Chromatin jets define the properties of cohesin-driven in vivo loop extrusion

Graphical abstract

Highlights

- Cohesin-powered chromatin jets emerge from focal areas of accessible chromatin
- Unless constrained by CTCF, jets can propagate symmetrically for 1–2 Mb
- CTCF can block jets or deflect the angle of jet propagation
- These data suggest that loop extrusion is controlled independently in both directions

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In brief

Cohesin organizes the genome in 3D nuclear space. In this issue of Molecular Cell, Guo et al. describe and characterize “jets,” an unusual form of cohesin-mediated chromatin interactions. Jets offer insights into the physiological behavior of cohesin-mediated loop extrusion and the principles that underlie genome organization.
Chromatin jets define the properties of cohesin-driven \textit{in vivo} loop extrusion

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SUMMARY

Complex genomes show intricate organization in three-dimensional (3D) nuclear space. Current models posit that cohesin extrudes loops to form self-interacting domains delimited by the DNA binding protein CTCF. Here, we describe and quantitatively characterize cohesin-propelled, jet-like chromatin contacts as landmarks of loop extrusion in quiescent mammalian lymphocytes. Experimental observations and polymer simulations indicate that narrow origins of loop extrusion favor jet formation. Unless constrained by CTCF, jets propagate symmetrically for 1–2 Mb, providing an estimate for the range of \textit{in vivo} loop extrusion. Asymmetric CTCF binding deflects the angle of jet propagation as experimental evidence that cohesin-mediated loop extrusion can switch from bi- to unidirectional and is controlled independently in both directions. These data offer new insights into the physiological behavior of \textit{in vivo} cohesin-mediated loop extrusion and further our understanding of the principles that underlie genome organization.

INTRODUCTION

The interphase genome is organized by the separation of chromatin states into active (A) and inactive (B) compartments and the formation of TADs, contact domains, and CTCF-based chromatin loops by cohesin and CTCF (Rao et al., 2014; Nora et al., 2017; Schwarzwar et al., 2017; Dekker and Mirny, 2016; Merkenschlager and Nora, 2016; Mirny et al., 2019). Chromatin loops form preferentially between CTCF molecules bound to DNA motifs in convergent orientation, suggesting that loop formation is based on one-dimensional (1D) tracking rather than three-dimensional (3D) diffusion (Rao et al., 2014; Vietri-Rudan et al., 2015; de Wit et al., 2015) and consistent with models of loop extrusion mediated by the structural maintenance of chromosomes (SMCs) protein complexes (Riggs et al., 1990; Kimura et al., 1999; Nasmyth, 2001; Alipour and Marko, 2012; Sanborn et al., 2015; Fedenberg et al., 2016, 2017; Banigan et al., 2022). SMC complexes translocate on DNA and can extrude DNA loops at rates of approximately 1 kb s$^{-1}$ (Gruber and Errington, 2009; Ganji et al., 2018; Golfier et al., 2020; Kong et al., 2020; Terakawa et al., 2017; Tran et al., 2017; Wang et al., 2017; Wang et al., 2018; Anchimiuk et al., 2021; Davidson et al., 2019; Kim et al., 2019). Structural studies provide evidence for a conserved cohesin interface with the N terminus of CTCF (Li et al., 2020; Nora et al., 2020; Pugacheva et al., 2020). It is thought that a process akin to loop extrusion underlies the formation of contact domains and CTCF-based loops: (1) loop domains reform rapidly after transient cohesin depletion, providing a minimal estimate of the speed of cohesin-mediated chromatin contact propagation (Rao et al., 2017). (2) Hi-C “stripes” connect CTCF domain
anchors with multiple points within domains, highly suggestive of loop extrusion (Vian et al., 2018). (3) Loops become longer when the residence time of cohesin on chromatin is increased by the removal of the cohesin unloading factor WAPL (Haahrus et al., 2017; Wutz et al., 2017; Liu et al., 2021a).

During DNA replication in bacteria, SMC complexes are loaded from predefined sites, providing evidence that individual SMC complexes can align entire chromosome arms comprising several million base pairs of DNA (Marbouty et al., 2015; Tran et al., 2017; Wang et al., 2017, 2018; reviewed by Banigan and Mirny, 2020). In contrast to bacteria, loading sites for SMC complexes are not well defined in mammalian cells. Active regulatory elements are enriched for cohesin and the cohesin component and loading factor NIPBL. ATAC-seq accessibility and the acetylation of histone H3 on lysine 27 (H3K27ac) are part of the chromatin signature of active regulatory elements and therefore considered as chromatin features of potential cohesin loading sites (Vian et al., 2018). Biochemical, functional, and genetic interactions of NIPBL/cohesin with H3K27ac-associated BET proteins (Olley et al., 2018; Luna-Peláez et al., 2019; Linares-Saldana et al., 2021), the enhancer-associated MLL3/4 complex (Yan et al., 2018), and the components of the transcriptional machinery (Kagey et al., 2010; Liu et al., 2021b; van den Berg et al., 2017) implicate enhancers and super-enhancers as candidate loading sites for cohesin within chromatin domains and TADs (Vian et al., 2018; Kagey et al., 2010). However, in the absence of direct assays for cohesin loading in living cells, the identity of in vivo cohesin loading sites remains uncertain. Moreover, chromatin regions that give rise to domains, loops, and stripes in mammalian genomes typically contain multiple potential cohesin loading sites. Therefore, it is unclear whether cohesin-dependent contacts are established by the continuous action of individual cohesin complexes or by a succession of multiple cohesin complexes.

Here, we describe cohesin-dependent chromatin contacts that originate from small, isolated sites of accessible chromatin and traverse adjacent B compartment regions in a jet-like fashion in primary mammalian cells. Jet origins feature local chromatin accessibility, NIPBL, cohesin, and H3K27ac. Our experimental and computational modeling suggest that lone accessible sites favor focal cohesin loading and chromatin jet formation, whereas broad or multiple accessible sites drive diffuse cohesin loading and favor the formation of contact domains (Fudenberg et al., 2016; Fudenberg et al., 2017; Banigan et al., 2022). Once initiated, jets can propagate symmetrically for 1–2 Mb. Unilateral CTCF encounters can convert bi- to unidirectional extrusion, indicating that both directions of extrusion are controlled independently. These data provide new insights into the physiological behavior of in vivo cohesin-mediated loop extrusion in unperturbed mammalian cells.

RESULTS

To elucidate the properties of cohesin-mediated chromatin contact propagation in vivo, we performed in situ Hi-C on wild-type, Ctf<sup>-/-</sup>, and Rad21<sup>-/-</sup> Ctf<sup>-/-</sup> primary DP thymocytes (Figure S1; Table S1). The efficiency of CTCF depletion was 80.0% ± 10.1% (n = 7), and the efficiency of RAD21 depletion was 83.3% ± 4.2% (n = 7; Figure S1C). We chose small DP thymocytes because they are quiescent and therefore require cohesin and CTCF for 3D genome organization in interphase but not for cell cycle-related functions such as DNA replication or chromosome segregation (Seitan et al., 2011; Nasmyth and Haering, 2009). We examined the resulting Hi-C maps for features consistent with loop extrusion. In addition to the familiar Hi-C patterns of ubiquitous pleat-like compartments and diamond-shaped domains (see Figure 1A for a schematic and illustrative example, Figure S1 for quantification), visual inspection revealed a smaller number of jet-like projections from the Hi-C diagonal (see Figure 1B for a schematic and illustrative example, Table S2 for quantification), reminiscent of Hi-C patterns that are formed by aligned chromosome arms in bacteria (Tran et al., 2017; Wang et al., 2017, 2018; reviewed by Banigan and Mirny, 2020). The examination of published Hi-C data (Kieffer-Kwon et al., 2017) shows that similar jets are present in B cells (Figure S2).

The majority of jets originated from small regions denoted as open chromatin, i.e., A compartment intervals based on Hi-C eigenvector. Jets projected into surrounding closed chromatin regions (B compartment intervals, Figure 1C; Table S2). We identified a set of n = 38 jets and developed a protractor tool to quantify jet strength and jet angles in individual Hi-C replicates (see method details section; Figures 1D and 1E; Table S2). Protractor scanning confirmed increased chromatin contacts around small A compartment intervals that contained visually identified jets compared with small A compartment intervals that lacked jets, including regions that contained contact domains, or neither jets nor contact domains (Figure 1F; Table S2). The propagation of chromatin jets perpendicular to the Hi-C diagonal is consistent with bidirectional loop extrusion in vivo (Figure 1G).

Jets can be constrained by CTCF and released by CTCF removal

In comparison with wild-type thymocytes (Figure 1), additional jets arise in Ctf<sup>-/-</sup> thymocytes (Figures 2A, 2B, and S3; Table S2). Aggregate subtraction plots confirmed that average jet strength increased in the absence of CTCF (Figures 2C and S4). Protractor scan quantification showed that a subset of jets became significantly stronger in CTCF-deficient (Ctf<sup>-/-</sup>) compared with wild-type DP thymocytes. Integration with CTCF chromatin immuno-precipitation sequencing (ChIP-seq) data showed that this increase in strength was confined to those jets that were targets of CTCF binding in wild-type DP thymocytes (Figure 2D). The loss of CTCF led to a significant decrease in the variance of jet length distributions (Figure 2E). This is consistent with the role of CTCF in blocking cohesin-mediated loop extrusion (Li et al., 2020) and indicates that CTCF is a major determinant of jet propagation. Polymer simulations in which CTCF stalls cohesin reproduced the experimentally observed impact of CTCF on the propagation of jets in the presence and the absence of CTCF (Figure 2F; see method details section and Table S3 for parameters). These data indicate that jet propagation can be constrained by CTCF and released by CTCF removal (Figure 2G).

Jets are cohesin dependent

To ask whether jet formation depends on cohesin, we compared Hi-C maps of wild-type, CTCF-deficient (Ctf<sup>-/-</sup>), and cohesin-deficient (Rad21<sup>-/-</sup> Ctf<sup>-/-</sup>) DP thymocytes.
We observed that jets were substantially weakened by the depletion of cohesin (Figures 1A and 2A; Table S2). The jet remnants visible in Hi-C maps of cohesin-deficient thymocytes are likely due to the imperfect depletion of the RAD21 protein (Figure S1C). The aggregate analysis and subtraction plots confirmed the weakening of jets in the absence of cohesin (Figure 3A). The protractor scan quantification of jet strength of wild-type, CTCF-deficient (Ctcf−/−), and cohesin-deficient (Rad21−/− Ctf2−/−) DP thymocytes showed that reduced jet strength after cohesin depletion was significant, whether or not CTCF was bound at jet origins (Figure 3B). The analysis of published dilution Hi-C data from wild-type and Rad21−/− DP thymocytes (Seitan et al., 2013) confirmed that depletion of cohesin alone was sufficient to weaken jet formation (Figure S5).

Jet origins have chromatin features consistent with cohesin loading sites

Compared with A compartment intervals genome wide, the sub-set of A compartment-like intervals that give rise to jets was...
distinctly smaller in size (Figure 4A). Among A compartment intervals <1 Mb, jets preferentially originated from smaller A compartment intervals compared with contact domains (Figure 4B). To identify chromatin features of A compartment intervals that form jets, we integrated Hi-C with ATAC-seq, which identifies accessible chromatin, ChIP-seq for the cohesin subunit RAD21 and NIPBL, as well as H3K27ac as a mark for active regulatory elements. H3K27ac-associated BET proteins (Olley et al., 2018; Luna-Peláez et al., 2019; Linares-Saldana et al., 2021), the enhancer-associated MLL3/4 complex (Yan et al., 2018), and components of the transcriptional machinery (Kagey et al., 2010; Liu et al., 2021b) form biochemical, functional, and genetic interactions of NIPBL and cohesin (Olley et al., 2018; Luna-Peláez et al., 2019; Linares-Saldana et al., 2021; Yan...
of the transcriptional machinery (Kagey et al., 2010; van den Berg et al., 2019; Linares-Saldana et al., 2021), and jet origins have chromatin features consistent with these studies.

Isolated focal areas of open chromatin favor jet formation

Jets showed a more focused distribution of ATAC-seq and H3K27ac signal than contact domains (Figure 5A; quantification in Figure 5B), indicating that jets are more likely to arise from isolated focal areas of open chromatin. We used polymer simulations to model the impact of narrow versus broad cohesin loading areas on the resulting chromatin contacts. In these simulations, 100 monomers represented narrow cohesin loading 2,000 monomers represented broad areas of cohesin loading. All other parameters such as total polymer length, number of cohesins, drift, and unloading probability were kept constant (see simulations 1 and 2 in Table S3). Narrow cohesin loading resulted in jet-like features (Figure 5C, left), whereas broad loading gave rise to contact domains instead (Figure 5C, middle; Fudenberg et al., 2016, 2017). Interestingly, additional loading along the length of the jet disrupted jet formation (Figure 5C, right).

Taken together, the dependence of jets on cohesin and their origin from isolated areas of open chromatin marked by H3K27ac, RAD21, and NIPBL, the preference of jets for narrow defined origins, are powered by cohesin, and are constrained by CTCF provides a unique opportunity to determine the properties of in vivo loop extrusion. We developed a stencil tool to quantify the range of jet propagation (see method details section, Figure 6A). We found that jets can propagate for ~1–2 Mb in wild-type and CTCF-deficient DP thymocytes (Figure 6B; Table S2) and for similar distances in resting B cells (Figure S2).

Jet angles are modulated by CTCF, providing experimental evidence for one-sided loop extrusion

We next analyzed the impact of CTCF on the angle of jet propagation. Jets that propagate perpendicular to the diagonal in a cohesin-dependent manner illustrate that in vivo loop extrusion. We developed a stencil tool to quantify the range of jet propagation (see method details section, Figure 6A). We found that a subset of jets deviates from the perpendicular (Figures 7A and S6; Table S2). The jet depicted in Figure 7A has a negative projection angle, indicating that this jet is deflected toward upstream sequences in wild-type cells. This jet is flanked by downstream CTCF sites. The jet depicted in Figure S6A has a positive projection angle in wild-type cells, indicating that it is deflected toward downstream sequences in wild-type cells. This jet is flanked by upstream CTCF sites.

In contrast to the jets shown in previous figures, these angled jets are not symmetrical in wild-type cells. Accordingly, the distance traveled by loop extrusion is greater in one direction (Figure 7C; Table S2). Interestingly, the projection angles of these deflected jets change in CTCF-deficient cells, and jets align more closely with the perpendicular (Figures 7A, 7B, 7A, and 7B; Table S2). Consequently, the range of jet angles is broader in wild-type than in CTCF-deficient cells (Figure 7D). We next counted the number of CTCF motifs with CTCF ChIP-seq signals than contact domains (Figure 5D).

Figure 3. Jets are cohesin dependent

(A) Mean Hi-C contacts for n = 38 jets in wild-type and CTCF KO (as in Figure 2C) with Rad21–/– Ctcf–/– DP thymocytes (DKO, right). The difference in mean Hi-C signal CTCF KO minus wild type (right), x and y axes are in Mb.

(B) Numerical comparison of jet strength in wild-type and Ctcf–/– (as in Figure 3D) with Rad21–/– Ctcf–/– (DKO) DP thymocytes n = 38. Jets with at least one CTCF ChIP-seq peak within 200 kb of the jet origin (CTCF bound, n = 19) and jets without a CTCF ChIP-seq peak within 200 kb of the jet origin (not CTCF bound, n = 18) were affected by the loss of cohesin (one-sided t test). The mean (red) and 95% confidence interval are shown (gray).
seq peaks in DP thymocytes as CTCF motif scores, where positive scores indicate CTCF downstream binding and negative motif scores with CTCF upstream binding. Positive CTCF motif scores were associated with jet deflection toward upstream sequences and negative CTCF motif scores were associated with jet deflection toward downstream sequences (Figure 7E).

The observation that CTCF can alter the projection angle of jets is informative with respect to the behavior of the loop extrusion machinery upon encounter with CTCF. The loop extrusion model posits that extrusion is blocked by CTCF bound to sites in convergent orientation (Fudenberg et al., 2016), and recent studies provide a mechanism for how the encounter of cohesin with the N terminus of CTCF arrests loop extrusion and stabilizes cohesin at CTCF sites (Li et al., 2020; Nora et al., 2020; Puga-cheva et al., 2020). If loop extrusion encounters CTCF in only one direction, extrusion may either stop completely or continue

Figure 4. Jet origins have chromatin features consistent with cohesin loading sites
(A) Length distribution of all A compartment intervals called genome wide versus A compartment intervals that give rise to jets. Based on n = 38 jets and n = 1,267 A compartment intervals.
(B) Length distribution of A compartment intervals <1 Mb that contain contact domains, jets, or neither.
(C) Strength of ATAC-seq (Miyazaki et al., 2020) and ChIP-seq signal for A compartment intervals <1 Mb that contain contact domains, jets, or neither.
(D) Quantification of ATAC-seq and ChIP-seq signals shown in (C).
in the direction that is not blocked by CTCF (Fudenberg et al., 2017; Figure 7F). Our observation that jets can be deflected by an encounter with CTCF indicates that loop extrusion can switch from bidirectional to unidirectional in a CTCF-dependent manner. Using polymer simulations to model how unilateral CTCF encounter affects jet angles, we find that models where a unilateral encounter with CTCF imposes a bidirectional block on cohesin-mediated loop extrusion do not reproduce the experimental observation of altered jet angles (Figure 7G). We next modeled CTCF effects on jet angles under the assumption that unilateral encounter with CTCF imposes a unidirectional block on cohesin-mediated loop extrusion (Figure 7H). The results of this simulation reproduce the experimental observations that (1) jet angles change in response to unilateral CTCF encounter and that (2) both arms of the jets show different lengths in response to unilateral CTCF encounter (Figures 7C and 7F; Table S2). These findings provide experimental evidence for previous assumptions that when blocked in one direction, loop extrusion continues in the other direction (Fudenberg et al., 2017). Taken together, our experimental data and in silico simulations support a model where cohesin-driven loop extrusion in vivo is by default symmetrical and therefore bidirectional. However, CTCF can significantly deflect the projection angle of cohesin-driven chromatin contacts, indicating that extrusion can switch from bidirectional to unidirectional.

Taken together, these data provide a mechanistic dissection of in vivo loop extrusion. A model for jet formation is depicted in Figure S7, including symmetrical extrusion in the absence of CTCF (Figure S7A) the impact of bi- (Figure S7B) and unidirectional (Figure S7C) CTCF encounters and propagation from narrow versus broad origins (Figures S7A and S7D).

**DISCUSSION**

Here, we describe and quantitatively characterize cohesin-dependent chromatin jets and provide insights into the physiological behavior of in vivo cohesin-mediated loop extrusion.

Loading sites for SMC complexes are well defined in bacteria (Gruber and Errington, 2009; Ganji et al., 2018; Golfier et al., 2020; Kong et al., 2020; Terakawa et al., 2017; Tran et al., 2017; Wang et al., 2017; Wang et al., 2018; Anchimiuk et al., 2021). In mammalian cells, active regulatory elements are enriched for cohesin and NIPBL, ATAC-seq accessibility and H3K27ac, and considered as chromatin features of potential cohesin loading sites (Vian et al., 2018; Olley et al., 2018; Luna-Pérez et al., 2019; Linares-Saldana et al., 2021; Yan et al., 2018; Kagey et al., 2010; Liu et al., 2021b; van den Berg et al., 2017). In mammalian cells, active regulatory elements are enriched for cohesin and NIPBL, ATAC-seq accessibility and H3K27ac, and considered as chromatin features of potential cohesin loading sites (Vian et al., 2018; Olley et al., 2018; Luna-Pérez et al., 2019; Linares-Saldana et al., 2021; Yan et al., 2018; Kagey et al., 2010; Liu et al., 2021b; van den Berg et al., 2017). However, the identity of in vivo cohesin loading sites has remained uncertain. Jets originate from isolated areas of accessible chromatin enriched for H3K27ac, NIPBL, and RAD21. Jets that project directly
from their origins and propagate perpendicular to the Hi-C diagonal to provide powerful evidence for the location of cohesin loading. Computational modeling supports the interpretation that cohesin is loaded at jet origins. In these simulations, narrow sites of cohesin loading give rise to jets. Previous simulations suggested that focal cohesin loading at promoters might generate jet-like features (Banigan et al., 2022), although these were not observed experimentally. In contrast to narrow loading sites, broad areas of cohesin loading lead to domain formation (Fudenberg et al., 2016, 2017). This interpretation is further strengthened by experimental observations that the likelihood of jet formation is inversely related to the size of the potential cohesin loading area. Jets echo the SMC-driven process that aligns bacterial chromosome arms, both in terms of Hi-C pattern and the requirement for defined loading sites (Gruber and Errington, 2009; Ganji et al., 2018; Marbouty et al., 2015; Golffier et al., 2020; Kong et al., 2020; Terakawa et al., 2017; Tran et al., 2017; Wang et al., 2017, 2018; Anchimiuk et al., 2021). Akin to the alignment of bacterial chromosome arms, jets may reflect the continuous, linear activity of stacked SMC complexes (Figure S7C). Indeed, our simulations indicate that the “re-loading” of additional cohesin complexes along the length of the jet would disrupt jet formation (Figure S7D).

Recent reports describe interesting chromatin features that are similar to jets in appearance and likely related in terms of mechanism. In yeast, pericentromeric jet-like features form when centromeres are engaged by mitotic spindles (Paldi et al., 2020), and the jet-like alignment of chromatin fragments are associated with double-strand break repair (Piazza et al., 2021; Arnould et al., 2021). These features are characterized by their short range of ~25 kb. Chromatin flares have been found to form in the specialized chromatin environment of zebrafish sperm (Wike et al., 2021). Flares show evidence of linear alignment of DNA sequences, but with an average length of ~175 kb, they are on a smaller scale than jets. Flare formation has not been linked to specific SMC complexes (Wike et al., 2021). A recent preprint observes the transient occurrence of cohesin-dependent jet-like chromatin features in response to the co-depletion of WAPL and CTCF in mouse ES cells, referred to as plumes (Liu et al., 2021b). As described here for jets, these structures originate from small accessible regions surrounded by B compartments (Liu et al., 2021b). However, because plumes form only after the depletion of the cohesin unloading factor WAPL, they cannot serve to establish the physiological range of cohesin-mediated loop extrusion.

Although WAPL depletion was required, it was not sufficient for plume formation unless CTCF was also depleted (Liu et al., 2021b), which masks unidirectional in vivo loop extrusion upon a one-sided CTCF encounter.

Similar to flares and plumes, jets arise from focal origins. However, unlike flares (Wike et al., 2021), jets form in somatic cells, rather than in the specialized chromatin environment of sperm, and are unequivocally cohesin dependent. Unlike plumes (Liu et al., 2021b), jets form in unperturbed primary cells at steady state and therefore reflect in vivo loop extrusion under conditions of physiological cohesin residence time. Therefore, jets allow measurements of the genomic distances traversed by cohesin complexes loaded at specific sites, and jet length may reflect the range of individual cohesin complexes on chromatin in vivo. From their origins at isolated areas of accessible chromatin, jets propagate across neighboring closed (B compartment) chromatin to cover distances of approximately 1–2 Mb.

Our findings are consistent with the assumption that cohesin-mediated loop extrusion is symmetrical in the absence of CTCF and therefore bidirectional by default (Banigan et al., 2020). We find that unilateral CTCF encounter deflects the angle of jet propagation. This provides strong evidence that cohesin-mediated loop extrusion can switch from bi- to unidirectional in vivo (Fudenberg et al., 2017; Vian et al., 2018) and therefore is independently controlled in both directions. Jets provide new insights into the physiological behavior of in vivo cohesin-mediated loop extrusion and further our understanding of the principles that underlie genome organization.

**Limitations of the study**

Although our data show that jets are cohesin dependent and can be blocked or deflected by CTCF, we do not yet fully understand the rules that govern jet formation and, in particular, why not all focal sites of accessibility give rise to jets. We do not know the residence time of cohesin in quiescent primary lymphocytes and, as a result, can only speculate on cohesin extrusion speeds during jet propagation. Estimates of cohesin residence times in interphase range between 10 and 25 min in other cell types (Gerlich et al., 2006; Tedeschi et al., 2013; Hansen et al., 2017; Wutz et al., 2020). Assuming that cohesin residence times are similar in quiescent primary lymphocytes, our data suggest that cohesin can traverse “closed” B compartment chromatin at speeds that are comparable with—and possibly in excess of—SMC complex extrusion speed in bacteria and cohesin-mediated extrusion in vitro (Ganji et al., 2018; Golffier et al., 2020; Kong et al., 2020; Terakawa et al., 2017; Tran et al., 2017; Wang et al., 2017, 2018; Davidson et al., 2019; Kim et al., 2019). Our current model for the formation of jets is that individual cohesin complexes mediate continuous
Figure 7. CTCF can modulate jet angles
(A) A jet that is deflected by CTCF (WT) and released by the removal of CTCF (CTCF KO).
(B) Replicate-based protractor scan quantification of the jet shown in (A).
(C) Cohesin travels the same distance in both directions for jets that are perpendicular to the diagonal (d1 = d2, blue). Cohesin travels further in one direction than in the other for jets that are not perpendicular to the diagonal (d1 ≠ d2, black).
(D) Summary of jet angles in the presence and absence of CTCF. 0° describes the perpendicular to the Hi-C diagonal. P indicates the significance of the change in variance (F test).
(E) Experimentally observed jet angles reflect CTCF binding. CTCF motif scores represent the number of CTCF ChIP-seq peaks with convergent motifs downstream within 150 kb of the jet origin minus the number of CTCF ChIP-seq peaks with convergent motifs upstream within 150 kb of the jet origin. Downstream > upstream, n = 8 jets; downstream < upstream, n = 10 jets. High CTCF motif scores indicate CTCF binding downstream and are associated with jet deflection toward upstream sequences and vice versa.
(F) Schematic. Unilateral CTCF encounter (encounter with CTCF in one direction) may inactivate cohesin-mediated loop extrusion completely (top) or convert bidirectional (two-sided) extrusion into unidirectional (one-sided) extrusion (bottom).
(G) Polymer simulation of how CTCF binding (200 monomers up- and/or downstream of the center of the loading area) affects jet angles under the assumption that unilateral encounter with CTCF imposes a bidirectional block on cohesin-mediated loop extrusion.
(H) Polymer simulation of how CTCF binding (200 monomers up- and/or downstream of the center of the loading area) affects jet angles under the assumption that unilateral encounter with CTCF imposes a unidirectional block on cohesin-mediated loop extrusion. CMs, chromatin monomers (see method details section).
cohesin-mediated loop extrusion is independently controlled in both directions but does not address whether cohesin complexes work as dimers (Kim et al., 2019), monomers (Yatskevich et al., 2019; Davidson et al., 2019), or multimers (Xiang and Koshland, 2021).

**STAR★METHODS**

Detailed methods are provided in the online version of this paper and include the following:

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.molcel.2022.09.003.

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**DECLARATION OF INTERESTS**

The authors declare no competing interests.

**INCLUSION AND DIVERSITY**

We support inclusive, diverse, and equitable conduct of research.

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**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE                          | SOURCE                                      | IDENTIFIER        |
|----------------------------------------------|---------------------------------------------|-------------------|
| Antibodies                                   |                                             |                   |
| Rabbit polyclonal anti-CTCF                  | Abcam                                       | Cat# ab70303; RRID:AB_1209546 |
| Rabbit polyclonal anti-H3K27ac               | Active Motif                                | Cat# 39133; RRID:AB_2561016 |
| Rabbit polyclonal anti-Rad21                 | Abcam                                       | Cat# ab992; RRID:AB_2176601 |
| Rabbit polyclonal anti-SMC3                  | Abcam                                       | Cat# ab9263; RRID:AB_307122 |
| Rabbit polyclonal anti-Histone H3            | Abcam                                       | Cat# ab1791; RRID:AB_302613 |
| Rat monoclonal anti-CD4                      | Biolegend                                   | Cat# 100514; RRID:AB_312717 |
| Rat monoclonal anti-CD8a                     | Thermo Fisher Scientific                    | Cat# 12-0081-85; RRID:AB_465532 |
| Arm.Hamster monoclonal anti-CD69             | BD Biosciences                              | Cat# 553236; RRID:AB_394725 |
| Chemicals, peptides, and recombinant proteins|                                             |                   |
| 16% Formaldehyde, Methanol-free              | Pierce                                      | Cat# 28908        |
| Deposited data                               |                                             |                   |
| Hi-C, ChIP-seq data (CTCF, Rad21, H3K27ac)   | This Study                                  | GEO:GSE199059     |
| ChIP-seq data (Nipbl)                        | Seitan et al., 2013                         | GEO:GSE48763      |
| Experimental models: Organisms/strains       |                                             |                   |
| Mouse: Rad21tm1.1Mmk                          | Our colony                                  | MGI:5293824       |
| Mouse: Ctcf conditional allele               | Our colony                                  | https://doi.org/10.1038/nature10312 |
| Mouse: Tg(Cd4-cre)1Cwi                       | Jax B6.Cg-Tg(Cd4-cre)1Cwi                   | MGI:2386448       |
| Critical commercial assays                   |                                             |                   |
| NEBNext ChIP-seq Library Prep Master Mix Set for Illumina | NEB                                           | E6240             |
| NEBNext Multiplex Oligos for Illumina        | NEB                                         | E7335, E7500      |
| AMPure XP                                   | Beckman Coulter                             | A63881            |
| Software and algorithms                      |                                             |                   |
| Bowtie2 (v2.3.2)                              | Langmead and Salzberg, 2012                 | http://bowtie-bio.sourceforge.net/bowtie2/index.shtml |
| Hi-C-Pro (v2.7.8)                            | Servant et al., 2015                        | https://github.com/nservant/Hi-C-Pro |
| Juicer (v0.7.5)                               | Durand et al., 2016                         | https://github.com/aidenlab/juicer |
| Juicebox                                     | Durand et al., 2016                         | https://github.com/aidenlab/Juicebox |
| R                                            | The R Foundation                            | https://www.r-project.org/ |
| Samtools (v1.4)                               | Li et al., 2009                              | http://samtools.sourceforge.net/ |
| Stencil tool                                 | This paper                                  | https://github.com/garciamillan/DNA_jets |
| Protractor tool                              | This paper                                  | https://github.com/garciamillan/DNA_jets |
| Polychrom                                    | Imakaev, 2019                               | Version v0.1.0, 10.5281/zenodo.3579473 |
| bowtie (v1.0.0)                               | Langmead, 2010                              | http://bowtie-bio.sourceforge.net/index.shtml |
| MACS2 (v2.1.0)                                | Zhang et al., 2008                          | https://github.com/macs3-project/MACS/wiki/Advanced%3A-Call-peaks-using-MACS2-subcommands |
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to the lead contact, Matthias Merkenschlager (matthias.merkenschlager@lms.mrc.ac.uk).

Materials availability
This study did not generate new unique reagents.

Data and code availability
In situ Hi-C data for Hi-C CD69+ DP wild-type, CD4 Cre Ctcf<sup>−/−</sup> (CTCF KO), and CD4Cre Rad21<sup>−/−</sup> Ctcf<sup>−/−</sup> (DKO) thymocytes and ChIP-seq data for RAD21, CTCF, and H3K27ac in wild-type DP thymocytes generated in this study have been deposited at the NCBI Gene Expression Omnibus (GEO) under accession number GSE199059 and are publicly available as of the date of publication. ATAC-seq data in wild-type DP thymocytes are publicly available at GEO GSE141223 (Miyazaki et al., 2020). Dilution Hi-C for wild-type and CD4 Cre Rad21<sup>−/−</sup> DP thymocytes and ChIP-seq data for NIPBL in wild-type DP thymocytes are publicly available at GEO GSE48763 (Seitan et al., 2013). Mouse B cell Hi-C data are publicly available at GEO GSE82144 (Kieffer-Kwon et al., 2017).

All original code has been deposited at Zenodo and is publicly available as of the date of publication. The DOI for code for jet simulations is https://doi.org/10.5281/zenodo.7028262. The DOI for code for jet quantification is https://doi.org/10.5281/zenodo.7034657.

Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals
Mice harbouring a conditional Ctcf allele (Ctcf<sup>lox/lox</sup>, Heath et al., 2008) were crossed with CD4Cre (Lee et al., 2001, RRID:IMSR_EM:01139). Mice harbouring a conditional Rad21 allele (Rad21<sup>lox/lox</sup>, Tachibana-Konwalski et al., 2010; Seitan et al., 2011, RRID:MGI:5293828) were crossed with CD4Cre (Lee et al., 2001, RRID:IMSR_EM:01139). Mice were maintained under specific pathogen free conditions, 12 hour light/dark cycle, food and water ad libitum, under the care of Imperial College London animal care staff and veterinary services. Males and females of the following genotypes were used between the ages of 6 and 12 weeks: Ctcf<sup>wt/wt</sup>, Rad21<sup>wt/wt</sup> CD4Cre (controls), Ctcf<sup>lox/lox</sup>, Rad21<sup>wt/wt</sup> CD4Cre (CTCF ko), Ctcf<sup>lox/lox</sup>, Rad21<sup>lox/lox</sup> CD4Cre (double ko). Experimental groups were defined by genotype. Mouse work was performed according to the Animals (Scientific Procedures) Act Mouse work was done under a project licence issued by the UK Home Office, UK following review by the Imperial College London Animal Welfare and Ethical Review Body (AWERB).

METHOD DETAILS

Mice and cells
Mouse strains harbouring conditional alleles for Ctcf (Ctcf<sup>lox/lox</sup>, Heath et al., 2008) and Rad21 (Rad21<sup>lox/lox</sup>, Tachibana-Konwalski et al., 2010; Seitan et al., 2011, RRID:MGI:5293828) were crossed with CD4Cre (Lee et al., 2001, RRID:IMSR_EM:01139) for lineage- and developmental stage-specific deletion at the transition to the CD4+ CD8+ (DP) stage of T cell development in the thymus (Lee et al., 2001) as described (Seitan et al., 2011). Quiescent CD4<sup>+</sup> CD8<sup>+</sup> CD69<sup>−</sup> DP thymocytes were isolated by flow cytometry (Figure S1B).

Hi-C and analysis
In situ Hi-C was performed as described (Rao et al., 2014) with the following modifications. Cells were cross-linked in 1% formaldehyde for 10 min at room temperature. Chromatin was digested with 100 units of Mbol overnight at 37 °C while shaking at 900 rpm. After overnight incubation, 100 units of Mbol were added for the second-round digestion for 2-4 hours. The DNA polymerase I, large (Klenow) fragment, was used to fill in Mbol-digested fragment overhangs in the presence of biotin-labelled dATP. Fragment ends were then ligated with 4000 units of T4 DNA ligase by incubating at room temperature for 4-6 h with rotation at 12 rpm. After ligation, cell nuclei were collected by centrifugation, and chromatin was reverse crosslinked overnight at 65 °C. RNase A was used to remove RNA after decrosslinking. Genomic DNA was isolated and precipitated by sodium acetate-ethanol precipitation, and then purified DNA was sheared for 8 min using a sonicator (Bioruptor) with high power setting, and 30 seconds on and 30 seconds off per minute. DNA fragments in the range of 300–500 bp were then selected using the AMPure XP beads (Beckmann Coulter). After biotin-labelled DNA fragments were captured on Dynabeads MyOne Streptavidin T1 (Thermo Fisher), fragment ends were repaired in a mixture of enzymes containing T4 polynucleotide kinase, T4 DNA polymerase I and DNA polymerase I, large (Klenow) fragment, and then NEBNext adaptors for Illumina sequencing were ligated to the da-tailed fragment ends. After the USER enzyme digestion, NEBNext oligos for Illumina sequencing were used for PCR for library preparation. A PCR titration was carried out to determine the lowest number of PCR cycles (8-10 cycles in this work). Final Hi-C sequencing libraries were quantified and checked for size...
distribution using Bioanalyzer (Agilent Technologies), and then sequenced on an HiSeq 2500 sequencer (Illumina). Hi-C raw
sequencing data (2 × 100 bp paired-end reads) were mapped using the bowtie 2 (version 2.3.2) and then processed by the HiC-
Pro (version 2.7.8) with default settings (Servant et al., 2015; Langmead and Salzberg, 2012). The reference genome used for map-
ping in this study was the UCSC assembly mm9, NCBI build 37. After converting using the script, HiC-Pro hicpro2juicebox.sh, valid
chromatin contacts were normalized and hic files were created by the Juicer Pre (Juicer tools version 0.7.5) (Durand et al., 2016).
Compartment were called by eigenvector decomposition, performed using the cooltools eigdecmod module (https://github.
com/open2c/cooltools) at 25kb resolution. A and B compartments were assigned by GC-content. Hi-C contact domains were called
using Arrowhead with default parameters at 5kb and 10kb resolution as described (Rao et al., 2014). For cases of proximal 5kb and
10kb domains (defined as <10kb Euclidean distance) the 10kb domains were removed. To reduce redundancy in downstream anal-
ysis nested domains were then pruned such that only internal and non-nested domains were retained.
Subcompartments were determined at 100kb resolution based on methods described (Rao et al., 2014) with the following alter-
ations: intra-chromosomal rather than inter-chromosomal contacts were used for classification, k-means clustering was used to
assign subcompartments instead of a hidden Markov mode, a 5Mb mask was applied around the diagonal of each Hi-C map to re-
move the impact of local interactions, and Hi-C matrices were interpolated to reduce noise. Hi-C data were visualised in Juicebox
(Robinson et al., 2018).

**ChiP-seq**

Cells were cross-linked in 1% formaldehyde solution for 10 minutes at room temperature and then the reaction was quenched
with 125 mM glycine at room temperature for 5 minutes. Cell nuclei were isolated using a RIPA lysis buffer (10mM Tris pH7.5, 1mM EDTA,
1% Triton X-100, 0.1% Sodium deoxycholate, 0.1% SDS, 0.15M NaCl, 1X Roche protease inhibitors). Following isolation of cell
nuclei, genomic DNA was sonicated in a high-salt RIPA buffer (10mM Tris pH7.5, 1mM EDTA, 1% Triton X-100, 0.1% Sodium de-
oxycholate, 1% SDS, 0.8M NaCl, 1X Roche protease inhibitors) for 30 minutes on a sonicator (Bioruptor) with high power setting,
and 30 seconds on and 30 seconds off per minute. Immunoprecipitation was performed in a RIPA buffer (10mM Tris pH7.5, 1mM
EDTA, 1% Triton X-100, 0.1% Sodium deoxycholate, 0.1% SDS, 0.15M NaCl, 1X Roche protease inhibitors), and antibody-antigen
complex was precipitated using Dynabead Protein G (Thermo Fisher). Protein was digested using Proteinase K, and then DNA-pro-
tein complex was reverse crosslinked overnight at 65°C. RNase A was added to remove RNA after decrosslinking. ChiP-seq library
preparation were carried out using the NEBNext ChiP-Seq Library Prep Master Mix Set for Illumina (NEB) according to the manufac-
turer’s protocol. Final sequencing libraries were sequenced on an HiSeq 2500 sequencer (Illumina). ChiP-seq raw reads (50 bp
single-end) were mapped using the bowtie (version 1.0.0) (Langmead, 2010). After alignment and duplicates removal, peaks were
identified using MACS2 (Feng et al., 2012).

**Centrality score**

The centrality score is the sum of ATAC-seq or ChiP-seq signal at the 50kb centred on the feature divided by the sum of the signal
from the 50-75kb upstream and downstream of the feature.

**In silico modelling**

We use simulations to investigate the effect of cohesin loop extrusion and CTCF binding on Hi-C maps. The simulations work in two
stages. One-dimensional movement of cohesin subunits across chromatin is followed by three-dimensional polymer folding simu-
lations to produce simulated Hi-C maps. To model chromatin folding we use the publicly available software polychrom (Version
ev0.1.0, 10.5281/zenodo.3579473, Imakaev et al., 2019) and manually add pairs of cohesin cuffs as extra bonds that define the struc-
ture, as described in the following.

Cohesin is modelled as a pair of two random walkers that move along a straight line representing chromatin, modelled as a chain of
monomers. Each monomer is a possible position for a walker. We initialise the simulation by placing with uniform probability both
walkers of a fixed number of cohesin molecules within a certain loading area. We may further place CTCF molecules along the poly-
mer. After this initial setup, the one dimensional simulation follows the process described in Algorithm 1 (Table S4). All of the actions in
the algorithm have predefined probabilities. This is explicitly shown for the unloading action, and for the rest it is abbreviated. We
model the movement of the two cuffs of each cohesin as a random walk with drift. The left cuff has a drift towards the left and the
other towards the right. The cuffs cannot switch places or bypass each other and may therefore stall others. They can further be
stopped and captured by CTCF molecules.

Two CTCF capture mechanisms and their impact on the formation of jets are investigated. In the first case, a suitably oriented
CTCF captures only the cuff that it encounters while the other cuff of the same cohesin is free to move and further extrude chromatin.
In the second case, when a CTCF captures one cuff of a cohesin, both cuffs are considered immobilized and cohesin stops extruding.
The drift is the difference between the jump probabilities of the cuffs. Cohesin is unloaded with different probabilities. Cohesin
captured by CTCF has a longer residence time on the polymer, which is reflected by the smaller unloading probability compared
to free cohesin. We allow for 0, 1 or 2 CTCF sites, at the positions indicated. At each of these sites, we place 5 CTCFs equidistantly,
every 50 monomers. Each CTCF may capture a cohesin cuff once it steps on the site with the probability stated and each may release
them again with the much smaller release probability indicated.
We chose a 5000-monomers-long polymer for our simulations. The Hi-C maps are based on typically about twenty polymer stochastic systems with the same settings. Once the one dimensional simulation has run, the trajectories of all of the cohesin cuffs, are input as extra bonds in a three dimensional polychrom simulation, from which finally the average contacts are calculated, and the contact map is generated. The three dimensional simulations are run on Nvidia RTX6000 GPUs and use a variable Langevin integrator. Model parameters are listed in Table S3.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Quantification of jets: The Protractor tool quantifies jet strength and jet angles, and the Stencil tool quantifies jet reach. Both use observed over expected Hi-C data with KR normalisation. We denote observed data at \((x,y) \in [0,L] \) by \(c_{xy}\) and expected data by \(e_{xy}\). Jet candidates are defined by start and end coordinates \((a_1, a_2)\) along a chromosome (equivalently \((a_2, a_1)\)). To determine jet strength and orientation we define the protractor as the set of points on the Hi-C map enclosed by a semi-circle, typically of a 2Mb radius, whose origin is placed at \((a_1, a_2)\). The straight side of the semi-circle is parallel to the Hi-C diagonal (Figure 1B). We divide the area of the protractor into \(n\) equal sectors (typically \(n = 80\)), labelled with a number \(i\). The observed Hi-C map is given by:

\[
\alpha_i = \frac{\sum_{(x,y) \in \Delta} (w(x) + w(y) - w(x)w(y))}{\text{expected data}}
\]

where \(w(x)\) is an indicator function of a stripe of missing data at \(x \in [0,L]\), defined as:

\[
w(x) = \begin{cases} 1 & \text{if } c_{xy} = 0 \text{ for all } y \in [0,L], \\ 0 & \text{otherwise}, \end{cases}
\]

so that \((w(x) + w(y) - w(x)w(y)) = 1 - (1 - w(x))(1 - w(y))\) in Equation 1 is unity whenever \(w(x) = 1\) or \(w(y) = 1\). Since the Hi-C map satisfies the symmetry \(c_{xy} = c_{yx}\), it is equivalent to identify stripes of missing data in the \(x\) coordinate or in the \(y\) coordinate, which is why \(w(x)\) in Equation 2 is defined by probing for missing data along \(y\) only. We define \(f_i\) as the intensity of observed relative to expected interactions within each sector \(i\):

\[
f_i = \left\{ \begin{array}{ll} N \sum_{(x,y) \in \Delta} (c_{xy} - e_{xy}) & \text{if } W_i < \tau, \\ \text{not defined} & \text{otherwise.} \end{array} \right.
\]

where \(N = 10^9/T\) and \(T\) is the total number of counts, and the tolerance is \(\tau = 0.1Q/n\), where \(Q\) is the total number of points in the protractor. We estimate the mean and standard deviation of \(f_i\) based on Hi-C replicates (Figure 1B). The values \(f_i\) may be subject to fluctuations inherent to the Hi-C data, the radius of the protractor, and the total number of sectors \(n\). To obtain robust estimates, we apply a Gaussian kernel to \(f_i\):

\[
\bar{f} = \frac{1}{B \sqrt{2\pi}} \int_{-\infty}^{x} e^{-\left(\frac{t^2}{2}\right)} dt
\]

where we set the bandwidth \(B = 1\). We determine the index \(M\) among those indices \(i\) such that \(\theta_i \in [-45^\circ, 45^\circ]\), where \(\theta\) reaches its maximum,

\[
\bar{f}_M = \max \left\{ f_i : i = \left\lceil \frac{n - 1}{2} \right\rceil, \ldots, \left\lfloor \frac{3n}{4} - \frac{1}{2} \right\rfloor \right\}
\]

which immediately gives the jet orientation \(\theta_M\). A nonzero orientation is indicative of asymmetric extrusion. We define the jet strength as

\[
S = 10^{-2} \bar{f}_M
\]

To measure the reach of jets that quantifies interactions between neighbouring regions of \((a_1, a_2)\), we devised the stencil, which is a rectangle 3.5Mb in length and 0.1Mb in width. The stencil is placed on the Hi-C map so that the mid-point of one of the short sides is at \((a_1, a_2)\) and its long sides are aligned with the jet orientation given by \(\theta_M\). We include in the stencil only contacts between locations outside the interval \([a_1, a_2]\). We divide the stencil along the direction \(\theta_M\) into \(m = 70\) equally sized pieces (Figure 6A). Similarly to Equations 1, 3, and 4, we define the functions

\[
W_i = \sum_{(x,y) \in \Delta} (w(x) + w(y) - w(x)w(y))
\]
\[ g_i = \begin{cases} 
\mathcal{N} \sum_{x \in \Delta_i} (c_{xy} - e_{xy}) & \text{if } W_i < \tau, \\
\text{not defined} & \text{otherwise}, 
\end{cases} \]  
(Equation 8)

\[ g_i = \sum_{j=0}^{m-1} g_j \frac{1}{B\sqrt{2\pi}} e^{-\frac{1}{2}(\frac{x - m}{B})^2}, \]  
(Equation 9)

where \( \Delta_i \) is the set of points in the Hi-C map that are contained in each piece \( i \) along the stencil, \( i \in \{0, \ldots, m-1\} \). We subtract the background given by \( \text{Rad21}^-\text{Ctcf}^- \) from the stencil \( g_i \) and find the piece \( i \) where the subtracted stencil falls below a threshold of 5. We then calculate the up- and downstream distances \( d_1 \) and \( d_2 \) (Figure 7C).

All statistical analysis and software used is listed in the Software and algorithms section of the STAR Methods table, and all statistical details of experiments can be found in the figure legends, figures, results, and method details sections, including the statistical tests used, the number of replicate experiments, definition of center, dispersion, and precision measures, and the definition of significance. Experimental and control groups were defined by genotypes, no randomization was performed, and no data were excluded from the analysis.