Mechanism of CRL4\textsuperscript{Cdt2}, a PCNA-dependent E3 ubiquitin ligase

Courtney G. Havens and Johannes C. Walter\textsuperscript{1}

Department of Biological Chemistry and Molecular Pharmacology, Harvard Medical School, Boston, Massachusetts 02115, USA

Eukaryotic cell cycle transitions are driven by E3 ubiquitin ligases that catalyze the ubiquitylation and destruction of specific protein targets. For example, the anaphase-promoting complex/cyclosome (APC/C) promotes the exit from mitosis via destruction of securin and mitotic cyclins, whereas CRL1\textsuperscript{Skp2} allows entry into S phase by targeting the destruction of the cyclin-dependent kinase (CDK) inhibitor p27. Recently, an E3 ubiquitin ligase called CRL4\textsuperscript{Cdt2} has been characterized, which couples proteolysis to DNA synthesis via an unusual mechanism that involves display of substrate degrons on the DNA polymerase processivity factor PCNA. Through its destruction of Cdt1, p21, and Set8, CRL4\textsuperscript{Cdt2} has emerged as a master regulator that prevents rereplication in S phase. In addition, it also targets other factors such as E2F and DNA polymerase \(\eta\). In this review, we discuss our current understanding of the molecular mechanism of substrate recognition by CRL4\textsuperscript{Cdt2} and how this E3 ligase helps to maintain genome integrity.

The regulated destruction of proteins is integral to the physiology of all eukaryotic cells. Thus, cell cycle transitions, the maintenance of genome integrity, signaling, and many other cellular processes involve controlled proteolysis. Regulated proteolysis is carried out by the ubiquitin–proteasome system. Ubiquitin is attached to proteins destined for destruction via an isopeptide bond between the C-terminal glycine of ubiquitin and one or more lysines of the target. This ubiquitin is then modified by additional ubiquitins that are connected to a lysine on the foregoing ubiquitin, thereby forming “ubiquitin chains.” Most ubiquitin chains target the substrate for destruction by the 26S proteasome. However, some proteins are mono-ubiquitylated or di ubiquitylated, while others are poly-ubiquitylated via Lys 63 chains that modulate protein function without causing destruction (Komander 2009; Ye and Rape 2009; Behrends and Harper 2011). Generally, the E3–substrate interaction involves the binding of a “substrate receptor” subunit of the E3 to a short stretch of amino acids called a “degron” motif within the substrate. The best understood degrons are six to eight amino acids long, transferable, and necessary and sufficient for binding to the substrate.

There are two major families of multiprotein E3 ubiquitin ligases in eukaryotes, which are characterized by the presence of RING or HECT domains (Ardley and Robinson 2005; Deshaies and Joazeiro 2009). The RING domain coordinates zinc ions in a protein fold that binds to an E2-conjugating enzyme. The cullin ring E3 ligases (CRLs) are the largest family of RING ubiquitin ligases. In CRLs, a cullin scaffold binds via its C terminus to the RING domain protein Rbx1, which recruits an E2 (Fig. 1A, E2 not shown). The N terminus of the cullin binds to an adaptor [e.g., Skp1], which in turn binds a substrate receptor that contacts the substrate, bringing it close to the E2 (Fig. 1A). In HECT domain ligases, the E2 enzyme transfers ubiquitin to a cysteine residue in the HECT domain, from which it is transferred to the substrate (data not shown).

Regulation of substrate recognition by E3 ubiquitin ligases

Many substrates of the ubiquitin proteasome system are not destroyed constitutively, but rather become unstable only in response to an endogenous or exogenous signal. Often, signaling leads to post-translational modification of the degron, which triggers binding to the ligase [Ravid and Hochstrasser 2008]. For example, the E3 ligase Cul1–Skp1–FBW7 [CRL1\textsuperscript{FBW7}; also known as SCF\textsuperscript{FBW7}] recognizes the Cyclin E degron by this mechanism. FBW7
contains eight WD40 repeats that fold into an eight-bladed β propeller. The bottom surface of the propeller connects to an F-box motif that interacts with Skp1. The top surface of the propeller forms a surface that interacts specifically with the doubly phosphorylated degron of Cyclin E. Interestingly, some substrates such as Cdc6 and Skp2 can be protected from destruction when their degrons are phosphorylated [Mailand and Diffley 2005; Gao et al. 2009]. Phosphorylation is not the only post-translational modification that regulates degrons. For instance, the E3 ligase CRL2^{HL} targets HIF1 [hypoxia-inducible factor-1] only when its degron motif is hydroxylated, ensuring that HIF1 protein levels increase under hypoxic conditions [van et al. 2001; Jaakkola et al. 2001; Hon et al. 2002; Min et al. 2002]. In another example, CRL1^{Fbx2} recognizes glycosylated proteins that are retrotranslocated from the ER into the cytosol and targets them for destruction [Yoshida et al. 2002; Mizushima et al. 2007]. Finally, some ubiquitin ligases interact preferentially with substrates that are sumoylated [Perry et al. 2008]. Thus, post-translational modification of substrates represents a common means to couple the activity of E3 ligases to signaling events that sense intracellular or extracellular conditions.

Notably, E3–substrate interactions are not always dependent on post-translation modification of the substrate. The anaphase-promoting complex/cyclosome [APC/C], a RING E3 ubiquitin ligase that promotes the exit from mitosis by destroying mitotic cyclins and other proteins, is itself phosphorylated. Phosphorylation of APC/C core subunits is required to allow activation of the complex and the interaction between APC/C and Cdc20 [Kraft et al. 2003], the WD40 substrate receptor that initially recruits APC/C substrates during mitotic exit [Pfleger et al. 2001]. Conversely, the other APC/C substrate receptor, Cdh1, is inactive when phosphorylated by CDKs, ensuring that it acts after APC/C-Cdc20 [Zachariae et al. 1998; Jaspersen et al. 1999; Kramer et al. 2000].

CRL1^{TIR1} [SCF^{TIR1}] is another example of a ubiquitin ligase that is modified to regulate destruction of a substrate, but in this case, the modification is noncovalent. In the absence of the plant hormone auxin, repressors block transcription of auxin-responsive factors. However, in the presence of auxin, the repressors are targeted for ubiquitin-mediated degradation. Strikingly, auxin binds directly to a pocket in the substrate receptor F-box protein TIR1, which stabilizes the interaction between CRL1^{TIR1} and its transcription repressor substrates, allowing them to be ubiquitylated and destroyed [Mockaitis and Estelle 2008; Tan and Zheng 2009].

In summary, the interaction of E3 ubiquitin ligases with their substrates is regulated in a number of different ways, most of which involve post-translation modification of the substrate or ligase, or employment of small molecule cofactors. In this review, we discuss in detail the E3 ubiquitin ligase CRL4^{Cdt2}, whose interaction with substrates depends on a novel strategy that involves display of a degron motif on a cell cycle-regulated protein scaffold, chromatin-bound PCNA.

The architecture of CRL4^{Cdt2}

CRL4 ubiquitin ligases consist of a cullin scaffold [Cul4], an adaptor protein (Ddb1), and a substrate receptor [DCAF [Ddb1- and Cul4-associated factor]] that binds directly to Ddb1. At least 20 bona fide DCAs are known in mammalian cells [Angers et al. 2006; He et al. 2006; Higa et al. 2006b; Jin et al. 2006; Higa and Zhang 2007; O’Connell and Harper 2007; Hu et al. 2008; Lee et al. 2008; McCall et al. 2008; Scrima et al. 2008; Choe et al. 2009; Jackson and Xiong 2009; Xu et al. 2010]. CRL4^{Cdt2} contains the DCAF Cdt2 [Cdc10-dependent transcript 2], also known as DCAF2, DTL, L2DTL, or RAMP, which was first discovered in fission yeast [Hofmann and Beach 1994]. The overall architecture of CRL4^{Cdt2} probably conforms to the basic modular structure established for cullin-based ligases [Fig. 1A; Zheng et al. 2002]. Like Cul1, Cul4 is an elongated α-helical protein that interacts through its C terminus with the ring finger protein Rbx1. The N terminus of Cul4 binds to Ddb1, which contains three β propellers [BPA, BPB, and BPC] that each comprise seven WD40 repeats or “blades.” BPA binds Cul4, whereas BPA and BPC form a clam-shaped structure that points toward the C terminus of Cul4. Although the structure of Cdt2 has not been solved, its sequence suggests that between residues 40 and 397 it contains a seven-bladed β propeller, as seen for many other CRL substrate receptors [data not shown]. The structure of CRL4^{Ddb2} is likely to be similar to that of CRL4^{Ddb2}, which promotes ubiquitylation of the nucleotide excision repair factor XPC in the context of UV
damage (Sugasawa et al. 2005) and whose architecture can be modeled from two previous crystal structures (Fig. 1B; Angers et al. 2006; Scrima et al. 2008). If Cdt2 is indeed structurally analogous to Ddb2 (Fig. 1B), an α helix near the “bottom” of its β propeller would bind the cleft formed by BPA and BPC, whereas the “top” would interact with substrates, positioning them for ubiquitin transfer from the E2 protein. The functions of the N-terminal and C-terminal domains of Cdt2 that reside outside the β propeller are unknown.

Cdt1 destruction by CRL4Cdt2 is coupled to DNA replication and repair via PCNA

Among eukaryotes, there are six confirmed substrates of CRL4Cdt2 (see Fig. 2, bold names), but the mechanism of ubiquitylation has been most intensively studied for the replication licensing factor Cdt1 (Cdc10-dependent transcript 1, no structural relationship with Cdt2) (Hofmann and Beach 1994). Cdt1 is required for the recruitment of the MCM2–7 helicase to origins of DNA replication in G1 (the “licensing” reaction), but it is destroyed in S phase (Nishitani et al. 2004), which ensures that each origin of DNA replication undergoes only one initiation event per cell cycle (Fig. 3; for review, see Arias and Walter 2007). Early experiments in worms showed that the destruction of Cdt1 in S phase is dependent on Cul4 (Zhong et al. 2003). Soon thereafter, experiments in mammalian cells demonstrated that Cul4 and Ddb1 promote Cdt1 destruction after DNA damage (Higa et al. 2003; Hu et al. 2004). Subsequently, multiple groups showed that Cdt2 is also required for Cdt1 destruction (Higa et al. 2006a; Jin et al. 2006; Ralph et al. 2006; Sansam et al. 2006). It is now clear that CRL4Cdt2 ubiquitylates Cdt1 in S phase and after DNA damage in all metazoans and in fission yeast. Cdt2 likely functions as the substrate receptor for CRL4Cdt2 (Abbas et al. 2008), although this remains to be formally proven. While budding yeast contains Cul4 and Ddb1-like proteins (Zaidi et al. 2008), it lacks an identifiable Cdt2 ortholog, and Cdt1 is not unstable in S phase of these cells, suggesting that budding yeast lacks CRL4Cdt2.

In a separate line of inquiry, experiments in *Xenopus* egg extracts showed that the ubiquitylation of Cdt1 during S phase occurs on chromatin and is strictly coupled to the process of DNA replication (Arias and Walter 2005). Analogously, after DNA damage, ubiquitylated Cdt1 also localizes to chromatin (Jin et al. 2006; Ishii et al. 2010; Roukos et al. 2011). Strikingly, all metazoan Cdt1 molecules contain a PCNA interaction protein motif (PIP box) at their extreme N termini (Fig. 2A,B), which binds to a hydrophobic pocket within PCNA that is composed of residues from the interdomain connector loop and the C terminus of PCNA (Fig. 4A). The PIP box is essential for Cdt1 destruction in S phase and after DNA damage (Arias and Walter 2005). Analogously, after DNA damage, ubiquitylated Cdt1 also localizes to chromatin (Jin et al. 2006; Ishii et al. 2010; Roukos et al. 2011). This binding is dependent on prior interaction of a CRL4Cdt2 substrate with PCNA on chromatin (Jin et al. 2006; Ishii et al. 2010; Roukos et al. 2011). This binding is dependent on prior interaction of a CRL4Cdt2 substrate with PCNA on chromatin (Jin et al. 2006; Ishii et al. 2010). If DNA replication or repair are inhibited upstream of PCNA loading, Cdt1 and CRL4Cdt2 do not bind to chromatin and Cdt1 is not destroyed (Arias and Walter 2005, 2006; Havens and Walter 2009; Guarino et al. 2011). Together, the data suggest a model wherein Cdt1 first uses its PIP box to dock onto PCNA that is engaged in replication or repair synthesis; subsequently, CRL4Cdt2 is recruited, whereupon Cdt1 is polyubiquitylated on chromatin and destroyed (Fig. 5).

**Figure 2.** CRL4Cdt2 degrons. (A) PIP box and PIP degron consensus. Canonical PIP box residues are shown in violet, and PIP degron-specific residues are shown in blue. The TD motif promotes high-affinity binding to PCNA, whereas the basic residue four amino acids downstream of the PIP box is required for docking of CRL4Cdt2 onto the PCNA–PIP degron complex. (B) The Cdt1 PIP degron sequence from various species. In all cases, the degron is at the extreme N terminus. Same color scheme as in A. (C) C-terminal and internal PIP degrons of various other substrates. Confirmed substrates are in bold. Asterisk denotes the C terminus.

**CRL4Cdt2 recognizes a ‘PIP degron’**

The discovery that Cdt1 destruction is dependent on the binding of its PIP box to PCNA raised the question of why most PIP box-containing proteins, such as DNA polymerases and DNA ligases, are not targeted by CRL4Cdt2. The answer emerged from a comparison between known CRL4Cdt2 substrates and canonical PIP box-binding proteins,
which showed that the former contain a specialized PIP box dubbed a “PIP degron” (Havens and Walter 2009; Michishita et al. 2011). The PIP degron contains at least two essential elements [Fig. 2A]. As discussed above, the first is a PIP box, which is crucial for binding to chromatin-bound PCNA (PCNA<sup>Ctld</sup>). Interestingly, most CRL<sup>Ctld</sup> substrates contain a TD motif at positions 5 and 6 of the PIP box [Fig. 2A], which forms a loop that protrudes off the surface of PCNA [Fig. 4A]. The TD motif is not usually considered part of the PIP consensus (Moldovan et al. 2007). However, in combination with the classic PIP box residues [Fig. 2A, pink residues], the TD motif confers unusually high-affinity PCNA binding (Nakanishi et al. 1995; Warbrick et al. 1995; Chuang et al. 2005; Havens and Walter 2009; Michishita et al. 2011). Thus, effective CRL<sup>Ctld</sup>-dependent destruction of substrates requires that they bind with high affinity to PCNA, perhaps to ensure processive ubiquitylation. The high-affinity binding might also ensure that CRL<sup>Ctld</sup> substrates are destroyed before other PCNA-dependent processes take place. The few CRL<sup>Ctld</sup> substrates that lack the TD motif [e.g., Dm Cdt1, Sp Cdt1, Ce polH, Dm E2F] [Fig. 2C] likely compensate with other residues to achieve efficient PCNA binding. It is interesting to note that most PIP box proteins that are not CRL<sup>Ctld</sup> substrates lack a TD motif and thus bind to PCNA with suboptimal affinity (data not shown). Indeed, addition of the TD motif to the PIP boxes of Fen1 or DNA ligase dramatically increases their affinities for PCNA (Havens and Walter 2009; Michishita et al. 2011). Suboptimal binding of canonical PIP box proteins to PCNA may be important to maintain appropriate exchange of PCNA binding proteins at the replication fork and during DNA repair (Fridman et al. 2010). In contrast, high-affinity binding of CRL<sup>Ctld</sup> substrates to PCNA is clearly not deleterious, likely because they are destroyed.

The second essential element of the PIP degron, which is conserved in all known CRL<sup>Ctld</sup> substrates, is a basic residue four amino acids downstream from the PIP box [Fig. 2A]. This residue, called “B+4,” is surface-exposed in the p21–PCNA cocrystal structure (Fig. 4B; Gulbis et al. 1996). Importantly, mutation of B+4 abolishes docking of CRL<sup>Ctld</sup> onto the PCNA–Cdt1 complex and inhibits Cdt1 destruction, but it has no effect on the binding of Cdt1 to PCNA (Havens and Walter 2009; Michishita et al. 2011). Thus, B+4 likely makes specific contact with acidic residues on the top face of Cdt2 [Fig. 4B, dashed line].

The PIP degron is portable, since canonical PIP box proteins such as Fen1 or DNA ligase can be converted into a CRL<sup>Ctld</sup> substrate by introducing the TD and B+4 motifs into its PIP box [Havens and Walter 2009; Michishita et al. 2011]. In addition, an ~25-amino-acid peptide containing the PIP box of human Cdt1 is sufficient to confer CRL<sup>Ctld</sup>-mediated destruction on GST [Nishitani et al. 2006; Senga et al. 2006]. Together, these results indicate that the PIP degron, which has the consensus Q-x-x-W-T-D-y-x-x-B [Fig. 2A], is necessary and sufficient to interact with CRL<sup>Ctld</sup> in the context of chromatin-bound PCNA and to promote protein destruction.

Other amino acids that are closely linked to the PIP box play roles in some CRL<sup>Ctld</sup> targets. For example, several substrates including the CDK inhibitors (CKIs) contain basic residues at the +3 and +5 positions [Fig. 2C], which interact with acidic residues within PCNA in the p21–PCNA crystal structure (Gulbis et al. 1996). Indeed, mutation of B+5 in Xenopus Cdt1 and human p21 reduced destruction due to impaired PCNA binding.
Mechanisms that couple CRL4\textsuperscript{Cdt2} activity to S phase and DNA damage

CRL4\textsuperscript{Cdt2} mediated proteolysis occurs only in S phase and after DNA damage because this ubiquitin ligase promotes ubiquitylation of substrates only on the DNA-bound fraction of PCNA [You et al. 2002, Arias and Walter 2005, 2006; Chuang and Yew 2005; Havens and Walter 2009; Guarino et al. 2011; Jorgensen et al. 2011]. What is the mechanistic basis for the coupling to PCNA\textsuperscript{DNA}, or, why does soluble PCNA not promote CRL4\textsuperscript{Cdt2} activity? In Xenopus egg extracts, endogenous Cdt1 does not interact with soluble PCNA [PCNA\textsuperscript{soluble}] [Fig. 5, double arrow #1] but only with PCNA that is bound to PCNA\textsuperscript{DNA} [Havens and Walter 2009]. As a result, Cdt1 and CRL4\textsuperscript{Cdt2} are only able to interact on PCNA\textsuperscript{DNA} [Fig. 5]. How the selective binding of Cdt1 to PCNA\textsuperscript{DNA} comes about is currently unclear. However, it does not require CDK, ATR, ATM, or DNA-PK, since none of these protein kinases are required for Cdt1 destruction [Higa et al. 2003; Arias and Walter 2006; Jin et al. 2006; Ralph et al. 2006]. Interestingly, an artificial PIP degron [Fen1 protein mutated to contain the TD and B\textsuperscript{4} motifs] was found to bind to PCNA\textsuperscript{soluble}. However, this interaction does not support CRL4\textsuperscript{Cdt2} recruitment, demonstrating that CRL4\textsuperscript{Cdt2} senses whether DNA is present in the PCNA–PIP degron complex [Havens and Walter 2009]. The requirement for DNA is not understood, but it could reflect direct interactions of CRL4\textsuperscript{Cdt2} with DNA or the involvement of other DNA-bound proteins. In summary, CRL4\textsuperscript{Cdt2} activity is coupled to PCNA\textsuperscript{DNA} by at least two mechanisms. First, some substrates, like Cdt1, bind only to PCNA\textsuperscript{DNA}. Second, a substrate–PCNA\textsuperscript{soluble} complex does not serve as a docking site for CRL4\textsuperscript{Cdt2}. In theory, steps even further downstream, such ubiquitin transfer to the substrate itself, might also be DNA-dependent, but this remains to be examined.

Mechanism of PCNA–PIP degron–CRL4\textsuperscript{Cdt2} complex assembly

The molecular architecture of the ternary PCNA\textsuperscript{DNA}–PIP degron–CRL4\textsuperscript{Cdt2} complex and the role of PCNA in promoting ubiquitin transfer are not well understood. As noted above, Xenopus Cdt1 does not interact with PCNA\textsuperscript{soluble} or CRL4\textsuperscript{Cdt2} in the absence of DNA [Fig. 5, gray double-headed arrows #1 and #2]. Instead, Cdt1 docks onto PCNA\textsuperscript{DNA}, and its B\textsuperscript{4} residue is then instrumental
in recruiting CRL4Cdt2 [Fig. 5]. Recently, we identified residues in the interdomain connector loop of PCNA that are essential for recruitment of CRL4Cdt2 to the PIP degron–PCNA complex but do not affect binding of the PIP degron to PCNA [CG Havens and JC Walter, unpubl.]. This argues that CRL4Cdt2 contacts residues not only in the PIP degron but also in PCNA. As such, CRL4Cdt2 effectively recognizes a composite interface formed by two distinct polypeptides: PCNA and the PIP degron of the substrate. Since soluble Cdt1 and CRL4Cdt2 do not interact, PCNA DNA functions as a matchmaker that brings these two components together.

Some studies have reported that certain substrates (like p21) interact with CRL4Cdt2 in the absence of PCNA [Fig. 5, gray double arrow #2; Hu et al. 2004; Liu et al. 2005; Higa et al. 2006b; Kim and Kipreos 2007; Abbas et al. 2008, 2010; Kim et al. 2008]. Since destruction of all substrates requires PCNA DNA, this result would imply that PCNA DNA acts on the preformed substrate–ligase complex to facilitate ubiquitin transfer. For example, PCNA DNA might induce an allosteric change in the substrate or CRL4Cdt2 that potentiates ubiquitin transfer. However, a caveat in virtually all experiments showing substrate–CRL4Cdt2 interaction in the absence of PCNA is that the binding is detected when one or both components are overexpressed. An alternative interpretation of the data is that the detected interactions contribute to the stability of the ternary complex, but only in the context of PCNA DNA.

Recent biochemical experiments suggest that the C-terminal half of Xenopus Cdt2 interacts directly with PCNA, independently of substrate [Fig. 5, gray double arrow #3; Kim et al. 2010], and imaging of Cdt2 in mammalian cells is consistent with this idea [Roukos et al. 2011]. Although deletion of the C-terminal 310 amino acids of Cdt2 disrupted the interaction with PCNA and inhibited CRL4Cdt2 function, this deletion likely caused large-scale disruption of the Cdt2 structure. In the future, targeted disruption of the CRL4Cdt2–PCNA interaction will be required to examine its relevance for CRL4Cdt2 function.

CRL4Cdt2 substrate ubiquitylation and beyond

The residues that are ubiquitylated by CRL4Cdt2 have been studied in three substrates: Cdt1, p21, and Xic1. A fragment of human Cdt1 containing amino acids 1–98 in which all the lysines were mutated to arginines was still destroyed, albeit slowly [Senga et al. 2006]. These data suggest that in the absence of lysine acceptor sites for ubiquitin, the N-terminal methionine could be used. In contrast, when all of the lysines in p21 were mutated to arginines, p21 was stabilized [Bendjennat et al. 2003; Stuart and Wang 2009]. In Xic1, mutation of five lysine residues just downstream from the PIP box reduced polyubiquitylation and destruction [Kim et al. 2010]. Given the ubiquitylation data from Xic1 and the fact that short peptides of CRL4Cdt2 substrates containing the PIP boxes were destroyed [You et al. 2002; Chuang and Yew 2005; Nishitani et al. 2006; Senga et al. 2006], it is likely that multiple lysines upstream of or downstream from the PIP box can function as ubiquitin acceptors. Determining the preferred sites of CRL4Cdt2-dependent ubiquitylation requires more work.

Recently, it was discovered that CRL4Cdt2 uses two separate E2 ubiquitin-conjugating enzymes to target different substrates [Shibata et al. 2011]. Thus, it cooperates with UBE2G to ubiquitylate Cdt1 and UBC8H8 to ubiquitylate Set8 and p21. E2s can regulate the specificity for ubiquitin nucleation on the substrate and chain formation on the preceding ubiquitins by recognizing the surface of ubiquitin or the substrate near the lysine to be ubiquitylated [Jin et al. 2008; Ye and Rape 2009; Rodrigo-Brenni et al. 2010; Saha et al. 2011]. We speculate that the location of the PIP box [N- vs. C-terminal] and possible contributions from surrounding residues in substrates might necessitate the use of two different E2 enzymes.

An important question is how ubiquitylated Cdt1 is removed from PCNA DNA. A recent genome-wide siRNA screen identified the ATPase p97 (also known as VCP or Cdc48p) and its cofactor, UFD1, as being essential for Cdt1 destruction [Raman et al. 2011]. Cells treated with siRNA against p97 or UFD1 accumulated polyubiquitylated Cdt1 and Set8 on chromatin. Therefore, p97-UFD1 likely functions downstream from CRL4Cdt2 ubiquitylation to extract substrates from the chromatin so they can be degraded by the proteasome.

The biology of CRL4Cdt2 targets

In addition to Cdt1, there are five confirmed substrates of CRL4Cdt2 [Fig. 2]. Below, we discuss these substrates, focusing on the biological importance of their destruction in S phase and after DNA damage.

Sed1

The first confirmed substrate of CRL4Cdt2 was the Schizosaccharomyces pombe protein Sed1 [Liu et al. 2003, 2005; Holmberg et al. 2005], an inhibitor of the ribonuclease reductase [RNR] enzyme that catalyzes the synthesis of dNTPs. Sed1 inhibits RNR through at least two mechanisms [Nestoras et al. 2010], and it is destroyed in S phase and after DNA damage [Liu et al. 2003]. Importantly, mutations in CRL4Cdt2 induce mutagenesis, cause slow growth, and prevent double-strand break repair in S. pombe, and these defects are partially or completely alleviated if Sed1 is also deleted [Liu et al. 2005; Moss et al. 2010]. Thus, a major function of CRL4Cdt2 in S. pombe is the elimination of Sed1 in S phase and after DNA damage so that dNTPs are available for replicative and repair DNA synthesis. Although Sed1 contains a sequence that conforms to the PIP degron consensus [Fig. 2C], its role in mediating the protein’s destruction has not been reported. Interestingly, S-phase-dependent destruction of Sed1 does not require checkpoint signaling; however, damage-dependent Sed1 destruction does require the checkpoint [Liu et al. 2003]. This is because transcriptional up-regulation of Cdt2 after DNA damage
involves checkpoint kinases [Liu et al. 2005], but the underlying mechanism is unknown. It seems that this transcriptional regulation of Cdt2 function is unique to S. pombe.

E2F

Most metazoan cells express two or more E2F family transcription factors. The “activator E2Fs” promote transcription of genes in G1 whose expression is essential to enter S phase. Activator E2F is negatively regulated in early G1 phase by the retinoblastoma gene product Rb. In addition, “repressor E2Fs” inhibit the activator E2F by competing for its binding partner, Dp, which is essential for E2F DNA binding. In Drosophila, the activator E2F, E2f1, is antagonized by Rb and the repressor E2f2, E2f12, but neither inhibitory pathway is essential for development, suggesting the existence of additional mechanisms that restrain E2f1 activity. Indeed, it was noted that E2f1 is specifically destroyed in S phase [Asano et al. 1996]. Duronio and colleagues [Shibutani et al. 2008] then discovered that this process requires a PIP degron in E2f1, as well as components of CRL4Cdt2. Interestingly, CRL4\(^{Cdt2}\)-dependent E2f1 destruction also requires Dp, suggesting that assembly of the PCNA–E2f1–CRL4Cdt2 complex might require DNA binding by E2f1–Dp. This could help link destruction to the DNA-bound form of PCNA. Compared with E2f1, ectopic expression of E2f1\(^{APIP}\) caused increased expression of an E2f1 target gene, a reduced G1 population, developmental defects, and apoptosis, indicating that E2f1\(^{APIP}\) functions as a gain-of-function allele in vivo. Given that E2fs in other species do not contain recognizable PIP degrons, it appears unlikely that CRL4\(^{Cdt2}\)-dependent destruction of activator E2Fs is conserved in other species. Instead, mammalian cells appear to contain multiple pathways to restrain E2F activity once cells have entered S phase [Dimova and Dyson 2005]. At present, it has not been examined whether Drosophila E2F is destroyed after DNA damage.

DNA polymerase (pol) \(\eta\)

Pol \(\eta\) is a translesion DNA polymerase that specializes in the bypass of UV-induced pyrimidine dimers. Interestingly, Caenorhabditis elegans pol \(\eta\) suppresses the DNA damage-induced cell cycle checkpoint during early embryogenesis, presumably by promoting by-pass of lesions before they lead to ATR checkpoint activation, a function that is shared by a sumo E3 ligase called gei-17 [Kim and Michael 2008]. The checkpoint is likely suppressed because embryogenesis is extremely sensitive to perturbations of cell cycle progression. Strikingly, in embryos lacking gei-17, DNA damage induces the efficient destruction of pol \(\eta\) by CRL4\(^{Cdt2}\)-dependent on a PIP degron within pol \(\eta\) [Fig. 2]. Gei-17 protects pol \(\eta\) from CRL4\(^{Cdt2}\) modification via direct, DNA damage-induced sumoylation. Importantly, when wild-type embryos are treated with high levels of DNA-damaging agents, CRL4\(^{Cdt2}\)-dependent pol \(\eta\) destruction become detectable. Based on these results, it was proposed that in early C. elegans embryos, Gei-17 normally sumoylates pol \(\eta\) at sites of DNA damage and thereby protects it from CRL4\(^{Cdt2}\)-mediated destruction. After lesion bypass, sumoylation is reversed, allowing targeting and subsequent eviction of the error-prone pol \(\eta\) by CRL4\(^{Cdt2}\). In the future, it will be important to test this interesting model explicitly.

Cdt1

As mentioned above, CRL4\(^{Cdt2}\) targets Cdt1 for destruction in S phase and after DNA damage. The function of the S-phase destruction is clearly to prevent rereplication. This is evident from the fact that overexpression of Cdt1 is sufficient to promote DNA rereplication in various systems [Vaziri et al. 2003; Arias and Walter 2005; Li and Blow 2005; Takeda et al. 2005; Yoshida et al. 2005; Kerns et al. 2007]. Moreover, Cdt1 lacking a PIP box is more potent than wild-type Cdt1 in promoting rereplication [Arias and Walter 2006]. Interestingly, expression of non-degradable Cdt1 in fly embryonic epidermal cells caused S-phase defects and failure of cells to enter mitosis, but in follicle cell endocycles, Geminin was able to restrain Cdt1 activity [Lee et al. 2010]. This result suggests a differential requirement for CRL4\(^{Cdt2}\) in maintaining genome stability in different tissues.

When asynchronously growing cells are UV-irradiated, the G1-phase population of Cdt1 is destroyed within minutes via CRL4\(^{Cdt2}\) [Higa et al. 2003]. It was proposed that this destruction represents a checkpoint to prevent S-phase entry when cells experience genotoxic stress. However, MCM2–7 complexes are already loaded onto DNA in telophase and Cdt1 is not required for initiation from loaded MCM2–7 complexes. Therefore, such a DNA damage-induced initiation checkpoint would only be effective if MCM complexes dissociate after Cdt1 destruction. Dissociation may not have to be complete to block S-phase entry given the proposal that cells contain a licensing checkpoint that blocks S phase when there are insufficient levels of chromatin-bound MCM2–7 complexes [Machida et al. 2005; Teer et al. 2006; Liu et al. 2009]. An alternative role for DNA damage-induced Cdt1 destruction is in the prevention of rereplication in G2-phase cells [Arias and Walter 2007]. Thus, when G2 cells experience replicative stress, CDKs are inhibited to block cell cycle progression. However, this likely removes an important barrier against rereplication. To compensate, cells might replace the CDK-dependent CRL1\(^{Skp2}\)-mediated destruction of Cdt1 that occurs in G2 with CRL4\(^{Cdt2}\)-mediated destruction, which is normally turned off at this time in the cell cycle [Nishitani et al. 2006]. Unfortunately, there is no established way to separate the function of CRL4\(^{Cdt2}\)-mediated destruction in S phase and after DNA damage, making it difficult to specifically address the function of UV-dependent Cdt1 destruction.

CKIs

Mammalian cells express three major CKIs, p21\(^{Cip1}\), p27\(^{Kip1}\), and p57\(^{Kip2}\), whose abundance during the cell cycle is carefully regulated. Thus, p21 is expressed in G1
and G2, but its level drops dramatically in S phase and after UV irradiation (Amador et al. 2007; Abbas et al. 2008; Kim et al. 2008). While several studies showed that p21 levels are negatively regulated by the APC/C in mitosis and by CRL1Skp2 (Yu et al. 1998; Bornstein et al. 2003; Wang et al. 2005; Amador et al. 2007), the disappearance of p21 in S phase remained unexplained. Moreover, conflicting theories accounted for the destruction of p21 after UV irradiation (for review, see Abbas et al. 2008). In 2008, four studies reported that human p21 is a CRL4Cdt2 target (Abbas et al. 2008; Kim et al. 2008; Nishitani et al. 2008; Stuart and Wang 2009). Thus, knockdown of Cul4, Ddb1, Cdt2, or PCNA inhibits p21 destruction in S phase and after damage, as does mutation of p21’s PIP box, which conforms perfectly to the PIP degron consensus (Fig. 2C). Moreover, purified CRL4Cdt2 is able to ubiquitylate p21 in vitro and in vivo. The destruction of CKIs by CRL4Cdt2 is highly conserved, as this phenomenon is observed for Xic1 [a Xenopus CKI] in frogs [You et al. 2002; Chuang and Yew 2005; Chuang et al. 2005; Kim et al. 2010], and the CKI in worms (CKI-1) [Kim et al. 2007, 2008]. Although it has not been confirmed, the Drosophila CKI Dacapo [Dap] is also a likely target, since it contains a nearly perfect PIP degron (Fig. 2C). While it appears that CRL4Cdt2 is solely responsible for the UV-induced destruction of p21, the relative roles of CRL4Cdt2 and CRL1Skp2 in the S-phase destruction of p21 remains to be fully resolved, although both ligases appear to make significant contributions (Abbas et al. 2008; Kim et al. 2008; Nishitani et al. 2008; Stuart and Wang 2009).

The first hints of why CKIs are targeted for destruction in S phase came from studies in worms. Kipreos and colleagues [Zhong et al. 2003; Kim and Kipreos 2007] showed that mutations in CRL4Cdt2 cause accumulation of Cdt1 and CKI-1, as well as massive rereplication. Importantly, this phenotype is largely reversed when CKI-1 or Cdt1 are co-silenced, suggesting that the destruction of either factor in S phase is sufficient to prevent rereplication. They then showed that CKI-1 exerts its effect through the licensing protein Cdc6. Thus, previous experiments had shown that Cdc6 is phosphorylated by Cdk2 in S phase, which leads to its nuclear export, but the relevance of this pathway in controlling replication had been unclear (for review, see Arias and Walter 2007). Kim et al. [2007] revealed that destruction of CKI-1 is required for the export of Cdc6, presumably because persistent CKI-1 inhibits Cdk2, blocking Cdc6 phosphorylation and export. Expressing nondegradable Cdt1 together with nonexportable Cdc6 was sufficient to cause rereplication [Kim et al. 2007]. These experiments support the model that CRL4Cdt2 inhibits licensing by two largely redundant means: (1) destruction of the licensing factor Cdt1 and (2) destruction of CKI-1, which maintains CDK2 activity during S phase, thus promoting the nuclear export of Cdc6 (Fig. 3). The CRL4Cdt2-dependent destruction of CKIs in S phase has been confirmed in mammalian cells [Kim et al. 2008; Nishitani et al. 2008], where it also prevents rereplication by promoting the nuclear export of Cdc6 [Kim et al. 2008].

The reason for p21 destruction after UV damage is not fully understood. One possibility is that p21 must be destroyed to alleviate its inhibition of PCNA so that PCNA can participate in repair. It is clear in vitro that p21 can inhibit DNA replication and repair through direct binding to PCNA via the PIP box in p21 [Chen et al. 1995; Cooper et al. 1999; Bendjennat et al. 2003]. However, what is unclear is whether this occurs in vivo. Some laboratories report inhibition of DNA repair synthesis or nucleotide excision repair in cultured cells upon overexpression of the p21 PIP box (Dayrol et al. 1998; Cooper et al. 1999; Bendjennat et al. 2003; Cazzalini et al. 2003), while others show no effect of p21 on nucleotide excision repair (Bates et al. 1998; Medema et al. 1998; Niculescu et al. 1998; Adimoolam et al. 2001; Soria et al. 2008). Other evidence indicates that the PIP box region of p21 can block pol η and PCNA foci from forming and therefore negatively regulates translesion DNA synthesis (Soria et al. 2008). Thus, although the CRL4Cdt2-dependent destruction of p21 after DNA damage is now firmly established, further work is needed to determine the precise role of this proteolysis event in cell cycle progression and DNA repair.

Interestingly, ATR is involved in DNA damage-induced p21 destruction through activation of GSK8, which phosphorylates p21 on S114 (Bendjennat et al. 2003; Lee et al. 2007). While a phospho-mimetic substitution of S114 enhanced p21 ubiquitylation by CRL4Cdt2 in vitro (Abbas et al. 2008), this phosphorylation event has so far not been directly linked to CRL4Cdt2 function in vivo. If CRL4Cdt2-dependent p21 destruction requires substrate phosphorylation by ATR, it will represent an interesting exception, since Cdt1 and Set8 destruction are independent of checkpoint signaling [Higa et al. 2003; Arias and Walter 2006; Ralph et al. 2006; Centore et al. 2010].

Set8

Set8 [PR-Set7 or KMT5A] is the sole histone methyltransferase that monomethylates histone H4 on Lys 20. In the absence of Set8, cells arrest in G2 and fail to condense their chromosomes [Houston et al. 2008]. Set8 expression is normally low in G1, ceases just before S-phase entry, remains absent in S phase, and then rises to high levels in the G2 phase and in mitosis [Rice et al. 2002; Yin et al. 2008; Oda et al. 2010]. Consistent with this expression profile of Set8, H4K20me1 levels are also low in S phase and rise late in G2/M [Rice et al. 2002]. The Set8 expression profile appears to result from the action of at least three distinct E3 ubiquitin ligases [for review, see Wu and Rice 2011]. Thus, APC/C_1Cdt1 targets Set8 during G1 [ Wu et al. 2010], and CRL1Skp3 further reduces its levels in late G1 [Yin et al. 2008]. The absence of Set8 in S phase and after DNA damage is due to the action of CRL4Cdt2 [Abbas et al. 2010; Centore et al. 2010; Oda et al. 2010; Tardat et al. 2010; Jorgensen et al. 2011], which recognizes a highly conserved PIP degron in Set8.

To determine why Set8 is eliminated in S phase by CRL4Cdt2, several groups overexpressed a PIP box mutant of Set8 (Set8APPI) that cannot be destroyed in S phase. Set8APPI expression caused elevated H4K20me1 in S phase, DNA damage, and checkpoint activation, as well
as G2 arrest and loss of cell proliferation [Abbas et al. 2010; Centore et al. 2010; Tardat et al. 2010; Jorgensen et al. 2011]. Various explanations for these phenotypes were put forward. Two reports observed that Set8$^{\Delta\text{PIP}}$ caused premature chromatin compaction in S phase, which was attributed to abnormal H4 methylation, and this compacted chromatin structure was proposed to underlie the observed DNA damage and G2 arrest [Centore et al. 2010; Jorgensen et al. 2011]. In contrast, another study found that Set8$^{\Delta\text{PIP}}$ expression causes loss of histone gene expression and chromatin decondensation, and these defects were invoked to explain the effect of Set8 on cell cycle progression [Abbas et al. 2010]. Interestingly, two groups also observed that Set8$^{\Delta\text{PIP}}$ caused substantial rereplication, as seen by the accumulation of cells with >4N DNA content [Abbas et al. 2010; Tardat et al. 2010]. Notably, Tardat et al. [2010] showed that Set8 binds to origins of replication in G1 and promotes local H4 monomethylation. Furthermore, silencing of Set8 inhibited MCM2–7 recruitment to origins. Thus, Set8 might be a bona fide licensing factor, which must be down-regulated in S phase to prevent rereplication. In this view, the primary defect of Set8$^{\Delta\text{PIP}}$ expression is rereplication, which is known to cause DNA damage and G2 arrest [Hook et al. 2007]. A potential complication with all of the above experiments that might explain some of the variability observed is that Set8$^{\Delta\text{PIP}}$ was expressed ectopically, in many cases from very strong promoters. Ideally, Set8$^{\Delta\text{PIP}}$ should be knocked into the endogenous Set8 locus and the resulting phenotype examined.

Like Spd1, E2F, Cdt1, and p21, Set8 is also targeted for destruction by CRL4$^{\text{Cdt2}}$ after DNA damage [Abbas et al. 2010; Centore et al. 2010; Oda et al. 2010; Jorgensen et al. 2011]. Interestingly, Set8 is proposed to play a positive role in the DNA damage response, as Set8 localizes to laser-induced DNA double-stranded breaks in a PCNA-dependent manner, and its catalytic activity is required for the recruitment of 53BP1 to the breaks [Oda et al. 2010]. This raises the question of why Set8 might be destroyed after damage. Notably, Set8 destruction occurs well after Set8 and 53BP1 are recruited to damage foci [Oda et al. 2010]. Thus, Set8 proteolysis might contribute to the completion of DNA repair and/or help turn off DNA damage signaling, but this model would have to be tested explicitly. It should be noted that for some CRL4$^{\text{Cdt2}}$ substrates that are destroyed in S phase and after DNA damage, only the S phase or DNA damage pathway may have functional consequences.

The role of Cdt2 in cell physiology

The disruption of Cdt2 expression in various systems has shed light on the role of CRL4$^{\text{Cdt2}}$ in cell physiology. Homozygous Cdt2$^{+/−}$ mouse embryos die at the two- to four-cell stage with an abnormal nuclear morphology whose cause is unknown [Liu et al. 2007]. This observation shows that Cdt2 is essential for cell proliferation and development, possibly due to critical roles in chromosome duplication. In tissue culture cells, silencing of Cdt2 via siRNA leads to G2 arrest, rereplication, and centrosome amplification [Jin et al. 2006; Sansam et al. 2006; Kim et al. 2008]. The G2 arrest and rereplication defect was also seen in fish embryos carrying a Cdt2 mutation [Sansam et al. 2006]. Since G2 arrest and DNA damage are observed during rereplication [Hook et al. 2007], the primary cause of these defects may be rereplication. This interpretation is bolstered by the fact that all three vertebrate CRL4$^{\text{Cdt2}}$ substrates identified (Cdt1, CKIs, and Set8) are implicated in preventing rereplication [see above]. Importantly, co-silencing of p21 with Cdt2 completely suppresses rereplication [Kim et al. 2008], likely because p21 inhibition allows Cdk2-mediated export of the licensing factor Cdc6 [see above]. Therefore, the destruction of p21 in S phase is sufficient to suppress measurable rereplication. Similarly, co-silencing of Cdt1 suppresses some of the phenotypes exhibited by Cdt2 mutant fish embryos [Sansam et al. 2006]. It will be interesting to determine the effect of co-silencing Set8 with Cdt2, as this will test the idea that CRL4$^{\text{Cdt2}}$-mediated destruction of p21, Cdt1, or Set8 in S phase is sufficient to block rereplication. In summary, by targeting these three distinct licensing activities, vertebrate CRL4$^{\text{Cdt2}}$ functions as a master regulator of rereplication. Importantly, as illustrated by E2F and DNA pol η, CRL4$^{\text{Cdt2}}$ likely also has important roles outside the suppression of rereplication.

Interestingly, aspects of CRL4$^{\text{Cdt2}}$ inhibition are nucopied by MLN4924, a new small molecule that inhibits the CRL NEDD8-activating enzyme [NAE] [Soucy et al. 2009]. Conjugation of the ubiquitin-like protein NEDD8 is required to activate CRL ligases [Duda et al. 2008; Deshaies et al. 2010]. Thus, inhibiting the NAE by MLN4924 prevents destruction of numerous CRL substrates involved in cell proliferation and cancer pathways, including Cdt1, p21, Cyclin D, and Cyclin E [Soucy et al. 2009; Lin et al. 2010; Milhollen et al. 2011]. Strikingly, treatment of cancer cells with MLN4924 triggers rereplication, DNA damage, G2 arrest, and apoptosis. However, when the cells were also treated with siRNA against Cdt1, rereplication is prevented [Lin et al. 2010; Milhollen et al. 2011]. MLN4924 is currently in clinical trials as an anti-cancer agent. In the future, it will be interesting to determine whether the stabilization of other CRL4$^{\text{Cdt2}}$ targets, principally p21 and Set8, contributes to the rereplication and anti-proliferative effects of MLN4924.

New substrates of CRL4$^{\text{Cdt2}}$

To fully understand the biology and mechanism of CRL4$^{\text{Cdt2}}$, it will be important to determine all the substrates that are regulated by this E3 ubiquitin ligase. In addition to the six proteins discussed in the previous section that are confirmed targets of CRL4$^{\text{Cdt2}}$, there are a few other likely substrates.

S. pombe Epe1 is an anti-silencing factor that contains a catalytically inactive JmjC-type demethylase domain. Epe1 is concentrated at the boundaries between heterochromatin and euchromatin and prevents the ectopic spread of heterochromatin [Braun et al. 2011]. Interestingly, CRL4$^{\text{Cdt2}}$ is required to prevent the accumulation...
of Epe1 in the body of heterochromatin and to maintain gene silencing. In the absence of CRL4<sup>Cdt2</sup>, the turnover of Epe1 decreased and its steady-state levels increased, including during S phase. Epe1 ubiquitylation also depended on Ddb1 (Cdt2 was not tested), and a two-hybrid interaction was detected between Epe1 and Cdt2. While these observations provide strong evidence that Epe1 is a CRL4<sup>Cdt2</sup> substrate, it will be important to identify the PIP degron in Epe1 and confirm that Epe1 proteolysis depends on PCNA. Interestingly, Epe1 contains two QxxL PIP boxes [Fig. 2C]. While they lack a B+4 residue, they both contain lysines at the +2 position, which is also observed for one S. pombe Cdt1 PIP box and for a putative Spd1 PIP box (Fig. 2C). We speculate that in fission yeast, CRL4<sup>Cdt2</sup> recognizes a B+2 and/or B+4 residue. If Epe1 is indeed targeted by the canonical PCNA and CRL4<sup>Cdt2</sup> pathway, it will be interesting to determine how Epe1 destruction is confined to heterochromatic regions.

In zebrafish and humans, deletion of Cdt2 causes failure of the G2/M checkpoint after ionizing radiation. In fish, this defect is not rescued when Cdt1 is also silenced (Sansam et al. 2006). These data raise the interesting possibility that Cdt2 is directly involved in initiating G2/M arrest, or that a protein other than Cdt1 must be destroyed by CRL4<sup>Cdt2</sup> to activate the checkpoint. Since it is not clear how destruction of any known CRL4<sup>Cdt2</sup> substrates would accomplish this function, there might exist a new CRL4<sup>Cdt2</sup> target that participates in regulation of the G2/M checkpoint.

Interestingly, when Cdt2 is silenced in mammalian tissue culture cells, the basal level of PCNA monoubiquitylation at K164 that is normally seen in the absence of DNA damage is diminished, and PCNA-dependent translesion DNA synthesis is reduced [Terai et al. 2010]. One interpretation of these data is that PCNA is ubiquitylated by CRL4<sup>Cdt2</sup>, which has docked onto PCNA via a bona fide PIP degron substrate. However, based on biochemical experiments, CRL4<sup>Cdt2</sup> might also recognize PCNA directly [Terai et al. 2010]. If this is the case, it raises the intriguing possibility that there exists a class of CRL4<sup>Cdt2</sup> substrates that lack PIP degrons.

One way to find new substrates of CRL4<sup>Cdt2</sup> is to search the database for matches to the PIP degron. Indeed, two groups used this approach to identify Set8 (Abbas et al. 2010; Li et al. 2002), which contains a PIP degron. It therefore seems likely that numerous proteins with nonideal PIP degrons are targeted for destruction by CRL4<sup>Cdt2</sup>. One potential approach to identify these is to isolate CRL4<sup>Cdt2</sup>, interacting proteins. However, this method suffers from the fact that key substrates like Cdt1 appear to bind CRL4<sup>Cdt2</sup> only in the context of PCNA DNA and are therefore unlikely to be isolated in this manner. Therefore, identification of new CRL4<sup>Cdt2</sup> substrates will likely require more sensitive methods (Yen and Elledge 2008; Yen et al. 2008).

**Regulation of CRL4<sup>Cdt2</sup> function by post-translational modifications**

An important question is whether any post-translational modifications are required for CRL4<sup>Cdt2</sup> function. While the destruction of Cdt1 and Set8 does not depend on the checkpoint protein kinases ATR and ATM [Higa et al. 2003; Arias and Walter 2006; Centore et al. 2010], p21 ubiquitylation and destruction after UV irradiation requires ATR-dependent phosphorylation on Ser 114 [Bendjennat et al. 2003; Lee et al. 2007; Abbas et al. 2008] for reasons that are currently unknown. Cdt2 is another potential target of modification, as Cdt2 is phosphorylated in S phase, and a hyperphosphorylated form of Cdt2 is enriched on chromatin after UV irradiation [Ishii et al. 2010]. However, the functional significance of this phosphorylation, as well as the relevant protein kinase(s), are unknown. Finally, chromatin-bound PCNA is phosphorylated on Tyr 211 [Wang et al. 2006] and ubiquitylated/SUMOylated on K164 [Hohe et al. 2002]. At present, there is no evidence that these modifications are required for CRL4<sup>Cdt2</sup> function. In summary, with the exception of p21 modification, there is no compelling evidence so far that other substrates, CRL4<sup>Cdt2</sup>, or PCNA must be modified to support this proteolysis pathway, but this picture could change upon further investigation.

There are a few instances in which post-translational modifications function to inhibit the CRL4<sup>Cdt2</sup> pathway. Thus, p21 phosphorylation on Thr 145 or Ser 146 within the PIP box (positions 2 and 3) stabilizes the protein after UV irradiation, and this effect is explained by the fact that this modification disrupts the interaction with PCNA [Scott et al. 2000; Li et al. 2002].

In worms, pol η is SUMOylated by GEI-17 after DNA damage [Kim and Michael 2008]. This modification stabilizes pol η and protects it from CRL4<sup>Cdt2</sup>-mediated destruction so the translesion polymerase can repair damaged DNA. A recent study in S. pombe suggests that the presence of ubiquitylated PCNA may interfere with Cdt1 destruction [Guarino et al. 2011].

**Temporal regulation of CRL4<sup>Cdt2</sup>-dependent ubiquitylation**

An intrinsic feature of CRL4<sup>Cdt2</sup>-mediated proteolysis is that the process is only initiated after DNA replication has begun, since PCNA loading requires new DNA synthesis. This temporal delay in the destruction of CRL4<sup>Cdt2</sup> substrates upon S-phase entry has interesting implications for the destruction of different substrates. For example, immediate destruction of Spd1 upon S-phase entry may not be essential. Thus, S. pombe cells likely use the existing pool of dNTPs to begin DNA synthesis, whereupon CRL4<sup>Cdt2</sup>-dependent destruction of Spd1 stimulates the production of additional dNTPs.
to enable progression through S phase. Similarly, the precise timing of E2F destruction may not be crucial, as long as E2F activity is attenuated in S phase. In contrast, a delay in Cdt1, Set8, or p21 destruction in S phase would not be ideally suited to achieve an absolute injunction against rerelication. Thus, there might exist a temporal window at the start of S phase when DNA replication has begun but before Cdt1, Set8, and p21 levels have dropped. During this window, licensing on the nascent DNA should be possible. There are several potential solutions to this dilemma. First, in humans, CRL1Skp2 might help destroy Cdt1, Set8, and/or p21 in late G1 or S early phase (Li et al. 2003; Kondo et al. 2004; Takeda et al. 2005; Nishitani et al. 2006). Second, other inhibitors of rerepllication, such as Geminin, might prevent licensing in early S phase when Cdt1 is still present (Arias and Walter 2005). Third, we speculate that CRL4Cdt2 might protect replicated DNA from reinitiation locally. This idea is based on the fact that a new PCNA molecule is loaded every 150 nucleotides during Okazaki fragment synthesis. If PCNA unloading is slow after Okazaki fragment maturation, a high concentration of PCNA molecules would accumulate on the lagging strand (Arias and Walter 2006), protecting it from relicensing. The high density of PCNA would promote destruction of Cdt1 near replicated DNA even before the total pool of Cdt1 has disappeared from the nucleus. This mechanism is plausible because in metazoans, there is a lag phase between Cdt1 binding to ORC and recruitment of MCM2–7 (Waga and Zembutsu 2006), allowing time for CRL4Cdt2-dependent destruction of any Cdt1 bound to replicated DNA. This mechanism might also protect the leading strand, which is held in close association with the lagging strand by cohesion. As such, PCNA might function as a “body armor” for replicated DNA. For individual substrates, it is interesting to consider how well the timing and function of CRL4Cdt2-dependent destruction are aligned, and in cases of misalignment, whether compensatory mechanisms might exist.

Conclusion and future perspectives
As discussed in this review, the study of CRL4Cdt2 is establishing new concepts in the field of proteolysis. CRL4Cdt2 appears to be unique among E3 ubiquitin ligases because there is little evidence that its interaction with substrates depends on post-translational modifications. Rather, binding to substrates depends on their prior interaction, through a PIP degron, with a cell cycle-regulated structure, chromatin-bound PCNA. The coupling of ubiquitin ligase function to a transient interaction, through a PIP degron, with a cell cycle-regulated structure, chromatin-bound PCNA, the coupling of ubiquitin ligase function to a transient interaction between the degron and another protein is a novel concept in ubiquitin-mediated proteolysis. This strategy represents an elegant means to couple proteolysis to other metabolic events, and it will be surprising if this mechanism is not used by other E3 enzymes.

The analysis of CRL4Cdt2 substrates has also revealed interesting new concepts in DNA replication and cell cycle regulation. Thus, the discovery of p21 as a CRL4Cdt2 target has provided the most compelling evidence to date that Cdc6 export is important to prevent relicensing in S phase. Similarly, the targeting of Set8 by CRL4Cdt2 led to the insight that Set8 participates in licensing and that certain forms of chromatin are incompatible with the G1- and S-phase programs. The identification of Epe1 as a likely CRL4Cdt2 target represents a fascinating example in which proteolysis sculpts chromatin structure. Given the crucial role of CRL4Cdt2 in chromosome maintenance, particularly the control of licensing, it is possible that this E3 ubiquitin ligase has tumor suppressor activity (Abbas and Dutta 2011).

Several important questions remain to be answered. First, how many CRL4Cdt2 substrates exist? Identification of new targets is of prime importance, since it will help refine our understanding of the PIP degron and teach us further lessons about the regulation of S phase, DNA repair, and other chromatin-related processes. Second, does PCNA function solely as a platform to bring together the substrate and ligase, or does it also modulate the activity of the ligase? Third, why is CRL4Cdt2 activity strictly dependent on chromatin-bound PCNA? This dependency is the crux of CRL4Cdt2’s temporal regulation during the cell cycle. Finally, are DNA-bound PCNA and a PIP degron sufficient to harness the activity of CRL4Cdt2, or are additional proteins and/or post-translational modifications required? Given the intense interest that has focused on CRL4Cdt2 in recent years, we hope the answers to these questions will soon emerge.

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References
Abbas T, Dutta A. 2011. CRL4Cdt2: master coordinator of cell cycle progression and genome stability. Cell Cycle 10:241–249.

Abbas T, Sivaprasad U, Terai K, Amador V, Pagano M, Dutta A. 2008. PCNA-dependent regulation of p21 ubiquitylation and degradation via the CRL4Cdt2 ubiquitin ligase complex. Genes Dev 22:2496–2506.

Abbas T, Shihata E, Park J, Jha S, Karnani N, Dutta A. 2010. CRL4(Cdt2) regulates cell proliferation and histone gene expression by targeting PR-Set7/Set8 for degradation. Mol Cell 40:9–21.

Adimoolam S, Lin CX, Ford JM. 2001. The p53-regulated cyclin-dependent kinase inhibitor, p21 [crip1, waf1, sdi1], is not required for global genomic and transcription-coupled nucleotide excision repair of UV-induced DNA photoproducts. J Biol Chem 276:25813–25822.

Amador V, Ge S, Santamaria PG, Guardavaccaro D, Pagano M. 2007. APC/C(Cdc20) controls the ubiquitin-mediated degradation of p21 in prometaphase. Mol Cell 27:462–473.

Angers S, Li T, Yi X, MacGoss MJ, Moon RT, Zheng N. 2006. Molecular architecture and assembly of the Ddb1–Cul4A ubiquitin ligase machinery. Nature 443:590–593.

Armley RG, Robinson PA. 2005. E3 ubiquitin ligases. Essays Biochem 41:15–30.

Arias EE, Walter JC. 2005. Replication-dependent destruction of Cdt1 limits DNA replication to a single round per cell cycle in Xenopus egg extracts. Genes Dev 19:114–126.
Arias EE, Walter JC. 2006. PCNA functions as a molecular platform to trigger Cdt1 destruction and prevent re-replication. Nat Cell Biol 8:84–90.

Arias EE, Walter JC. 2007. Strength in numbers: preventing rereplication via multiple mechanisms in eukaryotic cells. Genes Dev 21:427–518.

Asano M, Nevins JR, Wharton RP. 1996. Ectopic E2F expression induces S phase and apoptosis in Drosophila imaginal discs. Genes Dev 10:1422–1432.

Bates S, Ryan KM, Phillips AC, Vousden KH. 1998. Cell cycle arrest and DNA endoreduplication following p21Waf1/Cip1 expression. Oncogene 17:1691–1703.

Behrends C, Harper JW. 2011. Constructing and decoding unconventional ubiquitin chains. Nat Struct Mol Biol 18:520–528.

Bendennat M, Boulaire J, Jascur T, Brickner H, Barbier V, Sarasin A, Fotedar A, Fotedar R. 2003. UV irradiation triggers ubiquitin-dependent degradation of p21(WAF1) to promote DNA repair. Cell 114:599–610.

Bornstein G, Bloom J, Sitry-Shevah D, Nakayama K, Pagano M, Behrends C, Harper JW. 2011. Constructing and decoding un-conventional ubiquitin chains. Nat Struct Mol Biol 18:520–528.

Braun S, Garcia JF, Rowley M, Rougemaille M, Shankar S, Bendjennat M, Boulaire J, Jascur T, Brickner H, Barbier V, Sarasin A, Fotendar A, Fotendar R. 2003. UV irradiation triggers ubiquitin-dependent degradation of p21(WAF1) to promote DNA repair. Cell 114:599–610.

Braun S, Garcia JF, Rowley M, Rougemaille M, Shankar S, Bendjennat M, Boulaire J, Jascur T, Brickner H, Barbier V, Sarasin A, Fotendar A, Fotendar R. 2003. UV irradiation triggers ubiquitin-dependent degradation of p21(WAF1) to promote DNA repair. Cell 114:599–610.

Cayrol C, Knibiehler M, Ducommun B. 1998. p21 binding to cyclin-dependent kinase inhibitor, p27Xic1, is both necessary and sufficient for inhibiting invasion of a boundary-associated antiscarcinoma factor into heterochromatin. Mol Cell 2:351–356.

Chen J, Jackson PK, Kirschner MW, Dutta A. 1995. Separate domains of p21 involved in the inhibition of Cdk kinase and PCNA. Nature 374:386–388.

Choe KP, Przybysz AJ, Strange K. 2009. The WD40 repeat protein WDR-23 functions with the CUL4–DDB1 ubiquitin ligase to regulate nuclear abundance and activity of SKN-1 in Caenorhabditis elegans. Mol Cell Biol 29:2704–2715.

Chuang LC, Yew PR. 2005. PCNA recruits Cdk inhibitor Xic1 to DNA and couples its proteolysis to DNA polymerase switching. J Biol Chem 280:35299–35309.

Chuang LC, Zhu XN, Herrera CR, Tseng HM, Pfleger CM, Block K, Yew PR. 2005. The C-terminal domain of the Xenopus cyclin-dependent kinase inhibitor, p27Xic1, is both necessary and sufficient for phosphorylation-independent proteolysis. J Biol Chem 280:34290–35285.

Cooper MP, Balajee AS, Bohr VA. 1999. The C-terminal domain of p21 inhibits nucleotide excision repair in vitro and in vivo. Mol Biol Cell 10:2119–2129.

Deshaias RJ, Joazeiro CA. 2009. RING domain E3 ubiquitin ligases. Annu Rev Biochem 78:399–434.

Deshaias RJ, Emberly ED, Saha A. 2010. Control of cullin-ring ubiquitin ligase activity by neddylation. Subcell Biochem 45:41–56.

Dimova DK, Dyson NJ. 2005. The E2F transcriptional network: old acquaintances with new faces. Oncogene 24:2810–2826.

Duda DM, Borg LA, Scott DC, Hunt HW, Hammel M, Schulman BA. 2008. Structural insights into NEDD8 activation of cullin-RING ligases: conformational control of conjugation. Cell 134:995–1006.

Fridman Y, Palgi N, Lazar D, Ben-Aroya S, Hieter P, Aharoni A. 2010. Subtle alterations in PCNA–partner interactions severely impair DNA replication and repair. PLoS Biol 8:e1000507. doi: 10.1371/journal.pbio.1000507.

Gao D, Inuzuka H, Tseng A, Chin KY, Toker A, Wei W. 2009. Phosphorylation by Akt1 promotes cytoplasmic localization of Skp2 and impairs APC/Cdc1 mediated Skp2 destruction. Nat Cell Biol 11:397–408.

Guarino E, Shepherd ME, Salguero J, Hua H, Deegan RS, Kearsey SE. 2011. Cdt1 proteolysis is promoted by dual PIP degrons and is modulated by PCNA ubiquitylation. Nucleic Acids Res doi: 10.1093/nar/gkr222.

Guilbis JM, Kelman Z, Hurwitz J, O’Donnell M, Kuriyan J. 1996. Structure of the C-terminal region of p21(WAF1/CIP1) complexed with human PCNA. Cell 87:297–306.

Hao B, Oehlmann S, Sowa ME, Harper JW, Pavlitch N. 2007. Structure of a Fbw7–Skp1–cyclin E complex: multisite-phosphorylated substrate recognition by SCF ubiquitin ligases. Mol Cell 26:131–143.

Havens CG, Walter JC. 2009. Docking of a specialized PIP Box onto chromatin-bound PCNA creates a degron for the ubiquitin ligase CRL4Cdt2. Mol Cell 35:93–104.

He YJ, McCall CM, Hu J, Zeng Y, Xiong Y. 2006. DDB1 functions as a linker to recruit receptor WD40 proteins to CUL4–ROC1 ubiquitin ligases. Genes Dev 20:2949–2954.

Higa LA, Zhang H. 2007. Stealing the spotlight: CUL4–DDB1 ubiquitin ligase docks WD40-repeat proteins to destroy. Cell Div 2:5.

Higa LA, Mihaylov IS, Banks DP, Zheng J, Zhang H. 2003. Radiation-mediated proteolysis of CDT1 by CUL4–ROC1 and CSN complexes constitutes a new checkpoint. Nat Cell Biol 5:1008–1015.

Higa LA, Banks D, Wu M, Kobayashi R, Sun H, Zhang H. 2006a. L2DTL/CDT2 interacts with the CUL4/DDB1 complex and PCNA and regulates CDT1 proteolysis in response to DNA damage. Cell Cycle 5:1675–1680.

Higa LA, Wu M, Ye T, Kobayashi R, Sun H, Zhang H. 2006b. CUL4–DDB1 ubiquitin ligase interacts with multiple WD40-repeat proteins and regulates histone methylation. Nat Cell Biol 8:1277–1283.

Hoege C, Pfander B, Moldovan GL, Pyrowolakis G, Jentsch S. 2002. RAD6-dependent DNA repair is linked to modification of PCNA by ubiquitin and SUMO. Nature 419:135–141.

Holmman JF, Beach D. 1994. cdt1 is an essential target of the Cdc10/Ser1 transcription factor: requirement for DNA replication and inhibition of mitosis. EMBO J 13:425–434.

Hohmberg C, Fleck O, Hansen HA, Liu C, Slabey R, Carr AM, Nielsen O. 2005. Ddb1 controls genome stability and meiosis in fission yeast. Genes Dev 19:853–862.

Hon WC, Wilson MJ, Harlos K, Claridge TD, Schofield CJ, Pugh CW, Maxwell PH, Ratcliffe PJ, Stuart DI, Jones EY. 2002. Structural basis for the recognition of hydroxyproline in HIF-1α by pVHL. Nat Rev Cancer 1:975–978.

Hook SS, Lin J, Dutta A. 2007. Mechanisms to control rereplication and implications for cancer. Curr Opin Cell Biol 19:663–671.

Houston SI, McManus KJ, Adams MM, Sims JK, Carpenter PB, Hendzel MJ, Rice JC. 2008. Catalytic function of the PR-Set7 histone H4 lysine 20 monomethyltransferase is essential for mitotic entry and genomic stability. J Biol Chem 283:19478–19488.

Hu J, Xiong Y. 2006. An evolutionarily conserved function of proliferating cell nuclear antigen for cdt1 degradation by the
cul4–ddb1 ubiquitin ligase in response to DNA damage. 

J Biol Chem 281: 3753–3756.

Hu J, McCall CM, Ohta T, Xiong Y. 2004. Targeted ubiquitination of CDT1 by the DDB1–CUL4–ROC1 ligase in response to DNA damage. Nat Cell Biol 6: 1003–1009.

Hsu J, Zacharek S, He YJ, Lee H, Shumway S, Duronio RJ, Xiong Y. 2008. WD40 protein FBW5 promotes ubiquitination of tumor suppressor TSC2 by DDB1–CUL4–ROC1 ligase. 

Genes Dev 22: 866–871.

Ishii T, Shiono Y, Takami T, Murakami Y, Ohnishi N, Nishitani H. 2010. Proliferating cell nuclear antigen-dependent rapid recruitment of Cdt1 and CRL4Cdt2 at DNA-damaged sites after UV irradiation in HeLa cells. 

J Biol Chem 285: 41993–42000.

Ivan M, Kondo K, Yang H, Kim W, Valiando J, Omm M, Salic A, Asara JM, Lane WS, Kaelin WG Jr. 2001. HIFα targets for VHL-mediated destruction by proline hydroxylation: implications for O2 sensing. Science 292: 464–468.

Jaakkola P, Mole DR, Tian YM, Wilson ML, Giebert J, Gaskell SJ, Kriegsheim A, Hebestreit HF, Mukherji M, Schofield CJ, et al. 2001. Targeting of HIF-α to the von Hippel-Lindau ubiquitylation complex by O2-regulated prolyl hydroxylation. 

Science 292: 468–472.

Jackson S, Xiong Y. 2009. CRL4s: the CUL4–RING E3 ubiquitin ligase family. Trends Biochem Sci 34: 562–570.

Jaspersen SL, Charles JF, Morgan DO. 1999. Inhibitory phosphorylation of the APC regulator Hct1 is controlled by the kinase Cdc28 and the phosphatase Cdc14. 

Curr Biol 9: 227–236.

Jin A, Arias EE, Chen J, Harper JW, Walter JC. 2006. A family of diverse Cul4–Dbdl1-interacting proteins includes Cdt2, which is required for S phase destruction of the replication factor Cdt1. 

Mol Cell 23: 709–721.

Jin J, Williamson A, Banerjee S, Philipp I, Rape M. 2008. Mechanism of ubiquitin-chain formation by the human anaphase-promoting complex. 

Cell 133: 653–665.

Jorgensen S, Eskildsen M, Fugger K, Hansen L, Larsen MS, Kousholt AN, Syljuasen RG, Trelle MB, Jensen ON, Helin K, et al. 2011. SET8 is degraded via PCNA-coupled CRL4(CDT2) ubiquitylation in S phase and after UV irradiation. 

J Cell Biol 192: 43–54.

Kerns SL, Torke SJ, Benjamin JM, McGarry TJ. 2007. Geminin prevents rereplication during Xenopus development. 

J Biol Chem 282: S514–S521.

Kim Y, Kipreos ET. 2007. The Caenorhabditis elegans replication licensing factor CDT-1 is targeted for degradation by the CUL-4/DDB-1 complex. 

Mol Cell Biol 27: 1394–1406.

Kim SH, Michael WM. 2008. Regulated proteolysis of DNA polymerase γ during the DNA-damage response in C. elegans. 

Mol Cell 32: 757–766.

Kim J, Feng H, Kipreos ET. 2007. C. elegans CUL-4 prevents rereplication by promoting the nuclear export of CDC-6 via a CKI-1-dependent pathway. 

Curr Biol 17: 966–972.

Kim Y, Starostina NG, Kipreos ET. 2008. The CRL4Cdt2 ubiquitin ligase targets the degradation of p21Cip1 to control checkpoint-dependent requirements for CUL4–RING E3 ubiquitin ligases. 

EMBO J 24: 395–404.

Lee HO, Zacharek SJ, Xiong Y, Duronio RJ. 2010. Cell type-dependent requirement for pIP box-regulated Cdt1 destruction during S phase. 

Mol Cell Biol 21: 3639–3653.

Li A, Blow JJ. 2009. Cdt1 downregulation by proteolysis and geminin inhibition prevents DNA re-replication in Xenopus. 

EMBO J 24: 915–926.

Li Y, Dowbenko D, Lasky LA. 2002. AKT/PKB phosphorylation of p21Cip/WAF1 enhances protein stability of p21Cip/WAF1 and promotes cell survival. 

J Biol Chem 277: 11352–11361.

Li X, Zhao Q, Liao R, Sun P, Wu X. 2003. The SCF(Skp2) ubiquitin ligase complex interacts with the human replication licensing factor Cdt1 and regulates Cdt1 degradation. 

J Biol Chem 278: 30854–30858.

Lin Jj, Millolinnen MA, Smith PG, Narayanun U, Dutta A. 2010. NEDD8-targeting drug MLN4924 elicits DNA rereplication by stabilizing Cdt1 in S phase, triggering checkpoint activation, apoptosis, and senescence in cancer cells. 

Cancer Res 70: 10310–10320.

Liu C, Powell KA, Mundt K, Wu L, Carr AM, Caspari T. 2003. Cop9/signalosome subunits and Pcu4 regulate ribonucleotide reductase by both checkpoint-dependent and -independent mechanisms. 

Genes Dev 17: 1130–1140.

Liu C, Poitelea M, Watson A, Yoshida SH, Shimoda C, Holmberg C, Nielsen O, Carr AM. 2005. Transactivation of Schizosaccharomyces pombe cdt2+ stimulates a Pcu4–Dbdl1–CSN ubiquitin ligase. 

EMBO J 24: 3940–3951.

Liu CL, Yu IS, Pan HW, Lin SW, Hsu HC. 2007. L2dtl is essential for cell survival and nuclear division in early mouse embryonic development. 

J Biol Chem 282: 1109–1118.

Liu P, Slater DM, Lenburg M, Nevis K, Cook JG, Vaziri C. 2009. Replication licensing promotes cyclin D1 expression and G1 progression in untransformed human cells. 

Cell Cycle 8: 125–136.

Machida YI, Terc JK, Dutta A. 2005. Acute reduction of an origin-recognition complex (ORC) subunit in human cells reveals a requirement of ORC for Cdk2 activation. 

J Biol Chem 280: 27624–27630.

Mailand N, Diffley JF. 2005. CDKs promote DNA replication origin licensing in human cells by protecting Cdc6 from APC/C-dependent proteolysis. 

Cell 122: 915–926.

McCall CM, Miliani de Marval PL, Chastain PD 2nd, Jackson SC, He YJ, Koteake Y, Cook JG, Xiong Y. 2008. Human immunodeficiency virus type 1 Vpr-binding protein VprBP, a WD40 protein associated with the DDB1–CUL4 E3 ubiquitin ligase, is essential for DNA replication and embryonic development. 

Mol Cell Biol 28: 5621–5633.

Medema RH, Klompmaker R, Smits VA, Rijksen G. 1998. p21waf1 can block cells at two points in the cell cycle, but...
does not interfere with processive DNA-replication or stress-activated kinases. *Oncogene* 16: 431–441.

Michishita M, Morimoto A, Ishii T, Komori H, Shiomi Y, Higuchi Y, Nishitani H. 2011. Positively charged residues located downstream of PIP box, together with TD amino acids within PIP box, are important for CRL4(Cdt2)-mediated proteolysis. *Genes Cells* 16: 12–22.

Milhollen MA, Narayanay U, Soucy TA, Veiby PO, Smith PG, Min JH, Yang H, Ivan M, Gertler F, Kaelin WG Jr, Pavletich NP, Mizushima T, Yoshida Y, Kumanomidou T, Hasegawa Y, Suzuki Michishita M, Morimoto A, Ishii T, Komori H, Shiomi Y, Moldovan GL, Pfander B, Jentsch S. 2007. PCNA, the maestro of activated kinases. *Genes Cells* 16: 2396–2407.

Ralph E, Boye E, Kearsery SE. 2006. DNA damage induces Cdt1 proteolysis in fission yeast via a pathway dependent on Cdt2 and Ddb1. *EMBO Rep* 7: 1134–1139.

Raman M, Havens CG, Walter JC, Harper JW. 2011. A genome wide screen identifies p97 as an essential regulator of DNA damage-dependent CDT1 destruction. *Mol Cell* [in press].

Ravid T, Hochstrasser M. 2008. Diversity of degradation signals in the ubiquitin-proteasome system. *Nat Rev Mol Cell Biol* 9: 679–690.

Rice JC, Nishioka K, Sarma K, Steward R, Reinberg D, Allis CD. 2002. Mitotic-specific methylation of histone H4 Lys 20 follows increased PR-Set7 expression and its localization to mitotic chromosomes. *Genes Dev* 16: 2225–2230.

Rodrigo-Brenni MC, Foster SA, Morgan DO. 2010. Catalysis of lysine 48-specific ubiquitin chain assembly by residues in E2 and ubiquitin. *Mol Cell* 39: 548–559.

Roukos V, Kinkhabwala A, Colombelli J, Kotsantis P, Taraviras S, Nishitani H, Stelzer E, Bastiaens P, Lygerou Z. 2011. Dynamic recruitment of licensing factor Cdt1 to sites of DNA damage. *J Cell Sci* 124: 422–434.

Saha A, Lewis S, Kleiger K, Kuhlman B, Deshaies RJ. 2011. Essential role for ubiquitin-ubiquitin-conjugating enzyme interaction in ubiquitin discharge from Cdc34 to substrate. *Mol Cell* 42: 75–83.

Sansam CL, Shepard JL, Lai K, Ianari A, Daniellan PS, Amsterdam A, Hopkins N, Lees JA. 2006. DTL/CDT2 is essential for both CDT1 regulation and the early G2/M checkpoint. *Genes & Dev* 20: 3117–3129.

Scott MT, Morrice N, Ball KL. 2000. Reversible phosphorylation at the C-terminal regulatory domain of p21(Waf1/Cip1) modulates proliferating cell nuclear antigen binding. *J Biol Chem* 275: 28307–28316.

Shibata E, Abbas T, Huang X, Wohlschlegel JA, Dutta A. 2011. Selective ubiquitylation of p21 and Cdt1 by UBCH8 and UBE2G ubiquitin conjugating enzymes via the CRL4<sup>CAEB</sup> ubiquitin ligase complex. *Mol Cell Biol* doi: 10.1128/MCB.05496-11.

Shibutani ST, de la Cruz AF, Tran V, Turbyfill WJ III, Reis T, Shibutani ST, de la Cruz AF, Tran V, Turbyfill WJ III, Reis T, Edgar BA, Duronio R. 2008. Intrinsinc negative cell cycle regulation provided by PIP box- and Cull4Cdt2-mediated destruction of E2F1 during S phase. *Dev Cell* 15: 890–900.

Soria G, Speroni J, Podhajcer OL, Prives C, Gottifredi V. 2008. p21 differentially regulates DNA replication and DNA-repair-associated processes after UV irradiation. *J Cell Sci* 121: 3271–3282.

Souchy TA, Smith PG, Milhollen MA, Berger AJ, Gavin JM, Adhikari S, Brownell JE, Burke KE, Cardin DP, Critchley S, et al. 2009. An inhibitor of NEDD8-activating enzyme as a new approach to treat cancer. *Nature* 458: 732–736.

Stuart SA, Wang YJ. 2009. Ionizing radiation induces ATM-independent degradation of p21/Cip1 in transformed cells. *J Biol Chem* 284: 15061–15070.

Sugasawa K, Okuda Y, Saijo M, Nishi R, Matsuda N, Chu G, Mori T, Iwai S, Tanaka K, Tanaka K, et al. 2005. UV-induced
ubiquitylation of XPC protein mediated by UV-DDB-ubiquitin ligase complex. *Cell* **121**: 387–400.

Takeda DY, Parvin JD, Dutta A. 2005. Degradation of Cdt1 during S phase is Skp2-independent and is required for efficient progression of mammalian cells through S phase. *J Biol Chem* **280**: 23416–23423.

Tan X, Zheng N. 2009. Hormone signaling through protein destruction: a lesson from plants. *Am J Physiol Endocrinol Metab* **296**: E223–E227. doi: 10.1152/ajpendo.90807.2008.

Tardat M, Brustel J, Kirsh O, Lefevbre C, Callanan M, Sardet C, Julien E. 2010. The histone H4 Lys 20 methyltransferase PR-Set7 regulates replication origins in mammalian cells. *Nat Cell Biol* **12**: 1086–1093.

Terai K, Machida YJ, Lahit H, Novac O, Hyrien O, Marheineke K, Zannis-Hadjopoulos M, Dutta A. 2006. Proliferating human cells hypomorphic for origin recognition complex 2 and pre-replicative complex formation have a defect in p53 activation and Cdk2 kinase activation. *J Biol Chem* **281**: 6253–6260.

Terai K, Abbas T, Jazaeri AA, Dutta A. 2010. CRL4(Cdt2) E3 ubiquitin ligase monoubiquitinates PCNA to promote translesion DNA synthesis. *Mol Cell* **37**: 143–149.

Vaziri C, Saxena S, Jeon Y, Lee C, Murata K, Machida Y, Wagle N, Hwang DS, Dutta A. 2003. A p53-dependent checkpoint pathway prevents rereplication. *Mol Cell** **11**: 997–1008.

Waga S, Zembutsu A. 2006. Dynamics of DNA binding of replication initiation proteins during de novo formation of pre-replicative complexes in *Xenopus* egg extracts. *J Biol Chem* **281**: 10926–10934.

Wang W, Nacusi L, Sheaff RJ, Liu X. 2005. Ubiquitination of p21Cip1/WAF1 by SCFSkp2: substrate requirement and ubiquitination site selection. *Biochemistry** **44**: 14553–14564.

Wang SC, Nakajima Y, Yu YL, Xia W, Chen CT, Yang CC, McIntush EW, Li LY, Hawke DH, Kobayashi R, et al. 2006. Tyrosine phosphorylation controls PCNA function through protein stability. *Nat Cell Biol* **8**: 1359–1368.

Warbrick E, Lane DP, Glover DM, Cox LS. 1995. A small peptide inhibitor of DNA replication defines the site of interaction between the cyclin-dependent kinase inhibitor p21WAF1 and proliferating cell nuclear antigen. *Curr Biol* **5**: 275–282.

Wickliffe KE, Lorenz S, Wemmer DE, Kuriyan J, R ape M. 2011. The mechanism of linkage-specific ubiquitin chain elongation by a single-subunit E2. *Cell* **144**: 769–781.

Wu S, Rice JC. 2011. A new regulator of the cell cycle: the PR-Set7 histone methyltransferase. *Cell Cycle* **10**: 68–72.

Wu S, Wang W, Kong X, Congdon LM, Yokomori K, Kirschner MW, Rice JC. 2010. Dynamic regulation of the PR-Set7 histone methyltransferase is required for normal cell cycle progression. *Genes Dev** **24**: 2531–2542.

Xu H, Wang J, Hu Q, Quan Y, Chen H, Cao Y, Li C, Wang Y, He Q. 2010. DCAF26, an adaptor protein of Cul4-based E3, is essential for DNA methylation in *Neurospora crassa*. *PLoS Genet* **6**: e1001132. doi: 10.1371/journal.pgen.1001132.

Ye Y, Rape M. 2009. Building ubiquitin chains: E2 enzymes at work. *Nat Rev Mol Cell Biol** **10**: 755–764.

Yen HC, Elledge SJ. 2008. Identification of SCF ubiquitin ligase substrates by global protein stability profiling. *Science* **322**: 923–929.

Yen HC, Xu Q, Chou DM, Zhao Z, Elledge SJ. 2008. Global protein stability profiling in mammalian cells. *Science* **322**: 918–923.

Yin Y, Yu VC, Zhu G, Chang DC. 2008. SET8 plays a role in controlling G1/S transition by blocking lysine acetylation in histone through binding to H4 N-terminal tail. *Cell Cycle* **7**: 1423–1432.

Yoshida Y, Chiha T, Tokunaga F, Kawasaki H, Iwai K, Suzuki T, Ito Y, Matsuoka K, Yoshida M, Tanaka K, et al. 2002. E3 ubiquitin ligase that recognizes sugar chains. *Nature** **418**: 438–442.

Yoshida K, Takisawa H, Kubota Y. 2005. Intrinsic nuclear import activity of geminin is essential to prevent re-initiation of DNA replication in *Xenopus* eggs. *Genes Cells* **10**: 63–73.

You Z, Harvey K, Kong L, Newport J. 2002. Xic1 degradation in *Xenopus* egg extracts is coupled to initiation of DNA replication. *Genes Dev** **16**: 1182–1194.

Yu ZK, Gervais JL, Zhang H. 1998. Human CUL-1 associates with the SKP1/SKP2 complex and regulates p21(CIP1/WAF1) and cyclin D proteins. *Proc Natl Acad Sci** **95**: 11324–11329.

Zachariae W, Schwab M, Nasmyth K, Seufert W. 1998. Control of cyclin ubiquitination by CDK-regulated binding of Hct1 to the anaphase promoting complex. *Science** **282**: 1721–1724.

Zaidi IW, Rabut G, Poveda A, Scheel H, Malmstrom J, Ulrich H, Hofmann K, Pasero P, Peter M, Luke B. 2008. Rtt101 and Mms1 in budding yeast form a CUL4(DDB1)-like ubiquitin ligase that promotes replication through damaged DNA. *EMBO Rep* **9**: 1034–1040.

Zheng N, Schulman BA, Song L, Miller JJ, Jeffrey PD, Wang P, Chu C, Koepp DM, Elledge SJ, Pagano M, et al. 2002. Structure of the Cul1–Rbx1–Skp1–F boxSkp2 SCF ubiquitin ligase complex. *Nature** **416**: 703–709.

Zhong W, Feng H, Santiago FE, Kipreos ET. 2003. CUL-4 ubiquitin ligase maintains genome stability by restraining DNA-replication licensing. *Nature** **423**: 885–889.