Location, location, location: nuclear structure regulates gene expression in neurons
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Genome architecture plays a critical role in regulating the expression of genes that are essential for nervous system development. During neuronal differentiation, spatially and temporally regulated transcription allows neuronal migration, the growth of dendrites and axons, and at later stages, synaptic formation and the establishment of neuronal circuitry. Genome topology and relocation of gene loci within the nucleus are now regarded as key factors that contribute to transcriptional regulation. Here, we review recent work supporting the hypothesis that the dynamic organization of chromatin within the nucleus impacts gene activation in response to extrinsic signalling and during neuronal differentiation. The consequences of disruption of the genome architecture on neuronal health will be also discussed.

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Introduction
Development and function of the mammalian central nervous system (CNS) pose a great challenge for gene regulation. A fairly homogenous population of neuronal precursors gives rise to a dazzling array of cell types that include neurons, oligodendrocytes, astrocytes, microglia and endothelial cells [1]. Moreover, single cell sequencing experiments have revealed an increasing number of neuronal types with molecularly distinct features [2]. This multitude of neuronal and non-neuronal cell types is generated through careful control of cell proliferation, migration, and terminal differentiation. The CNS largely retains plasticity throughout the life of the organism and this ability to adapt to the external environment is necessary for learning, and ultimately survival.

In all eukaryotic cells, the genome is heavily compacted to fit inside the nucleus. Despite this compression, genes must remain accessible for transcription, DNA replication and other nuclear processes. In the nucleus, chromatin is highly organized within the three-dimensional (3D) space, allowing tight control of gene expression. Genome conformation is regulated at multiple levels, from the wrapping of DNA around histones to form nucleosomes, to epigenetic modifications of DNA and histones, and ATP-dependent histone sliding and exchange. In addition, a key role is played by the formation of higher order domains, and by the movement of genomic regions in relation to nuclear landmarks. Interplay within and between these levels of regulation allows precise gene regulation.

In this review, we will describe recent work demonstrating how genome topology and nuclear positioning influence gene transcription in neurons. We will illustrate the unforgiving nature of nuclear structure deregulation by describing neurological diseases resulting from mutations affecting genome architecture.

The nuclear lamina is a repressive compartment for transcription

The radial arrangement of the genome within the nucleus is highly ordered. The nuclear periphery is enriched in heterochromatin and is generally associated with gene-poor or transcriptionally silent regions of the genome; it is, therefore, considered a compartment that represses transcription. During neuronal development, certain genes move away from the lamina toward the nuclear interior, and this event is concomitant with transcriptional activation [3–5] (Figure 1a). Similarly, in the adult brain, Bdnf activation by kainate-induced seizures is accompanied by movement of the Bdnf gene away from the nuclear periphery [6]. Such movement may relocate genes away from a repressive environment, or toward sites rich in transcriptional apparatus. In embryonic cortical neurons for example, activity-dependent genes such as cFos and Gadd45b relocate to RNA polymerase II (RNAPII) foci in response to depolarization [7,8]

Lamina-associated domains (LADs) are large genomic regions that localize at the nuclear lamina and guide the spatial organization of the interphase nucleus [9]. LADs are predominantly gene-poor and contain silent genes bearing repressive histone modifications. Detailed mapping of LADs in mouse embryonic stem cells (ESCs), neuronal progenitor cells (NPCs) and astrocytes revealed...
that in most cases, dissociation from the nuclear lamina protein LaminB1 resulted in the transcription of genes involved in neuronal and astrocyte specification [5]. Some LADs are constitutive, but hundreds of genes change their lamina interaction during differentiation. Importantly, not only have correlations between radial location and gene expression been observed in a variety of cell types but tethering of genes to the lamina was found to induce gene silencing [10]. Conversely, forced activation of endogenous genes within a LAD was sufficient to relocate the region to the nuclear interior [11]. Interestingly, during the early stages of neuronal differentiation, a substantial group of genes detached from the lamina but remained inactive [5]. These findings suggest that while movement away from the lamina is often necessary for triggering transcription, it is not capable of inducing gene

Neuronal differentiation is marked by genome architecture changes.
There are marked differences at all hierarchical levels of genome architecture between pluripotent embryonic stem cells (ESCs), multipotent neural progenitor cells (NPCs) and terminally differentiated cortical neurons (CNs). (a) Following activation, many neuronal genes move radially toward the center and away from the repressive nuclear periphery. It should be noted that dissociation from the lamina often reflects the acquisition of transcriptional competence rather than gene activation. (b) At TAD level, changes in the boundaries of some TADs, depicted as triangles, lead to the formation of neural specific boundaries. Increasing colour intensity represents regions that show more frequent interaction. (c) At the finest level, dynamic changes in sub-TAD looping and cell type specific factors induce local loop formation.
activation per se (Figure 1a). Nonetheless, these genes were more prone to transcriptional activation at later differentiation stages [5], indicating that release from the lamina may prime them for rapid expression when additional extrinsic signals come into play. This is in accordance with data showing that decondensation of endogenous genes within a LAD was sufficient to relocate the region to the nuclear interior, without affecting transcription [11].

A striking example of how nuclear geometry can be regulated in the nervous system is provided by the dramatic folding of the nuclear membrane observed in hippocampal neurons [12]. The nuclear infoldings were dynamic, and increased in number following synaptic activation, leading to an increase in surface area of the transcriptionally repressive lamina. These nuclear membrane invaginations may facilitate the relay of calcium signals to the nucleus by generating smaller functional compartments, and may affect transcriptional regulation. Quiescent neural stem cells also develop nuclear envelope invaginations, which may contain telomeres and other heterochromatin domains [13]. Although it is unclear whether these invaginations are functionally related to the infoldings described in neurons, they also affect the surface area of a transcriptionally repressive compartment and may lead to the existence of signaling microdomains.

**Role of the nuclear lamina in the pathogenesis of neurological disorders**

The protein components of the lamina are critical for its regulatory function (Box 1), and disruption of them may cause neurological disease. LaminB1 is required for brain development [14,15], for dendritogenesis in cortical neurons [16], and for the regulation of gene expression during olfactory neuron differentiation [17]. Mice lacking the LaminB1 paralog, LaminB2, exhibit cortical development defects [14,15]. Importantly, loss of function mutations of LMNB1 in humans result in neural tube defects [18], whereas mutations of the LAMA gene gives rise to a group of disorders known as the laminopathies [19]. In addition to the well-known premature ageing disease Hutchinson–Gilford progeria syndrome, laminopathies include more than 15 distinct diseases characterized by muscular, metabolic and neurological symptoms. Although the mechanisms are undoubtedly complex and remain to be fully elucidated, genes and indeed entire chromosomes have been shown to change radial nuclear position, implying that transcriptional regulation is likely an important facet of the disease (Figure 3a).

**Chromosomes are organized into topologically associating domains (TADs)**

The genome is arranged in self-interacting genomic regions known as topologically associating domains (TADs) [20,21]. DNA sequences within a TAD interact more frequently with each other than with regions located outside. TAD boundaries function as transcriptional insulators. Chromosome conformation capture technologies, which use ligation to link regions that are close together in 3D space, have allowed identification of genomic contacts on a large scale. Importantly, the existence of TADs was recently confirmed genome-wide using a ligation-independent method that relies instead on genome amplification and DNA sequencing from nuclei dissected into ultrathin sections [22]. TADs have been identified in all cell types studied, including mouse and human brain [23**,24**]. TAD organization is generally stable, and some TAD boundaries are conserved across differentiation stages, cell types and even between species [21]. Multiple lines of evidence indicate that TADs are formed by loading of cohesin onto the genome, which uses energy derived from ATP hydrolysis to spool the DNA through its ring-like structure, forming a TAD [25] (Figure 2). Extrusion stops when cohesin encounters CTCF (CCCTC-binding factor) bound to the DNA, resulting in the majority of TAD boundaries containing both CTCF and cohesin.

A recent study performed ultradeep mapping of genome interactions in mouse ESCs, NPCs and cortical neurons (CNs) differentiated *in vitro*, as well as NPCs and CNs purified from embryonic cortex [23**]. The authors found that during neuronal differentiation, a global reorganization of chromatin takes place (Figure 1b). Although many TAD boundaries remain unchanged, in general TADs become fewer and larger, average intra-TAD contacts become stronger and average inter-TAD contacts are depleted. Importantly, formation of new TAD boundaries was observed close to developmentally regulated genes as they became transcriptionally activated during neuronal differentiation. However, forced activation of a
silent genomic region was insufficient to create a TAD boundary. This suggests that transcription alone is not sufficient to induce the structural changes associated with boundary formation, but rather the formation of new TAD boundaries during neuronal development may be permissive or instructive for the expression of specific genes.

**Regulation of TAD organization during neuronal development**

Boney et al. showed three distinct classes of TAD boundaries in NPCs and CNs: first, boundaries characterized by the presence of CTCF and cohesin; second, CTCF-negative, cohesin-positive boundaries that are close to active gene promoters; and finally, boundaries without active marks of transcription that are enriched for repeat elements [23**]. Below we will discuss the major players in TAD regulation and their role in neuronal gene regulation.

**CTCF**

CTCF is a DNA-binding protein that plays a critical role in determining looping interactions at various levels of the architectural hierarchy [26]. CTCF can regulate both inter-chromosomal and intra-chromosomal interactions and is integral in establishing TAD boundaries and sub-TAD looping. A recent study used an acute depletion system to remove CTCF from murine ESCs, NPCs and astrocytes [27**]. Dramatic loss of looping and insulation at TAD boundaries was observed upon CTCF depletion, which caused the merging of neighboring TADs and the formation of ectopic contacts across the original TAD borders. These changes were coincident with transcriptional dysregulation, although the defects were less severe in astrocytes than ESCs or NPCs, suggesting a less prominent role for CTCF in terminally differentiated cells. This is in keeping with an increasing proportion of CTCF-negative borders during neuronal differentiation [23**]. Interestingly, TAD insulation was re-established upon restoration of CTCF levels in ESCs and NPCs, but not in resting astrocytes, indicating that passage through the cell cycle may be indispensable for restoring TAD boundaries [27**].

In mice, increasing evidence indicates that CTCF influences the expression of neuronal genes [28,29*,30], including the protocadherin genes, whose stochastic variation in expression levels generates neuronal diversity [28,29*]. In the developing cortex, CTCF regulates NPC differentiation and survival [31], and at later stages, dendritic arborization and spine development [28]. In cortical interneurons, CTCF...
Mechanisms of disease caused by alteration of genome topology.
Disruption of chromosome organization is associated with the transcriptional dysregulation observed in many neurological disorders. (a) ‘Organizer’ genes encode proteins with roles in configuring the genome within the 3D nucleus. Mutation in ‘organizer’ genes causes loss of these important proteins, and thereby dramatic downstream effects on gene expression. Examples include mutations in cohesin, YY1, CTCF, and Lamins A and B1. (b) Mutations in TAD boundaries can cause neighboring TADs to merge, creating ectopic contacts such as enhancer–promoter (E–P) loops that induce inappropriate gene activation. Examples include TAD border deletion leading to aberrant activation of LMNB1 and FOXG1. (c) Mutations can prevent sub-TAD interactions such as E–P or active gene looping, abrogating gene activation.

regulates the expression of genes involved in fate determination, and mutations of the CTCF gene lead to defects in cell identity and neuronal migration [30]. Loss of CTCF in the hippocampus is associated with impaired long-term potentiation, reduced dendritic spine density, and defects in spatial and fear memory [29]. As well as finding many deregulated genes in the hippocampus of CTCF-knockout mice, including protocadherin and learning-related genes, the authors found a significant lack of upregulation of Bdnf and Arc genes in response to fear conditioning. The absence of induced Bdnf and Arc expression was associated with disruption of the genome architecture surrounding the gene loci. This suggests that stimuli-responsive gene expression is affected by CTCF-regulated genome architecture, as well as developmentally regulated gene programs.

Cohesin
Cohesin is a ring-like multiprotein complex initially described as part of the complex that holds sister chromatids together during anaphase and later found to have an additional role in genome architecture [32]. Cohesin is critical for genome looping in neural stem cells differentiated into astrocytes in vitro [33]. Loss of cohesin causes loss of TAD insulation, increasing inter-TAD interactions and resulting in widespread transcriptional dysregulation [33]. The transcriptional changes resulting from loss of cohesin may cause a number of downstream effects, depending on the cell type. The nervous system may be particularly sensitive to cohesin-mediated transcription changes as mutation of cohesin in mice induces behavioral defects, possibly due to transcriptional dysregulation of genes necessary for dendrite and synapse development [34,35]. Importantly, mutations of the cohesin pathway in humans cause Cornelia de Lange syndrome, a severe developmental disorder that manifests with neurological symptoms, including psychomotor delay and intellectual disability [36].

Topoisomerase IIβ (TOP2B)
TOP2B is a member of the DNA topoisomerase family responsible for relaxing the DNA supercoils that change DNA topology, and often co-occupies TAD boundaries.
with CTCF and cohesin [37]. An early study indicated that TOP2B plays an essential role in neuronal development, as deletion of the gene in mice induced profound defects in motor and sensory neuron innervation [38]. In cortical neurons, TOP2B and CTCF are found in a complex bound to many neuronal genes, in keeping with the idea that TOP2B may regulate genome structure in the brain [39] and that this may explain some of the effects of TOP2B loss on neuronal development. An intriguing finding of this study is that TOP2B promotes expression of activity regulated genes, at least in part, by causing DNA double strand breaks in response to neuronal activity, suggesting that relieving topological constraint may be critical for gene expression.

**Diseases caused by mutations in TAD boundaries**

TAD boundaries insulate genomic regions from each other, and, therefore, perturbation of TAD boundaries leads to inappropriate interactions between chromatin regions, including aberrant enhancer–promoter looping (Figure 3b). Recent technological advances have allowed sequencing of patient samples to go beyond the exome and identify more variants in the non-coding portion of the genome. As a result, it is increasingly recognized that mutations in TAD borders contribute significantly to both Mendelian heritable disorders and complex multifactorial diseases.

One example is provided by Fragile X syndrome, a disease marked by severe cognitive defects that is caused by a repeat expansion in the FMR1 gene. *FMR1* encodes Fragile X Mental Retardation Protein (FMRP), a translational repressor essential for normal cognitive development [40]. An interesting recent study indicated that the repeat expansion within *FMR1* interferes with CTCF binding sites, disrupting a TAD border and inhibiting *FMR1* expression [41]. The authors hypothesize that TAD disruption alters looping of the *FMR1* promoter to enhancers. Importantly, this disease mechanism may extend to other neurological disorders, as the authors found a striking correlation between disease-associated DNA repeats and TAD boundaries, at boundaries with high CpG island density [41]. A case study of the rare neurological disorder Autosomal Dominant adult-onset demyelinating LeukoDystrophy (ADLD) provides another example of how nuclear architecture may influence gene regulation [42]. The deletion observed in ADLD patients eliminates a TAD boundary, allowing an enhancer to contact the *LMNB1* gene, amplifying its expression. *LMNB1* encodes the LaminB1 protein that is an integral part of the nuclear lamina, itself a repressive compartment for transcription (Box 1). Therefore, this example highlights two points at which gene regulation through nuclear architecture can be manipulated in disease. *LMNB1* overexpression causes progressive CNS demyelination, autonomic dysfunction, ataxia and cognitive impairment.

A survey of 922 deletion sites found in the DECIPHER clinical genome database further revealed that many disease-associated deletions overlap TAD boundaries [43]. Analysis of enhancer usage and gene functions suggested that part of the disease phenotypes could be attributed to the adoption of new enhancers following boundary disruption, and consequent transcriptional changes. The phenotype of the disorders in this survey is complex and often involves several organs; however intellectual disability, along with congenital abnormalities, is one of the most common symptoms [43]. This suggests that the developing brain may be more sensitive to epigenetic insults, compared to other tissues. In one interesting example from this study, deletion of a boundary brings an ectopic foetal brain enhancer into the regulatory domain of *FOXG1* [43]. *FOXG1* encodes a transcription factor that is expressed in the developing nervous system from the earliest stages and is critical for regulating multiple important processes including dorsoventral patterning, neural precursor proliferation, neuronal differentiation, and neuronal cell fate [44]. Patients with the boundary deletion exhibit neurodevelopmental defects, with intellectual disability, developmental delay, and postnatal microcephaly [43], similar to that caused by mutations in *FOXG1* itself [45]. However, a recent study suggests that the deletion causes loss of *FOXG1* regulatory elements that are of more significance in the patient phenotype than the TAD boundary deletion [46].

**Dynamic looping generates sub-TADs**

Within the TADs themselves, highly variable restructuring of the chromatin architecture creates sub-TAD loops, which may link genes to their enhancers or to other co-regulated genes (Figure 2). Sub-TAD looping is tightly controlled to ensure correct spatiotemporal gene regulation, and varies markedly between cell types and developmental stages. Although CTCF is implicated in the formation and maintenance of sub-TADs, other mechanisms are also involved.

During neuronal differentiation, average intra-TAD contacts become stronger [23]. Interactions independent of CTCF binding were detected between promoters of active genes at all developmental stages and correlated with gene expression. Surprisingly, the authors identified very long-range (>30 MB) looping events between activated genes in ESCs, NPCs and CNs. These interactions span TADs, and even occur between genes on different chromosomes. Contact enrichment correlated with the number of exons, RNA splicing events and gene transcription levels. Contacts between transcribed genes have also been described in astrocytes, where the astrocyte-specific gene *Gap* has been found to associate with a number of other expressed genes [47]. One recent exciting discovery was that during olfactory neuron differentiation, olfactory receptor genes aggregate with each other and with their enhancers [48]. In progenitor cells this
clustering seems to be repressive for transcription. However, in mature olfactory sensory neurons, the transcribed olfactory receptor gene separates from the repressive cluster and forms a separate hub together with its enhancers. Thus, long range *trans* interactions create both an activating structure for a single gene, and a repressive structure to prevent expression of more than one olfactory receptor gene per cell [48**].

At a finer level, interactions were observed between the promoter and the entire transcribed region of expressed genes, as well as the termination site [23**](Figure 2). Such intra-gene contacts may be required for RNA splicing and RNAPII recycling. Looping events that allowed the establishment of enhancer–promoter contacts were primarily identified within TADs and, as for contacts between promoters, were predominantly independent of CTCF binding. As perhaps expected, enhancer–promoter contacts correlated with transcriptional activation, further indicating the impact of highly dynamic changes of chromatin architecture on gene expression during neuronal differentiation.

**Regulation of sub-TAD looping during neuronal development**

Proteins implicated in establishing or stabilizing enhancer–promoter loops include CTCF, YY1 [49,50], RNAPII [51], HNRNPU [52], the chromatin remodeler SMARCA4 [53,54], and the Mediator complex [55]. Neural-specific transcription factors have also recently been implicated in organizing genome topology [23**,48**]. Although CTCF does regulate sub-TAD looping, its role diminishes during neuronal differentiation, concomitant with a decrease in the number of CTCF binding sites across the genome [50]. Similarly, in the adult cortex, cerebellum and olfactory bulb, fewer CTCF binding sites were found when compared to the embryonic brain. Decreased CTCF binding correlated with lower levels of CTCF expression [29*,50], which also varied between cell types within the CNS [29*]. Interestingly, the developmental decrease of global CTCF and of CTCF binding was uniquely observed in the neural lineage [50]. ESCs and NPCs showed a similar number of looping events despite reduced CTCF occupancy in NPCs, demonstrating that a higher percentage of NPC-only loops were not anchored by CTCF [50]. This indicates that other proteins contribute to maintaining chromatin architecture in differentiated neurons.

Ying Yang 1 (YY1) is a zinc finger protein that is strongly enriched at interaction sites between NPC-specific genes and enhancers [50]. YY1 is critical for neuronal differentiation [56] and in humans, haploinsufficiency of YY1 results in loss of H3K27ac at enhancers, transcriptional dysregulation and intellectual disability [57]. Knockdown of YY1 in NPCs reduced the interaction frequency between the YY1-bound genes Sox2, Klf4, and Zfp462 and their enhancers, concomitantly inhibiting their expression levels [50]. Interestingly, another study has suggested that YY1 regulates enhancer–promoter looping not only in NPCs but also in pluripotent cells and in other differentiated cell types [49].

The lineage-specific factors that help to organize genome structure also undergo substantial changes during neuronal differentiation (Figure 1c). In ESCs, contacts occur between sites bound by Polycomb, and sites bound by pluripotency-specific transcription factors, such as Nanog. During neuronal differentiation, expression levels, chromatin occupancy and the contacts mediated by these factors all decrease. In NPCs, contacts occur between regions bound by the transcription factor Pax6 [23**], which is a master regulator of the NPC state. NeuroD2 and Tbr1 were also found at contact sites during neuronal development, with interactions increasing during differentiation to peak in mature CNs. Similarly, in olfactory neurons, the transcription factor Lhx2 and its cofactor Ldb1 were necessary for the interaction of olfactory receptor genes with each other and with enhancers [48**]. Taken together, these findings indicate that dynamic organization of chromatin contacts around developmental stage-specific transcription factors within TAD boundaries significantly affects gene expression.

**Consequences of alterations in enhancer–promoter looping**

Disruption of enhancer–promoter loops (Figure 3c) has been linked to a number of neurological diseases. For example, single nucleotide polymorphisms (SNPs) have been found at sites that physically interact with genes implicated in schizophrenia [24*,58]. SNPs that may disrupt chromatin looping have been identified at contact sites of the risk locus CACNA1C [58], the anti-psychotic drug target gene DRD2, several acetylcholine receptors, and genes involved in excitatory synaptic transmission [24*]. Although it is not clear whether all of these mutations interfere with expression of their partnered schizophrenia-related gene, at least in the case of *FOXG1* a mutation in a region found to loop to the promoter abrogated *FOXG1* expression [24*]. Similarly, SNPs associated with four neurodegenerative disorders (Frontotemporal lobar degeneration, Alzheimer’s disease, Parkinson’s disease, and Amyotrophic lateral sclerosis) significantly overlap with CTCF binding sites [59]. SNPs at interaction sites can stimulate inappropriate expression as well as abrogate expression. One SNP associated with Frontotemporal lobar degeneration was found to promote CTCF-mediated looping from cis-regulatory elements to the *TMEM106B* promoter, increasing its expression [59]. Disrupted looping in the absence of specific mutations has also been found in neurological disease, for example at the *GAD1* locus, which encodes the enzyme that synthesizes the neurotransmitter GABA, in schizophrenia [60].

Finally, it is tempting to speculate that changes in genome architecture could represent an early pathogenic
event associated with predisposition for neurological diseases, thereby suggesting that neuropsychological disorders that manifest later in life may be rooted into genome conformational changes that occurred at earlier developmental stages.

Conclusions

It is becoming increasingly clear that genome topology and radial nuclear location affect genome organization and spatiotemporal activation of gene expression in the nervous system. There are many factors that can influence transcription, including epigenetic modifications of DNA and histones, nucleosome remodeling, non-coding RNAs and region relocation to specific nuclear compartments. Many of these mechanisms intercalate with genome topology and exert changes through it. Folding of the genome into TAD and sub-TAD compartments plays a key role in the regulation of gene expression as this, among other events, limits the contacts between genes and their enhancers and co-regulated genes. Of note, genome ‘contacts’ are highly dynamic events that indicate proximity of sequences in 3D space and frequency of interaction, rather than fixed structures. Importantly, radial location of genes within the nuclear space affects both the chromatin environment and promoter accessibility to other genes and regulatory regions. Many neurological disorders can arise as a direct result of dysregulation of genome architecture, both through mutations of genes encoding proteins that have a crucial role in establishing or maintaining appropriate chromatin structure, or through mutations in the DNA sequences at critical contact points (Figure 3). Importantly, changes in genome topology may confer transcriptional priming, rather than activation, and potentially provide the biological basis for predisposition to neuropsychiatric disorders. Although the contribution of genome architecture to the pathogenesis of neurodevelopmental disorders awaits further investigation, technologies aimed at modifying chromatin structure may help restore appropriate gene expression in diseased neurons.

Conflict of interest statement

Nothing declared.

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References and recommended reading

Papers of particular interest, published within the period of review, have been highlighted as:

* of special interest
** of outstanding interest


Neural epigenetics


In this thorough study, the authors map genome topology in incredibly high resolution throughout murine neural development, both in vitro and in vivo. Integrating the resultant maps with additional epigenetic data enables them to make some novel insights into genome organization during neural differentiation, as well as providing an important resource for the neurosciences community.


The authors generate high resolution maps of chromosome contacts during human corticalogenesis, and use them to elucidate links between disease-risk SNPs and the sites they loop to.


Using amino-inducible degradation to dramatically reduce CTCF levels, the authors examine the effect on genome conformation in ES cells, NPCs and postmitotic astrocytes, elegantly addressing the role of CTCF in genome organization.


The authors demonstrate a critical role for CTCF in the genome structure and activation of activity regulated genes, and in memory formation.


Research linking disease-causing tandem repeats, including those in FMR1, to sites that act as TAD and subTAD boundaries.


This exciting recent study demonstrates the importance of trans interactions of olfactory receptor genes and their enhancers in forming hubs that select one olfactory receptor gene for expression and silence all others.


This study demonstrates a novel role for YY1 in chromosome looping in neuronal cells, and explores the decreasing role of CTCF during neuronal differentiation.


