# Three-dimensional chromatin interactions remain stable upon CAG/CTG repeat expansion

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# 22 Abstract

Expanded CAG/CTG repeats underlie thirteen neurological disorders, including myotonic 23 dystrophy (DM1) and Huntington's disease (HD). Upon expansion, CAG/CTG repeat loci acquire 24 heterochromatic characteristics. This observation raises the hypothesis that repeat expansion 25 provokes changes to higher order chromatin folding and thereby affects both gene expression in 26 cis and the genetic instability of the repeat tract. Here we tested this hypothesis directly by 27 performing 4C sequencing at the DMPK and HTT loci from DM1 and HD patient-derived cells. 28 Surprisingly, chromatin contacts remain unchanged upon repeat expansion at both loci. This was 29 true for loci with different DNA methylation levels and CTCF binding. Repeat sizes ranging from 30 15 to 1,700 displayed strikingly similar chromatin interaction profiles. Our findings argue that 31 extensive changes in heterochromatic properties are not enough to alter chromatin folding at 32 33 expanded CAG/CTG repeat loci. Moreover, the ectopic insertion of an expanded repeat tract did not change three-dimensional chromatin contacts. We conclude that expanded CAG/CTG 34 repeats have little to no effect on chromatin conformation. 35

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# 37 Introduction

The genome is organized into hierarchical topologically associated domains (TADs) (1). This 38 three-dimensional organization of chromatin in the nucleus has a profound impact on 39 transcription, DNA replication, recombination, and repair (2, 3). For instance, heterochromatic and 40 euchromatic loci are spatially separated and display distinct three-dimensional (3D) chromatin 41 interactions as gleaned by microscopy and chromosome conformation capture (3C)-based 42 techniques (2, 4, 5). How this higher-order chromatin structure impinges on biological functions, 43 what determines chromatin domain boundaries, and how it contributes to disease is unclear. 44 Expanded CAG/CTG repeats loci (hereafter referred to according to their mRNA sequence) are 45 ideal to address these questions because they cause diseases and they are associated with 46 changes in local chromatin structure, transcriptional output, and genetic instability (6). 47

CAG/CTG repeats underlie 13 different neurological and neuromuscular disorders including 48 myotonic dystrophy type 1 (DM1) and Huntington's disease (HD). They are part of a larger group 49 of diseases caused by the expansion of short tandem repeats (STRs) (7, 8). Disease-associated 50 STRs (daSTRs) are genetically unstable, especially once they surpass a critical threshold of about 51 35 to 50 units. Their expansion is associated with extensive chromatin remodeling of the 52 expanded loci (6, 9-17). Two such examples are fragile X syndrome (FXS), caused by the 53 expansion of a CGG repeat at the FMR1 gene located on the X chromosome (18-21), and 54 Friedreich's ataxia (FRDA), caused by a homozygous GAA expansion in the first intron of the 55 FXN gene (22). In the case of FXS, expansions beyond 200 CGGs are associated with promoter 56 silencing in *cis*. This locus accumulates high levels of heterochromatic marks including CpG 57 methylation, H4K20me3, H3K9me2/3, and H3K27me3 while losing euchromatin-associated 58 marks, such as H3 and H4 acetylation as well as H3K4me2 (14, 23-26). In FRDA patient-derived 59 cells, the FXN locus with a tendency towards heterochromatinization and loses CTCF binding in 60 cis (12, 27-30). 61

The shift from a euchromatic to a heterochromatic state upon repeat expansion has led to the hypothesis that there is a concurrent change in higher order chromatin folding. Indeed, 3C-based experiments have revealed an increase in the frequencies of 3D chromatin interactions surrounding expanded GAA and CGG repeats in patient-derived lymphoblastoid cell lines (LCLs) (*12*, *23*, *31*). This was interpreted as enhanced compaction around the *FXN* locus and as a rearrangement of chromatin contact domains at the *FMR1* locus in FXS cells, respectively. It was speculated that higher order chromatin folding might contribute to the silencing of the genes in

the vicinity of the expanded repeat (*12*, *23*, *31*). Importantly, daSTRs are found predominantly at
 TAD and sub-TAD boundaries, suggesting more generally that daSTR expansions may disrupt
 TADs (*31*). This may then contribute to gene silencing in cis and to the high levels of instability
 found at these sequences, ultimately altering disease progression (*31*).

The critical unknowns in this model are whether changes in higher-order chromatin structure are 73 confined to CGG and GAA repeats or if this is general to daSTRs and whether changes in 74 chromatin interactions cause alterations in gene expression and repeat instability. The later 75 hypothesis is especially appealing in the context of the expanded CTG repeats in the 3'UTR of 76 the DMPK gene in DM1 cells because, like expanded CGG and GAA repeat loci, this region 77 undergoes heterochromatinization upon repeat expansion. The changes observed at the DMPK 78 locus include the loss of CTCF binding and of a DNAsel hypersensitive site, an increase in DNA 79 and H3K9 methylation, as well as a loss of acetylated histone marks around the repeat tract (13, 80 16, 17, 32-34). Moreover, the ectopic introduction of an expanded CAG repeat in yeast, was 81 sufficient to relocate the locus to the nuclear periphery in S-phase budding yeast cells leading to 82 changes in repeat size (35). Importantly, expanded CAG/CTG repeats cause the majority of 83 daSTR disorders. Together, these observations prompted us to test the hypothesis that chromatin 84 conformation changes at expanded CAG/CTG repeats. We used 4C-seg in LCLs from DM1 and 85 HD individuals and in HEK293-derived cells harboring an ectopic CAG repeat expansion. We 86 found no evidence to support changes in chromatin interactions that would underlie the alterations 87 in transcriptional output or genetic instability of these sequences. This was consistent in different 88 cell types, genetic backgrounds, and in the presence of specific heterochromatin marks. We 89 conclude that changes in higher order chromatin folding do not contribute to the pathogenesis of 90 expanded CAG/CTG repeat disorders. 91

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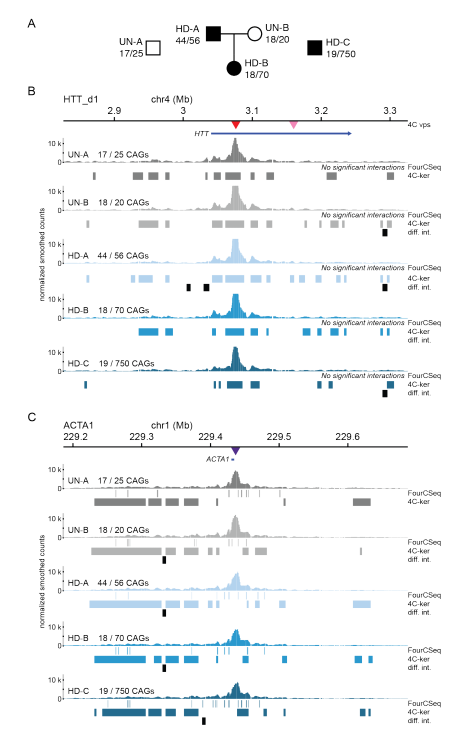
#### 93 **Results**

# 94 Chromatin conformation is stable upon repeat expansion at the *HTT* locus

To assess whether chromatin contacts change upon CAG repeat expansion at the *HTT* locus, we used a series of HD patient-derived LCLs (GM02164, GM03620, and GM14044, referred to as HD-A, HD-B, HD-C respectively) as well as two lines from unaffected individuals (GM04604 and GM02180, UN-A and UN-B respectively). Their family relationships and their repeat sizes are found in Fig. 1A, S1A, and Table S1.

To determine chromatin conformation, we used 4C-seg (36, 37) because it maximizes resolution 100 at the loci of interest. This allows a high sensitivity to small changes in conformation that may be 101 missed by other 3C-based methods. For the HTT locus, we used 2 viewpoints on chromosome 4 102 within the gene body – 1 kb (HTT d1) and 85 kb (HTT d85) downstream of the CAG repeats. To 103 control for a potential effect of the pathology on genome-wide chromatin conformation, we also 104 used a viewpoint located near the ACTA1 gene on chromosome 1. We obtained three replicates 105 of each 4C viewpoint and compared the DNA interaction profiles between the unaffected and HD 106 cell lines (Fig. 1B). Replicates from the same cell lines show good correlation in fragments with 107 more than 20 mapped reads (Fig. S2A-C). 108

We then identified fragments and regions that interact with the 4C viewpoint at frequencies higher 109 than expected (significant interactions, see methods). We also looked for regions that show 110 significant differences in interaction frequencies between patient-derived and unaffected cells 111 (differential interactions, see methods). To do so, we used two 4C-seq data analysis packages: 112 FourCSeq (38) and 4C-ker (39). We found that the interaction profiles were similar within a 2 Mb 113 region around the viewpoint (Fig. 1B, Fig. S3). Notably, the chromatin conformation remained 114 unaltered in HD-A, an HD patient cell line that has two expanded alleles (44/56 CAGs), as well 115 as in HD-B (18/70 CAGs) and HD-C (19/750 CAGs). The ACTA1 viewpoint also produced 116 indistinguishable interaction profiles between unaffected and HD patient LCLs (Fig. 1C). We 117 identified few small regions displaying differential interactions, but they were mainly outside of 118 regions with significant interactions (Fig. 1BC). In addition, most regions of differential interactions 119 were not exclusive to HD patient cells, as they were also found in comparisons between the two 120 unaffected cell lines. This suggests that the minor changes in interaction frequencies are due to 121 factors other than the presence of the expanded repeat tract in HTT, e.g. the different genetic 122 backgrounds. Taken together, these results show that a CAG repeat expansion at the HTT locus 123 does not cause significant alterations in chromatin conformation. 124



# Fig. 1. Chromatin interactions of the *HTT* locus in unaffected and HD cells.

(A) Pedigree of HD patient cell lines used in this study along with unrelated unaffected and HD cell lines (UN-A and HD-C respectively). (B) 4C-seg chromatin interaction profiles (average of triplicate smoothed and normalized counts) from the HTT d1 viewpoint (1 kb downstream of the CAG repeats - red central triangle) in two unaffected (UN-A and UN-B) and three HD patient cell lines (HD\_A, HD\_B, and HD\_C). The top blue bar represents the HTT gene and the triangles represent the location of both HTT viewpoints. The interaction profiles for the HTT\_d85 viewpoint (85 kb downstream of the CAG repeat) can be found in Fig. S3. (C) 4Cseq chromatin interaction profiles (average of triplicate smoothed and normalized counts) from the ACTA1 viewpoint (central purple triangle). For panels B and C, high-interacting regions were called using 4C-ker and significant interactions were called using FourCSeq. Regions of differential interactions are marked with black bars below each 4C-seq track and labeled as "diff. int.". The top blue bar represents the ACTA1 gene.

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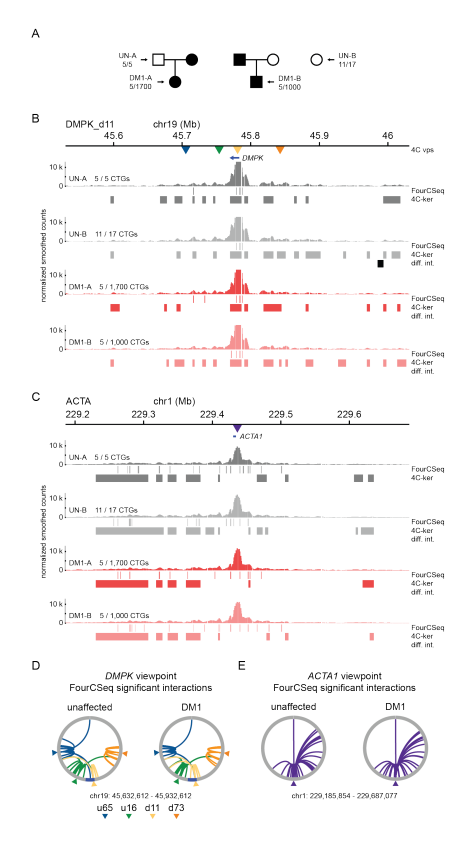
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# 127 Chromatin conformation is stable upon repeat expansion at the DMPK locus

Repeat expansion at the *HTT* locus is not known to be associated with significant changes in histone marks and chromatin accessibility, which might cause changes in chromatin conformation. To determine whether our findings applied in a case where chromatin structure is altered around expanded CAG/CTG repeats, we studied four viewpoints in the *DMPK* gene. We

used two DM1 patient-derived LCLs (GM06077 and GM04648, DM1-A and DM1-B respectively). 132 The DM1-A cell line harbored one expanded DMPK allele with 1,700 CTGs and the DM1-B cell 133 line, 1,000 CTGs (Fig. 2A and S1B). We found that DM1-A cells had increased CpG methylation 134 levels at two CTCF binding sites flanking the repeats (Fig. S4B), with a concomitant loss of CTCF 135 binding at these sites (Fig. S4C). In DM1-B cells we observed normal methylation levels at both 136 CTCF binding sites and slightly reduced CTCF binding (Fig S4B). Thus, DM1-A cells displayed 137 molecular signatures of congenital DM1 (40), whereas DM1-B cells have adult onset DM1 138 139 characteristics (Table S1).

We performed 4C-seg on the unaffected and DM1 cell lines using four different 4C viewpoints at 140 distinct distances away from the DMPK CTG repeats (Fig. 2A). Replicates from the same cell 141 lines also showed good correlation in 4C fragment counts above 20 (Fig. S2C-G). Similar to the 142 HD scenario, we observed strikingly similar chromatin interaction profiles between unaffected and 143 DM1 samples for all four viewpoints in the DMPK region (Fig. 2B, Fig. S5). As expected, the 144 ACTA1 viewpoint showed interaction profiles indistinguishable between unaffected and DM1 cell 145 lines (Fig. 2C). As with the HTT viewpoints, none of the DMPK or ACTA1 viewpoints in DM1 146 patient cells had significant interactions that were also called as regions of differential interaction 147 (Fig. 2B-E). Taken together, we found no evidence for large-scale changes in chromatin 148 interactions at the DMPK locus driven by CTG expansions, despite the changes in CTCF 149 occupancy and DNA methylation levels. 150



# Fig. 2. Chromatin interactions of the DMPK

#### locus in unaffected and DM1 cells.

(A) Pedigree of DM1 patient cell lines used in this study along with an unrelated unaffected cell line (UN-B). (B) 4C-seq chromatin interaction profiles (average of triplicate smoothed and normalized counts) from the DMPK d11 viewpoint (11kb downstream of the CTG repeats - yellow triangle) in two unaffected (UN-A and UN-B) and two DM1 patient cell lines (DM1-A and DM1-B). The top blue bar represents the DMPK gene and the triangles represent the location of the four DMPK viewpoints. The profiles for the three other viewpoints can be found in Fig. S5. (C) 4C-seq chromatin interaction profiles (average of triplicate smoothed and normalized counts) from the ACTA1 viewpoint (central purple triangle) in two unaffected and two DM1 patient cell lines. The top blue bar represents the ACTA1 gene. For panels B and C, highinteracting regions were called using 4C-ker and significant interactions were called using FourCSeq. Regions of differential interactions are marked with black bars below each 4C-seq track and labeled as "diff. int.". (D) Circos plot of the significant interactions called with FourCSeq (nominal p-value < .05) from four different viewpoints surrounding the CTG repeats of DMPK in the unaffected and DM1 cell lines (left and right respectively) (DMPK\_u65 in blue, DMPK\_u16 in green, DMPK d11 in yellow, and DMPK d73 in orange, 65 kb upstream, 16 kb upstream, 11 kb downstream, and 73 kb downstream, respectively). (E) Circos plot of the significant interactions called with FourCSeq (nominal pvalue < .05) from the control ACTA1 viewpoint in unaffected and DM1 cell lines (left and right respectively).

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# 155 Lack of allelic bias in chromatin interactions at expanded CAG/CTG repeats

DM1 and HD are both dominantly inherited disorders with individuals being heterozygous for the 156 expanded allele. Thus, one potential caveat in our data was that the presence of a normal-length 157 allele masks changes in 3D chromatin interactions made by the expanded allele. To evaluate this 158 possibility, we took advantage of the presence of at least one parental cell line for the DM1-A and 159 HD-B patient cell lines in our dataset (Fig. 1A and 2A) and identified allele-specific single 160 nucleotide polymorphisms (SNPs) in the 4C-seq data from these samples. We selected a subset 161 of the SNPs that could be assigned unambiguously to either the expanded or the normal allele 162 within 1 Mb of the 4C viewpoints (see methods). We reasoned that if chromatin contacts were 163 established without a systematic bias for either allele in DM1 and HD patient cells, the proportion 164 of 4C fragments in which the expanded chromosome had more reads than the normal-length one 165 would be close to 50%. We analyzed the sequencing coverage of the 4C data at these SNPs 166 positions and found that the viewpoints did not establish preferential contacts with a single 167 chromosome in neither the DM1-A nor HD-B patient cell lines (Table 1). These results are 168 consistent with the conclusion that chromatin interactions at both disease loci do not show allelic 169 bias. Together, these results corroborate the conclusion that expanded CAG/CTG repeats do not 170 significantly alter the DNA interactions at two expanded CAG/CTG repeat loci. 171

Cell line	Disease	Genomic region <sup>1</sup>	Total # of samples	# of samples with more normal chromosome reads	# of samples with more expanded chromosome reads	p-value <sup>2</sup>
DM1-A	DM1	ACTA1	6	1	5	0.22
		DMPK	28 *	18	9	0.09
HD-B	HD	ACTA1	5	2	3	1
		HTT	19	7	12	0.25

Table 1. Allele-specific interactions in 4C-seq data.

<sup>1</sup> 1Mb region around the 3'end of DMPK, 5' end of HTT, or the ACTA1 4C-seq viewpoint.

<sup>2</sup> p-value of an exact binomial test.

\* 1 sample had the same number of reads in both alleles, thus it was not included

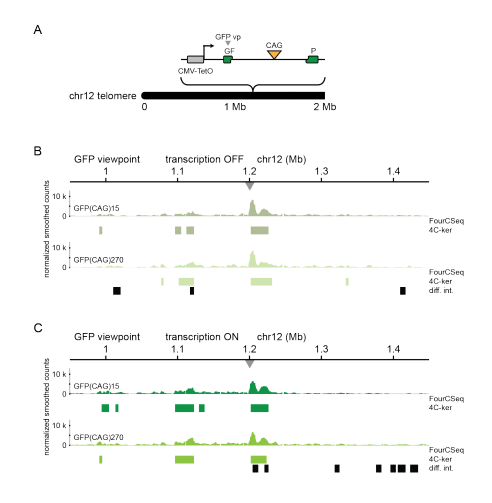
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#### 173 Chromatin conformation at an ectopic CAG repeat locus

It remained possible that the genetic background may have had a confounding effect on the chromatin interactions made at expanded CAG/CTG repeat loci. To test this, we compared the chromatin interactions of a hemizygous ectopic locus with either normal-length or expanded CAG/CTG repeats in otherwise isogenic cell lines. We obtained two clonal populations of Flp-In T-REx 293 cells that contain a single, stably integrated construct containing CAG repeats within

the intron of a GFP mini-gene (41, 42). One cell line contained 15 CAGs (GFP(CAG)<sub>15</sub>) whereas 179 the other had 270 CAGs (GFP(CAG)<sub>270</sub>) which is within the pathogenic range for both DM1 and 180 HD. The reporter sports a doxycycline-inducible promoter. Using targeted locus amplification (43) 181 and TAIL PCR, we mapped the insertion site to the p-arm of chromosome 12, 1.2 Mb from the 182 telomere (Fig 3A). We performed 4C-seg in both cell lines with a viewpoint located 1 kb upstream 183 of the CAG repeats. The chromatin interaction profiles of this ectopic CAG locus were also very 184 similar between the 15 and 270 CAG repeat cells, with few regions of differential interactions 185 overlapping with those displaying significant interactions using 4Cker. By contrast, FourCSeg did 186 not detect any changes that were significant (Fig. 3B). We concluded that CAG repeats at an 187 ectopic site causes few changes to chromatin conformation. 188

Some studies suggest that transcription may help define topological domain boundaries (44-47). 189 To determine whether transcription through expanded CAG repeats could lead to changes in 190 chromosome conformation, we induced transcription by culturing the GFP(CAG)<sub>n</sub> cells with 191 doxycycline for five days. We observed that only 2 of 17 regions of differential interactions 192 overlapped fully or in part with high-interacting regions over a 2 Mb region using 4Cker. 193 FourCSeq, on the other hand, found no significant interactions (Fig. 3C). These results suggest 194 that an ectopic expanded CAG repeat tract is not enough to alter the chromosome conformation 195 significantly at this locus, regardless of the transcriptional status. 196



# Fig. 3. Chromatin interactions of an expanded CAG repeat locus in isogenic cells.

(A) Diagram of the integration site of the CAG ectopic locus in GFP(CAG)<sub>n</sub> cells. (B) 4C-seq chromatin interaction profiles (average of triplicate smoothed and normalized counts) from the GFP viewpoint in GFP(CAG)<sub>15</sub> (top) and GFP(CAG)<sub>270</sub> (bottom). Cells were treated with doxycycline to induce transcription of the GFP mini-gene. (C) 4C-seq chromatin interactions from the GFP viewpoint in GFP(CAG)<sub>15</sub> (top) and GFP(CAG)<sub>270</sub> (top) when transcription was not induced.

# 199 Discussion

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Here we showed that chromatin interactions remain stable at expanded CAG/CTG repeat loci. 200 This was true for two disease loci and one transgene and in two different cell types. Our findings 201 are also supported by allele-specific analysis of 4C-seq interactions. Furthermore, CpG 202 methylation and CTCF binding at two sites flanking the CTG repeats of DMPK did not impact 203 chromosome conformation. This is especially relevant because CTCF is a key architectural 204 protein involved in the demarcation of TAD and sub-TAD boundaries (48). Additionally, when we 205 inserted a hemizygous transgene with CAG repeats, we found that expanded CAG repeats did 206 not induce a significant re-organization of the chromatin contacts established at this exogenous 207 CAG/CTG repeat locus. These results further show that an expanded CAG/CTG repeat tract is 208 not sufficient to change chromatin folding. 209

210 One possibility is that expanded CAG/CTG repeats lead to changes in chromatin conformation in 211 specific cell types that are especially vulnerable in DM1 or HD, for example cardiomyocytes or

212 medium spiny neurons, but not in LCLs. However, that would complicate the hypothesis by adding 213 a requirement for a cell-type specific factors to mediate these changes. If such a factor exists, 214 our conclusion remains unaffectcted: that the expansion of CAG/CTG repeats is not enough to 215 effect changes in chromatin conformation.

Our findings are in contrast to the effect of expanded CGG and GAA repeats on chromosome 216 conformation (12, 23, 31). In FXS patient-derived B-lymphoblast and fibroblasts, expanded CGG 217 repeats in *FMR1* were correlated with heterochromatic characteristics, decreased CTCF binding, 218 and ultimately a disruption of a TAD boundary near the expanded repeats (31). The DM1 patient 219 cells used here, especially the congenital DM1 patient cell line, share similar changes in chromatin 220 marks to FXS patient cells. And yet, these factors did not amount to an alteration of the 3D 221 chromatin interactions at the expanded DMPK locus. There are important differences between 222 FXS/FRDA and DM1/HD that could account for the differences. Apart from the nucleotide 223 composition of the repeat tract themselves, the disease loci and thus their flanking sequences 224 are different. Further investigations are needed to understand what accounts for the differences. 225

226 It was speculated that the changes in chromatin conformation found at expanded CGG repeats could lead to changes in repeat instability (31). Indeed, CGG repeat expansions beyond 200 units 227 are associated with promoter silencing, chromatin contact alterations and their instability is 228 lessened (49, 50). By contrast, long GAA and CAG/CTG repeats are more unstable (7, 51, 52), 229 yet the chromosome conformation at expanded GAA repeats is altered but not at expanded 230 CAG/CTG repeats. Thus, there does not seem to be a correlation between changes in 3D 231 chromatin contacts and repeat instability. One possibility is that the effect of chromosome 232 conformation on repeat instability is repeat-type specific. Another is that chromosome 233 conformation is unrelated to the repeat instability and is neither a cause nor a consequence. 234 Similarly, changes in transcriptional output around expanded repeats may be unrelated to 235 chromatin conformation and rather be the consequence of changes in local chromatin marks, or 236 they may be specific to the repeat sequence. 237

The expanded CAG/CTG repeat loci in DM1 and HD provide an endogenous genomic substrate to study the mechanisms necessary for the establishment of 3D chromatin domains in daSTRs given the involvement of CTCF binding, CpG methylation, and other chromatin remodeling events at these loci. Although CTCF is generally a major driver of genome organization, it is not the only driver. There are cases where CTCF binding abrogation does not alter an evolutionarily conserved TAD boundary, for instance at the *Firre* locus (53). Moreover, our results are in line with recent evidence in *Drosophila* arguing that genome topology is not predictive of transcriptional output genome-wide (*54*). Indeed, our data argue that changes in chromatin topology are unlikely to underpin the molecular pathology of expanded CAG/CTG repeat disorders.

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# 249 Materials and Methods

#### 250 Cell lines

All LCLs were obtained from the Coriell Institute for Medical Research Cell Repository. They were 251 grown at 37°C and 5% CO<sub>2</sub> in RPMI 1640 medium supplemented with 15% FBS, 2 mM L-252 glutamine, and 1% penicillin-streptomycin. Cells were counted and passaged every 3-4 days 253 depending on cell density. The GFP(CAG)<sub>270</sub> and GFP(CAG)<sub>15</sub> lines were previously 254 characterized (41). They were maintained at 37°C and 5% CO<sub>2</sub> in DMEM with Glutamax, 1% 255 penicillin-streptomycin, 15  $\mu$ g mL<sup>-1</sup> blasticidine, and 150  $\mu$ g mL<sup>-1</sup> hygromycin. Transcription of the 256 GFP mini-gene was activated by culturing GFP(CAG)<sub>n</sub> cells with doxycycline at a final 257 concentration of 2  $\mu$ g mL<sup>-1</sup> for five days. 258

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# 260 Repeat length determination and small-pool PCR

Genomic DNA was isolated from each LCL using the NucleoSpin Tissue kit (Macherey-Nagel). 261 PCR products containing the CTG repeats from *DMPK* were amplified with primers oVIN-1252 262 and oVIN-1251 (Table S3). PCR products with the CAG repeats from HTT were produced with 263 primers oVIN-1333 and oVIN-1334 (Table S3). For normal-length alleles, several PCRs were set 264 up with Mango Taq (Bioline) and the products were gel-extracted and Sanger-sequenced with the 265 same primers used for the amplification. For expanded alleles, small-pool PCR was performed 266 based on a previously described protocol (55). Briefly, the same primers were used for the 267 amplification of expanded DMPK and HTT alleles with 1 ng of genomic DNA per PCR. The 268 products were run on an agarose gel and transferred to a nylon membrane. An oligo made up of 269 10 CAGs was used to obtain a radioactive probe used for the visualization of the expanded alleles. 270 The number of repeats reported here is an estimation of the modal number of repeats. 271

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# **Bisulfite sequencing**

Bisulfite sequencing was performed according to a previously described method (*40*). Bisulfite conversion was performed with the EZ DNA Methylation kit (Zymo Research, D5001) using the standard protocol. 200 ng of genomic DNA were used for bisulfite conversion at 50°C for 12 hours.

Bisulfite-converted DNA was desulphonated, eluted, and immediately used for nested and hemi-277 nested PCR amplification of the upstream and downstream CTCF binding sites respectively using 278 previously described primers (40). 50 ng of bisulfite-converted DNA were used for the first PCR, 279 and 3 µL of the products were used for the second PCR. The final amplicons were purified with 280 the NucleoSpin PCR Clean-up kit (Macherey-Nagel, 740609) and used for 2x250 bp paired-end 281 sequencing (Illumina). The primers used for both rounds of PCR are found in Table S3. 282 Sequencing reads were preprocessed using TrimGalore (github.com/FelixKrueger/TrimGalore) 283 with the following parameters: -q 20 --length 20 --paired. Reads were aligned using QuasR 284 (Gaidatzis et al 2015, PMID 25417205) to human reference (GRCh38) DNA sequences 285 corresponding to the amplified PCR products. DNA methylation calls for each CpG were extracted 286 using the qMeth() function in QuasR. DNA methylation frequencies were calculated as methylated 287 CpGs / total covered CpG x 100 and plotted in R. 288

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# 290 CTCF ChIP-qPCR

Chromatin immunoprecipitation was performed according to the Diagenode AutoiDeal ChIP-291 292 gPCR kit (Diagenode, C01010181) standard protocol. Samples of 6x10<sup>6</sup> cells were sonicated using the Diagenode Bioruptor Pico (Diagenode, B01060010), with 10 cycles of 30 sec "on" and 293 30 sec "off". Correct DNA fragmentation was verified by agarose gel electrophoresis. 294 Immunoprecipitation was performed with  $4x10^6$  cells using a CTCF antibody (Diagenode, 295 C15410210) and the Diagenode IP Star Compact Automated System robot (Diagenode 296 B03000002). Results were analyzed using the StepOnePlus qPCR by Applied Biosystems 297 (ThermoFisher Scientific, 4376357) with Applied Biosystems SYBR<sup>™</sup> Green PCR Mastermix 298 (ThermoFisher Scientific, 4309155). Primer sequences for gPCR are listed in Table S3. 299

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# **4C-seq**

4C library preparation was performed based on a previously described protocol (56). 10<sup>7</sup> cells 302 were crosslinked in 2% formaldehyde for 10 minutes at room temperature and guenched with 303 glycine to a final concentration 0.13 M. Cells were lysed for 15 minutes on ice in lysis buffer (50 304 mM Tris-HCl pH 7.5, 150 mM NaCl, 5 mM EDTA, 0.5% NP-40, 1% Triton X-100). The first 305 digestion was performed at 37°C overnight with 600 U of DpnII (NEB) followed by DpnII heat 306 inactivation. The first ligation was performed at 16°C overnight with 50 U T4 DNA ligase (Thermo 307 Scientific) in 7 mL. Ligated samples were de-crosslinked with 30 µL Proteinase K (10 mg/mL) at 308 309 65°C overnight followed RNAse A treatment and phenol-chloroform purification. The second digestion was performed with 50 U of Bfal (NEB) at 37°C overnight followed by Bfal heat 310

inactivation. The second ligation was performed at 16°C overnight with 100 U T4 DNA ligase in
 14 mL. 4C template samples were purified with phenol-chloroform and QIAquick PCR purification
 kit columns. 4C libraries were generated by amplifying 1 µg total of purified 4C template with 4C
 viewpoint primers, pooling reactions and purifying the PCR products with AMPure XP beads to
 exclude products less than 130 bp. Single-end sequencing of pooled 4C libraries was performed
 on Illumina HiSeg 2500.

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# 318 4C-seq data analysis

Demultiplexing and mapping were performed using the BBCF HTSstation (57) according to (58). 319 The 4C fragments surrounding the viewpoints (±2.5 kb) were excluded from the rest of the 320 analysis. The demultiplexed reads were mapped to the human genome GRCh38 using bowtie2 321 (v 2.2) (59). Fragment counts were obtained using FourCSeq (v1.18.0) (38). The number of 322 mapped reads for each sample is found in Table S2. For plotting the data, fragment counts were 323 normalized (reads per million) and smoothed with a running mean (window size = 5 fragments). 324 The smoothed and normalized fragment counts were averaged among replicates of the same 4C 325 326 library samples and visualized with gFeatBrowser (http://www.gfeatbrowser.com). For the FourCSeq analysis, we defined significant interactions as fragments with a z-score equal to or 327 greater than 1.96 and a false-discovery rate of 0.1, using the following parameters: minCount=20 328 and fitFun="distFitMonotone" in getZScores function; zScoreThresh=1.96, fdrThresh = 0.1 in 329 addPeaks function. 4C-ker (v0.0.0.9000) uses a Hidder-Markov Model that accounts for 330 differences in coverage near 4C viewpoints to determine three types of domains: high-interacting, 331 low-interacting, and non-interacting domains. For each viewpoint, we used k=5 in the 332 nearBaitAnalysis function, and plotted the high-interacting regions. The difference between 333 significant interactions called with FourCSeg and 4C-ker is expected given that FourCSeg usually 334 identifies "peaks" of significantly interacting regions whereas 4C-ker identifies regions (60). The 335 analysis of differential interactions was performed with the differential Analysis function of 4C-ker, 336 which is based on the DESeq2 framework, using default parameters (including a p-value 337 threshold of 0.5). The UN-A cell line was used as the reference condition for all comparisons. 338

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#### 340 **4C SNP data analysis**

We called SNPs from 4C-seq data from the DM1-A and HD-B patient-derived samples using GATK (v3.7.0) (61) and samtools/bcftools (v1.5 and v1.4.1 respectively) (62). Biallelic SNPs located within a 1 Mb region of the 3' end of *DMPK*, the 5' end of *HTT*, and the *ACTA1* 4C viewpoint were selected. Among these, only SNPs that could unambiguously inform which

parental allele they came from were retained for downstream analysis. This required a 345 homozygous genotype in at least one of the parents' cell lines. We validated a subset of these 346 SNPs by PCR amplification of the genomic region encompassing the variants followed by library 347 preparation and paired-end sequencing with Illumina Miseg nano 2 x 250 bp. We then analyzed 348 the sequencing coverage at allele-specific SNP positions from the 4C-seq data from DM1-A and 349 HD-B patient cell lines. For samples with at least 10 reads per SNP position, we counted the 350 number of times the expanded allele had more mapped reads than the normal allele. We applied 351 an exact binomial test to statistically assess whether the proportion of cases where the expanded 352 allele had more reads than the normal allele was significantly different to 0.5, which represented 353 the null hypothesis of no allelic bias. 354

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# 356 Targeted Locus Amplification

Targeted locus amplification was performed based on a previously described protocol (43).  $10^7$ 357 cells were crosslinked in 2% formaldehyde for 10 minutes at room temperature and guenched 358 with glycine to a final concentration of 275 mM. Cell were lysed for 5 minutes at room temperature 359 360 in lysis buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 5 mM EDTA, 0.5% NP-40, 1% Triton X-100). Crosslinked samples were digested at 37°C overnight with 400 U NIaIII (NEB) followed by 361 NIaIII inactivation. Samples were ligated at room temperature for 2 hours with 20 U T4 DNA ligase 362 (Thermo Scientific) in 500 µL. Ligated samples were de-crosslinked with 5 µL Proteinase K (10 363 mg/mL) at 65°C overnight followed by RNAse A treatment and phenol-chloroform purification. 364 Samples then digested overnight at 37°C with 50 U Nspl (NEB) followed by Nspl inactivation. The 365 second ligation was performed overnight at 16°C with 100 U T4 DNA ligase in 14 mL. TLA 366 circularized templates were purified with QIAquick PCR purification kit columns. TLA libraries 367 were generated by amplifying 800 ng of purified TLA template with TLA viewpoint primers. Paired-368 end sequencing of pooled TLA libraries was performed on Illumina HiSeq 4000. Reads were 369 mapped using a custom TLA analysis pipeline using the BWA mapping software (v0.7.17) (63). 370 First, reads were mapped to the human genome GRCh38. Then, unaligned sequences are 371 digested *in-silico* with the NIaIII restriction site and remapped to the genome. The combined 372 373 mapping results were used to determine the integration site.

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#### 375 **References**

1. J. H. Gibcus, J. Dekker, The hierarchy of the 3D genome. *Mol. Cell.* **49**, 773–782 (2013).

 W. A. Bickmore, B. van Steensel, Genome architecture: domain organization of interphase chromosomes. *Cell.* **152**, 1270–1284 (2013).

- A. Pombo, N. Dillon, Three-dimensional genome architecture: players and mechanisms. *Nat. Rev. Mol. Cell Biol.* 16, 245–257 (2015).
- D. G. Lupiáñez, M. Spielmann, S. Mundlos, Breaking TADs: How Alterations of Chromatin Domains Result in Disease. *Trends Genet.* 32, 225–237 (2016).
- M. Spielmann, D. G. Lupiáñez, S. Mundlos, Structural variation in the 3D genome. *Nat. Rev. Genet.* **19**, 453–467 (2018).
- V. Dion, J. H. Wilson, Instability and chromatin structure of expanded trinucleotide repeats. *Trends Genet.* 25, 288–297 (2009).
- A. López Castel, J. D. Cleary, C. E. Pearson, Repeat instability as the basis for human diseases
   and as a potential target for therapy. *Nat. Rev. Mol. Cell Biol.* **11**, 165–170 (2010).
- 389 8. H. T. Orr, H. Y. Zoghbi, Trinucleotide repeat disorders. *Annu. Rev. Neurosci.* **30**, 575–621 (2007).
- C. Sandi, M. Sandi, S. A. Virmouni, S. Al-Mahdawi, M. A. Pook, Epigenetic-based therapies for
   Friedreich ataxia. *Front Genet.* 5 (2014), doi:10.3389/fgene.2014.00165.
- 392 10. D. Colak *et al.*, Promoter-Bound Trinucleotide Repeat mRNA Drives Epigenetic Silencing in Fragile
   393 X Syndrome. *Science*. 343, 1002–1005 (2014).
- 11. E. Tabolacci, P. Chiurazzi, Epigenetics, fragile X syndrome and transcriptional therapy. *Am. J. Med. Genet. A.* 161A, 2797–2808 (2013).
- P. K. Chan *et al.*, Heterochromatinization induced by GAA-repeat hyperexpansion in Friedreich's ataxia can be reduced upon HDAC inhibition by vitamin B3. *Hum. Mol. Genet.* 22, 2662–2675 (2013).
- A. López Castel *et al.*, Expanded CTG repeat demarcates a boundary for abnormal CpG
   methylation in myotonic dystrophy patient tissues. *Hum. Mol. Genet.* 20, 1–15 (2011).
- 401
   14. D. Kumari, K. Usdin, The distribution of repressive histone modifications on silenced FMR1 alleles 402 provides clues to the mechanism of gene silencing in fragile X syndrome. *Hum. Mol. Genet.* 19, 403 4634–4642 (2010).
- 404
   405
   15. D. Kumari, K. Usdin, Chromatin remodeling in the noncoding repeat expansion diseases. *J. Biol. Chem.* 284, 7413–7417 (2009).
- 406
   407
   408 16. D. H. Cho *et al.*, Antisense transcription and heterochromatin at the DM1 CTG repeats are constrained by CTCF. *Mol. Cell.* 20, 483–489 (2005).
- 408
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   18. A. J. Verkerk *et al.*, Identification of a gene (FMR-1) containing a CGG repeat coincident with a
   411
   breakpoint cluster region exhibiting length variation in fragile X syndrome. *Cell.* 65, 905–914 (1991).
- 412
   19. Y. H. Fu, D. Kuhl, A. Pizzuti, M. Pieretti, J. S. Sutcliffe, Variation of the CGG repeat at the fragile X site results in genetic instability: resolution of the Sherman paradox. *Cell.* 67, 1047–1058 (1991).
- 414 20. I. Oberlé *et al.*, Instability of a 550-base pair DNA segment and abnormal methylation in fragile X
   415 syndrome. *Science*. **252**, 1097–1102 (1991).
- 416
   417 J. P. Fryns, A. Kleczkowska, E. Kubien, H. Vandenberghe, Cytogenetic Findings in Moderate and 417 Severe Mental-Retardation - a Study of an Institutionalized Population of 1991 Patients. *Acta* 418 *Paediatr Scand Suppl.* **313**, 1–& (1984).
- 419 22. V. Campuzano *et al.*, Friedreich's ataxia: autosomal recessive disease caused by an intronic GAA

- 420 triplet repeat expansion. *Science*. **271**, 1423–1427 (1996).
- 421
   423. N. Gheldof, T. M. Tabuchi, J. Dekker, The active FMR1 promoter is associated with a large domain 422 of altered chromatin conformation with embedded local histone modifications. *Proc. Natl. Acad. Sci.* 423 *U.S.A.* 103, 12463–12468 (2006).
- 424 24. R. Pietrobono *et al.*, Molecular dissection of the events leading to inactivation of the FMR1 gene.
   425 *Hum. Mol. Genet.* 14, 267–277 (2005).
- 426
  425. B. Coffee, F. P. Zhang, S. Ceman, S. T. Warren, D. Reines, Histone modifications depict an
  427 aberrantly heterochromatinized FMR1 gene in fragile X syndrome. *Am. J. Hum. Genet.* **71**, 923–
  428 932 (2002).
- 429
   430
   26. B. Coffee, F. Zhang, S. T. Warren, D. Reines, Acetylated histones are associated with FMR1 in normal but not fragile X-syndrome cells. *Nat. Genet.* 22, 98–101 (1999).
- 431
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  28. S. Al-Mahdawi *et al.*, The Friedreich ataxia GAA repeat expansion mutation induces comparable epigenetic changes in human and transgenic mouse brain and heart tissues. *Hum. Mol. Genet.* **17**, 735–746 (2008).
- 437
   438
   29. E. Soragni *et al.*, Long intronic GAA\*TTC repeats induce epigenetic changes and reporter gene silencing in a molecular model of Friedreich ataxia. *Nucleic Acids Res.* 36, 6056–6065 (2008).
- 30. E. Greene, L. Mahishi, A. Entezam, D. Kumari, K. Usdin, Repeat-induced epigenetic changes in intron 1 of the frataxin gene and its consequences in Friedreich ataxia. *Nucleic Acids Res.* 35, 3383–3390 (2007).
- 442 31. J. H. Sun *et al.*, Disease-Associated Short Tandem Repeats Co- localize with Chromatin Domain
   443 Boundaries. *Cell*, 1–31 (2018).
- 32. J. R. Brouwer, A. Huguet, A. Nicole, A. Munnich, G. Gourdon, Transcriptionally Repressive
   Chromatin Remodelling and CpG Methylation in the Presence of Expanded CTG-Repeats at the
   DM1 Locus. *J Nucleic Acids*. 2013, 567435–16 (2013).
- 33. T. R. Klesert *et al.*, Mice deficient in Six5 develop cataracts: implications for myotonic dystrophy.
   *Nat. Genet.* 25, 105–109 (2000).
- 34. A. D. Otten, S. J. Tapscott, Triplet repeat expansion in myotonic dystrophy alters the adjacent chromatin structure. *Proc. Natl. Acad. Sci. U.S.A.* 92, 5465–5469 (1995).
- 35. X. A. Su, V. Dion, S. M. Gasser, C. H. Freudenreich, Regulation of recombination at yeast nuclear pores controls repair and triplet repeat stability. *Genes Dev.* 29, 1006–1017 (2015).
- 36. H. J. G. van de Werken *et al.*, Robust 4C-seq data analysis to screen for regulatory DNA
  interactions. *Nat. Methods.* 9, 969–972 (2012).
- 455 37. M. Simonis *et al.*, Nuclear organization of active and inactive chromatin domains uncovered by 456 chromosome conformation capture-on-chip (4C). *Nat. Genet.* **38**, 1348–1354 (2006).
- 457 38. F. A. Klein *et al.*, FourCSeq: analysis of 4C sequencing data. *Bioinformatics*. **31**, 3085–3091 (2015).
- 39. R. Raviram *et al.*, 4C-ker: A Method to Reproducibly Identify Genome-Wide Interactions Captured by 4C-Seq Experiments. *PLoS Comput Biol.* **12**, e1004780–23 (2016).
- 461 40. L. Barbé *et al.*, CpG Methylation, a Parent-of-Origin Effect for Maternal-Biased Transmission of

462	Congenital M	Intonic Dvg	strophy Am	I Hum	Conot 100	188_505 (2	017)
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- 463
   41. C. Cinesi, L. E. N. Aeschbach, Bin Yang, V. Dion, Contracting CAG/CTG repeats using the CRISPR-Cas9 nickase. *Nat Commun.* 7, 1–10 (2016).
- 465 42. B. A. Santillan, C. Moye, D. Mittelman, J. H. Wilson, GFP-based fluorescence assay for CAG 466 repeat instability in cultured human cells. *PLoS ONE*. **9**, e113952 (2014).
- 467 43. P. J. P. de Vree *et al.*, Targeted sequencing by proximity ligation for comprehensive variant detection and local haplotyping. *Nat. Biotechnol.* **32**, 1019–1025 (2014).
- 469 44. A. R. Barutcu, B. J. Blencowe, J. L. Rinn, Differential contribution of steady-state RNA and active 470 transcription in chromatin organization. *EMBO Rep.*, e48068 (2019).
- 471
  45. D. Racko, F. Benedetti, J. Dorier, A. Stasiak, Transcription-induced supercoiling as the driving force
  472 of chromatin loop extrusion during formation of TADs in interphase chromosomes. *Nucleic Acids*473 *Res.* 46, 1648–1660 (2018).
- 474 46. T. B. Le, M. T. Laub, Transcription rate and transcript length drive formation of chromosomal interaction domain boundaries. *EMBO J.* **35**, 1582–1595 (2016).
- 476 **47**. F. Le Dily *et al.*, Distinct structural transitions of chromatin topological domains correlate with coordinated hormone-induced gene regulation. *Genes Dev.* **28**, 2151–2162 (2014).
- 478
   48. M. Merkenschlager, E. P. Nora, CTCF and Cohesin in Genome Folding and Transcriptional Gene
   479
   Regulation. *Annu Rev Genomics Hum Genet.* **17**, 17–43 (2016).
- 480
  49. R. Adihe Lokanga, X.-N. Zhao, A. Entezam, K. Usdin, X inactivation plays a major role in the
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- 50. D. Wohrle, U. Salat, H. Hameister, W. Vogel, P. Steinbach, Demethylation, reactivation, and
   destabilization of human fragile X full-mutation alleles in mouse embryocarcinoma cells. *Am. J. Hum. Genet.* 69, 504–515 (2001).
- 486 51. C. T. McMurray, Mechanisms of trinucleotide repeat instability during human development. *Nat.* 487 *Rev. Genet.* **11**, 786–799 (2010).
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   53. A. R. Barutcu, P. G. Maass, J. P. Lewandowski, C. L. Weiner, J. L. Rinn, A TAD boundary is
   492 preserved upon deletion of the CTCF-rich Firre locus. *Nat Commun.* 9, 1444 (2018).
- 493 54. Y. Ghavi-Helm *et al.*, Highly rearranged chromosomes reveal uncoupling between genome
   494 topology and gene expression. *Nat. Genet.* 144, 327 (2019).
- 495
  495 55. V. Dion, Y. Lin, L. Hubert, R. A. Waterland, J. H. Wilson, Dnmt1 deficiency promotes CAG repeat 496 expansion in the mouse germline. *Hum. Mol. Genet.* **17**, 1306–1317 (2008).
- 497 56. E. Splinter, E. de Wit, H. J. G. van de Werken, P. Klous, W. de Laat, Determining long-range
   498 chromatin interactions for selected genomic sites using 4C-seq technology: from fixation to
   499 computation. *Methods.* 58, 221–230 (2012).
- 500 57. F. P. A. David *et al.*, HTSstation: a web application and open-access libraries for high-throughput sequencing data analysis. *PLoS ONE*. **9**, e85879 (2014).
- 502 58. D. Noordermeer *et al.*, The dynamic architecture of Hox gene clusters. *Science*. **334**, 222–225 (2011).

- 504 59. B. Langmead, S. L. Salzberg, Fast gapped-read alignment with Bowtie 2. *Nat. Methods*. **9**, 357– 505 359 (2012).
- 506 60. C. Walter, D. Schuetzmann, F. Rosenbauer, M. Dugas, Benchmarking of 4C-seq pipelines based 507 on real and simulated data. *Bioinformatics*. **46**, e91. (2019).
- 508 61. A. McKenna *et al.*, The Genome Analysis Toolkit: a MapReduce framework for analyzing next-509 generation DNA sequencing data. *Genome Res.* **20**, 1297–1303 (2010).
- 62. H. Li, A statistical framework for SNP calling, mutation discovery, association mapping and
   population genetical parameter estimation from sequencing data. *Bioinformatics*. 27, 2987–2993
   (2011).
- 513 63. H. Li, R. Durbin, Fast and accurate long-read alignment with Burrows-Wheeler transform. 514 *Bioinformatics*. **26**, 589–595 (2010).
- 515

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522

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529

# 530 Author contributions

G.A.R.B. and V.D. designed the experiments, prepared the figures, and wrote the manuscript with input from all the authors. G.A.R.B. performed the experiments and data analysis. M.L. and G.A.R.B. performed the analysis of the 4C-Seq data. F.M. performed the SNP calling and G.A.R.B. performed the statistical analysis. J.Y.T. and A.C.M. helped with mapping the insertion site. L.V. and R.M. performed the CTCF ChIP-qPCR. G.A.R.B. performed the bisulfite sequencing experiments and T.B. analyzed the data. I.X. supervised G.A.R.B. together with V.D. and helped in designing the experiments.

538

# 539 **Competing interests**

- 540 The authors declare no competing interests.
- 541

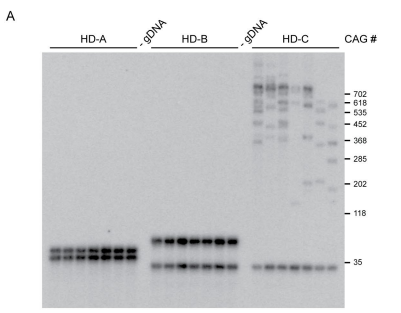
# 542 Data and material availability

543 All sequencing data underlying this study will be shortly deposited in the SRA of NCBI. All data

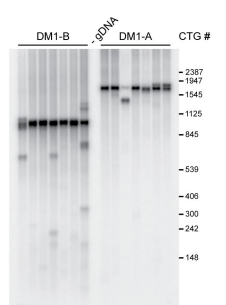
544 presented here are available from the corresponding author.

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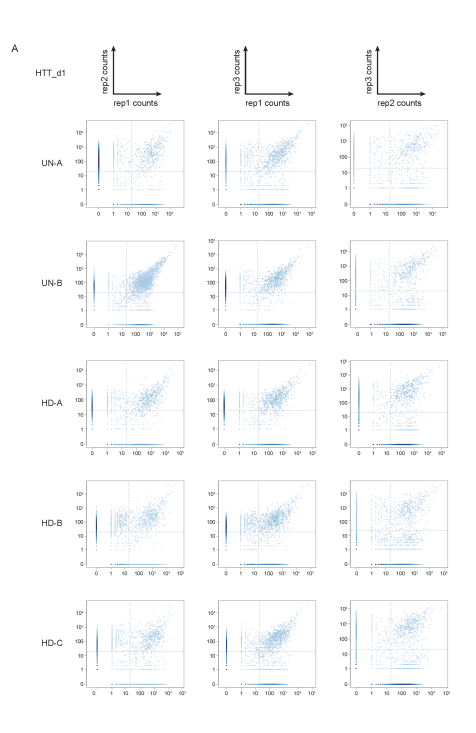
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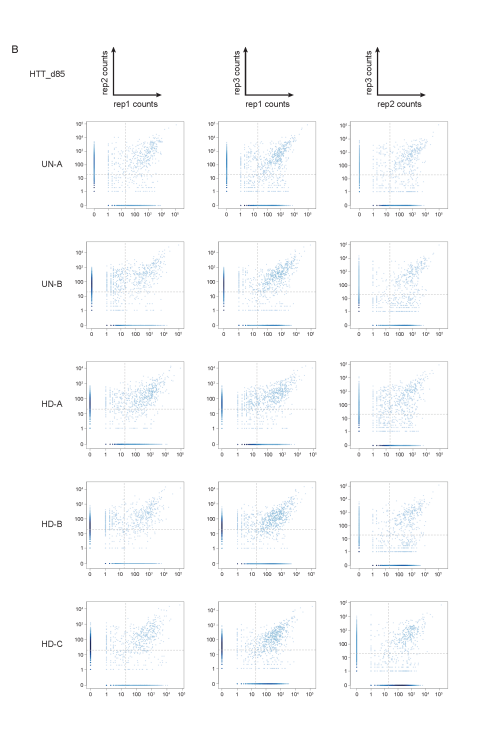
# 550 Fig. S1. Repeat sizes of HD and DM1 patient cell lines.

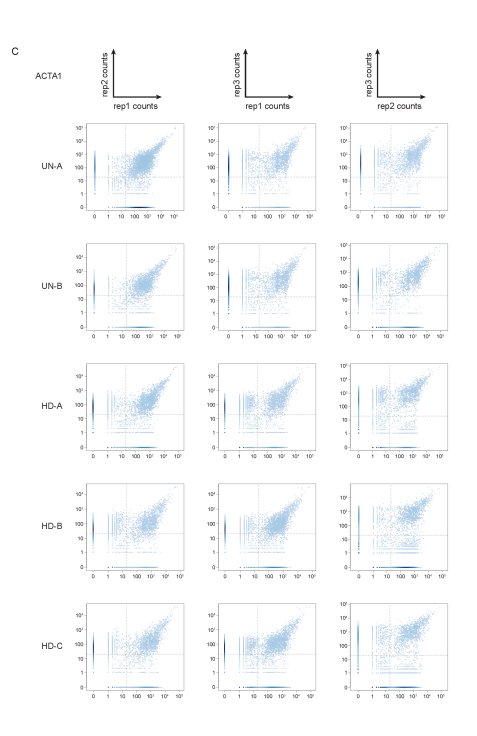
(A) Small-pool PCR of DNA isolated from HD-A, HD-B, and HD-C cells (shown: 1 ng gDNA per
 PCR). Lanes labeled "-gDNA" are control PCRs with no gDNA. (B) Small-pool PCR of DNA
 isolated from DM1-A and DM1-B cells (shown: 1 ng gDNA per PCR). Lanes labeled "-gDNA" are
 control PCRs with no gDNA.

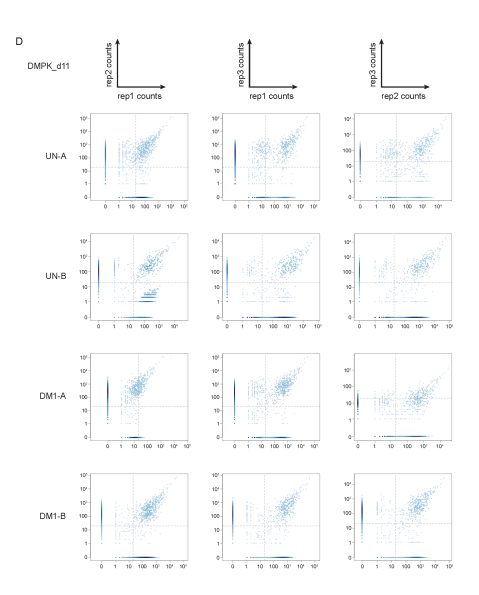
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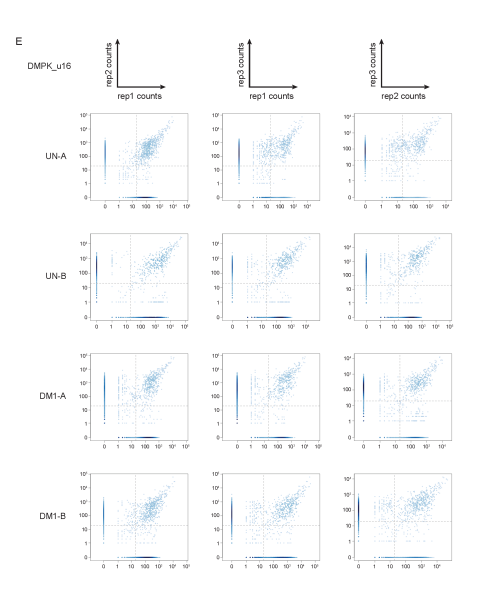
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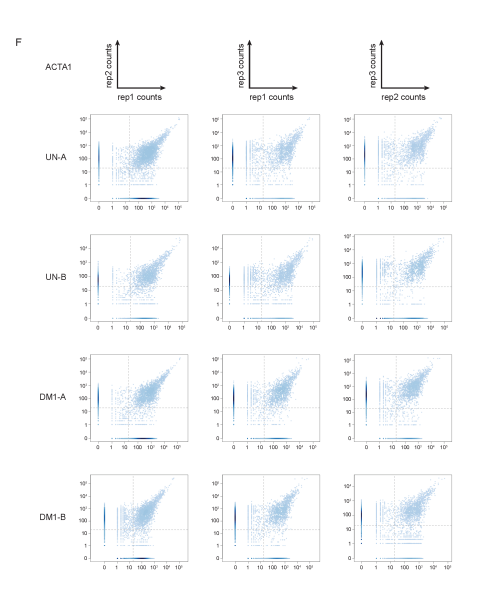


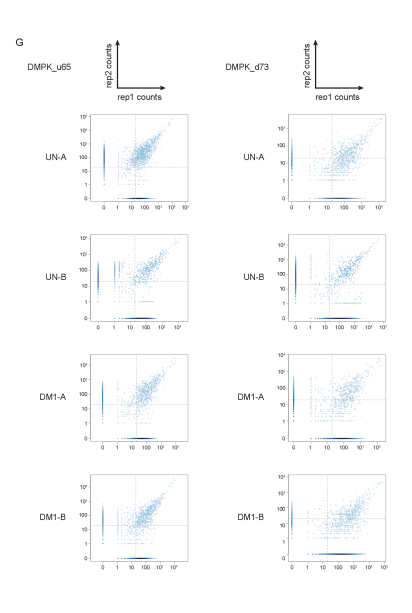


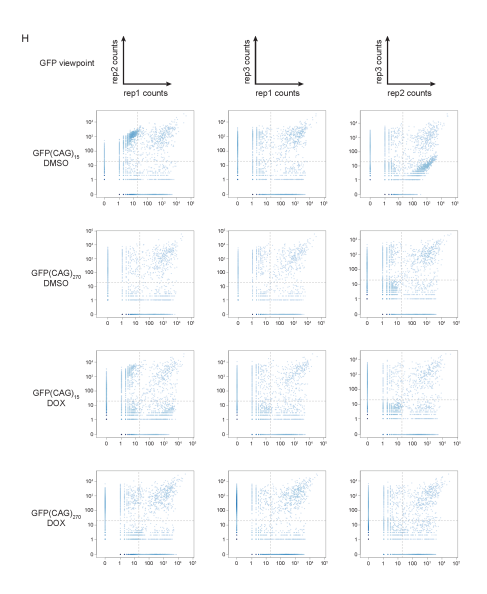








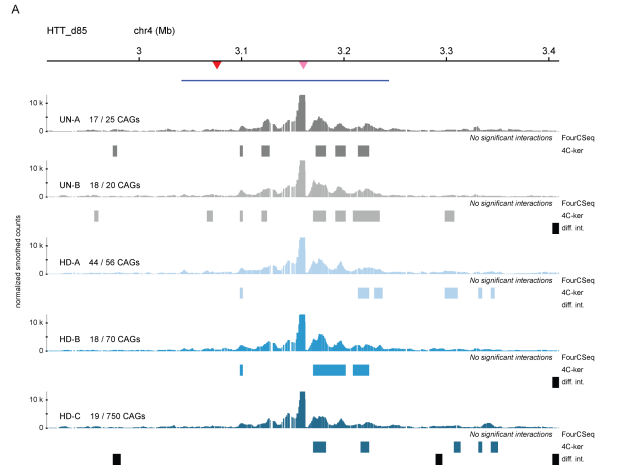




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# 566 Fig. S2. Correlation between 4C library replicates of the *HTT*, *DMPK*, *ACTA1*, and GFP 567 viewpoints.

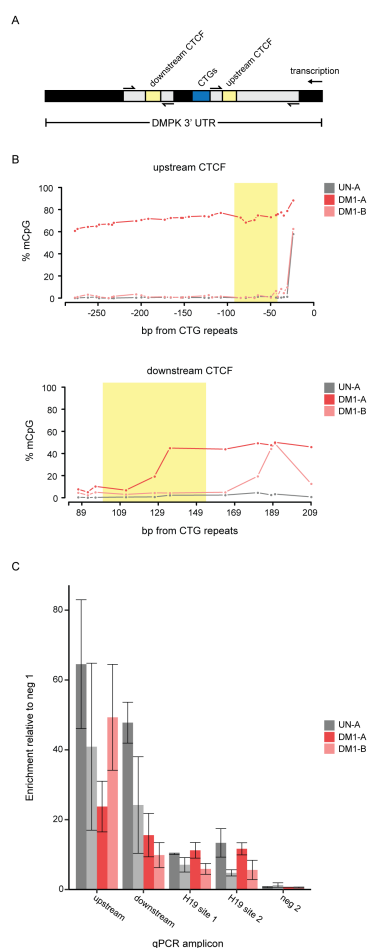
- 568 Scatter plots comparing the mapped read counts per 4C fragment between replicates from UN-
- A, UN-B, HD-A, HD-B, and HD-C cells (A-C); from UN-A, UN-B, DM1-A, and DM1-B cells (D-G); 569 and from GFP(CAG)<sub>15</sub> and GFP(CAG)<sub>270</sub> (H) separated by viewpoint: (A) HTT d1 viewpoint (1 570 kb downstream of the CAG repeat), (B) HTT d85 viewpoint (85 kb downstream of the CAG 571 repeat), (C) ACTA1 viewpoint, (D) DMPK d11 viewpoint (11 kb downstream of the CTG repeat), 572 (E) DMPK u16 viewpoint (16 kb upstream of the CTG repeat), (F) ACTA1 viewpoint, (G) 573 DMPK u65 and DMPK d73 viewpoints (65 kb upstream and 73 kb downstream of the CTG 574 repeat, respectively), and (H) GFP viewpoint in GFP(CAG)<sub>15</sub> and GFP(CAG)<sub>270</sub> cells with 575 transcription off (DMSO) or on (DOX). 576
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579 Fig. S3. Chromatin interactions of the HTT\_d85 viewpoint in unaffected and HD patient 580 cells.

(A) 4C-seq chromatin interaction profiles from the HTT\_d85 viewpoint (85 kb downstream of the
CAG repeat) in two unaffected (UN-A and UN-B) and three HD LCLs (HD-A, HD-B, and HD-C).
High-interacting regions were called using 4C-ker and significant interactions were called using
FourCSeq. Regions of differential interactions are marked with black bars below each 4C-seq
track and labeled as "diff. int.". The top blue bar represents the *HTT* gene. The triangles at the
top represent the location of the two *HTT* viewpoints.

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# Fig. S4. Heterochromatinization of expanded *DMPK* alleles in DM1 patient cells.

(A) Schematic of the 3' UTR of DMPK. The PCR amplicons used for bisulfite sequencing and CTCF ChIP-qPCR are indicated in grey. The CTCF binding sites are represented by the yellow boxes. (B) CpG methylation levels of two CTCF binding sites directly flanking the CTG repeats of DMPK using bisulfitesequencing (top: upstream site; bottom: downstream site). The specific CTCF binding sites are demarcated with yellow bars. (C) Chromatin immunoprecipitation of CTCF followed by qPCR at the cognate binding sites found in A and B. This was done in the UN-A, UN-B, DM1-A, and DM1-B LCLs (N=3 for all). CTCF was expected to be bound to the H19 gene, but not at two intergenic sites (neg 1 and neg 2). We normalized the enrichment at each site to neg 1. We found significant differences when no we normalized to the Neg2 site. The error bars represent the standard error of the mean.

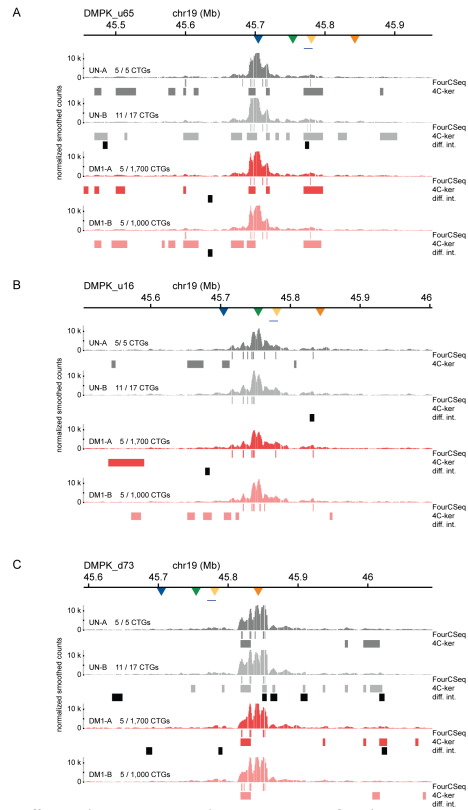


Fig. S5. Chromatin interactions at DMPK\_u65, DMPK\_u16, and DMPK\_d73 viewpoints in unaffected and DM1 patient cells.

(A) 4C-seq chromatin interaction profiles from the DMPK u65 viewpoint (65 kb upstream of the CAG repeat) in two unaffected (UN-A and UN-B) and two DM1 LCLs (DM1-A and DM1-B). Highinteracting regions were using called 4C-ker and significant interactions were called using FourCSeq. Regions of differential interactions are marked with black bars below each 4Cseq track and labeled as "diff. int.". The top blue bar represents the DMPK gene. The triangles at the top represent the location of the four DMPK viewpoints. (B) 4C-seq chromatin interaction profiles from the DMPK u16 viewpoint (16 kb upstream of the CAG repeat) in two

unaffected (UN-A and UN-B) and two DM1 LCLs (DM1-A and DM1-B). (C) 4C-seq chromatin
interaction profiles from the DMPK\_d73 viewpoint (73 kb downstream viewpoint in two unaffected
(UN-A and UN-B) and two DM1 LCLs (DM1-A and DM1-B).

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Table S1. Characteristics of cell lines used in this study.

Cell line	Code	Disease status <sup>1</sup>	Sex 1	Age at sampling (years) <sup>1</sup>	Repeat locus	# of repeats in wt allele <sup>2</sup>	Sizing method	# of repeats in expanded allele <sup>2</sup>	Sizing method
GM04604	UN-A	unaffected	Male	NA	DMPK	5	Sanger sequencing	5	Sanger sequencing
					HTT	17	Sanger sequencing	25	Sanger sequencing
GM02180	UN-B	unaffected	Female	51	DMPK	11	Sanger sequencing	17	Sanger sequencing
					HTT	18	Sanger sequencing	20	Sanger sequencing
GM06077	DM1-A	DM1	Female	4	DMPK	5	Sanger sequencing	1700	sp-PCR
GM04648	DM1-B	DM1	Male	23	DMPK	5	Sanger sequencing	1000	sp-PCR
GM02164	HD-A	HD	Male	58	HTT	44	sp-PCR	56	sp-PCR
GM03620	HD-B	HD	Female	29	HTT	18	Sanger sequencing	70	sp-PCR
GM14044	HD-C	HD	Male	16	HTT	19	Sanger sequencing	750 *	sp-PCR

<sup>1</sup> Information obtained from the Coriell Institute Human Genetic Cell Repository.

<sup>2</sup> For sp-PCR, the number of repeats is an estimation of the modal number of repeats.

NA: not available

sp-PCR: small pool-PCR

# Table S3. 4C-seq library statistics.

Sample	# of reads	# of mapped reads	% of total	# of reads mapped to cis chr	% of cis chr reads from total	# of reads mapped to valid 4C frags in cis chr	% of reads in 4C frags from total cis chr reads
DMPK_u65_GM04604_rep1	11'192'157	8'116'207	72.5	2'296'664	28.3	1'805'386	78.6
DMPK_u65_GM04604_rep2	1'688'853	1'035'697	61.3	606'020	58.5	513'147	84.7
DMPK_u65_GM02180_rep1	1'183'618	725'020	61.3	402'692	55.5	342'602	85.1
DMPK_u65_GM02180_rep2	1'461'840	888'805	60.8	466'817	52.5	406'381	87.1
DMPK_u65_GM06077_rep1	8'498'158	5'535'188	65.1	1'053'071	19	908'403	86.3
DMPK_u65_GM06077_rep2	1'586'296	966'005	60.9	574'031	59.4	515'578	89.8
DMPK_u65_GM04648_rep1	8'646'819	5'368'775	62.1	1'045'112	19.5	942'967	90.2
DMPK_u65_GM04648_rep2	2'167'574	1'221'577	56.4	668'154	54.7	588'914	88.1
DMPK_u16_GM04604_rep1	16'969'035	13'541'809	79.8	2'368'499	17.5	1'502'637	63.4
DMPK_u16_GM04604_rep2	8'471'505	1'910'469	22.6	611'665	32	391'201	64
DMPK_u16_GM04604_rep3	5'023'436	4'174'649	83.1	325'026	7.8	214'817	66.1
DMPK_u16_GM02180_rep1	4'967'776	3'272'865	65.9	1'156'773	35.3	807'405	69.8
DMPK_u16_GM02180_rep2	9'791'581	6'074'072	62	2'791'710	46	1'918'545	68.7
DMPK_u16_GM02180_rep3	3'499'397	2'404'710	68.7	743'880	30.9	494'040	66.4
DMPK_u16_GM06077_rep1	44'300'878	31'633'701	71.4	4'230'357	13.4	3'136'434	74.1
DMPK_u16_GM06077_rep2	7'516'556	2'615'231	34.8	791'991	30.3	512'797	64.7
DMPK_u16_GM06077_rep3	3'344'272	2'426'791	72.6	693'947	28.6	436'064	62.8

DMPK_u16_GM04648_rep1	14'372'890	11'367'448	79.1	1'616'380	14.2	1'080'070	66.8
DMPK_u16_GM04648_rep2	5'456'784	1'941'577	35.6	874'656	45	560'803	64.1
DMPK_u16_GM04648_rep3	8'354'943	5'260'739	63	2'365'753	45	1'549'651	65.5
DMPK_d11_GM04604_rep1	23'059'825	10'787'846	46.8	2'388'136	22.1	1'782'296	74.6
DMPK_d11_GM04604_rep2	8'042'866	717'825	8.9	355'974	49.6	251'617	70.7
DMPK_d11_GM04604_rep3	6'716'905	4'244'006	63.2	1'416'961	33.4	988'977	69.8
DMPK_d11_GM02180_rep1	4'296'953	866'253	20.2	442'187	51	330'779	74.8
DMPK_d11_GM02180_rep2	3'308'800	894'717	27	412'986	46.2	308'579	74.7
DMPK_d11_GM02180_rep3	7'282'955	4'517'085	62	2'175'116	48.2	1'518'060	69.8
DMPK_d11_GM06077_rep1	15'395'452	7'880'443	51.2	2'460'619	31.2	1'843'809	74.9
DMPK_d11_GM06077_rep2	774'586	109'054	14.1	56'680	52	41'488	73.2
DMPK_d11_GM06077_rep3	6'524'825	2'605'272	39.9	1'097'272	42.1	773'163	70.5
DMPK_d11_GM04648_rep1	8'560'029	4'972'802	58.1	1'413'824	28.4	979'700	69.3
DMPK_d11_GM04648_rep2	12'454'988	1'910'083	15.3	928'582	48.6	678'608	73.1
DMPK_d11_GM04648_rep3	8'259'430	3'846'132	46.6	1'529'463	39.8	1'063'240	69.5
DMPK_d73_GM04604_rep1	5'976'375	5'457'650	91.3	325'342	6	171'443	52.7
DMPK_d73_GM04604_rep2	4'710'733	2'209'035	46.9	816'530	37	507'446	62.1
DMPK_d73_GM02180_rep1	4'278'924	2'351'635	55	781'457	33.2	512'019	65.5
DMPK_d73_GM02180_rep2	6'211'002	4'265'102	68.7	1'463'307	34.3	918'581	62.8

DMPK_d73_GM06077_rep1	25'744'531	24'078'279	93.5	2'287'837	9.5	334'267	14.6
DMPK_d73_GM06077_rep2	5'923'808	2'670'770	45.1	871'927	32.6	565'589	64.9
DMPK_d73_GM04648_rep1	8'317'618	7'753'683	93.2	686'800	8.9	210'742	30.7
DMPK_d73_GM04648_rep2	13'833'430	8'816'046	63.7	3'351'872	38	2'184'265	65.2
ACTA1_d2_GM04604_rep1	17'462'688	9'831'359	56.3	5'908'533	60.1	4'186'550	70.9
ACTA1_d2_GM04604_rep2	20'016'202	13'201'152	66	9'429'955	71.4	6'749'626	71.6
ACTA1_d2_GM04604_rep3	14'370'212	9'150'078	63.7	6'493'921	71	4'854'805	74.8
ACTA1_d2_GM02180_rep1	4'445'588	2'894'160	65.1	2'047'454	70.7	1'448'565	70.7
ACTA1_d2_GM02180_rep2	9'724'309	6'455'364	66.4	4'490'505	69.6	3'238'325	72.1
ACTA1_d2_GM02180_rep3	14'471'204	9'623'984	66.5	6'832'538	71	5'111'636	74.8
ACTA1_d2_GM06077_rep1	11'677'329	7'881'675	67.5	4'502'685	57.1	3'148'013	69.9
ACTA1_d2_GM06077_rep2	20'390'273	13'386'906	65.7	9'414'304	70.3	6'751'808	71.7
ACTA1_d2_GM06077_rep3	6'528'425	4'468'502	68.4	3'101'465	69.4	2'186'147	70.5
ACTA1_d2_GM04648_rep1	28'697'528	18'150'768	63.2	9'891'236	54.5	7'190'802	72.7
ACTA1_d2_GM04648_rep2	6'247'015	4'135'124	66.2	2'850'483	68.9	2'025'028	71
ACTA1_d2_GM04648_rep3	7'615'653	5'226'254	68.6	3'625'835	69.4	2'493'472	68.8
ACTA1_d2_GM02164_rep1	4'982'938	3'226'684	64.8	2'262'970	70.1	1'598'786	70.6
ACTA1_d2_GM02164_rep2	14'175'182	9'482'669	66.9	6'432'203	67.8	4'689'799	72.9
ACTA1_d2_GM02164_rep3	8'985'756	6'149'600	68.4	4'314'765	70.2	3'032'024	70.3
ACTA1_d2_GM03620_rep1	3'392'689	2'252'717	66.4	1'594'521	70.8	1'125'432	70.6

ACTA1_d2_GM03620_rep2	10'153'899	6'911'566	68.1	4'682'012	67.7	3'412'228	72.9
ACTA1_d2_GM03620_rep3	10'314'868	6'898'449	66.9	4'788'393	69.4	3'452'374	72.1
ACTA1_d2_GM14044_rep1	3'864'340	2'522'916	65.3	1'719'045	68.1	1'218'110	70.9
ACTA1_d2_GM14044_rep2	13'566'864	8'987'115	66.2	6'029'854	67.1	4'498'420	74.6
ACTA1_d2_GM14044_rep3	10'730'254	7'441'130	69.3	5'128'011	68.9	3'821'275	74.5
HTT_d1_GM04604_rep1	4'212'566	2'457'650	58.3	1'404'970	57.2	1'103'125	78.5
HTT_d1_GM04604_rep2	10'749'708	1'832'850	17.1	888'698	48.5	639'666	72
HTT_d1_GM04604_rep3	5'921'707	3'530'799	59.6	2'010'544	56.9	1'556'448	77.4
HTT_d1_GM02180_rep1	2'641'610	1'678'631	63.5	975'311	58.1	687'885	70.5
HTT_d1_GM02180_rep2	4'261'764	2'775'669	65.1	1'331'970	48	972'583	73
HTT_d1_GM02180_rep3	6'713'195	3'878'698	57.8	2'112'225	54.5	1'644'256	77.8
HTT_d1_GM02164_rep1	1'547'564	1'006'519	65	550'667	54.7	378'050	68.7
HTT_d1_GM02164_rep2	6'426'843	4'114'843	64	2'049'276	49.8	1'475'404	72
HTT_d1_GM02164_rep3	4'561'062	2'649'147	58.1	1'529'524	57.7	1'141'625	74.6
HTT_d1_GM03620_rep1	2'799'272	1'795'212	64.1	1'031'035	57.4	704'440	68.3
HTT_d1_GM03620_rep2	5'475'190	3'487'162	63.7	1'558'114	44.7	1'137'993	73
HTT_d1_GM03620_rep3	7'098'155	3'663'907	51.6	2'006'314	54.8	1'576'201	78.6
HTT_d1_GM14044_rep1	2'595'315	1'668'384	64.3	945'665	56.7	659'221	69.7
HTT_d1_GM14044_rep2	6'212'554	3'953'292	63.6	1'666'102	42.1	1'189'773	71.4
HTT_d1_GM14044_rep3	5'199'603	2'755'357	53	1'459'570	53	1'128'283	77.3

HTT_d85_GM04604_rep1	3'626'944	2'088'253	57.6	1'300'447	62.3	850'403	65.4
HTT_d85_GM04604_rep2	4'387'441	2'691'863	61.4	1'062'983	39.5	637'080	59.9
HTT_d85_GM04604_rep3	4'022'177	2'243'116	55.8	1'343'423	59.9	882'695	65.7
HTT_d85_GM02180_rep1	2'884'811	1'333'135	46.2	773'017	58	467'647	60.5
HTT_d85_GM02180_rep2	4'805'661	3'283'731	68.3	1'132'457	34.5	719'942	63.6
HTT_d85_GM02180_rep3	3'937'446	2'278'255	57.9	1'335'882	58.6	909'086	68.1
HTT_d85_GM02164_rep1	2'065'583	773'630	37.5	440'336	56.9	279'660	63.5
HTT_d85_GM02164_rep2	4'271'456	2'674'960	62.6	1'320'008	49.3	846'436	64.1
HTT_d85_GM02164_rep3	3'365'643	1'762'199	52.4	1'048'467	59.5	674'809	64.4
HTT_d85_GM03620_rep1	3'134'531	654'436	20.9	397'137	60.7	245'017	61.7
HTT_d85_GM03620_rep2	3'232'273	2'024'049	62.6	978'716	48.4	577'818	59
HTT_d85_GM03620_rep3	4'074'032	2'290'232	56.2	1'356'058	59.2	898'455	66.3
HTT_d85_GM14044_rep1	2'057'402	798'621	38.8	453'565	56.8	276'126	60.9
HTT_d85_GM14044_rep2	5'576'017	2'543'284	45.6	956'326	37.6	583'814	61
HTT_d85_GM14044_rep3	3'538'845	2'077'028	58.7	1'191'091	57.3	772'206	64.8
GFP_CAG_15_DMSO_rep1	7'048'343	2'833'417	40.2	1'700'523	60	1'278'717	75.2
GFP_CAG_15_DMSO_rep2	7'487'213	2'887'212	38.6	1'638'799	56.8	1'284'684	78.4
GFP_CAG_15_DMSO_rep3	6'936'377	2'769'339	39.9	1'464'325	52.9	1'172'916	80.1
GFP_CAG_15_DOX_rep1	11'304'727	4'562'245	40.4	2'182'322	47.8	1'705'839	78.2

GFP_CAG_15_DOX_rep2	8'452'926	3'124'374	37	1'540'855	49.3	1'198'057	77.8
GFP_CAG_15_DOX_rep3	7'631'549	2'833'020	37.1	1'466'250	51.8	1'157'955	79
GFP_CAG_270_DMSO_rep1	8'320'728	3'099'916	37.3	1'907'976	61.5	1'512'580	79.3
GFP_CAG_270_DMSO_rep2	6'840'505	2'694'631	39.4	1'575'886	58.5	1'229'060	78
GFP_CAG_270_DMSO_rep3	7'338'988	2'673'561	36.4	1'569'149	58.7	1'225'397	78.1
GFP_CAG_270_DOX_rep1	9'301'866	3'679'030	39.6	2'078'639	56.5	1'620'884	78
GFP_CAG_270_DOX_rep2	8'353'878	3'289'609	39.4	1'735'082	52.7	1'333'717	76.9
GFP_CAG_270_DOX_rep3	7'577'290	3'045'799	40.2	1'588'724	52.2	1'267'573	79.8

Table S4. PCR primers used in this study.	

Name	Application	Viewpoint / amplicon	Primer type	5' - 3' primer sequence	Reference
oVIN-1252	DMPK CTG repeats	-	Forward	CACTTTGCGAACCAACGATA	This study
oVIN-1251	DMPK CTG repeats	-	Reverse	GAGCGTGGGTCTCCGCCCAG	This study
oVIN-1333	HTT CAG repeats	-	Forward	CCGCTCAGGTTCTGCTTTTA	This study
oVIN-1334	HTT CAG repeats	-	Reverse	CAGGCTGCAGGGTTACCG	This study
oVIN-1180	4C-seq	ACTA1	Reading	AGAACAGGCCCTTGAGGGAT	This study
oVIN-1181	4C-seq	ACTA1	Non-reading	GGGCCATCTCCCACAGTTTG	This study
oVIN-1146	4C-seq	DMPK_d11	Reading	CCCAAAGTTGTCCCTCCTGG	This study
oVIN-1147	4C-seq	DMPK_d11	Non-reading	CCAGGCTGCCCAGTTTAACA	This study
oVIN-1158	4C-seq	DMPK_d73	Reading	GCTGAGAGATGGAGAAATGCAGA	This study
oVIN-1159	4C-seq	DMPK_d73	Non-reading	GTTGTTGGCTGAGGAAGGGG	This study
oVIN-1136	4C-seq	DMPK_u65	Reading	GACATTGCATGACAAGGGGGA	This study
oVIN-1137	4C-seq	DMPK_u65	Non-reading	ACTGACACAACACAAACCACAGA	This study
oVIN-1128	4C-seq	DMPK_u16	Reading	ACAAAGAGGCCTGTGATTGAGG	This study
oVIN-1129	4C-seq	DMPK_u16	Non-reading	CTGGGTGACCTGGGCTTCTA	This study
oVIN-1380	4C-seq	GFP	Reading	ATGGTGGCGACCGGTGGATC	This study
oVIN-1381	4C-seq	GFP	Non-reading	TCTTAGGCACCTTTGTCCTA	This study
oVIN-943	4C-seq	HTT_d1	Reading	TAGGCTTAGATGAGCAGATC	van de Werken, H. J. G., et al. (2012). Nature Methods, 9(10), 969–972.

oVIN-944	4C-seq	HTT_d1	Non-reading	GTCAGGCTTGCCAGAATAC	van de Werken, H. J. G., et al. (2012). Nature Methods, 9(10), 969–972.
oVIN-1030	4C-seq	HTT_d85	Reading	TGGTTGTACATTATGAGATC	van de Werken, H. J. G., et al. (2012). Nature Methods, 9(10), 969–972.
oVIN-1031	4C-seq	HTT_d85	Non-reading	AAAGGGTTAAATGTCCATCA	van de Werken, H. J. G., et al. (2012). Nature Methods, 9(10), 969–972.
oVIN-2582	TLA	GFP_d3	Reading	CACAAATCAGCGATTTCCAT	This study
oVIN-2583	TLA	GFP_d3	Non-reading	TCAGAAGCCATAGAGCCC	This study
oVIN-2590	TLA	GFP_u4	Reading	ACACAGGAAACAGCTATGAC	This study
oVIN-2589	TLA	GFP_u4	Non-reading	TCCACAGAATCAGGGGATAA	This study
oVIN-2103	Bisulfite seq PCR 1	upstream CTCF site	Forward	TGTYGTYGTTTTGGGTTGTATTG	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2110	Bisulfite seq PCR 1	upstream CTCF site	Reverse	CAACATTCCYGACTACAAAAACCCCTT	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2107	Bisulfite seq PCR 1	downstream CTCF site	Forward	TTYGGTTAGGTTGAGGTTT	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.

oVIN-2108	Bisulfite seq PCR 1	downstream CTCF site	Reverse	TTAACAAAAACAAATTTCCC	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2111	Bisulfite seq PCR 2	upstream CTCF site	Forward	TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGGTT GTATTGGGTTGGTGGTTTA	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2112	Bisulfite seq PCR 2	upstream CTCF site	Reverse	GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCT ACAAAAACCCTTYGAACCC	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2113	Bisulfite seq PCR 2	downstream CTCF site	Forward	TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGTAA ATTGTAGGTTTGGGAAG	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2114	Bisulfite seq PCR 2	downstream CTCF site	Reverse	GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGTT AACAAAAACAAATTTCCC	Barbé, L., et al. (2017). American Journal of Human Genetics, 100(3), 488–505.
oVIN-2443	CTCF ChIP qPCR	upstream CTCF site	Forward	AAGGACCCTTCGAGCCCC	This study
oVIN-2444	CTCF ChIP qPCR	upstream CTCF site	Reverse	AGTTCACAACCGCTCCGAG	This study
oVIN-2445	CTCF ChIP qPCR	downstream CTCF site	Forward	GCAAAAGCAAATTTCCCGAGT	This study
oVIN-2446	CTCF ChIP qPCR	downstream CTCF site	Reverse	AAACTGCAGGCCTGGGAA	This study
oVIN-2447	CTCF ChIP qPCR	H19 site 1	Forward	CCCATCTTGCTGACCTCAC	ChIPAb+ CTCF - ChIP Validated Antibody and Primer Set

oVIN-2448	CTCF ChIP qPCR	H19 site 1	Reverse	AGACCTGGGACGTTTCTGTG	ChIPAb+ CTCF - ChIP Validated Antibody and Primer Set
oVIN-2556	CTCF ChIP qPCR	H19 site 2	Forward	TGTGGATAATGCCCGACCTGAAGATCTG	Yao, H., et al., (2010). Genes & Development, 24(22), 2543– 2555. <u>http://</u> <u>doi.org/10.1101/</u> <u>gad.1967810</u>
oVIN-2557	CTCF ChIP qPCR	H19 site 2	Reverse	ACGGAATTGGTTGTAGTTGTGGAATCGGAAGT	Yao, H., et al., (2010). Genes & Development, 24(22), 2543– 2555. <u>http://</u> <u>doi.org/10.1101/</u> <u>gad.1967810</u>
oVIN-976	CTCF ChIP qPCR	negative 1	Forward	CCTTCAGAACCCTTCAGTGC	This study
oVIN-977	CTCF ChIP qPCR	negative 1	Reverse	ATCACACCTGTGGCTTCCTC	This study
oVIN-978	CTCF ChIP qPCR	negative 2	Forward	TGCTTTTCTAGGAGGTAGGTGTC	This study
oVIN-979	CTCF ChIP qPCR	negative 2	Reverse	AGCCCAGCAGAAAGGTCTTC	This study