Cand1 Promotes Assembly of New SCF Complexes Through Dynamic Exchange of F-box Proteins

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SUMMARY

The modular SCF ubiquitin ligases feature a large family of substrate receptors that enable recognition of diverse targets. However, how the repertoire of SCF complexes is sustained remains unclear. Real-time measurements of formation and disassembly indicate that SCF(Fbxw7) is extraordinarily stable but, in the Nedd8-deconjugated state, is rapidly disassembled by the cullin-binding protein Cand1. Binding and ubiquitylation assays show that Cand1 is a protein exchange factor that accelerates the rate at which Cul1–Rbx1 equilibrates with multiple F-box Protein–Skp1 modules. Depletion of Cand1 from cells impedes recruitment of new F-box proteins to pre-existing Cul1 and profoundly alters the cellular landscape of SCF complexes. We suggest that catalyzed protein exchange may be a general feature of dynamic macromolecular machines and propose a hypothesis for how substrates, Nedd8, and Cand1 collaborate to regulate the cellular repertoire of SCF complexes.

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INTRODUCTION

Three enzymes work in succession to covalently attach ubiquitin and ubiquitin chains to target proteins: an ubiquitin activating enzyme (E1), an ubiquitin conjugating enzyme (E2), and an ubiquitin ligase (E3) (Dye and Schulman, 2007). The proteasome, a massive multi-subunit protease, recognizes and degrades proteins conjugated to lysine 48- or lysine 11-linked polyubiquitin chains containing at least four ubiquitins (Thrower et al., 2000; Wickliffe et al., 2011). Cullin–RING ubiquitin ligases (CRLs) are the largest family of E3s and are typified by the SCF complexes, which comprise four subunits: the scaffold Cul1, the RING domain protein Rbx1, the adaptor Skp1, and a substrate-binding F-box protein (Petroski and Deshaies, 2005; Dye and Schulman, 2007). Sixty-nine proteins in the human genome have F-box motifs, at least 42 of which form SCF complexes (Lee et al., 2011). Although this modularity of SCF complexes allows for recognition of diverse substrates, how SCF complex formation is regulated remains unclear.

Cullin-associated and neddylation-dissociated protein 1 (Cand1) is a Cul1 associated protein whose binding is mutually exclusive with the F-box Protein–Skp1 sub-complex and is also blocked by attachment of the ubiquitin-like protein Nedd8 to lysine 720 of Cul1 (Liu et al., 2002; Zheng et al., 2002a; Goldenberg et al., 2004; Bornstein et al., 2006; Duda et al., 2008; Siergiejuk et al., 2009). Neddylation of Cul1 induces a major conformational rearrangement in Cul1–Rbx1 that eliminates a Cand1 binding site and stimulates ubiquitin transfer from associated E2s to substrates (Duda et al., 2008; Saha and Deshaies, 2008). In vitro, Cand1 acts as a stoichiometric inhibitor of cullin neddylation, SCF assembly, and SCF ubiquitin ligase activity (Liu et al., 2002; Zheng et al., 2002a; Goldenberg et al., 2004; Bornstein et al., 2006; Duda et al., 2008; Siergiejuk et al., 2009). However, genetic evidence indicates that Cand1 is a positive regulator of SCF and other CRLs in vivo (Zheng et al., 2002a; Chuang et al., 2004; Feng et al., 2004; Lo and Hannink, 2006; Zhang et al., 2008; Bosu et al., 2010; Kim et al., 2010) suggesting that Cand1-mediated recycling of substrate receptor modules is important for proper CRL function (Liu et al., 2002; Cope and Deshaies, 2003; Dye and Schulman, 2007; Zhang et al., 2008; Schmidt et al., 2009). Here, we demonstrate a protein exchange factor activity for Cand1 that resolves the conflicting biochemical and genetic data, and together with other data on the Nedd8 conjugation cycle enables us to propose a specific model for how the cellular repertoire of CRL complexes is controlled.

RESULTS

Dynamics of SCF<sub>Fbxw7</sub> Assembly and Disassembly

To reconcile the conflicting observations on Cand1, we sought to characterize the assembly properties of SCF complexes by developing a real-time assay based on FRET that monitors the binding dynamics between F-box Protein–Skp1 and Cul1–Rbx1. Fbxw7 co-expressed with Skp1 was tagged at its C-terminus, via the Sortase reaction (Popp et al., 2009; Proft, 2009), with the peptide GGGGK conjugated to the fluorescent dye TAMRA, producing covalently labeled Fbxw7<sub>TAMRA</sub>. The trans-peptidation reaction was efficient (not shown) and the tag did not compromise ubiquitylation activity (Figure S1A). We observed FRET when Fbxw7<sub>TAMRA</sub>–Skp1 was mixed with Cul1–Rbx1 in which the N-terminus of Cul1 is fused to cyan fluorescent protein (CFP<sub>Cul1</sub>) (Figure 1A). The association rate constant ($k_{on}$) for complex assembly, $4 \times 10^6$ M$^{-1}$ s$^{-1}$, was determined by monitoring donor CFP<sub>Cul1</sub>–Rbx1 fluorescence at varying concentrations of Fbxw7<sub>TAMRA</sub>–Skp1 in a stopped flow apparatus (Figure 1B and Figure 1C). The FRET observed in our assay could be competed away by excess non-fluorescent Skp2–Skp1 (Figure 1D). Using this chase assay, we measured a dissociation rate constant ($k_{off}$) for SCF<sub>Fbxw7</sub> of $9 \times 10^{-7}$ s$^{-1}$, or 0.5 week$^{-1}$ (Figure 1E). These measurements revealed an extraordinarily tight complex with a $K_D$ of 200 fM. Neddylation of Cul1 did not affect the maximum FRET efficiency in our assay, nor
the rate of association or dissociation of SCFFbxw7, confirming that the neddylation-induced major conformational change in Rbx1 and Cul1’s C-terminal domain does not affect the binding interface of F-box Protein–Skp1 to Cul1’s N-terminal domain (Figure 1C and Figure S1B–S1D) (Zheng et al., 2002b; Petroski and Deshaies, 2005; Duda et al., 2008).

Cand1 Accelerates SCFFbxw7 Disassembly

F-box Protein–Skp1 and Cand1 antagonize each other’s binding to Cul1–Rbx1 (Liu et al., 2002; Zheng et al., 2002a; Goldenberg et al., 2004; Bornstein et al., 2006; Duda et al., 2008; Siergiejuk et al., 2009; Lee et al., 2011). To test if this is a general property, we prepared cell lysate from 293 cells that expressed tetracycline-inducible, FLAG-tagged Cu1 and were substantially depleted of Cand1 by stable expression of a lentiviral shRNA construct (Tet-FLAGCu1 Cand1kd cells). SCF complexes should be stable in this lysate due to the near absence of Cand1. This lysate was either mock-treated or supplemented with a large excess of pure Cand1. Addition of Cand1 resulted in reduced recovery of 20 of 21 F-box proteins observed in FLAGCu1 immunoprecipitates, whereas the level of Rbx1 association was not significantly affected (Figure S2; see Supplementary Discussion for a consideration of the one exception). To measure directly Cand1’s effect on SCF assembly in the absence of confounding factors that might be present in cell lysates, we added two-fold molar excess Cand1 to preformed pure SCFFbxw7 assembled from Fbxw7TAMRA–Skp1 and CFP Cul1–Rbx1. We observed a significant reduction in FRET after five minutes, indicating that Cand1 interfered with this assemblage (Figure 2A). This observation was not an artifact of our FRET assay, because when we repeated these experiments with neddylated CFP Cul1–Rbx1, Cand1’s effect was eliminated (Figure 2B).

The short time span used for Figure 2A is in direct contrast to the slow dissociation of Fbxw7TAMRA in the presence of excess unlabeled Skp2–Skp1 observed in Figure 1E, suggesting that Cand1 was not a conventional competitive inhibitor that trapped Cul1–Rbx1 as it dissociated from Fbxw7–Skp1, but instead actively promoted the disassembly of the SCFFbxw7 complex. To directly test this hypothesis, we measured the loss of FRET in real time upon addition of Cand1 to a preformed complex in a stopped flow apparatus (Figure 2C). Titration of Cand1 revealed increasingly rapid rates of SCFFbxw7 dissociation that followed saturation kinetics with a maximum rate of 1.3 s$^{-1}$ and a half maximal concentration ($K_M$) of 26 nM (Figure 2D). To eliminate interference from re-association of Fbxw7TAMRA–Skp1 with CFP Cul1–Rbx1, we repeated our measurements with unlabeled Skp2–Skp1 competitor in the reaction (Figure 2D). The maximal rate of Cand1-dependent dissociation remained unchanged while the $K_M$ increased to 53 nM (Figure 2D). In agreement with previous results, SCFFbxw7 formed with neddylated Cu1 showed no Cand1-induced dissociation over 30 seconds (Figure 2D). However, when the reactions were allowed to proceed for hours, we observed that saturating Cand1 accelerated disassembly of neddylated SCFFbxw7 by 45-fold (Figure 2E).

Four main points arise from this analysis. First, the $K_M$ of 53 nM sets an upper limit on the $K_D$ between Cand1 and CFP Cul1–Rbx1. Second, the saturation kinetics reveal the existence of a transient complex that contained Cand1, CFP Cul1–Rbx1, and Fbxw7TAMRA–Skp1. Third, the maximal observed rate of 1.3 s$^{-1}$ represents the rate of Fbxw7TAMRA–Skp1 dissociation from the transient complex, which is accelerated by >1 million-fold compared to the spontaneous dissociation of SCFFbxw7. Finally, neddlyation of Cu1 attenuated the effect of Cand1 by 30,000-fold.

These observations are reminiscent of a recent analysis that shows that binding of IκBα to NF-κB both slows the association of NF-κB with DNA and causes a significant increase in the rate of dissociation of NF-κB from DNA (Bergqvist et al., 2009). Thus, it seemed likely that Cand1’s dramatic effect on the dissociation rate of SCFFbxw7 would also be
accompanied by a decrease in the rate of assembly of SCF^{Fbxw7}. However, in the presence of saturating Cand1, we measured an association rate of $2.2 \times 10^6 \text{M}^{-1} \text{s}^{-1}$ for Fbxw7^{TAMRA}–Skp1 and CFP^{Cul1}–Rbx1 (Figure 2F). This is equivalent (within the error of our measurements) to the association rate without Cand1 (Figure 1C). As expected, when CFP^{Cul1} was modified with Nedd8, Cand1 also had no effect on the association rate (Figure 2F).

The above result has profound implications for building a model of Cand1’s function. First, Cand1 is not a competitive inhibitor. Whereas competitive inhibitors decrease $k_{on}$ and have no effect on $k_{off}$, Cand1 had no effect on $k_{on}$ but increased $k_{off}$. Second, unlike allosteric inhibitors, such as IκBα, that decrease the association rate of their targets in addition to increasing the dissociation rate, Cand1 only affected the dissociation rate of SCF^{Fbxw7}. This specific type of behavior is found in the mechanism of guanine nucleotide exchange factors (GEFs) that actively promote the exchange of GTP for GDP from target GTPases (Klebe et al., 1995; Goody and Hofmann-Goody, 2002; Guo et al., 2005). In these cases, GEF binding to its target GTPase increases the dissociation rate of guanine nucleotides, yet does not inhibit their association rate. This is accomplished by toggling between stable GEF–GTPase and GTPase–nucleotide complexes through a transient GEF–GTPase–nucleotide ternary complex in which the dissociation rates of both GEF and nucleotide are dramatically increased relative to their stable binary complexes (Klebe et al., 1995; Goody and Hofmann-Goody, 2002; Guo et al., 2005). Our results thus far conform to this model. Both Fbxw7–Skp1 and Cand1 formed tight complexes with Cul1–Rbx1 and there exists a transient ternary intermediate that exhibited a greatly increased rate of Fbxw7–Skp1 dissociation from Cul1–Rbx1.

**F-box Proteins Remove Cand1 from Cul1**

If Cand1 conforms to the behavior of GEFs, Cand1’s dissociation rate from Cul1–Rbx1 should be very slow but should increase dramatically in the presence of F-box Protein–Skp1 complexes. To measure this, we first developed a dissociation assay in which we monitored competitive displacement of Cand1^{TAMRA} from GST-Rbx1–Cul1 in the presence of 10-fold excess of unlabeled Cand1. Displacement was monitored either by native gel electrophoresis (Figure S3A and S3B) or rapid pull-down of GST on glutathione resin (Figure 3A). This protocol revealed that spontaneous dissociation of Cand1^{TAMRA} from Cul1 was extremely slow, with a rate of $1.2 \times 10^{-5} \text{s}^{-1}$ (Figure 3B). This explains why Ubc12 is unable to conjugate Nedd8 to Cul1 that is bound to Cand1 (Goldenberg et al., 2004; Bornstein et al., 2006; Duda et al., 2008; Siergiejuk et al., 2009).

Strikingly, this rate was greatly accelerated in the presence of Fbxw7–Skp1, such that displacement of Cand1^{TAMRA} was largely completed within 5 min (Figure 3C). This key result indicates that Fbxw7–Skp1 is sufficient to displace Cand1 from Cul1, without requiring assistance from the Nedd8 conjugation pathway. This is consistent with reports that loss of Nedd8 conjugation activity has very little effect on the steady-state repertoire of SCF ubiquitin ligases (Bennett et al., 2010; Lee et al., 2011). A similar enhancement of Cand1 displacement by 3 different F-box Protein–Skp1 complexes was observed using an indirect assay wherein removal of Cand1 was evidenced by the ability to conjugate Nedd8 to the liberated Cul1 (Figure S3C and S3D). Displacement of Cand1 by Fbxw7–Skp1 was very specific, because Cand1 was not displaced by either Skp1 alone nor Fbxw7 bound to a mutant of Skp1 (Skp1ΔΔ) used for crystallography that lacks two loops that would be expected to clash with Cand1 (Figure 3C and Figure S3B). Importantly, Skp1ΔΔ was fully able to sustain ubiquitylation, indicating that it was competent to bind Cul1 (Figure S3E).

Combining the results above with the data gathered so far, we constructed a kinetic framework similar to that of GEFs (Klebe et al., 1995; Goody and Hofmann-Goody, 2002;...
Guo et al., 2005) in which a transient ternary species comprising Fbxw7–Skp1, Cul1–Rbx1, and Cand1 rapidly collapses into sub-complexes that contain Cul1–Rbx1 bound by either Cand1 or Fbxw7–Skp1 (Figure 3D). We can build a similar cycle for neddylated Cul1–Rbx1 in which Cand1 rapidly dissociates from the ternary species, yet Fbxw7–Skp1 slowly dissociates form the ternary species (Figure 3D).

Our observations on Skp1ΔΔ, coupled with the report that deletion of a short β-hairpin from Cand1 allows formation of a stable Cand1–Skp1ΔΔ–Cul1–Rbx1 complex (Goldenberg et al., 2004) leads us to propose that the transient ternary complex we observed in Figure 2D is a high energy intermediate created by clashes involving the flexible acidic loops in Skp1 and the β hairpin in Cand1, enabling rapid dissociation of either Cand1 or F-box Protein–Skp1 from Cul1–Rbx1. In addition, we suggest that the interaction dynamics between Fbxw7–Skp1 and Cand1 should apply to most or all F-box proteins that form SCF complexes given that: (1) the main interaction of F-box proteins with Cul1 occurs through Skp1; (2) three different F-box proteins rapidly evicted Cand1 from Cul1–Rbx1 (Figure S3D); and (3) Cand1 was able to dislodge 20 of 21 F-box proteins from FLAGCul1 in cell lysates (Figure S2).

Cand1 Functions as an F-box protein Exchange Factor

Given that Cand1 can disrupt multiple SCF complexes in a cell lysate, we wondered if Cand1 can promote exchange of one Cul1-bound F-box protein for another. To test this directly, we measured the effect of Cand1 on the ability of Fbxw7TAMRA–Skp1 to gain access to Cul1 sequestered into complexes with β-TRCP–Skp1. As expected, pre-incubation of β-TRCP–Skp1 and CFP Cul1–Rbx1 diminished the observed association rate of 210 nM Fbxw7TAMRA–Skp1 with CFP Cul1–Rbx1 from 0.9 s⁻¹ to 5 × 10⁻⁵ s⁻¹, a reduction of 18,000 fold (Figure 4A). Remarkably, addition of 150 nM Cand1 to this assay increased the observed association rate to 0.07 s⁻¹, a 1,400-fold rescue (Figure 4B). Thus, Cand1 greatly reduced the timescale with which Cul1–Rbx1 equilibrates with the total population of β-TRCP–Skp1 and Fbxw7–Skp1. To determine whether SCFFbxw7 assembled in the presence of Cand1 was active, we measured ubiquitylation of a cyclin E peptide (CycE) that serves as an SCFFbxw7 substrate (Hao et al., 2007; Pierce et al., 2009). Addition of β-TRCP–Skp1 to a preformed SCFFbxw7 did not affect the rate of ubiquitylation of CycE (Figure 4C). However, switching the order in which β-TRCP–Skp1 and Fbxw7–Skp1 were incubated with Cul1–Rbx1 effectively inhibited the ubiquitylation of CycE, because β-TRCP–Skp1 formed stable complexes with Cul1, thereby denying access to CycE–Fbxw7–Skp1. Under these conditions, addition of Cand1 potently stimulated ubiquitylation of CycE (Figure 4C). The same was true if Cul1–Skp1 was preassembled with Skp2–Skp1 (Figure S3F). To our knowledge, all prior in vitro experiments implicated Cand1 as an inhibitor of SCF ubiquitin ligase activity. However, the design of our assay – which more faithfully mimics what happens in vivo as new F-box proteins are being synthesized – highlights the ability of Cand1 to activate an SCF complex by enabling the new F-box protein to gain access to Cul1 assembled into pre-existing SCF complexes.

Cand1 Modulates the SCF Repertoire in Cells

Consistent with the exchange activity observed with purified components, addition of excess purified β-TRCP–Skp1 to lysates of Tet-FLAGCul1 cells substantially reduced the recovery of 13 out of 15 endogenous F-box proteins detected in anti-FLAG immunoprecipitates (Figure S4). However, this effect was significantly attenuated for 11 of these F-box proteins by shRNA-mediated depletion of Cand1. In contrast, excess β-TRCP–Skp1 had no effect on recovery of Rbx1 regardless of the presence or absence of Cand1. Together with our biochemical data on purified proteins, these results point towards a general ability of Cand1
to act as a protein exchange factor that equilibrates Cul1–Rbx1 with the available pool of F-box Protein–Skp1 complexes.

To address whether Cand1’s exchange activity influences SCF complexes in cells, we quantified the steady state and dynamic nature of the Cul1-associated proteome in Tet.\textsuperscript{FLAG}Cul1 cells with or without Cand1 depletion. To evaluate the steady state architecture of the SCF proteome, we grew control cells in medium formulated with isotopically light lysine and arginine (‘light’ medium) and Cand1-depleted cells in medium formulated with isotopically heavy lysine and arginine (‘heavy’ medium). Both cultures were pulsed with tetracycline for 1 h at t=0 hours to induce transient synthesis of \textsuperscript{FLAG}Cul1. After t=25 hours \textsuperscript{FLAG}Cul1 immunoprecipitates were prepared, mixed, and analyzed by quantitative mass spectrometry. Analysis of isotopic ratios revealed that Cand1 depletion had no effect on Cul1 levels but exerted far-reaching effects on the SCF repertoire, with 14 of 34 SCF complexes increasing by ≥2-fold, and 2 decreasing by ≥1.5-fold (Figure 5A, black bars). Meanwhile, Cul1-bound Skp1 was increased 1.7-fold in Cand1-deficient cells. The net assembly of SCF complexes in Cand1-depleted cells is in agreement with the relative abundances of Cul1 (302 nM) and Cand1 (390 nM) in 293 cells (Bennett et al., 2010) and our biophysical data that excess Cand1 causes a net disassembly of SCF\textsuperscript{Fbxw7} (Figure 2A). In addition, the levels of Cul1-associated Nedd8 and CSN were increased by 2.4 and 1.5-fold, respectively. These changes are consistent with the increased overall assembly of SCF complexes in Cand1-depleted cells, and the ability of F-box proteins to stabilize binding of CSN (Enchev et al., 2012) while suppressing deneddylation of Cul1 (Emberley et al., 2012; Enchev et al., 2012). The effect of Cand1 on F-box proteins was specific, because the tightly-bound Cul1 partner Rbx1 exhibited very little change. These and other changes observed in Figure 5A were validated qualitatively by western blotting (Figure 5B). Importantly, mass spectrometry-based quantification of peptides diagnostic for a select group of F-box proteins revealed that the change observed in the steady-state repertoire of SCF complexes in Cand1-depleted cells was not a simple reflection of altered abundance of F-box proteins in total cell lysate (Figure 5A, white bars). To evaluate the relationship between altered abundance of SCF complexes and their role in protein degradation, we examined the degradation of an Fbxl3 substrate, cryptochrome Cry1 (Busino et al., 2007; Siepka et al., 2007). Whereas SCF\textsuperscript{Fbxl3} was more abundant in Cand1-depleted cells (Figure 5A), the rate of degradation of Cry1 was reduced (Figure 5C), as was the association between Cry1 and Cul1 (Figure S5A).

To connect our in vitro observations with what happens in cells, we sought to evaluate SCF dynamics in vivo. Control and Cand1-depleted cells growing in ‘light’ medium were pulsed with tetracycline for 1 h at t=0 hours to induce transient synthesis of \textsuperscript{FLAG}Cul1. At t=49 hours, the cells were pulsed with ‘heavy’ medium in the presence of the proteasome inhibitor epoxomicin to suppress apparent F-box protein exchange due to degradation. At t=61 hours, \textsuperscript{FLAG}Cul1 immunoprecipitates were prepared and analyzed by quantitative mass spectrometry to determine the fraction of each bound protein that was produced during the 12-hour pulse-label. In a parallel experiment, we monitored the heavy:light ratios in total cell lysate of peptides diagnostic for twelve different F-box proteins to evaluate the rate at which newly-synthesized forms of these proteins accumulated. As shown in Figure 5D, several newly-synthesized F-box proteins including Fbxw11, Fbxo11, and Fboxo21 exhibited reduced incorporation into the pool of pre-existing Cul1 in the absence of Cand1, consistent with Cand1 being an F-box protein exchange factor. Importantly, the isotopic ratios for F-box proteins in total cell lysate (including the three mentioned above) were not affected by the presence or absence of Cand1 (Figure S5B), indicating that low penetration of these pulse-labeled F-box proteins into the Cul1-bound pool of Cand1-depleted cells was not due to a reduced rate of synthesis.
DISCUSSION

Cand1 is an F-box Protein Exchange Factor and Modulates the Cellular Repertoire of SCF Complexes

Here we establish the first kinetic framework for the dynamic assembly of SCF complexes. Prior biochemical studies suggested that Cand1 is an inhibitor of SCF complexes, whereas genetic studies indicated that it promotes SCF function. Our results resolve this apparent paradox. We show that Cand1 can unambiguously stimulate SCF activity in vitro by enabling an F-box Protein–Skp1 complex to access Cul1 that was previously occupied by a different F-box Protein–Skp1 complex, and that Cand1 promotes assembly in vivo of new F-box proteins with pre-existing Cul1 molecules. We conclude that Cand1 serves as an exchange factor for F-box Protein–Skp1 complexes and conforms to the scheme originally proposed for GEFs (Klebe et al., 1995; Goody and Hofmann-Goody, 2002; Guo et al., 2005). However, unlike GEFs that exchange GDP for GTP, Cand1 promotes exchange of multiple F-box proteins on the Cul1 scaffold. Because Cand1 can bind other cullins (Lo and Hannink, 2006; Bosu et al., 2010; Chua et al., 2011) and influences ubiquitylation or degradation of multiple CRL substrates in vivo (Zheng et al., 2002a; Chuang et al., 2004; Feng et al., 2004; Lo and Hannink, 2006; Zhang et al., 2008; Bosu et al., 2010; Kim et al., 2010), we suggest that it serves as an exchange factor for CRL adaptors in general. In anticipation of a widespread role for Cand1 exchange activity in CRL biology and by analogy to guanine nucleotide exchange factors (GEFs), we suggest that it be referred to as a ‘substrate receptor exchange factor’ (SREF).

A perplexing feature of prior genetic studies is that Cand1 deficiency causes a partial reduction-of-function for some SCF complexes but not others (Chuang et al., 2004; Feng et al., 2004; Zhang et al., 2008; Bosu et al., 2010; Kim et al., 2010). We suggest that the impact of Cand1 on the steady state distribution of SCF complexes – with some complexes showing large changes in level whereas other complexes exhibit minimal perturbation (Figure 5A) – may partially account for this heretofore unexplained behavior. Another puzzling observation made in both human (Lo and Hannink, 2006) and Arabidopsis (Chuang et al., 2004) Cand1-deficient cells is that accumulation of a given CRL complex paradoxically correlates with stabilization of its substrate – a phenomenon also observed here. Our results suggest the interesting possibility that Cand1 biases the assembly of SCF complexes to favor F-box proteins for which substrates are available.

‘Substrate Sculpting’ of the CRL Repertoire: An Hypothesis

Based on the recent observations that substrates and other ligands can greatly reduce binding and deneddylation of CRLs by CSN (Fischer et al., 2011; Emberley et al., 2012; Enchev et al., 2012) we propose the hypothesis shown in Figure 6, which provides a potential mechanism for how the cellular repertoire of CRL complexes could be optimized to match substrate demand. In our model, which builds upon previous proposals (Cope and Deshaies, 2003; Schmidt et al., 2009), the exchange activity of Cand1 coupled with cycles of Nedd8 conjugation and de-conjugation enables CRL complexes to toggle between two radically different states—the Cand1 ‘exchange’ regime and the Nedd8 ‘stable’ state. When a cullin is conjugated with Nedd8, it can have extraordinary (sub-picomolar) affinity for its adaptor-bound substrate receptor (SR), which binds in a manner that is essentially irreversible (t1/2 ~9 days for Fbxw7–Skp1). Therefore, we envision that a CRL exists in a ‘stable’, active state when it is saturated with substrate, which occludes CSN binding. The Nedd8-conjugated CRL ubiquitylates substrate and can recruit downstream factors involved in substrate degradation (Bandau et al., 2012; den Besten et al., 2012). Once substrate is depleted, the ability of CSN to bind the CRL and remove Nedd8 is enhanced. In this metastable ‘intermediate’ state, the complex can either bind a new substrate and become
reactivated by Nedd8 conjugation to return to the ‘stable’ state, or it can bind Cand1 and enter the ‘exchange’ regime, resulting in up to a one million-fold increase in dissociation rate of adaptor–SR. The resulting Cand1–cullin–Rbx complex rapidly assembles with any of the available adaptor–SR complexes to form an unstable ternary intermediate that promptly decays to regenerate Cand1–cullin–Rbx or yield a new CRL complex. Neddylation of the latter species, which is stabilized by substrate, completes the cycle. We envision that substrates, by shielding CRL complexes from the actions of CSN and Cand1, help to sculpt the cellular repertoire of CRL complexes. Other factors are likely to contribute as well, including the rate of synthesis and degradation of substrate receptor proteins, as well as post-translational modifications and localization controls that modulate the access of CRL complexes to the Nedd8–Cand1 cycle.

An implication of our model is that in Cand1-deficient cells, the assembly state of CRLs would become uncoupled from substrate demand, such that the cullin could potentially become tied up in superfluous and non-productive CRL complexes. This situation may be tolerated in cells with constitutively high turnover of F-box and other substrate receptor proteins, but given the significant phenotypic consequences of Cand1 mutation (Chuang et al., 2004; Feng et al., 2004; Zhang et al., 2008; Bosu et al., 2010; Kim et al., 2010; Helmstaedt et al., 2011), it is apparent that ongoing synthesis and degradation of adaptor–SR modules (Bennett et al., 2010) or other mechanisms of SCF disassembly (Yen et al., 2012) by themselves do not suffice to sustain a fully functional CRL network.

Although other examples of factors that actively promote complex disassembly have been described (Klebe et al., 1995; Goody and Hofmann-Goody, 2002; Guo et al., 2005; Bergqvist et al., 2009; Chan et al., 2009), to our knowledge Cand1 is the first example of a factor that has the potential to promote equilibration of a protein scaffold with a large number of interacting partners. We suggest that ‘protein exchange factors’ that work analogously to Cand1 may play important roles in processes such as DNA replication, transcription, mRNA splicing, and vesicle trafficking that rely on protein machines that engage in transactions notable for their speed and affinity.

**EXPERIMENTAL PROCEDURES**

**FRET Assay**

Fluorimeter scans were performed on FluoroLog-3 (Jobin Yvon) in a buffer containing 30 mM Tris pH 7.6, 100 mM NaCl, 0.5 mM DTT, and 1 mg/ml Ovalbumin (Sigma) in a volume of 250 μl. Mixtures were excited at 430 nm and the emissions were scanned from 450 nm to 650 nm. Stopped flow reactions were performed on a Kintek stopped flow machine in the same buffer as the fluorimeter scans.

**Ubiquitylation Assay**

CycE was incubated with [γ-32P] ATP (132 nM) and Protein Kinase A for 45 min at 30°C to make radiolabeled CycE. Ubiquitylation reactions contained ATP (2 mM), Ubiquitin (60 μM), Ubiquitin E1 (1 μM), Cdc34b (10 μM), Cul1–Rbx1 (150 nM), and Fbxw7–Skp1 (varying concentrations). Additional proteins were included as mentioned in the text. Reactions were performed and quenched in buffers previously described for ubiquitylation assays (Pierce et al., 2009). Reactions were analyzed by running on 16% gels, drying, and quantifying with a phosphor screen (Molecular Devices).

**Cul1-Cand1 Dissociation Assay**

Cul1 with N-terminal tagged GST-Rbx1 (0.1 μM) was pre-incubated with Cand1 TAMRA (0.1 μM) for 10 min and captured on glutathione sepharose 4B resin. Aliquots of resin were
transferred to Micro Bio-Spin columns (Bio-Rad), resuspended in 1 μM of the indicated proteins, and incubated for 15 s or 5 min. Reactions were terminated by separation of beads and supernatant by centrifugation, and equivalent portions of each were fractionated by SDS-PAGE. Gels were scanned by a Typhoon™ scanner to quantify Cand1TAMRA. To analyze the dissociation rate of the Cul1-Cand1 complex, at various time points following the addition of 1 μM Cand1 to 0.1 μM GST-Rbx1–Cul1–Cand1TAMRA, an aliquot was withdrawn and incubated with glutathione sepharose 4B resin for 15-min and processed for SDS-PAGE analysis and fluorography. The gels were then stained by SilverQuest™ staining kit (Invitrogen) to detect the total Cand1 bands. The intensity of Cand1TAMRA and total Cand1 band was measured with ImageJ (NIH), and the intensity of Cand1TAMRA was normalized by the intensity of total Cand1 at each time point.

**Kinetic Analysis**

Regressions were performed in Matlab with the exception of Figure 3B, which was generated in Prism.

**Mass spectrometric analyses**

All data-dependent liquid chromatography-mass spectrometry and data analyses for Cul1-bound proteins were performed as described previously (Lee et al., 2011) with the following modifications. In experiments where we tested the effect of excess β-TrCP or Cand1 on SCF complex composition, purified β-TrCP or Cand1 was added to the lysate for 2 h at 23 °C, followed by immunoprecipitation for 15 min at the same temperature. To calculate the overall protein group ratio for an experiment, median evidence ratios were first calculated for each biological replicate, and overall ratio was calculated from the mean of the biological replicates. Standard errors of the overall protein group ratios were calculated using bootstrap analysis. PseudoMRM (Greco et al., 2010), also referred to as targeted peptide monitoring (Sandhu et al., 2008; Hewel et al., 2010) or peptide ion monitoring (Kulasingam et al., 2008) was used for the quantification of the selected peptides of F-box proteins in global lysates.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. FRET Reveals Properties of SCF Assembly

(A) Fluorescence emission spectra from excitation at 430 nm of 70 nM CFP\textsuperscript{Cul1–Rbx1}, 70 nM Fbxw7\textsuperscript{TAMRA–Skp1}, a mixture of the two, or buffer alone revealed FRET with 30\% efficiency upon complex formation. Signals were normalized to peak donor emission at 478 nm.

(B) The change in donor fluorescence versus time in a stopped flow apparatus with 5 nM CFP\textsuperscript{Cul1–Rbx1} and varying concentrations of Fbxw7\textsuperscript{TAMRA–Skp1}. Signal changes were fit to single exponential curves.

(C) The association rate constant for CFP\textsuperscript{Cul1–Rbx1} is $k_{on} = 4.2 \times 10^6$ M\textsuperscript{-1} s\textsuperscript{-1}, and $k_{on} = 4.5 \times 10^6$ M\textsuperscript{-1} s\textsuperscript{-1}.

(D) Fluorescence emission spectra from excitation at 430 nm of 70 nM CFP\textsuperscript{Cul1–Rbx1} and Fbxw7\textsuperscript{TAMRA–Skp1}.

(E) The dissociation rate constant for CFP\textsuperscript{Cul1–Rbx1} is $k_{off} = 8.5 \times 10^{-7}$ s\textsuperscript{-1}, and $K = 2 \times 10^{-13}$ M.

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(C) The rate of signal change in (B) versus the concentration of Fbxw7\textsuperscript{TAMRA}–Skp1. Fitting the data to \( k_{\text{obs}} = k_{\text{on}} \cdot [\text{Fbxw7}] + k_{\text{off}} \) gave \( k_{\text{on}} \) of \( 4 \times 10^6 \text{ M}^{-1} \text{ s}^{-1} \) regardless of Cul1’s neddylation status. Error bars: +/- SD, \( n \geq 3 \).

(D) 700 nM Skp2–Skp1 (chase) competed FRET away if pre-incubated with 70 nM Fbxw7\textsuperscript{TAMRA}–Skp1 before, but not after addition of 70 nM CFP\textsuperscript{Cul1} for 5 min.

(E) Fluorescence emission at 478 nm versus time after addition of chase to pre-incubated CFP\textsuperscript{Cul1–Rbx1} and Fbxw7\textsuperscript{TAMRA}–Skp1 normalized to peak donor emission in (D). Single exponential fit with a fixed end point of 1 gave \( k_{\text{off}} \) of \( 8.5 \times 10^{-7} \text{ s}^{-1} \). \( K_D \) is thus \( 2 \times 10^{-13} \text{ M} \). Error bars: +/- SD, \( n=3 \). See also Figure S1.
Figure 2. Cand1 Actively Removes Fbxw7–Skp1 From Cul1 by Altering Off Rate
(A) As in Figure 1A and Figure 1D except with the addition of 100 nM Cand1.
(B) As in (A), except using neddylated CFP-Cul1.
(C) The change in donor fluorescence versus time in a stopped flow apparatus upon addition of 150 nM Cand1 to 50 nM CFP-Cul1–Rbx1 pre-incubated with 50 nM Fbxw7 TAMRA–Skp1.
(D) The single exponential observed rates of SCF disassembly for various Cand1 concentrations mixed with 5 nM CFP-Cul1–Rbx1 or 5 nM neddylated CFP-Cul1–Rbx1 pre-incubated with 5 nM Fbxw7 TAMRA–Skp1. Chase indicates 700 nM Skp2–Skp1. Error bars: +/- SD, n ≥ 3.
(E) As in Figure 1E except with 150 nM or 300 nM Cand1 and 700 nM Skp2–Skp1 chase mixed with 70 nM neddylated CFP-Cul1 pre-incubated with 70 nM Fbxw7 TAMRA–Skp1. Error bars: range of values, n=2.
(F) As in Figure 1C, except with 150 nM Cand1 pre-incubated with 5 nM CFP-Cul1–Rbx1. Error bars: +/- SD, n ≥ 3. See also Figure S2.
Figure 3. F-box proteins Rapidly Remove Cand1 from Cul1

(A) GST-Rbx1–Cul1–Cand1TAMRA (100 nM) was supplemented with 1 μM Cand1. At indicated times, aliquots were removed and incubated with glutathione resin for 15 min. Resin-associated proteins were fractionated by SDS-PAGE and detected by fluorography. 

(B) The ratio of released Cand1TAMRA to total Cand1 over time was fit to a single exponential giving $k_{off}$ of $1.2 \times 10^{-5}$ s$^{-1}$. Error bars: +/- SD, n=3.

(C) GST-Rbx1–Cul1–Cand1TAMRA (100 nM) pre-incubated with glutathione resin was supplemented with buffer or 1 μM of indicated proteins. Bound and released proteins were collected at indicated times and distribution of Cand1TAMRA was evaluated as in (A).

(D) Summary of the rates measured here. Transient complexes are in brackets. See also Figure S3.
Figure 4. Cand1 Functions as an F-box protein Exchange Factor

(A) Fluorescence emission at 478 nm versus time after addition of 210 nM Fbxw7TAMRA–Skp1 to 70 nM CFP–Cul1 pre-incubated with 70 nM β-TRCP–Skp1. A single exponential fit gave $k_{off}$ of $5 \times 10^{-5}$ s$^{-1}$. Error bars: range of values, n=2.

(B) The change in donor fluorescence versus time in a stopped flow apparatus upon addition of 150 nM Cand1 to 70 nM CFP–Cul1–Rbx1 pre-incubated first with 70 nM β-TRCP–Skp1 and second with 210 nM Fbxw7TAMRA–Skp1.

(C) Cul1–Rbx1 (150 nM) was pre-incubated with 500 nM Fbxw7–Skp1 (lanes 1–6) or 660 nM β-TRCP–Skp1 (lanes 7–18) for 5 min, followed by addition of 600 nM radiolabeled cycE peptide substrate and either 660 nM β-TRCP–Skp1 (lanes 1–6) or 500 nM Fbxw7–Skp1 (lanes 7–18). Either buffer (lanes 1–12) or 200 nM Cand1 (lanes 13–18) were then added, and reactions were incubated an additional 5 min prior to initiation of an
ubiquitylation assay (all lanes) by supplementation of all lanes with 60 μM Ubiquitin, 1μM Ubiquitin E1, and 10 μM Cdc34b. See also Figure S4.
Figure 5. Cellular Cand1 Shapes Steady State and Dynamic SCF Landscape

(A) Tet-FLAG Cul controlkd and Tet-FLAG Cul1 Cand1kd cells (kd refers to knockdown) grown in medium with isotopically light or heavy lysine plus arginine, respectively, were induced with 1 μg/mL tetracycline for 1 hour and lysed in 1 μM MLN4924 and 2 mM o-phenanthroline 24 hours later. Two experiments were performed according to this protocol. In the first experiment, we used ‘pseudoMRM’ mass spectrometry to measure the relative amounts of 14 observable F-box proteins in total cell lysate from Cand1-depleted and control cells (white bars). In the second experiment, we retrieved FLAG Cul1 and measured the relative amounts of 34 F-box proteins in the immunoprecipitates (black bars). All isotopic ratios were normalized to FLAG Cul1’s (0.94), which was set to 1.0. For both experiments, results represent the ratio Cand1kd:controlkd of each protein in anti-FLAG IP measured by mass spectrometry. Each protein had ≥ 2 peptides. Error bars represent standard errors of overall protein group ratios, calculated from bootstrap analysis of two biological replicates (the second replicate was performed as a label-swap). Abundance changes in Cul1 IP for all the proteins listed in Fig5a except for Fbxo8, Fbxw9, and Fbxw4 achieved p-values < 0.05. Fbxo44a and b correspond to IPI00647771 and IPI00414844, respectively. Statistical analysis is provided in Table S1.

(B) Immunoblot validation with indicated antibodies of results in (A).

(C) The same cells used in (A) were transfected with a plasmid that encodes FLAG Cry1. Forty-eight hours later, a chase was initiated by addition of 40 μg/mL cycloheximide. Cells
were harvested at the indicated times and their content of \textit{\textbf{FLAG}}Cry1 and GAPDH was evaluated by SDS-PAGE and immunoblotting (left panel), and quantified (right panel). (D) The same cells used in (A) and grown in isotopically light lysine plus arginine were induced with 1 $\mu$g/mL tetracycline for 1 hour at $t=0$ hours, treated with 5 $\mu$M epoxomicin at $t=48$ hours, shifted to isotopically heavy lysine plus arginine at $t=49$ hours, and lysed at $t=61$ hours in 1 $\mu$M MLN4924 and 2 mM o-phenanthroline. Two experiments were performed according to this protocol. In the first experiment shown here, we used data-dependent mass spectrometry to discover and measure the fraction of F-box proteins in \textit{\textbf{FLAG}}Cul1 IPs that was heavy (\textit{i.e.} made in the 12 hours prior to lysis). In the second experiment (Figure S5B), we used pseudoMRM to target 9 F-box proteins (italicized) and measure the fraction of heavy-labeled species in total cell lysate from Cand1-depleted and control cells. Each protein had $\geq 2$ peptides. Error bars represent standard errors of overall protein group ratios, calculated from bootstrap analysis of two biological replicates. The F-box proteins shown to the left of the dotted line are those for which the different association observed in control and Cand1$^{kd}$ cells achieved a p-value <0.05. See also Figure S5 and Table S2.
Figure 6. Hypothesis for Control of CRL Assembly by Substrate, Cand1, and Nedd8
Rapid exchange of multiple CRL adaptor-bound substrate receptors occurs in the Cand1 exchange regime through the formation and decay of transient ternary complexes shown in brackets. Cand1 and adaptor are drawn as deformed in these complexes, to emphasize the proposal that they clash sterically, yielding an unstable state. In the presence of substrates, CRLs that pass through an intermediate state become neddylated and enter a stable state where ubiquitylation of substrates occurs. Loss of substrates facilitates recruitment of CSN, removal of Nedd8, and a return to the exchange regime effected by Cand1.