



Role of CTCF in DNA damage response

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ABSTRACT

CCCTC-binding factor (CTCF) is a highly conserved, ubiquitously expressed zinc finger protein. CTCF is a multifunctional protein, associated with a number of vital cellular processes such as transcriptional activation, repression, insulation, imprinting and genome organization. Emerging evidence indicates that CTCF is also involved in DNA damage response. In this review, we focus on the newly identified role of CTCF in facilitating DNA double-strand break repair. Due to the large number of cellular processes in which CTCF is involved, factors that functionally affect CTCF could have serious implications on genomic stability. It is becoming increasingly clear that exposure to environmental toxicants could have adverse effects on CTCF functions. Here we discuss the various ways that environmental toxicants could impact CTCF functions and the potential consequences on DNA damage response.

1. Introduction

Cells are constantly exposed to both endogenous and exogenous DNA damaging agents. Endogenous DNA damage is caused by spontaneous hydroxylation, deamination, DNA replication misincorporation and DNA topoisomerase errors [1–3]. In addition, reactive oxygen species (ROS) generated by normal physiological processes could also damage DNA [4,5]. Types of DNA damage induced by endogenous sources include base modifications [6], single-strand breaks (SSBs) and double-strand breaks (DSBs) [5,7]. Exogenous DNA damage can be caused by physical and chemical sources. Physical sources of DNA damage include ultraviolet (UV) light, which causes pyrimidine-pyrimidone (6-4) photoproducts, as well as ionizing radiation (IR), which causes DSBs [8–12]. Environmental toxicants such as heavy metals, pesticides, dioxins and particulate air pollutants are a major category of chemical agents that damage DNA. They induce several types of DNA damage including base modifications, DNA fragmentation, SSBs and DSBs [13–21].

Consequences of DNA damage can be diverse. If left unchecked, DNA damage could lead to increased risk of a number of diseases including cancer and hereditary disorders [3,22–24]. To repair DNA, the cells have evolved a number of mechanisms termed DNA damage response (DDR) [25]. Depending on the type of DNA damage, specific repair pathways are activated. SSBs and nucleotide errors are repaired by base excision repair (BER), nucleotide excision repair (NER), or mismatch repair (MMR) [26–30]. DSBs are repaired by two main mechanisms, non-homologous end-joining (NHEJ) and homology-directed

repair or homologous recombination (HR) [31]. NHEJ, which involves direct ligation of broken or damaged DNA strands, does not require homology between DNA strands and therefore, is error-prone. This pathway is active through all stages of the cell cycle. HR requires homologous pairing of the sister chromatids and is active during S and G2 phase [31]. Since HR relies on the homology between the two DNA strands, it is generally considered error-free [31,32].

DDR is a complex process involving a number of enzymatic activities. For efficient DNA repair, recruitment of repair factors to the damaged sites needs to be temporally regulated. In addition, it is important that the repair process is spatially constrained [33]. Access of repair factors to inappropriate regions can affect genomic integrity. In eukaryotes spatial constraining of DNA is achieved through its organization into higher-order chromatin structures. Recent genome-wide studies have identified such spatially constrained higher-order chromatin structures such as heterochromatin domains [34,35]. Furthermore, the advent of chromosome conformation capture (3C) and 3C based technologies such as chromosome conformation capture-on-chip (4C), chromosome conformation capture carbon copy (5C) and Hi-C have been instrumental in identifying chromatin organization and interactions in three-dimensional (3D) space [36–38]. Recently, 5C and Hi-C technologies have enabled the discovery of local interacting megabase sized regions of the genome called topologically associated domains (TADs) [39–47]. TADs are structural as well as functional components of chromatin. While interactions occur at high frequency within a given TAD, the interaction frequencies between TADs are low [39].

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Insulator binding proteins, which mediate long-range interactions via their ability to loop DNA through protein–protein interactions are essential for the establishment of higher order chromatin structures including heterochromatin domains and TADs [39,41,43,48]. A number of insulator binding proteins have been characterized in *Saccharomyces cerevisiae* and *Drosophila melanogaster* [49–52]. However, CTCF-binding factor (CTCF) is the only insulator binding protein identified in vertebrates [53,54]. Recent studies have uncovered a major role for CTCF in DNA double strand break response [55–57]. In this review, we discuss emerging evidence that have begun to shed light on the role of CTCF in DNA repair. We also evaluate the deleterious effects of environmental toxicants on CTCF, which could potentially impact DNA repair processes.

2. CTCF is a multifunctional protein

CTCF is a highly conserved, ubiquitously expressed protein. It consists of an N-terminal domain, a C-terminal domain and a central DNA-binding domain. CTCF binds to thousands of sites across the genome [34,58], and its binding is largely invariant among various cell-types. CTCF DNA binding domain contains 11 zinc fingers (ZFs): 10 Cys₂His₂ (C2H2) ZFs and 1 Cys₂His-Cys (C3H1) ZF. Studies on CTCF ZF mutants show that mutations in any of the 11 ZFs lowers its DNA-binding affinity. This suggests that all the 11 ZFs contribute to DNA binding. Furthermore mutations at the core ZFs (ZFs 4–7) impacted the DNA binding of CTCF considerably more than the peripheral ZFs [59,60]. Deletion of N- and C- terminal regions outside the ZF domain did not impact CTCF's DNA binding ability [61]. CTCF binding to DNA occurs sequence specifically and the core consensus CTCF DNA binding sequence is well characterized [34,58,59]. However, CTCF is known to recognize a diverse set of sequences through combinatorial use of its ZFs [59].

CTCF is a versatile protein with a wide array of functions. Originally identified as a transcriptional repressor, CTCF is now known to be involved in several processes including transcriptional activation, X-chromosome inactivation, V(D)J recombination, RNA polymerase II (Pol II) pausing and imprinting [60,62,63]. Of the many functions of CTCF, it is most well-characterized for its role in chromatin insulation [53,64,65]. Insulators are DNA sequence elements that play a key role in preventing inappropriate interactions between adjacent regions of the genome [53,64,65]. The insulator function is mediated by architectural proteins that bind the insulator sequences. While a number of insulator-binding proteins are known in yeast and flies, CTCF is the only known insulator-binding protein in vertebrates [49–54]. CTCF can function both as an enhancer-blocking insulator and domain-barrier insulator [53,54,64,65]. Through its enhancer-blocking function, CTCF prevents communication between enhancers and inappropriate promoters. Imprinting at the Igf2/H19 locus is a classic example of the enhancer blocking insulation function of CTCF [66–68]. The domain barrier function of CTCF enables organization of the genome into transcriptionally active and silent regions, which is necessary to prevent spreading of condensed chromatin, into the neighboring active regions [35]. CTCF also contributes to the formation of TADs by binding their boundaries and functioning as barriers. CTCF enables long-range interactions through looping of DNA [43,44,69]. This ability of CTCF to mediate long-range interactions and organize genome in three-dimensional (3D) nuclear space forms the basis for a number of its functions [43,44]. Through recent investigations, it is becoming increasingly clear that CTCF plays a major role in DSB repair [55–57].

3. H2AX phosphorylation

Phosphorylation of histone H2A variant H2AX, on serine 139, termed γ H2AX, is one of the first events in DNA damage response [70]. γ H2AX plays a key role in DDR. It is widely used as a surrogate marker of DSBs and as a biomarker for cancer therapy evaluation [71]. Upon

induction of DSB, H2AX is phosphorylated by the phosphoinositide-3-kinase-related protein kinase (PIKK) family, which includes ataxia telangiectasia mutated (ATM), DNA-PKcs, and ATM- and Rad3-related (ATR) [71]. Formation of γ H2AX initiates within 30 s of DNA damage and reaches maximal levels at 10–30 min [70]. γ H2AX spreads into megabase regions (0.5–2 Mb) surrounding DNA lesions and forms DNA repair foci [72]. One γ H2AX focus exists for each DSB. γ H2AX foci function as a signal for the recruitment of DDR proteins. γ H2AX also decondenses chromatin at the damage sites, thereby increasing accessibility, which enables assembly and retention of DNA repair factors [73]. Upon repair of DSBs, the γ H2AX foci decline and disappear. The half-life of γ H2AX foci ranges between 2 and 7 h [74,75].

3.1. CTCF delimits γ H2AX domains

CTCF is rapidly recruited to DSBs induced by several sources including γ -irradiation, IR and laser micro irradiation [55–57,76]. Recruitment of CTCF occurring within 30 s of DSB induction suggests this to be an early event in DDR [56]. Analysis of CTCF mutants show that the zinc finger domain of CTCF is important for its recruitment to DSBs [56]. Interestingly, the timing of CTCF recruitment to DSBs coincides with the phosphorylation of H2AX. This suggests that CTCF is recruited to DNA after DSB induction. However, analysis of CTCF binding sites that are conserved among different cell-types showed that the γ H2AX domains were flanked by CTCF before and during DDR [57]. Together, these studies suggest that upon DNA damage, both newly recruited, as well as constitutively DNA-bound CTCF flank γ H2AX domains. Whether this happens in a site-specific manner remains to be seen. CTCF ChIP-Seq signals at the boundaries of γ H2AX domains are stronger compared to those within the domains [57]. Given the role of CTCF in delimiting heterochromatin domains and TADs by binding their boundaries, it is reasonable to speculate that CTCF could be involved in confining γ H2AX foci to DSB sites. Supporting this notion, γ H2AX domains increase in size upon siRNA-mediated depletion of CTCF [56]. This suggests that γ H2AX can spread outside the domain boundaries and expand the DDR region in the absence of CTCF. In addition, CTCF depletion increased both the number of IR-induced γ H2AX foci (IRIF) in each cell as well as the number of foci-positive cells [55,56], and CTCF depleted cells display hypersensitiveness to IR [55,56]. Full-length CTCF could restore the IRIF formation [56]. This suggests that CTCF plays a key role in constraining γ H2AX domains by binding its boundaries and limiting its distribution to the vicinity of DSBs, thereby restricting the area of DDR (Fig. 1).

3.2. CTCF aids formation of γ H2AX nano-foci

Super-resolution microscopy imaging has shown that a single confocal microscopy identified γ H2AX focus can be further resolved into spatially clustered substructures [57]. These 200 nm diameter substructures named γ H2AX nano-foci have been suggested as the smallest γ H2AX modified chromatin regions [57]. ChIP-chip studies by Iacovoni et al. have revealed similar substructures within γ H2AX domains. This study showed that the γ H2AX domains are discontinuous and contains holes (no- γ H2AX regions) and peaks (high- γ H2AX regions) [72]. By profiling histone H3 using ChIP-chip, Iacovoni et al. demonstrate normal histone occupancy in the holes. Therefore, the γ H2AX-depleted holes are formed by reduction in the levels of phosphorylation [72] rather than decreased nucleosome occupancy [72]. Promoters and transcription start sites (TSS) within γ H2AX domains show reduction in γ H2AX levels [72]. Furthermore, RNA polymerase II (Pol II) is enriched at the holes, indicating active gene promoters. Moreover, highly expressed genes within γ H2AX domains were less enriched for γ H2AX [72]. This could suggest that γ H2AX depletion at active gene promoters is necessary to maintain gene transcription within γ H2AX foci during DDR.

Occurrence of γ H2AX as spatially proximal clusters at DSBs suggests

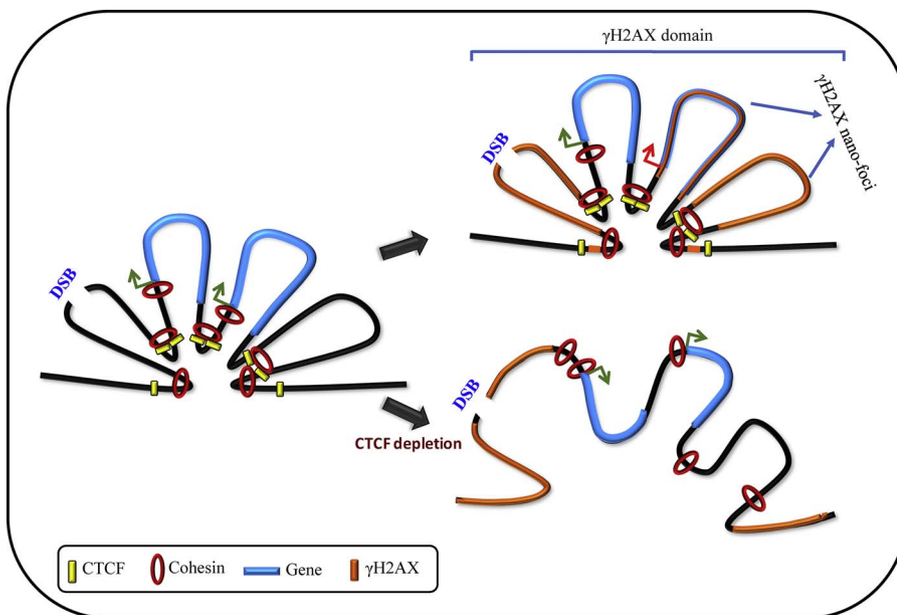


Fig. 1. CTCF establishes γ H2AX domains during DSB repair. Upon DNA damage, CTCF binds DSBs and facilitates γ H2AX domain formation by functioning as domain barrier. Cohesin likely interacts with CTCF during γ H2AX foci and nano-foci establishment. Cohesin binding at the promoters of active genes within domains inhibit γ H2AX establishment, thereby protecting transcription. Loss of cohesin binding at gene promoters cause γ H2AX spreading and gene silencing. CTCF depletion causes loss of higher-order chromatin structures resulting in the spreading of γ H2AX foci and impairment of DNA repair.

Green arrow: active gene; red arrow: repressed gene

higher-order chromatin organization. CTCF mediated long-range interactions form the basis of higher-order chromatin organization in vertebrates. Therefore, long-range interactions mediated by CTCF likely bring the edges of the DNA damaged regions into close proximity thereby restricting DDR to the damaged sites. Spreading of γ H2AX domains caused by CTCF depletion is indicative of the DDR regions spreading beyond the damages sites. Moreover, CTCF depletion resulted in reduction in the number and size of γ H2AX nano-foci [57]. This suggests that CTCF, through its ability to organize 3D structure of the genome, functions as a delimiter of γ H2AX domains and an organizer of γ H2AX nano-foci.

4. Role of cohesin in DSB repair

A number of functions mediated by CTCF involve interaction with its numerous protein partners. One of the major CTCF-interacting proteins is the cohesin complex. Cohesin is important for CTCF-mediated chromatin looping. Cohesin is a multi-subunit protein complex, which plays an essential role in DNA replication and sister chromatid cohesion. Cohesin complex consists of SMC1, SMC3, SCC1 (Mcd1 in yeast and Rad21 in humans) and SCC3 (IRR/SCC3 in yeast and SA1 and SA2 in humans). Cohesin is recruited to DNA upon DSB induction [72,77–79]. The extent of cohesin recruitment around DSBs is species specific. While in yeast cohesin binds to 50–100 kb regions around DSBs, its distribution is limited to 5 kb in humans. During HR, cohesin tethers sister chromatids, facilitating DNA repair [80–82]. Cohesin binding to DSBs is essential for HR and the cohesin subunit SA2 has actually been shown to antagonize the error-prone NHEJ and favour HR [78]. SA2 deletion, on the other hand, increased DNA end joining and promoted NHEJ [78].

4.1. Cohesin maintains H2AX profiles within domains

Recent studies have revealed an important role for cohesin in γ H2AX distribution within the domains. Distribution of γ H2AX within the domains is discontinuous and cohesin binding is enriched at the γ H2AX-depleted regions [83]. Depletion of cohesin subunit SCC1 resulted in γ H2AX spreading to the previously cohesin-bound regions [83]. This suggests a role for cohesin in the organization of γ H2AX within domains by preventing γ H2AX establishment at specific loci. The cohesin enriched/ γ H2AX depleted regions correlated with the promoters of active genes. SCC1 depletion caused downregulation of these

genes [83]. Interestingly, ChIP-Seq experiments before and after IR-induced DNA damage demonstrated that upon DNA damage, increase in cohesin binding occurred at the regions already bound by cohesin prior to damage [79]. This suggests that cohesin recruitment to active gene promoters occurs constitutively, and is not dependent on DSB induction. Upon DSB induction, levels of cohesin at the active gene promoters are further enriched, which inhibits γ H2AX establishment and helps maintain transcription [83]. Upon cohesin depletion, γ H2AX-depleted regions at the TSS of genes within the γ H2AX domains (holes) were undetectable [83]. Moreover, cohesin depletion increased γ H2AX levels on cohesin-bound genes resulting in decreased transcription after DSB induction [83]. This suggests that cohesin at active gene promoters function as a protector of transcription in the event of DNA damage, by preventing γ H2AX establishment (Fig. 1).

In addition to spreading γ H2AX within the domains, SCC1 depletion also induces γ H2AX spreading outside the domains [83], suggesting domain barrier function for cohesin. However, not all of the γ H2AX domains that spread upon cohesin depletion were associated with cohesin binding [83]. It has been suggested that γ H2AX spreading could merely be an outcome of SCC1 depletion-induced increase in the total levels of γ H2AX [83]. Therefore, while the role of cohesin in protecting transcription within γ H2AX domains during DSB repair is clear, its role in constraining γ H2AX domains remains inconclusive.

4.2. Cohesin cooperates with CTCF during establishment of long-range interactions

Cohesin shares the core consensus DNA binding sequence with CTCF [84,85]. Genome-wide studies show extensive colocalization of cohesin and CTCF [84,85]. Moreover, CTCF depletion abrogated cohesin binding at specific sites showing that CTCF is involved in its recruitment [86]. Therefore, it is likely that CTCF is required for cohesin recruitment at DSBs. Moreover, CTCF and cohesin cooperate during the organization of higher-order chromatin structures [86]. Therefore, it is plausible that CTCF and cohesin interact during the establishment and maintenance of γ H2AX foci, which requires higher-order chromatin organization. Regulation of γ H2AX domains by CTCF and cohesin is reminiscent of TADs. CTCF depletion results in increased inter-TAD interactions, suggesting its role as a barrier of TADs. On the other hand, cohesin depletion inhibited intra-TAD interactions, suggesting its role in chromatin organization within the TAD domains [86]. Similarly, CTCF borders γ H2AX domains and depletion of CTCF results in the

spreading of γ H2AX into the surrounding regions. However, cohesin depletion, while not always associated with γ H2AX domain spreading, is clearly associated with the increase in γ H2AX within the domain [83]. This suggests that while CTCF primarily functions as a γ H2AX domain barrier, cohesin plays a role in the maintenance of the nano-foci within γ H2AX domains (Fig. 1).

5. CTCF recruits BRCA2 to DSBs during HR

Hilmi et al. recently demonstrated a direct role of the CTCF in HR through its ability to recruit BRCA2 to DSBs [55]. *BRCA2* is a tumour suppressor gene. Mutations in *BRCA2* are associated with increased risk of breast and ovarian cancers. *BRCA2* deleted cells as well as mice are hypersensitive to radiation and genotoxic agents, and exhibit spontaneous chromosomal aberrations [87,88]. Moreover, *BRCA2* deficient cells display chromosomal instability and defects in homologous recombination [87,88]. Upon DNA damage, *BRCA2* is recruited to DNA, which in turn recruits RAD51. RAD51 filament formation, which allows strand invasion and homologous pairing [89–91], is an initial step in the process of HR.

Co-immunoprecipitation experiments demonstrate that exposure to multiple DNA damaging agents favor interaction between CTCF and *BRCA2* [55]. Furthermore, *BRCA2* recruitment to DSBs is diminished in CTCF-depleted cells, although *BRCA2* protein levels remained unaltered [55]. This shows that *BRCA2* is recruited to DSBs through its interaction with CTCF (Fig. 2). Interestingly, CTCF-mediated recruitment of *BRCA2* to DSBs is dependent on poly(ADP-ribosylation) (PARlation) of CTCF. PARP1 inhibitors decrease the interaction between CTCF and *BRCA2*. Moreover, CTCF PARlation-defective mutant cannot recruit *BRCA2* to DSBs although it can bind DNA efficiently [55]. Therefore, although PARlation of CTCF is not required for its DNA binding, it is essential for the recruitment of *BRCA2* to DSBs (Fig. 2). Interestingly, CTCF knock-down did not diminish the binding of NHEJ associated protein 53BP1 to DSBs [55].

6. Effects of environmental toxicants on CTCF

Given the key role of CTCF in DSB repair, factors that can affect the

expression and function of CTCF could have major implications in genomic stability. Here we discuss some of the factors that can adversely affect CTCF functions and potentially impact DNA repair processes.

6.1. Ni-exposure lowers the DNA-binding affinity of CTCF

Exposure of human lung epithelial cells to nickel (Ni), an environmental carcinogen associated with lung and nasal cancers in humans, diminishes the DNA-binding ability of CTCF [35]. The DNA-binding affinity of proteins is dependent on the underlying sequences, with the binding being strongest to the core consensus sequences. Variants of the consensus sequences, which are considered weaker binding sequences, bind the protein with lower affinities [92]. Ni inhibited CTCF DNA binding in a dose-dependent manner. At low concentrations, Ni disrupted CTCF binding only at the weakest sites. With increasing concentrations of Ni, binding was affected even at the stronger binding sites. Therefore, Ni impacts the DNA-binding affinity of CTCF in a dose- as well as binding sequence-dependent manner [35]. CTCF functions as a repressive domain barrier by binding the boundaries of domains marked by H3K27me3 and H3K9me2 [34,35,58]. Ni-exposure-induced impairment of CTCF binding caused disruption of H3K9me2 domain boundaries resulting in spreading of H3K9me2 into active chromatin regions causing gene silencing [35]. Therefore, it is clear that Ni-induced loss of CTCF DNA binding has functional consequences in terms of cellular regulation.

6.2. Zn^{2+} substitution by metal ions impairs DNA binding of zinc finger proteins (ZFPs)

Although the mechanisms underlying Ni-induced reduction in the DNA-binding affinity of CTCF is still not fully understood, extensive studies on the impact of toxic metals on the DNA binding of ZF-transcription factors, XPA and SP1, offer some clues. The DNA-binding domains of XPA and SP1 contain 1 C4-type ZF and 3 C2H2-type ZF, respectively [93]. Zn^{2+} in the ZFs of XPA and SP1 can be substituted by Ni^{2+} [94,95]. This substitution dramatically alters the structure of these proteins, impacting their DNA-binding affinities [94,95]. In

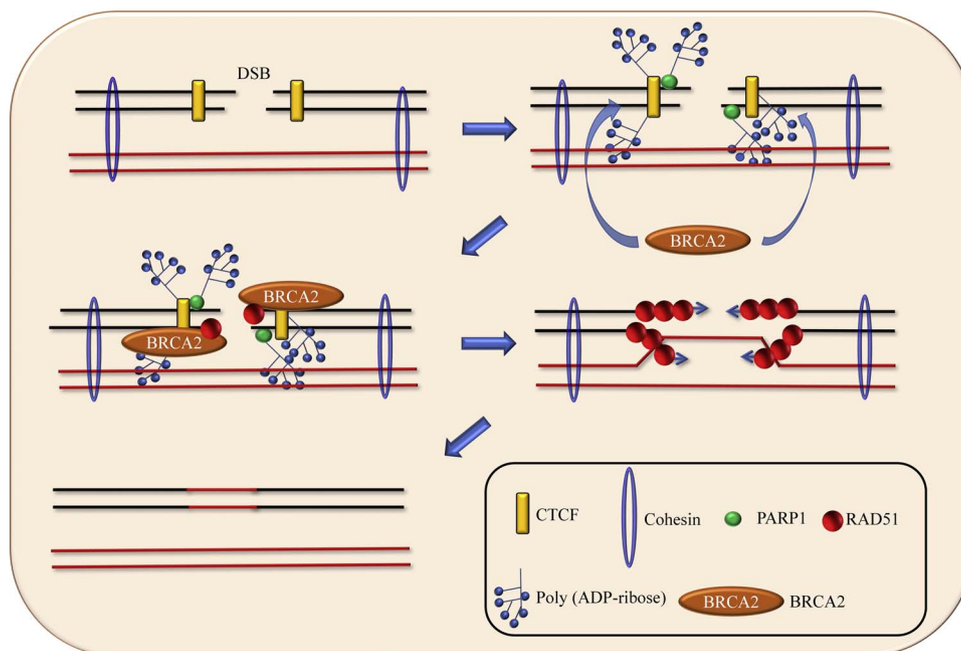


Fig. 2. CTCF recruits BRCA2 to DSBs during HR.

During DSB repair, CTCF recruits BRCA2 to DSBs in a PARlation dependent manner. BRCA2 in turn recruits RAD 51, which form filaments to allow strand invasion and homologous recombination.

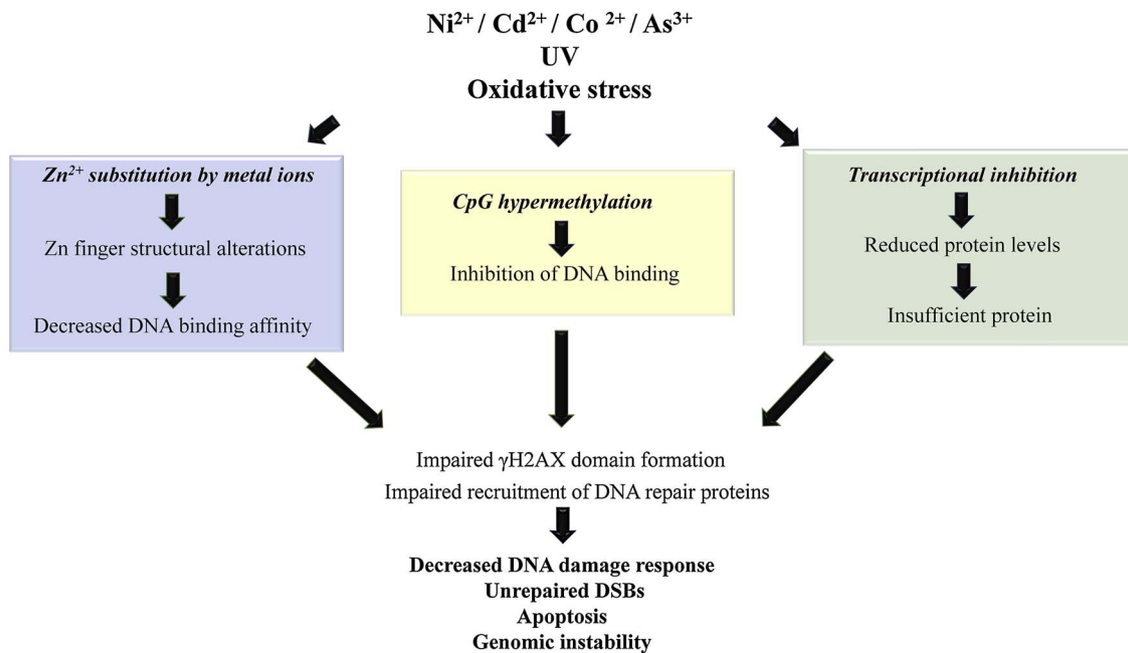


Fig. 3. Environmental exposures could functionally impact CTCF in multiple ways and potentially impair DNA repair function.

addition, As^{3+} , a major environmental carcinogen prevalent in air and drinking water can also substitute Zn^{2+} in the ZFs of XPA, thus interfering with its DNA-binding ability [96,97]. In addition, Co^{2+} and Cd^{2+} have also been shown to inhibit XPA activity by altering its ZF structure. It has been suggested that alteration in the ZF structure due to Zn^{2+} substitution by toxic metals could change the DNA-binding sequence preference of these proteins, thus functionally affecting them [94,95,98]. Furthermore, exposure to certain toxic metals has also been associated with reduction in the Zn content of several proteins. For instance, As^{3+} exposure caused decrease in the Zn content of PARP1 (C3H3-type ZF-protein) and XPA [96,99]. MMA^{3+} exposure decreased the Zn content in APTX (C2H2-type ZFP), PARP1 and XPA [96]. Based on these studies, it is clear that ZFs are major targets of several toxic metals.

Since ZFs are clearly important targets of several toxic metals, it is plausible that the DNA-binding affinity of CTCF, a ZFP, could be affected by environmental toxicants via similar mechanisms (Fig. 3). DSB repair studies in the presence or absence of Ni showed that HR was inhibited in cells exposed to 500 μM NiCl_2 [19]. However, DSB repair was not impacted at lower doses of Ni, suggesting dose-dependent effect of Ni on DNA repair processes. Moreover, all the tested doses of AsO_3 inhibited HR pathway and favored the error-prone NHEJ [19]. It is conceivable that inhibition of DNA repair processes due to exposure to toxic metals could be, at least in part, due to the impact of metal ions on the DNA-binding affinity of CTCF.

6.3. CTCF DNA binding is CpG methylation sensitive

CTCF core consensus DNA binding sequence contains CpG, which can be methylated. A large number of studies have shown that CTCF cannot bind CpG methylated DNA [60,66,67,100]. Negative correlation between DNA methylation and CTCF binding is in fact an important gene regulatory mechanism. Although CTCF binding sites are largely invariant between various cell-types, a subset of CTCF binding sites have been shown to be cell-type specific. Interestingly, 41% of these cell-type specific sites exhibit differential DNA methylation profiles [101]. Therefore, DNA methylation is a key factor in determining the target specificity of CTCF. Imprinting at the *Igf2/H19* locus is one of the most well-understood processes involving interplay between DNA methylation and CTCF binding [66,67]. Maternally inherited imprinting

control region (ICR) at the *Igf2/H19* is hypomethylated, which favors CTCF binding [66,67]. This insulates the *IGF2* promoter from the distal enhancer thus ensuring its silencing. On the other hand, the paternally inherited ICR is methylated, which prevents CTCF binding. This promotes interaction of *IGF2* with the distal enhancers enabling its transcription.

In addition to being a normal process, interplay between CTCF and DNA methylation has also been implicated in pathogenic processes [102]. Expression of *BCL6* oncogene is associated with B cell lymphoma. Lai et al. showed that in lymphomas, *BCL6* is hypermethylated within its first intron, which prevented CTCF binding. Lack of CTCF binding abrogated the CTCF-mediated silencing of *BCL6* and resulted in its increased expression [103]. Similarly, increased expression of *PDGFRA*, a glioma oncogene has also been shown to be caused by the CpG methylation-induced loss of CTCF binding [104]. Hypermethylation at a TAD boundary, which causes loss of CTCF binding, allows activation of *PDGFRA* by enabling interaction with a previously blocked constitutive enhancer [104]. Collectively, these studies demonstrate CpG methylation as one of the most important factors associated with CTCF function.

6.4. DNA methylation can be altered by environmental agents

DNA methylation status is thus a major factor that can influence DNA binding of CTCF. Therefore, exposure to environmental chemicals that can alter DNA methylation profiles could potentially affect CTCF DNA binding and impair DNA repair outcome (Fig. 3). Incidentally, DNA methylation is the most extensively studied epigenetic mark in cells and animals exposed to a multitude of environmental toxicants [105–107]. A large number of studies have shown that a plethora of environmental carcinogens such as Ni, As, Cr, Pb, bisphenol A and persistent organic pollutants alter DNA methylation profiles [105,107]. For instance, Ni-exposure causes extensive DNA hypermethylation [108]. Silencing of *p16* by DNA hypermethylation has been observed in tumors of mice exposed to NiS [108]. In Pb^{2+} exposed individuals, dose-dependent *p16* hypermethylation has been observed [109]. Moreover, extensive DNA hypermethylation has been observed in mammalian cells and in workers with chromate-induced lung cancers [110]. Oxidative stress induced by exposure to a number of environmental toxicants could also potentially play a major role in altering

DNA methylation profiles.

6.5. Environmental toxicants affect CTCF transcription

In addition to impaired DNA binding, another way that CTCF could be affected by environmental agents is through transcriptional regulation (Fig. 3). Cells exposed to inorganic arsenic (iAs^{3+}) show decreased levels of CTCF [111]. Human corneal epithelial and human hematopoietic myeloid cells exposed to UV display decrease in the mRNA and protein levels of CTCF [112]. In PPC-1 prostate cancer cells, oxidative stress induced by H_2O_2 resulted in increased binding of NF- κ B to CTCF promoter causing its repression [113]. Consequently, loss of imprinting was observed at the IGF2 locus [113].

7. Conclusion

Several years of intensive investigations on CTCF have identified a multitude of processes in which it is involved. Although the association of CTCF with DSB repair has been detected only recently, it is already clear that it plays a critical role in this process. CTCF is a genome organizer. It has the ability to mediate long-range interactions and establish higher-order chromatin structures. This ability of CTCF to organize the genome through long-range interactions underlies most of its well-known functions such as transcriptional activation, insulation and imprinting. Indeed, the genome organization function of CTCF appears to play a major role in influencing DNA repair processes as well. CTCF likely constrains the DNA repair sites by functioning as a barrier, thus enabling spatial restriction of DNA repair associated processes such as H2AX phosphorylation, chromatin structural changes and assembly of repair factors. In addition, CTCF could also potentially be involved in restricting DNA damage-induced loss of torsional stress to a single domain [114,115]. Cellular processes such as replication and transcription generate DNA supercoiling and torsional stress within a topological domain [114,115]. Increase in torsional stress is inhibitory to transcription [116]. Single or double strand break could release this torsional stress [117,118]. Although DNA damage is generally considered a harmful process, programmed DNA damage and subsequent repair is currently being recognized as a facilitator of transcription due its role in relieving torsional stress [117,119]. Whether CTCF has any role in restricting loss of topological stress to a single domain during programmed DNA damage needs to be investigated. However, CTCF's association with DNA repair processes may not be limited to its genome organization function. It could facilitate recruitment of DNA repair proteins to their target sites, as exemplified by its role in recruiting BRCA2 to DSBs.

Due to the large number of processes in which CTCF is involved, factors that functionally affect CTCF could have catastrophic consequences on genomic stability. Harmful effects of environmental toxicants on CTCF are currently beginning to be understood. CTCF is particularly vulnerable to environmental factors due to the number of ways that it could be affected: alterations to its ZF structure due to Zn^{2+} substitution by metal ions could affect its DNA binding affinity; CpG hypermethylation due to exposure to a multitude of environmental toxicants could also inhibit its DNA binding; finally, environmental toxicants could decrease its levels through transcriptional inhibition. Since CTCF's DNA-binding ability is critical for its functions, any CTCF dependent process including DNA repair could be impaired by environmental toxicants (Fig. 3). Environmental toxicants that impair CTCF's DNA repair function could particularly be deleterious to genomic stability since many of these toxicants could also induce DNA damage, thereby leaving the cell with damaged DNA and weakened DDR.

The mechanistic details of how CTCF regulates DSB repair is still not fully understood. Moreover, how environmental agents impact CTCF function in terms of DNA repair and the long-term consequences of environmental exposures on DNA repair needs to be investigated in

much greater detail.

Conflicts of interest statement

The authors declare that there are no conflicts of interest.

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