DNA damage and decisions: CtIP coordinates DNA repair and cell cycle checkpoints

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Maintenance of genome stability depends on efficient, accurate repair of DNA damage. DNA double-strand breaks (DSBs) are among the most lethal types of DNA damage, with the potential to cause mutation, chromosomal rearrangement, and genomic instability that could contribute to cancer. DSB damage can be repaired by various pathways including nonhomologous end-joining (NHEJ) and homologous recombination (HR). However, the cellular mechanisms that regulate the choice of repair pathway are not well understood. Recent studies suggest that the tumor suppressor protein CtIP controls the decision to repair DSB damage by HR. It does so by regulating the initiation of DSB end resection after integrating signals from the DNA damage checkpoint response and cell cycle cues.

CtIP links cell cycle control, DNA damage checkpoints and repair

CtIP, the CtBP (carboxy-terminal binding protein) interacting protein, was initially characterized for its role in transcription – first, as a cofactor for the transcriptional repressor CtBP, and also as a binding partner for other proteins including the cell cycle regulators retinoblastoma protein (Rb) and breast cancer 1 (BRCA1) [1–4]. In the G1 phase of the cell cycle, CtIP associates with Rb, allowing CtIP to bind its own promoter as well as the promoters of other E2F target genes such as Cyclin D1. This releases Rb-mediated transcriptional repression and increases expression of genes required for S phase entry [5]. CtIP and Rb could also directly regulate the initiation of DNA synthesis by interaction with MCM7, a component of the replicative helicase [6]. CtIP therefore has both transcription-dependent and -independent roles in cell cycle progression [7,8]. CtIP also plays a central role in the cell cycle checkpoint response to DNA DSBs, with new evidence demonstrating that CtIP controls the choice of DSB repair pathway (Figure 1).

DNA DSBs can be induced in cells by ionizing radiation, treatment with radiomimetic chemicals and perturbations in DNA replication. DSBs induce activation of the G1/S checkpoint, intra-S checkpoint or G2/M checkpoint, depending on when the DNA damage occurs (Figure 1). In the S and G2 phases of the cell cycle, CtIP undergoes CDK-dependent phosphorylation, which promotes its binding to the tumor suppressor protein BRCA1 and the Mre11–Rad50–NBS1 (MRN) complex [9–11]. CtIP undergoes additional phosphorylation by the checkpoint protein kinase ATM in response to DSB damage [12]. Although the underlying mechanism remains controversial, ATM phosphorylation of CtIP leads to transcription of cell cycle inhibitor genes, such as p21 and Gadd45 [12,13]. CtIP also promotes checkpoint signaling and subsequent cell cycle arrest by the Chk1 protein kinase [9,11]. Chk1 activation requires phosphorylation by the ATM-related checkpoint protein kinase ATR, which is activated on single-stranded DNA (ssDNA) coated with RPA protein [14,15]. The role of CtIP in Chk1 activation could be indirect, however; CtIP promotes formation of ssDNA and subsequent ATR activation through its role in DSB resection, a process that digests 5’ termini of DNA ends (see below) [10,16–18]. This model also explains why CtIP is required for Chk1 phosphorylation and checkpoint activation in response to DSBs, but not to stalled replication forks [11]. At stalled replication forks, the ssDNA structure that activates ATR is generated through the uncoupling of the DNA helicase activity from the DNA polymerase activity [19], a mechanism that is distinct from DSB resection.

Recent studies have revealed that DSB repair is a key function of CtIP. DSB damage can be repaired by different pathways, determined in part by the cell cycle stage in which the damage occurs [20]. In G1 phase of the cell cycle, DSBs are mainly repaired by the nonhomologous end-joining (NHEJ) pathway, which re-ligates broken DNA ends. In the S and G2 phases, owing to the presence of sister chromatids, DSBs can also be repaired by homologous recombination (HR) [20]. A fundamental difference between HR and NHEJ is that HR-mediated repair requires substantial DSB resection (approx. 100–200 nucleotides), which generates ssDNA required for homology searching and strand invasion [21]. CtIP is required for DSB resection, and therefore is required for HR [10,16–18,22]. Interestingly, CtIP can also participate in DSB repair by microhomology-mediated end-joining (MMEJ) during the G1 phase [22]. In contrast to NHEJ, MMEJ requires limited resection of DSB ends (four to six nucleotides) to provide short, complementary sequences that stabilize DNA end ligation. CtIP might promote this limited DSB resection, thereby allowing cells to repair DSB damage by MMEJ [22] (Figure 1).

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Despite limited sequence homology, evidence strongly suggests that mammalian CtIP is the functional equivalent of *S. cerevisiae* Sae2 and *S. pombe* Ctp1 [17,23,24]. CtIP orthologs have also been identified in *C. elegans* (COM-1), *Xenopus* (xCtIP), chicken, mouse and *A. thaliana* (AtGR1) [18,22,25–27]. The main regions of similarity of the proteins are in the carboxyl terminus, which includes a conserved CDK phosphorylation site, T847 [17,23,26–28], and in the amino terminus, which is believed to mediate dimerization or oligomerization of these proteins [29,30]. Studies on these CtIP orthologs all support the idea that these proteins are required for DSB resection (Table 1).

**CtIP promotes DSB resection together with MRN**

CtIP is required, but not sufficient, for DSB end resection – this process also requires the MRN complex [31–33]. The MRN complex is highly conserved and is involved in nearly every aspect of the DSB damage response, including DNA damage sensing, signaling and repair [34]. In response to DSBs, MRN acts to recognize damaged chromatin, and binds broken DNA ends [35–37]. Subsequently, MRN initiates checkpoint signaling through recruitment and activation of ATM on damaged chromatin [38,39]. MRN together with activated ATM promotes DSB end resection, and subsequent activation of ATR and repair by HR [40–43]. The Mre11 subunit of MRN harbors DNA-binding, exonuclease and endonuclease activities [34]. It also displays limited DNA unwinding capability in the presence of Rad50 and NBS1 [44,45]. The endonuclease activity of Mre11 is apparently directly involved in DSB resection; however, this activity is not required for MRN function in DSB sensing or for recruitment and activation of ATM [46,47]. These findings suggest the possibility that association of additional proteins with MRN at DSB sites redirects its function from damage sensing and checkpoint initiation to DNA end resection and repair.

CtIP physically interacts with MRN, and both the N-terminus and C-terminus of CtIP have been shown to interact with the NBS1 subunit of MRN [10,16,17]. CtIP also colocalizes with MRN at DSB ends, suggesting that CtIP could influence MRN function [10,16–18]. The molecular nature of the CtIP–MRN interaction has been best characterized in fission yeast. Ctp1 is phosphorylated at SXT/SDT repeats (potential CK2 phosphorylation sites) which mediate binding to the FHA motif of NBS1. This interaction is important for Ctp1 localization to DSBs and Ctp1-dependent resistance to DNA damaging agents [48,49]. It has not been determined whether these CK2 phosphorylation sites are conserved in CtIP proteins in metazoans. However, a similar mechanism mediates the association of the mammalian checkpoint mediator protein MDC1 with NBS1, leading to the retention of the MRN complex on chromatin [50–53].

Recruitment of metazoan CtIP to DSBs also requires MRN [18]. However, CtIP damage recruitment is significantly delayed (by 5–15 min) compared with that of NBS1, suggesting that CtIP is not brought to DSBs passively through its interaction with MRN [18]. Given that CtIP
damage recruitment also requires ATM kinase activity, and that ATM activation depends on MRN, MRN could play an indirect role in the initial damage recruitment of CtIP by activating ATM [18]. MRN could also facilitate CtIP damage relocation by establishing a specific chromatin structure at the DSB ends. Consistent with this idea, S. cerevisiae MRX (Mre11-Rad50-Xrs2, the counterpart of MRN in metazoans) interacts with components of the chromatin remodeling complexes RSC and INO80 to promote histone removal at DSB ends [54,55]. However, it is formally possible that the CtIP-MRN interaction directly mediates CtIP damage recruitment at a later stage. MRN might also be required to retain CtIP at DSBs after its damage recruitment.

Emerging evidence suggests the hypothesis that CtIP regulates MRN function at DSBs, and redirects MRN function from DNA damage sensing to end resection. Consistent with this idea, purified recombinant human CtIP protein stimulates the nuclease activity of a Mre11–Rad50 complex towards a closed ssDNA substrate in vitro [17]. This finding is also consistent with the observation that purified CtIP can interact with Mre11 and Rad50 [16]. Interestingly, yeast Sae2 has been shown to possess endonuclease activity itself in vitro [56]. This raises the intriguing possibility that both Sae2/CtIP and MRX/N directly participate in DSB end processing.

Recently, a two-step model has been proposed for DSB end resection, based on studies in budding yeast [32,57–59]. In this model, Sae2 and MRX initiate the DSB resection process to remove 50–100 nucleotides from the 5’ termini of the DNA. Following this first step, the Exo1 exonuclease and Sgs1/DNA2 helicase/endonuclease function redundantly to carry out further resection to generate long 3’ ssDNA tails [32,57–59]. Notably, functional homologs of all of these proteins are present in fission yeast and metazoans, suggesting that the model for DSB end resection could be conserved in higher organisms (Figure 2). Consistent with this idea, human Exo1 and the Sgs1 ortholog BLM have been shown to be involved in DSB resection [60–62]. Another Sgs1 ortholog, the WRN helicase, might promote the initial unwinding of DNA ends at the DSB, together with the activity of the MRN complex [45,63]. CtIP together with MRN might then cleave the 5’ ssDNA flap at the ssDNA–dsDNA junction to initiate DSB resection. This endonucleolytic cleavage of 5’ termini is absolutely essential for resection of DNA ends that are blocked to processing by a covalently attached protein, such as the DSB ends generated during meiosis that are linked to the Spo11 protein, or DSB ends bound to topoisomerases, which remain after treatment with topoisomerase inhibitors [64–70]. This endo-cleavage step might occur repeatedly, or could be switched to resection by Exo1 or by BLM-DNA2 (Figure 2).

### Regulation of CtIP by CDK, BRCA1 and ATM

The function of CtIP is highly regulated to ensure that DSB resection and HR occur in appropriate stages of the cell cycle (Figure 3). The kinase activity of CDK (CDK1 in yeast...
and CDK2 in mammals) in the S and G2 phases is required for DSB resection [42,71,72]. In addition, Sae2/CtIP is a target of CDK [9,28,73]. These findings provide a mechanistic explanation for why HR primarily operates in the S and G2 phases of the cell cycle. Two CDK consensus phosphorylation sites, S327 and T847, have been identified in CtIP (Figure 4) [9,73]. CDK-dependent phosphorylation of S327 mediates CtIP association with the C-terminal BRCT domains of BRCA1 as well as with MRN [9,10,74,75]. This interaction promotes the ubiquitination of CtIP by the N-terminal ubiquitin-ligase activity of BRCA1 [76]. CtIP phosphorylation at S327 and ubiquitination are required for the damage localization of CtIP and therefore also needed for its function in DSB end resection [76]. Consistent with this idea, CtIP is reported to be an essential gene in chicken DT40 cells, and the human CtIP(S327A) mutant introduced into CtIP-knockout DT40 cells is deficient in supporting HR [22]. However, a more recent study reported that phosphorylation of chicken CtIP at S332 (equivalent to S327 in human CtIP) is not required for DSB resection or HR, and that CtIP-knockout chicken DT40 cells are viable [77]. Further work is needed to resolve these discrepancies. It is possible that phosphorylation of chick CtIP by CDK at other sites plays a...

**Figure 2.** A model for the role of CtIP in DSB end resection. After DSB damage, CtIP is phosphorylated by CDK and ATM, and ubiquitinated by BRCA1, leading to its recruitment to the DNA ends. CtIP damage recruitment commits cells to DNA end resection and repair by HR. CtIP together with MRN initiate DSB resection by promoting endonucleolytic cleavage of the 5′-DNA termini. The WRN helicase might promote this initial DSB resection by unwinding the DNA ends. The 5′ termini are then further resected by the overlapping activities of Exo1 and DNA2/BLM. The ssDNA generated from the DSB end processing is bound by RPA, and the RPA-coated ssDNA structure triggers activation of the ATR checkpoint protein kinase and DNA repair by HR.

**Figure 3.** Cell cycle regulation of CtIP. CtIP function is regulated both at the level of protein expression and post-translational modifications. CtIP protein levels are low in G1 phase and high in the S, G2 and M phases, although the gene encoding CtIP is transcribed throughout the cell cycle [82]. Modification of CtIP by CDK2 and BRCA1 occurs in the S/G2 phase and is required for CtIP function in HR. CDK2-mediated phosphorylation of CtIP at S327 promotes its association with BRCA1 and the MRN complex [9,10,74,75]. CtIP also is ubiquitinated by BRCA1 [76]. CtIP phosphorylation and association with BRCA1 are required for CtIP localization to damaged chromatin [76]. Additional phosphorylation of CtIP at T847 by CDK2 could facilitate CtIP function in DSB resection [73]. Although the mechanism of CtIP function in DNA resection remains to be determined, CtIP modification or binding to MRN could influence MRN enzymatic activity. CtIP might also promote limited DNA resection in G1 in the absence of CDK2 or BRCA1.
redundant role in supporting CtIP damage recruitment and its subsequent role in DSB resection. Interestingly, the human CtIP(S327A) mutant introduced into CtIP-knockout chicken DT40 cells is proficient in supporting MMEJ in the G1 phase [22]. This suggests that human CtIP can translocate to DSBs and promote limited DSB resection in the absence of S327-phosphorylation and BRCA1-dependent ubiquitination, at least during the G1 phase. To date, it is not clear how BRCA1-mediated ubiquitination facilitates association of human CtIP with DNA damage sites. One possibility is that the polyubiquitin chains on CtIP interact with other DSB-associated proteins such as RAP80/Abaxas that contain an ubiquitin-interacting motif [78–80]. It will be important to understand how CtIP activity in DSB end resection is regulated in the G1 phase, and whether CDK and BRCA1 control the extent of CtIP-dependent DNA end resection in the S and G2 phases.

CDK-dependent phosphorylation of CtIP at T847 is also required for DSB resection and subsequent HR in the S and G2 phases [73]. The non-phosphorylatable CtIP(T847A) mutant is still recruited to DNA damage sites, suggesting that CDK2 phosphorylation at T847 regulates the function of CtIP in DSB resection after damage recruitment. The corresponding CDK phosphorylation site in S. cerevisiae Sae2, S267, is also required for DSB resection and HR, indicating a conserved mechanism of regulation [28]. However, this site is not present in S. pombe Ctp1. Unlike the other CtIP orthologs, Ctp1 is regulated at the level of transcription by the MBF transcription factor that controls S phase gene expression. The absence of Ctp1 gene expression in the G1 phase prevents DSB resection and HR in the G1 phase of the cell cycle in fission yeast [23]. In budding yeast, where Sae2 is expressed throughout the cell cycle, CDK-dependent, post-translational regulation is required to restrict Sae2 function in DSB resection to the S and G2 phases. The activity of human CtIP in DSB resection is regulated by both CDK activity and the cell cycle-dependent oscillation of CtIP protein abundance (Figure 3) [9,73,81,82].

In addition to CDK, efficient DSB resection and subsequent ATR activation also require ATM kinase activity [40–43]. Although the contribution of ATM kinase activity to DSB resection has not been clear, our recent study indicates that ATM promotes CtIP recruitment to DSBs, suggesting that CtIP is a target of ATM in DNA end resection [18]. Thus, CtIP plays a crucial role in linking ATM activation to ATR activation after DNA damage [18,83]. Two ATM phosphorylation sites, S664 and S745, have been identified in human CtIP (Figure 4) [12]. However, these sites apparently are dispensable for CtIP recruitment to DSBs [18], suggesting that phosphorylation at other sites by ATM or another protein downstream of ATM facilitates CtIP damage relocation. One possible role of ATM kinase activity in CtIP damage recruitment is to promote conformational changes in CtIP that lead to exposure of a DNA-binding motif in a central region of CtIP (Figure 4) [18]. The CtIP DNA binding motif is required for its recruitment to DSBs, probably by directly binding to DNA at the damage sites [18]. Another possibility is that CtIP phosphorylation by ATM or a kinase downstream of ATM such as Chk2 primes CtIP for CDK-dependent phosphorylation, which facilitates CtIP interaction with the FHA domain of NBS1, and association with DNA damage sites. Future studies are needed to define the functional relationship between CDK, BRCA1, ATM and the DNA-binding activity in CtIP in the regulation of CtIP damage recruitment and its function in DNA end resection after damage recruitment.

CtIP and cancer

CtIP is an essential gene in mammalian cells, with homozygous inactivation causing embryonic lethality in mice [25]. Inactivation of just one CtIP allele predisposes mice to multiple types of cancers, particularly lymphomas, suggesting that CtIP functions as a tumor suppressor protein [25]. Mutations in CtIP have also been identified in human cancer cell lines [4]. In colorectal cancer cells...
with mutations in the mismatch repair (MMR) pathway, the A9 repeat in the middle region of the gene encoding CtIP is frequently mutated as a result of microsatellite instability, resulting in translation of a truncated CtIP protein of 357 amino acids [84]. Although no biallelic mutations of CtIP have been identified in cancers, the haploinsufficiency observed in mice suggests that mutation of a single allele of CtIP might promote genome instability and tumorigenesis. Interestingly, epigenetic inactivation of CtIP has been reported in breast cancer, and is associated with increased resistance to tamoxifen treatment [85].

Disruption of CtIP function could also be the underlying mechanism for cancer susceptibility in cells with certain BRCA1 mutations. BRCA1 mutations are associated with hereditary predisposition to breast and ovarian cancer, with the most common mutations occurring in the C-terminal BRCT tandem repeats [86]. BRCA1 forms distinct complexes in cells, owing to the mutually exclusive binding of its BRCT domains with phosphorylated proteins including CtIP, BACH1/BRIP1/FancJ and RAP80/Abraxas [74,79,80,87]. Structural studies have demonstrated that two cancer-associated mutations in the BRCT domains of BRCA1, M1775R and R1699W, directly interfere with the hydrophobic pocket to which phosphorylated CtIP binds [75,88]. Defects in the BRCA1–CtIP interaction and the DNA damage response have also been observed in HCC1937 breast cancer cells that harbor a BRCA1 mutant protein lacking a BRCT repeat, consistent with the notion that a functional CtIP–BRCA1 complex is important for tumor suppression [4,74].

Although inactivation and mutation of CtIP and BRCA1 promote cancer formation and development, these might also be exploited for targeted cancer therapy on the basis of synthetic lethality [89]. Recently, inhibitors of the base excision repair enzyme PARP1 have been used to preferentially kill BRCA1-deficient cancer cells [90,91]. This selective killing effect is believed to occur as a result of the defects in HR-mediated repair of the DSBs that are converted from the unrepaird single-strand breaks induced by PARP1 inhibition in the BRCA1-deficient cells [90,91]. Synthetic lethality also was observed between PARP1 inhibitors and mutation in other genes in the HR pathway [92], or mutation of genes such as PTEN that influence HR gene expression [93]. Given the crucial role of CtIP in HR and its intimate relationship with BRCA1, PARP1 inhibitors could be also efficacious for treatment of cancers where CtIP is mutated or epigenetically inactivated.

Concluding remarks
Recent studies on CtIP place this protein at the intersection of cell cycle control, checkpoint signaling and DNA repair. After DNA damage, CtIP receives signals from both the cell cycle kinase CDK and checkpoint kinase ATM that activate its function in DNA damage signaling and repair. CtIP is then recruited to DSB damage sites, where it interacts with the MRN complex to mediate the initiation step of DNA end resection. In so doing, CtIP drives the transition from ATM-dependent checkpoint signaling to ATR-dependent checkpoint signaling. Importantly, the engagement of CtIP-dependent DNA end resection commits cells irreversibly to repair by the HR pathway. Although much has been learned about what regulates CtIP relocalization to DNA damage sites, many questions remain as to how CtIP facilitates DSB resection after damage recruitment (Box 1). In particular, it will be important to determine whether CtIP, as with yeast Sae2, functions as an endonuclease, and if so, whether and how this activity is regulated by CDK, BRCA1 and ATM. The mechanism of cooperation between CtIP and MRN, and the functional relationship between the potential nuclease activity of CtIP and that of MRN in DSB resection also remain to be delineated. Furthermore, it remains to be investigated whether the two-step model for DSB resection proposed in budding yeast applies to higher organisms, and if it does, how the switch is made from the initial MRN-CtIP dependent endonucleolytic cleavage step to the second resection step that is potentially carried out by Exo1 and BLM-DNA2. Finally, it will be interesting to find out whether the DNA binding motif identified in CtIP is required for the transcription function of CtIP, and whether CtIP promotes transcription of additional genes that are involved in DNA damage signaling and repair. Future studies on CtIP function and regulation will provide insights into how defects in the DSB damage response predispose patients to cancer, and might help identify and treat patients that could best respond to cancer therapies that cause DNA damage or target the DNA damage checkpoint response pathway.

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