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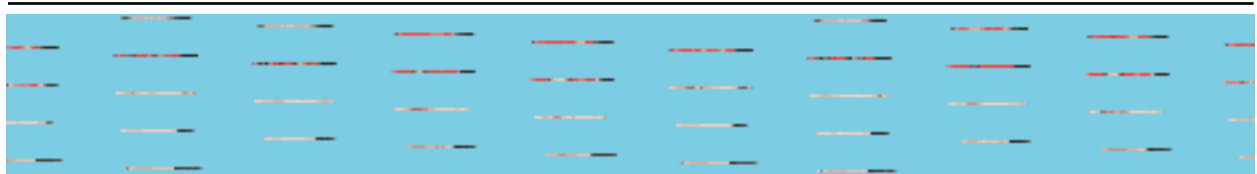
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# Loss of multi-level 3D genome organization during breast cancer progression

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## Abstract

Breast cancer entails intricate alterations in genome organization and expression. However, how three-dimensional (3D) chromatin structure changes in the progression from a normal to a breast cancer malignant state remains unknown. To address this, we conducted an analysis combining Hi-C data with lamina-associated domains (LADs), epigenomic marks, and gene expression in an *in vitro* model of breast cancer progression. Our results reveal that while the fundamental properties of topologically associating domains (TADs) are overall maintained, significant changes occur in the organization of compartments and subcompartments. These changes are closely correlated with alterations in the expression of oncogenic genes. We also observe a restructuring of TAD-TAD interactions, coinciding with a loss of spatial compartmentalization and radial positioning of the 3D genome. Notably, we identify a previously unrecognized interchromosomal insertion event, wherein a locus on Chromosome 8 housing the *MYC* oncogene is inserted into a highly active subcompartment on Chromosome 10. This insertion is accompanied by the formation of *de novo* enhancer contacts and activation of *MYC*, illustrating how structural genomic variants can alter the 3D genome during oncogenesis. In summary, our findings provide evidence for the loss of genome organization at multiple scales during breast cancer progression revealing novel relationships between genome 3D structure and oncogenic processes.

## 46 Introduction

47 The mammalian genome is folded into large-scale dynamic 3-dimensional (3D) chromatin  
48 conformations that provide a framework for regulated gene expression. In the interphase  
49 nucleus, chromosomes exist as territories (Rowley and Corces 2018; Cremer and Cremer  
50 2010) within which gene-rich and active A-compartments are segregated from more gene-  
51 poor and inactive B compartments (Misteli 2004; Crosetto and Bienko 2020; Lieberman-Aiden  
52 et al. 2009), both ranging in size from single promoters up to 2 megabase-pairs (Mbp) (Sood  
53 and Misteli 2022; Harris et al. 2023). Compartments consist of multiple subcompartments that  
54 differ in their chromosomal contact frequencies, enrichment in histone modifications and  
55 chromatin-binding proteins (Spracklin et al. 2023). At a resolution ranging from ~10-800  
56 kilobase-pairs (kbp), topologically associated domains (TADs) emerge as local, highly  
57 interacting domains (Rao et al. 2014; Nora et al. 2012; Dixon et al. 2012). TADs are further  
58 organized into sub-TADs and support the formation of internal smaller chromatin loops, which  
59 includes enhancer-promoter interactions (Berlivet et al. 2013). TADs and sub-TADs seemingly  
60 form independently of compartments (Rowley and Corces 2018). It was initially proposed that  
61 TADs provide a platform for the coordinated regulation of subsets of genes by enhancers  
62 located within the boundaries of the TAD. However, disrupting TAD structure minimally affects  
63 gene expression (Nora et al. 2017; Rao et al. 2017; Schwarzer et al. 2017), suggesting that  
64 other levels of genome organization have a more important regulatory role.

65 An intriguing possibility is that long-range TAD-TAD interactions, forming TAD cliques  
66 (Zhao et al. 2023; Pei et al. 2022; Li et al. 2022; Arnould et al. 2023; Paulsen et al. 2019),  
67 sculpt the 3D genome into distinct functional domains. TAD cliques are enriched in B  
68 compartments, and can gain or lose TADs during differentiation, generally in correlation with  
69 gene repression or activation, respectively (Paulsen et al. 2019). Additionally, the nuclear  
70 envelope imposes positional constraints on chromatin by anchoring heterochromatin through  
71 lamina-associated domains (LADs) of ~0.1-10 Mbp in size (Gonzalez-Sandoval and Gasser  
72 2016; Rønningen et al. 2015; van Steensel and Belmont 2017). Accordingly, genes located at  
73 the nuclear periphery are repressed or expressed at lower levels than genes localized towards  
74 the nuclear center (Misteli 2004; Crosetto and Bienko 2020). Integrating Hi-C and LAD data  
75 allows the generation of 3D structural genome models that accurately recapitulate the radial  
76 nuclear positions of TADs (Paulsen et al. 2017, 2018; Li et al. 2017; Boninsegna et al. 2022),  
77 and provide an enhanced understanding of the link between chromatin architecture and gene  
78 regulation in disease contexts (Paulsen et al. 2017).

79 Indeed, transcriptional dysregulation has been attributed to alterations in 3D genome  
80 organization in diseases, including cancer (Feng and Pauklin 2020; Osman et al. 2022). Yet,  
81 analysis of multiple cancers shows that TAD boundary deletions only rarely change gene  
82 expression, emphasizing their rare involvement in gene expression dysregulation in cancer  
83 (Akdemir et al. 2023). Even with shifts in A and B compartments between colon tumors and  
84 normal intestinal cells, TADs remain relatively unchanged (Johnstone et al. 2020),  
85 suggesting compartment switching significantly impacts gene expression control (Sood and  
86 Misteli 2022; Ibrahim and Mundlos 2020).

87 Breast cancer is the most common cancer in women. These cancers have been classified  
88 into five subtypes based on the expression of receptors for human epidermal growth factor,  
89 estrogen and progesterone, and clinical features (Sørlie et al. 2001). Among these subtypes,  
90 triple-negative breast cancers, which do not express any of these receptors, are the most  
91 aggressive (Bobbitt et al. 2023). Few studies have examined the chromatin architectural  
92 features of breast cancer cells or tissues (Dozmorov et al. 2023; Kim et al. 2022; Wang et al.

2022; Zhou et al. 2019). Nonetheless, one study reports that triple-negative breast cancer cells display the most severe disruption of the 3D genome, including weakening of TAD borders, loss of 3D chromatin interactions leading to fewer chromatin loops, and dynamic compartmental changes (Kim et al. 2022). However, how 3D chromatin organization is associated with the progression from a normal to a breast cancer malignant state remains unexplored. Here, we rely on an *in vitro* isogenic breast cancer progression model (Dawson et al. 1996; Santner et al. 2001) to investigate 3D genome organization changes in this process.

## Results

### TAD properties are conserved across breast cancer stages

We performed duplicate Hi-C experiments at three stages of an *in vitro* human breast cancer progression model, corresponding to a nonmalignant state (using MCF10A cells; “10A” from here on), a premalignant state (MCF10AT1 cells; “T1”) and a malignant tumorigenic state (MCF10Ca1a cells; “C1”). We obtained an average of >560 million pairwise interactions after filtering (Supplemental Table S1), supporting analyses in the range of 1-10 kbp bin resolution (Lajoie et al. 2015; Ay and Noble 2015). Filtering statistics and the relative fraction of intra- (cis) and inter- (trans) chromosomal contacts indicate high quality libraries (Lajoie et al. 2015), and samples show high reproducibility between replicates (Supplemental Fig. S1 and S2) (Yang et al. 2017). We identified and masked out regions with translocations and used ICE-normalized interactions which adjusts for bias resulting from copy-number alterations (see Methods) (Kim et al. 2022).

TAD comparisons across samples and replicates (Fig. 1A) show similar TAD numbers (~4000 TADs) and genomic size (~0.6 Mbp; Supplemental Fig. S3), TAD border insulation strength ( $R^2 = 0.77-1.00$ ; Supplemental Fig. S4) and intra-TAD contact frequencies ( $R^2 = 0.91-1.00$ ; Supplemental Fig. S5), and genomic positions of both TADs (Jaccard Index [JI] = 0.76-0.80; called at 50kb resolution), and sub-TADs (JI = 0.71-0.75; 10kb resolution) (Supplemental Fig. S6). Inspecting TADs surrounding known breast-cancer related genes (Lee and Muller 2010) reveals minor differences that were not consistently reproducible across replicates (Supplemental Fig. S7-S13). We conclude that TADs are overall similar across the three cell types and are therefore likely not the main oncogenic drivers in this progression model of breast cancer.

### Compartments and subcompartments engage in major switching events during breast cancer progression

To map differences in long-range genome contacts in our progression model, we searched for any A/B compartment switching, a property reported to be implicated in breast cancer (Barutcu et al. 2015). Most of the genome (2124 Mbp; ~78%) remains in the same compartment across stages (Fig. 1B). However, for the fraction that switches between stages, we note 385 Mbp (~14%) in 10A switching to either B (162 Mbp; ~6%) or A (222 Mbp; ~8%) compartments in T1 (Fig. 1B; Supplemental Table S2). Most of the switched compartments remain in their switched state in C1, but a fraction switches back to B (76 Mbp; ~3%) or A (61 Mbp; ~2%). The extent of switching observed in our model is comparable to that found in other cellular contexts involving changes in cell fate (Vilarrasa-Blasi et al. 2021; Dixon et al. 2015; Liu et al. 2021), indicating that compartment switching is a prominent feature of breast cancer progression.

139 We next used dHiC to identify subcompartments (Fig. 1C) (Chakraborty et al. 2022). We  
 140 predict four A subcompartments (A0-A3) and four B subcompartments (B0-B3), with A1, A2  
 141 and B1 being the most prominent and A0 and B0 the least prominent for all cell stages  
 142 (Supplemental Fig. S14; Supplemental Table S3). Overlaying a range of active and repressive  
 143 histone modifications from public ChIP-seq datasets onto the subcompartments reveals a  
 144 gradual enrichment of the ratio of heterochromatin to euchromatin features, going from B3 to  
 145 A3 subcompartments (Supplemental Fig. S15). From this analysis, B2 and B3 show features  
 146 of heterochromatin, with enrichment of repressive histone marks (H3K27me3) and Lamin B1.  
 147 Compared to the other subcompartments, A0 and B0 are the least enriched in active and  
 148 repressive histone marks respectively, raising the interesting possibility that they could be  
 149 more easily remodeled into other stronger euchromatic or heterochromatic subcompartments.  
 150 In contrast, A3 and B3 display the strongest active and inactive states respectively, which  
 151 would arguably be more resistant to configuration changes (Supplemental Fig. S15).

152 Indeed, we find that (i) weaker, intermediate subcompartments (A0, A1, B0, B1) undergo  
 153 more dynamic switches than more prominent and stronger subcompartments (A2, A3, B2, B3)  
 154 (Fig. 1D; Supplemental Fig. S16-18). However, (ii) switching between B0 and A0 ( $B0 \leftrightarrow A0$ ) is  
 155 minimal; whereas (iii)  $B0 \leftrightarrow A1$  is the most prominent inter-compartment switch (Fig. 1D;  
 156 Supplemental Fig. S16-18), indicating specific switching between weakly heterochromatic and  
 157 euchromatic states. A further characterization of paths of subcompartment switching across  
 158 the three stages shows that (iv) a similar fraction of 24-26% of subcompartments ends up in  
 159 either a more closed (and thus more B-like) subcompartment in C1, or a more open (A-like)  
 160 subcompartment. Further, reverting back to the initial 10A-subcompartment landscape in C1  
 161 rarely happens (Supplemental Fig. S19). Lastly, (v) switches mainly involve a transition  
 162 between consecutive subcompartment strengths, occurring with similar frequencies towards  
 163 both more open (e.g.  $A2 \rightarrow A3$ ) and less open (e.g.  $B2 \rightarrow B3$ ) subcompartments (Supplemental  
 164 Fig. S19). Comparing the switching levels between the two replicates shows a high degree of  
 165 consistency, with coefficients of determination ( $R^2$ ) ranging from 0.87 to 0.94 ( $P < 10^{-20}$ ; see  
 166 Supplemental Fig. S20).

167 To determine implications of subcompartment switching on 3D genome organization, we  
 168 generated 3D genome models (Paulsen et al. 2017) integrating Hi-C data with nuclear lamina-  
 169 chromatin interactions (LADs) mapped by ChIP-seq of lamin B1 in each cell type (see  
 170 Methods). The resulting models (Fig. 1E; Supplemental Fig. S21-23) display expected  
 171 genomic features where more gene-poor chromosomes are located more frequently towards  
 172 the nuclear periphery (Supplemental Fig. S24-26). Our 3D modeling of all three cell types  
 173 reveals major differences between 10A cells versus T1 and C1 cells. As expected, there is a  
 174 gradual increase in nuclear peripheral localization in subcompartments going from A3 towards  
 175 B3 in 10A cells. This, however, is less apparent in T1 and C1 cells (Fig. 1F). Moreover, A3  
 176 subcompartments significantly shift away from the nuclear center in T1 and C1 cells,  
 177 compared to 10A (Fig. 1F; Supplemental Table S4). B3 subcompartments appear to be slightly  
 178 shifted towards the nuclear interior, which concurs with a reduction in LAD coverage in T1 and  
 179 C1 cells (Supplemental Fig. S27). Taken together, our data indicate that subcompartment  
 180 switching characterizing the 10A-T1-C1 cell transition is accompanied by a partial loss of radial  
 181 disposition of chromatin.

182

### 183 **Subcompartment switching reflects changes in gene expression**

184 We next investigated whether the 3D genome structural changes observed were linked to  
 185 transcriptional changes. Differential gene expression analysis from RNA-seq data between  
 186 the cell types reveals 3180 differentially expressed (DE) genes between 10A and T1 cells, and

187 8362 DE genes between 10A and C1 cells ( $|\text{LFC}| > 0.5$ ;  $P < 0.01$ ; Supplemental Table S5).  
 188 Analysis of disease-linked gene set enrichment among DE genes in T1 and C1, reveals  
 189 “breast carcinoma” as the most enriched disease term, followed by other cancer types. “Organ  
 190 system benign neoplasm” is the second most associated term for T1, whereas this term is  
 191 absent for C1, reflecting the expected transcriptomic differences between the transformed (T1)  
 192 versus the malignant (C1) cell line (Supplemental Fig. S28). Overlaying the RNA-seq data  
 193 onto Hi-C-based subcompartments in all cell types reveals anticipated associations where  
 194 gene expression levels gradually increase from B3 to A3 (Fig. 2A, Supplemental Fig. S29),  
 195 reinforcing the validity of the subcompartment classification and the RNA-seq data in our  
 196 model.

197 Next, we analyzed up- and downregulated DEGs in conjunction with the corresponding  
 198 subcompartment switches. As expected, and as previously reported (Nagai et al. 2019; Liu et  
 199 al. 2021), this analysis shows that at specific chromosomal regions (Fig. 2B), or at genome-  
 200 wide levels (Fig. 2C-F), there are correlated patterns of subcompartment switching and  
 201 differential gene regulation. Inspecting the presence of downregulated genes in T1 relative to  
 202 10A reveals that B1(10A)→B2(T1) is the most frequent switch for these genes (Fig. 2C left  
 203 panel). Switching between B0/A0(10A)→B1(T1) or from A1(10A)→B1/B0/A0(T1) is enriched  
 204 relative to what is seen for non-differentially expressed genes (Fig. 2C middle panel) or  
 205 upregulated genes (Fig. 2C right panel).

206 To quantify whether up- and downregulated genes were associated with expected  
 207 subcompartment switches towards more open (or A-like), or closed (or B-like)  
 208 subcompartments, respectively, we analyzed enrichment of DE genes in each  
 209 subcompartment switch relative to a switch in opposite direction. This analysis revealed a  
 210 significant enrichment of downregulated genes in switching of B1→B2, B0→B1, A0→B1 and  
 211 A1→B0 between 10A and T1 (all  $P < 0.05$  [binomial test]; Supplemental Table S6). Upregulated  
 212 genes were significantly enriched in B1→A1, B1→B0, and A1→A3 (all  $P < 0.05$ ; Fig. 2C;  
 213 Supplemental Table S6). Comparing 10A and C1, downregulated genes were enriched in  
 214 A0→B2, A0→B1, B1→B2, and A1→A0 (all  $P < 0.05$ ), whereas upregulated genes were  
 215 enriched in B3→B1, B1→A1, B1→A2, A0→A2, A1→A2, A1→A3 (all  $P < 0.05$ ; Fig. 2E;  
 216 Supplemental Table S7). This suggests that specific subcompartment switching towards more  
 217 open or closed states is associated with up- or downregulated DEGs, respectively.

218 Further, we computed the log-ratio of subdiagonal sums in the upper- compared to the  
 219 lower triangular matrices from Fig. 2C. When this log-ratio is negative, DE genes are more  
 220 present in 10A, and when it is positive, DE genes are more present in T1 (Fig. 2D) or C1 (Fig.  
 221 2F). We confirm switching trends towards more open subcompartments for upregulated  
 222 genes, and the opposite for downregulated genes (Fig. 2D right) ( $P = 4.04 \times 10^{-9}$ ; Fisher's exact  
 223 test; Supplemental Table S8), while no trend is seen for non-differentially expressed genes  
 224 (Fig 2D middle). When contrasting 10A and C1, similar trends are seen (Fig. 2E-F;  $P = 6.66 \times$   
 225  $10^{-13}$ ; Supplemental Table S9). Thus, subcompartment switching aligns with changes in gene  
 226 expression.

227

### 228 **Reorganization of TAD-TAD interactions coincides with subcompartment alterations**

229 Beyond their subcompartment organization, we have previously shown that TADs can arrange  
 230 in densely interconnected long-range contact configurations termed TAD cliques (Paulsen et  
 231 al. 2019). To identify these cliques, each TAD is represented as a node in a graph, with edges  
 232 indicating statistically significant Hi-C interactions between pairs of TADs. We define a "TAD  
 233 clique" as a maximal, fully-connected subset of nodes that cannot be expanded without losing  
 234 full connectivity, which is identified using the Bron-Kerbosch algorithm (Bron and Kerbosch

235 1973). The size of a TAD clique is the number of nodes in a maximal clique (see Fig. 3A inset,  
236 and Supplemental Fig. S30). We computed TAD cliques in 10A, T1 and C1 cells (see Fig. 3A-  
237 B), and found that the distribution of maximal clique sizes overall show an expected decrease  
238 in their frequency as their size increases (Fig. 3A; Supplemental Fig. S31A). Notably, in 10A  
239 TAD cliques of size 3 are 17% less frequent than in T1 ( $P=0.023$ ; Suppl. Table S10), and 24%  
240 less frequent than in C1 ( $P=7.3 \times 10^{-8}$ ; Suppl. Table S10). Larger TAD cliques (size $>4$ ) on the  
241 other hand seem generally more enriched in 10A, than in T1 and C1. T1 generally exhibits  
242 fewer TAD cliques than 10A and C1 (Fig. 3A). Therefore, larger clique sizes are more apparent  
243 in normal 10A cells. These observed trends and differences are likely not related to structural  
244 variants in the T1 and C1 cells, as masking these out reveals similar distributions as in non-  
245 masked data (Supplemental Fig. S31).

246 We overlapped TAD cliques of different sizes with subcompartments, and observed that  
247 association of B-type subcompartments increases with TAD clique size (Supplemental Fig.  
248 S32). To characterize TAD clique dynamics, we analyzed alterations in TAD maximal clique  
249 sizes across cell types (Fig. 3C-D; Supplemental Fig. S33-S39). A large fraction of TADs do  
250 not engage in cliques (Fig. 3C). However, those TADs that do form cliques grow or decrease  
251 in size (Fig. 3C-D; Supplemental Fig. S33-S39) across all stages. Thus, like  
252 subcompartments, TAD cliques also reconfigure extensively in this model system of breast  
253 cancer progression.

254 Our previous observations do not exclude the possibility that growing TAD cliques involve  
255 TADs of various chromatin composition, thus belonging to various subcompartments. To  
256 explore this, we used HDBSCAN (McInnes et al. 2017; McInnes and Healy 2017) to cluster  
257 each clique based on their TAD-wise enrichment in subcompartments. Clustering was  
258 performed targeting clusters with a minimum cluster size of 200 TAD cliques, plus a cluster  
259 dedicated to collect outliers (see Methods for more details). The resulting nine clusters (Fig.  
260 3E) show that individual TAD cliques frequently involve multiple types of subcompartments  
261 (Supplemental Fig. S40), and thus represent a level of TAD association where  
262 subcompartments of different types potentially intermix.

263 To investigate the dynamics of TAD clique clusters, we explored the frequency of each of  
264 the nine TAD clique clusters in each cell type (Fig. 3F; Supplemental Table S11). We find that  
265 most clusters display a cancer progression stage-specific association, such that a cluster  
266 tends to be more enriched in one or two of the three cell types examined here (Fig. 3E,F).  
267 Some clusters involving B-type subcompartments (Cluster 0 and 2) appear to be significantly  
268 more prevalent in T1 and C1 compared to 10A. Conversely, some clusters with A-type  
269 subcompartments (Cluster 1, 5-7) seem to be diminished in T1, but exhibit partial recovery in  
270 C1 (Fig. 3F; Supplemental Table S12). These are also clusters enriched in weaker,  
271 intermediate subcompartments, which could reflect their increased switching.

272 In summary, TAD cliques bring together TADs overlapping with various subcompartments,  
273 which differ in 10A, T1 and C1 cells. Overall, this signifies the role of subcompartment  
274 alterations and intermixing during breast cancer progression in this model system.

275

### 276 **Interchromosomal insertion of the *MYC* locus coincides with de novo enhancer** 277 **contacts and oncogene activation**

278 As expected, the cell types used in our breast cancer progression model harbor structural  
279 variations (SVs), including copy number alterations, insertions, deletions and translocations.  
280 We exploited the ability of Hi-C to detect many of these SVs, to call SVs genome wide  
281 (Supplemental Fig. S41-S43). We manually inspected the called variants and confirmed  
282 known SVs (Santner et al. 2001), including t(3;9), t(3;5), t(3;17), t(6;19). In addition, we

283 identified previously undescribed SVs, including t(7;9) and t(10;17). Analyzing the 3D  
284 structural consequences of all SVs in these cell lines is computationally infeasible.  
285 Nevertheless, to investigate 3D genome consequences of select SVs, we focused on a striking  
286 pair of regions on Chromosome 8 (126330000-128235000 bp; hg38) (Fig. 4A) and  
287 Chromosome 10 (71280000-73310000 bp; hg38) (Fig. 4B). Both regions show a gradual and  
288 coordinated increase in copy number, going from T1 to C1 (Fig. 4A,B). A large region at the  
289 end of Chromosome 8 is amplified in 10A, but not in T1 and C1 (Fig. 4A). The region on  
290 Chromosome 8 is well-characterized (Visscher et al. 1997; Grisanzio and Freedman 2010)  
291 and contains several breast cancer related genes. These notably include *MYC*, an oncogenic  
292 transcription factor with a pivotal role in breast cancer progression (Liao and Dickson 2000)  
293 and whose amplification and overexpression is a marker of aggressive and invasive breast  
294 cancer (Corzo et al. 2006; Berns et al. 1992).

295 Further inspection of the Hi-C data at the intersection of these two regions in the Hi-C  
296 matrix reveals, after ICE balancing, an enrichment of contacts of the region towards the entire  
297 Chromosome 10, clearly indicating that the *MYC* locus is inserted in the Chromosome 10  
298 region (Fig. 4C). Enriched contacts with other chromosomes are also seen, due to the  
299 amplification of the locus (Fig. 4C). Zooming in on the specific interchromosomal contacts  
300 between the *MYC*-locus and the locus on Chromosome 10, we note Hi-C patterns that typically  
301 suggest chromatin looping and extrusion of intra-chromosomal DNA (Fig. 4D). This again  
302 indicates an insertion event. Investigating the region using a different balancing scheme which  
303 explicitly models and accounts for copy number alterations (Servant et al. 2018) reveals similar  
304 trends. This indicates that the copy number alterations themselves are not distorting the Hi-C  
305 data in this region (Supplemental Fig. S44). Comparing the contact maps between the three  
306 cell types reveals that the insertion is novel in the T1 and C1 cells (Supplemental Fig. S45).

307 We next examined the chromatin context of this insertion event. Inspection of  
308 subcompartments in this region shows that the insertion site on Chromosome 10 is nearly  
309 entirely covered by A3-subcompartments (Supplemental Fig. S46A) in all three cell types,  
310 indicating that the *MYC* locus is inserted into a highly active genome region on Chromosome  
311 10. Initially, the *MYC* gene resides in an A2 subcompartment on Chromosome 8 in the 10A  
312 cell line. However, it switches to a more active or “open” state (A2→A3) in T1 and C1  
313 (Supplemental Fig. S46B).

314 Next, we assessed the nature of the *de novo* contacts detected between the *MYC* locus  
315 and sites on the host Chromosome 10. To this end, we overlaid positions of MCF10A  
316 enhancers predicted from EnhancerAtlas 2.0 (Gao and Qian 2020) onto the Chromosome 8  
317 and 10 amplification unit (Fig. 4D; green segments). The four distinct *MYC* interactions  
318 detected in the Hi-C map (Fig. 4D, circles) evidently corresponds to (i) two *de novo* contact  
319 points between enhancer elements on Chromosome 10 and the *MYC* promoter, and (ii) an  
320 interaction with enhancer elements ~1 Mbp upstream of the *MYC* promoter (Fig. 4D).  
321 Generating a virtual 4C plot (Sexton et al. 2012) (see methods) anchored at the *MYC* locus,  
322 confirms that the enrichment of contacts towards these putative enhancer elements is  
323 specific to T1 and C1 (Fig 4D; bottom track).

324 To validate the *de novo* interactions (proximity) of the *MYC* gene with elements on  
325 Chromosome 10 at the cellular level, we carried out a fluorescence in situ hybridization (FISH)  
326 analysis of the relative positions of the *MYC* gene (*MYC* probe on Chromosome 8 at position  
327 127543567-127702694) and either a *de novo* interacting enhancer on Chromosome 10  
328 (“Chr10 enhancer” probe at position 73117711-73267436), or another site on Chromosome  
329 10 ~500 kbp downstream (“Chr10 downstream” probe at position 73794997-73992612; see  
330 Methods). The data show that both the “*MYC*-Chr10 enhancer” and the “*MYC*-Chr10 distal”

331 distances are significantly shorter in C1 cells than in 10A cells ( $P=1.9 \times 10^{-14}$  and  $1.3 \times 10^{-7}$   
332 respectively; Fig. 4E,F; Supplemental Fig. S47A-D), yet the *MYC* enhancer contact is  
333 significantly shorter than the distal interaction ( $P=9.3 \times 10^{-6}$ ). These findings support the view  
334 that the *MYC* locus has undergone an insertion into Chromosome 10 in C1 cells.  
335 Furthermore, this region harbors enhancer elements which may interact with *MYC*,  
336 though resolution limitations of our FISH probes prevent a more detailed characterization.

337 Furthermore, to provide an element of functional significance to this *MYC* insertion, we  
338 examined RNA-seq data for these two regions. The data show an upregulation of breast  
339 cancer related genes in T1 and C1 relative to 10A, including *LRATD2*, *PCAT1*, *CASC19*,  
340 *CASC8*, *POU5F1B*, *PVT1* and *MYC* on Chromosome 8. Several breast cancer related genes  
341 in the Chromosome 10 region are also upregulated, such as *UNC5B*, *CDH23*, *PSAP*,  
342 *SPOCK2*, *ASCC1*, *DDIT4*, *NUDT13*, *ECD*, *DNAJC9* and others (Supplemental Table S13  
343 and S14).

344 To dissect the functionality of the *MYC* insertion we designed 5' and 3' sgRNA pairs to  
345 target the de novo putative enhancer interactions (Supplemental Fig. S48A). The *MYC* de  
346 novo contacts are proximal to the *DNAJC9* locus on Chromosome 10 (Figure 4D), where  
347 three putative cis regulatory elements (REs) enriched in ENCODE H3K27ac and annotated  
348 CREs are found (REs 1-3; Supplemental Fig. S48B). Virtual 4C-track overlays reveal  
349 enriched contacts in T1 and C1 cells compared to 10A within this region, with a peak aligning  
350 with RE1 (Supplemental Fig. S48B; top). C1 cells were double transfected with  
351 pLentiMultiCRISPR vectors where Cas9 is coexpressed with puromycin and each  
352 sgRNA pair to create six different genomic deletions spanning RE 1-3 (Supplemental  
353 Fig. S48B) or sgRNAs for mCherry or EGFP as controls. The Cas9 protein was equally  
354 expressed from all vectors in C1 cells compared to GAPDH (Supplemental Fig. S48C). After  
355 two days of puromycin selection for Cas9 expressing C1 cells, we observe a subtle, yet not  
356 significant, decrease in *DNAJC9* expression upon Cas9 targeting of the three REs in the  
357 locus (Supplemental Fig. S48D). *MYC* expression slightly decreased in C1 cells expressing  
358 guide pairs for deletion of RE1.1 and RE1.2, and showed a subtle increase in expression  
359 for sgRNAs surrounding RE2.1-2 and RE3.1-2, yet not statistically significant  
360 (Supplemental Fig. S48E). We did not observe any change in expression of *POLR2A*,  
361 encoding for the largest subunit of RNA polymerase II on Chromosome 17 upon targeting of  
362 Cas9 to REs 1-3 or controls (Supplemental Fig. S48F). To evaluate Cas9 editing efficiency,  
363 we amplified genomic DNA from puromycin-selected, doubly transfected C1 cells using  
364 primers targeting regions surrounding sites recognized by sgRNAs in RE1, 2 or 3  
365 (Supplemental Fig. S49A). PCR products were analyzed by gel electrophoresis, purified,  
366 and sequenced (Supplemental Fig. S49B). This evaluates the frequency of indels  
367 surrounding each of the sgRNA breakpoints, which was quantified using the TIDE method  
368 (Brinkman et al. 2014), and found to be relatively low (0.8–7.6%; Supplemental Table S19).  
369 This may partially explain the subtle effect on *MYC* expression and low Cas9 deletion  
370 frequency of RE1-3. However, the genomic complexity and redundancy of the *MYC* locus  
371 make definitive conclusions challenging.

372 Collectively, these observations indicate that the *MYC* locus has not only been amplified  
373 but also inserted into a transcriptionally active region of Chromosome 10 in C1 cells, with the  
374 formation of de novo cis-contacts between *MYC* and regulatory elements in the host  
375 chromosome. Upregulation of *MYC* and of other breast cancer-related genes in malignant  
376 C1 cells speculatively suggests functionality of these de novo interactions, yet determination  
377 of any involved enhancer elements remains. Our results highlight how structural variants in  
378 the genome may cooperate with 3D genomic states during oncogenesis.

## 379 Discussion

380 The intricate 3D organization of the genome plays a pivotal role in gene regulation and cell  
381 function. Using a multilayered (epi)genomic approach, we report here for the first time, to our  
382 knowledge, significant alterations in higher-order 3D chromatin organization in an *in vitro*  
383 breast cancer progression model.

384 First, major compartment and subcompartment switching emerges as a prominent  
385 dynamic feature, which as expected based on previous studies (Nagai et al. 2019; Liu et al.  
386 2021), correlates with altered expression of genes implicated in breast cancer. Second, we  
387 identify a reconfiguration of preferred associations between TADs, or TAD cliques (Paulsen et  
388 al. 2019), which bring together new A- and B-type subcompartment associations. Third, spatial  
389 subcompartment organization changes in the pre-malignant and malignant states potentially  
390 also contribute to the loss of gene expression control. Collectively, there is a coordinated and  
391 progressive reorganization of higher-order genomic structure to an abnormal state. However,  
392 casual relationships remain to be determined.

393 The extent of compartment switching observed in our model (22%) is similar to what is  
394 seen during B cell differentiation (28.1%) (Vilarrasa-Blasi et al. 2021), although the latter  
395 includes an additional intermediate compartment, inflating the percentage. In contrast, 7.5%  
396 of the genome harbors statistically significant compartment switches during differentiation of  
397 mouse embryonic stem cells (ESCs) into neuronal progenitors (Chakraborty et al. 2022). A/B-  
398 compartment switching varies between 8-36% comparing various ESC-derived cell types  
399 (Dixon et al. 2015; Liu et al. 2021). Comparatively, this indicates that the level of switching  
400 observed in our oncogenic model is substantial.

401 Our study did not uncover involvement of substantial TAD border dynamics in the *in vitro*  
402 progression model of breast cancer. However, we cannot entirely rule out its subtle  
403 involvement due to minor differences in consistency across replicates. Further investigation of  
404 these differences is needed to uncover any potential functional consequences. Additionally,  
405 the potential role of cell heterogeneity in cancer cells cannot be dismissed, given that our data  
406 represent an overall behavior in bulk cell populations (Zhu et al. 2023). This heterogeneity is  
407 expected to primarily affect 3D genome models, but single-cell copy number variations could  
408 also influence estimates of overall copy numbers.

409 A new observation from our data is a previously unknown insertion of the *MYC* locus  
410 (normally localized on Chromosome 8) into highly active A3-subcompartments on  
411 Chromosome 10. This is accompanied by *de novo* interactions correlating with not only  
412 elevated *MYC* expression in T1 and C1 cells, but also with enhanced expression of other  
413 breast cancer related genes on both chromosomes. Our FISH analysis supports our Hi-C  
414 findings of *de novo* interactions between *MYC* and a region harboring enhancer elements on  
415 Chromosome 10 in C1 cells, but not in 10A cells. Furthermore, a complementary study has  
416 also classified this insertion as an enhancer hijacking event (Wang et al. 2021). Notably, in  
417 murine B cells, *MYC* has been shown to relocate to active transcription factories in proximity  
418 to *IGH*, a frequent translocation partner (Osborne et al. 2007). However, our FISH results  
419 indicate that in 10A cells, *MYC* is not proximal to the Chromosome 10 locus, suggesting *MYC*  
420 is not part of a transcription factory with this locus prior to the insertion event. Our analyses  
421 did not test for changes in *MYC* location in response to signaling pathway activation as in  
422 (Osborne et al. 2007), and thus refer to a "baseline" situation in these cells.

423 Identifying enhancer candidates from Hi-C interactions is challenging due to the data's low  
424 genomic resolution in pinpointing enhancer-promoter contacts. Tools like 4C and Micro-C can  
425 provide higher-resolution insights into specific chromatin interactions. However, these

426 techniques do not guarantee that detected contacts correspond to functional interactions, as  
427 spatial proximity does not inherently imply biological significance (Splinter et al. 2012; Hsieh  
428 et al. 2020).

429 Focusing on three regulatory elements (REs1-3) in the *DNAJC9* locus on Chromosome  
430 10 intersecting the central part of the “dot” observed in our Hi-C data (Fig. 4D), we utilized a  
431 CRISPR-Cas9 strategy to analyze consequences of deleting these putative enhancer  
432 elements in C1 cells. From this, we observed changes in expression of *DNAJC9* and *MYC*  
433 compared to sgRNA controls, yet these changes were not statistically significant. Studies have  
434 shown that deleting individual enhancers can have minimal impact due to redundancy  
435 among enhancers and may cause only transient effects (Osterwalder et al. 2018; Diao et al.  
436 2016), possibly explaining why targeting individual *DNAJC9* REs may have low effect  
437 on *MYC* expression. These effects may, however, also be masked by the large copy  
438 numbers of *MYC* and *DNAJC9* loci (Fig. 4A,B). Moreover, the low editing efficiency of  
439 individual sgRNAs in pool C1 cells after transient transfection and puromycin selection  
440 suggest a low frequency of the RE1-3 deletions (Supplemental Table S19). Therefore, we  
441 cannot definitively determine the functional significance of the observed enhancer  
442 interactions.

443 The detailed spatial configuration of this insertion remains unclear, but there are several  
444 possibilities. The mirror-symmetry and isolated contact map of the Chromosome 10 segment  
445 with specific regions of the Chromosome 8 segment (see Fig. 4D), featuring stripes,  
446 could suggest that the region exists as circular, extrachromosomal DNA (ecDNA). The  
447 formation of ecDNA structures is a common form of oncogenic amplification, and often occurs  
448 around *MYC* (Hung et al. 2022; Hoff et al. 1988) to promote functional *cis*-regulatory  
449 contacts (Morton et al. 2019). However, the focal and limited *MYC* probe signals  
450 (Supplemental Fig. S47A-B) are inconsistent with an ecDNA event (Hung et al. 2021),  
451 leaving the structural interpretation unresolved.

452 Beyond our identified interchromosomal insertion of the *MYC* locus, the intricate  
453 aneuploidy patterns commonly observed in cancer pose a challenge that existing standard  
454 software and pipelines struggle to effectively address. Promising strides have been  
455 made using long-read Hi-C technology (Garg 2023) or statistical approaches (Brunette et al.  
456 2024) to alleviate this issue. Nevertheless, a significant gap remains in the  
457 availability of computational pipelines tailored for comprehensive downstream analysis of  
458 reordered cancer genome data relative to a karyotypically normal reference genome.  
459 This needs to be addressed if further progress in understanding the 3D cancer genome is  
460 to be achieved.

461 In conclusion, our findings provide new insights into the complex genomic structural  
462 changes that underlie breast cancer metastatic progression. The interplay between  
463 subcompartment switching, LAD-linked subcompartment spatial nuclear reorganization, and  
464 TAD clique dynamics represents a multifaceted mechanism by which the 3D genome can be  
465 abnormally and dynamically reconfigured during breast cancer development. Targeting  
466 these 3D genome alterations could potentially lead to improved or new therapeutic  
467 strategies to better treat breast cancer in the future (Park et al. 2023).

## 468 **Methods**

### 469 **Cells**

471 MCF10A, MCF10AT1, and MCF10Ca1a cell lines were grown in DMEM/Nutrient F12  
472 (DMEM/F12) media supplemented with 5% horse serum (MCF10A and MCF10Ca1a) or 2.5%  
horse serum (MCF10AT1), 14 mM NaHCO<sub>3</sub>, 10 µg/mL insulin, 2 mM L-glutamine, 20 ng/mL  
human epidermal growth factor, 500 ng/mL Hydrocortisone and 100 ng/mL cholera Toxin.

473 MCF10A cells were obtained from the American Type Culture Collection (CRL-10317).  
 474 MCF10AT1 and MCF10Ca1a cells were obtained from the Barbara Ann Karmanos Cancer  
 475 Institute (Detroit, Michigan).

476

#### 477 **Fluorescence in situ hybridization (FISH)**

478 FISH was done as exactly described by us previously (Paulsen et al. 2017). In short, MCF10A  
 479 and MCF10Ca1a cells were incubated in a hypotonic buffer, fixed in ice-cold methanol:acetic  
 480 acid and dropped on glass slides. BAC FISH probe DNA (BacPac Resource Center) was  
 481 labeled using a Nick Translation Kit (Roche). The following probes were used:

- 482 • 'MYC' gene probe: clone ID RP11-1136L8; Chr.8; position 127543567-127702694
- 483 • 'Chr. 10 enh.' enhancer probe: clone ID RP11-152N13; Chr.10; position 73117711-  
 484 73267436
- 485 • 'Chr. 10 distal' probe: clone ID RP11-390A15; Chr.10; position 73794997-73992612

486 The MYC gene probe was labeled with Digoxigenin-11-dUTP (Roche). The Chr.10 enhancer  
 487 and Chr.10 distal probes, both positioned to mark sites expected to be proximal to the *MYC*  
 488 gene on Chromosome 10) were labeled with Biotin-16-dUTP (Roche). For each slide, 200 ng  
 489 of each 'MYC gene' + 'Chr.10 enhancer' probe, or 'MYC gene' + 'Chr.10 distal' probe were  
 490 mixed with 30 µg of Cot-1 DNA and 150 µg salmon sperm DNA and precipitated. DNA was  
 491 dissolved in hybridization mix and pre-annealed for 1 h. Slides were RNase-treated, washed,  
 492 dehydrated in ethanol, denatured, and dehydrated again.

493 Probes were denatured, pre-annealed and applied onto cells for overnight hybridization at  
 494 37°C. Slides were then washed in 2× SSC and in 0.1× SSC, blocked in 5% skim milk and  
 495 incubated with Anti-Digoxigenin (Roche; mouse; 0.4 µg/ml). Slides were washed, incubated  
 496 with Avidin Alexa Fluor 488 conjugate (Invitrogen; 1.7 µg/ml) and Alexa Fluor® 594-  
 497 conjugated Anti-Mouse (Jackson ImmunoResearch; rabbit; 2.5 µg/ml), washed and incubated  
 498 with Biotinylated Anti-Avidin D conjugate (Vector; goat; 1.0 µg/ml) and Alexa Fluor® 594-  
 499 conjugated Anti-Rabbit (Jackson ImmunoResearch; donkey; 2.5 µg/ml). Slides were washed  
 500 and incubated 30 min with Avidin Alexa Fluor 488 conjugate (Invitrogen) (1.7 µg/ml). Slides  
 501 were mounted with 0.2 µg/ml DAPI in Dako Fluorescent Mounting Medium. Images taken  
 502 under a 100× objective (numerical aperture 1.4) mounted on an IX71 inverted microscope  
 503 (Olympus) fitted with the DeltaVision wide-field imaging station (GE Healthcare).

504

#### 505 **Hi-C**

506 HiC Libraries were prepared in duplicates for each cell type using the Arima-HiC+ kit (Arima  
 507 Genomics, USA) according to manufacturer instructions. For each cell line (10A, T1, C1) 1  
 508 million cells were used per HiC reaction. Cells were fixed with 1.2% formaldehyde for 12 min.  
 509 All subsequent steps were carried out according to the Arima protocol. Resulting libraries were  
 510 amplified with 5 PCR cycles and sequenced on an Illumina NovaSeq 6000 instrument in  
 511 paired-end run with 2×101 bp.

512

#### 513 **RNA-seq**

514 All mRNA-seq experiments were performed in triplicate. Total RNA was isolated using the  
 515 Qiagen RNeasy kit following manufacturer's instructions. Stranded mRNAseq libraries were  
 516 constructed using Illumina mRNA Prep kit, following vendor protocol with poly(A) enrichment  
 517 (Illumina). Libraries were sequenced with 2×75bp paired-end on an Illumina NextSeq 500.

518

#### 519 **CRISPR-Cas9 Regulatory element deletions**

520 Paired sgRNAs (5' and 3'- sgRNAs) were designed to target both ends of selected candidate  
521 regulatory elements (REs) in the human *DNAJC9* locus to create a deletion. sgRNA with on-  
522 target high predicted cleavage and low number of off-targets were carefully selected using  
523 UCSC Genome Browser (hg38) (Perez et al. 2025) CRISPR Target track based on  
524 CRISPOR (Haeussler et al. 2016). The sgRNAs were cloned stepwise into Lenti-multi-  
525 CRISPR plasmid that contains human codon-optimized *Streptococcus pyogenes* Cas9  
526 (spCas9) followed by a *porcine teschovirus-1* 2A peptide (P2A) self-cleavage sequence  
527 upstream of the puromycin sequence (Addgene plasmid # 85402) (Cao et al. 2016). The  
528 first gRNA was cloned by insertion of two annealed complementary oligonucleotides into  
529 the BsmBI site using the Infusion cloning approach (TaKaRa Clontech, Supplemental  
530 Table S20). The plasmid containing one guide was opened with NheI (NEB) and a  
531 DNA fragment containing U6 promoter, sgRNA and scaffold (IDT technologies) was  
532 inserted by infusion cloning (Supplemental Table S19). A total of two sgRNA pairs per  
533 three *DNAJC9* locus REs were cloned. sgRNAs for mCherry and EGFP were designed  
534 as negative controls and inserted separately as annealed oligonucleotides by infusion  
535 cloning into the BsmBI site in two separate Lenti-multi-CRISPR plasmids. All  
536 plasmids were fully sequenced by Oxford Nanopore technology (Eurofins Genomics)  
537 and the sequences manually verified.

538 A total of 600,000 C1 cells were seeded per six well in DMEM/F12 supplemented media  
539 as described above. After 24 hours, three technical replicates of six wells were transfected  
540 with 2.5 µg of Lenti-multi-CRISPR plasmid containing 5' and 3'-sgRNA RE pair or EGFP  
541 sgRNA or Cherry sgRNA using Lipofectamine 3000 (Thermo Fisher Scientific). At 24 hours  
542 post transfection, each 6 well was transfected again with 2.5 µg of the same sgRNA-Lenti-  
543 multi-CRISPR plasmid using Lipofectamine 3000. After 24 hours, the double transfected C1  
544 cells were selected with 4 µg/ml of puromycin for 48 hours. On day five, the media was  
545 changed and the cells left to recover. On day six, the six wells were washed in 1 x PBS and  
546 either collected in 500 µl TRIzol Reagent (Thermo Fisher Scientific) for total RNA or trypsinized  
547 and pelleted for genotyping. Genomic fragments spanning sgRNAs in RE1, 2 or 3 were  
548 amplified by Q5 polymerase (NEB), single bands were purified and sanger sequenced  
549 (Eurofins) (Supplemental Table S20). The TIDE web tool (version 3.3.0) (Brinkman et al. 2014)  
550 was used to determine the efficiency of individual guide RNAs in generating targeted indel  
551 mutations in a cell population at 25 nucleotides from the breakpoint (Supplemental Table S19).

552 To verify Cas9 expression, whole cell lysates of C1 were lysed in 2x SDS loading dye  
553 and loaded on a 4-20% SDS PAA gel (BioRad) after lipofectamine transfection  
554 with pLentiMultiCRISPR (addgene #85402) with sgRNA pairs spanning RE1.1, RE1.2,  
555 RE2.1, RE2.2, RE3.1, RE3.2, and guide controls against EGFP and mCherry. Whole cell  
556 lysates from C1 cells transfected with a vector with humanized Cas9 (hCas9, addgene  
557 #41815) was loaded as control (Mali et al. 2013). The proteins were transferred to a PVDF  
558 membrane (Millipore) and detected with anti-Cas9 (Cas9-clone 7A9 Cat # MAC133, Sigma  
559 Aldrich, expected size 160 kda) and anti-GAPDH (Novus biologicals, expected size 37  
560 kda) as a loading control followed by secondary IR800 donkey anti-mouse and IR680  
561 donkey anti-goat (LicorBio) and detected by an OdysseyDLx imager.

562 Total RNA from the three six wells for each 5' and 3'- sgRNA pair or control EGFP/  
563 mCherry sgRNAs Lenti-multi-CRISPR plasmid transfected cells was purified using a RNeasy  
564 kit with an on-column DNase I treatment according to manufacturer's protocol (Qiagen). RNA  
565 was eluted in 25 µL ddH<sub>2</sub>O quantified using NanoDrop and quality controlled by agarose gel  
566 electrophoresis. A total of three different replicates of double transfections with puromycin  
treatment conducted at different times were processed. 2 mg of total RNA was

567 used for generation of cDNA primed with oligo(dT) according to manufacturer's instructions  
568 (AffinityScript cDNA synthesis kit Cat # 600559, Agilent).

569 The endogenous *MYC*, *DNAJC9*, *RPL30* and *POL2RA* mRNA expression levels were  
570 measured by quantitative RT-PCR using EvaGreen dye (Biotium) on a BioRad CFX Real-  
571 Time System. A standard curve was generated for each primer set with Thermal cycle  
572 conditions for 40 PCR cycles (95° for 12 min, 95° for 15s, 65° for 30s, 72° for 30s)  
573 (Supplemental Table S19). The Real-Time Efficiency (%) for each primer pair was calculated  
574 according to  $E = -1 + 10(-1/\text{slope})$  (Rasmussen 2001) and converted primer efficiency =  $E$   
575  $(\%/100 + 1)$ .

576 Three technical replicates were run per qRT-PCR (n=3 rounds of CRISPR; (n=9 technical  
577 replicates)). Relative gene expression was analyzed using Pfaffl's mathematical model (Pfaffl  
578 2001) for 5' and 3' - sgRNA RE pair or mCherry sgRNA versus EGFP sgRNA transfected cells  
579 normalised to housekeeping gene RPL30 as reference:

580

$$581 \text{ Ratio (RQ)} = (E_{\text{target gene}})^{\Delta C(\text{target gene})} / (E_{\text{reference}})^{\Delta C(\text{reference})}$$

582

583  $\Delta C(\text{target}) = \text{Ct}(\text{Mean target gene in CRISPR control gRNA(EGFP) cells}) - \text{Ct}(\text{target gene in}$   
584  $\text{CRISPR RE or mCherry sgRNA cells})$

585  $\Delta C(\text{reference}) = \text{Ct}(\text{reference gene in CRISPR control gRNA(EGFP) cells}) - \text{Ct}(\text{reference}$   
586  $\text{gene in in CRISPR RE or mCherry sgRNA cells})$

587

588 Ratio was calculated for average of technical Cp values per replicate, then these were  
589 averaged and STDEV calculated. The control sample was from pooled cells transfected with  
590 EGFP gRNA. Samples are RE = Pool of cells with guide pair for RE1.1; RE1.2; RE2.1; RE2.2;  
591 RE3.1 or RE3.2. A pool of cells with mCherry gRNA acts as a negative CRISPR control. Target  
592 genes are *MYC*, *DNAJC9* and *POLR2A* (internal control target gene on chromosome 17,  
593 assumed not affected by CRISPR deletions). qRT-PCR data statistical analysis and plots were  
594 performed on GraphPad Prism 10.4.1. For statistical evaluations of the determined CP  
595 variations and calculated relative expression variations, data were analysed for significant  
596 differences by one-way ANOVA using Dunnett's multiple comparisons test between mCherry  
597 sgRNA and each of different RE guide pairs were performed on GraphPad Prism v 10.4.1  
598 (Supplemental Table S20, the confidence level was set to 0.05).

599

### 600 **ChIP-seq of lamin B1 and mapping of lamina-associated domains (LADs)**

601 ChIP of lamin B1 was done as described by us (Rønningen et al. 2015). In short, cells were  
602 cross-linked with 1% formaldehyde, lysed in 50 mM Tris-HCl pH 7.5, 10 mM EDTA, 1% SDS  
603 and protease inhibitors, and sonicated in a Bioruptor (Diagenode) into ~200-bp fragments.  
604 After sedimentation, the supernatant was diluted tenfold in the RIPA buffer. Chromatin was  
605 incubated overnight at 4°C with antibodies to lamin B1 (10 µg per 10 million cells; Abcam,  
606 ab16048), coupled to Invitrogen Dynabeads protein A/G (Thermo-Fisher). ChIP samples were  
607 washed four times in ice-cold RIPA buffer, after which cross-links were reversed and the DNA  
608 was eluted for 6 h at 68°C. DNA was purified using phenol–chloroform isoamylalcohol and  
609 dissolved in H<sub>2</sub>O. ChIP-seq libraries were prepared using the Diagenode Microplex library  
610 preparation kit v2 and TruSeq LT indexes, and samples were sequenced (single-end) on an  
611 Illumina HiSeq 4000.

612 ChIP sequence reads were mapped to hg38 with Bowtie 2 v2.4.1  
613 (<https://github.com/BenLangmead/bowtie2>) (Langmead and Salzberg 2012) after removing

614 duplicates using Picard MarkDuplicates (<http://broadinstitute.github.io/picard/>). Reads from  
 615 both input samples were merged, mapped to hg38 and duplicates removed as above. To  
 616 alleviate normalization bias, each pair of mapped ChIP and input read files contained the same  
 617 read depth after down-sampling reads for each chromosome. Mapped reads were used to call  
 618 LADs from ten consecutive runs of Enriched Domain Detector (EDD)  
 619 (<http://github.com/CollasLab/edd>) (Lund et al. 2014) with auto-estimation of GapPenalty and  
 620 BinSize, and mean GapPenalty and BinSize values were used for a final EDD run (Forsberg  
 621 et al. 2019). Final LADs for each cell type were the union of the three replicates.

622

### 623 **Structure of code used for data analyses**

624 Source code used to produce all results presented in this paper (except LADs analysis and  
 625 downstream analysis of Chrom3D models) is hosted on GitHub at  
 626 [github.com/paulsengroup/2022-mcf10a-cancer-progression](http://github.com/paulsengroup/2022-mcf10a-cancer-progression) and is archived on Zenodo at the  
 627 following DOI: [10.5281/zenodo.17121063](https://doi.org/10.5281/zenodo.17121063).

628 Most of the computation is organized into Nextflow workflows, with each workflow performing  
 629 a subset of the data analysis (e.g. the differential expression and comparative analyses are  
 630 performed by two different workflows).

631 All workflows mentioned in the method sections are hosted on the GitHub repository inside  
 632 folder workflows/. Scripts and Jupyter notebooks are found under folders bin/ and notebooks/,  
 633 respectively.

634 Some of the analysis steps required us patching third party tools. The patches are described  
 635 in section Software patches and patch files are available on the GitHub repository under  
 636 container/patches/. All our patches were contributed upstream. Most of our patches were  
 637 accepted by upstream and are already part of a stable release. Figures were produced directly  
 638 by workflows or using scripts and notebooks under bin/plotting and notebooks/, respectively.  
 639 The final version of figures were produced by assembling draft images with Inkscape and  
 640 Omnigraffle.

641 Data analysis workflows were run on a HPC cluster using Apptainer (Kurtzer et al. 2017).

642

### 643 **Hi-C preprocessing**

644 We used a modified version of nf-core/hic v2.0.0 (<https://zenodo.org/records/2669513>) (Ewels  
 645 et al. 2020; Servant et al. 2015; Ewels et al. 2016; da Veiga Leprevost et al. 2017) for quality  
 646 control, sequence mapping and filtering using hg38.p14 (International Human Genome  
 647 Sequencing Consortium 2001). The workflow was run using restriction enzymes for the 2-  
 648 enzyme Arima Kit (`--restriction_site="^GATC,G^ANTC"` and `--  
 649 ligation_site="GATCGATC,GANTGATC,GANTANTC,GATCANTC"`). The following optional  
 650 flags were specified: `--skip_maps`, `--skip_dist_decay`, `--skip_tads`, `--skip_compartments`, `--  
 651 skip_balancing`, `--skip_mcool`, `--split_fastq=false`. The .chrom.sizes file given to nf-core/hic  
 652 was filtered to only retain chromosome sequences using `grep '^chr[[:digit:]]XY\|+[:space:]'`.  
 653 Chromosomes were sorted by name using `gnu sort -V`. The output of nf-core/hic was  
 654 compressed using workflow `robomics/compress-nfcore-hic-output v0.0.1`  
 655 (<https://zenodo.org/records/7949266>). Hi-C contact matrices in multi-resolution Cooler format  
 656 were generated using a patched version of cooler v0.9.1 (Abdennur and Mirny 2020). We used  
 657 cooler cloud to ingest interactions in .validPair format into a .cool file at 1 kbp resolution. We  
 658 then converted the .cool file to .mcool file format using cooler zoomify. Finally, matrices were  
 659 balanced with several methods using juicer\_tools v2.20.00 (Durand et al. 2016) (VC, KR,  
 660 SCALE methods using intra-chromosomal, inter-chromosomal and genome-wide  
 661 interactions), and cooler (ICE method using intra-chromosomal, inter-chromosomal and

662 genome-wide interactions) (Abdennur and Mirny 2020). Regions overlapping centromeres and  
 663 assembly gaps for hg38 were masked before balancing (files retrieved from UCSC FTP server  
 664 on 2023/02/24). Downstream analyses were performed using cis-only, ICE balanced matrices  
 665 unless otherwise specified. For some analyses, replicates for the same condition were merged  
 666 to generate deeper Hi-C matrices using cooler merge (Abdennur and Mirny 2020). The  
 667 resulting matrices were then coarsened and balanced cooler and juicer\_tools as outlined  
 668 above (Abdennur and Mirny 2020; Durand et al. 2016). All the above steps were performed  
 669 by running script run\_nfcore\_hic.sh, which runs workflow postprocess\_nfcore\_hic.nf after  
 670 running nf-core/hic. Fig. S1 was generated by workflow postprocess\_nfcore\_hic.nf based on  
 671 the output of nf-core/hic. Fig. S2 was generated by workflow comparative\_analysis\_hic.nf by  
 672 running the Python implementation of HiCRep v0.2.6 (Lin et al. 2021; Yang et al. 2017).  
 673 Correlation values shown in the plot were computed as the weighted average of the correlation  
 674 coefficient of each individual chromosome using chromosome sizes as weights. Individual  
 675 correlation coefficient values are available in Supplemental File S1.  
 676 In addition, interactions involving Chromosome 8 and Chromosome 10 were balanced with  
 677 LOIC using the iced package (v0.5.10) (Servant et al. 2018). LOIC balancing is performed in  
 678 one of the steps of the comparative\_analysis\_hic.nf workflow.

679

### 680 **RNA-seq preprocessing**

681 We used nf-core/mnaseq v3.12.0 (<https://zenodo.org/records/7998767>) to perform quality  
 682 control, trimming, alignment and generating the gene expression matrix (Ewels et al. 2020; Di  
 683 Tommaso et al. 2017) (Ewels et al. 2020; Di Tommaso et al. 2017); This includes the following  
 684 tools: featureCount (Liao et al. 2014); GffRead (Pertea and Pertea 2020); MultiQC (Ewels et  
 685 al. 2016); preseq (Daley and Smith 2013); Qualimap 2 (Okonechnikov et al. 2016) ; RSeQC  
 686 (Wang et al. 2012); Salmon (Patro et al. 2017); SortMeRNA (Kopylova et al. 2012); STAR  
 687 (Dobin et al. 2013); StringTie2 (Kovaka et al. 2019); UCSC Tools (Kent et al. 2010); DESeq2  
 688 (Love et al. 2014); dupRadar (Sayols et al. 2016); Tximeta (Love et al. 2020); Bioconda  
 689 (Grüning et al. 2018) ; Biocontainers (da Veiga Leprevost et al. 2017) and Apptainer (Kurtzer  
 690 et al. 2017). nf-core/mnaseq was run using hg38.p14 as reference genome (International  
 691 Human Genome Sequencing Consortium 2001). The FASTA file used as reference was first  
 692 filtered using SeqKit v2.5.1 (Shen et al. 2016) to remove mitochondrial chromosomes and  
 693 unplaced scaffolds by using the following pattern '^chr[XY\d]+\$. We used the genome  
 694 annotation for hg38 from GENCODE v43 (Frankish et al. 2019). The pipeline was launched  
 695 using the following optional flags: --aligner=star\_salmon, --pseudo\_aligner=salmon, --  
 696 extra\_salmon\_quant\_args='--seqBias --gcBias', --gencode=true.

697 The workflow was run by launching script run\_nfcore\_mnaseq.sh.

698

### 699 **ChIP-seq analyses of public datasets**

700 We used nf-core/chipseq v2.0.0 to perform quality control, mapping, filtering and peak calling  
 701 of publicly available ChIP-seq datasets from MCF10A cells (Ewels et al. 2020; Di Tommaso  
 702 et al. 2017). This includes the following tools: BWA (Li and Durbin 2009); BEDTools (Quinlan  
 703 and Hall 2010); BamTools (Barnett et al. 2011); deepTools (Ramírez et al. 2016);  
 704 featureCounts (Liao et al. 2014); HOMER (Heinz et al. 2010); MultiQC (Ewels et al. 2016);  
 705 phantompeakqualtools (Landt et al. 2012); preseq (Daley and Smith 2013); SAMtools (Li et  
 706 al. 2009); UCSC Tools (Kent et al. 2010); DESeq2 (Love et al. 2014); Bioconda (Grüning et  
 707 al. 2018); BioContainers (da Veiga Leprevost et al. 2017) and Apptainer (Kurtzer et al. 2017).  
 708 The pipeline was run using hg38.p14 as reference genome from UCSC (International Human  
 709 Genome Sequencing Consortium 2001) and the gene annotation from GENCODE v43

710 (Frankish et al. 2019). BWA was used for mapping and effective genome size for MACS2 was  
711 computed using unique-kmers.py from khmer v2.1.1 (Li and Durbin 2009; Zhang et al. 2008;  
712 Crusoe et al. 2015; Döring et al. 2008). Option --narrow\_peak was specified when processing  
713 ChIP-seq datasets for CTCF and TP53.

714 The workflow was launched by executing script run\_nfc\_core\_chip.sh.

715

### 716 **Calling of copy number variations**

717 Copy number variations (CNVs) were called from Hi-C data using HiNT-CNV v2.2.8 (Wang et  
718 al. 2020).

719 The list of restriction sites for the Arima 2-enzyme kit was generated by running script  
720 bin/compute\_restriction\_sites\_for\_hint.py on hg38.14 (International Human Genome  
721 Sequencing Consortium 2001).

722 Reference data and background matrices used by HiNT were downloaded from  
723 [compbio.med.harvard.edu/hint](https://compbio.med.harvard.edu/hint) and have been archived on Zenodo at the following DOI:  
724 [10.5281/zenodo.13255302](https://doi.org/10.5281/zenodo.13255302).

725 Data analysis steps are defined in workflow detect\_structural\_variants which was run by  
726 launching script run\_detect\_structural\_variants.sh.

727

### 728 **Annotation of structural variations**

729 Large structural variations such as translocations were manually annotated by us by looking  
730 for square regions of enriched interactions and sharp transitions in the trans portion of the Hi-  
731 C matrix (see Supplemental Tables S15-17).

732

### 733 **TAD analyses**

734 We determined genomic positions of topologically associated domains (TADs) in all samples  
735 with HiCEXplorer v3.7.2 using default parameters on matrices at 10, 20, 50 and 100 kbp  
736 resolutions (Ramírez et al. 2018; Wolff et al. 2018, 2020). Workflow tad\_analysis.nf was used  
737 to generate the draft figures that went into Supplemental Fig. S3-6. The same workflow was  
738 also used to run HiCEXplorer. Draft figures for Fig. 1A was generated using notebook  
739 bin/plotting/plot\_tads\_highlass.ipynb. Supplemental Fig. S5 was generated by fetching  
740 interactions at 50 kbp resolution for each 10A TAD across cell types, masking values  
741 overlapping the first 150 kbp around the diagonal. Finally interactions were summed and  
742 normalized by TAD size (as in, the number of pixels overlapping a TAD). To generate  
743 Supplemental Fig. S6, TADs from two cell types were paired based on the highest overlap  
744 (Jaccard Index of genomic coordinates), and the distribution of pairwise overlaps were plotted.  
745 TADs overlapping or around regions masked by matrix balancing were not considered, as the  
746 insulation score can be unreliable in these regions.

747 The analysis is encapsulated by workflow tad\_analysis.nf which was run by launching script  
748 run\_tad\_analysis.sh.

749

### 750 **TAD clique analyses**

751 The TAD annotation generated by HiCEXplorer was given as input to workflow  
752 [github.com/robomics/call\\_tad\\_cliques](https://github.com/robomics/call_tad_cliques) v0.5.1 ([doi.org/10.5281/zenodo.12689353](https://doi.org/10.5281/zenodo.12689353)) to identify  
753 cliques of TADs as outlined in Paulsen et al. (Paulsen et al. 2019). TAD cliques were called  
754 across all three cell types using TADs from 10A. This is a necessary simplification required to  
755 allow comparing cliques across cell types in downstream analyses. The output of  
756 [github.com/robomics/call\\_tad\\_cliques](https://github.com/robomics/call_tad_cliques) was used as input for workflow

757 postprocess\_call\_tad\_cliques.nf, which generated the draft version of Fig. 3B, 3C, 3D and  
758 Supplemental Figs. S31,S33-39.

759 Workflows were launched by running script run\_call\_tad\_cliques\_workflow.sh.

760

### 761 **Subcompartment analyses**

762 Subcompartments were determined using a patched version of dcHiC d4eb244 (Chakraborty  
763 et al. 2022). A and B compartment annotation was generated from subcompartments by  
764 aggregating A-subcompartments and B-subcompartments. The dcHiC processing pipeline  
765 was wrapped in workflow compartment\_analysis.nf, which converts matrices in .mcool file to  
766 a format understood by dcHiC and runs all analysis steps except those for --pcatype=trans, -  
767 -pcatype=fithic, --pcatype=dloop, and --pcatype=enrich. When appropriate, the reference  
768 genome assembly, annotation and .chrom.sizes for hg38.p14 were provided to dcHiC through  
769 the --gfolder option. We fixed the seed used by dcHiC to ensure our results are reproducible  
770 by others. Subcompartments were called at the following resolutions: 10 kbp, 20 kbp, 50 kbp,  
771 100 kbp, 200 kbp and 500 kbp. However, only subcompartments at 10 kbp resolution were  
772 used for further data analyses. The same workflow was used to generate the draft figures for  
773 Fig. 1B and 1D as well as Supplemental Figs. S16-19. Fig. 1C was generated using the output  
774 of the viz step of dcHiC as the starting point.

775 Figure S9-11 were generated by tracking subcompartment assignments across cell types.  
776 Bins were paired based on their genomic coordinates, then subcompartment switches were  
777 classified into 5 classes: neutral (no switch), reverted (e.g. switch occurred in T1 but was  
778 reverted in C1), transition to open/close and other switches (e.g. partial reversion:  
779 A3→A1→A2). Subcompartment switches were further classified using a delta score  $\Delta_{ij}$  defined  
780 as follows. We quantify the degree of switching between two stages, i and j ( $\Delta_{ij}$ ), by assigning  
781 ranks to the subcompartments (B3=0...A3=7) and calculating the rank difference. A negative  
782 rank difference ( $\Delta_{ij}$ ) thus indicates a transition towards B-type subcompartments. Conversely,  
783 a positive value signifies a shift towards A-type compartments.

784 Data analysis steps are defined in workflow compartment\_analysis which was run by  
785 executing script run\_compartment\_analysis.sh.

786

### 787 **Virtual 4C**

788 Draft virtual 4C figures (Fig. 4D; bottom and Suppl. Fig. S48B) have been generated using  
789 notebook virtual\_4c\_myc.ipynb.

790 The two figures were generated by fetching and normalizing interactions for the following  
791 regions from the merged Hi-C maps for 10A, T1, and C1:

792 ○ Fig. 4D; bottom:

793 "chr8:127736000-127737000" (MYC promoter) and

794 "chr10:71280000-73310000" (10kb resolution)

795 ○ Suppl. Fig. 48B:

796 "chr8:127736000-127737000" (MYC promoter) and

797 "chr10:73240000-73270000" (5kb resolution)

798

### 799 **Differential expression analyses**

800 Differential expression analysis was performed using DESeq2 v1.38.0 and apeglm v1.20.0  
801 (Love et al. 2014; Zhu et al. 2019).

802 DESeq2 was called with default parameters using the raw count table produced by nf-  
803 core/rnaseq as input. Log<sub>2</sub>-fold-change shrinkage estimation was computed with the lfcShrink

804 function from DESeq2 using apeglm as shrinkage estimator and the following log2  
805 FoldChange cutoffs: 0.0, 0.1, 0.25, 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5, 5.0.

806 The analysis was automated using workflow `diff_expression_analysis.nf`, which was launched  
807 with `run_diff_expression_analysis.sh`.

808 Supplemental Fig. S28 was generated by running script `run_cluster_profiler_do.py`. The script  
809 uses clusterProfiler v4.8.1 (Wu et al. 2021; Yu et al. 2012) and the disease ontology (DO)  
810 database from the DOSE v3.26.1 package (Yu et al. 2015) to perform over-representation  
811 analysis of DO terms (Schriml et al. 2023). Package enrichplot v1.20.0 (Wu et al. 2021) was  
812 used for visualization. The IDs of differentially expressed genes were provided as input to  
813 clusterProfiler. Genes were considered as differentially expressed based if they exhibit an  
814 absolute log2FoldChange value of 0.5 or greater and a p-value of 0.01 or smaller (correcting  
815 for multiple testing). Furthermore, clusterProfiler was run using a q-value of 0.05.

816

### 817 **Comparative analyses**

818 All comparative analyses were performed using workflow `comparative_analysis.nf`.

819 Fig. S15 was generated by overlapping subcompartment labels with several different  
820 epigenetic markers. First, each ChIP-seq peak was assigned a score by summing the raw  
821 ChIP-seq signal over each peak, then each peak was assigned to a subcompartment. Finally  
822 peaks were grouped by subcompartment label and the mean signal was computed for each  
823 subcompartment.

824 Fig. 2A and Supplemental Fig. S29 were generated by overlapping expression levels in  
825 TPMs with subcompartment labels overlapping genes from GENCODE v43 for hg38. In case  
826 a gene was tagged with multiple subcompartment labels, the gene was assigned the label with  
827 the largest coverage. In case of coverage tie, the gene was discarded (note that this is an  
828 extremely rare occurrence).

829 Figure 2B was generated using pyGenomeTracks (Ramírez et al. 2018; Lopez-Delisle et  
830 al. 2021) to visualize up/down regulated genes overlapping a region involved in  
831 subcompartment switching.

832 Figures 2C-F were generated by overlapping subcompartments with differentially  
833 expressed genes across all three cell types. First genes were assigned a subcompartment  
834 label by overlapping their TSS with the subcompartment annotation generated by dcHiC. Next  
835 genes were grouped based on log2FoldChange and p-value in downregulated, non-  
836 differentially expressed and upregulated genes ( $lfc=2.0$ ;  $pvalue=0.01$ ). Finally each group of  
837 genes was plotted as heatmaps showing subcompartment switches across pairs of cell types  
838 (genes not involved in subcompartment switches are not shown). Figures 2D and 2F were  
839 generated from the table underlying heatmaps 2C and 2E as follows: compute the sum of  
840 genes for each diagonal  $i$ ; starting from diagonal  $i = 1$ , pair diagonal  $i$  with diagonal  $-i$ ;  
841 finally compute the  $\log_2$  ratio between the sum of diagonal  $i$  and the sum of diagonal  $-i$ .  
842 Positive values indicate a positive correlation between one of the classes of differentially  
843 expressed genes and switches towards open chromatin while negative values indicate a  
844 positive correlation with subcompartment switches towards close chromatin.

845 The draft figure for Fig. 3A was generated with `bin/plotting/plot_tad_cliques.py` using TAD  
846 cliques, TADs, compartment PCA and GENCODE v43 gene annotation as input.

847 The draft figure for Supplemental Fig. S32 was generated by overlapping subcompartment  
848 states with TADs annotated with the size of the largest clique to which they belonged to.

849 The draft figures for Fig. 3F-G and Supplemental Fig. S40 were generated by clustering  
850 TADs and TAD cliques using HDBSCAN (McInnes et al. 2017; McInnes and Healy 2017)  
851 based on their subcompartment composition.

852 First, domains (that is TADs or TAD cliques) were annotated with their subcompartment state  
853 composition by overlapping subcompartment states with the domains using  
854 `annotate_domains_with_subcompartments.py`. This resulted in a count matrix with one row  
855 per domain, where each row counts the number of bins labeled for each subcompartment  
856 state. For example, given a domain overlapping 3 A1 bins, 10 A2 bins and 2 A3 bins, the entry  
857 in the count matrix corresponding to this domain would be 0, 0, 0, 0, 0, 3, 10, 2.

858 Next, this count matrix was given as input to `cluster_domains_by_subcompartment_state.py`,  
859 which clustered domains using HDBSCAN.flat based on their similarity in subcompartment  
860 composition. Clustering was performed using the following settings: `n_clusters=9`;  
861 `metric=euclidean`; `min_cluster_size=200`; `min_samples=5`; `cluster_selection_method=leaf`.  
862 Finally, clusters were visualized using `plot_domain_subcompartment_clusters.py`.

863 All comparative analyses are defined in workflow `comparative_analysis.nf` which was  
864 launched with script `run_comparative_analysis.sh`.

865

### 866 **3D genome modeling**

867 A patched version of Chrom3D v1.0.2 was used to generate 3D genome models for 10A, T1  
868 and C1 (Paulsen et al. 2017). Briefly, this Nextflow workflow runs [NCHG](https://doi.org/10.5281/zenodo.12680450)  
869 ([doi.org/10.5281/zenodo.12680450](https://doi.org/10.5281/zenodo.12680450)) to identify statistically significant cis and trans interactions.  
870 Translocated regions were excluded when identifying statistically significant trans interactions.  
871 For intra-chromosomal interactions we used a log-ratio of 1.2 and adjusted-pvalue of 0.01 as  
872 cutoffs, while for inter-chromosomal interactions a log-ratio of 0.25 and adjusted-pvalue of  
873 0.01 were used as cutoffs. The workflow then uses statistically significant interactions in  
874 Chrom3D simulations as spatially proximal interaction constraints. Furthermore, the workflow  
875 matches LADs to the corresponding TAD domains such that these domains can be used as  
876 peripheral sub-nuclear constraints in the simulations. A total of 100 Chrom3D simulations were  
877 run for each condition with a nuclear occupancy of 0.15, at TAD resolution and 2 million  
878 simulation steps (Paulsen et al. 2017).

879 ChimeraX was used to visualize the simulated Chrom3D models (Pettersen et al. 2021). The  
880 median distance of each chromosome from the nuclear center was calculated for each  
881 Chrom3D simulation and visualized using `ggplot2` (Villanueva and Chen 2019). The median  
882 distance and the standard deviation of each subcompartment were likewise evaluated for each  
883 Chrom3D simulation and visualized using `ggplot2`. Subcompartment distances from the  
884 nuclear center were statistically evaluated and compared within conditions using the Wilcoxon  
885 rank sum test in R (R Core Team 2024) with the function `wilcox.test`.

886

### 887 **FISH analysis**

888 Analysis of FISH data is automated using workflow `fish.nf`. Draft plots have been generated  
889 using notebooks `plot_fish_blob_distance_stats.ipynb` and  
890 `plot_fish_blob_radial_positioning.ipynb`.

891

892 The FISH data analysis consists of three steps:

- 893 - Nuclei segmentation
- 894 - Probe localization
- 895 - Plot generation and statistical tests

896

897 Nuclei segmentation

898 Nuclei segmentation is performed using DAPI signal (i.e. the blue channel of the RGB image  
899 produced by the microscope).

900 First, RGB images containing one or more nuclei are converted to grayscale by extracting the  
901 blue channel from the images. Grayscale images are then sharpened using unsharp masking.  
902 This is done to improve the separation of the profiles of nuclei that are located very close to  
903 one another. Next, foreground objects (i.e. nuclei profiles) are highlighted using Otsu  
904 thresholding (Otsu 1979). Objects that are significantly smaller than plausible nuclei profiles  
905 are masked out. Next, we compute the contour of each individual object using the “marching  
906 squares” method. Finally, object contours are used to crop out individual nuclei.  
907 To properly deal with various edge cases (such as mitotic nuclei and overlapping nuclei),  
908 nuclei whose contour significantly deviates from an ellipsoid are flagged so that they can be  
909 skipped in subsequent analysis steps. This is achieved by first fitting an ellipse to the contour  
910 detected using the first algorithm described in (Fitzgibbon and Fisher). Next, we compute the  
911 overlap coefficient (OC) between the fitted ellipse and the nucleus contour, and mask nuclei  
912 with an OC of 0.95 or lower. Finally, nuclei that are only partially visible in an image are also  
913 flagged.

914

#### 915 Probe localization

916 For each nucleus, we extract the red and green channels from the RGB image produced by  
917 the segmentation step. The resulting grayscale images are processed independently as  
918 follows.

919 Given a probe and a cell line, we can estimate the number of probe signals we expect to find  
920 in each nucleus. This information is used to guide the blob (i.e. probe signal) detection using  
921 the Laplacian of Gaussian (LoG) method. We perform a sweep of LoG parameters, starting  
922 from parameters that are suitable to detect large blobs in an image. If the first run of the LoG  
923 algorithm detects  $N$  or more blobs, where  $N$  is the number of expected probe signals, then the  
924 parameter sweep is halted. Otherwise, the coordinates of detected blobs (if any) are stored,  
925 and the LoG algorithm is re-run using parameters to detect progressively smaller blobs. Every  
926 iteration, the coordinates of newly detected blobs are merged with the coordinates of blobs  
927 detected in previous runs of the algorithm. The parameter sweep stops as soon as  $N$  or more  
928 blobs are detected, or when the parameter space is exhausted.

929

#### 930 Plot generation and statistical tests

931 The coordinates of blobs identified in the previous step are finally used to generate Fig. 4F  
932 and Supplemental Fig. S47.

933

934 The number of nuclei and observations for each condition is shown in Suppl. Table S18.

935

#### 936 Color mapping strategy

937 We provide colorblind-friendly versions of the raw and processed FISH images.  
938 These images were generated using script `color_mapping_fish.py`.  
939 In brief, this script applies a local linear mapping to convert the original colors (blue for DAPI  
940 and red/green for the FISH probes) to different colors that can be more easily distinguished  
941 by those affected by common forms of colorblindness.  
942 The mapping is local because we only map colors for pixels whose original color is  
943 perceptually different from black and white.

944

#### 945 **Software used throughout data analysis code**

946 The following software packages were used throughout the code used for data analysis:

- 947 ● BEDTools (Quinlan and Hall 2010) - to perform common set operations on genomic  
948 intervals.
- 949 ● bedGraphToBigWig (Kent et al. 2010) - to convert bedGraph files to bigWig format.
- 950 ● bioframe (Open2C et al. 2024) to perform common set operations on genomic  
951 intervals.
- 952 ● hickpy (Rossini and Paulsen 2024) - to perform certain input-output operations on  
953 .cool files.
- 954 ● Matplotlib (Hunter May-June 2007) - to generate plots.
- 955 ● NumPy (Harris et al. 2020) - to efficiently perform arithmetic operations on vector data.
- 956 ● OpenCV - to manipulate and process microscopy images.
- 957 ● pandas (McKinney 2010) - to read, write and manipulate tabular data using  
958 dataframes.
- 959 ● pyBigWig - to read and write bigWig files.
- 960 ● SAMtools (Danecek et al. 2021) - to index FASTA files and perform IO operations on  
961 SAM, BAM and CRAM files.
- 962 ● scikit-image (van der Walt et al. 2014) - to manipulate and process microscopy  
963 images.
- 964 ● SciPy (Virtanen et al. 2020) - to compute Pearson's correlation values and perform  
965 statistical tests.
- 966 ● seaborn (Waskom 2021) - to generate plots.

967

968 **Software patches**969 This section describes the purpose of the patches we applied to Chrom3D, cooler, dcHiC, and  
970 HiNT. Patch files are available under containers/patches.

971

972 Patch for nf-core/hic v2.0.0 (<https://zenodo.org/records/2669513>)973 Our patch involved updating the pipeline to correctly handle the --restriction\_site and --  
974 ligation\_site parameters.975 Our patch was accepted by upstream and is now part of v2.1.0 of the pipeline. See [nf-  
976 core/hic/pull/153](https://nf-core.org/hic/pull/153) for more details.

977

978 Patches for dcHiC d4eb244 (Chakraborty et al. 2022)

979 We patched dcHiC as follows:

- 980 ● Define a default seed and introduce a --seed CLI option to ensure results are  
981 reproducible across runs
- 982 ● Wrap calls to depmixS4::fit into a try-catch block to attempt model fitting up to 5 times  
983 in case of spurious failures
- 984 ● Properly handle the possibility that a chromosome does not have any bin overlapping  
985 one or more subcompartment type.

986

987 Our patches were accepted by upstream but are not yet part of a stable release. See [ay-  
988 lab/dcHiC/pull/59](https://ay-lab.org/dcHiC/pull/59), [ay-lab/dcHiC/pull/60](https://ay-lab.org/dcHiC/pull/60), [ay-lab/dcHiC/pull/62](https://ay-lab.org/dcHiC/pull/62) and  
989 [containers/patches/dchic.patch](https://containers.cshlp.org/patches/dchic.patch) for more details.

990

991 Patch for cooler v0.9.1 (Abdennur and Mirny 2020)

992 We patched cooler to ensure that convergence of cooler balance using cis-only interactions  
993 was correctly reported for all chromosomes instead of just the last one.

994 When balancing interactions using the --cis-only flag, cooler balances interactions for each  
995 chromosome independently. Given that there is no guarantee that ICE will converge within the  
996 given number of iterations, cooler balance stores an attribute inside balanced cooler files to  
997 report whether balancing was successful (i.e. convergence was achieved).  
998 In cooler v0.9.1 there is a bug in the logic that writes this attribute that causes the convergence  
999 for the last chromosome balanced to be reported instead of the convergence status for each  
1000 chromosome. Our patch addresses this bug such that the convergence status of matrices  
1001 balanced with --cis-only can be assessed correctly.  
1002 The patch also addressed some minor issues related to detecting pandas' version at runtime  
1003 and handling of bin sizes represented using unsigned integers.  
1004 Our patches were accepted by upstream and are now part of cooler v0.9.2.  
1005 See [open2c/cooler/pull/313](https://open2c/cooler/pull/313), [open2c/cooler/pull/323](https://open2c/cooler/pull/323), [open2c/cooler/pull/324](https://open2c/cooler/pull/324) and  
1006 [containers/patches/cooler.patch](https://containers/patches/cooler.patch) for more details.

1007  
1008 Patch for Chrom3D v1.0.2 (Paulsen et al. 2017)

1009 We patched Chrom3D to support building the project using CMake and addressing a few  
1010 compiler warnings and errors that were preventing us from compiling Chrom3D with a modern  
1011 C++ compiler toolchain. For more details, refer to the following two patch files available at  
1012 [robomics/chrom3d-nf](https://robomics/chrom3d-nf) ([doi.org/10.5281/zenodo.12687842](https://doi.org/10.5281/zenodo.12687842)) [chrom3d-cmake.patch](https://chrom3d-cmake.patch), [chrom3d-](https://chrom3d-fix-warnings.patch)  
1013 [fix-warnings.patch](https://chrom3d-fix-warnings.patch).

1014  
1015 Patch for HiNT v2.2.8 (Wang et al. 2020)

1016 We patched HiNT to support processing Hi-C matrices using Arima restriction enzymes. See  
1017 [containers/patches/hint.patch](https://containers/patches/hint.patch) for more details.

## 1019 Data Access

1020 All raw and processed sequencing data generated in this study have been submitted to the  
1021 NCBI Gene Expression Omnibus (GEO; <https://www.ncbi.nlm.nih.gov/geo/>) under the  
1022 following accession numbers: GSE246689, GSE246599, GSE247171, and GSE246947.

1023 All raw and processed microscopy data generated in this study have been submitted to  
1024 Zenodo (<https://zenodo.org/>) and are available for download at the following DOI:  
1025 [10.5281/zenodo.15167335](https://doi.org/10.5281/zenodo.15167335).

## 1027 Competing Interests

1028 None declared

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1046

## 1047 **Authors' contributions**

1048 JP, PC, DT conceived and designed the study; JP, RR and MO designed data analyses; RR  
 1049 and MO analyzed data; RR processed Hi-C, RNA-seq, ChIP-seq, and FISH data and did  
 1050 downstream analyses; MO performed 3D genome modeling; MN did Hi-C experiments; RD  
 1051 and YD did cell culture work and RNA-seq; AB and MA analyzed LADs; RE conceived and  
 1052 designed CRISPR-Cas9 RE strategy, verified clones, analyzed data and generated figures;  
 1053 ML performed cloning of Lenti-multi-CRISPR constructs and PCR; RA did CRISPR-Cas9 time-  
 1054 course, RNA and genomic extraction, qRT-PCR, PCR and analysis; JP, PC, DT, RE  
 1055 supervised the work. All authors read and approved the final manuscript.

1056

1057

## 1058 **References**

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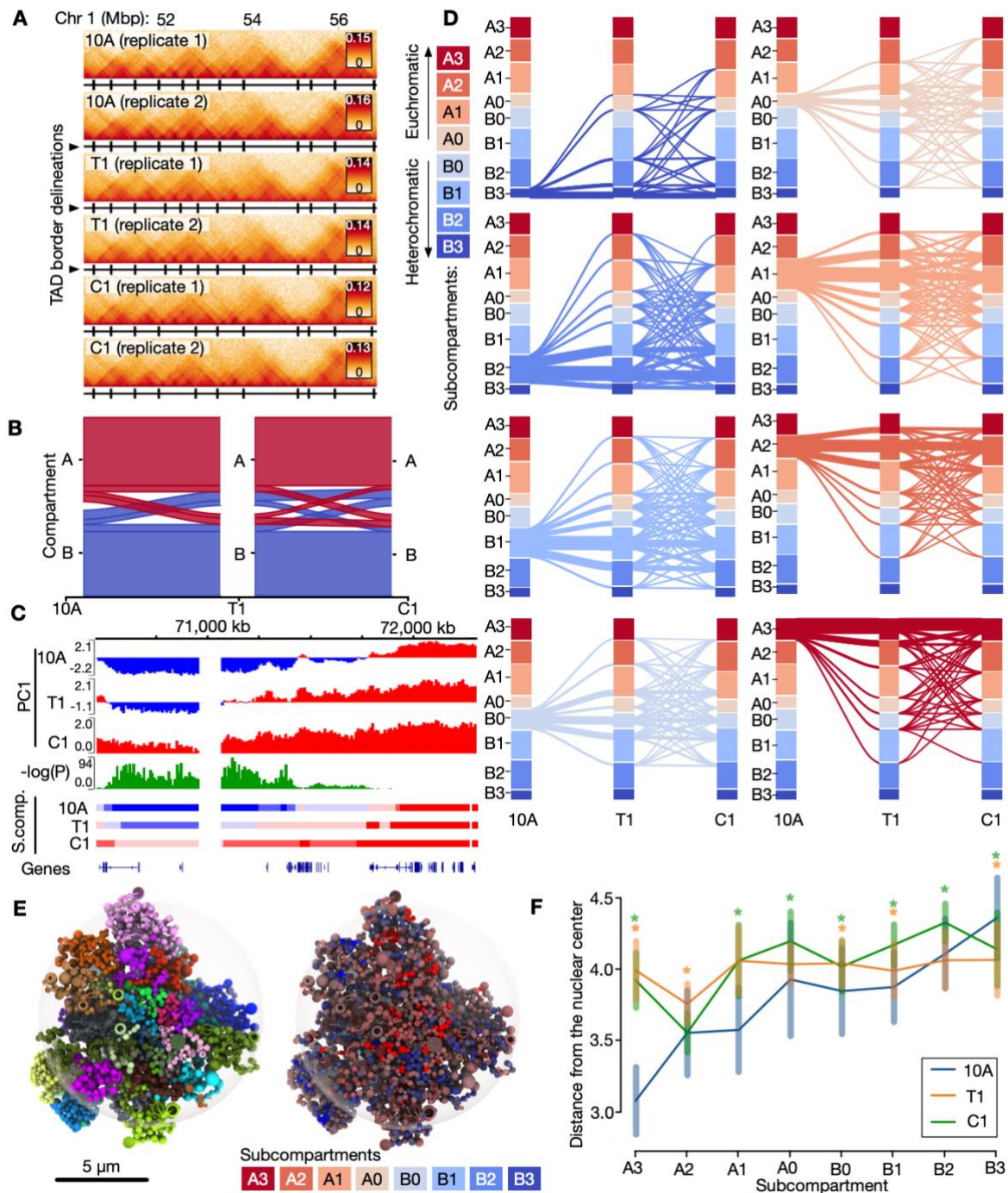
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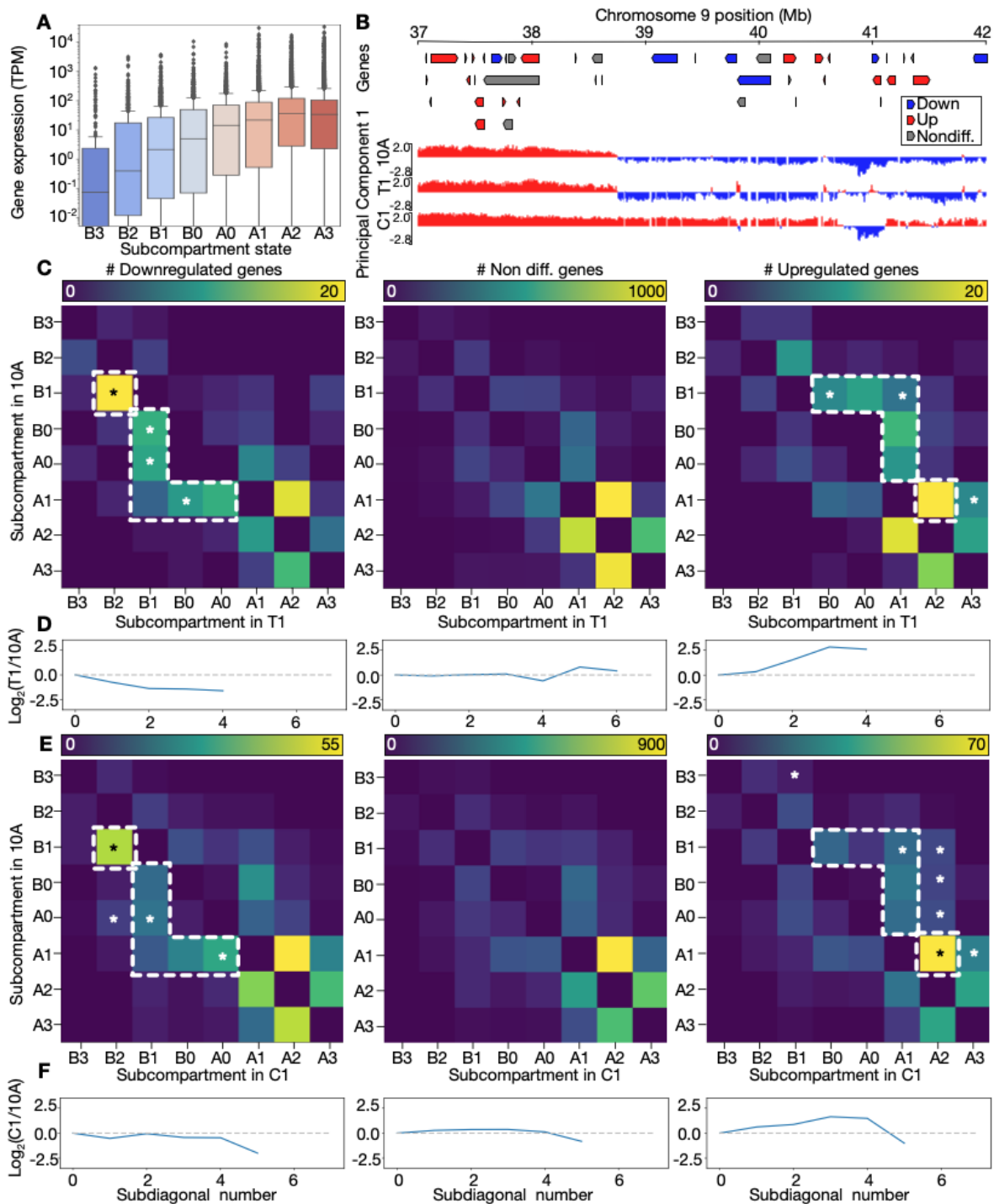
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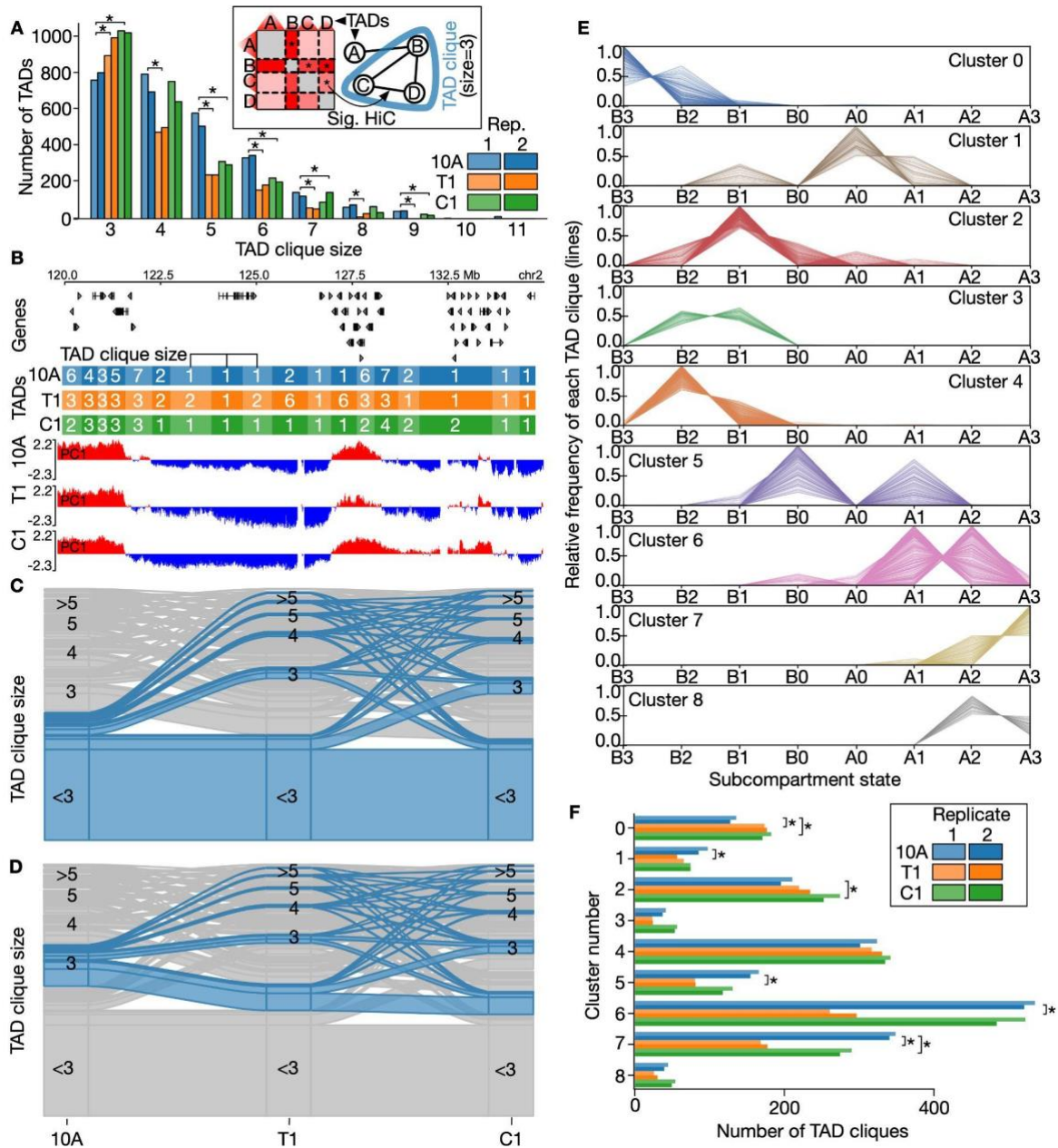
1452 **Figures**

**Fig. 1.** Alterations in genome compartment properties during breast cancer progression stages. **A:** Example Hi-C data for a region on Chromosome 1. TAD delineations are shown as black vertical lines below each track. **B:** Alluvial plot showing A/B compartment conservation and switching during progression from 10A (left), via T1 (middle) and to C1 (right). **C:** Example region on Chromosome 11 showing a statistically significant switch in the first principal component (PC1) in the three stages. Corresponding subcompartments in the three stages (10A, T1, C1) are shown below. **D:** Switching between four A-type subcompartments (A0-A3), and four B-type (B0-B3) called from the Hi-C data (shown in distinct colors). Subcompartment switches across 10A, T1 and C1 shown as separate alluvial plots starting at each of the eight different subcompartments in 10A. **E:** Left: Tomographic view of exemplary Chrom3D model from 10A cells with chromosomes colored individually. Right: The same model with regions colored by their subcompartment associations. **F:** Median distance (with standard deviation) from the nuclear center for each subcompartment in each condition. Asterisks indicate significant differences in T1 and C1 compared to 10A (see Supplemental Table S4)

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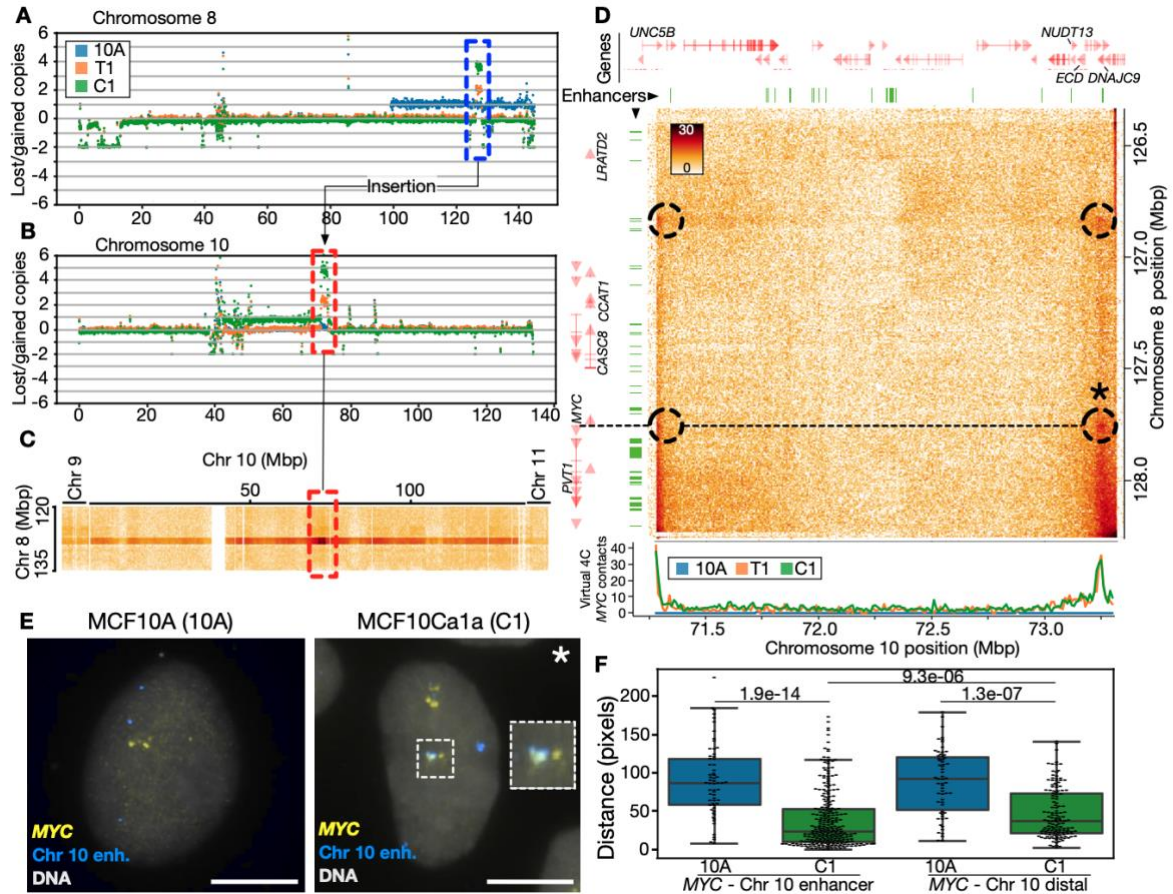


**Fig. 2.** Coordinated changes in subcompartments and gene expression. **A:** Gene expression levels (TPM) by subcompartment state (protein-coding genes; MCF10A [replicates merged]). Vertical axis in log scale. **B:** Example of subcompartment switching on Chromosome 9. Up- and down-regulated genes in C1 vs. 10A indicated in red and blue, respectively. **C:** Heatmaps showing number of DE downregulated genes (left panel), non-differential genes (middle panel) and upregulated genes (right panel) in subcompartments switching between 10A (vertical axis) and T1 (horizontal axis). Dotted lines highlight regions with enrichment relative to non-differential and upregulated genes. Asterisks indicate significant differences (see Suppl. Table S6) **D:**  $\text{Log}_2$ -ratio of number of genes in subdiagonal sums in the upper vs. lower triangular of the corresponding heatmap from C. Ratio not displayed for subdiagonal sums that are equal to zero. **E:** Heatmaps as in C but contrasting 10A with C1. Dotted lines highlight regions with enrichment relative to non-differential and downregulated genes. Asterisks indicate significant differences (see Suppl. Table S7) **F:**  $\text{Log}_2$ -ratio plots as in D, but contrasting 10A with C1.

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**Fig. 3.** TAD clique dynamics during breast cancer progression stages. **A:** Absolute number of TADs in cliques with maximal size ranging from 3 to 11 shown for each of the three breast cancer stages. Asterisks indicate significant differences (see Suppl. Table S10). Inset illustrates how “TAD cliques” are identified from the statistically significant HiC-interactions between TADs. **B:** Example region on Chromosome 2 showing changes in TAD clique sizes and compartment eigenvectors. **C:** Alluvial plots highlighting the alluvial path of a non-clique in 10A across the three cancer stages. **D:** Alluvial path of a TAD clique of size 3 in 10A. **E:** TAD clique clustering into 9 clusters based on their enrichment in subcompartments in the same cell type (vertical axis). Each line represents a TAD clique. **F:** Numbers of TAD cliques belonging to each of the clusters in E. Asterisks indicate significant differences (see Suppl. Table S12). See Suppl. Fig. S40 for numbers in the “mixed” cluster.

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**Fig. 4.** *MYC* locus insertion on Chromosome 10 is accompanied by *de novo* contacts with a potential enhancer element. **A:** Lost and gained copies for the entire Chromosome 8 in 10A, T1, and C1 cells. The dotted blue box highlights specific amplification of the Chromosome 10 region. **B:** Lost and gained copies for the entire Chromosome 10, with a dotted red box highlighting specific amplifications. **C:** Interchromosomal C1 Hi-C contacts between the region on Chromosome 8 (vertical axis) and the entire Chromosome 10. Highlighted region from B indicated in red. End of Chromosome 9 and beginning of Chromosome 11 shown on the left and right side, respectively. **D:** Top: Zoom-in on the C1 Hi-C map of the Chromosome 8/10 amplification unit. Dotted circles indicate enriched “dots” of contacts involving enhancers and *MYC*. Asterisk (\*) marks the enhancer-interaction validated in FISH in panels E-F. The dotted line shows the position of the *MYC* gene on Chromosome 8 relative to contacts on Chromosome 10 within the amplification unit. Positions of enhancers from MCF10A indicated as green segments (top and left). Genes shown in red (top and left), with names for selected genes. Bottom: Virtual 4C track (see Methods) showing Hi-C contacts from *MYC* towards the chromosome 10 region for 10A (blue), T1 (yellow) and C1 (green). **E:** FISH image of the *MYC* gene (yellow) and the Chromosome 10 enhancer (blue; Chr. 10 position 73117711-73267436) expected to interact with *MYC* from the Hi-C data, in 10A and C1 cells. Bars, 7  $\mu$ m. **F:** Quantification of probe-probe distances in pixels, for the *MYC*-Chr10 enhancer probe pair (left; n = 62 [10A] and 269 [C1] observations) and the *MYC*-Chr10 distal probe pair (position 73794997-73992612; n = 62 [10A] and 140 [C1] observations); N = 28-95 nuclei analyzed; P-values are shown (Kolmogorov-Smirnov test).