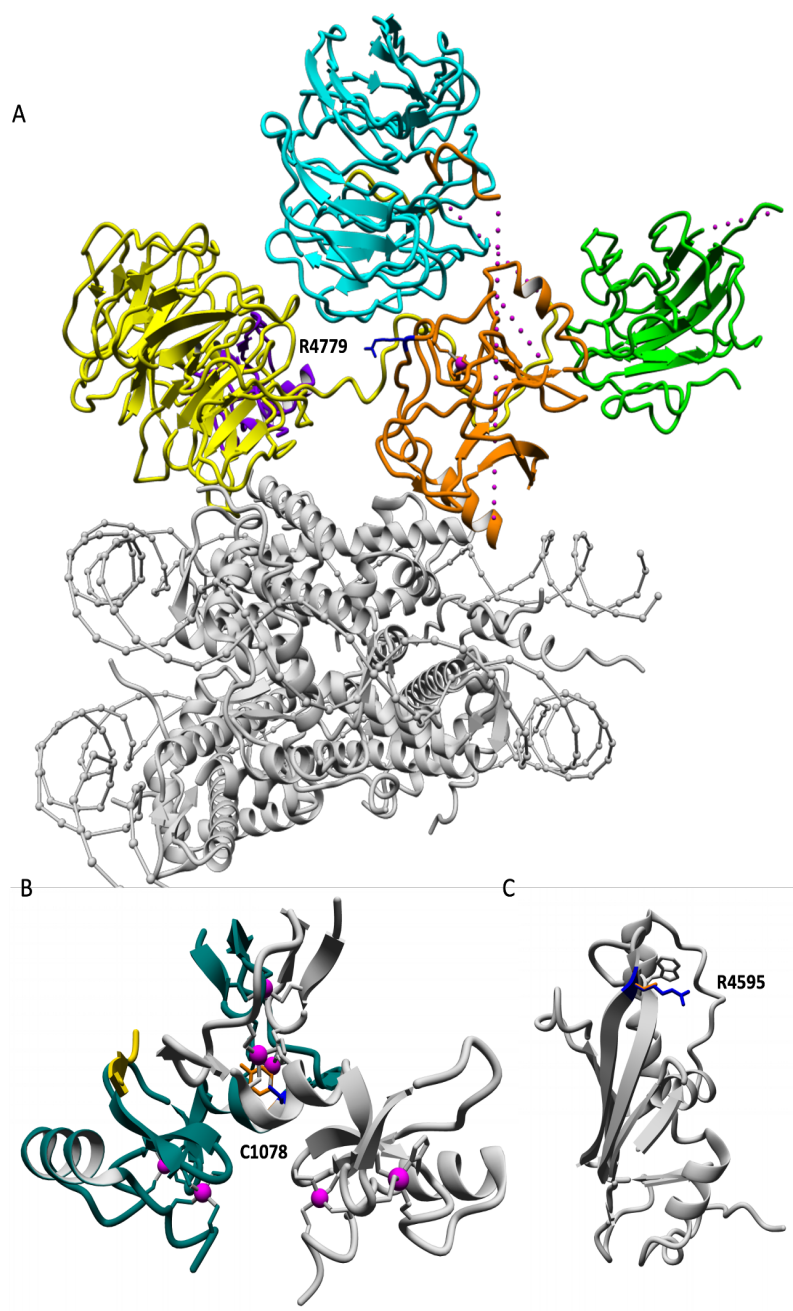
900

901 **Figure S1.** Distribution of the PTVs across *KMT2C* transcripts with their relative expression in
902 the gnomAD database.



903
904 **Figure S2.** *KMT2C*-related NDD facial gestalt heatmap showing significant facial areas most
905 different from controls.





912

913 *A) Arg4779 is located on a surface of the SET domain that interacts with WDR5 (PDB:5F6K).*
914 *Arg change to Gly is a radical change and will likely affect the interaction that is required for*
915 *the KMT2C activity. Previous study (25561738) showed that Arg4779Pro affected the*
916 *methyltransferase activity of KMT2C. B) Cys1078 is binding to Zn in ePHD6 to ensure the*
917 *correct folding and structure of the domain (PDB:6MLC); a change to Tyr would disrupt the*
918 *Zn binding and, therefore, domain structure. C) Arg4595 is located on a surface of the FYR*
919 *domain (AlphaFold2 model) and interacts with Trp4987, so change to smaller Cys would*
920 *affect local aminoacid packing and interactions, and as result might destabilize the domain or*
921 *affect binding to other proteins or binding between FYRN and FYRC domain parts.*