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Mutation-specific pathophysiological mechanisms define different neurodevelopmental disorders associated with SATB1 dysfunction

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1 Mutation-specific pathophysiological mechanisms define different neurodevelopmental

2 disorders associated with SATB1 dysfunction

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156 Abstract

Whereas large-scale statistical analyses can robustly identify disease-gene relationships, they 157 do not accurately capture genotype-phenotype correlations or disease mechanisms. We use 158 multiple lines of independent evidence to show that different variant types in a single gene, 159 SATB1, cause clinically overlapping but distinct neurodevelopmental disorders. Clinical 160 evaluation of 42 individuals carrying SATB1 variants identified overt genotype-phenotype 161 relationships, associated with different pathophysiological mechanisms, established by 162 functional assays. Missense variants in the CUT1 and CUT2 DNA-binding domains result in 163 stronger chromatin binding, increased transcriptional repression and a severe phenotype. In 164 contrast, variants predicted to result in haploinsufficiency are associated with a milder clinical 165 presentation. A similarly mild phenotype is observed for individuals with premature protein 166 truncating variants that escape nonsense-mediated decay, which are transcriptionally active 167 but mislocalized in the cell. Our results suggest that in-depth mutation-specific genotype-168 phenotype studies are essential to capture full disease complexity and to explain phenotypic 169 170 variability.

171 Main text

SATB1 encodes a dimeric/tetrameric transcription factor¹ with crucial roles in development and maturation of T-cells²⁻⁴. Recently, a potential contribution of *SATB1* to brain development was suggested by statistically significant enrichment of *de novo* variants in two large neurodevelopmental disorder (NDD) cohorts^{5; 6}, although its functions in the central nervous system are poorly characterized.

177 Through international collaborations⁷⁻⁹ conforming to local ethical guidelines and the 178 declaration of Helsinki, we identified 42 individuals with a rare (likely) pathogenic variant in SATB1 (NM 001131010.4), a gene under constraint against loss-of-function and missense 179 variation (pLoF: o/e=0.15 (0.08-0.29); missense: o/e=0.46 (0.41-0.52); gnomAD v2.1.1)¹⁰. 180 Twenty-eight of the SATB1 variants occurred de novo, three were inherited from an affected 181 parent, and five resulted from (suspected) parental mosaicism (Figure S1). Reduced 182 penetrance is suggested by two variants inherited from unaffected parents (identified in 183 individual 2 and 12; Table S1A), consistent with recent predictions of incomplete penetrance 184 being more prevalent in novel NDD syndromes⁶. Inheritance status of the final four could not 185 186 be established (Table S1A). Of note, two individuals also carried a (likely) pathogenic variant 187 affecting other known disease genes, including NF1 (MIM #162200; individual 27) and FOXP2 (MIM #602081; individual 42) which contributed to (individual 27) or explained (individual 42) 188 189 the observed phenotype (Table S1A).

190 Thirty individuals carried 15 unique SATB1 missense variants, including three recurrent 191 variants (Figure 1A), significantly clustering in the highly homologous DNA-binding domains CUT1 and CUT2 (p=1.00e-7; Figure 2A, Figure S2)^{11; 12}. Ten individuals carried premature 192 protein truncating variants (PTVs; two nonsense, seven frameshift, one splice site; Table S1A, 193 194 Table S2), and two individuals had a (partial) gene deletion (Figure S3). For 38 affected individuals and one mosaic parent, clinical information was available. Overall, we observed a 195 broad phenotypic spectrum, characterized by neurodevelopmental delay (35/36, 97%), ID 196 (28/31, 90%), muscle tone abnormalities (abnormal tone 28/37, 76%; hypotonia 28/37, 76%; 197 spasticity 10/36, 28%), epilepsy (22/37, 61%) behavioral problems (24/34, 71%), facial 198

dysmorphisms (24/36, 67%; Figure 1B-1C, Figure S4A), and dental abnormalities (24/34, 71%) 199 (Figure 1D, Table 1, Figure S4B, Table S1). Individuals with missense variants were globally 200 201 more severely affected than those with PTVs: 57% of individuals with a missense variant had 202 severe/profound ID whereas this level of ID was not observed for any individuals with PTVs. 203 Furthermore, hypotonia, spasticity and (severe) epilepsy were more common in individuals 204 with missense variants than in those with PTVs (92% versus 42%, 42% versus 0%, 80% versus 205 18%, respectively) (Figure 1F, Table 1, Table S1A). To objectively quantify these observations, 206 we divided our cohort into two variant-specific clusters (missense versus PTVs) and assessed the two groups using a Partitioning Around Medoids clustering algorithm¹³ on 100 features 207 derived from standardized clinical data (Human Phenotype Ontology (HPO); Figure S5A and 208 Suppl. JSON)¹⁴. Thirty-eight individuals were subjected to this analysis, of which 27 were 209 classified correctly as either belonging to the PTV or missense variant group (p=0.022), 210 confirming the existence of at least two separate clinical entities (Figure 1G, Figure S5B). 211 Moreover, computational averaging of facial photographs¹⁵ revealed differences between the 212 213 average facial gestalt for individuals with missense variants when compared to individuals with 214 PTVs or deletions (Figure 1B-E, Figure S4, Table S1B).

We performed functional analyses assessing consequences of different types of 215 SATB1 variants for cellular localization, transcriptional activity, overall chromatin binding, and 216 217 dimerization capacity. Based on protein modeling (Figure 2, Suppl. Notes), we selected five 218 missense variants (observed in 14 individuals) in CUT1 and CUT2 affecting residues that 219 interact with, or are close to, the DNA backbone (mosaic variant c.1220A>G; p.Glu407Gly and de novo variants c.1259A>G; p.Gln420Arg, c.1588G>A; p.Glu530Lys, c.1588G>C; 220 221 p.Glu530Gln, c.1639G>A; p.Glu547Lys), as well as the only homeobox domain variant 222 (c.2044C>G; p.Leu682Val, de novo). As controls, we selected three rare missense variants from the UK10K consortium, identified in healthy individuals with a normal IQ: c.1097C>T; 223 p.Ser366Leu (gnomAD allele frequency 6.61e-4), c.1555G>C; p.Val519Leu (8.67e-6) and 224 c.1717G>A; p.Ala573Thr (1.17e-4) (Figure 1A, Table S3)¹⁶. When overexpressed as YFP-225 fusion proteins in HEK293T/17 cells, wildtype SATB1 localized to the nucleus in a granular 226

pattern, with an intensity profile inverse to the DNA-binding dye Hoechst 33342 (Figure 3A-B).
In contrast to wildtype and UK10K control missense variants, the p.Glu407Gly, p.Gln420Arg,
p.Glu530Lys/p.Glu530Gln and p.Glu547Lys variants displayed a cage-like clustered nuclear
pattern, strongly co-localizing with the DNA (Figure 3A-B, Figure S6).

To assess the effects of SATB1 missense variants on transrepressive activity, we used 231 a luciferase reporter system with two previously established downstream targets of SATB1, 232 233 the *IL2*-promoter and IgH-MAR (matrix associated region)¹⁷⁻¹⁹. All five functionally assessed 234 CUT1 and CUT2 missense variants demonstrated increased transcriptional repression of the IL2-promoter, while the UK10K control variants did not differ from wildtype (Figure 3C). In 235 assays using IgH-MAR, increased repression was seen for both CUT1 variants, and for one of 236 the CUT2 variants (Figure 3C). The latter can be explained by previous reports that the CUT1 237 domain is essential for binding to MARs, whereas the CUT2 domain is dispensable^{20; 21}. Taken 238 together, these data suggest that etiological SATB1 missense variants in CUT1 and CUT2 239 lead to stronger binding of the transcription factor to its targets. 240

To study whether SATB1 missense variants affect the dynamics of chromatin binding 241 242 more globally, we employed fluorescent recovery after photobleaching (FRAP) assays. Consistent with the luciferase reporter assays, all CUT1 and CUT2 missense variants, but not 243 the UK10K control variants, affected protein mobility in the nucleus. The CUT2 variant 244 245 p.GIn420Arg demonstrated an increased half time, but showed a maximum recovery similar to 246 wildtype, while the other CUT1 and CUT2 variants demonstrated both increased halftimes and 247 reduced maximum recovery. These results suggest stabilization of SATB1 chromatin binding for all tested CUT1 and CUT2 variants (Figure 3D). 248

In contrast to the CUT1 and CUT2 missense variants, the homeobox variant p.Leu682Val did not show functional differences from wildtype (Figure 3A-D, Figure S6), suggesting that, although it is absent from gnomAD, highly intolerant to variation and evolutionarily conserved (Figure S2, Figure S7A-B), this variant is unlikely to be pathogenic. This conclusion is further supported by the presence of a valine residue at the equivalent position in multiple homologous homeobox domains (Figure S7C). Additionally, the mild

phenotypic features in this individual (individual 42) can be explained by the fact that the individual carries an out-of-frame *de novo* intragenic duplication of *FOXP2*, known to cause NDD through haploinsufficiency²².

We went on to assess the impact of the CUT1 and CUT2 missense variants (p.Glu407Gly, p.Gln420Arg, p.Glu530Lys, p.Glu547Lys) on protein interaction capacities using bioluminescence resonance energy transfer (BRET). All tested variants retained the ability to interact with wildtype SATB1 (Figure 3E), with the potential to yield dominant-negative dimers/tetramers *in vivo* and to disturb normal activity of the wildtype protein.

The identification of SATB1 deletions suggests that haploinsufficiency is a second 263 underlying disease mechanism. This is supported by the constraint of SATB1 against loss-of-264 function variation, and the identification of PTV carriers that are clinically distinct from 265 individuals with missense variants. PTVs are found throughout the locus and several are 266 predicted to undergo NMD by in silico models of NMD efficacy (Table S4)²³. In contrast to these 267 predictions, we found that one of the PTVs, c.1228C>T; p.Arg410*, escapes NMD (Figure S8A-268 269 B). However, the p.Arg410* variant would lack critical functional domains (CUT1, CUT2, 270 homeobox) and indeed showed reduced transcriptional activity in luciferase reporter assays when compared to wildtype protein (Figure S8), consistent with the haploinsufficiency model. 271

Four unique PTVs that we identified were located within the final exon of SATB1 (Figure 272 273 1A) and predicted to escape NMD (Table S4). Following experimental validation of NMD 274 escape (Figure 4A-B), three such variants (c.1877delC; p.Pro626Hisfs*81, c.2080C>T; 275 p.Gln694* and c.2207delA; p.Asn736llefs*8) were assessed with the same functional assays that we used for missense variants. When overexpressed as YFP-fusion proteins, the tested 276 277 variants showed altered subcellular localization, forming nuclear puncta or (nuclear) 278 aggregates, different from patterns observed for missense variants (Figure 4C, Figure S9A-B). In luciferase reporter assays, the p.Pro626Hisfs*81 variant showed increased repression of 279 both the *IL2*-promoter and IgH-MAR, whereas p.Gln694* only showed reduced repression of 280 281 IgH-MAR (Figure 4D). The p.Asn736llefs*8 variant showed repression comparable to that of wildtype protein for both targets (Figure 4D). In further pursuit of pathophysiological 282

mechanisms, we tested protein stability and SUMOylation, as the previously described 283 p.Lys744 SUMOylation site is missing in all assessed NMD-escaping truncated proteins 284 (Figure 4A)²⁴. Our observations suggest the existence of multiple SATB1 SUMOylation sites 285 286 (Figure S10) and no effect of NMD-escaping variants on SUMOylation of the encoded proteins (Figure S10) nor any changes in protein stability (Figure S9C). Although functional assays with 287 NMD-escaping PTVs hint towards additional disease mechanisms, HPO-based phenotypic 288 289 analysis or qualitative evaluation could not confirm a third distinct clinical entity (p=0.932; 290 Figure S5, Figure S11, Table S5).

Our study demonstrates that while statistical analyses^{5; 6} can provide the first step towards 291 identification of new NDDs, a mutation-specific functional follow-up is required to gain insight 292 into the underlying mechanisms and to understand phenotypic differences within patient 293 cohorts (Table S6). Multiple mechanisms and/or more complex genotype-phenotype 294 correlations are increasingly appreciated in newly described NDDs, such as those associated 295 with RAC1, POL2RA, KMT2E and PPP2CA²⁵⁻²⁸. Interestingly, although less often explored, 296 297 such mechanistic complexity might also underlie well-known (clinically recognizable) NDDs. For instance, a CUT1 missense variant in SATB2, a paralog of SATB1 that causes Glass 298 syndrome through haploinsufficiency (MIM #612313)²⁹, affects protein localization and nuclear 299 300 mobility in a similar manner to the corresponding SATB1 missense variants (Figure S12, Figure 301 S13)³⁰. Taken together, these observations suggest that mutation-specific mechanisms await 302 discovery both for new and well-established clinical syndromes.

In summary, we demonstrate that at least two different previously uncharacterized NDDs are caused by distinct classes of rare (*de novo*) variation at a single locus. We combined clinical investigation, *in silico* models and cellular assays to characterize the phenotypic consequences and functional impacts of a large patient series uncovering distinct pathophysiological mechanisms of the *SATB1*-associated NDDs. This level of combined analyses is recommended for known and yet undiscovered NDDs to fully understand disease etiology.

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328

329 Conflict of interest

KM, TBP, and TSS are employees of GeneDx, Inc. KR is employee of Ambrygen Genetics.

332 References

Wang, Z., Yang, X., Chu, X., Zhang, J., Zhou, H., Shen, Y., and Long, J. (2012). The
 structural basis for the oligomerization of the N-terminal domain of SATB1. Nucleic
 Acids Res 40, 4193-4202.

336	2. Alvarez, J.D., Yasui, D.H., Niida, H., Joh, T., Loh, D.Y., and Kohwi-Shigematsu, T. (2000).
337	The MAR-binding protein SATB1 orchestrates temporal and spatial expression of
338	multiple genes during T-cell development. Genes Dev 14, 521-535.
339	3. Cai, S., Lee, C.C., and Kohwi-Shigematsu, T. (2006). SATB1 packages densely looped,
340	transcriptionally active chromatin for coordinated expression of cytokine genes. Nat
341	Genet 38, 1278-1288.
342	4. Kitagawa, Y., Ohkura, N., Kidani, Y., Vandenbon, A., Hirota, K., Kawakami, R., Yasuda,
343	K., Motooka, D., Nakamura, S., Kondo, M., et al. (2017). Guidance of regulatory T cell
344	development by Satb1-dependent super-enhancer establishment. Nat Immunol 18,
345	173-183.
346	5. Satterstrom, F.K., Kosmicki, J.A., Wang, J., Breen, M.S., De Rubeis, S., An, J.Y., Peng,
347	M., Collins, R., Grove, J., Klei, L., et al. (2020). Large-Scale Exome Sequencing
348	Study Implicates Both Developmental and Functional Changes in the Neurobiology of
349	Autism. Cell 180, 568-584.e523.
350	6. Kaplanis, J., Samocha, K.E., Wiel, L., Zhang, Z., Arvai, K.J., Eberhardt, R.Y., Gallone, G.,
351	Lelieveld, S.H., Martin, H.C., McRae, J.F., et al. (2020). Evidence for 28 genetic
352	disorders discovered by combining healthcare and research data. Nature.
353	7. Sobreira, N., Schiettecatte, F., Valle, D., and Hamosh, A. (2015). GeneMatcher: a
354	matching tool for connecting investigators with an interest in the same gene. Hum
355	Mutat 36, 928-930.
356	8. Thompson, R., Johnston, L., Taruscio, D., Monaco, L., Beroud, C., Gut, I.G., Hansson,
357	M.G., t Hoen, P.B., Patrinos, G.P., Dawkins, H., et al. (2014). RD-Connect: an
358	integrated platform connecting databases, registries, biobanks and clinical
359	bioinformatics for rare disease research. J Gen Intern Med 29 Suppl 3, S780-787.
360	9. Firth, H.V., Richards, S.M., Bevan, A.P., Clayton, S., Corpas, M., Rajan, D., Van Vooren,
361	S., Moreau, Y., Pettett, R.M., and Carter, N.P. (2009). DECIPHER: Database of
362	Chromosomal Imbalance and Phenotype in Humans Using Ensembl Resources. Am
363	J Hum Genet 84, 524-533.

364	10. Karczewski, K.J., Francioli, L.C., Tiao, G., Cummings, B.B., Alföldi, J., Wang, Q., Collins,
365	R.L., Laricchia, K.M., Ganna, A., Birnbaum, D.P., et al. (2020). The mutational
366	constraint spectrum quantified from variation in 141,456 humans. Nature 581, 434-
367	443.
368	11. Lelieveld, S.H., Reijnders, M.R., Pfundt, R., Yntema, H.G., Kamsteeg, E.J., de Vries, P.,
369	de Vries, B.B., Willemsen, M.H., Kleefstra, T., Lohner, K., et al. (2016). Meta-analysis
370	of 2,104 trios provides support for 10 new genes for intellectual disability. Nat
371	Neurosci 19, 1194-1196.
372	12. Lelieveld, S.H., Wiel, L., Venselaar, H., Pfundt, R., Vriend, G., Veltman, J.A., Brunner,
373	H.G., Vissers, L., and Gilissen, C. (2017). Spatial Clustering of de Novo Missense
374	Mutations Identifies Candidate Neurodevelopmental Disorder-Associated Genes. Am
375	J Hum Genet 101, 478-484.
376	13. Kaufman L., R.P.J. (1987). Clustering by means of medoids
377	https://wis.kuleuven.be/stat/robust/papers/publications-1987/kaufmanrousseeuw-
378	clusteringbymedoids-I1norm-1987.pdf.
379	14. Köhler, S., Carmody, L., Vasilevsky, N., Jacobsen, J.O.B., Danis, D., Gourdine, J.P.,
380	Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of
380 381	Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic
380 381 382	Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027.
380 381 382 383	 Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027. 15. Reijnders, M.R.F., Miller, K.A., Alvi, M., Goos, J.A.C., Lees, M.M., de Burca, A.,
380 381 382 383 384	 Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027. 15. Reijnders, M.R.F., Miller, K.A., Alvi, M., Goos, J.A.C., Lees, M.M., de Burca, A., Henderson, A., Kraus, A., Mikat, B., de Vries, B.B.A., et al. (2018). De Novo and
380 381 382 383 384 385	 Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027. 15. Reijnders, M.R.F., Miller, K.A., Alvi, M., Goos, J.A.C., Lees, M.M., de Burca, A., Henderson, A., Kraus, A., Mikat, B., de Vries, B.B.A., et al. (2018). De Novo and Inherited Loss-of-Function Variants in TLK2: Clinical and Genotype-Phenotype
380 381 382 383 384 385 386	 Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027. 15. Reijnders, M.R.F., Miller, K.A., Alvi, M., Goos, J.A.C., Lees, M.M., de Burca, A., Henderson, A., Kraus, A., Mikat, B., de Vries, B.B.A., et al. (2018). De Novo and Inherited Loss-of-Function Variants in TLK2: Clinical and Genotype-Phenotype Evaluation of a Distinct Neurodevelopmental Disorder. Am J Hum Genet 102, 1195-
380 381 382 383 384 385 386 387	 Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027. 15. Reijnders, M.R.F., Miller, K.A., Alvi, M., Goos, J.A.C., Lees, M.M., de Burca, A., Henderson, A., Kraus, A., Mikat, B., de Vries, B.B.A., et al. (2018). De Novo and Inherited Loss-of-Function Variants in TLK2: Clinical and Genotype-Phenotype Evaluation of a Distinct Neurodevelopmental Disorder. Am J Hum Genet 102, 1195-1203.
380 381 382 383 384 385 386 386 387 388	 Gargano, M., Harris, N.L., Matentzoglu, N., McMurry, J.A., et al. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. Nucleic Acids Res 47, D1018-d1027. 15. Reijnders, M.R.F., Miller, K.A., Alvi, M., Goos, J.A.C., Lees, M.M., de Burca, A., Henderson, A., Kraus, A., Mikat, B., de Vries, B.B.A., et al. (2018). De Novo and Inherited Loss-of-Function Variants in TLK2: Clinical and Genotype-Phenotype Evaluation of a Distinct Neurodevelopmental Disorder. Am J Hum Genet 102, 1195-1203. 16. Walter, K., Min, J.L., Huang, J., Crooks, L., Memari, Y., McCarthy, S., Perry, J.R., Xu, C.,

health and disease. Nature 526, 82-90.

17. Pavan Kumar, P., Purbey, P.K., Sinha, C.K., Notani, D., Limaye, A., Jayani, R.S., and
Galande, S. (2006). Phosphorylation of SATB1, a global gene regulator, acts as a
molecular switch regulating its transcriptional activity in vivo. Mol Cell 22, 231-243.

18. Kumar, P.P., Purbey, P.K., Ravi, D.S., Mitra, D., and Galande, S. (2005). Displacement

- 395 of SATB1-bound histone deacetylase 1 corepressor by the human immunodeficiency
- 396 virus type 1 transactivator induces expression of interleukin-2 and its receptor in T
- 397 cells. Mol Cell Biol 25, 1620-1633.
- 19. Siebenlist, U., Durand, D.B., Bressler, P., Holbrook, N.J., Norris, C.A., Kamoun, M., Kant,
- 399 J.A., and Crabtree, G.R. (1986). Promoter region of interleukin-2 gene undergoes

400 chromatin structure changes and confers inducibility on chloramphenicol

401 acetyltransferase gene during activation of T cells. Mol Cell Biol 6, 3042-3049.

- 402 20. Ghosh, R.P., Shi, Q., Yang, L., Reddick, M.P., Nikitina, T., Zhurkin, V.B., Fordyce, P.,
- 403 Stasevich, T.J., Chang, H.Y., Greenleaf, W.J., et al. (2019). Satb1 integrates DNA
 404 binding site geometry and torsional stress to differentially target nucleosome-dense
 405 regions. Nat Commun 10, 3221.
- 406 21. Dickinson, L.A., Dickinson, C.D., and Kohwi-Shigematsu, T. (1997). An atypical
- 407 homeodomain in SATB1 promotes specific recognition of the key structural element in
 408 a matrix attachment region. J Biol Chem 272, 11463-11470.
- 409 22. MacDermot, K.D., Bonora, E., Sykes, N., Coupe, A.M., Lai, C.S., Vernes, S.C., Vargha-

410 Khadem, F., McKenzie, F., Smith, R.L., Monaco, A.P., et al. (2005). Identification of

- 411 FOXP2 truncation as a novel cause of developmental speech and language deficits.
 412 Am J Hum Genet 76, 1074-1080.
- 23. Lindeboom, R.G.H., Vermeulen, M., Lehner, B., and Supek, F. (2019). The impact of
 nonsense-mediated mRNA decay on genetic disease, gene editing and cancer
 immunotherapy. Nat Genet 51, 1645-1651.
- 24. Tan, J.A., Sun, Y., Song, J., Chen, Y., Krontiris, T.G., and Durrin, L.K. (2008). SUMO
 conjugation to the matrix attachment region-binding protein, special AT-rich

- sequence-binding protein-1 (SATB1), targets SATB1 to promyelocytic nuclear bodies 418 where it undergoes caspase cleavage. J Biol Chem 283, 18124-18134. 419 420 25. Haijes, H.A., Koster, M.J.E., Rehmann, H., Li, D., Hakonarson, H., Cappuccio, G., Hancarova, M., Lehalle, D., Reardon, W., Schaefer, G.B., et al. (2019). De Novo 421 Heterozygous POLR2A Variants Cause a Neurodevelopmental Syndrome with 422 Profound Infantile-Onset Hypotonia. Am J Hum Genet 105, 283-301. 423 424 26. O'Donnell-Luria, A.H., Pais, L.S., Faundes, V., Wood, J.C., Sveden, A., Luria, V., Abou 425 Jamra, R., Accogli, A., Amburgey, K., Anderlid, B.M., et al. (2019). Heterozygous Variants in KMT2E Cause a Spectrum of Neurodevelopmental Disorders and 426 Epilepsy. Am J Hum Genet 104, 1210-1222. 427 27. Reynhout, S., Jansen, S., Haesen, D., van Belle, S., de Munnik, S.A., Bongers, E., 428 Schieving, J.H., Marcelis, C., Amiel, J., Rio, M., et al. (2019). De Novo Mutations 429 Affecting the Catalytic Calpha Subunit of PP2A, PPP2CA, Cause Syndromic 430 Intellectual Disability Resembling Other PP2A-Related Neurodevelopmental 431 Disorders. Am J Hum Genet 104, 139-156. 432 28. Reijnders, M.R.F., Ansor, N.M., Kousi, M., Yue, W.W., Tan, P.L., Clarkson, K., Clayton-433 Smith, J., Corning, K., Jones, J.R., Lam, W.W.K., et al. (2017). RAC1 Missense 434 Mutations in Developmental Disorders with Diverse Phenotypes. Am J Hum Genet 435 436 101, 466-477. 437 29. Zarate, Y.A., Bosanko, K.A., Caffrey, A.R., Bernstein, J.A., Martin, D.M., Williams, M.S., 438 Berry-Kravis, E.M., Mark, P.R., Manning, M.A., Bhambhani, V., et al. (2019). Mutation update for the SATB2 gene. Hum Mutat 40, 1013-1029. 439 30. Lee, J.S., Yoo, Y., Lim, B.C., Kim, K.J., Choi, M., and Chae, J.H. (2016). SATB2-440
- 441 associated syndrome presenting with Rett-like phenotypes. Clin Genet 89, 728-732.

443 Figure legends

Figure 1. Clinical evaluation of SATB1 variants in neurodevelopmental disorders. A) 444 Schematic representation of the SATB1 protein (NM 001131010.4/NP 001124482.1), 445 446 including functional domains, with truncating variants labeled in cyan, truncating variants predicted to escape NMD in orange, splice site variants in purple, missense variants in 447 magenta, and UK10K rare control missense variants in green. Deletions are shown in dark 448 blue below the protein schematic, above a diagram showing the exon boundaries. We obtained 449 450 clinical data for all individuals depicted by a circle. B-C) Facial photographs of individuals with (partial) gene deletions and truncations (B), and of individuals with missense variants (C). All 451 depicted individuals show facial dysmorphisms and although overlapping features are seen, 452 453 no consistent facial phenotype can be observed for the group as a whole. Overlapping facial 454 dysmorphisms include facial asymmetry, high forehead, prominent ears, straight and/or full eyebrows, puffy eyelids, downslant of palpebral fissures, low nasal bridge, full nasal tip and 455 full nasal alae, full lips with absent cupid's bow, prominent cupid's bow or thin upper lip 456 457 vermillion (Table S1B). Individuals with missense variants are more alike than individuals in 458 the truncating cohorts, and we observed recognizable overlap between several individuals in 459 the missense cohort (individual 17, 27, 31, 37, the siblings 19, 20 and 21, and to a lesser extent individual 24 and 35). A recognizable facial overlap between individuals with (partial) gene 460 461 deletions and truncations could not be observed. Related individuals are marked with a blue 462 box. D) Photographs of teeth abnormalities observed in individuals with SATB1 variants. 463 Dental abnormalities are seen for all variant types and include widely spaced teeth, dental fragility, missing teeth, disorganized teeth implant, and enamel discoloration (Table S1B). E) 464 Computational average of facial photographs of 16 individuals with a missense variant (left) 465 and 8 individuals with PTVs or (partial) gene deletions (right). F) Mosaic plot presenting a 466 selection of clinical features. G) The Partitioning Around Medoids analysis of clustered HPO-467 standardized clinical data from 38 individuals with truncating (triangle) and missense variants 468 469 (circle) shows a significant distinction between the clusters of individuals with missense variants (blue) and individuals with PTVs (red). Applying Bonferroni correction, a p-value 470

smaller than 0.025 was considered significant. For analyses displayed in (F) and (G),
individuals with absence of any clinical data and/or low level mosaicism for the *SATB1* variant
were omitted (for details, see Suppl. Materials and Methods).

474

Figure 2. 3D protein modeling of SATB1 missense variants in DNA-binding domains. A) 475 Schematic representation of the aligned CUT1 and CUT2 DNA-binding domains. CUT1 and 476 477 CUT2 domains have a high sequence identity (40%) and similarity (78%). Note that the 478 recurrent p.Q402R, p.E407G/p.E407Q and p.Q525R, p.E530G/p.E530K/p.E530Q variants affect equivalent positions within the respective CUT1 and CUT2 domains, while p.Q420R in 479 CUT1 and p.E547K in CUT2 affect cognate regions. B) 3D-model of the SATB1 CUT1 domain 480 (left; PDB 2O4A) and CUT2 domain (right; based on PDB 2CSF) in interaction with DNA 481 (yellow). Mutated residues are highlighted in red for CUT1 and cyan for CUT2, along the ribbon 482 visualization of the corresponding domains in burgundy and dark blue, respectively. C) 3D-483 homology model of the SATB1 homeobox domain (based on PDB 1WI3 and 2D5V) in 484 interaction with DNA (yellow). The mutated residue is shown in light gray along the ribbon 485 486 visualization of the corresponding domain in dark gray. **B-C**) For more detailed descriptions of 487 the different missense variants in our cohort, see Suppl. Notes.

488

489 Figure 3. SATB1 missense variants stabilize DNA binding and show increased 490 transcriptional repression. A) Direct fluorescence super-resolution imaging of nuclei of 491 HEK293T/17 cells expressing YFP-SATB1 fusion proteins. Scale bar = $5 \mu m$. B) Intensity profiles of YFP-tagged SATB1 and variants, and the DNA binding dye Hoechst 33342. The 492 493 graphs represent the fluorescence intensity values of the position of the red lines drawn in the 494 micrographs on the top (SATB1 proteins in green, Hoechst 33342 in white, scale bar = $5 \mu m$). For each condition a representative image and corresponding intensity profile plot is shown. 495 496 C) Luciferase reporter assays using reporter constructs containing the *IL2*-promoter region and 497 the IgH matrix associated region (MAR) binding site. UK10K control variants are shaded in green, CUT1 domain variants in red, CUT2 domain variants in blue and the homeobox variant 498

499 in gray. Values are expressed relative to the control (pYFP; black) and represent the mean \pm S.E.M. (n = 4, p-values compared to wildtype SATB1 (WT; white), one-way ANOVA 500 501 and post-hoc Bonferroni test). D) FRAP experiments to assess the dynamics of SATB1 chromatin binding in live cells. Left, mean recovery curves ± 95% C.I. recorded in HEK293T/17 502 503 cells expressing YFP-SATB1 fusion proteins. Right, violin plots with median of the halftime 504 (central panel) and maximum recovery values (right panel) based on single-term exponential 505 curve fitting of individual recordings (n = 60 nuclei from three independent experiments, p-506 values compared to WT SATB1, one-way ANOVA and post-hoc Bonferroni test). Color code 507 as in C. E) BRET assays for SATB1 dimerization in live cells. Left, mean BRET saturation curves ± 95% C.I. fitted using a non-linear regression equation assuming a single binding site 508 (y = BRETmax * x / (BRET50 / x); GraphPad). The corrected BRET ratio is plotted against the 509 ratio of fluorescence/luminescence (AU) to correct for expression level differences between 510 conditions. Right, corrected BRET ratio values at mean BRET50 level of WT SATB1, based 511 512 on curve fitting of individual experiments (n = 4, one-way ANOVA and post-hoc Bonferroni test, 513 no significant differences). Color code as in C. A-E) When compared to WT YFP-SATB1 or 514 UK10K variants, most variants identified in affected individuals show a nuclear cage-like localization (A), stronger co-localization with the DNA-binding dye Hoechst 33342 (B), 515 increased transcriptional repression (C), reduced protein mobility (D) and unchanged capacity 516 517 of interaction with WT SATB1 (E).

518

519 Figure 4. SATB1 frameshift variants in the last exon escape NMD. A) Schematic overview of the SATB1 protein, with truncating variants predicted to escape NMD that are included in 520 521 functional assays labeled in orange. A potential SUMOylation site at position p.K744 is 522 highlighted. B) Sanger sequencing traces of patient-derived EBV-immortalized lymphoblastoid 523 cell lines treated with or without cycloheximide (CHX) to test for NMD. The mutated nucleotides are shaded in red. Transcripts from both alleles are present in both conditions showing that 524 these variants escape NMD. C) Direct fluorescence super-resolution imaging of nuclei of 525 HEK293T/17 cells expressing SATB1 truncating variants fused with a YFP-tag. Scale bar = 5 526

µm. Compared to WT YFP-SATB1, NMD-escaping variants show altered localization forming nuclear puncta or aggregates. **D**) Luciferase reporter assays using reporter constructs containing the *IL2*-promoter and the IgH matrix associated region (MAR) binding site. Values are expressed relative to the control (pYFP; black) and represent the mean ± S.E.M. (*n* = 4, *p*values compared to WT SATB1 (white), one-way ANOVA and *post-hoc* Bonferroni test). All NMD-escaping variants are transcriptionally active and show repression of the *IL2*-promoter and IgH-MAR binding site.

Table 1. Summary of clinical characteristics associated with (*de novo***) SATB1 variants**

	All individuals		Individuals with PTVs and (partial) gene deletions		Individuals with missense variants	
	%	Present / total	%	Present / total	%	Present / total
	78	assessed	70	assessed	70	assessed
Neurologic		0.0 /0 /		0// 0		00/0/
Intellectual disability	90	28/31	80	8/10	95	20/21
Normal	10	3/31	20	2/10	5	1/21
Borderline	0	0/31	0	0/10	0	0/21
Mild	26	8/31	60	6/10	10	2/21
Moderate	10	3/31	10	1/10	10	2/21
Severe	19	6/31	0	0/10	29	6/21
Profound	19	6/31	0	0/10	29	6/21
Unspecified	16	5/31	10	1/10	19	4/21
Developmental delay	97	35/36	100	12/12	96	23/24
Motor delay	92	34/37	92	11/12	92	23/25
Speech delay	89	32/36	83	10/12	92	22/24
Dysarthria	30	6/20	9	1/11	56	5/9
Epilepsy	61	22/36	18	2/11	80	20/25
EEG abnormalities	79	19/24	29	2/7	100	17/17
Hypotonia	76	28/37	42	5/12	92	23/25
Spasticity	28	10/36	0	0/12	42	10/24
Ataxia	22	6/27	17	2/12	27	4/15
Behavioral disturbances	71	24/34	58	7/12	77	17/22
Sleep disturbances	41	12/29	27	3/11	50	9/18
Abnormal brain imaging	55	17/31	43	3/7	58	14/24
Regression	17	6/35	8	1/12	22	5/23
Growth						
Abnormalities during pregnancy	24	8/33	27	3/11	23	5/22
Abnormalities during delivery	32	10/31	55	6/11	20	4/20
Abnormal term of delivery	6	2/31	10	1/10	5	1/21
Preterm (<37 weeks)	6	2/31	10	1/10	5	1/21
Postterm (>42 weeks)	0	0/31	0	0/10	0	0/21
Abnormal weight at birth	16	5/32	22	2/9	13	3/23
Small for gestational age (<p10)< td=""><td>9</td><td>3/32</td><td>11</td><td>1/9</td><td>9</td><td>2/23</td></p10)<>	9	3/32	11	1/9	9	2/23
Large for gestational age (>p90)	6	2/32	11	1/9	4	1/23
Abnormal head circumference at birth	7	1/14	17	1/6	0	0/8
Microcephaly (<p3)< td=""><td>0</td><td>0/14</td><td>0</td><td>0/6</td><td>0</td><td>0/8</td></p3)<>	0	0/14	0	0/6	0	0/8
Macrocephaly (>p97)	7	1/14	17	1/6	0	0/8
Abnormal height	21	6/29	9	1/11	28	5/18
Short stature (<p3)< td=""><td>14</td><td>4/29</td><td>0</td><td>0/11</td><td>22</td><td>4/18</td></p3)<>	14	4/29	0	0/11	22	4/18
Tall stature (>p97)	7	2/29	9	1/11	6	1/18
Abnormal head circumference	26	7/31	11	1/9	32	6/22
Microcephaly (<p3)< td=""><td>26</td><td>7/31</td><td>11</td><td>1/9</td><td>32</td><td>6/22</td></p3)<>	26	7/31	11	1/9	32	6/22
Macrocephaly (>p97)	0	0/31	0	0/9	0	0/22
Abnormal weight	48	13/27	11	1/9	67	12/18
Underweight (<p3)< td=""><td>22</td><td>6/27</td><td>11</td><td>1/9</td><td>28</td><td>5/18</td></p3)<>	22	6/27	11	1/9	28	5/18
Overweight (>p97)	26	7/27	0	0/9	39	7/18
Other phenotypic features						
Facial dysmorphisms	67	24/36	64	7/11	68	17/25
Dental/oral abnormalities	71	24/34	55	6/11	78	18/23
Drooling/dysphagia	38	12/32	25	3/12	45	9/20
Hearing abnormalities	7	2/30	18	2/11	0	0/19
Vision abnormalities	55	17/31	73	8/11	45	9/20
Cardiac abnormalities	19	6/32	27	3/11	14	3/21
Skeleton/limb abnormalities	38	13/34	18	2/11	48	11/23
Hypermobility of joints	30	8/27	30	3/10	29	5/17
Gastrointestinal abnormalities	53	17/32	27	3/11	67	14/21
Urogenital abnormalities	17	5/30	0	0/11	26	5/19
Endocrine/metabolic abnormalities	30	9/30	0	0/11	47	9/19
Immunological abnormalities	32	8/25	25	2/8	35	6/17
Skin/hair/nail abnormalities	24	8/34	9	1/11	30	7/23
Neoplasms in medical history	0	0/34	0	0/11	0	0/23







