Visualizing the complex functions and mechanisms of the anaphase promoting complex/cyclosome (APC/C)

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The anaphase promoting complex or cyclosome (APC/C) is a large multi-subunit E3 ubiquitin ligase that orchestrates cell cycle progression by mediating the degradation of important cell cycle regulators. During the two decades since its discovery, much has been learnt concerning its role in recognizing and ubiquitinating specific proteins in a cell-cycle-dependent manner, the mechanisms governing substrate specificity, the catalytic process of assembling polyubiquitin chains on its target proteins, and its regulation by phosphorylation and the spindle assembly checkpoint. The past few years have witnessed significant progress in understanding the quantitative mechanisms underlying these varied APC/C functions. This review integrates the overall functions and properties of the APC/C with mechanistic insights gained from recent cryo-electron microscopy (cryo-EM) studies of reconstituted human APC/C complexes.

1. The anaphase promoting complex or cyclosome regulates cell cycle transitions

The anaphase promoting complex or cyclosome (APC/C) is a multi-subunit cullin-RING E3 ubiquitin ligase that functions to regulate progression through the mitotic phase of the cell cycle and to control entry into S phase [1–4]. The APC/C also plays a role in regulating meiosis, and has been implicated in post-mitotic functions including dendrite formation in neurons, as well as metabolic, learning and memory processes [5–10]. APC/C-mediated coordination of cell cycle progression is achieved through the temporal and spatial regulation of APC/C activity and substrate specificity. The APC/C becomes activated at the onset of mitosis, and ubiquitinates Nek2A and cyclin A (an S- and M-phase cyclin) at prometaphase. At metaphase, the APC/C targets for degradation two inhibitors of the anaphase transition, namely, securin and cyclin B (M-phase cyclin) [11,12]. Securin is a protein inhibitor of separase, a protease that cleaves the cohesin subunit kleisin [13]. Cleavage of kleisin disassembles cohesin to trigger sister chromatid segregation and the onset of anaphase [14–16], reviewed in Nasmyth [17]. Reduced cyclin B levels are also required for entry into anaphase, since Cdk1 (cyclin-dependent kinase 1)-cyclin B1 inhibits separase [18–20]. After anaphase, cyclin destruction continues to maintain negligible Cdk activity, necessary for the cell to disassemble the mitotic spindle and exit mitosis [12,21–25]. During G1, the main role of the APC/C is to sustain low levels of mitotic Cdk activity to allow for resetting of replication origins as a prelude to a new round of DNA replication in S phase [26,27].

The temporal regulation of APC/C activity is achieved through a combination of two structurally related coactivator subunits, Cdc20 and Cdh1 [28–38], coupled to protein phosphorylation, APC/C inhibitors and differential affinity for APC/C substrates. The two APC/C coactivators have opposing activity profiles. Cdc20 activates the APC/C during early mitosis when the APC/C is phosphorylated and Cdh1 activity is low due to its Cdk-dependent phosphorylation, whereas
APC/C-Cdc20-mediated reduction of Cdk activity stimulates Cdh1. In turn, APC/C-Cdh1 ubiquitinates Cdc20, leading to APC/C-Cdc20 inactivation (with Cdc20 auto-ubiquitination also playing a role [39]). Thus, Cdc20 activates Cdh1 that in turn antagonizes Cdc20 activity. The switching between APC/C-Cdc20 and APC/C-Cdh1 fulfils two main functions. First, APC/C-Cdc20 and APC/C-Cdh1 have overlapping but never-related protein Emi2 (XErp1) regulates the APC/C in CCdh1 during interphase [42–44], whereas Acm1 inhibits APC/CCdh1 substrates complex was published [86]. At 7.4 Å

2. The APC/C is a multi-subunit cullin-RING E3 ligase

The large size and complex architecture of the APC/C is intimately linked to its regulatory mechanisms involving control by reversible phosphorylation, the SAC, Emi1 and interchangeable coactivator subunits. These regulatory mechanisms ensure the APC/C is controlled in a cell-cycle-dependent manner and that its substrate specificity is also modulated throughout the cell cycle.

Subunit composition. The APC/C comprises the core complex (14 subunits in metazoans, 13 in yeast) [48–59], together with the interchangeable coactivator subunits (either Cdc20 or Cdh1) [28,29,31] (table 1). APC/C subunits are functionally and structurally organized into three classes: the catalytic module, the substrate recognition module and the scaffolding module (table 1). The catalytic module comprises Apc11, the RING domain subunit [61–63] and Apc2, the cullin subunit [50,51,63]. These two subunits are orthologues of Rbx1 and the cullin subunit of cullin-RING ligases (CRLs), respectively. In both the APC/C and CRLs, an N-terminal β-propeller domain of the RING domain subunit is integrated within the β-sheet of the C-terminal domain (CTD) of the cullin subunit. As discussed below, the catalytic module incorporates two conformationally-variable domains, the RING domain of Apc11 (Apc11RING) and the WHB domain of Apc2 (Apc2WHB), both attached to the CTD of Apc2 (Apc2CTTD) by flexible linkers. The conformational flexibilities of Apc2WHB and Apc11RING have important implications for APC/C catalysis and regulation.

Together, the coactivators and Apc10 form the substrate recognition module, with the coactivator’s WD40 β-propeller domain being primarily responsible for mediating degron recognition (D box, KEN box and ABBA motifs) [64–71]. Optimal D-box recognition requires the core APC/C subunit Apc10 (Doc1 in S. cerevisiae) [54,72,73]. The substrate recognition and catalytic modules represent the key functional subunits of the APC/C, reflected in their high degree of conservation. It is striking that these two functional modules represent only 15% of the total mass of the molecule. Most of the APC/C mass is conferred by the seven large scaffolding subunits, four of which form homo-dimers—further contributing to the high relative mass of the scaffolding module [74]. Remarkably, the majority of APC/C subunits, particularly the scaffolding subunits, are composed of multiple repeat motifs. Five scaffolding proteins are tetratricopeptide repeat (TPR) proteins, being composed of 13–14 TPR motifs arranged in contiguous arrays. TPR proteins, ubiquitous in all three domains of life, were first discovered in what were later identified as yeast APC/C subunits [75–78]. Their presence in multiple protein complexes of diverse functions such as the APC/C indicates a role in mediating protein–protein interactions and the assembly of multi-protein complexes [79]. Later, atomic resolution structural analysis of the APC/C provided a mechanistic rationale for many of the previously characterized TPR mutations [80–83].

The four canonical TPR proteins (Apc3, Apc6, Apc7, Apc8) are structurally highly homologous, being composed almost entirely of 14 TPR motifs. These self associate to form homo-dimers [81–83]. Apc1, the largest APC/C subunit, features another type of motif that is only observed in Apc1 and the Rpn1 and Rpn2 subunits of the 19S regulatory subunit of the proteasome (in exactly the same number and arrangement) [84]. These approximately 40-residue motifs are termed the PC (proteasome-cyclosome) repeat [85]. Although not discernible in sequence, cryo-electron microscopy (cryo-EM) studies revealed that Apc1 contains an N-terminal seven-bladed β-propeller domain [80,86]. Apc4 also comprises a β-propeller domain [87]. Finally, four small intrinsically disordered subunits (vertebrate Apc12, Apc13, Apc15, Apc16) function as TPR-accessory subunits. These subunits interact with TPR subunits and, as explained later, Apc12, Apc13 and Apc16 stabilize TRP subunits and mediate inter-TPR interactions [51,54,56,80,86,88,89]. Apc15 is not required for APC/C assembly. It functions to negatively regulate the SAC by controlling the stability of the Cdc20 subunit of the MCC through APC/C-dependent auto-ubiquitination [59,90–95].

Structural investigations of the APC/C were initiated some 18 years ago, shortly after its discovery in 1995 [1–3]. Initial efforts focused on a complementary approach of crystallography of individual APC/C subunits and small sub-complexes and homologous proteins [71,81–84,87,96–100], together with single particle cryo-EM studies of the intact complex that represented various functional states of the complex purified from endogenous sources: budding yeast, fission yeast, Xenopus and human [73,101–107]. A combination of crystallography of individual APC/C subunits, native mass spectrometry [74] and electron microscopy provided information on the subunit stoichiometry of the APC/C (table 1).

The recent progress in understanding the structure and mechanisms of the APC/C through atomic resolution structures, atomic resolution structures, and atomic resolution structures of various functional states of the complex from technical developments in reconstituting the recombinant APC/C [74,91,108] together with recent advances in single particle cryo-electron microscopy (direct electron detectors and software for image analysis and 3D-reconstructions) [109]. Recent EM studies have focused on reconstituted human APC/C complexes [80,86,92,93,99,110–112].

In 2014 a 7.4 Å resolution structure of the reconstituted APC/C Cdh1 substrates complex was published [86]. At 7.4 Å
Table 1. Subunits of the human anaphase promoting complex/cyclosome (AP/C). Alternative *S. cerevisiae* subunits in parenthesis.

| subunit | length (aa) | stoichiometry | location | domain/Region 1 | domain/Region 2 | domain/Region 3 | phosphorylation sites (from ref. [60]) |
|---------|-------------|---------------|----------|-----------------|-----------------|-----------------|---------------------------------------|
| ApC1    | 1944        | 1             | scaffolding module platform | WD40 domain (1 – 612) | mid-N (613 – 986) | PC domain (1013 – 1616) | 60,65,202,233,286,291,297,298,299,300,313,316,317,341,343,351,355, 362,364,372,373,377,378,389,394,416,501,518,520,522,524,530,535,6,537, 542,547,555,565,566,569,576,582,600,688,689,701,703,731,916, 920,921,922,1001,1347,1349 |
| ApC2    | 822         | 1             | catalytic module | NTD (1 – 432) | cullin repeats | CTD (433 – 822) | including WHB domain | 205,218,314,466,470,474,532,534,732,736,738,742 |
| ApC3A   | 824         | 2             | scaffolding module TPR lobe | TPR dimer interface | TPR motifs 1 – 7 (1 – 535) | TPR superhelix | — | 183,185,186,192,194,200,203,205,209,219,220,222,225,228,230,231,233, 237,241,244,251,252,255,264,267,276,279,281,289,291,302,304,310,312, 313,327,329,331,334,336,338,349,351,352,356,357,358,364,366,368, 369,383,384,386,387,388,389,419,426,430,434,435,438,439,444,446, 761,800,803,806,807,809,814,821 |
| ApC3B   | 824         | 2             | scaffolding module TPR lobe | TPR dimer interface | TPR motifs 8 – 14 (536 – 824) | TPR superhelix | — | — |
| ApC4    | 808         | 1             | scaffolding module platform | WD40 domain/4HBD | — | — | 199,469,488,757,758,777,779 |
| ApC5    | 755         | 1             | scaffolding module platform | NTD (1 – 169) | TPR superhelix | TPR motifs 1 – 13 (206 – 755) | — | 15,110,178,179,195,221,228,232,674 |
| ApC6A   | 620         | 2             | scaffolding module TPR lobe | TPR dimer interface | TPR motifs 1 – 7 (1 – 261) | TPR superhelix | TPR motifs 8 – 14 (262 – 620) | 112,559,573,577,580,584,585,592,599,607,614 |
| ApC6B   | 620         | 2             | scaffolding module TPR lobe | TPR dimer interface | TPR motifs 1 – 7 (1 – 261) | — | — | 119,120,123,125,126,573,582,584 |
| ApC7A   | 599         | 2             | scaffolding module TPR lobe | TPR dimer interface | TPR motifs 1 – 3 (21 – 166) | TPR superhelix | TPR motifs 4 – 7 (167 – 359) | 562,565,582,584,588,593,596 |
| ApC7B   | 597         | 2             | scaffolding module TPR lobe | TPR dimer interface | TPR motifs 1 – 7 (1 – 287) | — | — | 821 |
| ApC10   | 185         | 1             | degron recognition module | Doc homology (2 – 162) | IR tail (163 – 185) | — | — |
| ApC11   | 84          | 1             | catalytic module | Ω-strand (1 – 18) | RING domain (21 – 84) | — | — | (Continued.) |
The secondary structural architecture can be defined. Alpha-helices are resolved as rod-like structures, whereas β-sheets are visualized as planar structures. The subunit assignment of the electron microscopy (EM) density map was determined based on two approaches. One was a subunit deletion approach where the structures of reconstituted APC/C complexes lacking defined subunits were compared with the wild-type complex [74]. Difference density due to the deleted subunit could be assigned to a specific subunit. In a related approach, comparing two complexes that share a common subunit allows its identification. However Apc1, an essential subunit required for APC/C stability, which therefore cannot be deleted without disrupting the entire complex, was identified based on a process of elimination and by recognizing architectural features of the PC domain in the EM density map [80,86]. Finally, Apc13 in *S. cerevisiae* was identified through locating GFP fused to its C-terminus [74]. Importantly, EM density for Apc2CTD was weak and diffuse whereas that for Apc11RING and Apc2WHB was not visible, indicating a high degree of conformational flexibility of the catalytic module. Conformational heterogeneity was also confirmed through 3D classification of the cryo-EM data set [86]. Altogether, the EM studies revealed a striking degree of structural conservation from yeast to metazoan. The APC/C of higher eukaryotes differs from yeast because of an additional TPR subunit (Apc7) situated on the top of the TPR lobe that interacts only with Apc3 (table 1). The role of Apc7 has yet to be defined.

The 7.4 Å resolution structure of the APC/C was soon followed by a near-atomic resolution structure of the complex of APC/C<sub>Cdh1</sub> with the inhibitor Emi1 (APC/C<sub>Cdh1.Emi1</sub>) [80]. This structure was at 3.6 Å resolution overall, but a local resolution map showed that the more rigid regions of the map were closer to 3.2 Å resolution. Two regions in particular were recovered at lower resolution (approx. 5 Å) due to their higher relative flexibility. These were the catalytic module formed of Apc11 and Apc2CTD, and the coactivator Cdh1.

The 3.6 Å resolution cryo-EM map of APC/C<sub>Cdh1.Emi1</sub> provided the basis for understanding the detailed architecture of the APC/C and served as a template for understanding subsequent different functional states, some at lower resolution. Building of the atomic-resolution model was based on fitting of atomic coordinates of X-ray structures of most of the large subunits and close homologues. For Apc1, fitting to the N-terminal WD40 domain and densities adjacent to its central PC domain (Apc1PC) that lack structural homologues was performed *ab initio* [80]. The TPR accessory subunits Apc13, Apc15 and Apc16 were also built *ab initio* [80].

The APC/C adopts a triangular shape delineated by a lattice-like shell organized into two sub-structures (figure 1) [80,86]. The back and top of the complex is formed from a bowl-shaped TPR lobe, an assembly of the four canonical TPR proteins (Apc3, Apc6, Apc7, Apc8) and three TPR accessory subunits (table 1). The base of the APC/C comprises the platform subunits Apc4 and Apc5, together with two (non-PC) domains of Apc1. Apc1PC extends from the platform to contact the TPR lobe. Together, the TPR lobe and platform sub-structures define a central cavity. The degron recognition module of coactivator and Apc10 is located at the top of the cavity with Apc10 interacting extensively with Apc1PC. The catalytic module of Apc2-Apc11 is positioned at the periphery of the platform such that Apc2CTD and associated Apc11 are at the front of the cavity situated directly below Apc10 and Cdh1.

### Table 1.

| subunit  | length (aa) | stoichiometry | stockchemistry | location | domain/Region 1 | domain/Region 2 | domain/Region 3 | phosphorylation sites | (from ref. [60]) |
|----------|-------------|---------------|----------------|----------|-----------------|-----------------|-----------------|---------------------|-------------------|
| Apc12A   | 85          | 2             | 1              | scaffolding module | TPR lobe | extended chain, short α-helix | (1 – 25) Extended chain, short α-helix | 76 S80, 98 | |
| Apc12B   | 85          | 2             | 1              | scaffolding module | TPR lobe | extended chain, short α-helix | (1 – 25) Extended chain, short α-helix | 76 S80, 98 | |
| Apc13    | 74          | 1             | 1              | scaffolding module | TPR lobe | extended chain, short α-helix | (1 – 25) Extended chain, short α-helix | 76 S80, 98 | |
| Apc15    | 121         | 1             | 1              | scaffolding module | platform | extended chain and α-helix | (1 – 56) Extended chain and α-helix | 76 S80, 98 | |
| Apc16    | 110         | 1             | 1              | scaffolding module | TPR lobe | extended chain and α-helix | (1 – 56) Extended chain and α-helix | 76 S80, 98 | |
| Cdc20/Cdh1 | 499/496    | 1             | 1              | degron recognition module | TPR lobe | extended chain and α-helix | (168 – 471/172 – 473) | 493 S498 | |
| UbcH10   | 179         | 1             | 1              | catalytic module | TPR lobe | extended chain and α-helix | (168 – 471/172 – 473) | 493 S498 | |
The canonical TPR proteins form structurally related V-shaped homo-dimers [81–83]. Each subunit comprises an α-helical solenoid with two turns of TPR helix. Whereas the N-terminal TPR helix forms the homo-dimer interface, the C-terminal TPR helix creates a protein-binding groove. Apc6 binds its accessory subunit Apc12 through this groove (figure 1) [82,89], stabilizing Apc6 [89], whereas the Apc3 and Apc8 homo-dimers use one of their dyad-related C-terminal grooves to engage the coactivator subunits (either Cdc20 or Cdh1) (figures 1 and 2) [60,80,86]. Within the TPR lobe, the four canonical TPR proteins stack in a parallel array generating a left-handed super-helix that adopts pseudo dyad-symmetry. Together the TPR accessory subunits Apc13 and Apc16 (and presumably Apc9 in S. cerevisiae) interact with structurally and symmetry related sites on seven of the eight TPR subunits to stabilize the TPR lobe and contribute to defining the order of TPR protein assembly [80].

Apc10 and both coactivators share structurally related C-terminal Ile-Arg motifs (IR tails) that interact with the C-terminal TPR motifs of Apc3 (figures 1 and 2a,b) [66,88,96,113,114]. Additionally, coactivators comprise a C-box motif within their N-terminal domain (NTD) [68] that mediates interactions with the APC/C [68,113], dependent on Apc8 [115]. Due to the presence of multiple binding sites on the TPR lobe, the pseudo dyad-symmetry of the TPR lobe has important consequences for mechanisms of interaction with coactivators and substrates. Not only does the dyad symmetry of each TPR protein mean that there is multiplication of protein/ligand binding sites (for example the common IR tails of coactivator and Apc10 interact with separate subunits of the Apc3 homo-dimer (figure 2a,b)), but also the IR-tail binding site on Apc3 is structurally related to the C-box binding site on Apc8 [80]. Because of this, the structurally equivalent C-box binding site on Apc8A is capable of binding the IR tail of Cdc20MCC (in the APC/CMCC complex) [92,93]. A conformational transition involving the C-terminal TPR motifs of Apc3A occludes the coactivator IR-tail binding pocket in the absence of the IR-tail ligand [60,80,100]. Finally, regions of the NTD of coactivator also interact with Apc1PC (figure 3c). Thus the degron-recognition WD40 domains of the coactivators are connected to the APC/C scaffold through three sites, attached through flexible linkers. This allows for conformational flexibility of the WD40 domain.

In the platform, analogous to the Apc6–Apc12 interaction, the C-terminus of Apc15 inserts into the TPR groove of Apc5 as an extended chain, with its N-terminal α-helix (Apc15VTH) bridging Apc5 and Apc8 [80].

3. Coactivators are primarily responsible for degron recognition

The APC/C recognizes and ubiquitinates a variety of cell cycle substrates in a cell-cycle-dependent manner. Selection of substrates in a temporal manner is dependent on a variety of factors, but critical among these is the role of coactivators [29]. The APC/C is inactive without coactivator. One function of coactivators is to provide degron recognition sites that engage degrons present in most APC/C substrates [66,69–71], thereby recruiting substrates to the APC/C (figures 1, 3a,b, 4 and 5). In a few exceptions, for example Nek2A, the core APC/C recognizes substrates, bypassing degron recognition sites on the coactivator. However, Nek2A ubiquitination still relies upon the coactivator-induced stimulation of UbcH10-binding to the APC/C [86,117].

Due to the critical role coactivators play in defining APC/C activity, regulation of APC/C activity by phosphorylation and
inhibitory complexes such as the MCC, Emi1 and Acm1 is exerted primarily at the level of coactivators, either by controlling their interaction with the APC/C or by controlling coactivator interaction with degrons. Cdc20 activates the APC/C from early mitosis to anaphase after which Cdh1 binds to the APC/C through to late G1. Switching of these two highly structurally conserved and related coactivators at anaphase changes the substrate specificity and regulatory properties of the APC/C. Cdc20 is thought to recognize a restricted set of substrates (specifically cyclin A, cyclin B and securin), whereas Cdh1 is proposed to have a broader substrate specificity, being able to ubiquitinate all Cdc20 substrates, and in addition recognizes the Aurora A and B kinases, which are not substrates of APC/C-Cdc20 [118]. Aurora kinases are recognized by APC/C-Cdh1 through their essential N-terminal A box motif [119]. The role of the C-terminal D box of Aurora kinases is disputed, as discussed in Davey & Morgan [120]. Both coactivators mediate interactions of substrates harbouring D-box and KEN-box motifs to the APC/C. Optimal interactions of the D box also require the Apc10 subunit [54,72,73]. The ABBA motif is recognized by vertebrate Cdc20 [121], and S. cerevisiae Cdh1 [71] and Cdc20 [120,122]. In S. cerevisiae a specific coactivator termed Ama1 controls meiosis [123,124] that in turn is antagonized by the Mnd2 (Apc15) subunit [125,126].

To understand structurally how coactivators recognize D-box and KEN-box substrates, advantage was made of the fact that many APC/C inhibitors incorporate pseudo-substrate motifs that mimic D-box and KEN-box degrons in order to block substrate recognition. These inhibitors interact with higher affinity with coactivators than do substrates, thereby facilitating the biochemical isolation and crystallization of these complexes. A structure of the MCC from Schizosaccharomyces pombe, a complex of Cdc20, Mad2 and BubR1/Mad3, revealed how a KEN box and D box present in BubR1/Mad3 interact with their respective binding sites on the β-propeller domain of Cdc20 [69]. These findings were confirmed and extended in a structure of the β-propeller domain of Cdh1 in complex with Acm1, a Cdh1 specific inhibitor from S. cerevisiae [71]. The latter structure also revealed how the ABBA motif (A motif in Acm1 terminology [127]) interacts with Cdh1 (figures 3 and 4). A further study in which human Cdc20 was crystallized with a peptide modelled on the BubR1 KEN box also revealed details of Cdc20 interactions with the KEN-box motif [70].

**Figure 2.** The IR-tail and C-box binding sites of Apc3 and Apc8 respectively, are homologous. (a) Cdh1 IR-tail binding site. (b) Apc10 IR-tail binding site. (c) C-box binding site on Apc8B. The Ile and Arg side chains of the IR tail of both Cdh1 and Apc10 interact with a site on Apc3 that is homologous to the binding sites for Arg(47) and Ile(49) of the Cdh1 C box on Apc8B. The C box (DF[Y/F][W]R) forms additional contacts to Apc8B as shown. PDB 4UI9, from Chang et al. [80].
The classical APC/C degron is the destruction box or D box, a ten-residue motif (RxxLx[D/E][Ø]xN[N/S]) (figure 5a,c) first characterized in B-type cyclins as being necessary and sufficient for APC/C mediated ubiquitination [128–130]. Mutation of any of the three most highly conserved residues, Arg (P1), Leu (P4) or Asn (P9), ablated the destruction signal [128]. The D box binds in a mainly extended conformation to a shallow groove at the side of the b-propeller, found between the two b-blades 1 and 7 (figures 3b and 4a). The essential Leu (P4) residue anchors the D box to the channel within a hydrophobic pocket, whereas the N-terminal Arg (P1) residue interacts with an acidic pocket at the N-terminus of the channel (figure 4a). A conserved acidic residue at P6 interacts with an invariant Arg, whereas a hydrophobic residue at P7, conserved in many D-box motifs, interacts with a hydrophobic surface of the b-propeller (figures 4a and 5a,c) [69,71]. The side chain of P3 abuts a conserved Phe of the coactivator, likely accounting for the high occurrence of residues with small unbranched side chains at this D-box position (figures 4a and 5a,c). Although Arg and Leu are strongly preferred at P1 and P4, respectively, even these two residues are not strictly necessary. For example, in *Drosophila melanogaster* cyclin A [131] and *Homo sapiens* cyclin B3 [132], Phe is substituted for Leu.

Significantly, the conserved C-terminal hydrophilic residues (P8 to P10) do not interact with the coactivator, however the cryo-EM structure of APC/C-Cdh1.Emi1 (where the inhibitor Emi1 incorporates a D box) showed clear EM density extending from the P7 residue of the D box (interacting with the D-box site on Cdh1) to Apc10 [80]. This showed that the C-terminus of the D box interacts with a hydrophilic surface of Apc10 [80,96,97] involving polar and charged residues on two surface-exposed loops (the 80s and 140s loops) (figure 4b). This highly conserved region is required for D-box-dependent APC/C E3 ligase activity, and this potentially dynamic hydrophilic surface may allow for the accommodation of a variety of small polar residues at D-box positions P8 to P10 (figure 5a). Disruption of the 140s loop impairs D-box-dependent substrate recognition [133] and Ala substitutions of Ser88 and Asn147 of Apc10 attenuated APC/C-Cdh1 activity [80]. Although Apc10 primarily interacts with D-box residues P8 to P10, its 80s loop also contacts N-terminal residues of the D box (figure 4b). For example, the side chain of Glu87 (invariant in Apc10

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**Figure 3.** Coactivators interact with Apc1 and Apc3 and create a D-box co-receptor with Apc10. (a) Overview of the APC/C with the Cdh1 coactivator subunit. Based on the APC/C-Cdh1.Emi1 coordinates (PDB 4UI9) [80] with the KEN box and ABBA motif modelled on the *S. cerevisiae* Cdh1–Acm1 complex (PDB: 4BH6) [71]. Except for the D box, Emi1 coordinates are not shown. (b) Close-up view of the D-box co-receptor formed from Cdh1 and Apc10. (c) Cdk1-dependent phosphorylation of the NTD of Cdh1 blocks its binding to the APC/C. Red spheres indicate sites of inhibitory phosphorylation.
orthologues) is sandwiched between P2, P5 and P7 of the D box, perhaps explaining the occurrence of basic residues at these positions, especially for the non-canonical D-box sequences (discussed below) (figure 5a,c). Notably, Cdk1-phosphorylation at P2 (Pro is common at P3) negatively regulates APC/C-dependent substrate ubiquitination, for example Dbf4 [122], possibly due to the electrostatic repulsion between a phosphate group at P2 and Glu87. Thus, the D box is a bipartite degron comprising a coactivator-interacting N-terminal (RxxLx[D/E][Ø]) motif and a hydrophilic C-terminal-Apc10 binding segment. Coactivator and Apc10 create a D-box co-receptor for recognition of the bipartite degron. The atomic resolution structures of D-box motifs engaged by coactivators alone [69,71] and in complex with APC/C-coactivator complexes [60,80] rationalize the residue preferences at all 10 positions of the D box. Moreover, the preferences for an acidic residue at P6 and basic residue at P2 are consistent with the promotion of substrate ubiquitination by D-box phosphorylation at P6 [134] and substrate stabilization by phosphorylation at P2 [122].

**KEN box.** Another APC/C degron, the KEN motif ([DNE]-KENxxP), is commonly present in APC/C substrates usually in addition to the D box [135]. Efficient ubiquitination by either APC/C/Cdc20 or APC/C/Acm1 of substrates harbouring both D and KEN boxes is dependent on both degrons [54,64]. By forming a 310 helix, the three consecutive residues of the KEN box face in the same orientation and engage the top surface of the β-propeller (figures 3b and 4c) [69–71]. The KEN box is usually immediately C-terminal to acidic residues (figure 5b), and the structure of the KEN box–coactivator complex suggested that these would engage a positively charged patch on the β-propeller. A frequently observed Asp or Asn residue at P-1 stabilizes the KEN box conformation by forming a hydrogen bond to the Asn of the KEN box (figure 4c) [71]. Proline residues one to two residues C-terminal of the KEN box would direct the polypeptide chain away from the surface of the β-propeller.

**ABBA motif.** The A motif was discovered in the *S. cerevisiae* Cdh1 inhibitor Acm1 [127,136]. Later bioinformatics studies identified the ABBA motif as a general class that includes the A motif as a six-residue motif (Fx[ILV][FY][DE]) common to vertebrate cyclin A (and *S. cerevisiae* Clb5), BubR1, Bub1 and Acm1 [120–122]. Although the A motif was originally thought to confer specificity for *S. cerevisiae* Cdh1 [71,127], the situation is more complicated. Cdc20 also binds the ABBA motif—variations in non-consensus residues confer the specificity for *S. cerevisiae* Cdh1. Glu65(P5) of the ABBA motif of Acm1

![Figure 4. Substrate recognition is mediated by coactivators and Apc10. (a) D-box receptor on Cdh1, (b) D-box co-receptor (Cdh1 and Apc10), (c) KEN-box receptor on Cdh1, (d) ABBA-motif interactions with Cdh1. Coordinates in (a,c,d) are based on the *S. cerevisiae* Cdh1–Acm1 complex (PDB: 4BH6) [71]. (b) Based on APC/C(Cdh1Emi1) complex (PDB 4UI9) [80].](image-url)
contacts Lys333 in *S. cerevisiae* Cdh1 that is a Thr in *S. cerevisiae* Cdc20 [121]. Residues of human Cdc20 required for ABBA motif binding are not conserved in human Cdh1 (although are conserved in *S. cerevisiae* Cdh1), explaining the inability of human Cdh1 to recognize the ABBA motif [121]. A structure of Acm1 in complex with *S. cerevisiae* Cdh1 revealed that the ABBA motif forms an extended structure and binds to the inter-blade groove between β-blades 2 and 3, through a related mechanism to the D box (figure 4d). The side-chains of the three conserved non-polar residues anchor the ABBA motif to the ABBA-motif binding groove, with the Asp at P6 forming a salt-bridge with an Arg of blade 2 [71].

**Non-canonical degrons.** In addition to the D box, KEN box and ABBA motif, non-canonical degrons have also been identified (figure 5c). However, some of these are likely to be variants of the well-characterized D box and KEN box degrons [71,120]. For example, the conserved Arg (P1) at the N-terminus of the D box can be substituted with Lys, His or Gln although this is often accompanied by a Lys at P7 which can interact with the acidic patch at the N-terminus of the D-box binding channel [71]. The O box identified as an APC/C degron in Orc1 closely matches the D-box consensus [137], suggesting it may interact with the D-box receptor of APC/C*Cdh1* [71]. In mammals, the CRY box (CRYxPS) within the NTD of Cdc20 mediates APC/C*Cdh1*-dependent Cdc20 destruction in oocytes and embryos [140]. Insights into how the CRY box might interact with Cdh1 were provided by cryo-EM structures of the APC/C*CMcC* [92,93] (discussed in §8). These showed that the CRY box of the MCC Cdc20 subunit interacts with the WD40 domain of Cdc20 of APC/C*Cdc20* in proximity to the D-box binding site.

In addition to modulation of APC/C–substrate affinities by substrate phosphorylation in or adjacent to the degron, ubiquitination of Lys residues within or in close proximity to degrons may influence APC/C–substrate affinities. One example of this is that the KEN-box Lys residue is one of the most frequently ubiquitinated sites in vivo [141]. Modification of the KEN box would be expected to reduce APC/C–substrate affinities.

Discovery of new APC/C substrates will be facilitated by high-throughput automated approaches based on protein micro-arrays such as the extract-based functional assays [142,143].

### 4. The APC/C pairs with two E2s to assemble polyubiquitin chains

The APC/C is a RING domain E3 ligase. RING domains interact directly with their canonical E2s and bring these into close proximity with substrates bound to degron recognition sites situated elsewhere on the E3 ligase [144]. Metazoan APC/C assembles atypical Lys11-linked chains to promote proteolysis
and mitotic exit [145,146], in a process involving two distinct E2 activities. Chain formation is initiated with the E2 UbcH10 (also termed Ube2C) [147,148], whereas Ube2S is primarily responsible for chain extension [149–152]. Ube2S interacts with the acceptor ubiquitin to generate Lys11-linked chains through a substrate-assisted catalytic mechanism in which Gln34 on the acceptor ubiquitin activates and orients the target Lys11 to attack the donor ubiquitin conjugated to Ubc2S [152]. UbcH10 and Ube2S act in concert to generate branched chains (mixed K11 and K48 linkages). The ubiquitin chain topology determines the efficiency of proteasome-dependent proteolysis of the ubiquitinated substrate [153–156]. UbcH10 alone is competent to generate short ubiquitin chains of mixed K11, K48 and K63 linkage [157,158]. Neither UbcH10 nor Ube2S are essential, suggesting an alternative E2 can function in place of UbcH10 in vivo, likely to be UbcH5 [159]. However, Ube2S is essential for optimal release from a SAC-dependent arrest, possibly due to its role in reactivating the APC/C on cessation of SAC signalling [149–151]. In S. cerevisiae the APC/C generates canonical Lys48-linked chains also using two E2s: the initiating E2 Ubc4 and the elongating E2 Ubc1 [160]. A UBA domain in Ubc1 is required for processivity [160] by enhancing Ubc1 association with the APC/C in competition with Ubc4 [161].

4.1. Monoubiquitination catalysed by UbcH10

Cryo-EM studies of human APC/C<sub>Cdh1</sub> in complex with UbcH10 and Ube2S with and without ubiquitin have provided detailed mechanistic insights into the process of substrate ubiquitination [80,99,111,112]. UbcH10 is a canonical E2 that interacts with the RING domain of Apc11 [80,111]. In human APC/C, the catalytic module is a region of conformational flexibility [60,86]. Binding of UbcH10, but not Ube2S, is dependent on a conformational change mediated by the coactivator subunit (figures 6 and 7) [86,110]. Thus, coactivators are required for both substrate recognition and for stimulating the catalytic activity of the APC/C [117]. This conformational change involves a movement of the catalytic module from a ‘down’ to an ‘up’ position. In the ‘down’ position, Apc11<sup>RING</sup> is in contact with Apc5 of the platform, blocking the UbcH10-binding site. On conversion to the coactivator-bound state, movement of the catalytic module to an upward position exposes the UbcH10-binding site on Apc11<sup>RING</sup>-Apc2<sup>WHB</sup>, resulting in at least a 10-fold increased affinity for UbcH10 [86]. In this state the catalytic module is flexible with weak density recovered and conformational heterogeneity for a variety of ternary complexes [60,86]. Coactivators also increased the catalytic efficiency of S. cerevisiae APC/C (decrease in K<sub>m</sub> and increase in V<sub>max</sub>) [163], although this may result from a mechanism other than a coactivator-induced conformational change (D Barford & E Vázquez Fernández 2017, unpublished data).

The interaction of the zinc binding region (ZBR) domain of the inhibitor Emi1 with Apc11<sup>RING</sup> stabilizes the conformation of the catalytic module because the ZBR domain bridges Apc11<sup>RING</sup> with Apc11<sup>RING</sup>- and Apc2<sup>CTD</sup> (figure 1a). This allowed definition of Apc11<sup>RING</sup> and Apc2<sup>CTD</sup> to a local resolution of approximately 6 Å [80] and it showed for the first time how Apc11<sup>RING</sup> interacts with Apc2<sup>CTD</sup>. The juxtaposition of Apc11<sup>RING</sup> and Apc2<sup>CTD</sup> is similar to the swung out conformation of Rbx1<sup>RING</sup> in activated Cul5-Rbx1 [164]. Engagement of UbcH10 with Apc11<sup>RING</sup> is essentially similar to other RING domain–E2 interactions (figure 6b) [80,111].

Density for UbcH10 was poorly resolved, probably due to the low stoichiometry of UbcH10–APC/C interactions and conformational flexibility of the catalytic module. The Apc11<sup>RING</sup>-UbcH10 interface was confirmed by a detailed mutagenesis study by Schulman and colleagues [111]. On interacting with UbcH10, the catalytic module rotates by 12° relative to its position in the APC/C<sub>Cdh1-Emi1</sub> complex [80]. Importantly no EM density was visible for ubiquitin in the APC/C<sub>Cdh1</sub>-UbcH10~ubiquitin cryo-EM maps [80,111]. This would indicate that the ubiquitin moiety must be mobile, and only transiently adopts the closed E2–ubiquitin conformation that primes the E2~ubiquitin thioester bond to stimulate the intrinsic catalytic activity of E2~ubiquitin [152,165–169]. Formation of the closed E2–ubiquitin conformation, where the ubiquitin moiety interacts with the RING domain through its Ile36 and E2 through its Ile44, as a requirement for optimal substrate ubiquitination, is based on the finding that mutating either Ile36 or Ile44 residues in ubiquitin virtually eliminated ubiquitination of APC/C substrates [80]. The APC/C is reminiscent of other single domain RING and U-box E3s that bias the E2~ubiquitin conformation from multiple extended states to the closed state [168,170]. As discussed elsewhere [80,111], an interesting possibility is that substrate initiation motifs that promote lysine ubiquitination [158] may induce a closed UbcH10~ubiquitin conformation.

The study of Schulman and colleagues revealed that Apc2<sup>WHB</sup> forms an unusual interaction with the backside of UbcH10 [111]. This interaction follows a rigidification of the WHB domain (which is mobile in UbcH10-free structures) induced upon UbcH10 binding (figure 7c). Apc2<sup>WHB</sup> is essential and specific for APC/C-UbcH10-dependent substrate modification, but is dispensable for UbcH5 activity. The activity of Ube2S, which does not interact with Apc2<sup>WHB</sup>, is also independent of Apc2<sup>WHB</sup> [111]. Apc2<sup>WHB</sup> both enhances APC/C-UbcH10 affinity, but importantly also greatly stimulates (by more than 100-fold) the catalytic activity of UbcH10, likely by stabilizing the E2~ubiquitin closed conformation through an allosteric mechanism. Since the WHB-binding interface of UbcH10 differs substantially from its counterpart in UbcH5, similar interactions between UbcH5 and Apc2<sup>WHB</sup> are not possible, thus explaining how Apc2<sup>WHB</sup> contributes to UbcH10 specificity [111].

4.2. Polyubiquitination catalysed by Ube2S

The processive ubiquitination reaction catalysed by Ube2S involves modification of a constantly changing substrate that is the growing distal ubiquitin moiety of the polyubiquitin chain. Biochemical studies showed that UbcH10 and Ube2S do not compete for the same binding site on the APC/C [150,152], suggesting that Ube2S differs from canonical E2s by not interacting with the RING domain of Apc11, a notion also consistent with the observation that Ube2S catalyses formation of unattached K11-linked polyubiquitin chains [171]. APC/C–Ube2S interactions are dependent on the C-terminal LRRL motif of Ube2S [86,154,172,173]. The APC/C dramatically improves the catalytic efficiency of Ube2S-mediated Lys11-linked chain assembly [99,173]. This stimulatory effect of the APC/C requires a surface centred on Ala46 of the acceptor ubiquitin, indicating that APC/C tracks the distal ubiquitin of a growing ubiquitin chain [173]. This finding explains how the APC/C generates ubiquitin chains without altering its interactions with substrates and E2s.
Brown and colleagues [99] in agreement with Kelly et al. [173] showed that the APC/C increased Ube2S catalytic efficiency to massively increase polyubiquitination. Although this catalytic enhancement requires Apc11RING, two lines of evidence suggested that this did not involve the canonical E2-binding surface on Apc11RING. Mutagenesis studies identified a novel surface on Apc11RING (termed the exosite) required for Ube2S activity, a result complemented by NMR data showing chemical shift perturbations in this region of Apc11RING in the presence of ubiquitin. Conversely, acceptor ubiquitin mutants with specific defects in APC/C-Ube2S-dependent ubiquitination [99,152,173] map to a RING-binding surface on ubiquitin identified by NMR [99]. In a subsequent study, the structural basis for Ube2S-catalysed ubiquitin chain extension was defined [112]. A cryo-EM reconstruction of APC/Cdh1 in complex with Ube2S revealed that the Ube2S UBC (ubiquitin conjugating) domain interacts with Apc2, rationalizing the deleterious effects of mutations of the αC and αD helices (figure 6c) [99,112,173]. Its LRRL C-terminus interacts at a site between Apc2 and Apc4, as previously determined for the Emi1 LRRL tail in the APC/C/Cdh1.Emi1 structure [80]. The distal (acceptor) ubiquitin moiety of the ubiquitinated substrate engages the repurposed exosite on Apc11RING, following a conformational change of Apc11RING, presenting its K11 residue to undergo nucleophilic attack onto the donor ubiquitin conjugated to Ube2S. Thus the Apc11RING exosite captures the tip of the growing polyubiquitin chain promoting its reaction with Ube2S ≏ ubiquitin bound to Apc2 (figure 7e).

Figure 6. APC/C ubiquitination reaction. (a) Apo APC/C. In the absence of coactivator the catalytic module adopts a ‘down’ inactive conformation. UbcH10 binding to Apc11RING is blocked by Apc5, and Apc5 prevents the correct location of Apc2 WHB required to engage UbcH10. EM density for Apc11RING is weak indicating RING domain flexibility. PDB 5G05 from Zhang et al. [60]. (b) Complex of APC/C-Cdh1-UbcH10-ubiquitin conjugate. Apc2 WHB becomes ordered and engages UbcH10. dUb: modelled donor ubiquitin conjugated to UbcH10. The C-terminus of dUb is indicated with a red sphere. PDB 5A31, from Chang et al. [80]. PDB for Apc2 WHB 4YII Chang et al. [111]. (c) APC/C-Cdh1-Ube2S-ubiquitin conjugate. Ube2S is partially built. aUb: acceptor ubiquitin bound to the Apc11RING exosite. PDB 5L9T, from Brown et al. [112]. The figure is based on previous work [60,80,111,112].
substrate-recognition module, facing into the central cavity, and this is consistent with the relatively close proximity of the preferred target lysines to APC/C degrons (figures 6b and 7c). In contrast, Ube2S is sited on the periphery of the molecule, able to accept the distal ubiquitin moiety on the polyubiquitin chain. The growing polyubiquitin chain can then be easily accommodated on the outside of the molecule (figures 6c and 7e).

4.3. Multiubiquitination catalysed by UbcH10

The repurposing of Apc11RING that stimulates Ube2S-catalysed ubiquitin chain extension also plays a role in protein multiubiquitination catalysed by UbcH10 through its interaction with the canonical E2-binding site on Apc11RING. A cryo-EM structure of a monoubiquitinated substrate bound to APC/C-Cdh1–UbcH10 ~ ubiquitin showed that the substrate-conjugated ubiquitin moiety interacted with the Apc11RING exosite [112], a finding supported by mutagenesis data revealing that multiubiquitination catalysed by UbcH10 was defective in the Apc11RING exosite mutant. The structure suggests a model for how an interaction between the Apc11 exosite and a substrate conjugated ubiquitin would increase substrate affinity and hence processivity (figure 7d).

Importantly, these data are consistent with the proposal that substrate ubiquitination primes APC/C substrates for further ubiquitination through a mechanism termed processive affinity amplification [174].

The inherent weak affinities between the APC/C–substrate complex and the E2s UbcH10 and Ube2S were overcome by employing artificial reinforcement of these interactions through a three-way chemical linkage involving the substrate, ubiquitin and E2 [111,112]. The interactions between the Apc11RING exosite and ubiquitin were strengthened by generating a ubiquitin variant (Ubv) with substantially increased affinity for Apc11RING [112]. In another approach to stabilize APC/C–substrate interactions with UbcH10, either UbcH10 was directly fused to the C-terminus of Apc11 or the LRRL tail of Ube2S was fused to the C-terminus of UbcH10, enhancing its affinity 10-fold [80].

5. The APC/C controls cell-cycle-dependent substrate degradation

The capacity of the APC/C to control the degradation of regulatory proteins in a cell-cycle-dependent manner defines the ordered progression through distinct phases of the cell cycle.
cycle. The factors that affect differential rates of protein degradation during the cell cycle depend upon both changes in the composition and conformation of the APC/C itself as well as direct changes to individual substrates, and their intrinsic processivity. Switching between Cdc20 and Cdh1 contributes to altering APC/C substrate specificity. Cdh1 directs APC/C-mediated ubiquitination of the Aurora kinases [118], which are not substrates of APC/C<sub>Cdc20</sub>. Nevertheless, apart from this example, there are relatively few instances known where the timing of substrate degradation can be directly explained by the switch of coactivator. Apart from coactivator switching, the two best-characterized regulatory mechanisms for determining the cell cycle order of APC/C-regulated substrate degradation are the spindle assembly checkpoint and substrate phosphorylation.

5.1. Substrate degradation at the spindle assembly checkpoint

A few APC/C substrates are degraded in early mitosis (prometaphase), for example Nek2A, cyclin A and Hox10, during an active SAC [175–181]. Thus, ubiquitination of these substrates is not inhibited by the SAC. These substrates differ from the canonical D-box and KEN-box-dependent substrates cyclin B and securin whose ubiquitination is inhibited by the MCC [24,177,178]. This implies that these early substrates would incorporate additional novel APC/C-recognition motifs that do not rely on binding to D-box and KEN-box receptors. Indeed, in the case of Nek2A, its interaction with the APC/C occurs in the absence of coactivators [182,183], through a C-terminal Met-Arg (MR) tail motif that mimics the IR tail of coactivator and Apc10 [182,183]. However, coactivators are required to mediate Nek2A ubiquitination [117,181] by inducing a UbcH10-binding site on the APC/C [86]. For Nek2A to be degraded during an active checkpoint it requires both its C-terminal MR tail and the adjacent leucine zipper, implying a requirement for Nek2A dimerization. Deletion of either motif shifts the degradation to anaphase that is KEN-box dependent [181,184]. Nek2A binds to apo APC/C, but not APC/C<sub>C</sub>MCC [181], and its binding required the C-box site of Apc8, likely through its MR tail (since the IR tail of Cdc20 of the MCC binds to the C-box binding site of Apc8A [92,93]).

Cyclin A is degraded soon after nuclear envelope breakdown (NEBD) in prometaphase some 20 min before cyclin B. Importantly cyclin A degradation is not inhibited by an active SAC, although its degradation is affected by the SAC [121,176–178]. When the SAC is repressed by the over-expression of a dominant negative BubR1 mutant, cyclin B1 is degraded shortly after NEBD, similar to cyclin A [177]. In further support that the SAC is a major cause of the difference in timing of cyclin A and cyclin B degradation, inactivating the SAC using the Mps1 kinase inhibitor reresverine caused premature cyclin B degradation, with kinetics similar to cyclin A, and importantly no longer dependent on Apc13 [59], which is required to reactiviate APC/C<sub>C<sub>Cdc20</sub></sub> when the SAC is switched off.

Both the N-terminal 165 residues of cyclin A and the Cks subunit are necessary and sufficient to confer the SAC-resistant degradation of cyclin A [178,185,186]. Deletion of the cyclin A D box does not stabilize the protein at prometaphase, or affect degradation timing later in mitosis, questioning the importance of this motif in APC/C-dependent recognition [177,178,182,187]. A region of cyclin A (residues 98–165) C-terminal to the D box contributes to the degradation timing and this region (which incorporates the ABBA motif [121]) binds directly to Cdc20, competing with BubR1 [186]. An ABBA motif also contributes to the early timing of Clb5 degradation in S. cerevisiae compared with securin and Dbf4 [122]. However, unlike vertebrate cyclin A2, Clb5 degradation is sensitive to the SAC although there exists a lower rate of Clb5 degradation during a SAC that depends on the ABBA motif [122].

The ABBA motif clearly plays a role in determining the early destruction of cyclin A2 and Clb5 relative to cyclin B and securin. However, this may not be entirely due to the ability of the ABBA motif to overcome the SAC-induced inhibition of D-box and KEN-box-dependent substrates. One possibility is that cyclin A2 is a more processive substrate. This could be explained if cyclin A2 has a relatively higher affinity for the APC/C, thus competing effectively for binding sites on the APC/C. The ABBA motif may contribute to the higher affinity. However, against the competition argument is the finding that in S. cerevisiae over-expression of Clb5 did not alter the relative timing of destruction of the later substrate securin [188]. It is also interesting that in inactivated Xenopus egg extracts (where there is a weak checkpoint), mutation of the ABBA motif (F<sub>x</sub>[L/V]Y/F/YkXV: residues mutated in bold) to Ala had no to little effect on cyclin A degradation [189]. The Cks subunit of the Cdk1–cyclin B1–Cks complex recruits the complex to the checkpoint-inhibited phosphorylated APC/C at prometaphase, but ubiquitination of cyclin B1 is blocked by the MCC. This prior binding renders cyclin B1 a better APC/C substrate in metaphase [25].

5.2. Phosphorylation can regulate the timing of substrate ubiquitination

Phosphorylation of D box and KEN box degrons has important consequences for controlling the timing of APC/C-mediated protein degradation. Cdk1-dependent phosphorylation of the P2 site of Dbf4 suppresses its destruction [122], contributing to the timing of its destruction in mitosis. A bulky negatively-charged residue at P2 interferes with D-box binding to the D-box receptor of the coactivator whereas phosphorylation at the P6 position promotes human securin degradation [134]. The structural explanation for this was discussed in §3. In contrast, Cdk1-mediated phosphorylation of S. cerevisiae securin in close proximity to the KEN box (17 residues C-terminal) and D box (14 residues N-terminal) reduces the rate of APC/C-dependent securin ubiquitination some 5–10 fold [190]. Dephosphorylation of these sites by Cdc14 therefore promotes securin degradation. Interestingly, since active separase (produced as a result of securin degradation) stimulates Cdc14, a positive feedback loop is generated involving Cdc14-mediated dephosphorylation of securin. Together with the partial inactivation of CdkS at metaphase due to APC/C<sub>C<sub>Cdc20</sub></sub>-mediated destruction of mitotic cyclins, it increases the rate of securin degradation and the abruptness of anaphase onset [122,190]. In S. cerevisiae, one factor delaying securin degradation relative to Clb5, even in the absence of the SAC, is Cdk1-dependent phosphorylation of residues proximal to its KEN box.

At S-phase, Cdk-dependent phosphorylation of amino acids in the immediate vicinity of the D box of Cdc6 blocks binding to the APC/C, thereby protecting Cdc6 from ubiquitination, and promoting DNA replication origin licensing [191]. In another example, Aurora A-kinase phosphorylation of the
D-box P3 residue stabilizes geminin [192], likely because the P3 position has a preference for non-bulky residues.

5.3. Substrate ubiquitination topology may affect timing of proteolysis

The pattern of substrate ubiquitination (multi, poly and branched chains) that favours proteasome-dependent proteolysis (and possibly inhibition of DUB activity) would also contribute to more effective substrate destruction [153,154]. Processively polyubiquitinated substrates are degraded earlier in the cell cycle [122,155,193,194]. It is possible that the position of degrons relative to target lysines affects the efficiency and type of protein ubiquitination.

Finally, in mitosis, the mitotic spindle regulates the timing of spindle assembly factor (SAF) degradation through the microtubule-mediated protection of SAF ubiquitination [195].

6. Phosphorylation regulates APC/C activity at multiple levels

6.1. APC/C phosphorylation promotes Cdc20 association and activation

APC/C activity is entirely dependent on its association with either of the two coactivators Cdc20 and Cdh1, with the APC/C being activated early in mitosis (after NEBD—prometaphase), remaining active until late G1. Although high mitotic Cdk activity is required to stimulate the APC/C in mitosis, the APC/C remains active after mitotic cyclin degradation. This is due to the reciprocal effects of Cdk phosphorylation on the activities of Cdc20 and Cdh1 through affecting their affinity for the APC/C. The association of Cdc20 and Cdh1 with the APC/C is controlled at the level of both the core APC/C and coactivator phosphorylation. Cdk-dependent phosphorylation of core APC/C subunits activates APC/C[Cdc20] [196–201] by promoting Cdc20 association [60,199,201–203], whereas Cdh1 binding does not require APC/C phosphorylation [60,198]. Simultaneously, Cdk phosphorylation of Cdh1 completely blocks its capacity to bind and activate both mitotic and interphase APC/C [32,80,198,204]. As Cdk activity declines at anaphase due to APC/C[Cdc20]-mediated ubiquitination of cyclin A and cyclin B, both the APC/C and Cdh1 become dephosphorylated. This inactivates Cdc20, but allows binding of Cdh1 to generate APC/C[Cdhl]. Cdh1 is inactivated in late G1 due to S-phase cyclin-dependent phosphorylation and Emi1.

Multiple APC/C subunits are phosphorylated in early mitosis associated with activation of APC/C[Cdc20]. Apcl and Apcc are hyper-phosphorylated, with Apcc phosphorylation readily detected by its retarded mobility on SDS-PAGE. Phosphorylation mapping by mass spectrometry of endogenous APC/C defined multiple phosphosites on Apcl and Apcc [201,202,205–207], findings confirmed by in vitro APC/C phosphorylation analysis using purified Cdk and Plk1 [60]. Two hyper-phosphorylated regions of Apcl and Apcc are the 300s loop of the Apcl WD40 domain (Apcl300s loop), and a 300-residue segment in Apcc.

In 2016 three studies provided insights into mechanisms of activation of vertebrate (human and Xenopus) APC/C[Cdc20] by mitotic phosphorylation. These studies revealed that phosphorylation-dependent APC/C[Cdc20] activation primarily involves phosphorylation of the Apcl300s loop that relieves an auto-inhibitory segment within the Apcl300s loop, thereby enabling Cdc20 association [60,202,203]. Introducing phophomimetics into this loop stimulated the ability of Cdc20 to activate the APC/C [60,202,203] and promoted Cdc20 binding [202] in the absence of APC/C phosphorylation. In contrast, mutating phosphosites to Ala ablated Cdc20-dependent APC/C activation [202,203] and Cdc20 binding [203].

To understand the molecular basis for how phosphorylation activates APC/C[Cdc20], a cryo-EM structure of phosphorylated APC/C[Cdc20] was determined [60]. The structure of phosphorylated APC/C[Cdc20] is very similar to that of unphosphorylated APC/C[Cdhl] (figures 1 and 8). Cdc20...
interacts with the APC/C through three motifs: the C box to Apc8B (augmented by the KILR motif [208]), the IR tail to Apc3A and a region contacting Apc1PC. Relative to Cdh1 the contacts are fewer. Strikingly, EM density corresponding to phosphorylated regions could not be observed, indicating that phosphorylated regions of the APC/C do not directly or indirectly contribute to increasing the affinity of the APC/C for Cdc20. This implied that APC/C phosphorylation would remove an inhibitory segment from a Cdc20 binding site. To explore this possibility, the structures of phosphorylated and unphosphorylated apo APC/C were compared. The two structures were very similar, except that in the unphosphorylated apo structure, a segment of EM density occupies the C-box binding site (figure 9). The proximity of this unassigned EM density to the disordered 300s loop of the Apc1 WD40 domain (Apc1 WD40) suggested that this segment corresponded to a region of the Apc1300s loop. In a structure determined with this loop deleted, the C-box binding site was devoid of EM density [60]. Deletion of the Apc1300s loop constitutively activated APC/C [60].

The AI segment includes an Arg-Phe dipeptide, analogous to the Arg-Tyr motif of the C box. Modelling of the AI segment into EM density showed that the Arg side chain of the AI segment mimics the Arg of the Arg-Tyr motif of the C box, anchoring the AI segment to the C-box binding site (figure 9c). Mitotic phosphorylation of sites flanking the Arg-Phe motif would destabilize interactions between the AI segment and the C-box binding site through steric hindrance and charge repulsion, leading to the displacement and disordering of the AI segment and relief of auto-inhibition. These findings that an auto-inhibitory segment within the Apc1300s loop blocks Cdc20 activation and that its mitotic phosphorylation relieves this auto-inhibition are in agreement with biochemical data [202,203] (figure 10). Fujimitsu and colleagues [203] showed that Apc1300s bound to the APC/C in an anaphase extract, whereas the phosphomimetic mutants abolished this interaction, highlighting how the interaction of Apc1300s with the APC/C is dependent on its phosphorylation status.

The data of Zhang et al. [60] indicated that the critical determinant of activation of APC/C-Cdc20 by mitotic phosphorylation was displacement of the AI segment to relieve auto-inhibition. However, Apc3 is also highly phosphorylated in mitosis [201,202,205–207] and Cks stimulates both Cdk-dependent activation of APC/C-Cdc20 [197,209] and Apc1 and Apc3 phosphorylation [60,209], and interacts with Apc3 [25,203, 209,210]. Deletion of the hyperphosphorylated Apc3

Figure 9. Control of APC/C-Cdc20 by phosphorylation. (a) In the unphosphorylated state an auto-inhibitory segment (AI; dark green) within the Apc1300s loop of Apc1 WD40 mimics the Cdc20 C-box motif and binds to the C-box binding site, blocking Cdc20 association. The AI segment is located on the same face of the APC/C as the hyperphosphorylated Apc3 loop. (b) Zoomed view of the AI segment of Apc1 WD40 associated with the C-box binding site of Apc8B. (c) Superposition of the AI segment with the Cdc20 C box (purple) shows that a conserved Arg residue anchors both the C box and the AI segment to the C-box binding site. Sites of mitotic phosphorylation present within the AI segment that activate APC/C-Cdc20 are depicted as red spheres. From Zhang et al. [60].
Figure 10. Schematic of control by phosphorylation. In the unphosphorylated state an auto-inhibitory (AI) segment of Apc1WD40 mimics the Cdc20 C-box motif and binds to the C-box binding site, blocking Cdc20 association. Initial Cdk-dependent phosphorylation of a kinase loop recruits Cdk-cyclin-Cks to the APC/C to facilitate intramolecular phosphorylation of the AI segment (when transiently displaced from the C-box binding site). The phosphorylated AI segment is stably displaced from the C-box binding site, permitting Cdc20 association to generate APC/C-Cdc20. This scheme indicates the relay mechanism by which initial phosphorylation of exposed consensus Cdk1 sites on Apc3 allows recruitment of Cks-Cdk-cyclin to the APC/C to promote intramolecular phosphorylation of Apc1. From Zhang et al. [60].

loop reduced both Apc1 AI segment phosphorylation [60] and APC/C-Cdc20 activation [60,203], as well as disrupting interactions between the APC/C and Cdk–cyclin A–Cks [60,203]. The phosphorylated Apc3 loop (residues 202–342) directly binds Cks [203]. Thus a likely explanation for these findings, and for the lag phase that accompanies APC/C activation by Cdk-1–cyclin B–Cks [197], is that Apc3 phosphorylation recruits Cdk–cyclin–Cks via a relay mechanism. Cdk–cyclin–Cks association with the Apc3 loop would allow for a kinetically more efficient intra-molecular phosphorylation of the Apc1 auto-inhibitory segment that only becomes accessible to Cdk when transiently displaced from the C-box binding site. Phosphorylation of the Apc1300s loop stably displaces the AI segment from the C-box binding site (figure 10). Intra-molecular phosphorylation of the Apc1300s loop is associated with relaxed Cdk specificity and the phosphorylation of non-consensus Cdk sites [60]. Interestingly these sites are not evolutionarily conserved, suggesting that the exact location of the phosphorylation sites with Apc1300s is not critical to their capacity to displace the AI segment.

Cdh1 and Cdc20 bind to common sites on the APC/C, yet only APC/C-Cdc20 is activated by phosphorylation [80,60]. The phosphorylation-independent activity of Cdh1 is due to the increased affinity of Cdh1 for unphosphorylated apo APC/C, which overcomes the inhibition from the unphosphorylated AI segment. The increased affinity results from the more extensive contacts formed between the APC/C and Cdh1 relative to Cdc20. This also explains why the APC/C inhibitor TAME [211], which interacts with both the IR tail and C-box binding sites through structural mimicry of the IR tail and C-box, is a more potent inhibitor of APC/C-Cdc20 than APC/C-Cdh1 [60].

6.2. Cdk phosphorylation of Cdh1 and Cdc20 inhibits APC/C association

Binding of Cdh1 to the APC/C is negatively regulated by phosphorylation. Based on the structure of APC/C-Cdh1-Emi1, the four phosphorylation sites (Ser40, Thr121, Ser151 and Ser163 of human Cdh1) that suppress Cdh1 activity [80,212] can be rationalized (figure 3c). Phosphorylation of individual sites only partially suppresses APC/C activity, whereas phosphorylation of all four sites would destabilize Cdh1NTD–APC/C interactions through electrostatic repulsion and steric clashes. Ser40 is immediately N-terminal to the C box, whereas the side-chains of Ser151 and Ser163 flank the KLLR motif [80].

Cdk phosphorylation of the Cdc20 NTD also negatively regulates Cdc20 activation and its binding to the APC/C [115,213,214]. Cdk2-cyclin A2 phosphorylation of Cdc20 at interphase is proposed to prevent premature activation of APC/C [80], thereby stabilising cyclin B1 and promoting mitotic entry [214]. In mitosis Cdk1-cyclin B1 may contribute to Cdc20 phosphorylation [214]. The Cdk phosphorylation sites are close to the N-terminus of the C box (Thr55, Thr59 and Thr70 in human Cdc20, with C box comprising residues Asp77 to Arg83) [60], thus phosphorylation may block C-box binding to the Apc8B C-box binding site, reminiscent of Cdh1 inhibition by Cdk phosphorylation [80]. However, it should be noted that in the APC/C-Cdc20 structure residues N-terminal to Ser72 are largely disordered [60], making it unclear mechanistically how phosphorylation of Cdc20 N-terminal to the C box inhibits its activity. Significantly, mutation of Thr55, Thr59 and Thr70 to Ala produced no cellular phenotype [214], suggesting that multiple Cdk phosphorylation sites on Cdc20 contribute to its inactivation. These may involve mechanisms in addition to directly inhibiting its association with the APC/C. For instance, it is possible that Cdc20 NTD phosphorylation affects the structure of free Cdc20, possibly promoting a closed conformation that cannot bind the APC/C [215]. PP2A has been suggested as the Cdc20 phosphatase [115,214] and possibly binds directly to the APC/C mediated by PP2Aβ56 [216]. A recent study in Caenorhabditis elegans showed that kinetochore-associated PP1 also contributes to dephosphorylation of Cdc20 Cdk phosphorylations (with Thr32, equivalent to human Thr70), being a key site responsible for the control of C. elegans Cdc20 by phosphorylation) through a mechanism by which Cdc20 is recruited to kinetochore by the ABBB motif of Bub1 [217]. Thus, depending on the status of their microtubule attachment, kinetochores either inactivate (via the SAC) or activate (via Cdc20 dephosphorylation) APC/C-Cdc20. This explains the paradox that the checkpoint proteins Bub1/Bub3 promote anaphase onset independently of the checkpoint [218,219]. While Ala substitution of Cdk sites within the NTD of C. elegans Cdc20 accelerated normal mitosis,
this mutant retained the ability to significantly delay mitosis in the presence of unattached kinetochores [217], indicating that Cdk phosphorylation of Cdc20 does not contribute to the SAC.

Plk1 (mediated through a scaffolding role of Bub1) phosphorylation of human Cdc20 on Ser92 impaired the assembly of polyubiquitin chains in vitro, mainly through inhibition of Ube2S [220] by preventing the association of Ube2S to the APC/C [220,221]. Analysis of the APC/C/Cdc20 structure indicates that Ser92 is in contact with Apc8B [60], remote from the Ube2S binding site [112]. Thus the molecular mechanism by which Ser92 phosphorylation inhibits Ube2S is not currently clear. Ser92 phosphorylation does not affect the MCC-mediated inhibition of APC/C/Cdc20, revealing that Bub1-Plk1 directly inhibits APC/C/Cdc20 through a mechanism that is independent of the MCC [220]. The inhibitory phosphorylation on Cdc20 is removed by PP2A [86], a kinetochore-bound phosphatase [220,221] and PP1 [217].

6.3. Substrate phosphorylation can regulate association with the APC/C

Direct phosphorylation of substrates provides a third level of APC/C control by protein phosphorylation, discussed above.

7. Emi1 inhibits APC/C/Cdh1

In vertebrates, Emi1 functions as an antagonist of APC/C/Cdh1 during G2 [42,43]. Four functional elements of Emi1 mediate APC/C/Cdh1 inhibition [80,107,172,222]. Similar to the MCC, Emi1 blocks D-box recognition by APC/C-coactivator complexes and also antagonizes the two E2s UbcH10 and Ube2S. A D-box motif that occludes substrate recognition is connected through a linker to a zinc-binding region (ZBR) (Emi1ZBR), that interferes with UbcH10-dependent APC/C activity [107,172,222] (figure 1a). A C-terminal LRLL sequence (LR tail: Emi1LR), identical to the LRLL motif required for Ube2S-dependent synthesis of polyubiquitin chains on APC/C substrates [150,151], and its association with the APC/C [86,154,172], antagonizes Ube2S [107,172] by interacting with the Ube2S LRLL-tail binding site on Apc4 [80].

8. Reciprocal regulation of the spindle assembly checkpoint and APC/C/Cdc20

To ensure the fidelity of the inheritance of genetic information, the cell has evolved cell cycle checkpoints that control progression through cell cycle transitions that are dependent on the successful completion of a preceding event. The spindle assembly checkpoint (SAC), also known as the mitotic checkpoint and kinetochore checkpoint, coordinates sister chromatid segregation at the metaphase to anaphase transition with the correct bipolar attachment of sister chromatids to the mitotic spindle [40,41]. The SAC is exerted by the mitotic checkpoint complex (MCC), a multi-protein complex that functions to repress APC/C activity. Generation of the MCC is catalysed by unattached kinetochores whose structural and biochemical properties are becoming well defined [223,224]. MCC assembly occurs on the outer regions of the kinetochore, specifically the KMN (Kn1-Mis12-Ndc80) network which functions as a recruiting site for multiple checkpoint components. Key among these are the Mad and Bub proteins, identified over 25 years ago in genetic screens for SAC components [225,226]. A checkpoint cascade results in the assembly of a molecular scaffold that catalyses conversion of O-Mad2 (open state of Mad2) to C-Mad2 (closed state of Mad2), in a process that requires the kinetochore-associated C-Mad2. In the template-assisted mechanism [227], the kinetochore-associated C-Mad2, bound to the kinetochore through Mad1, interacts with O-Mad2 to promote its conversion to C-Mad2, a reaction catalysed by Mps1 [224,228]. C-Mad2 captures the N-terminus of Cdc20 and the resultant C-Mad2-Cdc20 binary complex interacts rapidly with BubR1-Bub3 to generate the tetrameric MCC (C-Mad2-Cdc20-BubR1-Bub3) [229]. The MCC is a potent APC/C inhibitor, some 3000-fold more potent than Mad2 alone [229]. The target of the MCC is APC/C/Cdc20 [230].

The structural mechanisms underlying how the APC/C and the MCC are reciprocally regulated in the context of the SAC were derived from cryo-EM reconstructions of APC/C/Cdc20 in complex with the MCC (APC/C/MCC) [92,93]. These studies explained how the MCC blocks D-box- and KEN-box-dependent substrates from interacting with APC/C/Cdc20, and also surprisingly revealed how the MCC interferes with the initiating E2, UbcH10.

Overall structure of the APC/C/MCC. Recombinant reconstituted APC/C/MCC comprises two Cdc20 subunits, consistent with the notion that the MCC interacts with APC/C/Cdc20 [92,93,230–232]. Importantly, the overall structure is essentially identical to the endogenous APC/C/MCC isolated from check-point-arrested HeLa cells (at much lower resolution) [105]. This validated the notion that the physiologically relevant form of APC/C/MCC includes two Cdc20 molecules (termed Cdc20APC/C and Cdc20MCC for the Cdc20 subunits of APC/C/Cdc20 and MCC, respectively) [230,231]. In the APC/C/MCC reconstruction, a large density element termed the MCC-Cdc20 module (MCC and Cdc20APC/C) occupies the central APC/C cavity extending from the front side of the platform domain. The core MCC elements comprising Cdc20MCC, the TPR domain of BubR1 and C-Mad2 resemble their counterparts in the fission yeast MCC structure [69]. Although present in the reconstituted complex, no EM density was visible for BubR1’s C-terminal regions (that includes a pseudo-kinase domain) and its associated Bub3 subunit. Mad2 adopts the closed conformation with its safety belt entrapping the N-terminal KILR motif of Cdc20. The MCC docks into the central cavity of the APC/C contacting Cdc20APC/C and ApccWTH (figure 11). ApccWTH rigidifies and reposition (relative to APC/C/Cdh1-UbcH10 [111]) to engage BubR1. Contacts between the two Cdc20 molecules are mainly mediated by BubR1 that intertwines between them. Through extensive contacts between BubR1 and the two Cdc20 molecules, BubR1 obstructs degron dependent binding to both coactivator subunits. This is achieved because BubR1 incorporates two copies of both the D-box (D1, D2) and KEN-box motifs (K1, K2) and three copies of the ABBA motif (A1–A3) (figure 11b). ApccWTH binds to both Cdc20APC/C and then fold back to contact the A2 and D2 motifs of Cdc20APC/C. This interaction drives BubR1 to intertwine around Cdc20APC/C and then fold back to contact the A2 and D2 motifs of Cdc20APC/C...
sites on Cdc20MCC, forming a lariat-like structure (figure 11a,c,d). The contacts between D1, A1 and K2 of BubR1 and Cdc20APC/C explain why these three motifs are critical to APC/C–MCC interactions and function to sustain the checkpoint response [121,208,233–236]. In contrast, the contacts between A2 and D2 with Cdc20MCC are not as critical for MCC stability and APC/C–MCC interactions, hence the more modest effects of disrupting the checkpoint when these motifs are deleted. D1, A1, K2, A2 occur in an evolutionarily conserved cassette, suggesting the mechanism for inhibiting the APC/C is conserved in all major eukaryotic super groups over one billion years of evolution [233,236]. In addition to directly blocking degron recognition sites on Cdc20APC/C, (although not necessarily Cdc20MCC), the APC/C is conserved in all major eukaryotic super groups dependent on the disorder-to-order transition of Apc15NTH. In support of this notion, deletion of Apc15 locks the Cdc20MCC-binding interface on the platform releases the MCC to rotate away from the catalytic site of the APC/C (figure 12a,b).

Having effectively shut down degron recognition by Cdc20APC/C, (although not necessarily Cdc20MCC), the MCC also represses APC/C’s E3 ligase catalytic activity. In the majority of APC/C MCC molecules (in the Alfieri et al. study [92]) APC/C MCC adopts a closed conformation (APC/C MCC-closed) whereby MCC, through the TPR domain of BubR1, contacts Apc2WHB (figures 11 and 12a). This obstructs the UbcH10 binding site on the catalytic module. APC/C MCC-closed is accompanied by an order-to-disorder transition of the Apc15 N-terminal helix (Apc15NTH) due to the binding of Cdc20MCC to the platform region. This induces an upward movement of the Apc4 helix bundle domain (Apc4 HB) and its adjacent Apc5 N-terminal domain (Apc5 NT), disrupting their contacts to Apc15NTH (figure 12a,b). Interestingly, in a small population of APC/C MCC, the molecule adopts an open state (APC/C MCC-open) whereby MCC has rotated away from the catalytic module exposing the UbcH10 binding site on Apc2WHB (figure 12b). This large repositioning of the MCC is dependent on the disorder-to-order transition of Apc15NTH. On transition from APC/C MCC-closed to APC/C MCC-open, Apc15NTH rebinds to Apc5 NT. This induces a downward rotation of Apc5 NT and downward translation of Apc4 HB, displacing the Cdc20MCC binding site on Apc4 HB by 10 Å. Loss of the Cdc20MCC-binding interface on the platform releases the MCC to rotate away from the catalytic site of the APC/C (figure 12a,b).

In the study of Schulman and colleagues the open APC/C MCC conformation predominates [93]. Whatever the cause of difference in the open-closed ratio between the two studies (possibly due to differences in APC/C MCC reconstitution approaches), open APC/C MCC is associated with an ordered conformation of Apc15NTH, suggesting that the order-to-disorder transition of Apc15NTH influences the open-closed transition. In support of this notion, deletion of Apc15 locks all APC/C MCC molecules into the closed state, consequently repressing Cdc20MCC auto-ubiquitination [92,93].

The open APC/C MCC conformation would suggest that UbcH10 has the capacity to interact with the APC/C MCC, a proposal indeed verified by structures of APC/C MCC–UbcH10 complexes (figure 12a) [92,93]. In these complexes, UbcH10 is bound to the Apc2WHB–Apc11RING catalytic module, as in previous APC/C–UbcH10 complexes [80,111]. The MCC adopts the open conformation with Apc15NTH ordered. Strikingly, the C-terminus of Cdc20MCC engages the catalytic site of UbcH10 with two Lys residues of Cdc20MCC (K485

**Figure 11.** APC/C regulation by the MCC. (a) Atomic structure of APC/C MCC. Ordered regions of BubR1 C-terminal to the TPR domain are shown in space-filling representation, as is the NTD and IR tail of Cdc20MCC. (b) Schematic of BubR1 showing positions of D-box (D1, D2), KEN-box (K1, K2) and ABBA motifs (A1–A3) and schematic of Cdc20. (c) Details of the MCC-Cdc20APC/C module with BubR1 forming extensive interactions with Cdc20MCC and Cdc20APC/C. (d) Schematic of interactions formed by Cdc20MCC and Cdc20APC/C with BubR1 and APC/C subunits. PDB SLCOV, from Alfieri et al. [92].
Figure 12. **APC/C-MCC** adopts open and closed states that allows for reciprocal control by the MCC. (a) In **APC/C-MCC-closed**, the MCC inhibits both substrate (for example, securin and cyclin B) and UbcH10 recognition. Upper panel: overall **APC/C-MCC-closed** structure. Lower panel: shows how binding of the MCC in **APC/C-MCC-closed** causes an upward movement of Apc5NTD and Apc4HBD (compare with b) and concomitant disordering of the N-terminal helix of Apc15 (Apc15NTH). (b) In **APC/C-MCC-open**, the catalytic module is exposed, Apc5NTD rotates, Apc4HBD translates down by 10 Å, and Apc15NTH and Apc11RING domain are indicated with arrows. (c) In the **APC/C-MCC-UbcH10** complex, **APC/C-MCC** adopts the open conformation with Apc15NTH ordered and UbcH10 docking to its canonical position on Apc11RING and Apc2WHB. The C-terminal tail of Cdc20MCC engages the catalytic site of UbcH10 for auto-ubiquitination of Lys485 and Lys490. In **APC/C-MCC-closed**, the C-terminal IR tail of Cdc20MCC engages the C-box binding site of Apc8A. From Alfieri et al. [92].

and K490) accessible to the catalytic site, and which in the reconstituted APC/C-MCC are auto-ubiquitinated in an Apc15- and UbcH10-dependent process [92]. Destabilizing closed APC/C-MCC, either through disrupting the Apc2WHB interface on BubR1 or by deleting the IR tail of Cdc20MCC (which binds Apc8A in closed APC/C-MCC), promotes Cdc20MCC auto-ubiquitination, even in the absence of Apc15 [92,93].

These structural and biochemical data show that Cdc20MCC is auto-ubiquitinated by UbcH10 in the context of the open APC/C-MCC conformation. This requires Apc15 and thus explains how Apc15 deletion suppresses Cdc20 auto-ubiquitination in a SAC-dependent manner. Because Apc15 deletion blocks progression into anaphase after release from the SAC [59,90,91], it suggests that one mechanism by which the amount of MCC dissociated from APC/CMCC after a mitotic arrest is the requirement for Apc15 [241]. The situation may differ in vivo because the Cdc20MCC-K485R/K490R mutant did not prevent MCC release from APC/C-MCC after a mitotic arrest [59]. However, it is possible that in the K485R/K490R mutant alternative lysines in Cdc20MCC could be ubiquitinated in the context of the APC/C-MCC, as Schulman and colleagues observed [93].

To ensure efficient MCC release at anaphase onset, additional MCC subunits might also be ubiquitinated. A good candidate is BubR1 that is ubiquitinated in the context of purified APC/C-MCC and UbcH10 [241]. Supporting the idea that MCC ubiquitination is required for its release from APC/C-MCC, depletion of either Apc11 or UbcH10 decreased the amount of MCC dissociated from APC/C-MCC after a mitotic arrest [59]. In addition, APC/C regulation of BubR1 homeostasis is essential for correct mitotic timing [242,243]. In APC/C-MCC-open the lysine-containing region of BubR1 proximal to D1 is close to the catalytic module of the APC/C, consistent with a model whereby BubR1 is auto-ubiquitinated in the context of APC/C-MCC [92] (figures 12c and 13).

The architecture and composition of APC/C-MCC complexes are conserved in *S. pombe*. However in fission yeast, although Apc15 is also required for Cdc20MCC auto-ubiquitination [232], it additionally functions to exert a checkpoint arrest by stabilizing APC/C–MCC interactions [232,235].
Cdc20MCC auto-ubiquitination is an important event because it contributes to the reciprocal regulation of the APC/C and MCC (figure 13). MCC binding to the APC/C represses APC/C activity. However, in a competing process, dependent on the open conformation of APC/CMCC, APC/CMCC is disassembled due to Cdc20MCC auto-ubiquitination [94] and APC/C-Cdc20 is reactivated. As long as the SAC is active and new MCC is generated, APC/CMCC will reform to suppress APC/C activity. When the SAC is turned off, MCC assembly at the kinetochore stops, and Cdc20MCC auto-ubiquitination [94,244] (although not in vivo [241]), and/or Cdc20 phosphorylation are candidates. Furthermore, USP44 catalyses Cdc20MCC deubiquitination, thereby antagonizing Cdc20MCC ubiquitination to stabilize the APC/CMCC, thus sustaining the SAC [95]. Moreover, a ubiquitination-independent pathway functions to disassemble the free MCC through the activities of the AAA+ ATPase TRIP13 [245–247], in conjunction with the SAC antagonist p31Comet [248], a C-Mad2 binding protein [249]. p31Comet extracts Mad2 from the MCC [250] and targets C-Mad2-Cdc20 to TRIP13 [251], and competes with BubR1 for C-Mad2 [252].

Errors in controlling accurate chromosome segregation due to defects in the SAC underlie aneuploidy and chromosome instability (CIN), and cause tumour heterogeneity and drug resistance [253]. However, extreme CIN correlates with improved cancer outcome, possibly because karyotypic diversity is required to adapt to selection pressures. A study from Swanton and colleagues found that partial APC/C dysfunction caused adaptation to Mps1 inhibitors, suggesting a likely resistance mechanism to therapies targeting the SAC [254].

9. Spatial regulation of the APC/C

Specifics of the intracellular location and spatial control of the APC/C are relatively little understood [255]. The APC/C is localized to centrosomes through Emi1–NUMA complexes [256], the recently identified KIAA1430 protein [257] and to the chromosomes through interactions mediated by the Ska3 complex [258,259].

10. Conclusion and perspectives

An enduring question asked about the APC/C is why is it so large? The simplest response must be that its size reflects its
central role in coordinating critical transitions during the cell cycle. This requires control at multiple levels (through interconvertable coactivator subunits, phosphorylation, the SAC and Emi1, often exerted through allosteric conformational changes of the APC/C), and the capacity of the APC/C to change its substrate specificity during different phases of the cell cycle, transition through which is controlled by the APC/C. Thus, the APC/C possesses intrinsic self-control mechanisms. This is probably best exemplified by the auto-ubiquitination of Cdc20MCC that enables spontaneous activation of the APC/C at anaphase only when all chromosomes have achieved correct bipolar attachment to the mitotic spindle. The large size of the APC/C is contributed by the seven scaffolding proteins, four of which form structurally related homo-dimers, which stack in parallel to create the pseudo-dyad symmetric TPR lobe. This TPR lobe, together with Apc1, Apc4 and Apc5 of the platform, assembles the scaffold to juxtaposition the catalytic and substrate recognition modules. The structural symmetry of the TPR lobe generates multiple structurally related binding sites that engage the C box and IR tail of coactivators and IR tail of Apc10. The structural equivalence of the C-box binding site of Apc8B and the IR-tail binding sites of Apc3 is illustrated by the engagement of the Cdc20MCC IR tail with the site of Apc8B and the IR-tail binding sites of Apc3 is illustrated by the engagement of the Cdc20MCC IR tail with the C-box binding site of Apc8A [92,93]. The APC/C is a distant ancestor of the large CRL family of E3 ligases. Both share a conserved catalytic module of RING and cullin subunits. How the APC/C evolved from the simpler CRLs is not clear because there are no obvious intermediate complexes. All the large scaffolding subunits comprise multiple repeat motif domains. TPR and WD40 repeat motifs are ubiquitous in proteins involved in protein–protein interactions, and it is intriguing that the only other known instances of the PC repeat domain are the Rpn1 and Rpn2 subunits of the 19S regulatory particle of the proteasome [85].

Of interest is the realization that the APC/C functions by engaging coactivators, substrates and inhibitors through recognition of short linear sequence motifs, for example degrons, C box, IR tail and MR tail [120]. Conformational variability of small domains attached to the APC/C scaffold through flexible linkers (Apc2WIIH, Apc1RING) and the WD40 domains of coactivators) has important implications for mediating catalysis and regulation.

Despite the huge progress in understanding the function and mechanism of the APC/C during the two decades since its discovery, much needs to be explored. We still have little molecular understanding of how the APC/C selects different substrates during the cell cycle. This will require cryo-EM structures of different states of the APC/C in complex with full-length intact substrates, in addition to more quantitative determinations of the affinities of different APC/C complexes for their cognate substrates. To what extent APC/C–substrate affinities compared with the catalytic efficiency of lysine ubiquitination (determined by substrate–lysine proximity to the E2 catalytic site and the competing rates of deubiquitination) controls the rate of substrate degradation is unclear. The level to which the intracellular location of APC/C complexes controls their various functions and how this is subject to cell-cycle regulation are also still largely unexplored.

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