

## Transcription-independent TFIIIC-bound sites cluster near heterochromatin boundaries within lamina-associated domains in *C. elegans*

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### KEYWORDS

20 Transcription Factor III C (TFIIIC), RNA Polymerase III, *C. elegans*, Insulator, Chromatin, Chromosome, lamina-associated domain, nuclear periphery, LEM-2, Histone H3 Lysine 9 methylation

## ABSTRACT

**BACKGROUND:** Chromatin organization is central to precise control of gene expression. In vertebrates, the insulator protein CTCF plays a central role in organizing chromatin into topologically associated domains (TADs). In nematode *C. elegans*, however, a CTCF homolog is absent, and pervasive TAD structures are limited to the dosage-compensated sex chromosome, leaving the principle of *C. elegans* chromatin organization unclear. Transcription Factor III C (TFIIIC) is a basal transcription factor complex for RNA Polymerase III (Pol III), also implicated in chromatin organization. TFIIIC binding without Pol III co-occupancy, referred to as extra-TFIIIC binding, has been implicated in insulating active and inactive chromatin domains in yeasts, flies, and mammalian cells. Whether extra-TFIIIC sites are present and contribute to chromatin organization in *C. elegans* remain unknown.

**RESULTS:** We identified, in *C. elegans* embryos, 504 TFIIIC-bound sites absent of Pol III and TATA-binding protein co-occupancy characteristic of extra-TFIIIC sites. Extra-TFIIIC sites constituted half of all identified TFIIIC binding sites in the genome. Unlike Pol III-associated TFIIIC sites predominantly localized in the sex chromosome, extra-TFIIIC sites were highly over-represented within autosomes. Extra-TFIIIC sites formed dense clusters in *cis*. The autosomal regions enriched for extra-TFIIIC site clusters presented a high level of heterochromatin-associated histone H3K9 trimethylation (H3K9me3). Furthermore, extra-TFIIIC site clusters were embedded in the lamina-associated domains. Despite the heterochromatin environment of extra-TFIIIC sites, the individual clusters of extra-TFIIIC sites were devoid of and resided near the individual H3K9me3-marked regions.

**CONCLUSION:** Clusters of extra-TFIIIC sites were pervasive near the outer boundaries of H3K9me3-marked regions in *C. elegans*. Given the reported activity of extra-TFIIIC sites in heterochromatin insulation in yeasts, our observation raised the possibility that TFIIIC may also demarcate heterochromatin in *C. elegans*.

## 45 BACKGROUND

Eukaryotic genomes are organized into domains of various chromatin features including actively transcribed regions, transcription factor-bound regions, and transcriptionally repressed regions [1–4]. Demarcation of chromatin domains is central to precise control and memory of gene expression patterns. Several proteins have been proposed to have activity in demarcating chromatin domains by acting as a physical boundary [5,6], generating nucleosome depleted regions [7], mediating long-range chromatin interactions [8,9], or tethering chromatin to the nuclear periphery [10]. Despite intense studies [11–15], how chromatin domains are demarcated remains poorly understood.

The genome of nematode *Caenorhabditis elegans* has served as a model to study chromatin organization [3,16–18]. The highest level of chromatin organization in *C. elegans* is the chromatin feature that distinguishes between the X chromosome and the autosomes. The X chromosome in *C. elegans* hermaphrodites is organized into topologically associated domains (TADs) [19], the architectural unit mediated by long-range chromatin interactions and commonly seen in metazoan genomes [20–22]. The five autosomes, however, lack robust TAD organization [19]. Instead, each autosome can be subdivided into three, megabase-wide domains, the left arm, the right arm, and the center [23]. The center domains display a low recombination rate [24,25], a high density of essential genes [26], and low heterochromatin-associated histone modifications [16,27]. Autosome arms are rich in repetitive elements [23] and heterochromatin-associated histone modifications [16,27], and are associated with the nuclear membrane [18,28,29]. Within these generally euchromatic centers and heterochromatic arms lie kilobase-wide regions of various chromatin states including transcriptionally active and inactive regions [3,4]. While condensins, a highly conserved class of architectural proteins [30], define half of TAD boundaries in the X chromosome [19], their contribution to autosomal chromatin organization is unclear [14,31]. Furthermore, CTCF, another conserved architectural protein central to chromosome organizations in vertebrates, is thought to be lost during the *C. elegans* evolution [32]. How chromatin domains and chromatin states in the *C. elegans* autosomes are demarcated remains an area of active investigation [4,28,33,34].

The transcription factor IIIC complex (TFIIIC) is a general transcription factor required for recruitment of the RNA Polymerase III (Pol III) machinery to diverse classes of small non-coding RNA genes [35]. TFIIIC has also been implicated in chromatin insulation [36,37]. TFIIIC binds DNA sequence elements called the Box-A and Box-B motifs [35]. When participating in Pol III-dependent transcription, TFIIIC binding to Box-A and Box-B motifs results in recruitment of transcription factor IIIB complex (TFIIIB), which then recruits Pol III [35,38]. By yet unknown mechanisms, however, TFIIIC is also known to bind DNA without further recruitment of TBP and Pol III [37]. These so-called “extra-TFIIIC sites,” or “ETC,” have been identified in various organisms including yeast [39,40], fly [41], mouse [42], and human [43]. In *Saccharomyces cerevisiae* and *Schizosaccharomyces pombe*, extra-TFIIIC sites exhibit chromatin boundary functions both as heterochromatin barriers and insulators to gene activation [40,44,45]. In addition, extra-TFIIIC sites in these yeast species have been observed at the nuclear periphery, suggesting a contribution to spatial organization of chromosomes [40,46]. In fly, mouse, and human genomes, extra-TFIIIC sites were found in close proximity to architectural proteins including CTCF, condensin, and cohesin [41–43,47]. These studies collectively suggest a conserved role for extra-TFIIIC sites in chromatin insulation and chromosome organization. However, whether extra-TFIIIC sites exist in the *C. elegans* genome is unknown.

In this study, we unveiled extra-TFIIIC sites in the *C. elegans* genome. Extra-TFIIIC sites were highly over-represented within a subset of autosome arms that presented a high level of heterochromatin-associated histone H3K9 trimethylation. Extra-TFIIIC sites formed dense clusters *in cis* and were embedded in the lamina-associated domains. Despite the heterochromatin environment of extra-TFIIIC sites however, the individual clusters of extra-TFIIIC sites were devoid of and resided near the boundaries of H3K9me3-marked regions. Our study thus raised the possibility that, like extra-TFIIIC sites in other organisms, *C. elegans* extra-TFIIIC sites may also have an activity to demarcate chromatin domains.

## RESULTS

### Half of *C. elegans* TFIIIC binding sites lack Pol III co-occupancy

The TFIIIC complex is a general transcription factor required for the assembly of the RNA Polymerase III (Pol III) machinery at small non-coding RNA genes such as tRNA genes (**Fig. 1A**). Extra-TFIIIC sites are TFIIIC-bound sites lacking Pol III co-occupancy and are implicated in insulating genomic domains and spatially organizing chromosomes [48]. To determine whether the *C. elegans* genome includes extra-TFIIIC sites, we analyzed the ChIP-seq data published in our previous study [49] for TFIIIC subunits TFTC-3 (human TFIIIC63/GTF3C3 ortholog) and TFTC-5 (human TFIIIC102/GTF3C5 ortholog) (**Fig. 1B**), the Pol III catalytic subunit RPC-1 (human POLR3A ortholog; “Pol III” hereafter), and the TFIIIB component TBP-1 (human TBP ortholog; “TBP” hereafter) in mixed-stage embryos of the wild-type N2 strain *C. elegans*. We identified 1,029 high-confidence TFIIIC-bound sites exhibiting strong and consistent enrichment for both TFTC-3 and TFTC-5 (**Fig. 1C**). tRNA genes were strongly enriched for TFTC-3, TFTC-5, Pol III, and TBP as expected (**Fig. 1D**). However, we also observed TFIIIC-bound sites with low or no Pol III and TBP enrichment (**Fig. 1D**). Of the 1,029 TFIIIC sites, 525 TFIIIC sites (51%) showed strong Pol III enrichment (“Pol III-associated TFIIIC sites” hereafter), whereas 504 TFIIIC sites (49%) showed no or very low Pol III and TBP enrichment (“extra-TFIIIC sites” hereafter; **Fig. 1E, F**).

The lack of Pol III and TBP binding in the subset of the TFIIIC sites may represent a premature Pol III preinitiation complex assembled at Pol III-transcribed non-coding RNA genes [50]. Alternatively, they could be unrelated to Pol III transcription and similar to extra-TFIIIC sites reported in other organisms. To distinguish these two possibilities, we examined the presence of transcription start sites (TSSs) of non-coding RNA genes near TFIIIC-bound sites. We first confirmed that almost all Pol III-associated TFIIIC sites (464 of 525 sites, 88%) harbored non-coding RNA gene TSSs within 100 bp, and the vast majority of these genes encoded tRNAs (376 sites, 72%) or snoRNAs (52 sites, 10%) (**Fig. 1G**). In contrast, only 4% of extra-TFIIIC sites (20/504) harbored non-coding RNA gene TSSs within 100 bp (**Fig. 1G**). Thus, these Pol III-independent TFIIIC binding events in the *C. elegans*

genome are unlikely to participate in local Pol III-dependent transcription, a characteristic behavior of extra-TFIIC sites reported in other organisms [37].

125 ***C. elegans* extra-TFIIC sites possess strong Box-A and Box-B motifs**

The TFIIC complex binds the Box-A and Box-B DNA motifs [35] (**Fig. 1A**). However, the majority of extra-TFIIC sites in yeast and human possess only the Box-B motif [39,40,43]. To determine whether *C. elegans* extra-TFIIC sites contain Box-A and Box-B motifs, we performed *de novo* DNA motif analyses at TFIIC sites. Almost all of the 504 extra-TFIIC sites in *C. elegans* harbored both the Box-A and Box-B motifs (90% with Box-A,  $E=1.9 \times 10^{-1568}$ ; 87% with Box-B,  $E=1.5 \times 10^{-1602}$ ; **Fig. 1H**). The pervasiveness of these motifs in extra-TFIIC sites was comparable to that in Pol III-associated TFIIC binding sites (94% with Box-A,  $E=1.1 \times 10^{-589}$ ; and 92% with Box-B,  $E=3.5 \times 10^{-1308}$ ) (**Fig. 1H**).

Because the Box-A and Box-B motifs constitute the gene-internal promoter for tRNA genes (**Fig. 1A**), we hypothesized that extra-TFIIC sites correspond to genetic elements similar to tRNA genes. In *C. elegans*, a class of interspersed repetitive elements called CeRep3 has been suspected as tRNA pseudogenes [51]. To determine whether extra-TFIIC sites coincide with repetitive elements, we surveyed the overlap between TFIIC sites and all annotated repetitive elements (**Fig. 1I**). Strikingly, 44.6% (225 sites) of extra-TFIIC sites overlapped repetitive elements (permutation-based empirical  $P < 0.001$ ), and almost all of the overlapped elements (95.1%; 214 sites) were the CeRep3 class of repetitive elements (**Fig. 1I**). In contrast, although significant, only 8.2% (43 sites) of Pol III-associated TFIIC sites overlapped repetitive elements of any class (permutation-based empirical  $P = 0.001$ ). Therefore, unlike extra-TFIIC sites in yeast and humans, *C. elegans* extra-TFIIC sites harbored both the Box-A and Box-B motifs; furthermore, a large fraction of these sites corresponded to a class of putative tRNA pseudogenes CeRep3.

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### ***C. elegans* extra-TFIIC sites are not associated with regulatory elements for protein-coding genes**

Previous studies in human and *S. cerevisiae* reported that extra-TFIIC sites are overrepresented near protein-coding gene promoters, proposing a potential role in protein-coding gene regulation [39,43]. To determine whether *C. elegans* extra-TFIIC sites were located near protein-coding genes, we measured distance to nearest protein-coding gene TSSs. Unlike extra-TFIIC sites in other organisms, *C. elegans* extra-TFIIC sites were not located closer to protein-coding gene TSSs than Pol III-associated TFIIC sites (Mann Whitney *U* test,  $P=0.01$ ) or than randomly permuted extra-TFIIC sites (Mann-Whitney *U* test,  $P=2 \times 10^{-5}$ ; **Fig. 2A**). To further investigate the relationship between extra-TFIIC sites and regulatory elements for protein-coding genes, we examined the chromatin states defined by a combination of histone modifications in early embryos [4]. Consistent with the distance-based analysis, extra-TFIIC sites were not overrepresented within “promoter” regions (14 sites, 2.8%, permutation-based empirical  $P=0.4$ ) (**Fig. 2B**). Instead, as expected from the CeRep3 repeat overrepresentation, extra-TFIIC sites were highly overrepresented among the chromatin states associated with repetitive elements including “Transcription elongation IV: low expression and repeats” (51 sites, 10.5%), “Repeats, intergenic, low expression introns” (85 sites, 17.5%), and “Repeat, RNA pseudogenes, H3K9me2” (192 sites, 39.5%) (permutation-based empirical  $P<0.001$ ; **Fig. 2B**). Extra-TFIIC sites were also overrepresented within the “Enhancer II, intergenic” chromatin state (40 sites, 8.2%, permutation-based empirical  $P<0.001$ ) (**Fig. 2B**). However, we did not observe robust enhancer-related H3K27ac or H3K4me1 signals at the extra-TFIIC sites (**Fig. 2C**). Therefore, *C. elegans* extra-TFIIC sites were not associated with promoter or enhancer regions for protein-coding genes.

### ***C. elegans* extra-TFIIC sites are densely clustered in the distal arms of autosomes**

The lack of association with local regulatory features led us to hypothesize that extra-TFIIC sites were related to large-scale organization of chromosomes as in the case of yeasts [40,52]. Because tRNA genes are highly over-represented in the X chromosome [23], we asked whether extra-TFIIC sites were also distributed unevenly in the genome. In stark contrast to Pol III-associated TFIIC sites, which

were highly over-represented in the X chromosome (202 of the 525 sites; permutation-based empirical  $P < 0.001$ ), extra-TFIIC sites were under-represented in the X chromosome (18 of the 504 sites; permutation-based empirical  $P < 0.001$ ; **Fig. 3A**). Extra-TFIIC sites were instead highly overrepresented in chromosome V (195 of the 504 sites; permutation-based empirical  $P < 0.001$ ; **Fig. 3A**). *C. elegans* autosomes can be subdivided into three domains of similar size (left arm, center, and right arm) based on repetitive element abundance, recombination rates, and chromatin organization [17,23,25]. We found that most extra-TFIIC sites were located in autosome arms (486 of the 504 sites; **Fig. 3B**). In addition, extra-TFIIC sites were overrepresented in only one of each autosome's two arms (overrepresented in the right arm of chromosome I; left arm of chromosome II; left arm of chromosome III; left arm of chromosome IV; right arm of chromosome V; permutation-based empirical  $P < 0.001$ ; **Fig. 3C**). Furthermore, within autosomal arms, extra-TFIIC sites were locally densely clustered (**Fig. 3D**), with a median interval between neighboring extra-TFIIC sites of 1207 bp (Mann Whitney *U*-test vs. within-arm permutations  $P = 2 \times 10^{-16}$ ; **Fig. 3E**). Among the autosome arms, the chromosome V right arm contained the largest number of extra-TFIIC sites (188 sites) with extensive clusters (**Fig. 3B, D, E**). Thus, *C. elegans* extra-TFIIC sites were fundamentally different from Pol III-associated TFIIC sites in their genomic distribution and highly concentrated at specific locations within autosomal arm domains.

### 190 ***C. elegans* extra-TFIIC sites intersperse H3K9me3-marked heterochromatin domains**

The autosome arms in *C. elegans* exhibit high levels of H3K9me2 and H3K9me3, histone modifications associated with constitutive heterochromatin [16,17]. Furthermore, in each autosome, H3K9me2 and H3K9me3 signals are known to be stronger in one arm than the other [16,27]. Because extra-TFIIC sites have been implicated in heterochromatin insulation [53,54], we hypothesized that extra-TFIIC sites were located near H3K9me2 or H3K9me3-marked regions. To test this hypothesis, we compared TFIIC-bound sites with the locations of H3K9me2 and H3K9me3-enriched regions identified in early embryos [3]. Strikingly, the chromosome arms in which extra-TFIIC sites were overrepresented coincided with the arms that exhibited strong H3K9me2 and H3K9me3 enrichment (**Fig. 4A, B**). Despite this domain-scale co-localization, however, 88% of extra-TFIIC sites resided in regions not



200 enriched for H3K9me3 or H3K9me2 at the local level (**Fig. 4C**). In particular, extra-TFIIC sites were strongly underrepresented in H3K9me3-enriched regions (only 2% in H3K9me3-enriched regions; permutation-based  $P < 0.001$ ; **Fig. 4D**). However, extra-TFIIC sites were located significantly closer to H3K9me2-enriched regions (median distance 3.1 kb; **Fig. 4E**) and H3K9me3-enriched regions (median distance 12.5 kb; **Fig. 4F**) compared with Pol III-associated TFIIC sites (H3K9me2, median distance 34.8 kb, Mann Whitney  $U$ -test  $P < 2 \times 10^{-16}$ ; H3K9me3, median distance 50.1 kb,  $P < 2 \times 10^{-16}$ ) or extra-TFIIC sites permuted within autosomal arms (H3K9me2, median distance 11.3 kb, Mann Whitney  $U$ -test  $P = 1 \times 10^{-14}$ ; H3K9me3, median distance 28.9 kb,  $P = 7 \times 10^{-14}$ ). Our analysis thus revealed that *C. elegans* extra-TFIIC sites were located close to, but not overlapped with, H3K9me2 and H3K9me3-enriched regions within autosomal arm domains.

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### ***C. elegans* extra-TFIIC sites were located within nuclear membrane-associated domains**

In *S. pombe* and *S. cerevisiae*, extra-TFIIC sites are localized at the nuclear periphery and thought to regulate spatial organization of chromosomes [40,46]. tRNA genes in *S. pombe* are associated with nuclear pores [55]. In *C. elegans*, Pol III-transcribed genes such as tRNA genes are associated with the nuclear pore component NPP-13 [49] (**Fig. 5A**). Owing to the similarity of extra-TFIIC sites to Pol III-transcribed genes, we hypothesized that extra-TFIIC sites associated with NPP-13. To test this hypothesis, we compared the location of extra-TFIIC sites with that of NPP-13-bound sites identified in mixed-stage embryos [49]. As expected, a large fraction of Pol III-associated TFIIC sites (215 sites, 41%) overlapped NPP-13-bound sites (permutation-based  $P < 0.001$ ; **Fig. 5B**). In contrast, while statistically significant (permutation-based  $P < 0.001$ ), only 6 of the 504 extra-TFIIC sites (1.2%) overlapped NPP-13-bound sites (**Fig. 5A**). Thus, extra-TFIIC sites were not likely to be associated with the nuclear pore.

Another mode of chromatin-nuclear envelope interactions in *C. elegans* is mediated by nuclear membrane-anchored, lamin-associated protein LEM-2 [28] (**Fig. 5A**). LEM-2 associates with the specific regions within the arms of chromosomes called “LEM-2 subdomains” interspersed by non-associated “gap” regions of various sizes [28]. We therefore investigated the location of extra-TFIIC

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230 sites with respect to that of LEM-2 subdomains and gaps (**Fig. 5C**). Strikingly, 441 of the 504 extra-TFIIC sites (88%) were located within LEM-2 associated subdomains (permutation-based  $P < 0.001$ , permutation performed within chromosomal domains) (**Fig. 5C, D**). In contrast, Pol III-associated TFIIC sites were underrepresented within LEM-2 associated domains (permutation-based  $P < 0.001$ , permutation performed within chromosomal domains). Our results suggest that *C. elegans* extra-TFIIC sites are localized at the nuclear periphery and intersperse H3K9me3-marked heterochromatin regions (**Fig. 5D**).

## 235 DISCUSSION

The TFIIIC complex is responsible for recruiting TBP and Pol III for transcription of small noncoding RNAs, such as tRNAs [35]. In addition to its transcriptional role, the TFIIIC complex has been known to bind genomic locations devoid of Pol III-mediated transcription in yeast [39,40], fly [41], mouse [42], and human [43]. These sites have been termed extra-TFIIIC sites [39,53]. However, whether such extra-TFIIIC sites are present in *C. elegans* had been unknown. Our data demonstrated that half of all TFIIIC-bound sites in *C. elegans* embryos lack RNA Pol III binding, TBP binding, and nearby noncoding RNA genes, revealing pervasive extra-TFIIIC sites in the *C. elegans* genome.

Previous studies have suggested that extra-TFIIIC sites act as genomic insulators by blocking enhancer activity or heterochromatin spreading [40,54,56], or mediating three-dimensional genome interactions [41,56]. Some of the genomic and chromatin features of *C. elegans* extra-TFIIIC sites reported in this paper resemble characteristics of extra-TFIIIC sites participating in chromatin insulation. First, similar to arrays of TFIIIC-bound sites capable of insulating heterochromatin and enhancer activities in *S. pombe* and human cells [40,56], *C. elegans* extra-TFIIIC were densely clustered in *cis*. Second, similar to the observation that some extra-TFIIIC sites are located at the boundaries of heterochromatin [40,54,56], *C. elegans* extra-TFIIIC sites were located close to, but not within, H3K9me3-marked regions. Third, similar to yeast extra-TFIIIC sites localized to the nuclear periphery [40,46], *C. elegans* extra-TFIIIC sites coincided with genomic regions known to be associated with nuclear membrane protein LEM-2 [28]. These observations raise the possibility that *C. elegans* extra-TFIIIC sites may act as a chromatin insulator.

There are also differences between *C. elegans* extra-TFIIIC sites and those in other organisms. First, unlike yeast and human extra-TFIIIC sites that only possess the Box-B motif [39,43], *C. elegans* extra-TFIIIC sites possess both the Box-A and Box-B motifs. Second, unlike human, fly, and mouse extra-TFIIIC sites that are located near regulatory elements for RNA Polymerase II (Pol II) transcription [41–43], *C. elegans* extra-TFIIIC sites were not located near gene promoters or enhancers or associated with chromatin features of Pol II-dependent regulatory regions. Third, unlike human and mouse extra-TFIIIC sites that are located near CTCF binding sites [42,43], *C. elegans* extra-TFIIIC sites

are unrelated to CTCF binding because the genome does not encode CTCF [32]. Our study could thus offer an opportunity for a comparative analysis of extra-TFIIC functions across eukaryotic species.

The molecular mechanisms underlying the chromatin organization of *C. elegans* autosomes remain poorly understood. Unlike the X chromosome organized into topological associated domains (TADs), the autosomes do not present strong and pervasive TADs [19]. The condensin binding sites that could create TAD boundaries in the X chromosome did not do so in the autosomes [14]. Several mechanisms for autosome chromatin organization have been proposed. These mechanisms include the antagonism between H3K36 methyltransferase MES-4 and H3K27 methyltransferase RPC2 that defines active vs. repressed chromatin boundaries [4,34]; small chromatin loops emanating from the nuclear periphery that allow active transcription within heterochromatin domains [28]; and active retention of histone acetylase to euchromatin that prevents heterochromatin relocalization [33]. Our data that extra-TFIIC sites are highly overrepresented in autosome arms and cluster in *cis* near the boundaries of H3K9me3-marked regions warrant future investigation of whether TFIIC proteins participate in chromatin insulation in *C. elegans* autosomes.

How strategies to demarcate chromatin domains have evolved in eukaryotes remain unclear. In vertebrates, CTCF has a central role in defining TAD boundaries by mediating DNA loops and is essential for development [11,57,58]. In *D. melanogaster*, CTCF is essential but does not appear to define TAD boundaries, and instead acts as a barrier insulator [59–61]. In the non-bilaterian metazoans, some bilaterian animals (such as *C. elegans*), plants, and fungi, CTCF orthologs are absent [32,62]. In contrast to CTCF, the TFIIC proteins are conserved across eukaryotes [63] and extra-TFIIC sites have been reported in human, [43], mouse [42], fly [41], *C. elegans* (this study), and yeast [39,40]. Whether extra-TFIIC is an evolutionary conserved mechanism for demarcating chromatin domains in eukaryotes, including those lacking a CTCF ortholog, will be an interesting subject of future studies.

## CONCLUSIONS

We identified TFIIIC-bound sites not participating RNA Polymerase III-dependent transcription in the *C. elegans* genome. These “extra-TFIIIC” sites were highly over-represented in the arm domains of the autosomes interacting with the nuclear lamina. Extra-TFIIIC sites formed dense clusters in *cis* near the outer boundaries of individual H3K9me3-marked heterochromatin regions. These genomic features of *C. elegans* extra-TFIIIC sites resemble extra-TFIIIC sites reported in other organisms that have activities in insulating heterochromatin. Our study warrants future investigation of whether TFIIIC proteins participate in heterochromatin insulation in *C. elegans*.

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## METHODS

### ChIP-seq dataset

ChIP-seq of TFTC-3, TFTC-5, RPC-1, and TBP-1 were performed in chromatin extracts of the mixed-stage N2-strain embryos in duplicates and have been reported in our previous publication [49]. These  
300 data sets are available at Gene Expression Omnibus (GEO; <http://www.ncbi.nlm.nih.gov/geo/>) with accession numbers GSE28772 (TFTC-3 ChIP and input); GSE28773 (TFTC-5 ChIP and input); GSE28774 (RPC-1 ChIP and input); GSE42714 (TBP-1 ChIP and input). ChIP-seq datasets for H3K4me3, H3K4me1, H3K27ac, H3K9me2, and H3K9me3, performed in chromatin extracts of early-stage N2-strain embryos [3], were downloaded from ENCODE website  
305 (<https://www.encodeproject.org/comparative/chromatin/>).

### Reference genome

The ce10 reference sequence was used throughout. The chromosomal domains (left arm, center, and the right arm) defined by recombination rates [24] was used.  
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### Gene annotation

The genomic coordinates and the type of *C. elegans* transcripts were downloaded from the WS264 annotation of WormMine. The WS264 genomic coordinates were transformed to the ce10 genomic coordinates using the liftOver function (version 343) with the default mapping parameter using the  
315 *ce11ToCe10.over.chain* chain file downloaded from the UCSC genome browser.

### TFIIIC site definition

MACS2 identified 1,658 TFTC-3-enriched sites. Of those, sites that had the TFTC-3 fold-enrichment (FE) score greater than 5, harbored TFTC-5-binding sites within 100 bp, and were located in the  
320 nuclear chromosomes were considered “high-confidence” TFIIIC-bound sites (1,029 TFIIIC sites). The “center” of each TFIIIC site was defined by the position of the base with the largest TFTC-3 FE score. Of the 1,029 TFIIIC sites, those with the maximum Pol III (RPC-1) FE greater than 20 within +/-250 bp of the site center were defined as “Pol III-associated TFIIIC” sites (525) and the remaining sites were

defined as “extra-TFIIC” sites (504). The genomic coordinates for Pol III-associated TFIIC sites and  
325 extra-TFIIC sites are listed in **Table S1**.

### Heatmap

For the heatmaps around TFIIC sites, a set of 20-bp windows with a 10-bp offset that covered a 2-kb  
region centered around the center of TFIIC sites was generated for each site. For each window, the  
330 mean of fold-enrichment score was computed from replicate-combined input-normalized fold  
enrichment bedgraph files. The signals were visualized using the ggplot2’s geom\_raster function  
(version 2.2.1) in R.

### Non-coding RNA genes

335 The genomic location and classification of non-coding RNA genes was described in the *Gene  
Annotation* section. For each TFIIC site extended +/-100 bp from the site center, whether the region  
contained the transcription start site (TSS) of non-coding RNA genes was assessed using the Bedtools  
*intersect* function [64] (version 2.26.0).

### 340 DNA motif analysis

To find DNA motifs *de novo*, 150-bp sequences centered around the center of the TFIIC-bound sites  
were analyzed by MEME (v4.11.3) [65] with the following parameters: minimum motif size, 6 bp;  
maximum motif size, 12 bp; and the expected motif occurrence of zero or one per sequence (-mod  
zoops) and with the 1st-order Markov model (i.e. the dinucleotide frequency) derived from the ce10  
345 genome sequence as the background.

### Genomic intersection and permutation

Unless otherwise noted, the overlap between the 1-bp center of each of the TFIIC sites and genomic  
features of interest (with size  $\geq 1$  bp) was assessed using the Bedtools *intersect* function [64] (version  
350 2.26.0). To estimate the probability of observing the overlap frequency by chance given the frequency,

location, and size of the features of interest and TFIIIC sites, the TFIIIC sites were permuted using the Bedtools *shuffle* function (version 2.26.0). The TFIIIC sites were shuffled across the genome, or within the chromosomes, or within chromosomal domains in which they reside, as described in each analysis section. After each permutation, the permuted set of TFIIIC sites were assessed for the  
355 overlap with the features of interest. This permutation was repeated 2,000 times to assess the frequency at which the number of intersections for the permuted set of the TFIIIC sites was greater or less than the number of intersections for the observed TFIIIC sites. If none of the 2,000 permutations resulted in the number of overlaps greater or less than the observed number of overlaps, the observed degree of overlaps was considered overrepresented or underrepresented, respectively, with the  
360 empirical P-value cutoff of 0.001. The mean number of overlaps after 2,000 permutations was computed for visualization.

### **Repetitive element analysis**

The ce10 genomic coordinate and classification of repetitive elements, compiled as the  
365 “RepeatMasker” feature, were downloaded from the UCSC genome browser. The intersection between repetitive elements and TFIIIC sites was assessed as described in the *Genomic intersection and permutation* section. The permutation of TFIIIC sites was performed within the chromosomes in which they resided.

### **370 Protein-coding gene distance**

The genomic location of protein-coding RNA genes was described in the *Gene Annotation* section. For each TFIIIC site, the absolute distance between the center of the TFIIIC site and the closest TSS of a protein-coding gene was obtained using the Bedtools *closest* function [64] (version 2.26.0). To assess the probability of observing such distance distribution by chance given the frequency and location of the  
375 TSSs and TFIIIC sites, the TFIIIC sites were permuted once using the Bedtools *shuffle* function (version 2.26.0) such that the TFIIIC sites were shuffled within the chromosomes, and the distance between the permuted TFIIIC sites and closest protein-coding gene TSS was obtained. Mann-



Whitney U test, provided by the *wilcox.test* function in R, was used to assess the difference of the distribution of the TFIIIC-TSS distances between groups.

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### **Chromatin state analysis**

The chromatin state annotations for autosomes and the X chromosome are reported previously [4]. The intersection between chromatin state annotations and TFIIIC sites was assessed as described in the *Genomic intersection and permutation* section. The permutation of TFIIIC sites was performed  
385 within the chromosomal domains (see *Reference genome*) to account for the difference of the chromatin state representation among different chromosomal domains.

### **Chromosomal distribution of TFIIIC sites**

The number of TFIIIC sites in each chromosome was assessed as described in the *Genomic  
390 intersection and permutation* section. The permutation of TFIIIC sites was performed across the genome. The number of TFIIIC sites in each chromosomal domain (see *Reference genome*) was assessed as described in the *Genomic intersection and permutation* section. The permutation of TFIIIC sites was performed within chromosomes.

### **395 Extra-TFIIIC site interval**

For each chromosomal domain, the genomic distance between every pair of two neighboring extra-TFIIIC sites (center-to-center distance) was computed in R. To estimate the degree of closeness between extra-TFIIIC sites only explained by the frequency of extra-TFIIIC sites within chromosomal domains, the extra-TFIIIC sites were permuted 2,000 times within chromosomal domains as  
400 described in the *Genomic intersection and permutation* section. In each of the 2,000 permutations, the genomic distance between two neighboring permuted extra-TFIIIC sites was computed, and the mean of the distances was computed. The distribution of the 2,000 means (by 2,000 permutations) was compared with the distribution of observed distribution of TFIIIC interval sizes by Mann-Whitney *U* test.

#### 405 **Analysis of H3K9me2 and H3K9me3 regions**

To define H3K9me2-enriched and H3K9me3-enriched regions, the genome was segmented into 1-kb windows, and the mean fold-enrichment score of H3K9me2 and H3K9me3 (see *ChIP-seq dataset*) was computed for each window using the Bedtools *map* function (version 2.26.0). Windows with the mean fold-enrichment score greater than 2.5 (1.5x standard deviation above mean for H3K9me2; and 1.3x  
410 standard deviation above mean for H3K9me3) were considered enriched for H3K9me2 or H3K9me3 and merged if located without a gap. This yielded 2,967 H3K9me2-enriched regions (mean size, 2.1 kb) and 1,331 H3K9me3-enriched regions (mean size, 4.9 kb).

The intersection between H3K9me2-enriched or H3K9me3-enriched regions and TFIIIC sites was assessed as described in the *Genomic intersection and permutation* section. The permutation of  
415 TFIIIC sites was performed within the chromosomal domains.

For each TFIIIC site, the absolute distance between the center of the TFIIIC site and the closest H3K9me2-enriched and H3K9me3-enriched regions was obtained using the Bedtools *closest* function [64] (version 2.26.0). To assess the probability of observing such distance distribution by chance given the frequency, location, and size of H3K9me2-enriched and H3K9me3-enriched regions and TFIIIC  
420 sites, the TFIIIC sites were permuted once using the Bedtools *shuffle* function (version 2.26.0). For the analysis of the distance to H3K9me2-enriched regions, this permutation was performed within the chromosomal domains. For the analysis of the distance to H3K9me3-enriched regions, permutation was performed with the chromosomal domains but excluding the H3K9me3-enriched regions themselves because the TFIIIC sites were strongly underrepresented in the H3K9me3-enriched  
425 regions.

#### **TFIIIC sites in NPP-13-binding sites**

The 223 NPP-13-binding sites identified in mixed-stage N2-stage embryos are previously reported [49]. The genomic coordinates were converted to the ce10 genomic coordinates using the UCSCtools  
430 *liftOver* function (version 343). The intersection between NPP-13-binding sites and TFIIIC sites was

assessed as described in the *Genomic intersection and permutation* section. The permutation of TFIIIC sites was performed within the chromosomal domains.

### **TFIIIC sites in LEM-2 subdomain and gaps**

435 The LEM-2 subdomains and gaps between LEM-2 subdomains identified in mixed-stage N2-stage embryos are previously reported [28]. The genomic coordinates were converted to the ce10 genomic coordinates using the UCSCtools *liftOver* function (version 343). The intersection between LEM-2 subdomains or gaps of variable size classes and TFIIIC sites was assessed as described in the *Genomic intersection and permutation* section. The permutation of TFIIIC sites was performed within  
440 the chromosomal domains.

### **LIST OF ABBREVIATIONS**

*C. elegans*: *Caenorhabditis elegans*

*D. melanogaster*: *Drosophila melanogaster*

445 *S. cerevisiae*: *Saccharomyces cerevisiae*

*S. pombe*: *Schizosaccharomyces pombe*

ChIP-seq: chromatin immunoprecipitation followed by sequencing

Pol II: RNA polymerase II

Pol III: RNA polymerase III

450 TBP: TATA-binding protein

TFIIIB: Transcription factor III B

TFIIIC: Transcription factor III C

TSS: Transcription start site(s)

455

## **DECLARATIONS**

### **ETHICS APPROVAL AND CONSENT TO PARTICIPATE**

Not applicable.

### 460 **CONSENT FOR PUBLICATION**

Not applicable.

### **AVAILABILITY OF DATA AND MATERIALS**

ChIP-seq datasets are available at Gene Expression Omnibus with accession numbers GSE28772,  
465 GSE28773, GSE28774, GSE42714.

### **COMPETING INTERESTS**

The authors declare no competing interests.

### 470 **FUNDING**

K.I., A.S., and V.B. are supported by National Institutes of Health grant R21 AG054770-01A1. A.S. was supported in part by National Institutes of Health grant R25 GM5533619. A.L. was supported by the Princeton University Program in Quantitative and Computational Biology and the Lewis-Sigler Richard Fisher `57 Fund.

475

### **AUTHOR CONTRIBUTIONS**

K.I. conceived the study. K.I., A.S., A.L., and V.B. analyzed the data. K.I., A.S., A.L., and V.B. wrote the manuscript.

### 480 **ACKNOWLEDGEMENTS**

We thank Jason D. Lieb, Sevinc Ercan, and Sebastian Pott for discussion.

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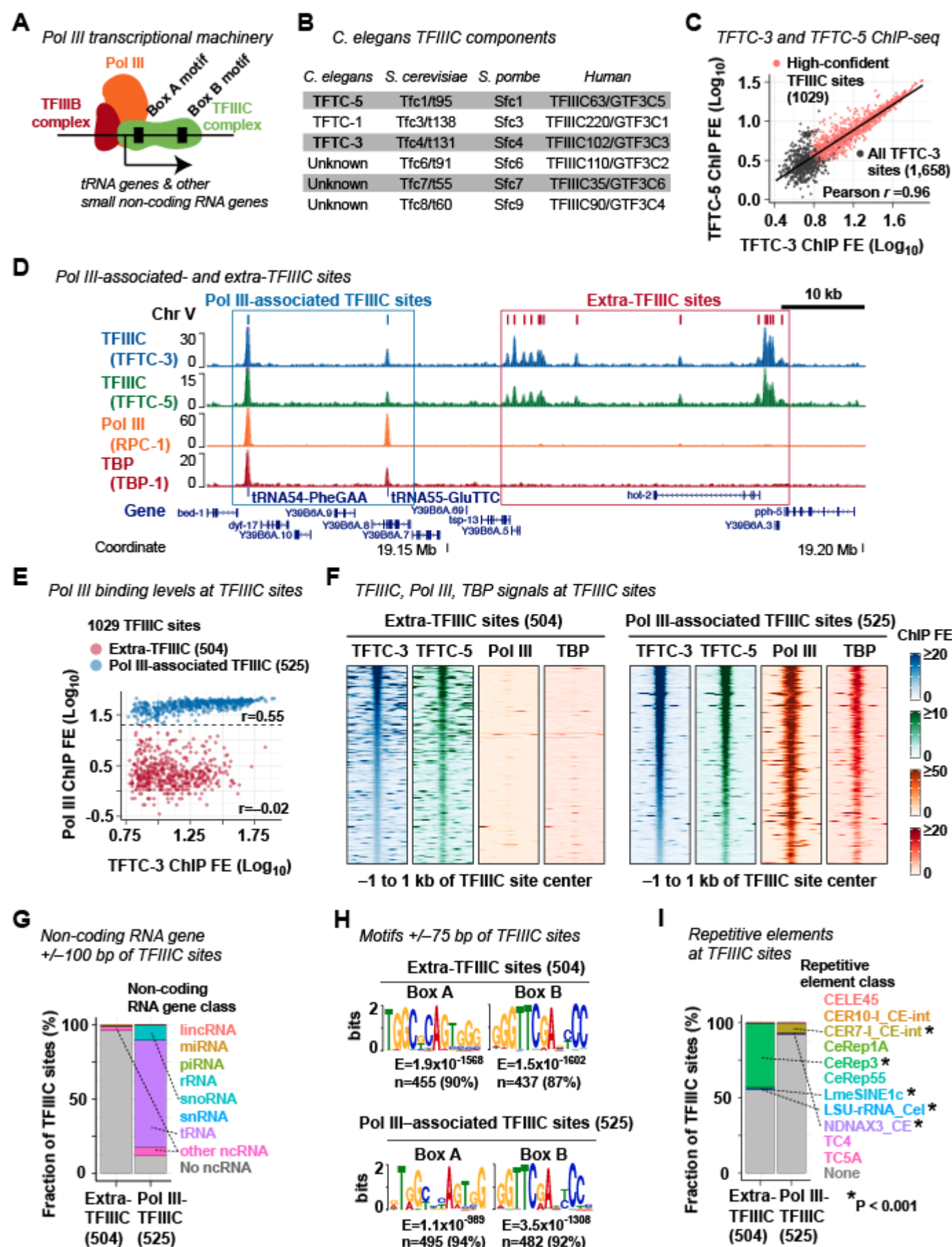
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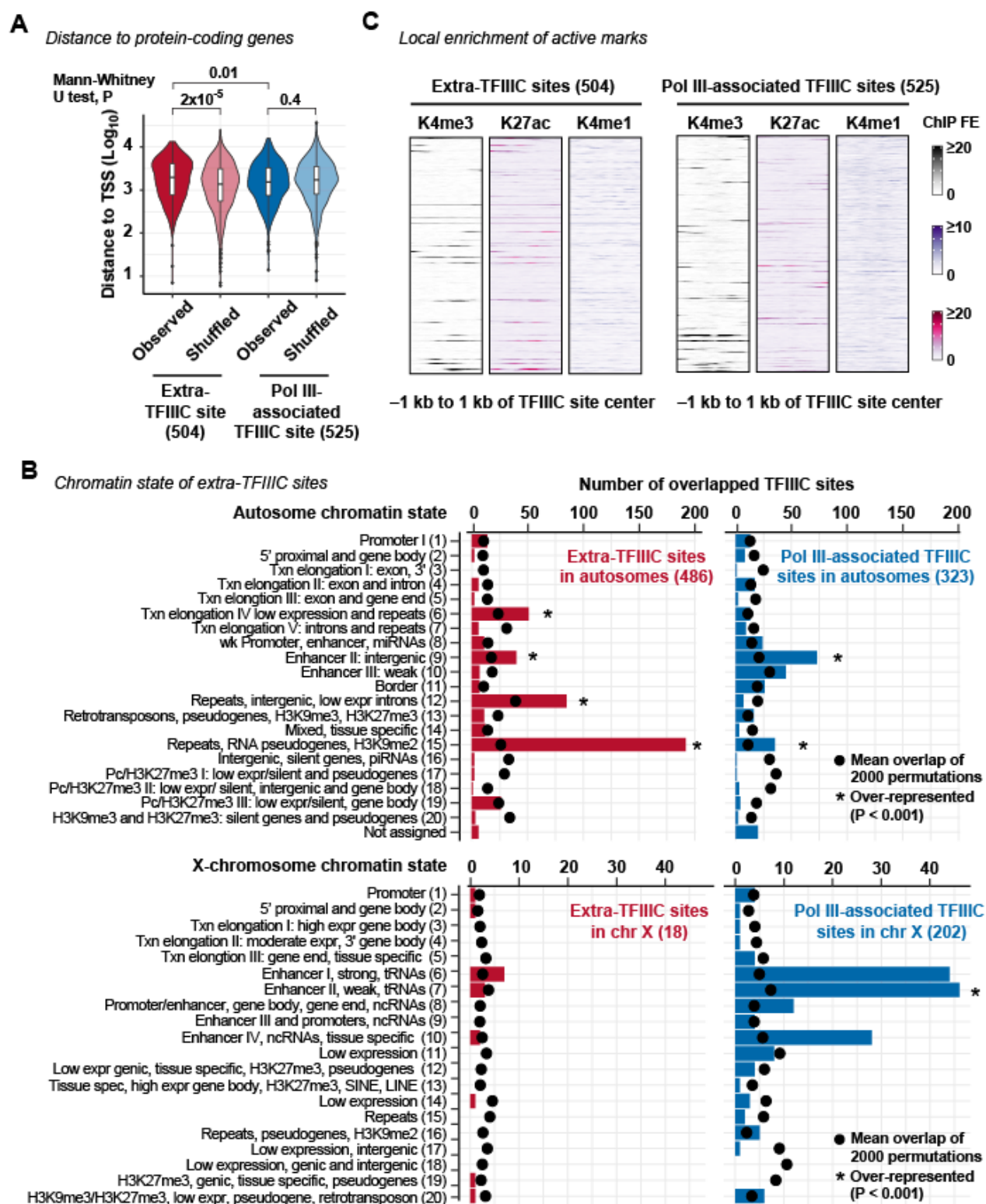
## Figure 1



## FIGURE 1. Identification of extra-TFIIC sites in the *C. elegans* genome

**(A)** A schematic of the RNA polymerase III transcriptional machinery. **(B)** *C. elegans* TFIIC complex proteins and their yeast and human orthologs for reference. **(C)** Correlation between TF3C-3 and TF3C-5 ChIP-seq fold enrichment (FE) scores at the 1,658 TF3C-3-binding sites. The 1,029 high-confident TFIIC sites (see **Methods**) are indicated. **(D)** A representative genomic region showing extra-TFIIC sites and Pol III-associated TFIIC sites. **(E)** TF3C-3 and Pol III (RPC-1) FE scores at the 1,029 TFIIC sites. Extra- and Pol III-associated TFIIC sites are indicated (see **Methods**).  $r$ , Pearson correlation coefficient within the TFIIC subclasses. **(F)** TF3C-3, TF3C-5, Pol III (RPC-1), TBP (TBP-1) ChIP-seq FE signals at extra- and Pol III-associated TFIIC sites. **(G)** Fraction of TFIIC sites harboring the transcription start site of non-coding RNA genes within  $\pm 100$  bp of the TFIIC site center. Dotted lines indicate non-coding RNA gene classes found in  $\geq 10$  TFIIC sites. Pol III-TFIIC, Pol III-associated TFIIC sites. **(H)** DNA sequence motifs found in  $\pm 75$  bp of TFIIC site centers. **(I)** Fraction of TFIIC sites overlapping repetitive elements.  $P$ , empirical  $P$ -value based on 2,000 permutations of the TFIIC sites. Extra-TFIIC and Pol III-associated were individually permuted within chromosomes. Dotted lines indicate repetitive element classes with  $P < 0.001$ .

**Figure 2**

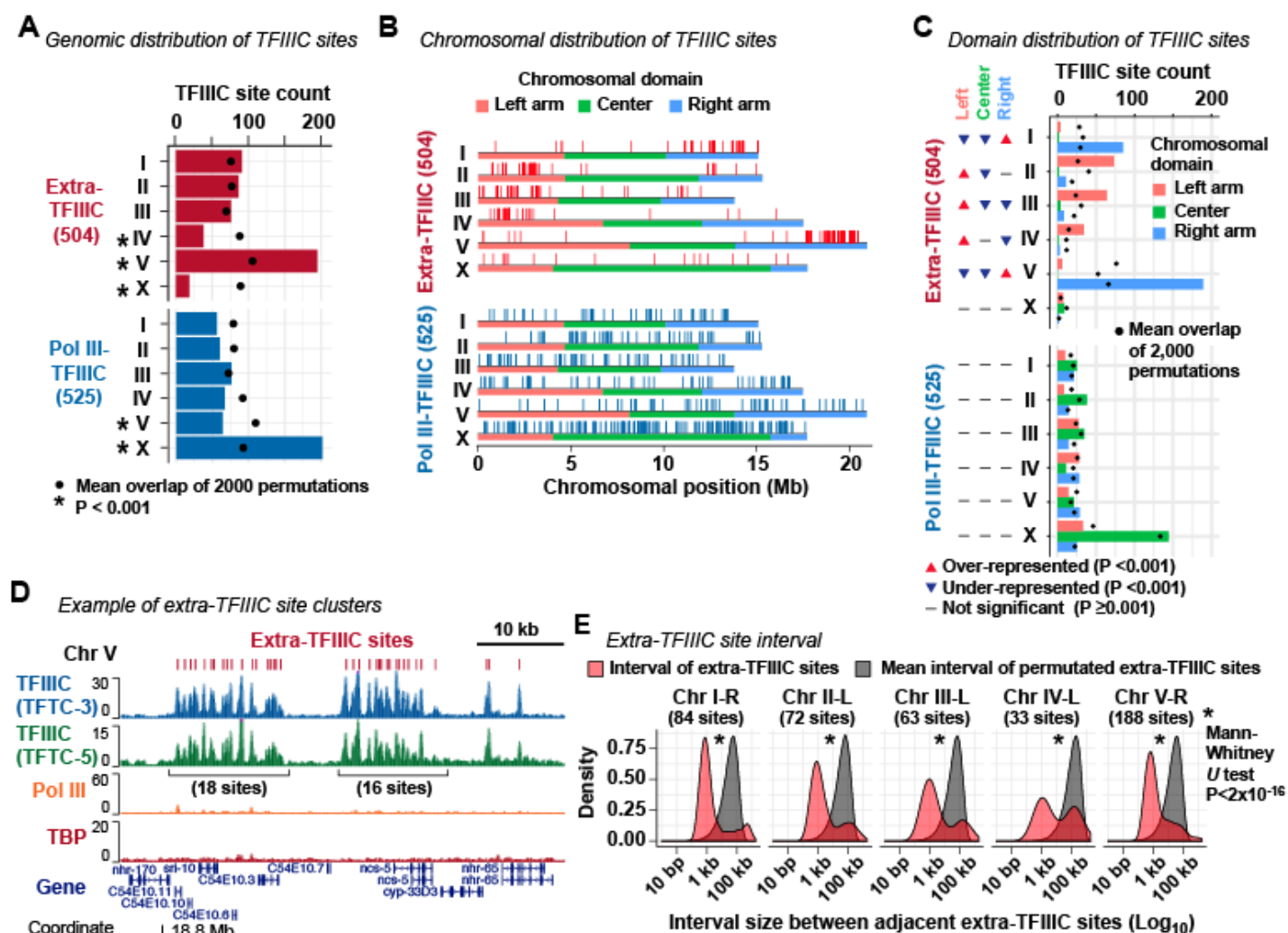


**FIGURE 2. Extra-TFIIC sites do not coincide with protein-coding gene promoters or enhancers**

(A) Distance between TFIIC site center and the transcription start site of protein-coding genes. Observed, the observed distance. Shuffled, the distance of a permuted set of TFIIC sites. (B) Chromatin state annotation of TFIIC sites. The chromatin state annotation reported by Evans et al. (2016) was used. To obtain mean overlap of permutations, extra- and Pol III-associated TFIIC sites were individually permuted 2,000 times within

chromosomal domains. P, empirical P-value based on 2,000 permutations of the TFIIIC sites. **(C)** H3K4me3, H3K27ac, and H3K4me1 fold-enrichment scores (FE) at TFIIIC sites.

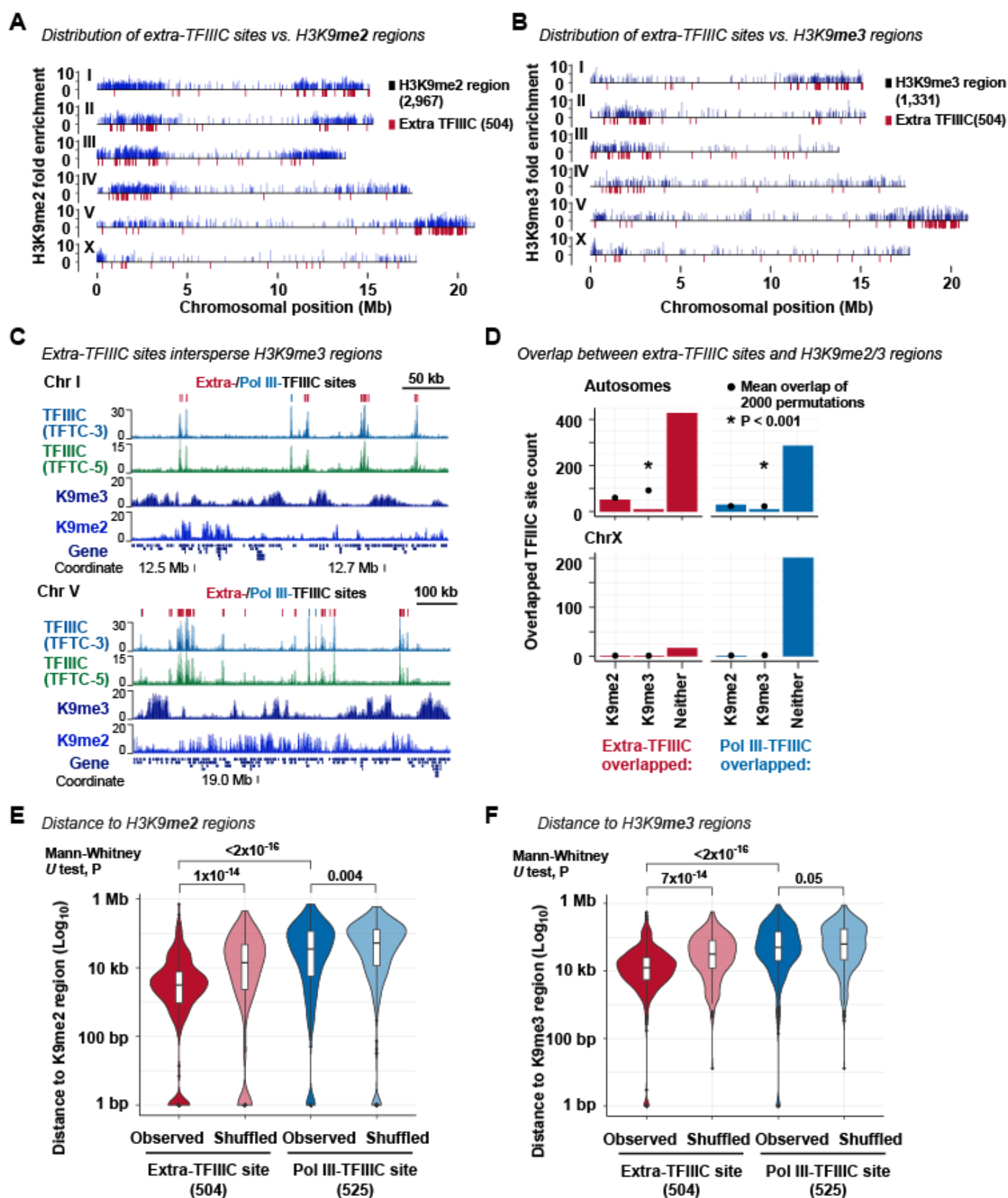
### Figure 3



#### FIGURE 3. Extra-TFIIC sites are clustered in autosomal arms

(A) Distribution of TFIIC sites across chromosomes. To obtain mean overlap of permuted TFIIC sites, extra- and Pol III-associated TFIIC sites were individually permuted 2,000 times across the genome. P, empirical P-value based on 2,000 permutations of the TFIIC sites. (B) Distribution of TFIIC sites within chromosomes. Horizontal color bars indicate chromosomal domains defined by Rockman and Kruglyak (2009). (C) Fraction of TFIIC sites within chromosomal domains. To obtain mean overlap of permuted TFIIC sites, extra- and Pol III-associated TFIIC sites were individually permuted 2,000 times within chromosomes. P, empirical P-value based on 2,000 permutations of the TFIIC sites. (D) A representative genomic region including clusters of extra-TFIIC sites. (E) Red, distribution of the interval size between two adjacent extra-TFIIC sites. Gray, distribution of the mean of the interval sizes of permuted TFIIC sites (2,000 means representing 2,000 permutations). Permutations were performed within chromosomal domains.

## Figure 4

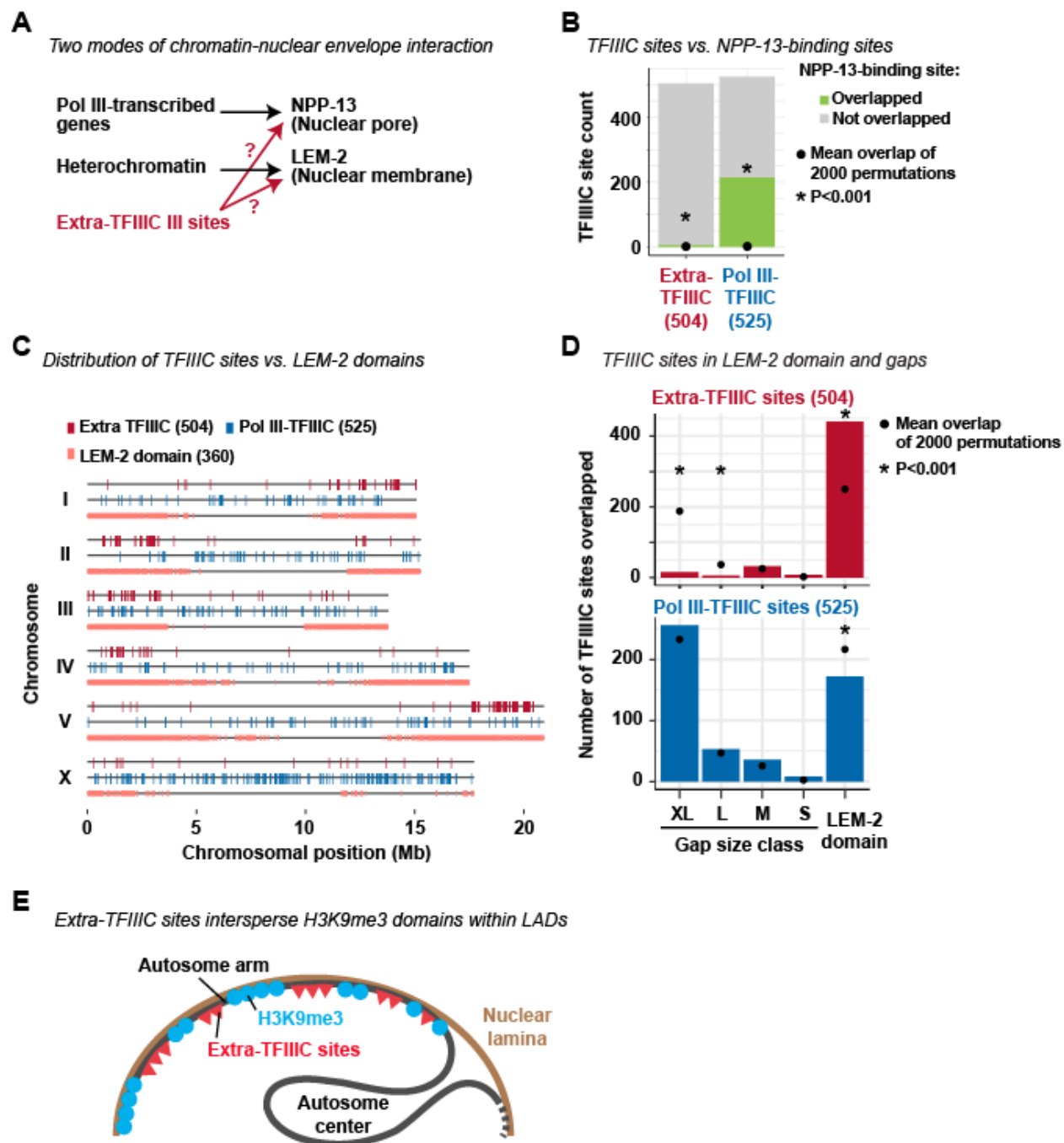


**FIGURE 4. Extra-TFIIC sites are located adjacent to H3K9me3-enriched regions**

(A) Distribution of TFIIC sites along with that of H3K9me2-enriched regions. (B) Distribution of TFIIC sites along with that of H3K9me3-enriched regions. (C) Representative genomic regions showing extra-TFIIC sites

interspersing H3K9me3-enriched regions. **(D)** Fraction of TFIIIC sites overlapping H3K9me2/H3K9me3-enriched regions. Because the H3K9me2/H3K9me3-enriched regions are infrequent in the X chromosome, the X chromosome is plotted separately. To obtain mean overlap of permuted TFIIIC sites, extra- and Pol III-associated TFIIIC sites were individually permuted 2,000 times within chromosomal domains. P, empirical P-value based on 2,000 permutations of the TFIIIC sites. **(E)** Distance between TFIIIC site center and H3K9me2-enriched regions. Observed, the observed distance. Shuffled, distance between a permuted set of TFIIIC sites and H3K9me2-enriched regions. Permutation was performed within chromosomal domains. **(F)** Same as (E), except that distance to H3K9me3-enriched regions is plotted and that permutation was performed excluding H3K9me3-enriched regions to reflect the lack of observed TFIIIC sites within H3K9me3 regions (D).

## Figure 5



### FIGURE 5. Extra-TFIIC sites reside in LEM-2-associated domains

(A) Two hypothetical scenarios in which extra-TFIIC sites associate with the nuclear periphery. (B) Fraction of TFIIIC sites overlapping NPP-13-binding sites defined by Ikegami and Lieb (2013). To obtain mean overlap of permuted TFIIIC sites, extra- and Pol III-associated TFIIIC sites were individually permuted 2,000 times within chromosomal domains. P, empirical P-value based on 2,000 permutations of the TFIIIC sites. (C) Distribution of TFIIIC sites along with that of LEM-2 domains defined by Ikegami et al. (2010). (D) Fraction of TFIIIC sites overlapping LEM-2 domains and gaps of various sizes between LEM-2 domains. To obtain mean overlap of permuted TFIIIC sites, extra- and Pol III-associated TFIIIC sites were individually permuted 2,000 times within chromosomal domains. P, empirical P-value based on 2,000 permutations of the TFIIIC



sites. **(E)** Model for chromosomal organization where extra-TFIIC sites are clustered in the autosome arms corresponding LEM-2 domains and intersperse H3K9me3-enriched regions.