1 Structural and kinetic analysis of the COP9-Signalosome activation and the

2 cullin-RING ubiquitin ligase deneddylation cycle

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27 Abstract

The COP9-Signalosome (CSN) regulates cullin–RING ubiquitin ligase (CRL) activity and assembly 28 29 by cleaving Nedd8 from cullins. Free CSN is autoinhibited, and it remains unclear how it becomes 30 activated. We combine structural and kinetic analyses to identify mechanisms that contribute to 31 CSN activation and Nedd8 deconjugation. Both CSN and neddylated substrate undergo large 32 conformational changes upon binding, with important roles played by the N-terminal domains of 33 Csn2 and Csn4 and the RING domain of Rbx1 in enabling formation of a high affinity, fully active 34 complex. The RING domain is crucial for deneddylation, and works in part through conformational 35 changes involving insert-2 of Csn6. Nedd8 deconjugation and re-engagement of the active site zinc 36 by the autoinhibitory Csn5 glutamate-104 diminish affinity for Cul1/Rbx1 by ~100-fold, resulting 37 in its rapid ejection from the active site. Together, these mechanisms enable a dynamic 38 deneddylation-disassembly cycle that promotes rapid remodeling of the cellular CRL network.

39 Introduction

Cullin–RING ubiquitin ligases comprise one of the largest families of regulatory enzymes in eukaryotic cells (Deshaies and Joazeiro, 2009). With as many as 240 different enzyme complexes, these E3s control a broad array of biological processes (Skaar et al., 2013). CRLs comprise seven distinct cullin–RING cores, each of which interacts with its own dedicated set of adaptor–substrate receptor complexes. Although ubiquitination by CRL enzymes is often regulated by covalent modifications of the substrate that stimulate binding to the substrate receptor, the CRL enzymes themselves are also subject to regulation.

47 A key mechanism that controls the activity of all known CRLs is the conjugation of the 48 ubiquitin-like protein Nedd8 to a conserved lysine residue in the cullin subunit (e.g. K720 in 49 human Cul1)(Enchev et al., 2015). The available structural and biochemical data indicate that 50 Nedd8 conjugation (neddylation) stabilizes a profound conformational change in the C-terminal 51 domain of the cullin. It loosens the interaction of the WHB domain with the RING subunit, allowing both of them to sample a greater conformational space (Duda et al., 2008), thereby enhancing the 52 53 ability of the RING domain to promote ubiquitin transfer to substrate (Duda et al., 2008; Saha and 54 Deshaies, 2008; Yamoah et al., 2008).

55 In addition to direct effects on ubiquitin ligase activity, Nedd8 also protects Skp1/Cul1/F-56 box (SCF) complexes from the substrate receptor exchange factor (SREF) Cand1 (Pierce et al., 57 2013; Schmidt et al., 2009; Wu et al., 2013; Zemla et al., 2013). Cand1 binds unmodified SCF 58 complexes and promotes rapid dissociation of the F-box protein (FBP)/Skp1 substrate receptor-59 adaptor module from the Cul1/Rbx1 core. Cand1 can subsequently be dissociated from Cul1 by a 60 different FBP/Skp1 complex, and as a result Cand1 functions as an SREF that accelerates the rate at which Cul1/Rbx1 comes to equilibrium with different FBP/Skp1 substrate receptor-adaptor 61 62 complexes (Pierce et al., 2013). Importantly, the SREF activity of Cand1 is tightly restricted by 63 Nedd8. Cand1 is not able to bind stably to Cul1 and promote dissociation of FBP/Skp1 when Cul1 64 is conjugated to Nedd8 (Liu et al., 2002; Pierce et al., 2013). These observations underscore the 65 importance of neddylation not only for controlling the enzymatic activity of CRLs, but also 66 potentially for controlling the repertoire of assembled CRLs.

67 The key role of Nedd8 in CRL biology highlights the importance of the enzymatic pathways 68 that attach and remove Nedd8 (Enchev et al., 2015). Of particular significance is the rate of Nedd8 69 deconjugation, because it serves as the gateway for the exchange cycle; once Nedd8 is removed, a 70 CRL complex is susceptible to the potent SREF activity of Cand1, and its substrate receptor can be exchanged (Pierce et al., 2013). Deconjugation of Nedd8 is mediated by the COP9-signalosome 71 72 (CSN), which is an eight-subunit Nedd8 isopeptidase (Lyapina et al., 2001). The enzymatic activity 73 of CSN resides in its Csn5 subunit, which contains a metalloprotease active site referred to as the 74 'IAMM' domain (Cope et al., 2002). The IAMM domain has the general structure E₇₆-Xn-H₁₃₈-X-75 H_{140} -X10-D₁₅₁ (the subscripts refer to the sequence position of these residues in human Csn5), 76 wherein the H and D residues coordinate a zinc ion. The fourth zinc-coordination site is occupied 77 by a water molecule that that also forms a hydrogen bond to E76 (Ambroggio et al., 2004; Sato et 78 al., 2008; Tran et al., 2003). Deneddylation of CRLs by CSN is rapid but can be regulated by CRL 79 substrates (Emberley et al., 2012; Enchev et al., 2012; Fischer et al., 2011). Structural analysis 80 suggests that a CRL ubiquitination substrate bound to a substrate receptor sterically prevents concurrent binding of CSN (Enchev et al., 2012; Fischer et al., 2011). This suggests a model 81 82 wherein a CRL complex has a higher probability of being conjugated to Nedd8 (and therefore of 83 being shielded from Cand1) as long as it is bound to substrate. Upon dissociation of substrate, a 84 race ensues between binding of either a new substrate or CSN. If CSN wins, Nedd8 is removed, 85 paving the way for Cand1 to initiate substrate receptor exchange.

Recently, a crystal structure of free CSN was determined (Lingaraju et al., 2014). A major insight to emerge from the structure was the unexpected finding that Csn5 was present in an

88 autoinhibited state, wherein a glutamate (Csn5-E104) within the 'insert-1' (INS1) sequence 89 common to JAMM family members (Sato et al., 2008) forms a fourth ligand to the zinc, displacing 90 the catalytic Csn5-E76-bound water molecule and shifting Csn5-E76. Csn5-E104 is found in all 91 Csn5 orthologs, but not in other JAMM proteins, suggesting that this mode of regulation is 92 conserved but unique to CSN. Comparison of the structure of free CSN to the structure of a 93 catalytically-dead mutant CSN bound to Nedd8-conjugated SCF^{Skp2} determined by negative stain electron microscopy (Enchev et al., 2012) implied that binding of substrate to CSN may induce 94 95 several conformational changes in the latter, including movement of the N-terminal domains 96 (NTD) of Csn2 and Csn4 towards the cullin. The latter movement, in turn, might be further 97 propagated to the Csn5/6 module (Lingaraju et al., 2014). Moreover, it is reasonable to expect that 98 during catalysis INS1 moves out of the active site and Csn5-E76 adopts a position similar to that 99 observed in a crystallographic structure of Csn5 in isolation (Echalier et al., 2013). Interestingly, if 100 Csn5-E104 is mutated to an alanine. CSN more rapidly cleaves the simple model substrate 101 ubiquitin-rhodamine (Lingaraju et al., 2014). This was interpreted to mean that the primary 102 reason for the autoinhibited state is to keep CSN off until it binds a physiologic substrate, which would prevent spurious cleavage of non-cullin Nedd8 conjugates and possibly even ubiquitin 103 104 conjugates. However, the full extent of the conformational changes required to form an activated 105 complex between CSN and its neddylated substrate, as well as the detailed molecular basis for 106 these changes remain to be established. Therefore, at present, the mechanism of how CSN is 107 switched on and off and the significance of this switching behavior remain unknown.

108 **Results**

Structural insights from cryo EM and single particle analysis of a CSN-SCF-Nedd8^{Skp2/Cks1} complex

111 To gain detailed insights into the molecular determinants underlying activation of CSN, we 112 performed cryo electron microscopy (cryo EM) and single particle analysis of CSN^{5H138A} (we use 113 the nomenclature CSN^{#x} where # refers to subunit number and x to the specific mutation) in complex with neddylated SCF^{Skp2/Cks1} (the sample is described in (Enchev et al., 2012) (Figure 1A, 114 115 Figure 1-figure supplements 1A-D). The Csn5-H138A mutant lacks one of the JAMM ligands that 116 coordinate the catalytic zinc. This mutant forms a normal CSN complex that has been extensively 117 characterized (Enchev et al., 2012). We used ~75000 single molecular images for the final three-118 dimensional reconstruction and the structure was refined to a nominal resolution of 7.2 Å,

according to the 'gold standard' criterion of a Fourier shell correlation (FSC) of 0.143 (Rosenthal
and Henderson, 2003; Scheres and Chen, 2012) (Figure 1–figure supplements 1C–D). However,
some regions in the density map were better defined than others (see below). To avoid overinterpretation, for the subsequent analysis we low-pass filtered the map to 8.5 Å, according to the
more stringent criterion of an FSC of 0.5.

124 The cryo EM structure reported here, alongside the available crystal structure of CSN 125 (Lingaraju et al., 2014), enabled us to visualize a broad array of conformational changes that take 126 place upon complex formation in both CSN and neddylated Cul1/Rbx1, well beyond what was 127 possible with the prior lower resolution model based on negative stain EM (Figure 1). Specifically, 128 this allowed us to describe movements of the N-terminal domains of Csn2 and Csn4, the MPN 129 domains of Csn5 and Csn6. Moreover, in contrast to our previous work, we could locate the RING 130 domain of Rbx1, as well as Nedd8 and the winged-helix B (WHB) domain of Cul1 relative to Csn5. 131 Nevertheless, the present resolution precludes the determination of the exact orientations of the 132 latter domains but notably, the relative positions of the RING, WHB and Nedd8 reported here have 133 not been reported in any structural model of a cullin, and strongly suggest that both the enzyme 134 and substrate undergo significant conformational rearrangements to enable catalysis.

135 To obtain the model shown in Figure 1, we initially docked the crystal structure of CSN 136 (Lingaraju et al., 2014) and a model of Cul1-Nedd8/Rbx1/Skp1/Skp2/Cks1 (Enchev et al., 2012) 137 as rigid bodies into the electron density map (Figure 1-figure supplements 1E-H). We observed 138 very good matches between the respective map segments and the atomic coordinates for the 139 scaffold subunits Csn1, Csn3, Csn7 and Csn8, the winged-helix domains of Csn2 and Csn4 (Figure 140 1-figure supplement 1E), and the helical bundle formed by the C-termini of all eight CSN subunits 141 (Figure 1-figure supplement 1F) as well as the expected recovery of secondary structure at this 142 resolution. Similarly, there was a very good overlap between the coordinates of Cul1 (with the 143 exception of helix29 and the WHB domain, see below) and Skp1 and the corresponding electron 144 density segments (Figure 1-figure supplement 1G). However, the local resolution was lower 145 without recovery of secondary structure in the N-terminal domain of Cul1. Moreover, the density of the substrate receptor Skp2/Cks1 was poorly defined (Figure 1-figure supplement 1G). 146 147 indicating a potential flexibility in this region. Since the presence of Skp1/Skp2 had modest effects 148 on the affinity and deneddylation activity (see below), we did not interpret this observation 149 further.

150 In contrast to the large segments of CSN that were unaltered upon binding substrate, there 151 was nearly no overlap between the EM density map and the N-terminal portions of Csn2 and Csn4, 152 as well as the MPN-domains of Csn5 and Csn6, the RING domain of Rbx1, the WHB domain of Cul1, 153 and Nedd8 (Figure 1-figure supplement 1H). We thus docked these domains individually (Figure 154 1B, Movie 1). A Csn2 N-terminal fragment encompassing the portion between its 155 crystallographically resolved N-terminus (amino acid 30) through to a flexible loop at amino acid 156 180, was docked as a rigid body (Figure 1-figure supplement 1), positioning it close to the four-157 helical bundle and helix 24 of Cul1 (Zheng et al., 2002). An N-terminal fragment of Csn4, spanning 158 amino acids 1 to 295, which ends in a previously reported hinge loop (Lingaraju et al., 2014), was 159 also docked independently as a rigid body (Figure 1-figure supplement 1]). The resulting 160 conformation of Csn4 resembles a crystal form of Csn4 observed in isolation (Lingaraju et al., 161 2014). The two N-terminal helical repeat motifs of Csn4 make contacts with the winged-helix A 162 domain of Cul1 (Figure 1B and Figure 1-figure supplement 1], right hand panel, red arrow and 163 green circle). Moreover, these positions of Csn2 and Csn4 delineated a density in the map, which 164 could accommodate the RING domain of Rbx1 (Figure 1B and Figure 1–figure supplement 1J, right 165 hand panel, black ellipse), with the RING proximal to two conserved helices between amino acids 166 160 and 197 of Csn4 (Figures 1B and Figure 1-figure supplement 1K, black arrow) and a loop in 167 Csn2 located between residues 289 and 306. The exact orientation of the RING domain awaits a 168 structure at higher resolution.

169 To improve the fit of Csn5 and Csn6, we moved their MPN domains as rigid bodies into the 170 neighboring map segment of similar shape and dimensions (Figure 1-figure supplement 1L). The 171 local resolution in this region was lower, presumably due to higher flexibility around the catalytic 172 site. Importantly, after docking Csn5, we observed two empty neighboring densities (Figure 1– 173 figure supplement 1L, right hand panel, circles), which accommodated the two yet undocked 174 protein components – Nedd8 and the WHB domain of Cul1 (Figure 1-figure supplement 1M). The 175 docking of the latter was enabled by allowing helix 29 and the WHB, amino acids 690 to the C-176 terminus of Cul1, to move as a rigid body towards Csn5. However, we did not observe an electron 177 density around helix 29 of Cul1, consistent with a structural flexibility in this region. This model 178 places the neddylated WHB domain in close binding proximity to the RING domain, as well as both 179 INS1 and INS2 of Csn5. The hydrophobic patch of Nedd8 is facing INS1, and not the WHB domain, 180 as has been reported for the isolated neddylated C-terminal domain of Cul5 (Duda et al 2008). 181 Similar to the RING domain, we cannot be fully certain about the exact orientations of Nedd8 and

the WHB domain at the present resolution. Nevertheless, to further substantiate this docking, we mutagenized conserved charged residues in the INS2 domain of Csn5 as well as the WHB domain, and as expected all of these constructs showed reduced catalytic activity in deneddylation assays (Figure 1–figure supplement 1N, 0).

186 We sought orthogonal experimental validation for the molecular docking of the individual 187 subunits and domains in the electron density map by performing cross-linking coupled to mass 188 spectrometric analysis of the cross-linked peptides (Leitner et al., 2014) following the procedure 189 described in Birol et al. (2014) (supplementary files 1-6). For the cross-linker used in this study 190 (disuccinimidylsuberate H_{12}/D_{12}), the maximum predicted distance between two cross-linked lysine residues is generally accepted to be below ~ 30 Å (Politis et al., 2014). As shown in 191 192 supplementary files 1, out of the 39 high-confidence inter-subunit cross-links detected within the CSN^{5H138A}–N8-SCF^{Skp2/Cks1} complex at a false discovery rate (FDR) of 5 percent, the great majority 193 194 was within regions of modeled atomic structure and only six links exhibited a distance larger than 195 30 Å when mapped onto our model. However, all of these larger-distance links are connected to 196 the flexibly positioned Skp2 density. Moreover, we further performed similar cross-linking 197 experiments on a number of different CSN-CRL complexes, varying the substrate receptor, the 198 cullin and the neddylation state (supplementary files 2-6). All results were consistent with the 199 architecture proposed here for CSN^{5H138A}-N8-SCF^{Skp2/Cks1}. Intriguingly, when taking into account 200 cross-links with an FDR of up to 0.25 (supplementary files 2 and 3), we found two cross-links that support proximity of K290 in Csn4 and K89 in the RING domain (supplementary file 2), as well as 201 202 K32 in Csn4-NTD and K587 in Cul1, which is in the immediate vicinity of the WHA domain of Cul1 203 (supplementary file 3), as suggested by our EM reconstruction.

Development and validation of an assay to measure binding of CSN to substrate and product.

To understand how the structure of CSN and the CSN–SCF complex relates to substrate binding and the mechanism of deneddylation, we sought to develop quantitative binding assays to measure interaction of CSN with its substrates and products. To this end, the environmentallysensitive dye dansyl was conjugated to the C-terminus of Cul1 using 'sortagging' (Theile et al., 2013) to generate dansylated Cul1/Rbx1 (Cul1^d/Rbx1)(Figure 2–figure supplement 2A). Cul1^d/Rbx1 exhibited normal E3 activity (Figure 2–figure supplement 2B) and bound CSN with an affinity similar to Cul1/Rbx1 based on their IC₅₀ values for competitive inhibition of a

213 deneddylation reaction (Figure 2-figure supplement 2C; Emberley et al., 2012). When Cul1^d/Rbx1 214 was incubated with CSN (all CSN preparations used in this work are shown in Figure 2-figure 215 supplement 2D), we observed an increase in dansyl fluorescence (Figure 2A). This signal was due 216 to specific binding because it was chased upon addition of excess unlabeled Cul1/Rbx1 (Figure 2A, 217 titration shown in Figure 2-figure supplement 2E) or Cand1 (Figure 2-figure supplement 2F), 218 which competes for substrate deneddylation by CSN (Emberley et al., 2012; Enchey et al., 2012). 219 Thus, we concluded that the increase in dansyl fluorescence accurately reported on the interaction 220 of CSN with Cul1^d/Rbx1. Using this assay we determined that CSN bound Cul1^d/Rbx1 with a *K*_d of 221 310 nM (Figure 2B). Cul1^d/Rbx1 binding to CSN was only modestly affected by the addition of free 222 Nedd8 (Figure 2-figure supplement 2G) or assembly with Skp2/Skp1 (Figure 2-figure 223 supplement 2H) or Fbxw7/Skp1 (Figure 2-figure supplement 2I).

224 We next sought to measure binding of neddylated Cul1^d/Rbx1 (Cul1^d-N8/Rbx1) to CSN but 225 it was not possible because the substrate was rapidly deneddylated. To circumvent this problem, 226 we performed binding assays with the extensively characterized inactive mutant CSN^{5H138A} (assay 227 confirming loss of activity is shown in Figure 3-figure supplement 3A). Remarkably, CSN^{5H138A} 228 bound Cul1^d-N8/Rbx1 ~200-fold more tightly than CSN bound Cul1^d/Rbx1 (K_d 1.6 nM vs. 310 nM; 229 Figures 3A–B). Note that the estimated K_d falls well below the fixed concentration of Cul1^d-230 N8/Rbx1 used in the assay. This introduces greater uncertainty into our estimate but nevertheless 231 we can conclude with confidence that the binding of substrate to CSN^{5H138A} is very tight (≤ 5 nM; 232 see Materials and Methods for further discussion of this matter). As reported above for CSN 233 binding to product, addition of Skp2/Skp1 or Fbxw7/Skp1 had comparatively minor effects on 234 affinity (Figure 3-figure supplements 3B–C). Thus, for the sake of simplicity, we used Cul1^d/Rbx1 235 heterodimer for the remaining binding experiments.

236 The strikingly high affinity we observed for binding of CSN^{5H138A} to Cul1^d-N8/Rbx1 led us to 237 question whether it was mainly due to Nedd8 or whether the H138A mutation might also enhance affinity. To this end, we measured binding of CSN^{5H138A} to Cul1^d/Rbx1 and observed an 238 239 unexpectedly low K_d of ~10 nM (Figure 3B, Figure 3–figure supplement 3D), which was confirmed 240 with an independent preparation of CSN^{5H138A} (Figure 3–figure supplement 3E). Thus, neddylation 241 improved affinity of Cul1^d/Rbx1 for CSN^{5H138A} by ~6-fold, whereas the Csn5-H138A mutation 242 improved affinity for Cul1^d/Rbx1 by ~30-fold. The high affinity binding of CSN^{5H138A} to substrate was supported by an orthogonal competition experiment in which 100 nM CSN^{5H138A} completely 243

blocked deneddylation of 75 nM Cul1-N8/Rbx1 (Figure 3-figure supplement 3A). We considered
the possibility that the Csn5-H138A mutation might enable formation of an aberrant, super-tight
enzyme:substrate ([ES]) complex that does not normally form between the wild type proteins.
However, as will be described later on, this hypothesis was rejected based on kinetic arguments.

We next sought to determine whether the large differences we observed in K_d values were 248 due to differences in k_{on} or k_{off} . Remarkably, despite a 200-fold difference in K_d for CSN^{5H138A} 249 250 binding to substrate compared to CSN binding to product, the *k*_{on} values for formation of these 251 complexes were nearly identical (2.0 x 10⁷ M⁻¹ sec⁻¹ for CSN-product and 2.2 x 10⁷ M⁻¹ sec⁻¹ for 252 CSN^{5H138A}-substrate; Figure 3C). This suggested that the difference in affinity was driven by a 253 large difference in k_{off} . To test this hypothesis, we directly measured k_{off} values for select [ES] and 254 enzyme-product complexes by pre-forming the complex and then adding excess unlabeled 255 Cul1/Rbx1 chase and monitoring the reduction in dansyl fluorescence over time (Figure 3C and 256 Figure 3-figure supplement 3F-I; for this and a subsequent experiment in Figure 4B, we used 257 CSN^{5E76A/5H138A} in one of the assays instead of CSN^{5H138A}; the double mutant behaved like CSN^{5H138A} 258 in that it bound Cul1^d/Rbx1 with the same affinity as shown in Figure 3-figure supplement 3I.). 259 Consistent with the predictions from the K_d and k_{on} values, substrate dissociated very slowly from 260 CSN^{5E76A,5H138A}, whereas product dissociated ~65-fold faster from CSN. This suggests that as 261 substrate is deneddylated to product, its affinity for CSN is strongly reduced and its k_{off} speeds up.

The N-terminal domains of Csn2 and Csn4 and the RING of Rbx1 promote enzyme-substrate interaction.

264 Armed with assays to measure binding and deneddylation of substrate, we next sought to 265 test the implications that emerged from our structural analysis of the CSN^{5H138A}–N8-SCF^{Skp2/Cks1} 266 complex. First, we investigated the roles of the NTDs of Csn2 and Csn4, both of which, upon 267 binding substrate, underwent conformational changes and made contact with Cul1 and the RING 268 domain of Rbx1 (Figures 1B-C, Figure 1-figure supplements 1I-K)(Lingaraju et al., 2014). To 269 measure the effect of these mutations on binding to Cul1^d-N8/Rbx1, we combined them with 270 Csn5-H138A to prevent deneddylation. Deletion of the first 269 amino acids of Csn2, observed to 271 interact with Cul1 but not the RING domain of Rbx1, caused a massive loss in binding to substrate 272 $(K_d > 1300 \text{ nM}; \text{Figure 3B}, \text{Figure 3-figure supplement 3K})$. Thus, the contact we observed between Csn2-NTD and N8-SCF^{Skp2/Cks1} was critical to formation of the [ES] complex. By contrast, 273 274 deletion of the first 297 amino acids NTD of Csn4 ($4\Delta N$), a portion which was observed to form

interfaces with both Cul1 and the RING domain of Rbx1, had a relatively modest effect; CSN^{4 Δ N,5H138A} bound Cul1^d/Rbx1 and Cul1^d-N8/Rbx1 with *K_d* values of > 750 nM and 20 nM, respectively (Figure 3B, Figure 3–figure supplement 3L-M).

278 In addition to the motions of the Csn2 and Csn4 NTDs, our structural analysis revealed 279 formation of substantial interfaces between CSN and the RING domain of Rbx1. To test the role of 280 the RING domain in complex formation, we generated both Cul1/Rbx1 and Cul1^d/Rbx1 in which 281 the RING domain can be deleted by introducing a TEV protease cleavage site (Dougherty et al., 1989) after residue 37 of Rbx1 to generate Cul1 (or Cul1^d)/Rbx1^{TEV} (Figure 4A). This was 282 283 essential, because it would not be possible to conjugate Nedd8 to Cul1/Rbx1 expressed as a 284 mutant lacking the RING domain. After conjugating Nedd8 to the purified complex, we treated it 285 with TEV protease to remove the RING domain. vielding Cul1 (or Cul1^d)-N8/Rbx1^{Δ RING} (Figure 4A). 286 The truncated Cul1/Rbx1^{ARING} was inactive in an ubiquitylation assay (Figure 4–figure supplement 287 4A) but behaved as a monodisperse sample with the expected hydrodynamic radius upon size 288 exclusion chromatography (Figure 4–figure supplement 4B). Notably, Cul1^d-N8/Rbx1^{ΔRING} bound CSN^{5E76A,5H138A} and CSN^{5E76A} with affinities (12 nM and 13 nM respectively: Figure 3B, Figure 3– 289 290 figure supplement 3N) similar to that observed for binding of wild type Cul1^d-N8/Rbx1 to 291 $CSN^{4\Delta N,5H138A}$. Given the similar effects of the Csn4- Δ NTD and Rbx1- Δ RING mutations on complex 292 formation, we next tested whether their effects arose from loss of the interface that forms 293 between these domains (Figure 1-figure supplement 1K). However, double mutant analysis 294 suggested that the Csn4- Δ N and Rbx1- Δ RING mutations had largely independent effects on 295 binding (Figure 4B). The overall picture that emerged from these studies in light of the structural 296 data is that the interaction of Csn2-NTD with neddylated substrate makes a large contribution to 297 binding energy, with modest enhancements independently provided by the Csn4-NTD and Rbx1-298 RING domains.

299 The 'E-vict' enables efficient clearance of product from the CSN active site

The striking difference in the K_d for CSN^{5H138A} binding to substrate compared to CSN binding to product suggested that a conformational rearrangement of the [ES] complex occurs upon cleavage of the isopeptide bond, resulting in a large increase in the product k_{off} , thereby preventing the enzyme from becoming product-inhibited. However, we were puzzled by the relatively minor impact of Nedd8 on the affinity of Cul1^d/Rbx1 for CSN^{5H138A}; whereas substrate bound with K_d of 1.6 nM, product binding was only ~6-fold weaker (Figure 3B). Why, then, did 306 CSN bind so much less tightly to product? We reasoned that a key difference between CSN^{5H138A} 307 and CSN is the absence of the active site zinc from CSN^{5H138A}, which prevents formation of a stable 308 apo-CSN complex in which E104 of the INS1 domain of Csn5 is bound to the active site zinc. If this 309 conjecture is correct, it makes the prediction that CSN^{5E104A}, which should also be unable to form 310 stable apo-CSN, should likewise exhibit high affinity for product. This was confirmed: CSN^{5E104} 311 bound Cul1^d/Rbx1 with a *K_d* of 26 nM (Figure 3B, Figure 3–figure supplement 30). Furthermore, 312 measurement of k_{off} values revealed that product dissociated from CSN^{5H138A} and CSN^{5E104A} about 313 8-fold more slowly than it dissociated from CSN (Figure 3C). Based on these observations, we 314 propose the 'E-vict' hypothesis, which is described in more detail in the Discussion. The essence of 315 this hypothesis is that, following cleavage of the isopeptide bond and dissociation of Nedd8, INS1 316 of Csn5 engages the active site zinc. This accelerates the rate of dissociation of deneddylated 317 Cul1/Rbx1, thereby preventing CSN from becoming clogged with product. We note that Csn5-E76 318 also contributes to the operation of this mechanism, because CSN^{5E76A} bound tightly to product 319 (Figure 3-figure supplement 3P). We speculate that engagement of the active site zinc by Csn5-320 E104 forces Csn5-E76 into a configuration that promotes egress of product. Further insights into 321 the exact sequence of events that accelerates product dissociation await high-resolution 322 structures of CSN bound to Cul1/Rbx1 in various states.

323 Kinetic effects of binding-defective mutations on substrate deneddylation

324 We next sought to address the effects of the enzyme and substrate mutations described in 325 the preceding sections on the deneddylation reaction. We previously showed that CSN^{2ΔN} has 326 severely reduced catalytic activity (Enchev et al., 2012), which is consistent with the binding data 327 reported here. CSN^{4ΔN} exhibited a 20-fold defect in substrate cleavage (Figure 5A, Figure 5–figure 328 supplement 5A). Meanwhile, the k_{cat} for cleavage of Cul1-N8/Rbx1^{Δ RING} by CSN was reduced by a 329 staggering ~18,000-fold relative to wild type substrate (Figures 5A–B). Given that the neddylated 330 Δ RING substrate bound to CSN with only modestly reduced affinity, we surmised that the principal 331 defect of this mutant might be its failure either to induce the activating conformational change in 332 CSN, and/or to position accurately the isopeptide bond in the active site. Although we do not have 333 the tools to address the latter point, we queried the former by examining the Csn6- Δ INS2 334 mutation, which partially mimics the effect of substrate binding in that it destabilizes the 335 autoinhibited state (Lingaraju et al., 2014). The Csn6- Δ INS2 mutation slightly weakened binding 336 to wild type product (Figure 3B, Figure 3–figure supplement 3Q) but completely suppressed the

modest binding defect of the neddylated Δ RING substrate (Figure 3B, Figure 3–figure supplement 3R) and promoted an ~8-fold increase in its deneddylation rate (Figure 5–figure supplement 5B). This partial suppression effected by Csn6- Δ INS2 suggests that the RING domain contributes to the constellation of conformational changes in CSN that occur upon substrate binding. Note that the CSN^{6 Δ INS2} enzyme nevertheless exhibited a > 1,000-fold defect towards the Cul1-N8/Rbx1 Δ RING substrate, strongly indicating further functions of the RING domain, which may include a potential role in substrate positioning as well.

A noteworthy feature of the deneddylation reactions carried out with CSN^{4 Δ N} enzyme or ARING substrate is that although k_{cat} was reduced in both cases, K_M was also reduced (Figure 5A). Whereas these results imply that deletion of the Csn4-NTD or Rbx1-RING improved affinity of the [ES] complex, our direct binding measurements indicated this was not the case. To understand

Equation 1: Cul1-N8/Rbx1 deneddylation by CSN. The vertical red bar indicates the cleaved bond.

$$[CSN] + [Cul1-N8/Rbx1] \xrightarrow{k_{on}} [CSN-Cul1-N8/Rbx1] \xrightarrow{k_{cat}} [CSN-Cul1 | N8/Rbx1]$$

348 this apparent paradox, it is essential to consider the kinetic behavior of CSN-mediated 349 deneddylation. The formal definition of K_M for a deneddylation reaction (Equation 1), as stipulated 350 by Briggs and Haldane (Briggs and Haldane, 1925), is: $K_M = (k_{off} + k_{cat})/k_{on}$. In the special case of Michaelis-Menten kinetics, which is based on the assumption that k_{off} is much larger than k_{cat} the 351 352 expression simplifies to k_{off}/k_{on} , or K_d . However, k_{cat} for CSN (~1.1 sec⁻¹) is actually much faster than k_{off} measured for dissociation of substrate from the CSN^{5E76A, 5H138A} mutant (0.017 sec⁻¹). The 353 implication of this is that almost every binding event between CSN and substrate results in 354 355 catalysis, and K_M (200 nM; Figure 5A and (Emberlev et al., 2012) is much larger than K_d (1.6 nM, Figure 3B). But, if k_{cat} is reduced by mutation, the Briggs-Haldane equation predicts that K_M should 356 357 approach K_d . Indeed, this is exactly what we see for reactions that exhibit reduced k_{cat} , including 358 reactions with mutant CSN^{4 Δ N} enzyme or mutant Δ RING substrate (Figure 5A). In the slowest 359 reaction (cleavage of Cul1/Rbx1^{Δ RING} by CSN) the K_M (5 nM) is in the same range as the K_d with which this substrate bound to CSN^{5E76A, H138A} (12 nM; Figure 3B), and approaches the *K*_d measured 360 for binding of substrate to CSN^{5H138A} (1.6 nM). This provides strong support for our proposal that 361 362 the CSN^{5H138A}–Cul1^d/Rbx1 complex is representative of the affinity that develops during normal 363 catalysis.

364 **Functional significance of Csn5 INS1** *in vitro* and in cells

365 To understand the significance of the E-vict mechansim to CSN function in vitro and in cells, 366 we measured the k_{cat} for CSN^{5E104A} and observed that it is 2.5-fold slower than for CSN (Figure 5A, Figure 5-figure supplements 5C-E). This was unexpected, because it was reported that the Csn5-367 368 E104A mutation enhances the catalytic activity of CSN towards an unnatural substrate (Lingaraju 369 et al., 2014). Interestingly, a similar reduced rate was observed in both single- and multi-turnover 370 reactions, indicating that under our specific reactions conditions, the activating conformational 371 changes/chemical step are affected at least as much as product dissociation. This may not be the 372 case in cells, where substrate receptors and other factors may further stabilize product binding.

373 To test if Csn5-E104 contributes to CSN function *in vivo*, we generated a partial knockout of 374 Csn5 in HEK293T cells using CRISPR/Cas9 (Shalem et al., 2014a). This cell line expressed severely 375 reduced levels of Csn5 and consequently displayed hyper-accumulation of Nedd8-conjugated 376 endogenous Cul1 (Figure 6A), but retained sufficient protein to survive. We introduced either an 377 empty retrovirus or retroviruses coding for Flag-tagged wild type or mutant Csn5 proteins into 378 these cells, and then monitored the Cul1 neddylation status by immunoblotting. In contrast to wild type FlagCsn5, cells expressing FlagCsn5-E104A, H138A or E76A did not regain a normal pattern of 379 380 Cul1 neddylation (Figure 6A). The same was observed for Cul2. Cul3. Cul4A. and Cul5 (Figure 6– 381 figure supplement 6A). Consistent with reduced CSN activity, as revealed by increased cullin 382 neddylation, Skp2 levels were reduced in cells expressing mutant Csn5 proteins (Figure 6A)(Cope 383 and Deshaies, 2006; Wee et al., 2005). To test whether mutations in the catalytic site of Csn5 384 resulted in increased affinity for Cul1, we immunoprecipitated wild type and mutant FlagCsn5 proteins and probed for co-precipitation of endogenous Cul1. In addition to the mutants described 385 386 above, we surveyed a much broader panel of catalytic site substitutions to determine whether the 387 results observed in our *in vitro* experiments were specific to the mutations employed or were a 388 general consequence of disrupting the active site. As shown in Figure 6B, the results were 389 concordant with what was observed *in vitro*. On the one hand, ^{Flag}Csn5–H138A retrieved high 390 levels of Cul1-N8. The same was true for FlagCsn5 carrying mutations in other core residues of the 391 JAMM domain (e.g. H140 and D151)(Cope et al., 2002). On the other hand, FlagCsn5-E104A 392 retrieved high levels of unmodified Cul1. We propose that this arises from its ability to bind and 393 deneddylate substrate (albeit at a reduced rate), but then remain tightly bound to the product due 394 to loss of the E-vict mechanism.

395 The unexpected reduction in activity observed for Csn5-E104A both in vitro and in cells 396 (Figures 5A, 6A) suggested that the adjacent residue, T103, may also be important for 397 deneddylation. A T103I mutation in Drosophila melanogaster impedes proper interaction of 398 photoreceptor neurons with lamina glial cells in the developing brain. If this mutant also causes a loss of CSN deneddylase activity, it would explain the recessive nature of this mutation in flies 399 400 (Suh et al., 2002). Indeed, like Csn5-E104A. Csn5-T103I did not restore a normal Cul1 neddylation 401 pattern when expressed in Csn5-depleted cells (Figure 6C) and exhibited low deneddylase activity 402 *in vitro* (Figures 5A, Figure 5–figure supplement 5F). In contrast to CSN^{5E104A}, however, CSN^{5T103I} 403 bound Cul1^d/Rbx1 product with low affinity, both *in vitro* (Figure 3B, Figure 3–figure supplement 3S) and in cells (Figure 6B). Therefore, although CSN^{5E104A} and CSN^{5T103I} both have diminished 404 405 catalytic activity, their defects appear to have distinct molecular bases. To further explore the 406 divergent effects of Csn5-E104A and Csn5-T103I mutations on Cul1 binding, HEK293T stably 407 expressing wild type or mutant versions of FlagCsn5 were grown in 'heavy' SILAC medium (wild 408 type), or 'light' SILAC medium (mutants). Each mutant lysate was individually mixed with wild 409 type lysate, and then subjected to immunoprecipitation and SILAC mass spectrometry. Whereas 410 all CSN subunits exhibited light:heavy ratios of ~ 1 (Figure 6-figure supplement 6B), the FlagCsn5-411 E104A pull-down showed elevated levels of all cullins compared to wild type, whereas FlagCsn5-412 T103I pulled down cullins at levels equal to or less than wild type FlagCsn5 (Figure 6D).

413 **Discussion**

414 'Induced fit' underlies interaction of substrate with CSN and triggers enzyme activation

415 Figure 7 displays a model that incorporates published data and data presented in this 416 manuscript. Panels A-C shows a schematic view of the structural transitions that occur upon 417 substrate binding, and collectively contribute to efficient catalysis, whereas panel D provides the 418 rate constants for the deneddylation cycle. We tentatively propose the following sequence of 419 events. Free CSN exists in an inactive state in which E104 of Csn5-INS1 forms a fourth ligand to 420 zinc (Figure 7A)(Lingaraju et al., 2014). In this state the NTDs of both Csn2 and Csn4 are in "open" 421 conformations relative to the cullin substrate, and the MPN domains of Csn5/Csn6 are in a distal 422 position relative to it. Substrate binds this state rapidly (Figure 7B), likely driven by electrostatic 423 interactions between Cul1 and Csn2-NTD. This would account for the similar, extremely fast k_{on} 424 values that we measured for different combinations of Cul1^d/Rbx1 and CSN. Binding of CSN to 425 neddylated substrate results in a series of conformational changes in both complexes (Figure 7B).

426 These include (i) the translocation of the N-terminal helical repeats of Csn2 towards the CTD of 427 Cul1, (ii) the movement of the NTD of Csn4 towards the RING domain of Rbx1 and the WHA 428 domain of Cul1, (iii) the translocation of the MPN domains of Csn5-Csn6 towards the neddylated 429 WHB domain of Cul1, (iv) movement of the WHB domain towards Csn5, (v) the opening of the 430 interface between Nedd8 and the WHB domain, and (vi) the formation of a new interface between 431 Csn5 and Nedd8 probably involving the hydrophobic patch of Nedd8 and neighboring residues, as 432 well as a tenuous interface between the WHB and the Rbx1 RING domain. Furthermore, although 433 not structurally resolved in the present study, movements of Csn5-E76 and E104 towards and 434 away from the zinc atom (vii), respectively, probably similar to the conformation reported in 435 (Echalier et al., 2013), must occur to enable catalysis. Finally, a series of other unresolved 436 movements are likely to be germane including (viii) positioning of the extended C-terminus of 437 Nedd8, and the corresponding portion of the WHB domain for catalysis as well as contacts 438 between the INS1 and INS2 domains of Csn5 and the WHB domain of Cul1.

439 To probe the significance of the conformational changes summarized above, we generated and analyzed mutant enzymes. Deletion of Csn2-NTD virtually eliminated substrate binding 440 441 (Figure 3-figure supplement 3K), suggesting that movement of this domain (motion i) enables a 442 high affinity interaction between CSN and neddylated CRLs. Meanwhile a mutant lacking Csn4-NTD, CSN^{4 Δ N,5H138A</sub>, bound Cul1-N8/Rbx1 ~10-fold less tightly than CSN^{5H138A}, albeit still with a} 443 444 relatively high affinity (20 nM, Figure 3B). A similar effect on binding affinity was seen with a 445 substrate lacking the RING domain of Rbx1 (Figure 3B). Even though the RING and Csn4-NTD domains are adjacent in the enzyme-substrate [ES] complex (Figure 1), double mutant analysis 446 447 suggests that they make substantially independent contributions to binding energy (Figure 4B). 448 Interestingly, enzyme assays revealed a much greater effect of deleting the RING than deleting the 449 Csn4-NTD, suggesting that the RING domain makes a profound contribution to catalysis in a 450 manner that does not depend on its proximity to Csn4-NTD. We do not know the extent to which 451 the reduced catalytic rates for these mutants arise from defects in enzyme activation versus 452 substrate positioning, but we note that cleavage of Δ RING substrate was accelerated by ~8-fold 453 upon deletion of Csn6-INS2, suggesting that at least a small part of the problem with this substrate 454 is that it failed to properly trigger activating conformational changes in CSN.

In addition to the movements of individual domains, formation of the [ES] complex is accompanied by wholesale translocation of the Csn5 and Csn6 subunits. We suggest that this

457 motion contributes primarily a k_{cat} effect, because deletion of Csn6-INS2, which is proposed to 458 facilitate this motion, enhanced *k*_{cat} but had no noteworthy impact on binding to substrate (Figure 459 3B), and removal of Csn5 from the complex did not substantially affect CSN assembly with 460 substrate (Enchev et al., 2012). We cannot conclude much about the other motions enumerated 461 above, but we note that mutations that are predicted to reside near the interface of the Csn5-INS2 462 and Cul1-WHB domains cause significant reductions in substrate deneddylation (Figure 1-figure 463 supplements 1N–0). In addition, reorientation of Nedd8 away from Cul1-WHB and towards Csn5 464 as predicted here is consistent with the prior observation that the hydrophobic patch of Nedd8 465 recruits UBXD7 to neddylated CRLs (den Besten et al., 2012). Presumably, the conformational 466 changes that occur during the activation process are connected in some manner. Interestingly, 467 CSN^{5E104A} and $CSN^{6\Delta INS2}$ both cleave ubiquitin-rhodamine at 0.04 sec⁻¹ (which is ~6-fold faster than 468 wild type CSN), but CSN^{5E104A,6 Δ INS2</sub> is yet 5-fold faster (0.2 sec⁻¹) than either single mutant} (Lingaraju et al., 2014). The activities of the single and double mutants imply that the Csn6-ΔINS2 469 470 mutation must destabilize binding of Csn5-E104 to the catalytic zinc, but only in a small fraction 471 $(\leq 20\%)$ of complexes. Meanwhile, movements at the Csn4/6 interface must do more to the active 472 site than simply disrupt the interaction of Csn5-E104 with the catalytic zinc, implying the 473 existence of at least two inputs to CSN activation. Resolving how binding of substrate is connected 474 to enzyme activation awaits high-resolution structural analyses of the enzyme and substrate in 475 various states.

476 A kinetic model for the CSN enzyme cycle reveals an essential role for the E-vict mechanism 477 in sustaining rapid catalysis

478 Upon formation of an [ES] complex, the conformational changes that occur in both CSN and 479 substrate culminate in cleavage of the isopeptide bond that links Nedd8 to cullin. Although we 480 don't know the microscopic rate constants for the various conformational changes and bond 481 cleavage, all evidence points to the former being slower than the latter, which can occur with $k \ge k$ 6.3 sec⁻¹, based on the k_{cat} for cleavage of N8-CRL4A^{DDB2} by CSN^{6ΔINS2} (Lingaraju et al., 2014). The 482 483 actual cleavage may be even faster because this measurement was made under multi-turnover 484 conditions, in which case product dissociation may have been rate limiting. Regardless, the sum 485 total rate of the activating conformational motions plus isopeptide bond cleavage reported here (~1 sec⁻¹) is considerably faster than substrate dissociation from CSN^{5H138A} (~0.017 sec⁻¹), 486 487 indicating that CSN conforms to Briggs-Haldane kinetics and essentially every [ES] complex that

forms proceeds to cleavage, the physiological implications of which are considered in the nextsection.

490 Cleavage of the isopeptide bond initiates a series of events leading to product release. 491 Removal of Nedd8 increases dissociation of Cul1/Rbx1 by ~7-10 fold. We propose that 492 dissociation of the cleaved Nedd8 also removes an impediment to Csn5-INS1, which can now bind 493 the catalytic site zinc via E104 to return CSN to its apo state. This engagement, which we refer to 494 as the 'E-vict' mechanism, is a critical step in what is likely to be a series of conformational 495 rearrangements that include repositioning of Csn5-E76. Collectively, these movements reduce the 496 affinity of CSN for product and accelerate its rate of dissociation by an additional order of magnitude. The removal of Nedd8 and E-vict together bring about an \sim 100-fold loss in affinity of 497 498 Cul1/Rbx1 for CSN. The slow dissociation of product from CSN mutants that were unable to 499 undergo E-vict $(0.12-0.16 \text{ sec}^{-1})$; Figure 3C) suggests that this mechanism is important for 500 maintaining physiological rates of CRL deneddylation. This is further supported by the 501 observation that Csn5-E104A, but not wild type Csn5, co-precipitates substantial amounts of 502 deneddylated Cul1 from cells (Fig. 6B). Slow clearance of product could explain, in part, the failure 503 of this mutant to complement a Csn5 deficiency (Fig. 6A). The E-vict mechanism presents an 504 elegant solution to a fundamental challenge facing enzymes: how to achieve high specificity 505 without compromising rapid turnover.

We note that the product k_{off} for Cul1^d/Rbx1 (1.1 sec⁻¹) is similar to the k_{cat} we measured for both single- and multi-turnover reactions. This suggests that depending on the exact structure of the neddylated CRL substrate, the rate-limiting step may vary from one deneddylation reaction to another. Regardless, our biochemical and cell-based data suggest that if the E-vict mechanism did not exist, product dissociation would become the Achilles heel of deneddylation reactions.

511 **CSN in its cellular milieu**

The kinetic parameters reported here coupled with quantitative measurements of protein concentrations by selected reaction monitoring mass spectrometry ((Bennett et al., 2010) and J.R. and R.J.D., unpublished data) allow a preliminary estimate of the steady-state distribution of CSN in cells. The total cullin concentration in the 293T cell line used in this work is ~2200 nM. Meanwhile, the CSN concentration is ~450 nM. Although the total amount of Nedd8-conjugated cullins was not measured, immunoblot data suggest that ~1000 nM is a reasonable estimate. The

518 K_d reported here for the [ES] complex (~2 nM), thus predicts potentially near-complete saturation 519 of the cellular CSN pool with neddylated cullins. This implies that formation of new [ES] 520 complexes is limited by the slowest step in the catalytic cycle, i.e. either the conformational 521 rearrangements or product dissociation. In vitro, CSN follows Briggs-Haldane kinetics and cleaves 522 Nedd8 off nearly every neddylated CRL that it binds. Because CSN is not in equilibrium with its 523 substrates in our simplified *in vitro* system, it cannot rely on differences in substrate K_d to achieve specificity. Thus, differences in k_{off} on the order of ≤ 10 -fold, which might occur with different 524 525 cullins or substrate adaptors, would be predicted to have minimal effects on catalytic efficiency 526 provided that k_{on} remains roughly the same, as was observed for different configurations of 527 substrate and product in this study. Importantly, this parameter can potentially be profoundly 528 altered by ubiquitylation substrates, E2 enzyme, or other in vivo binding partners of Nedd8-529 conjugated CRLs, which compete with CSN (Emberley et al., 2012; Enchev et al., 2012; Fischer et 530 al., 2011) and thus should reduce its apparent k_{on} . It is also conceivable that binding partners 531 might alter the partitioning of the CSN–N8-CRL complex either by increasing k_{off} and/or reducing 532 k_{cat} , such that N8-CRL bound to an ubiquitylation substrate dissociates prior to completion of the 533 conformational rearrangements that culminate in its deneddylation.

534 Based on measurements reported here, it is likely that CSN complexes in cells are 535 constantly undergoing catalysis, dissociating rapidly from product, and rebinding other CRLs on 536 the time-scale of a few seconds or less. Consistent with this picture, addition of a Nedd8 537 conjugation inhibitor to cells leads to nearly complete disappearance of neddylated cullins within 538 5 minutes, and this does not account for the time it takes the drug to equilibrate across the 539 membrane and deplete the cellular pool of Nedd8~Ubc12 thioesters (Soucy et al., 2009). The 540 dynamic properties of CSN measured here reveal a CRL network of extreme plasticity that can be 541 reconfigured in minutes to respond to changing regulatory inputs. Although quantitative studies 542 of CRL network dynamics remain in their infancy, it is evident that the tools are at hand to begin to 543 understand how these remarkable enzymes function and are regulated within cells.

544 Materials and Methods

545 *Cloning* All eight wild type CSN subunits were cloned into a single pFBDM baculovirus transfer 546 MultiBac vector (Berger et al., 2004). His₆-Csn5 was inserted into the first multiple cloning site (MCS1) 547 of a pFBDM vector using NheI/XmaI and Csn1 was put into MCS2 of the same vector with BssHII/NotI. 548 Similarly, Csn2 was inserted into a second pFBDM vector using BssHII/NotI and StrepII^{2x}-Csn3,

549 containing an N-terminal PreScission-cleavable StrepII^{2x}-tag, using NheI/XmaI. From this plasmid the Csn2/StrepII^{2x}-Csn3 gene cassette was excised out with AvrII/PmeI and inserted into pFBDM^{Csn1/His6Csn5}. 550 551 whose multiplication module had been linearized with BstZ17I and SpeI, yielding pFBDM^{Csn1/His6-} ^{Csn5/Csn2/StrepII2x-Csn3}. A pFBDM^{Csn4/Csn7b} vector was generated using BssHII/NotI to insert Csn4 and 552 NheI/XmaI for Csn7b, and the resultant gene cassette was inserted into linearized pFBDM^{Csn1/His6-} 553 Csn5/Csn2/StrepII2x-Csn3, resulting in pFBDM^{Csn1/His6-Csn5/Csn2/StrepII2x-Csn3/Csn4/Csn7b}. Finally, a pFBDM^{Csn6/Csn8} 554 555 vector was generated using BssHII/NotI for Csn6 and NheI/XmaI for Csn8 insertion. Once again the resultant gene cassette was inserted into linearized pFBDM^{Csn1/His6-Csn5/Csn2/StrepII2x-Csn3/Csn4/Csn7b}, vielding 556 the full wild type CSN vector pFBDM^{Csn1/His6-Csn5/Csn2/StrepII2x-Csn3/Csn4/Csn7b/Csn6/Csn8}. A similar cloning 557 strategy was applied for the generation of CSN^{5E76A}, CSN^{5E76A, H138A}, CSN^{5E212R, D213R} and CSN^{4ΔN1-297}. 558 except that site-directed mutageneses were performed on pFBDM^{Csn1/His6Csn5} and pFBDM^{Csn4/Csn7b} 559 respectively. CSN^{5E104A} and CSN^{5T103I} were generated with the same general approach, except that that 560 561 site-directed mutagenesis and sequence validation were performed on a pCRIITOPO plasmid (Invitrogen) containing StrepII^{2x}-Csn5. Those mutants were then ligated into a MCS1 linearized 562 pFBDM^{Csn1} plasmid. For the production of CSN^{6ΔIns2} we used co-expression from two separate viruses. 563 To this end we applied site-directed mutagenesis on the pFBDM^{Csn6/Csn8} vector to delete amino acids 564 174-179 in Csn6, generating pFBDM^{Csn6\DeltaIns2/Csn8}. The gene cassette of the latter was excised out using 565 pFBDM^{Csn4/Csn7b}. 566 AvrII/PmeI and inserted into BstZ17I/SpeI linearized vielding pFBDM^{Csn4/Csn7b/Csn6\DeltaIns2/Csn8}. The resultant bacmid was used together with a bacmid generated from 567 pFBDM^{Csn1/His6-Csn5/Csn2/StrepII2x-Csn3} in order to generate two baculoviruses, which were used for co-568 infection to generate CSN^{6ΔIns2}. An analogous strategy was applied to generate CSN^{4ΔN/6ΔIns2}. 569 $CSN^{5H138A/6\Delta Ins2}$ and $CSN^{5H138A/4\Delta N}$ 570

The TEV site in Rbx1 as well as mutations in the WHB domain of Cul1 were obtained by site-directed mutagenesis on the pFBDM-Cul1/Rbx1 vector described in (Enchev et al., 2010), which further contained a C-terminal sortase tag described in the next section. Cloning of Cul3/Rbx1 used in the crosslinking/mass spectrometry experiments, Nedd8-pro-peptide-StrepII^{2x} and StrepII^{2x}-Den1 are described in (Orthwein, 2015). Recombinant bacmid and virus generation as well as protein expression proceeded as described in (Enchev et al., 2012). All genes were validated by sequencing as wild type or mutant.

Protein Purification and modifications CSN and its mutant forms were purified as described in Enchev
et al. (2012). Nedd8-activating and conjugating enzymes were purified as described in Emberley et al.
(2012) and Enchev et al. (2012). Fluorescently-labeled Cull substrates were conjugated with untagged

581 Nedd8. Cul1-sortase was designed with GGGGSLPETGGHHHHHH inserted after the final amino acid 582 of Cull into the pGEX vector described in Emberley et al. (2012). All sortase reactions were done at 583 30 °C overnight with 30 µM Cul1/Rbx1, 50 µM Sortase and 250 µM GGGGK-dansyl in 50 mM Tris pH 584 7.6, 150 mM NaCl and 10 mM CaCl₂ and purified by size exclusion chromatography to yield 585 Cull^d/Rbx1. Cull^d/Rbx1 was neddylated and purified as in Emberley et al. (2012) to yield Cull^d-586 N8/Rbx1. Cand1 and Sortase were purified as described in Pierce et al. (2013). Production of Cul1/Rbx1 587 and Cul3/Rbx1 baculovirus constructs used for electron microscopy and crosslinking mass spectrometry, bacterial split-and-co-express Cul1/Rbx1 Δ^{RING} , Nedd8 with native N- and C-termini, used for electron 588 microscopy and crosslinking mass spectrometry and for the experiments involving Cul1/Rbx1^{TEV}. Den1 589 590 as well as the respective preparative neddylation were performed as described in Enchev et al. (2012) 591 and (Orthwein et al., 2015). Den1 was used in 1:50 ratio for 10 min at 25 °C to remove poly-neddylation. 592 Cull/Rbx1 complexes with mutations in the WHB domain of Cul1 (Figure 1-figure supplement 1N, O) and Cul1/Rbx1^{TEVARING} were purified from High Five insect cells as described in Enchev et al. (2010). 593 594 Dansylation of Cul1/Rbx1 variants expressed in insect cells was performed for 8 to 12 h at 30 °C while 595 spinning at 5000 g, and purified by passing the dansylation reaction through a 5 ml HisTrap FF column (GE Healthcare) in 50 mM Tris-HCl, pH 7.6, 400 mM NaCl, 20 mM imidazole. The Cull^d/Rbx1-596 597 containing flow through was concentrated, neddylated (if required), and further purified over a Superdex 598 200 size exclusion column (GE Healthcare) equilibrated with 15 mM HEPES, pH 7.6, 150 mM NaCl, 2 mM DTT, 2 % (v/v) glycerol. Neddylation of Cul1/Rbx1^{TEVARING} was performed at 25 °C for 12-14 h in 599 600 50 mM Tris-HCl, pH 7.6, 100 mM NaCl, 2.5 mM MgCl₂, 150 µM ATP, spinning at 2000 g, and was 601 followed by 30 min incubation with 1:50 (w/w) Den1 to remove poly-neddylation. The reaction was 602 purified over a Strep-Tactin Superflow Cartridge (QIAGEN), and eluted in 15 mM HEPES, pH 7.6, 250 603 mM NaCl, 2 mM DTT, 2 % (v/v) glycerol, 2.5 mM d-desthiobiotin. RING cleavage was performed for 604 12-14 hours at 25 °C, spinning at 2000 g, in the presence of 100 mM EDTA, pH 8 and 1:1 (w/w) TEV. 605 Dansylation proceeded as described above.

Deneddylation Assays All deneddylation assays were performed in a buffer containing 25 mM Tris-HCl, pH 7.5, 100 mM NaCl, 25 mM trehalose, 1 mM DTT, 1 % (v/v) glycerol, 0.01 % (v/v) Triton X-100 and 0.1 mg/ml ovalbumin or BSA. Radioactive deneddylation reactions with bacterially expressed substrates were done as described (Emberley et al., 2012). Radioactive deneddylation reactions with substrates expressed in insect cells were performed at 24 °C with 0.5 nM CSN (Figure 2–figure supplement 2C) or 2 nM CSN (Figure 5B). All remaining radioactive deneddylation reactions were performed with bacterially expressed Cul1-N8/Rbx1 substrates (50 nM) and 2 nM CSN unless otherwise noted. Single-

turnover reactions were done with 25 nM Cull substrates and 1 μ M CSN on a Kintek RQF-3 Rapid Quench Flow at 24 °C. Single-turnover data were fit to one phase decay function: Y=(Y0 - EP)*exp(k_{cat} *X) + EP (where EP corresponds to reaction end point value), to determine the k_{cat} . Deneddylation assays in Figure 1–figure supplement 1N, O were performed with 800 nM substrate and 20 nM enzyme and visualized by Coomassie stain. Depending on the exact protein preparations used and the laboratory, we observed rates for the wild type reaction ranging from 1.1-2.6 sec⁻¹.

619 Fluorescence Assays All assays were performed in a buffer containing 30 mM Tris pH 7.6, 100 mM 620 NaCl, 0.25 mg/ml ovalbumin or BSA and 0.5 mM DTT with 30 nM dansyl-labeled Cul1/Rbx1 and 621 titrated concentrations of CSN. The mixtures were allowed to reach equilibrium by incubation at room 622 temperature for ~ 10 minutes prior to measurements. Equilibrium binding assays using Cul1/Rbx1 623 variants expressed in insect cells (Figure 2, Figure 2–figure supplement 2E, Figure 3–figure supplement 624 3N, 3R, Figure 4B) were read at 530 nm on a CLARIOstar plate reader (BMG Labtech) in 384-well 625 plates (Corning, low flange, black, flat bottom), 90 ul per well, while binding assays using bacterially 626 expressed Cull/Rbx1 variants were performed on a Fluorolog-3 (Jobin Yvon) (all other binding data figures). Binding assay with Cull^d-N8/Rbx1 (substrate) and CSN^{5E76A} were allowed to equilibrate for 627 only 45 seconds, because although this mutant exhibited an ~300-fold decrease in activity (data not 628 629 shown) the residual activity was high enough to cause substantial deneddylation in a 10 minute incubation. It should be noted that several of the K_d values reported for CSN binding to Cull^d-N8/Rbx1 630 or Cull^d/Rbx1 are below the concentration of the dansylated ligand (30 nM). While this is generally not 631 632 the preferred approach, we found that 30 nM was the lowest concentration that consistently yielded 633 highly reproducible results. The estimated K_d is very sensitive to the density of data points at the 634 inflection point of the curve, and thus these estimates can be more prone to error. Nevertheless, different investigators in Zurich and Pasadena have consistently obtained an estimate of 1.6-5 nM for binding of 635 CSN^{5H138A} to Cull^d-N8/Rbx1 and of 9-13 nM for binding to CSN^{5H138A} to Cull^d/Rbx1, using different 636 protein preparations. To estimate K_d , all data points were fitted to a quadratic equation, Y = Y0+(Ymax-637 638 Y0)*(K_d +A+X-sqrt((K_d +A+X)^2-4A*X))/2*A where A equals concentration of labeled protein, using 639 Prism (Graph Pad). On-rate and off-rate measurements were performed on a Kintek Stopped-flow SF-640 2004 by exciting at 340 nM and collecting emissions through a 520 +/- 20 nm filter. For off-rate 641 measurements, the concentrations of proteins used in each reaction are provided in the legend of Figure 642 3-figure supplements 3F-I. Off rate data were fit to one phase decay function: Y=(Y0 - EP)*exp(k_{off} *X) + EP (where EP corresponds to reaction end point value). Whereas K_d , on-rate, and off-rate 643 644 measurements with different configurations of Cul1 or different CSN mutants are directly comparable,

off-rate measurements are not directly comparable to k_{cat} measurements and may differ from expectation by a few fold because different buffers were used, the Cul1/Rbx1 preparations were from different sources (bacterial for k_{cat} , baculoviral for k_{off}), and the Cul1/Rbx1 preparations carried different labels (dansylated Cul1 for k_{off} , [³²P]-Nedd8 for k_{cat} .

649 Cell Culture and SILAC Mass Spectrometry Cells were grown in Lonza DMEM containing 10% FBS 650 (Invitrogen). Transient transfections were done with FugeneHD per the manufacturers instructions 651 (Roche). Flag-tagged CSN5 coding sequences were cloned into a modified MSCV-IRES-GFP vector 652 (containing a pBabe multiple cloning site) via BamHI and EcoRI. Lenti-CRISPR constructs were made 653 as described (Shalem et al., 2014b) using the targeting sequences 5'-654 CACCGCTCGGCGATGGCGGCGTCC - 3' and 3' - AAACGGACGCCGCCATCGCCGAGC - 5'. 655 Lenti- and retroviruses were produced in 293T cells and the supernatant subsequently used for 656 transduction to establish stable cell lines. For Western Blot analysis cells were directly lysed in 2X SDS 657 sample buffer. Lysates were sonicated for 15 seconds at 10 % of maximum amplitude using a Branson Digital Sonifier and boiled for 10 minutes at 100 °C. SILAC labeling was in Invitrogen DMEM 658 containing 10% FBS and ¹³C₆¹⁵N₂-lysine and ¹³C₆-arginine from Cambridge Isotope Laboratory. For 659 immunoprecipitations, cells were lysed in Pierce Lysis Buffer containing cOmplete Protease Inhibitor 660 661 Cocktail (Roche) and lysates were sonicated for 10 seconds at 10 % of maximum amplitude using a Branson Digital Sonifier. After a 5 minute clearing at 18000 x g at 4°C, proteins were 662 immunoprecipitated with M2 Flag agarose beads (Sigma) for 30 minutes and prepared for mass 663 664 spectrometry as described in Pierce et al. (2013). Samples were analyzed using an EASY-nLC 1000 665 coupled to an Orbitrap Fusion and analyzed by MaxQuant (v 1.5.0.30).

666 Digested peptides (250 ng) were loaded onto a 26-cm analytical HPLC column (75 µm ID) 667 packed in-house with ReproSil-Pur C₁₈AQ 1.9 µm resin (120 Å pore size, Dr. Maisch, Ammerbuch, 668 Germany). After loading, the peptides were separated with a 120 min gradient at a flow rate of 350 669 nL/min at 50°C (column heater) using the following gradient: 2-6% solvent B (7.5 min), 6-25% B (82.5 670 min), 25-40% B (30min), 40-100% B (1min), and 100% B (9 min) where solvent A was 97.8% H₂O, 2% 671 ACN, and 0.2% formic acid) and solvent B was 19.8% H₂O, 80% ACN, and 0.2% formic acid. The 672 Orbitrap Fusion was operated in data-dependent acquisition (DDA) mode to automatically switch 673 between a full scan (m/z=350-1500) in the Orbitrap at 120,000 resolving power and a tandem mass 674 spectrometry scan of Higher energy Collisional Dissociation (HCD) fragmentation detected in ion trap 675 (using TopSpeed). AGC target of the Orbitrap and ion trap was 400,000 and 10,000 respectively.

676 SILAC MS data analysis. Thermo RAW files were searched with MaxQuant (v 1.5.3.8) (Cox and Mann, 677 2008; Cox et al., 2011) Spectra were searched against human UniProt entries (91 647 sequences) and a 678 contaminant database (245 sequences). In addition, spectra were searched against a decoy database 679 (generated by reversing the target sequences) to estimate false discovery rates. Trypsin was specified as 680 the digestion enzyme with up to two missed cleavages allowed. Variable modifications included 681 oxidation of methionine and protein N-terminal acetylation. Carboxyamidomethylation of cysteine was 682 specified as a fixed modification. SILAC was specified as the quantitation method with Arg6 and Lys8 683 specified as the heavy labeled amino acids. Precursor mass tolerance was less than 4.5 ppm after 684 recalibration and fragment mass tolerance was 0.5 Da. False discovery rates at the peptide and protein 685 levels were less than 1% as estimated by the decoy database search. Ratios were calculated for proteins 686 quantified in at least two of the four biological replicates. 95% confidence intervals and adjusted p-687 values were calculated using the R package limma (Ritchie et al., 2015)

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689 *Cross-linking coupled to mass spectrometry (XL-MS)* Chemical cross-linking of purified complexes was 690 performed using DSS H_{12}/D_{12} (Creative Molecules) as cross-linking agent and as previously described 691 (Birol et al., 2014). Subsequent MS analysis and cross-link assignment and detection were carried 692 out essentially as described (Leitner et al., 2014) on an Orbitrap Elite (Thermo Scientific) using the 693 *xQuest/xProphet* software pipeline.

Western Blot Analysis Proteins were separated by SDS-PAGE gel electrophoresis and transferred to a
nitrocellulose membrane by wet blot. Primary antibodies used for detection were: anti-CSN5 mouse
monoclonal Santa Cruz Biotechnology sc-393725, anti-Cul1 mouse monoclonal Santa Cruz
Biotechnology sc-17775, anti-Cul2 rabbit polyclonal Thermo Scientific #51-1800, anti-Cul3 rabbit
polyclonal Cell Signaling #2769, anti-Cul4A rabbit polyclonal Cell Signaling #2699, anti-Cul5 rabbit
polyclonal Bethyl Laboratories A302-173A, anti-β-actin mouse monoclonal Sigma A5316, anti-GFP
mouse monoclonal Clontech #632381.

Sample preparation for electron microscopy and data collection. CSN^{SH138A} -SCF-Nedd8^{Skp2/Cks1} samples for cryo-electron microscopy were generated by pre-incubating the purified components as described in Enchev et al (2012) and ran over a Superose 6 increase 3.2/300 column (GE Healthcare) at 4 °C, eluting 50 µl fractions in 15 mM HEPES, pH 7.6, 100 mM NaCl, 0.5 mM DTT. The sample was kept on ice and its homogeneity and mono-dispersity from the peak elution was immediately confirmed by visualization in negative stain. For cryo EM preparation, the sample was diluted to 0.1 mg/ml and 2 µl were applied to Quantifoil grids (R1.2/1.3 Cu 400 mesh), freshly coated with an extra layer of thin carbon and glow-

discharged for 2 min at 50 mA and 0.2 mbar vacuum. The grids were manually blotted to produce a thin sample film and plunge-frozen into liquid ethane. Data were collected automatically using EPU software in low dose mode on a Titan Krios transmission electron microscope, equipped with a Falcon II direct electron detector (FEI), and operated at 300 kV, an applied nominal defocus from -2.5 to - 5.0 μ m in steps of 0.25 μ m, and 80,460-fold magnification, resulting in a pixel size of 1.74 Å on the sample scale. Images were collected as seven separate frames with a total dose of 25 e⁻/Å².

714 *Electron microscopy data analysis.* CTF-estimation and subsequent correction were performed using 715 RELION (Scheres, 2012) and CTFFIND3 (Mindell and Grigorieff, 2003). All micrographs were initially 716 visually inspected and only those with appropriate ice thickness as well as Thon rings in their power 717 spectra showing regularity and extending to 6 Å or beyond were used for subsequent analysis. In order 718 to generate 2D references for automated particle selection, $\sim 4,000$ single particles were manually 719 picked and subjected to 2D classification in RELION. Six well-defined 2D class averages were selected, 720 low-pass filtered to 35 Å to prevent reference bias, and used as references. Approximately 150,000 721 single particles were automatically selected and subjected to reference-free 2D and 3D classification, in 722 order to de-select the particles, which resulted in poorly defined or noisy averages. Approximately half 723 of these single particles resulted in a well-defined 3D class average, which resembled the previously 724 published negative stain EM map of the same complex (Enchev et al., 2012). This dataset was subject to 725 3D auto-refinement in RELION, using a version low-pass filtered to 50 Å as an initial reference. The 726 converged map was further post-processed in RELION, using MTF-correction, FSC-weighting and a 727 soft spherical mask with a 5-pixel fall-off.

Modeling, docking and visualization. Csn7b was modeled using Csn7a as a template on the Phyre2
server (Kelley et al., 2015) and the modeled coordinates were aligned to Csn7a in PDB ID 4D10
(Lingaraju et al., 2014), effectively generating a CSN atomic model for the Csn7b-containing complex.
Model visualization, molecular docking, distance measurements and morph movie generation were
performed with UCSF Chimera (Pettersen et al., 2004).

Accession code. The cryo electron microscopy density map of CSN^{Csn5H138A}-SCF-Nedd8^{Skp2/Cks1} is
 deposited in the Electron Microscopy Data Bank under accession code EMD-3401.

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866 Figure Legends

Figure 1: Cryo-electron microscopy of a CSN-SCF complex. (A) Molecular model of CSN^{5H138A-}
SCF-N8^{Skp2/Cks1} docked into the cryo-electron density map (gray mesh). (B) Close-up view of the
model, showing the observed conformations of Csn2, Csn4, Rbx1, Csn5/6 and WHB-Nedd8 and (C)
a cartoon representation of the differences between the apo CSN and substrate-bound state.

871 Figure 2: Development and validation of a binding assay for CSN-Cul1/Rbx1 interaction. (A) 872 Equilibrium binding of CSN to Cul1^d/Rbx1 and competition by unlabeled Cul1/Rbx1. The indicated 873 proteins were mixed and allowed to equilibrate prior to determination of dansyl fluorescence in a 874 fluorometer. Note that Cul1/Rbx1 blocks the fluorescence enhancement caused by CSN. CSN, 875 Cul1^d/Rbx1, and Cul1/Rbx1 were used at 350, 30, and 4000 nM, respectively. (B) Equilibrium 876 binding of CSN to Cul1^d/Rbx1. Cul1^d/Rbx1 (30 nM) was mixed with increasing concentrations of 877 CSN and the proteins were allowed to equilibrate prior to determining the change in dansyl 878 fluorescence in triplicate samples. Error bars represent standard deviation.

879 Figure 3: Quantitative determination of enzyme-substrate binding affinities for wild type and mutant proteins. (A) Tight binding of CSN^{5H138A} to substrate. Cul1^d-N8/Rbx1 and CSN^{5H138A} 880 881 were mixed and allowed to equilibrate prior to determining the change in dansyl fluorescence. (B) Summary of K_d measurements for the indicated CSN complexes tested against unmodified 882 883 Cul1^d/Rbx1, Nedd8-conjugated Cul1^d-N8/Rbx1 or Cul1^d-N8/Rbx1^{ΔRING} ligand. Boxes shaded in 884 gray indicate combinations that could not be analyzed due to deneddylation during the binding 885 reaction. For some complexes that bound weakly it was not feasible to titrate to saturation and so a lower boundary for K_d is indicated. N.D., not determined. *, due to the configuration of our assay, 886 887 extremely low K_d values cannot be reliably determined. (C) Summary of k_{on} and k_{off} measurements for the indicated CSN complexes tested against Cul1^d/Rbx1 or Cul1^d-N8/Rbx1. Each reported k_{off} is 888 For comparison, k_{off} values calculated from k_{on} and K_d 889 the mean of at least 8 replicates. measurements are also shown. For cases where k_{on} was not measured (marked with asterisks), 890 we assumed a value that was the average (1.83 x 10^7 M⁻¹ sec⁻¹) of the three measured k_{on} values. 891 892 Boxes shaded in gray indicate combinations that could not be analyzed due to deneddylation upon 893 complex formation. N.D., not determined.

894 Figure 4: The N-terminal domains of Csn2 and Csn4 and the RING domain of Rbx1 play key

roles in substrate binding and deneddylation. (A) Generation of Cul1-N8/Rbx1^{ΔRING}. Top: a TEV

896 protease site was engineered between the N-terminal β -strand and the RING domain of Rbx1 as

indicated. Only the first 50 amino acids of Rbx1 are shown. Bottom: Purified protein was subjected to the indicated treatments (see Materials and Methods for details) and reactions were fractionated by SDS-PAGE and stained with Coomassie Blue. (B) Deletion of the Csn4-NTD and Rbx1-RING domains independently reduce affinity of CSN for substrate. The indicated proteins were mixed and allowed to equilibrate prior to determining the change in dansyl fluorescence in triplicate samples. Error bars represent standard deviation. K_d values measured in this experiment are also reported in Figure 3B.

904 Figure 5: The N-terminal domains of Csn2 and Csn4 and the RING domain of Rbx1 are 905 important for CSN-mediated deneddylation. (A) Summary of kinetic parameters for the 906 indicated CSN mutants in multi- or single-turnover deneddylation reactions with Cul1-N8/Rbx1 or 907 Cul1-N8/Rbx1^{Δ RING} substrate. Note that there may be modest discrepancies between these k_{cat} values and k_{off} values due to differences in assay configurations as described in Materials and 908 909 Methods. The Δ RING substrate used here and in panel B contains the sortase sequence at the C-910 terminus of Cul1 that was used for generation of dansylated Cul1. Control experiments revealed 911 that this tag, with or without dansylation, reduced k_{cat} by ~4-fold. In addition the wild type control 912 for the Δ RING reaction exhibited k_{cat} of 2.6 sec⁻¹. The rates shown have been correspondingly 913 adjusted to normalize them to other rates reported here. *, This rate is estimated from Figure 5-914 figure supplement 5B. (B) Kinetic analysis of deneddylation of Cul1-N8/Rbx $1^{\Delta RING}$ by CSN.

915 Figure 6: Functional analysis of Csn5 active site and INS1 mutants in biochemical and 916 cellular assays. (A) Csn5-E104 is important for CSN function in cells. CSN5 alleles in HEK293T 917 cells were partially knocked out (KO) by CRISPR/Cas9 to yield a major decrease in Csn5 that was 918 nonetheless compatible with viability. Wild type and the indicated Flag-tagged CSN5 mutants were 919 reintroduced by transduction of recombinant retroviruses that co-expressed GFP. Lysates of 920 transduced cells were separated by SDS-PAGE and blotted with antibodies to the indicated 921 proteins. CSN5 long refers to a long exposure of the Csn5 blot, captured to reveal residual Csn5 in the knock-out cells. # refers to transduced FlagCsn5 and * refers to endogenous Csn5. (B) Any 922 923 mutation of a core JAMM domain residue in Csn5 results in enhanced binding to Cul1. Same as (A), 924 except additional Csn5 mutants were tested and the cell lysates were immunoprecipitated with 925 anti-Flag and the immunoprecipitates were blotted for the indicated proteins. (C) Csn5-T103 is 926 important for CSN function in cells. Same as (A) except that the Csn5-T103I mutant was analyzed 927 in parallel with Csn5-E104A and wild type. (D) SILAC mass spectrometry of endogenous proteins

928 bound to FlagCsn5-E104A or FlagCsn5-T103I, relative to wild type FlagCsn5. Cells expressing mutant 929 and wild type FlagCsn5 proteins were grown in light and heavy medium, respectively. L:H ratios >1 930 indicate higher recovery of the listed protein from cells expressing mutant FlagCsn5, whereas ratios 931 <1 indicate higher recovery from cells expressing wild type FlagCsn5. Gray bars: FlagCsn5-E104A; 932 black bars: FlagCsn5-T103I. Error bars represent the 95% confidence interval as calculated by 933 limma (Smyth, 2005). Each protein was quantified in at least two of the four biological replicates 934 and error bars represent standard deviations. Ratios indicated by * differed significantly from 1.0 935 (p<0.05). For CSN, only Csn5 is shown; the remainder is shown in Figure 6–figure supplement 6B.

- 936 Figure 7: Structural and kinetic models for CSN activation and the CSN enzyme cycle. (A-C) 937 Proposed conformational changes that precede substrate cleavage. (D) Kinetic model for the 938 deneddylation cycle. Substrate cleavage is indicated by the slash between N8 and SCF. The asterisk 939 denotes the activated form of CSN. Numbers in red. black, green, and blue represent k_{off} (sec⁻¹), k_{on} 940 $(M^{-1} \text{ sec}^{-1})$, k_{cat} (sec⁻¹), and conformational change (sec⁻¹) rates, respectively. For rates >1, the 941 actual rate has not been measured but it is inferred to be >1 sec⁻¹ because the overall rate for 942 multiturnover catalysis is at least 1.1 sec⁻¹ and thus all sub-steps must be at least this fast. The k_{off} 943 of SCF from CSN varied depending upon whether the rate was measured directly or inferred from K_d and k_{on} (see Fig. 3C). The arrow connecting CSN and N8-SCF•CSN* combines two separate 944 945 steps: binding of N8-SCF to CSN, and activation of CSN to CSN*.
- 946 Figure 1-figure supplement 1: Crvo-electron microscopy and single particle analysis of a 947 **CSN**^{5H138A}-**N8**-SCF^{Skp2/Cks1} complex. (A) A representative cryo-electron micrograph of a 948 CSN^{5H138A}-N8-SCF^{Skp2/Cks1} complex with some single molecular views indicated by white circles 949 (left) and a power spectrum indicating Thon rings reaching 6 Å (right). Scale bar is 200 Å. (B) 950 Representative two-dimensional class averages from the curated dataset, used for the subsequent 951 analysis. Scale bar is as in (A). (C) Surface views of the final, post-processed cryo-electron map. (D). 952 Resolution estimate according to the FSC criteria of 0.143 and 0.5. (E) Fit of the PCI-domain 953 containing CSN subunits in the cryo-electron density map. Csn1,3,7, and 8 match the density very 954 well but the N-terminal domains of Csn2 and Csn4 do not, but their winged-helix domains fit well. 955 The horseshoe arrangement of the six winged-helix domains is indicated with a dotted black line. 956 (F) All the C-terminal helices of the CSN subunits match well the electron density map. (G) Fit of 957 SCF in the electron density map. (H) Same view as in Fig 1B but prior to flexible docking of the N-958 terminal domains of Csn2 and Csn4, the MPN domains of Csn5&6, the WHB domain of Cul1, and

959 Nedd8. (I) Movement of the N-terminus of Csn2 from its crystallographically-determined position 960 (left) into the EM density map (right). (J) Movement of the N-terminus of Csn4 from its 961 crystallographically-determined position (left) into the EM density map (right). The two N-962 terminal helical repeats of Csn4, red arrow, are in close proximity to the WHA domain of Cul1 963 (green circle). (K) Localization of the RING domain of Rbx1. The unfilled density that is indicated 964 by a black ellipse in the right-hand panel of S1I accommodates Rbx1 (shown in red). The helices of 965 Csn4 in close proximity to the RING domain of Rbx1 are indicated by a black arrow. (L) Re-966 localization of Csn5/6. Comparing the left and right panels, Csn5/6 move leftward to occupy 967 unfilled density. The tan and green circles below Csn5/6 indicate densities that are occupied by 968 Nedd8 and the WHB domain, as depicted in (M). (N, O) Deneddylation assays with (N) wild type 969 Cul1-N8/Rbx1 and indicated CSN variants and (O) wild type CSN and mutant Cul1 variants. Note 970 that all Cul1 constructs have an uncleaved C-terminal sortase tag, which is the reason for slower 971 deneddylation of wild type Cul1-N8/Rbx1 by wild type CSN relative to the kinetics reported 972 elsewhere in this work.

973 Figure 2-figure supplement 1: Supporting data for development and validation of CSN-974 **Cul1**^d/**Rbx1 binding assay.** (A) Dansylation of Cul1/Rbx1 constructs. Upper panel: dansylation of 975 bacterially expressed and purified Cul1/Rbx1. Lower panel: dansylation of Cul1/Rbx1 expressed and purified from insect cells. For details, see Materials and Methods. (B) Ubiquitination of ³²P-976 labeled β -catenin substrate peptide by dansvlated SCF^{β -TrCP} was monitored as described (Saha and 977 978 Deshaies, 2008). The k_{cat} measured here (0.048 min⁻¹) compares favorably with that previously 979 determined for wild type unmodified SCF (0.054 min⁻¹) (Saha and Deshaies, 2008). (C) IC_{50} study 980 of the inhibitory effects of unlabeled (red) or dansylated (black) product. Cul1/Rbx1 and 981 Cul1^d/Rbx1 were separately titrated into a deneddylation reactions containing 50 nM Cul1-982 ^{[32}P]N8/Rbx1 substrate and 0.5 nM CSN, and the resulting reaction rate was measured. (D) CSN 983 preparations used in this study. 600 ng of each sample were fractionated by SDS-PAGE and stained 984 with SYPRO Ruby. (E) *IC*₅₀ for competitive inhibition of CSN-Cul1^d/Rbx1 complex formation by 985 unlabeled Cul1/Rbx1 (~ 390 nM) agrees with the K_d measured for binding of Cul1^d/Rbx1 to CSN (310 nM). (F) Equilibrium binding of 100 nM CSN to 50 nM Cul1^d/Rbx1 and competition by 500 986 987 nM Cand1. The indicated proteins were mixed and allowed to equilibrate prior to determination of 988 dansyl fluorescence. (G-I) Free Nedd8 and F-box box proteins do not appreciably change affinity of 989 Cul1^d/Rbx1 for CSN. Same as Figure 2C, except that either 5 µM free Nedd8 (G), 100 nM

Skp2/Skp1 (H) or 100 nM Fbxw7/Skp1 (I) was included in the binding reaction. All binding and
activity measurements reported in this legend were carried out in triplicate and error bars
represent standard deviation.

993 Figure 3-figure supplement 1: Supporting experiments and titration curves for binding 994 data in Figures 3B-C. (A) CSN^{5H138A} is inactive and is a dominant-negative inhibitor of 995 deneddylation. CSN, CSN^{5H138A}, and substrate were used at 2 nM, 100 nM, and 75 nM, respectively. 996 For reactions containing with CSN and CSN^{5H138A}, mutant enzyme was preincubated with substrate 997 for 30 sec prior to initiating time-course by adding CSN. (B-E): The indicated proteins were mixed 998 and allowed to equilibrate prior to determining the change in dansyl fluorescence. (B) CSN^{5H138A} 999 and dansylated, Nedd8-conjugated SCF^{Skp2}. (C) CSN^{5H138A} and dansylated, Nedd8-conjugated 1000 SCF^{Fbxw7}. Note that addition of Fbxw7–Skp1 greatly increased the variability in the measurement for unknown reasons. (D) CSN^{5H138A} (first prep) and Cul1^d/Rbx1, (E) CSN^{5H138A} (second prep) and 1001 1002 Cul1^d/Rbx1, (F-I): The indicated CSN complexes were preincubated with Cul1^d/Rbx1 for 10 min, 1003 followed by addition of unlabeled Cul1/Rbx1 chase and measurement of the decay in dansyl 1004 fluorescence over time. Final protein concentrations are listed for each experiment. (F) CSN (2000 nM), Cul1^d/Rbx1 (200 nM), and Cul1/Rbx1 (3000 nM), (G) CSN^{5E104A} (600 nM), Cul1^d/Rbx1 (200 1005 1006 nM), and Cul1/Rbx1 (3000 nM), (H) CSN^{5E76A,5H138A} (400 nM), Cul1^d/Rbx1 (200 nM), and Cul1/Rbx1 (3000 nM). (I) CSN^{5E76A,5H138A} (200 nM). Cul1^d-N8/Rbx1 (100 nM). and Cul1/Rbx1 1007 1008 (1500 nM), (I-S): The indicated proteins were mixed and allowed to equilibrate prior to determining the change in dansyl fluorescence. (J) CSN^{5E76A, 5H138A} and Cul1^d/Rbx1, (K) CSN^{5H138A} 1009 or CSN^{2ΔN,5H138A} and Cul1^d-N8/Rbx1, (L) CSN^{4ΔN,5H138A} and Cul1^d/Rbx1, (M) CSN^{4ΔN,5H138A} and 1010 1011 Cul1d-N8/Rbx1, (N) CSN^{5E76A} or CSN^{5E76A,5H138A} and Cul1d-N8/Rbx1^{ΔRING}, (O) CSN^{5E104A} and 1012 Cul1^d/Rbx1, (P) CSN^{5E76A} and Cul1^d/Rbx1, (O) CSN^{5H138A,6ΔINS2} and Cul1^d/Rbx1, (R) CSN^{5H138A,6ΔINS2} 1013 and Cul1^d-N8/Rbx1^{ΔRING}, (S) CSN5^{T103I} and Cul1^d/Rbx1. All measurements in panels B-E and J-S 1014 were carried out in triplicate and error bars represent standard deviation. The measurement in 1015 panel P was performed in duplicates but the experiment was repeated on three independent 1016 occasions, obtaining similar results. Several of these results were independently confirmed in 1017 Zurich and Pasadena including panels J, M, O, P, and Q.

Figure 4-figure supplement 1: Biochemical characterization of Cul1/Rbx1^{TEVΔRING} proteins.
(A) Ubiquitination assay using the indicated Cul1-N8/Rbx1 variants (500 nM each) as an E3. Each
reaction contained, in addition, 100 nM Ube1, 1000 nM Cdc34b, 750 nM Skp1/Fbxw7 and 4000

nM CyclinE phosphopeptide, labeled with FAM. The samples were incubated at 25°C for the
indicated time points, analyzed by SDS PAGE and visualized by excitation at 473 nm. (B) Overlay
of Superdex 200 size exclusion profiles of purified Cul1/Rbx1 variants isolated from insect cells.

1024 Figure 5-figure supplement 1: Kinetic analysis of deneddylation. (A) Deneddylation reactions 1025 were carried out in triplicate with CSN^{4ΔN} at varying concentrations of Cul1-[³²P]N8/Rbx1 substrate and quantified to generate the curve shown. Estimates of k_{cat} and K_M are indicated. (B) 1026 1027 Deneddylation assays of Cul1-N8/Rbx1^{ΔRING} (100 nM), incubated with CSN (200 nM, upper panel) 1028 or CSN64INS2 (200 nM, lower panel). Samples were taken at the indicated time points, and 1029 visualized by SDS PAGE and Sypro Ruby staining. Note that the Δ RING substrate contained an unreacted Sortase tag at the C-terminus of Cul1 that reduced k_{cat} by ~4-fold. (C) Multi-turnover 1030 deneddvlation reactions were carried out with CSN or CSN^{5E104A} and Cul1-[³²P]N8/Rbx1. 1031 1032 Substrate was assaved at 1 and 1.3 uM to confirm that saturation was achieved. (D) Single-1033 turnover deneddylation reactions were carried out with CSN on Cul1-[32P]N8/Rbx1 +/-Skp1/Skp2, and with CSN^{5E104A} on Cul1-[³²P]N8/Rbx1. (E) Same as panel D except that CSN^{6ΔINS2} 1034 1035 was also evaluated. (F) Multi-turnover deneddylation reactions were carried out in triplicate with CSN^{5T1031} at varying concentrations of Cul1-[³²P]N8/Rbx1 substrate and quantified to generate the 1036 1037 curve shown. Estimates of k_{cat} and K_M are indicated.

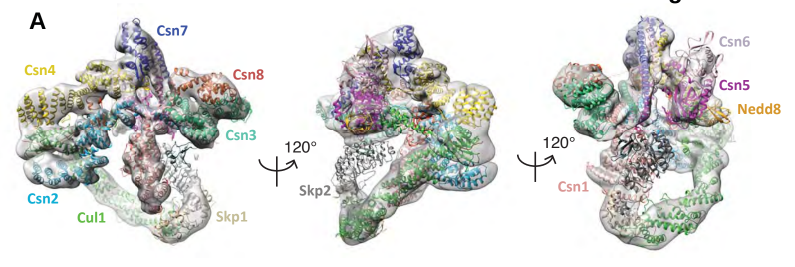
Figure 6-figure supplement 1: Time course and titration data for Figure 6A and supplementary immunoblot for Figure 6B. (A) Same as Figure 6B except that samples were immunoblotted for different cullins. (B) SILAC data for CSN subunits from pull-down analysis shown in Figure 6D.

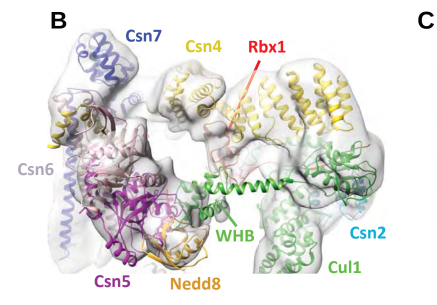
Supplementary File 1: Table S1. Cross-links within CSN5H138A-SCF-N8Skp2/Cks1. "Id" gives the 1042 1043 amino acid sequence of the cross-linked peptides and the exact position of the two cross-linked 1044 lysine residues is indicated by the numbers of the letters *a* and *b* respectively for the first and 1045 second peptide. "Protein1" and "Protein2" denote the cross-linked protein names and "Residue 1" and "Residue 2" respectively defines the position of the cross-linked lysine within the 1046 1047 sequence of the protein. "deltaS" is the delta score of the respective cross-link, which serves as a 1048 measure for how close the best assigned hit was scored in regard to the second best. "Id_Score" is 1049 a weighted sum of four subscores: xcorrc, xcorrx, match-odds and TIC that is used to assess the 1050 quality of the composite MS2 spectrum as calculated by *xQuest*. "FDR" denotes the false-discovery

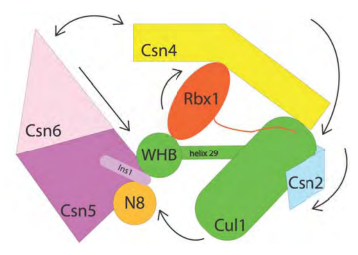
- 1051 rate as calculated by *xProphet*. The measured distance in Å is given for all cross-links, which fall
- 1052 within modeled residues.
- 1053 Table S2. Cross-links within CSN5^{H138A}-SCF^{Skp2/Cks1}
- 1054 Table S3. Cross-links within CSN-Cul1/Rbx1
- 1055 Table S4. Cross-links within CSN- SCF-N8^{Fbxw7FL}
- 1056 Table S5. Cross-links within CSN- SCF-N8^{Fbxw7trunc}
- 1057 Table S6. Cross-links within CSN- Cul3/Rbx1

Movie 1: Morphing CSN and Cul1-N8/Rbx1 conformational changes, occurring upon binding.
 Color code as in Figure 1.

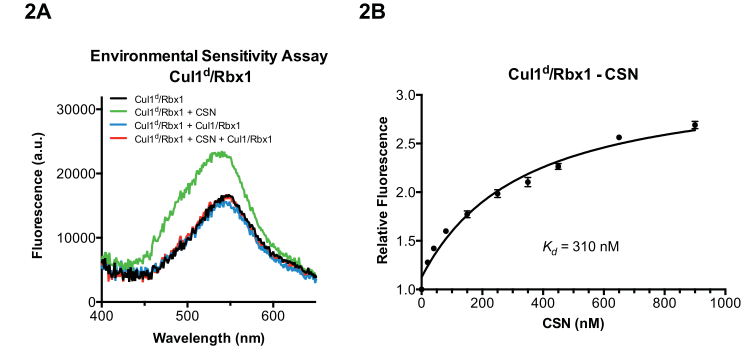
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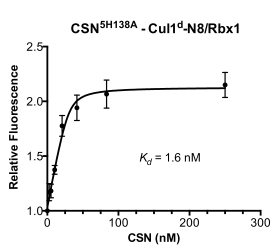






3B





| Ligand | Cul1 ^d /Rbx1 | Cul1 ^d -N8/Rbx1 | Cul1 ^d -N8/Rbx1 ^{ARING} | |
|-------------------------------------|-------------------------|--|---|--|
| wt | 310 | | | |
| 5 ^{H138A} | 10 | 1.6 | N.D. | |
| 5 ^{E76A,H138A} | 13 | N.D. | 12 | |
| 5 ^{E76A} | 13.5 | 0.2* | 13 | |
| 5 ^{E104A} | 26 | | | |
| 2 ^{AN} ,5 ^{H138A} | N.D. | > 1300 | N.D. | |
| 4AN,5H138A | > 750 | 20 | 292 | |
| 5H138A,6AINS2 | 23 | N.D. | 0.4* | |
| 5T1031 | 390 | And Address of the other sectors of the other secto | | |

(nAA)

1.6

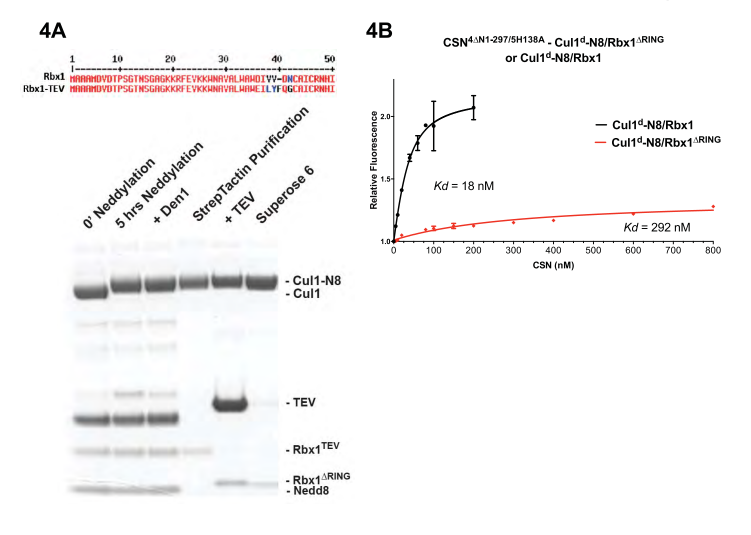
kon (107 M-1 sec-1)

calculated koff (sec-1)

measured koff (sec-1)

| Ligand | Cul1 ^d /Rbx1 | Cul1d-N8/Rbx1 | Cul1 ^d /Rbx1 | Cul1 ^d -N8/Rbx1 | Cul1 ^d /Rbx1 | Cul1 ^d -N8/Rbx1 |
|--------------------|-------------------------|---------------|-------------------------|----------------------------|-------------------------|----------------------------|
| wt | 2 | | 6.2 | | 1.1 | |
| 5 ^{E104A} | N.D. | | 0.48* | | 0.13 | |
| 5 ^{H138A} | 1.3 | 2.2 | 0.13 | 0.035 | 0.16 | N.D. |
| 5E76A,H138A | N.D. | N.D. | 0.26* | N.D. | 0.12 | 0.017 |

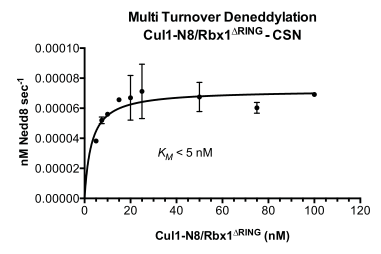
Figure 4

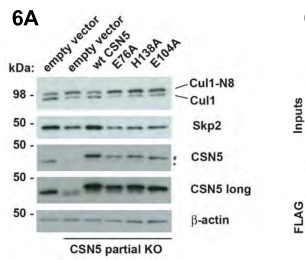


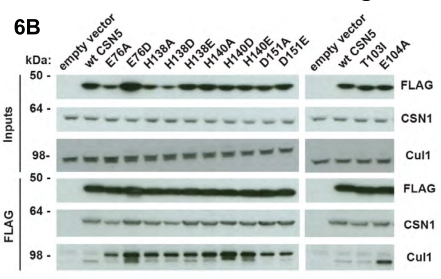
5A

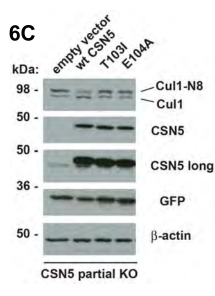
| CSN | k _{cat} (sec ⁻¹) | <i>К_М</i> (nM) | k _{cat} (sec ⁻¹) | k _{cat} (sec ⁻¹) |
|--------------------|---------------------------------------|---------------------------|---------------------------------------|---------------------------------------|
| wt | 1.1 | 200 | 1.1 | 0.00006 |
| 4∆N | 0.05 | 40 | N.D. | N.D. |
| 6∆INS2 | N.D. | N.D. | 1.7 | 0.00048* |
| 5 ^{E104A} | 0.4 | N.D. | 0.4 | N.D. |
| 5 ^{T103I} | ≤ 0.09 | 90 | N.D. | N.D. |
| turnover | ти | ılti | single | multi |
| substrate | | Cul-N8/Rbx1 | | |

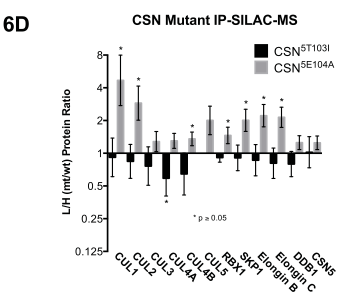


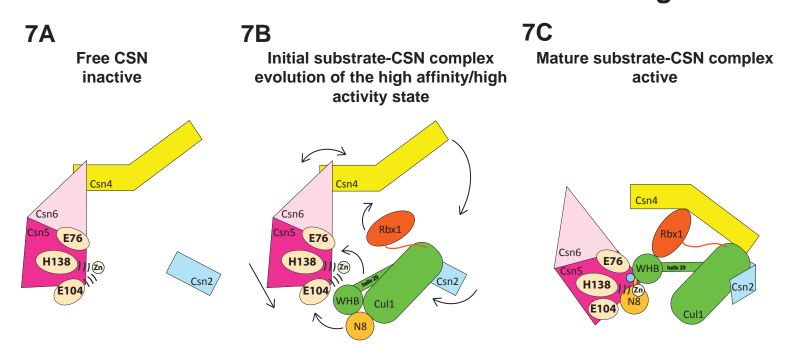




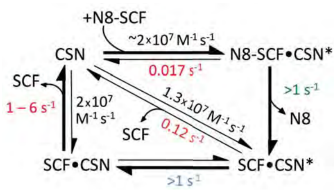




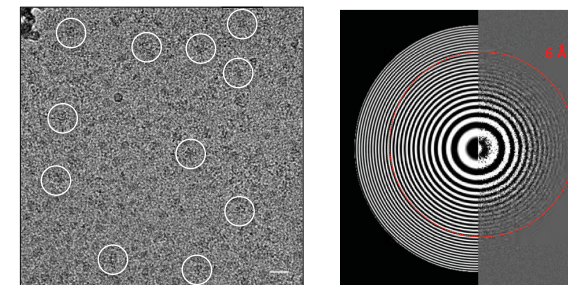




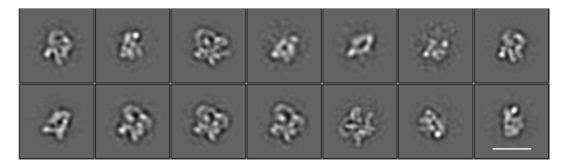
7D







S1B





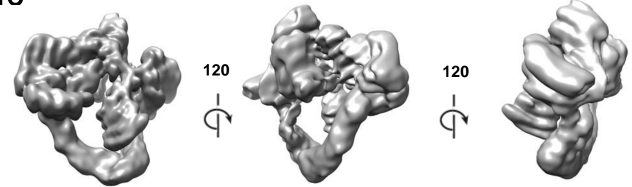
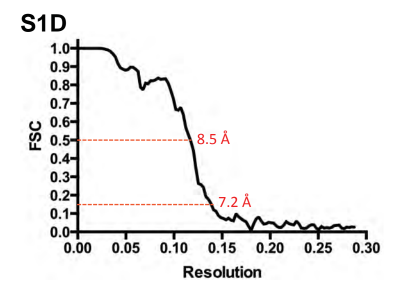
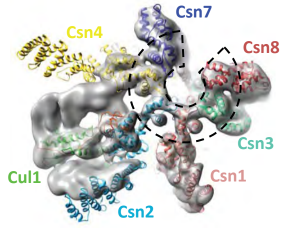


Figure Supplement 1



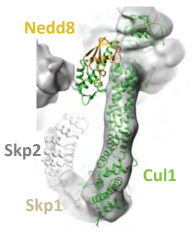
S1E

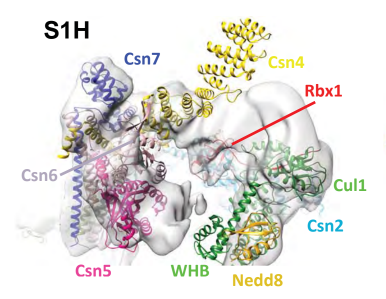












S1I

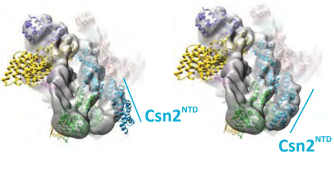
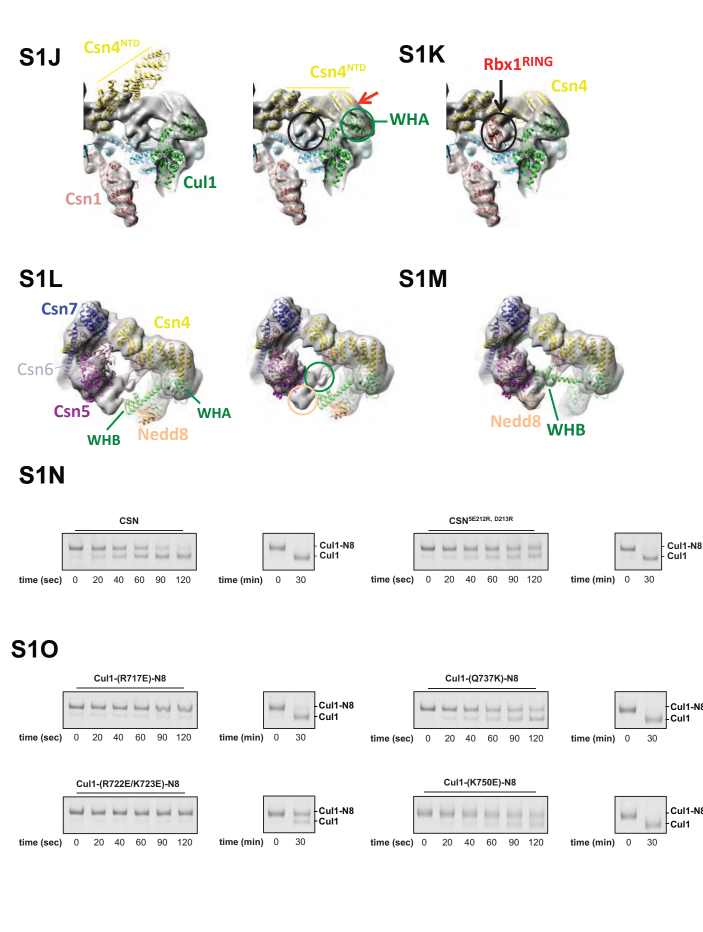
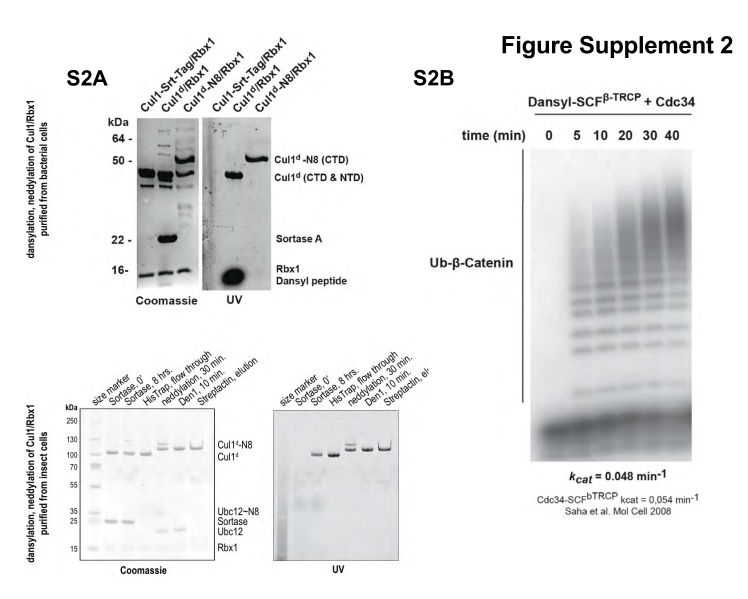


Figure Supplement 1

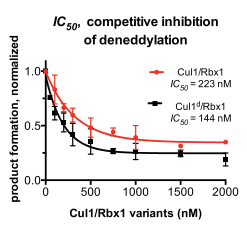
Cul1-N8

Cul1-N8





S₂C



S2D

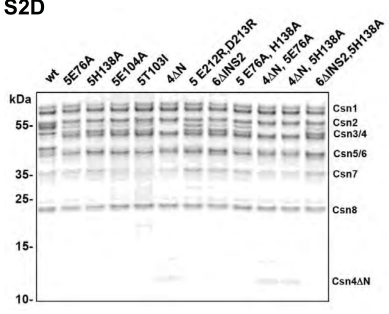
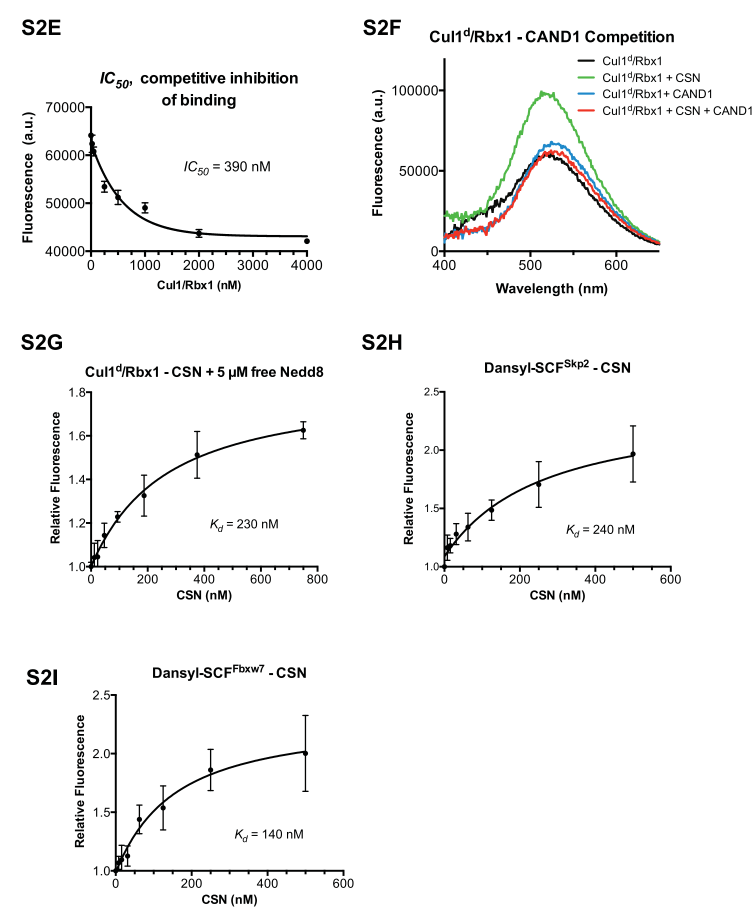
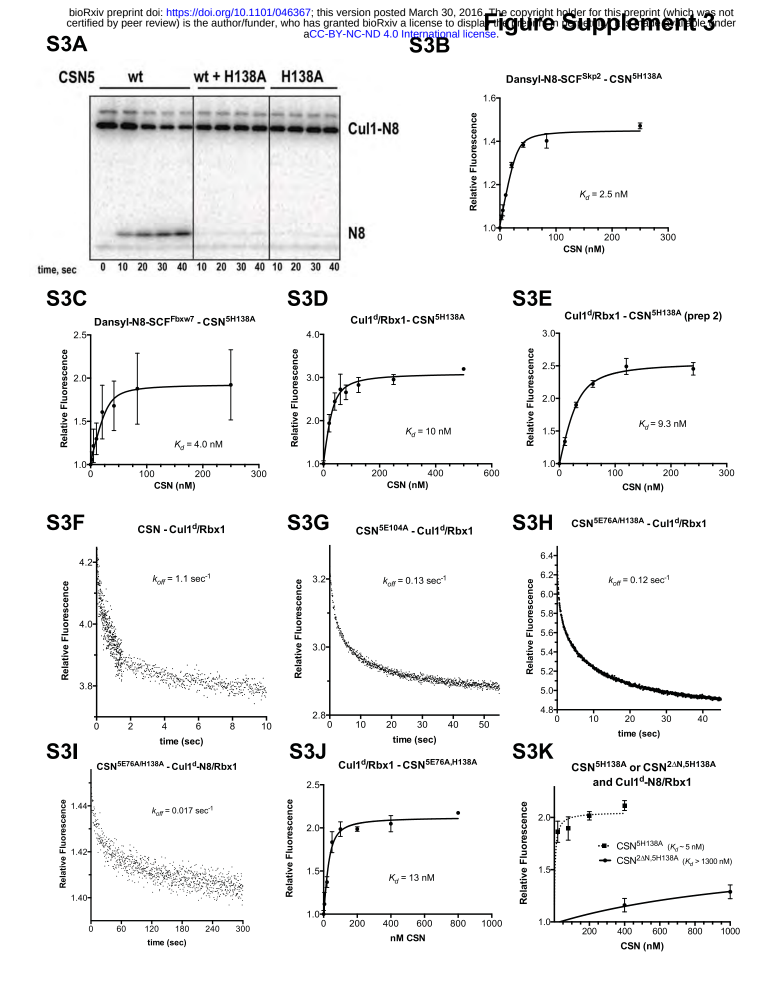
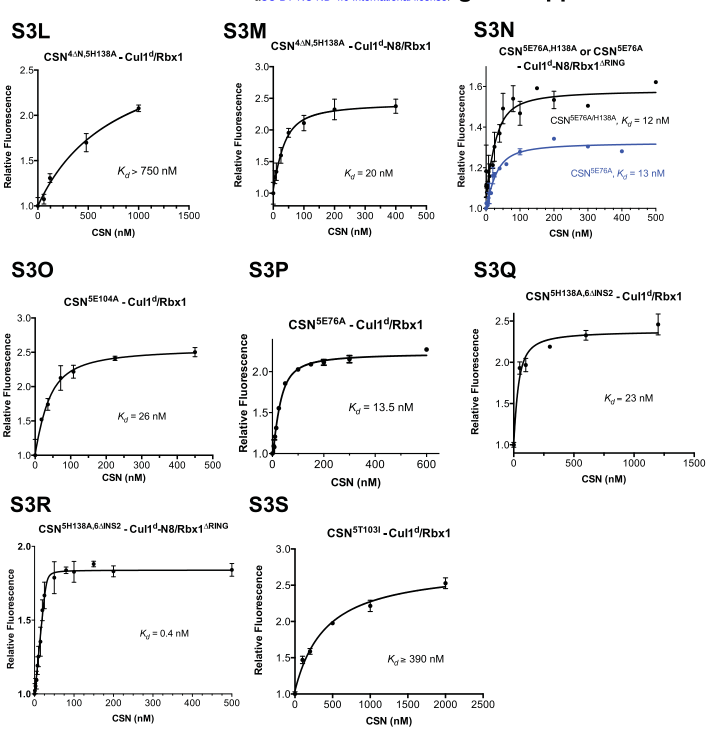


Figure Supplement 2

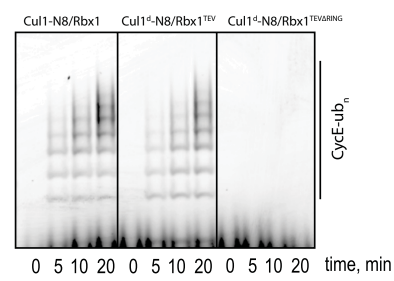




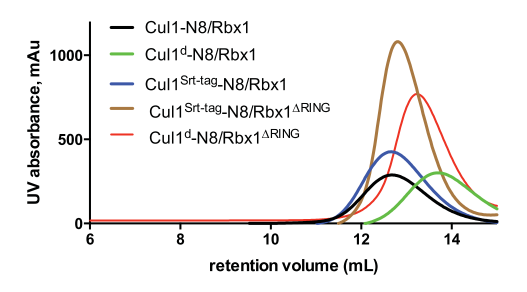
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S4A







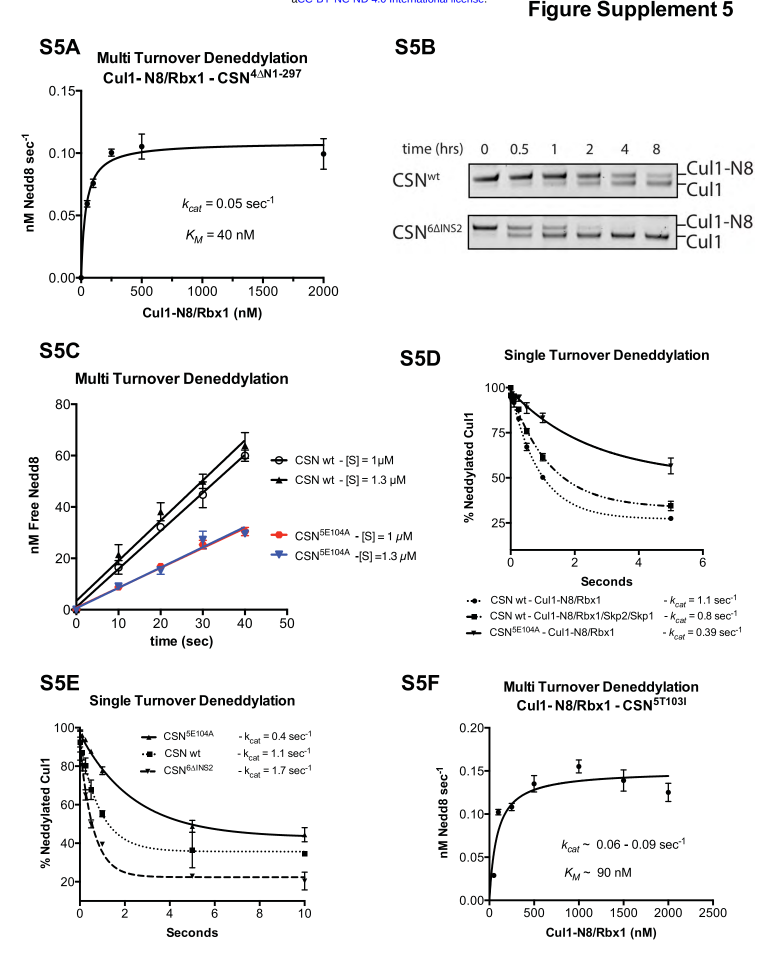


Figure Supplement 6

CSN^{5T103I}

CSN^{5E104A}

CSNIB CSNIR

C_{SN3}

