## 1 High-resolution CTCF footprinting reveals impact of chromatin state on

## 2 cohesin extrusion dynamics

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- 26

## 27 Abstract

28

29 DNA looping is vital for establishing many enhancer-promoter interactions. While CTCF is

30 known to anchor many cohesin-mediated loops, the looped chromatin fiber appears to

- 31 predominantly exist in a poorly characterized actively extruding state. To better characterize
- 32 extruding chromatin loop structures, we used CTCF MNase HiChIP data to determine both
- 33 CTCF binding at high resolution and 3D contact information. Here we present *FactorFinder*, a
- 34 tool that identifies CTCF binding sites at near base-pair resolution. We leverage this substantial
- advance in resolution to determine that the fully extruded (CTCF-CTCF) state is rare genome-
- 36 wide with locus-specific variation from  $\sim$ 1-10%. We further investigate the impact of chromatin
- 37 state on loop extrusion dynamics, and find that active enhancers and RNA Pol II impede cohesin
- **38** extrusion, facilitating an enrichment of enhancer-promoter contacts in the partially extruded loop
- 39 state. We propose a model of topological regulation whereby the transient, partially extruded
- 40 states play active roles in transcription.

#### 41 Background

42

43 Topologically associated domains (TADs) and regulatory enhancer-promoter chromatin loops

44 are largely formed by the cohesin complex through the process of CTCF-mediated loop

45 extrusion<sup>1,2</sup>. Topological alterations and subsequent changes in enhancer-promoter (EP) contacts

46 can modify gene expression<sup>3,4</sup> and cause aberrant phenotypes<sup>5–8</sup>. CCCTC-binding factor (CTCF)

47 can act as an extrusion barrier through its ability to bind and stabilize cohesin on DNA, serving

to preferentially localize and anchor one or both ends of cohesin loops. Genes with promoter proximal CTCF binding sites have been shown to have increased dependence on distal

proximation of on the proximation of t

51

52 Although conventional 3C techniques give an impression of static loops, cohesin-mediated

53 chromatin loops are actually dynamic with an extrusion rate of  $\sim 1$ kb/s<sup>12</sup>. Recent live cell-

54 imaging studies of two TADs found that the fully extruded state with a loop formed between two

55 convergent CTCF-bound anchors was present only 3-30% of the time<sup>13,14</sup>. While these findings

suggest that CTCF loops spend the vast majority of their time partially-extruded, the partially-

57 extruded state has not yet been well characterized.

58

59 Several studies have found evidence of promoter-proximal CTCF binding sites (CBS) having

60 large impacts on EP contact frequencies and transcription<sup>9–11</sup>. Putting this together with the high

61 prevalence of partially extruded CTCF-mediated loops, we hypothesize that promoter-proximal

62 CTCF sites enable gene regulation by halting cohesin on one side while cohesin continues to

extrude on the other side. Enhancers then slow down extrusion, thus enabling an increase in EP

64 contacts without requiring a fully extruded loop. The relationship between EP contacts and

65 transcription can be nonlinear such that small increases in EP contacts may cause large changes

66 in transcription<sup>3,4</sup>. As a result, even minor decreases in extrusion rate through enhancer regions

- 67 may affect gene expression.
- 68

69 The ability of MNase to efficiently digest naked DNA while sparing protein-bound DNA has

70 been employed in various strategies to footprint the binding sites of proteins such as transcription

factors with near base-pair resolution 15-18. A key advantage of using MNase over sonication-

72 based protocols is the shorter fragment size obtained, which directly leads to higher resolution

73 TF binding site identification. More recently, MNase DNA fragmentation has also been applied

to proximity ligation assays to map 3D genome architecture with nucleosome (~150 bp)

resolution, enabling precise characterization of 3D architecture including at TAD boundaries and

76 punctate enhancer-promoter interactions<sup>19–22</sup>. Since MNase HiChIP enables precise

characterization of both TF-binding and 3D contacts, it is uniquely poised to define how CTCF

renables 3D contacts.

79

- 80 To better characterize the partially extruded chromatin loop state, we first develop a
- 81 computational technique for high-resolution footprinting of CTCF using MNase HiChIP data.
- 82 We then employ this to study how, through its interaction with the looping factor cohesin, CTCF
- 83 can facilitate long-range DNA contacts. We further characterize how the length of loops
- 84 extruded by cohesin is affected by local chromatin state factors such as enhancer and RNA Pol II
- 85 density.
- 86

### 87 **Results**

88

89 <u>MNase HiChIP generates short, TF-protected and longer, histone-protected DNA</u>

90 *fragments* 

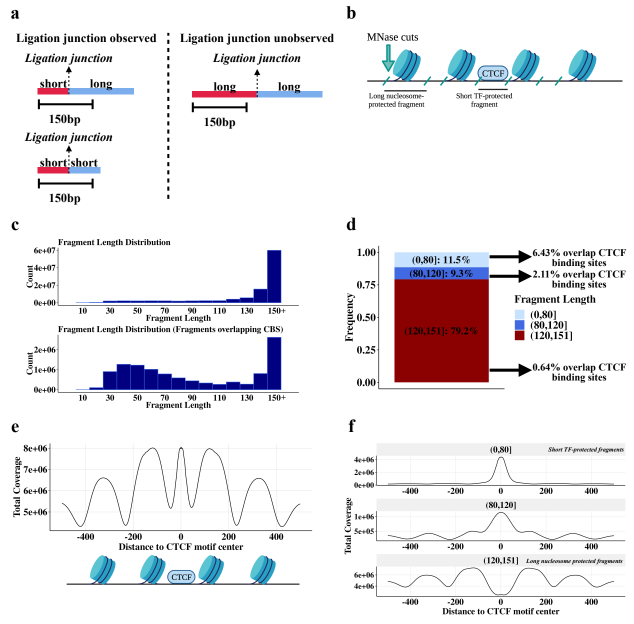
91 We used Micrococcal nuclease (MNase) HiChIP<sup>23</sup> with a CTCF antibody to profile 3D

- architecture in K562 cells, generating 150 bp reads with over 380 million unique pairwise
- 93 contacts across four replicates. Briefly, following cell fixation with DSG and formaldehyde,
- 94 chromatin is digested by MNase, immunoprecipitated to enrich for CTCF-bound DNA, and free
- 95 ends are then ligated. After reverse-crosslinking, the resulting ligation products are sequenced
- from both ends and the mapping locations of the paired reads can be used to infer chromosomal
- 97 locations of the physically interacting loci. In cases where the pre-ligation fragments are shorter
- than the read length it is also possible to infer the fragment length as the ligation junction
- 99 position will be observed within one or both of the reads. If multiple fragments within a read are 100 short enough to be aligned to distinct genomic locations, this is termed an 'observed ligation'
- 101 (Fig. 1a, Supp Fig 1).
- 102

As expected, due to the preference of MNase to selectively cleave DNA not shielded by boundproteins and the high abundance of histones in chromatin (Fig. 1b), the predominant fragment

- 105 length is approximately 150 bp, indicative of cuts between nucleosomes<sup>24</sup> (Fig 1c). We also
- noted a distribution of shorter fragment lengths, with 20% representing lengths shorter than 120
- bp (Fig. 1d). A metaplot centered on CTCF binding site motifs shows an enrichment of 30-60 bp
- 108 fragments suggesting that these shorter fragments represent CTCF-bound DNA (Fig. 1c) $^{2,25,26}$ .
- 109 Consistent with this, we find that short (<80 bp) fragments have a 10-fold higher overlap
- 110 frequency with CTCF motifs than long (>120 bp) fragments (Fig. 1d). This is similar to data
- 111 from the MNase-based CUT&RUN assay that also results in short fragments protected by small
- 112 proteins such as transcription factors $^{17}$ .
- 113
- 114 Fragment pileups at CTCF motif loci (Fig. 1e) show a strong enrichment of short fragments
- 115 centered on the CTCF motif sequence, and a concomitant depletion of long fragments at motifs
- 116 (Fig. 1f). Long fragments, in contrast, show peaks with a strong ~200 bp periodicity adjacent to
- the central CTCF binding site (Fig. 1f). This is consistent with the ability of CTCF to precisely
- 118 position a series of nucleosomes adjacent to its binding site<sup>25</sup>. Note that while long (>120 bp)
- 119 fragments are depleted at CTCF binding sites, they still represent a significant fraction of reads at

- 120 these sites (Fig. 1c). This likely reflects that CTCF motif loci without a bound CTCF are
- 121 frequently instead occupied by histones<sup>25</sup>, and even CTCF motifs with very strong CTCF ChIP-
- 122 seq signal are not always occupied by a CTCF.
- 123
- 124 In summary, long fragments correspond to nucleosome-protected DNA whereas short fragments
- arise from TF-protected DNA. This is due to the different sizes of CTCF and histone octamers,
- with nucleosomes protecting about twice the amount of DNA that CTCF protects<sup>25</sup>. Since
- 127 MNase cuts around bound proteins, the different protein sizes directly translate to different
- 128 fragment lengths. Accordingly, we next filter out long, nucleosome-protected fragments and
- 129 focus on short, TF-protected fragments to identify CBS.





131 Fig. 1 MNase CTCF HiChIP data contains short (~ <80 bp) CTCF-protected fragments and

- 132 longer (~>120 bp) nucleosome-protected fragments. **a** Schematic illustrating relationship
- between short fragments and observed ligations. **b** Schematic illustrating how the fragment
- 134 length results from MNase cutting around bound proteins of different sizes. **c** Fragment length
- distribution for all fragments (top plot) and fragments overlapping occupied CTCF motifs (lower
- 136 plot). Occupied CTCF motifs are defined here as CTCF motifs within 30 bp of a CTCF ChIP-seq
- peak summit. d Boxplot quantifying the frequency of different fragment lengths genome-wideand how often each fragment length group overlaps an occupied CTCF motif. Occupied CTCF
- 139 motifs are defined here as CTCF motifs within 30 bp of a CTCF ChIP-seq peak summit. e
- Fragment coverage metaplot +/- 500 bp around CTCF binding sites. Schematic below the
- 141 coverage metaplot illustrates the proteins producing these peaks. **f** Plot (**e**) stratified by fragment
- 142 length.
- 143

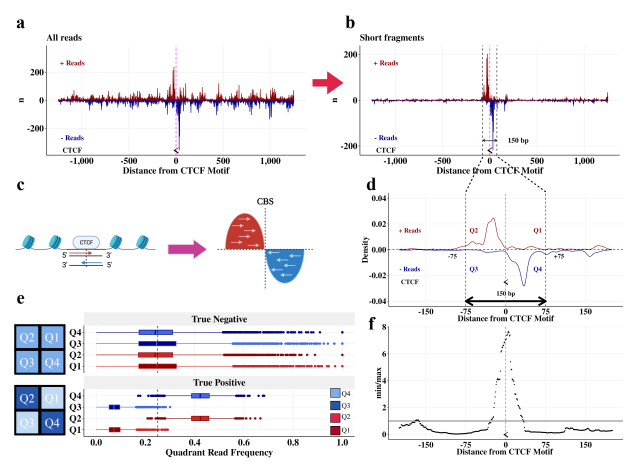
# 144 <u>FactorFinder leverages the strand-specific bimodal distribution of short fragments</u> 145 around CBS to obtain precise detection of CTCF binding

- 146 In order to characterize CTCF-mediated chromatin loop interactions, we first set out to map
- 147 CTCF loop anchors with high resolution. We take advantage of the difference in fragment
- 148 lengths associated with CTCF-bound vs nucleosome-bound DNA to focus only on likely CTCF-
- bound fragments. Fragment lengths can be determined for all fragments with length less than 150
- bp; the 150 bp read length results in censoring of fragments longer than 150 bp. While exact
- 151 fragment lengths can be obtained for all fragments shorter than 150 bp, observed ligations
- require a shorter fragment length. This is because observed ligations require distinct mapping of
- 153 fragments on either side of the ligation junction. Since at least  $\sim$ 25 bp are required to align a
- sequence to the reference genome, this results in fragments characterized as observed ligations
- having a maximum fragment length of  $\sim$ 125 bp, sufficient for the identification of most CTCF-
- 156 protected DNA fragments. Consequently, the fraction of informative, CTCF-protected fragments
- decreases with shorter sequencing read length (Supp Fig 1). The effect of subsetting the CTCF
  HiChIP dataset to only short fragments (<125 bp, identified by the proxy of an observed</li>
- 159 ligation), is shown in Fig 2a,b. These shorter, presumably CTCF-protected fragments, are
- 160 overwhelmingly located immediately adjacent to CTCF motifs.
- 161
- 162 Sequencing of short, CTCF-protected fragments results in a bimodal read distribution centered
- 163 on the CBS, with read 5' location peaks observed upstream (positive strand) and downstream
- 164 (negative strand) of the CBS (Fig. 2c). We refer to these regions as quadrants 2 and 4 (Q2 and
- 165 O4) respectively (Fig. 2d, e). In contrast, reads from the positive strand downstream of the CBS
- 166 (Q1) and negative strand upstream of the CBS (Q3) correspond to fragments with MNase cut
- sites underneath CTCF-protected DNA, and therefore reflect a lack of CTCF occupancy. CTCF
- 168 binding therefore produces an enrichment of reads in Q2,Q4 and a depletion of reads in Q1,Q3
- 169 (Fig. 2e). At sites without protein binding, MNase can cut at any location resulting in no
- 170 enrichment of reads in Q2 and Q4 compared to Q1 and Q3 (Fig. 2e). As a result, we can

- determine CTCF binding by testing if there are significantly more reads in Q2 and Q4 than Q1
- 172 and Q3 (Fig. 2f).
- 173
- 174 We can consider each read as an independent draw from a multinomial distribution with four
- 175 categories corresponding to the four quadrants. Under the null hypothesis, each read has equal
- probability of belonging to any of the four quadrants  $Q_i$ ,  $i \in \{1,2,3,4\}$ . Because true CTCF
- 177 binding induces a strong read pile-up in *both* quadrants 2 and 4 in addition to a depletion of reads
- 178 in quadrants 1 and 3 (Fig. 2d, e, f), we test for an enrichment of reads in Q2 and Q4 compared to
- 179 Q1 and Q3 by estimating the *FactorFinder* statistic  $\hat{\alpha} = \frac{\min(n_2, n_4)}{\max(n_1, n_3)}$ , where  $n_i$  is the number of
- 180 reads in  $Q_i$ . We then test if  $\hat{\alpha}$  is significantly greater than 1. Note that min and max are used to
- 181 enforce that both quadrants 2 and 4 must have more reads than both quadrants 1 and 3; using the
- 182 average would enable read pile-ups that occur in quadrant 2 or 4 (but not both) to be spuriously
- 183 called as CTCF binding events.
- 184
- 185 To evaluate the significance of  $\hat{\alpha}$  at a particular total read count  $N = \sum_{i=1}^{4} n_i$ , we simulated 100
- 186 million samples under the null hypothesis that each fragment is equally likely to occur in any of
- 187 the four quadrants. This was done at each total read count ranging from 5 to 500. P-values at read 188 counts beyond 500 are very similar to those at 500, so 500+ read counts are treated as bins with
- 189 500 total read count (Supp Fig 2). The empirical CDF of the 100 million  $log2(\hat{\alpha})$  at a given total
- read count was then computed and used to evaluate the probability of observing a value more
- 191 extreme than  $log2(\hat{\alpha})$  under the null hypothesis. The empirical CDF was evaluated at a sequence
- 192 of possible  $log2(\hat{\alpha})$  between 0 and 5 at step sizes of 0.01(this corresponds to  $\hat{\alpha} \in [1,32]$ .) This
- approach produces the same p-values as using  $\hat{\alpha}$  instead of  $log2(\hat{\alpha})$ , but using the log enables
- 194 smaller step size at large values of  $\hat{\alpha}$ . After acquiring the grid of p-values for each  $\hat{\alpha}$  at a given 195 read count *N*, we match the observed  $\hat{\alpha}$  at a read count of *N* with the corresponding p-value from
- 196 the table. Because this approach only requires quadrant-specific read counts to match with the
- 197 given table of p-values, it is very computationally efficient. Furthermore, by using the
- 198 multinomial framework we place no assumptions on the reads within each quadrant being
- 199 distributed as poisson, negative binomial, or another distribution. The only assumption we make
- 200 is that in the event of no CTCF binding, the reads are equally distributed amongst the four
- 201 quadrants. We have shown this assumption holds in Figures 2c, d, e.
- 202

In brief, we have shown that short fragments exhibit a strand-specific, bimodal distribution centered on the CBS. This distribution arises from MNase cutting around a bound CTCF and

- subsequent sequencing 5' to 3' of the DNA. Significance is assessed through a multinomial
- framework, which has the advantage of not placing any assumptions on the distribution of reads
- within each quadrant. Now that we have explored the theory behind *FactorFinder*, we
- 208 demonstrate its ability to identify CBS with high resolution and accuracy.



#### 209

Fig. 2 True CTCF binding sites have a bimodal strand-specific distribution centered on the 210 CTCF motif. a Unfiltered reads +/- 1250 bp around a CTCF binding site located on the negative 211 212 strand (chr1: 30,779,763 - 30,779,781). The midpoint of the CTCF motif is marked with the symbol "<", representing that it is on the negative strand, and a pink line. **b** Plot (a) filtered to 213 observed ligations (equivalently, short fragments.) c Schematic demonstrating the bimodal read 214 215 pile-up around a CTCF binding site. d Plot (b) as a density plot and zoomed in on the CTCF motif, with quadrant annotations. e Distributions of reads in quadrants for true negative and true 216 positive CTCF binding sites in DNA loop anchors. True positives are defined as CTCF motifs 217 that are the only CTCF motif in a loop anchor and within 30 bp of a CTCF ChIP-seq peak. True 218 219 negatives are areas of the loop anchors with one CTCF motif that are at least 200 bp from the 220 CTCF motif. Schematics of the quadrant read pile-up patterns are shown next to the corresponding true positive and true negative boxplots. **f** FactorFinder statistic ( $\hat{\alpha} = \frac{\min(n_2, n_4)}{\max(n_1, n_2)}$ ) 221 222 for plot (d) peaks at the CTCF motif.

223

#### 224 <u>Model evaluation</u>

- 225 FactorFinder uses a biologically-informed model that takes advantage of the distribution of short
- 226 fragments around a CTCF binding site to pinpoint CTCF binding. Additionally, our use of a
- 227 multinomial framework for significance evaluation avoids placing any distributional assumptions

- on the reads within a quadrant. We then sought to benchmark our CTCF binding site
- identification performance using CTCF motif locations<sup>27</sup>, CTCF ChIP-seq peaks<sup>28</sup>, and loop
   anchors identified by FitHiChIP at 2.5kb resolution<sup>29</sup>.
- 231
- 232 We define a high stringency true positive set of CTCF binding sites as CTCF motifs in loop
- anchors that are located within 30 bp of a CTCF ChIP-seq peak summit. To avoid ambiguity due
- to multiple closely spaced motifs, we further selected only those motifs that are unique within a
- 235 2.5kb loop anchor. Using this true positive set, we observe that the *FactorFinder* statistic,

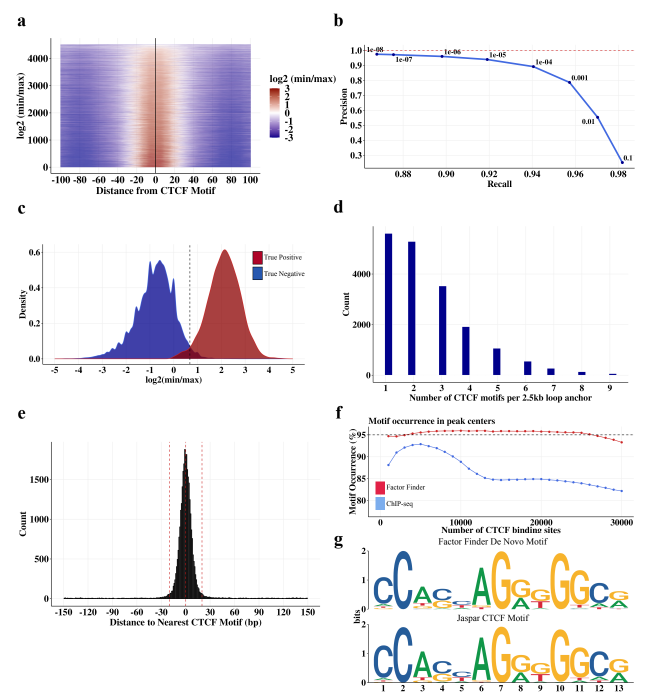
236  $log2(\hat{\alpha}) = log2(\frac{min(n_2, n_4)}{max(n_1, n_3)})$  has signal greater than 0 (equivalently,  $\hat{\alpha} > I$ ) almost exclusively

- within 20 bp of the CTCF motif center and centered on 0 bp from the CTCF motif center (Fig.
- 238 3a). Using this same set of true positive sites (false negatives are the regions of the loop anchors
- 239 >200 bp from a CTCF motif), we achieve > 90% precision and > 90% recall at a p-value
- threshold of 1e-05, and maintain high recall and precision at all p-value thresholds < 1e-05 (Fig.
- 241 3b). This high level of recall and precision is achieved because of the very different
- 242 *FactorFinder* statistic distributions for true positives and true negatives (Fig. 3c).
- 243

244 Because 70% of loop anchors defined with 2500 bp resolution contain multiple CTCF motifs

245 (Fig. 3d), higher levels of precision are often needed to determine the specific CTCF motif(s)

- 246 mediating a CTCF loop. Examining the effectiveness of *FactorFinder* genome-wide, we observe
- that almost all *FactorFinder* peak summits (93%) are within 20 bp of a CTCF motif center, with
- a median separation of 5 bp (Fig. 3e). Quantifying accuracy using motif occurrence within 20 bp
- 249 of a peak summit, we find that *FactorFinder* maintains ~95% motif occurrence while ChIP-seq
- declines to less than 85% motif occurrence (Fig. 3f). Applying the motif discovery tool
- 251 STREME<sup>30</sup> to 30 bp sequences centered on the *FactorFinder* peak summit produces a motif
- sequence that exactly matches the core JASPAR CTCF motif (Fig. 3g), further supporting
- 253 *FactorFinder*'s ability to identify true CTCF binding sites.



254

Fig. 3 CTCF binding sites identified by FactorFinder with single basepair resolution in MNase 255 K562 CTCF HiChIP data. a Heatmap of log2(min/max) as a function of distance between 256 FactorFinder peak center and CTCF motif center within loop anchors. Only CTCF motifs that 257 258 are unique within a loop anchor and within 30 bp of a CTCF ChIP-seq peak are used. **b** Precision 259 recall curve for true negative and true positive CTCF binding sites in DNA loop anchors. True positives are defined as in (a). True negatives are areas of the loop anchors in (a) that are at least 260 200 bp from the one CTCF motif. Precision is calculated as TP / (TP + FP), recall is calculated 261 as TP / (TP + FN). c FactorFinder statistic density plots using the same set of true positives and 262

- true negatives as (b). d Distribution of the number of CTCF motifs in a 2.5kb loop anchor. e
- Histogram with 1 bp bin size depicting *FactorFinder* resolution for all peaks genome-wide (not
- 265 just in loop anchors). f Motif occurrence in ChIP-seq and FactorFinder peak centers genome-
- wide. Motif occurrence is calculated as % peak centers within 20 bp of CTCF motif. Only peak
- centers within 150 bp of a CTCF motif are used for this figure. **g** 30 bp sequences centered on
- genome-wide *FactorFinder* peak centers produce a de novo motif (top) that matches the coreJASPAR CTCF motif (bottom).
- 270

## 271 <u>CTCF and Cohesin occupancy footprints</u>

We next examined the length characteristics of MNase HiChIP fragments overlapping individual CTCF motifs, to infer the presence and identity of the protein occupying the locus. For motifs with non-zero coverage, we observed long, 150+ bp fragments, as shown for three representative motifs in Figure 4a. These fragments likely represent cells with a nucleosome located at the motif locus, and are observed at CTCF motifs genome-wide (Fig. 1c). In addition, for a large subset of CTCF motifs, we also observed short, sub-nucleosome sized (<115 bp) fragments (Fig. 4a, Fig. 1c), likely instead representing DNA protected by CTCF.

279

A closer examination of the TF-scale fragments at *FactorFinder*-identified bound motifs reveals that they tend to exhibit a skew towards the downstream side of the CTCF motif (Fig. 4a, b, c), suggesting a preferred location for the protein(s) protecting the region from MNase cleavage. We considered cohesin as a potential candidate, given a recent finding that cohesin is stabilized on DNA through a specific interaction with the N terminus of the CTCF protein<sup>2</sup>, which localizes to the downstream side of the CTCF binding site.

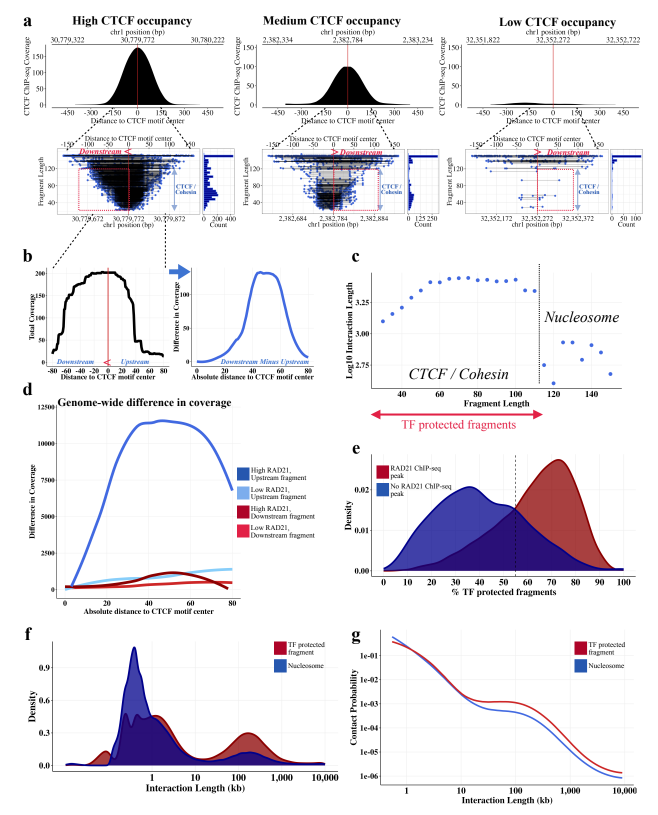
286

287 Given CTCF's role in mediating DNA looping we investigated whether the CTCF-adjacent protected footprint might relate to 3D architecture within the cell. We used HiChIP pairwise 288 289 interaction data where each ligation event reflects a single-cell point-to-point contact, to classify 290 each CTCF motif-overlapping fragment as either 'upstream" or 'downstream', depending on its 291 relationship to its interaction partner. Upstream fragments have long range contacts downstream 292 of the motif, and therefore have looping contacts in the same direction as a chromatin loop mediated by cohesin bound to the N terminus of the CTCF protein. Examining the difference in 293 294 coverage downstream and upstream of CBS genome-wide, we observe that upstream fragments

- 294 coverage downstream and upstream of CBS genome-wide, we observe that upstream mag
   295 overlapping CBS with an adjacent strong RAD21 ChIP-seq peak have substantially more
- adjacent coverage in the ~60 bp region downstream compared to upstream of the motif, while
   downstream fragments and CBS with weak adjacent RAD21 ChIP-seq peaks exhibit no
- difference (Fig. 4d). This finding further suggests that the CTCF-adjacent factor is associatedwith loop formation.
- 300

To further investigate whether the TF footprints identified at CTCF motifs might relate to anarchitectural role, we used HiChIP data to characterize their interaction patterns. We found that

- 303 TF-protected fragments (<115 bp) had contacts at substantially longer genomic distances than
- 304 nucleosome-protected fragments (Fig. 4c), suggesting that the TF presence may facilitate long
- 305 range interactions. Furthermore, we computed the frequency of TF-protected fragments at all
- 306 *FactorFinder*-identified CTCF bound sites, and found that it is strongly associated with the
- 307 presence of a RAD21 ChIP-Seq peak at the motif  $^{28}$  (Fig 4e).
- 308
- 309 Examination of the interaction length distribution shows that, as expected, the majority of
- 310 interactions occur within a linear separation of less than 10kb. The fraction of long-range
- 311 (>10kb) interactions, however, is significantly enriched (3.5-fold,  $p < 10^{-10}$ ) for short TF-
- 312 protected fragments as would be expected if these footprints represent CTCF/cohesin (Fig. 4f).
- 313 Similarly, an examination of the P(s) curve, showing contact probability as a function of linear
- distance, reveals a decreased attenuation in contact probability at longer interaction lengths (Fig.
  315 4g). Taken together, these findings suggest that we can classify CTCF HiChIP interaction data
- 316 based on footprint/fragment size as involving either unoccupied CTCF sites that tend to have
- 317 short-range chromatin interactions, or CTCF/cohesin occupied sites that, presumably through
- 318 loop extrusion, are able to make long-range contacts.





**Fig. 4** Cohesin and CTCF-protected fragments identified in CTCF MNase HiChIP. **a** High,

321 medium, and low CTCF occupied motifs. Cohesin footprint is observed downstream of the CBS

322 for high and medium CTCF occupancy motifs. For each occupancy level, CTCF ChIP-seq (top) 323 and all fragments overlapping the CTCF motif (bottom left) are depicted, along with the 324 corresponding fragment length histogram (bottom right). b Locus-specific high CTCF occupancy 325 figure from (a) as a coverage plot (left figure), difference in coverage between downstream and 326 upstream coverage (right figure). c Plotting median log10 interaction length as a function of 327 fragment length suggests presence of nucleosome vs TF-protected fragments. Only left fragments overlapping CTCF (+) motifs with start and end at least 15 bp from the CTCF motif 328 329 were included in this graph to remove confounding by MNase cut site. Using this figure, we are 330 approximating CTCF +/- cohesin-protected fragments as those with fragment length < 115, start and end at least 15 bp from the motif center. **d** Difference in coverage (downstream - upstream) 331 332 across all CBS shows an increase in coverage downstream of the CTCF motif for upstream 333 fragments underlying CBS with a strong adjacent RAD21 ChIP-seq peak. e CTCF motifs that 334 have a nearby RAD21 ChIP-seq peak (within 50 bp) have a larger proportion of TF-protected fragments. f TF-protected fragments have a noticeably larger bump in density of long range 335 336 interactions compared to nucleosome-protected fragments. Fragments were first filtered to those 337 with start and end at least 15 bp from the motif. TF-protected fragments were then defined as 338 fragments with length < 115 bp while nucleosome-protected fragments are fragments with length 339 at least 115 bp. g P(S) curve for fragments depicted in (f).

340

341 Active enhancers and gene transcription hinder cohesin-mediated loop extrusion

Using the techniques described above, MNase HiChIP enables us to simultaneously locate CBS
at high resolution, identify footprints of bound proteins, and interrogate specific chromatin
contacts at the single molecule level. We next sought to leverage these data to characterize
cohesin extrusion dynamics in a range of genomic contexts.

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We first estimated the frequency of fully extruded CTCF-CTCF chromatin loops genome-wide.
By obtaining fragments overlapping CTCF binding sites and estimating the fraction of
interaction partners overlapping a downstream convergent CTCF motif, we obtain 5% as the

350 genome-wide frequency of the fully extruded CTCF-CTCF state.. We also find a wide CBS to

- 351 CBS variability with an estimated range of  $\sim 1-10\%$  (Fig. 5a). This suggests that most CTCF-
- anchored chromatin contacts at the single-cell level are in the 'extruding' state, rather than joining two CTCF sites. These ranges are consistent with two recent locus-specific live cell imaging studies, which found that the fully extruded loop state is rare at the *Fbn2* TAD<sup>13</sup> and an
- engineered TAD on chr15<sup>14</sup>, occurring  $\sim$ 3-6%<sup>13</sup> and  $\sim$ 20-30% of the time<sup>14</sup> respectively. Note that the 20-30% estimate corresponds to a loop existing between any combination of three CBS
- 357 (+) and three CBS (-).
- 358

359 We next sought to use our data to examine how cohesin extrusion is impacted by chromatin

- 360 context. Since HiChIP libraries are a snapshot of millions of cells, we can estimate dynamic
- 361 extrusion parameters (primarily the average loop size extruded by cohesin<sup>31</sup>) from the interaction

- 362 length distribution. To determine the impact of chromatin state on cohesin extrusion, we first
- annotated the 1 MB regions downstream of *FactorFinder* identified CBS with ChromHMM
- 364 states<sup>32</sup> (Fig. 5b) to characterize the DNA through which a cohesin anchored at the CBS would
- 365 extrude through. Due to the highly correlated nature of ChromHMM annotations (Fig. 5c, d), we
- then divided the genome into three main chromatin state categories to uniquely classify each 1
- 367 MB region as either active, polycomb/bivalent or quiescent (Fig. 5d). CTCF/cohesin-protected
- 368 fragments overlapping CBS were accordingly annotated with the corresponding motif-level
- 369 chromatin state group, and extruded loop size estimates were obtained for each chromatin state370 based on the fragment-level interaction lengths.
- 371

372 Interestingly, we find that cohesin extrudes 1.75 times further through quiescent regions (252kb) 373 than through active regions (144kb), corresponding to a difference in average extruded loop size of ~110kb,  $p < 10^{-10}$  (Fig. 5e, Supp Fig. 3, Supp Fig. 4 right). The P(s) curve, a plot of interaction 374 375 decay with distance, confirms a depletion of the longest-range interactions in active regions (Fig 5f). This estimate for quiescent regions is consistent with a live cell imaging study of the Fbn2 376 locus in the absence of transcription that estimated a processivity of 300kb<sup>13</sup>. As quiescent 377 regions are characterized by low TF binding, low transcription, and minimal histone 378 modifications<sup>33</sup>, we hypothesized that the substantial difference in extruded loop size relates to 379 380 gene activity and enhancer density obstructing loop extrusion. Consistent with this, we found 381 that higher levels of H3K27ac and RNA Pol II binding in the 1MB region downstream of the 382 CBS strongly correlate with lower average extruded loop size (Fig. 5g).

383

384 We sought to establish that the observed differences in loop extrusion length as a function of chromatin state are not confounded by locus-specific effects on cohesin extrusion. Each CBS has 385 locus-specific genetic architecture and a different number of overlapping fragments, so we fit a 386 linear mixed effects model to account for this group-level heterogeneity. Specifically, we 387 388 compute the 'cohesin effect' on loop length, defined as the increase in average interaction length 389 for CTCF/cohesin bound fragments compared to nucleosome bound fragments for each CBS. 390 Controlling for the background interaction frequency of a region in this way confirms that 391 cohesin-associated loops are significantly shorter in active chromatin (Supp Fig. 4 left). Taken 392 together, these findings imply that gene and enhancer activity impede cohesin translocation (Fig. 393 5h).

394

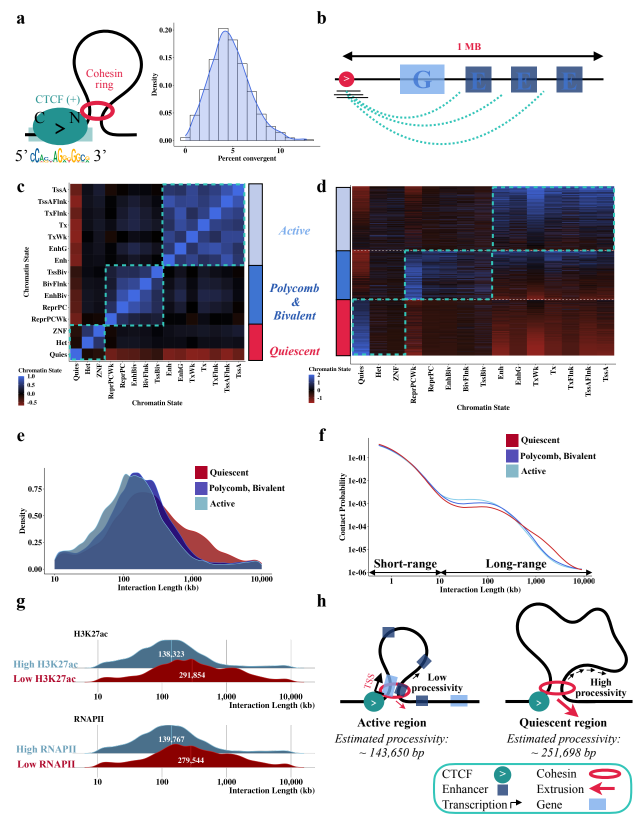




Fig. 5 Cohesin extrudes further through quiescent regions than active regions. a Most CTCF mediated looping contacts do not reflect the fully extruded state. Estimate is obtained using left

398 TF-protected (start and end at least 15 bp from motif center, length < 115) fragments that overlap 399 FactorFinder identified CBS (+) and have an interaction length greater than 10kb. For each CBS 400 with at least 50 long-range TF-protected fragments overlapping the motif, % convergent is 401 calculated as the number of interaction partners overlapping CTCF (-) motifs / total number of 402 fragments at motif. Because this estimate is conditional on CTCF binding at the anchor, we divide estimates by two to account for the  $\sim$ 50% occupancy of CTCF<sup>34</sup>. **b** Depiction of how 403 404 regions were annotated using ChromHMM. Correlation (c) and fragment (d) heatmaps for 405 ChromHMM annotated unique 1 MB regions downstream of left fragments overlapping CTCF (+) binding sites. All other plots in this figure are filtered to TF-protected (fragment length < 115 406 407 bp, start and end at least 15 bp from motif center) fragments. Density (e) and P(S) curves (f) for 408 chromatin state clusters shown in (c,d), filtered to the top 20%. Chromatin annotations making 409 up each cluster are added together and quantiles are obtained to determine fragments in the top 410 20% of active chromatin, quiescent chromatin, and bivalent / polycomb chromatin. g Ridge plots 411 for the bottom 10% quantile ("Low") and top 10% quantile ("High") of H3K27ac bp and number of RNAPII binding sites. ChIP-seq from ENCODE was used to annotate 1 MB downstream of 412 413 left fragments overlapping CBS (+) for this figure. h Diagram illustrating differences in 414 extrusion rates between active and quiescent chromatin states, with numbers obtained from Supp 415 Fig. 3.

416

#### 417 **Discussion**

418

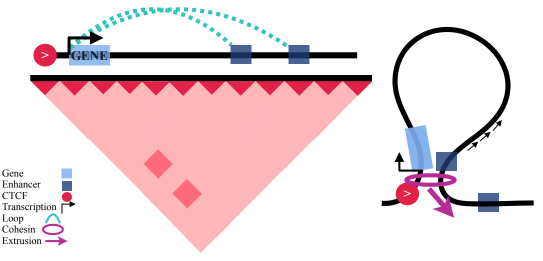
419 We have developed FactorFinder, a transcription factor footprinting method for MNase HiChIP 420 data and used it to identify CTCF binding sites with near base-pair resolution. We show that the 421 DNA protection footprints of nucleosomes and transcription factors can be readily distinguished 422 based on pre-ligation fragment size and strand origin and use these features to identify CTCF 423 binding sites. Significance is then assessed through a multinomial approach, which avoids 424 placing distributional assumptions on read counts. Using this method, the median distance 425 between FactorFinder peak summits and motif center is 5 bp, with 93% of peak summits 426 identified within 20 bp of a CTCF motif center.

427

428 We then leverage this methodological advance to investigate how chromatin state affects cohesin 429 extrusion dynamics. A close examination of CTCF-protected fragments revealed an additional CTCF-adjacent footprint downstream of the CBS, which we propose represents cohesin given its 430 431 positioning relative to looping orientation as well as its strong association with both long range 432 interactions and cohesin occupancy. We estimated the frequency with which a CTCF bound 433 locus forms a loop with a downstream CTCF site and found that it varies considerably from CBS to CBS, with a genome-wide range from ~1-10%. This is consistent with recent live-cell imaging 434 work that found that CTCF-mediated loops predominantly exist in the partially extruded state at 435

- 436 two studied loci<sup>13,14</sup>.
- 437

- 438 We next sought to characterize how cohesin impacts genome contacts in different chromatin
- 439 contexts. To this end, we employed our high-resolution *FactorFinder* identified CBS and
- 440 HiChIP 3D contact information to look at differences in extruded loop size in regions with
- different chromatin states. We observe an approximately 2-fold increase in extruded loop size
- 442 comparing quiescent chromatin to active chromatin, and this effect is similarly observed when
- examining the impact of H3K27ac and RNAPII binding. Our finding that RNAPII binding
- 444 obstructs cohesin-mediated loop extrusion is consistent with two recent studies that investigated
- 445 RNAPII's impact on cohesin through RNAPII and enhancer perturbations<sup>35</sup> as well as polymer
- 446 simulations, CTCF depletion, and Wapl knockout experiments<sup>36</sup>. These substantial differences in
- 447 average extruded loop size observed for different levels of RNAPII binding and H3K27ac
- suggest that gene and enhancer activity obstruct cohesin-mediated loop extrusion.
- 449
- 450 The obstruction of cohesin by gene and enhancer activity implies a model of CTCF-mediated
- 451 gene regulation where a fully extruded, stable, and convergent CTCF-CTCF loop is not required
- 452 for CTCF to mediate enhancer-promoter contacts. Instead, a promoter-proximal CTCF can halt
- 453 cohesin next to the TSS of a gene while cohesin continues to extrude on the other side,
- 454 effectively behaving as an enhancer recruiter. Cohesin slowing down through enhancer regions
- 455 would then enable an enrichment of enhancer-promoter contacts without requiring a stable
- 456 CTCF-CTCF loop (Fig. 6). This attenuation in cohesin extrusion may also provide a mechanism
- 457 relating gene regulation to the presence of RNAPII at enhancers<sup>37</sup>.



- 458
- 459 Fig. 6 Schematic of proposed model whereby single promoter-proximal CTCF sites enable an460 enrichment of enhancer-promoter contacts.
- 461
- 462 The dynamic CTCF-mediated enhancer-promoter contact model proposed here is consistent with
- 463 recent findings that promoter proximal CTCFs have important roles in gene regulation<sup>9-11</sup>, that
- 464 enhancer-promoter contacts are unstable<sup>38,39</sup>, and that CTCF and cohesin-mediated chromatin
- loops are dynamic<sup>13,14</sup>. The dynamic nature of EP contacts has contributed to the development of
- 466 the "kiss and kick" model<sup>40</sup> as a potential explanation for how enhancers and promoters come

- into contact but move away from each other at the time of transcription. Our findings are
- 468 compatible with the "kiss and kick" model, but additionally suggest a potential mechanism by
- 469 which distal enhancers can locate gene promoters without being stuck in a stable conformation.
- 470 This model would use promoter- or enhancer-proximal CTCF sites to enable distal enhancers to
- both come into contact with gene promoters and subsequently disengage during transcription. In
- this way, CTCF's role in long-range enhancer promoter contact would be as a dynamic
- 473 functional element recruiter instead of mediating continual stable contact between distal
- 474 enhancers and gene promoters.
- 475

## 476 Materials and methods

- 477 478 *CTCF MNase HiChIP*
- Four MNase K562 CTCF HiChIP (150 bp paired-end) libraries were generated using the Cantata
  Bio / Dovetail Genomics MNase HiChIP kit. CTCF MNase HiChIP was performed as described
  in the Dovetail HiChIP MNase Kit protocol v.2.0. Briefly, 5 million K562 cells per sample were
- 482 crosslinked with 3mM DSG and 1% formaldehyde and digested with 1ul MNase ("YET"
- 483 samples) or 2ul MNase ("GW" samples) in 100ul of 1X nuclease digestion buffer. Cells were
- 484 lysed with 1X RIPA containing 0.1% SDS, and CTCF ChIP was performed using 1500ng of
- chromatin (40-70% mononucleosomes) and 500 ng of CTCF antibody (Cell Signaling, cat #:
- 486 3418). Protein A/G beads pull-down, proximity ligation, and library preparation were done
- 487 according to the protocol. Libraries were sequenced to a read depth of  $\sim$ 172 million paired end
- reads per sample on the Illumina Nextseq 2000 platform.
- 489
- 490 <u>Software implementation</u>
- 491 Preprocessing, analysis and figure code used in this paper are available at
- 492 https://github.com/aryeelab/cohesin\_extrusion\_reproducibility. Data figures in this paper were
- 493 made in R v.4.1.2 using ggplot.
- 494
- 495 <u>Data availability</u>
- 496 Raw and Processed HiChIP data produced in this study will be uploaded to NCBI GEO (GSE
- 497 Record ID pending).
- 498 K562 ChIP-seq RAD21 BED file (Accession ID: ENCFF330SHG), CTCF BED file (Accession
- 499 ID: ENCFF736NYC), CTCF bigWig signal value (Accession ID: ENCFF168IFW), RNAPII
- 500 BED file (Accession ID: ENCFF355MNE), and H3K27ac BED file (Accession ID:
- 501 ENCFF544LXB) were obtained from ENCODE, and CTCF motifs were obtained from the R
- 502 package *CTCF*<sup>27</sup> (annotation record: AH104729, documentation:
- $\label{eq:solution} 503 \qquad https://bioconductor.org/packages/release/data/annotation/vignettes/CTCF/inst/doc/CTCF.html).$
- 504
- 505 Methods

- 506 Data Processing
- 507 4 replicates of K562 MNase CTCF HiChIP data were aligned to the reference genome using the
- 508 BWA-MEM algorithm<sup>41</sup>. Ligation events were then recorded using pairtools parse v.  $0.3.0^{42}$ ,
- 509 PCR duplicates were removed, and the final pairs and bam files were generated. HiChIP loop
- calls were then made using FitHiChIP Peak to Peak<sup>29</sup> with 2.5kb loop anchor bin size. The
- 511 MNase HiChIP processing protocol is based on guidelines from
- 512 <u>https://hichip.readthedocs.io/en/latest/before\_you\_begin.html</u>. Reproducible code is available at
- 513 https://github.com/aryeelab/cohesin\_extrusion\_reproducibility.
- 514

## 515 Identification of significant motifs

- 516 We use CTCF motifs identified as significant (p < 1e-05) by *FactorFinder* as the set of CTCF
- 517 binding sites. This p-value threshold was chosen based on the precision recall curve (Fig. 3b),
- and corresponds to a maximum FDR q-value of 3e-04.
- 519

## 520 <u>Multiple Testing</u>

- 521 For genome-wide footprinting analysis adjustment for multiple testing, CTCF motifs are
- 522 assigned the p-value of the closest *FactorFinder* sliding window. The Benjamini-Hochberg
- 523 method<sup>43</sup> was used to obtain q-values.
- 524

## 525 <u>Estimating cohesin footprints</u>

- 526 The cohesin footprint is observed by obtaining motif-level coverage estimates +/- 80 bp around
- 527 CBS, summing up the coverage across all motifs (within strata), and subtracting the upstream
- 528 coverage from the downstream (downstream coverage upstream coverage) at each base pair.
- 529 Note that downstream and upstream are defined relative to the motif strand, so downstream is to
- 530 the "left" of CBS (-) and to the "right" of CBS (+) in terms of reference genome base pairs. The
- aforementioned strata are defined by RAD21 ChIP-seq signal level (high vs low) and whether
- the fragment is the upstream or downstream interaction partner in its pair. RAD21 ChIP-seq high
- and low correspond to the top 25% and bottom 25% of ChIP-seq signal value of the adjacent
- 534 (within 50 bp of CBS) RAD21 ChIP-seq peak. Note that only mid-size (fragment length between
- 80 and 120), long range fragments (interaction length > 10kb) are used for this analysis.
- 536

## 537 <u>Estimating the fully extruded state</u>

- 538 We estimated a genome-wide range for the fully extruded state by obtaining CTCF/cohesin-
- 539 protected upstream fragments overlapping CBS (+) and estimating the fraction of interaction
- 540 partners overlapping a downstream convergent negative strand CTCF motif. CBS (+) were
- 541 required to have at least 50 CTCF/cohesin-protected upstream fragments overlapping the motif
- to enable sufficient sample size for the motif-specific percent convergent calculation. We then
- 543 accounted for CTCF occupancy (estimated as  $\sim 50\%$ )<sup>34</sup> by dividing this estimate by two. The
- 544 point estimate (5%) is the number of interaction partners overlapping a downstream convergent

negative strand CTCF motif genome-wide / the total number of fragments genome-wide, and the

- range (1-10%) are the 1st and 99th percentile of the CBS-level CTCF-CTCF chromatin loop
- 547 estimate.
- 548
- 549 Determining extruded loop size as a function of chromatin state
- 550 We used upstream fragments overlapping CTCF binding sites (+) for this analysis. 1 MB regions
- downstream of the CBS (+) were annotated using ChromHMM<sup>32</sup> to quantify the percentage of bp
- assigned to each of the 15 chromatin states. To simplify annotation, we grouped the 15
- chromatin states into three categories (quiescent, polycomb/bivalent, and active) based on their
   correlation (Fig 5c). Regions were clustered using Ward's hierarchical clustering method<sup>44</sup> (Fig
- 555 5d.). For extrusion dynamics analyses (Fig 5e,f,h), each of the three chromatin categories was
- represented by the 20% of regions with the highest fraction of DNA in this state. Extruded loop
- size was then estimated as the average log10 interaction length for each annotation. Only long
- range TF-protected fragments (start and end at least 15 bp from the motif center, length < 115,
- 559 interaction length > 10kb) were included in this estimate.
- 560

Similarly, high/low H3K27ac corresponds to the top 10% and bottom 10% of the number of
basepairs covered by H3K27ac ChIP-seq peaks in the 1 MB regions downstream of CBS (+).
High/low RNAPII corresponds to the top 10% and bottom 10% of the number of RNAPII ChIP-

- seq peaks located in the 1 MB regions downstream of CBS (+). Extruded loop size estimates
- were obtained in the same way for these annotated regions; long range TF-protected fragments
- were used to estimate the average log10 interaction length.
- 567
- 568 Directionality of CBS-adjacent nucleosome position signal
- 569 Interestingly, the strength of the nucleosome positioning signal is related to the orientation of the 570 DNA contact. Stratifying nucleosome-bound fragments based on whether they are the upstream
- 571 or downstream long-range (>10kb) fragment in a pair (effectively single-cell left or right loop
- 572 anchor) produces a differential nucleosome signal inside and outside the loop (Supp Fig. 5). For
- both upstream and downstream nucleosome-bound fragments, the nucleosome closest to the
- 574 CTCF binding site and inside the loop exhibits a substantially stronger signal than the closest
- nucleosome outside the loop. HiChIP ligations are unlikely to fully account for this signal as a
- 576 previous study using MNase-seq also showed a directional nucleosome preference around CBS
- 577 (see Fig. 1a), although this result was not noted in the text $^{25}$ .
- 578

## 579 **Disclosures**

- 580
- 581 Dovetail Genomics/Cantata Bio provided reagents and sample processing for HiChIP
- 582 experiments. M.B. and M.S.B were employees at Dovetail Genomics during the course of this
- research. M.J.A has financial and consulting interests unrelated to this work in SeQure Dx and
- 584 Chroma Medicine. M.J.A's interests are reviewed and managed by Dana Farber Cancer Institute.

585 J.K.J. is a co-founder of and has a financial interest in SeQure, Dx, Inc., a company developing

- technologies for gene editing target profiling. JKJ also has, or had during the course of this
- 587 research, financial interests in several companies developing gene editing technology: Beam
- 588 Therapeutics, Blink Therapeutics, Chroma Medicine, Editas Medicine, EpiLogic Therapeutics,
- 589 Excelsior Genomics, Hera Biolabs, Monitor Biotechnologies, Nvelop Therapeutics (f/k/a ETx,
- Inc.), Pairwise Plants, Poseida Therapeutics, and Verve Therapeutics. J.K.J.'s interests were
   reviewed and are managed by Massachusetts General Hospital and Mass General Brigham in
- reviewed and are managed by Massachusetts General Hospital and Mass Generaaccordance with their conflict of interest policies.
- 593

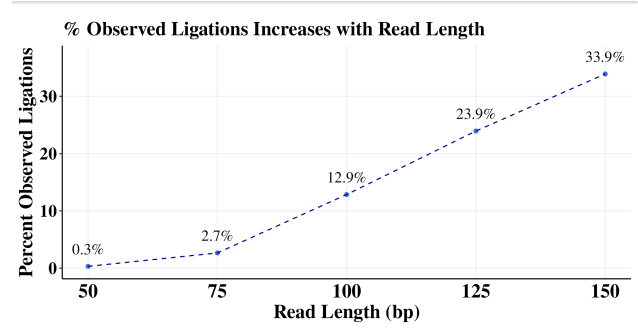
595

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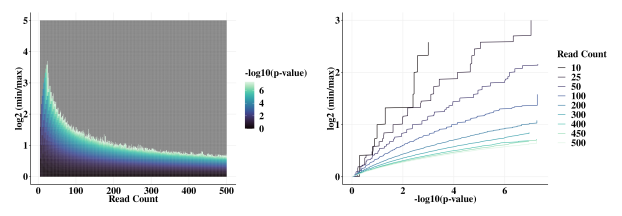
- 598 American Society of Gene & Cell Therapy (YET). The content is solely the responsibility of the
- authors and does not necessarily represent the official views of the American Society of Gene &
- 600 Cell Therapy. Dovetail Genomics / Cantata Bio supported data generation costs.
- 601



## 602 Supplementary Figures

603

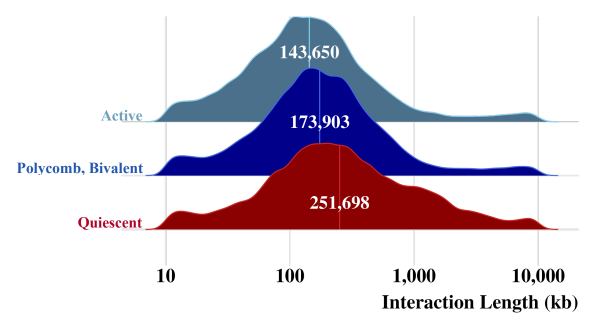
604
605 Supplementary Figure 1. Percent observed ligations increases with read length.





607 Supplementary Figure 2. The probability of observing a high *FactorFinder* statistic under the608 null hypothesis is higher at low read counts.

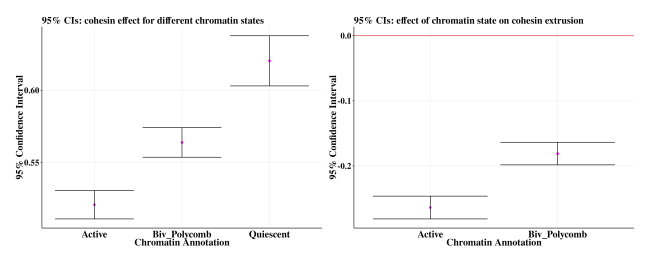




610

611 Supplementary Figure 3. Cohesin extrudes significantly further through quiescent regions than

- 612 active regions.
- 613

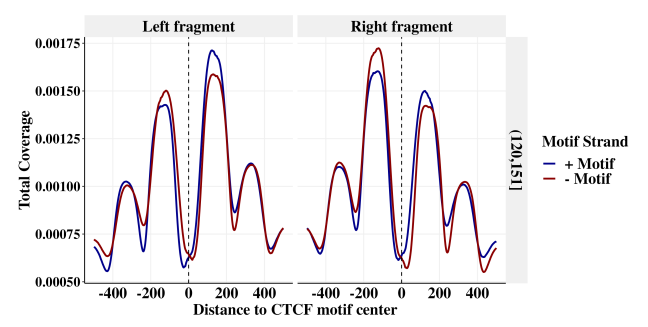




615 Supplementary Figure 4. Controlling for locus-specific variation with linear mixed models

616 does not attenuate the relationship between chromatin state and extruded loop size. Note that for

- 617 the figure on the right, the group that active and bivalent polycomb are being compared to is618 quiescent.
- 619



- 620
- 621 Supplementary Figure 5. Nucleosomes are preferentially positioned inside the loop.
- 622

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