Synergistic recruitment of UbcH7~Ub and phosphorylated Ubl domain triggers parkin activation

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Abstract

The E3 ligase parkin ubiquitinates outer mitochondrial membrane proteins during oxidative stress and is linked to early-onset Parkinson's disease. Parkin is autoinhibited but is activated by the kinase PINK1 that phosphorylates ubiquitin leading to parkin recruitment, and stimulates phosphorylation of parkin's N-terminal ubiquitin-like (pUbl) domain. How these events alter the structure of parkin to allow recruitment of an E2-Ub conjugate and enhanced ubiquitination is an unresolved question. We present a model of an E2-Ub conjugate bound to the phosphorylubiquitin-loaded C-terminus of parkin, derived from NMR chemical shift perturbation experiments. We show the UbcH7~Ub conjugate binds in the open state whereby conjugated ubiquitin binds to the RING1/IBR interface. Further, NMR and mass spectrometry experiments indicate the RING0/RING2 interface is re-modelled, remote from the E2 binding site, and this alters the reactivity of the RING2(Rcat) catalytic cysteine, needed for ubiquitin transfer. Our experiments provide evidence that parkin phosphorylation and E2-Ub recruitment act synergistically to enhance a weak interaction of the pUbl domain with the RING0 domain and rearrange the location of the RING2(Rcat) domain to drive parkin activity.

Keywords dynamics; E2 conjugating enzyme; E3 ubiquitin ligase; Parkinson's disease; ubiquitination

Subject Categories Post-translational Modifications, Proteolysis & Proteomics; Structural Biology

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Introduction

Parkinson's disease is the second most common neurodegenerative disease estimated to affect 1% of the population over 60 years of age (Tysnes & Storstein, 2017). The disease is believed to be a result of genetic predisposition or environmental factors (Corrigan et al., 1998) that lead to oxidative damage of mitochondrial proteins (Alam et al., 1997) and subsequent mitochondrial dysfunction (Schapira et al., 1990) and is characterized in patients by the loss of dopaminergic neurons in the substantia nigra of the midbrain (Hornykiewicz, 1966; Riederer & Wuketich, 1976). In addition to sporadic Parkinson's disease, there are also genetic forms of the disease that account for approximately 10% of all cases. In particular, mutations in the genes for PARK2 and PARK6 give rise to early-onset or autosomal recessive juvenile parkinsonism (ARJP) forms of the disease that have similar symptoms including rigidity, Bradykinesia and postural instability (Jankovic, 2008) but affect individuals at a much younger age. PARK2 encodes the E3 ubiquitin ligase parkin (Kitada et al., 1998) where mutations account for 50% of all ARJP cases. Along with the PTEN-induced kinase (PINK1) translated from PARK6, these proteins use the ubiquitin degradation pathway to turnover damaged mitochondria and maintain mitochondrial homeostasis, especially under conditions of oxidative stress.

Parkin is a member of the RBR E3 ligase family that also includes the human homolog of Ariadne (HHARI) and HOIL-1 interacting protein (HOIP; Spratt et al., 2014). These enzymes have a characteristic RBR motif comprising RING1, in-between-RING and RING2 (Rcat) domains that distinguish them from HECT and RING classes of E3 enzymes in terms of structure, mechanism and functionality. In particular, RBR E3 ligases incorporate a hybrid ubiquitination mechanism (Wenzel et al., 2011) whereby an E2 conjugating enzyme is recruited to the RING1 domain (similar to RING E3 ligases) and ubiquitin (Ub) is transferred from the E2~Ub conjugate to a catalytic cysteine in the RING2(Rcat) domain (similar to HECT E3 mechanisms) prior to labelling of a substrate lysine. RBR E3 ligases and RING E3 ligases have RING domains that are structurally similar and are expected to recruit E2 enzymes in a similar fashion (Budhidarmo et al., 2012), as recently shown in crystal structures of the RBR E3 ligases HHARI with UbcH7~Ub (Dove et al., 2017; Yuan et al., 2017) and HOIP with UbcH5b-Ub (Lechtenberg et al., 2016). However, a distinguishing feature of the HHARI and HOIP RBR E3 ligases is their ability to recognize an extended (“open”) form of the E2~Ub conjugate similar to that used by HECT E3 enzymes. This E2~Ub arrangement promotes a conformation susceptible to the

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tron, the other hand has been shown to function with a variety of E2 enzymes including UbcH7 and UbcH5b (Chaugule et al., 2011; Wenzel et al., 2011) although it appears that UbcH7 is the optimal E2 enzyme owing to its preference for ubiquitin transfer to cysteine, a requirement for RBR E3 ligases. While the formation of the UbcH7–Ub conjugate during recruitment by parkin is unknown, it has been established that a cryptic Ub binding site within the RING1–IBR interface is only uncovered upon pUb binding to parkin and this has been proposed to help coordinate E2–Ub recruitment (Kumar et al., 2017).

All RBR E3 ligases identified to date, including parkin, appear to be uniquely regulated (Spratt et al., 2014; Walden & Rittinger, 2018). Parkin is normally autoinhibited by an accessory ubiquitin-like (Ubl) domain (Chaugule et al., 2011) that blocks both the E2 and cryptic ubiquitin sites. In addition, structures show that another accessory module, the RING0 domain, partially obscures the catalytic cysteine in the RING2(Rcat) domain protecting this site from Ub transfer. At least two steps have been identified for the activation of parkin both as a result of phosphorylation by PINK1. Under oxidative stress conditions, PINK1 is activated and phosphorylates ubiquitin (pUb) near the outer mitochondrial membrane. This in turn helps recruit parkin to the membrane through binding of pUb to the RING1–IBR region of the E3 ligase (Sauvé et al., 2015; Wauer et al., 2015; Kumar et al., 2017) and subsequent phosphorylation of parkin’s Ubl (pUb) domain (Ordureau et al., 2014). These two events greatly stimulate ubiquitination activity (Kondapalli et al., 2012; Shiba-Fukushima et al., 2012; Kane et al., 2014; Kazlauskaité et al., 2014; Koyano et al., 2014) through an allosteric displacement mechanism of the pUb domain from parkin (Kumar et al., 2015; Sauvé et al., 2015). What is less clear is how parkin positions the E2–Ub conjugate to enable transfer of the Ub molecule to the RING2 (Rcat) domain as a necessary step for catalysis. Current crystal structures of parkin show the proposed E2 binding site on the RING1 domain is > 50 Å from the catalytic site (C431) in the RING2 (Rcat) domain suggesting a significant conformational rearrangement is needed (Riley et al., 2013; Trempe et al., 2013; Wauer & Komander, 2013; Kumar et al., 2015, 2017; Sauvé et al., 2015; Wauer et al., 2015). A similar dilemma arises from recent structures of HHARI in complex with UbcH7-ub that show the ubiquitin molecule is 47–53 Å from the catalytic site (Dove et al., 2017; Yuan et al., 2017). Alternatively, structures of HOIP:E2-Ub and parkin:pUb complexes assembled from domain-swapping or symmetry-related molecules raise the possibility of co-operation between multiple E3 ligase molecules to promote ubiquitin transfer (Lechtenberg et al., 2016; Kumar et al., 2017).

In this work, we identify how the pUb domain and E2–Ub conjugate co-operate to regulate parkin activity. We use NMR spectroscopy and chemical shift perturbation experiments to determine a model of pUb-bound parkin in complex with its biological UbcH7–Ub conjugate. The structure shows that a non-hydrolysable UbcH7-Ub conjugate binds in an altered “open” conformation with its thioester linkage poised for Ub transfer to the catalytic cysteine of the RING2(Rcat) domain. We show that E2-Ub recruitment to parkin results in two distinct types of NMR chemical shift changes: one set that is consistent with the E2–Ub binding site and a second set that corresponds to residues near the RING0/RING2 interface, indicating this region is re-modelled during E2–Ub recruitment. We show that the reactivity of the catalytic cysteine (C431) in the RING2(Rcat) domain is sensitive to both parkin phosphorylation and E2-Ub binding. Further, we use NMR spectroscopy and hydrogen–deuterium exchange (HDX) mass spectrometry experiments to show that the pUb domain undergoes transient interaction with the RING0 domain that is enhanced upon E2–Ub recruitment by parkin. These events also result in large changes in the exposure of the RING2(Rcat) domain consistent with its rearrangement. Overall, our work provides a dynamic picture of parkin activation whereby PINK1 phosphorylation of parkin and E2–Ub recruitment co-operate to drive parkin activity.

Results

Dual phosphorylation leads to dynamic repositioning of the pUbl domain

Multiple high-resolution structures of parkin have provided significant insights into its E3 ligase function. Structures of near full-length parkin (Kumar et al., 2015; Sauvé et al., 2015) show the C-terminal RING0–RING1–IBR–RING2(Rcat) domains (termed “R0RBR”) form a compact unit whereby the N-terminal Ubl domain interacts with the RING1 and IBR domains and portions of the tether region (Fig 1A). This mode of interaction confirmed that the Ubl domain exerted its previously identified autoinhibitory effect (Chaugule et al., 2011) by blocking the expected binding site on the RING1 domain from an E2 conjugating enzyme resulting in negligible ubiquitination activity. Despite the apparent well-folded, compact nature of parkin in this state, a wealth of flexibility exists within the E3 ligase that is not obvious from the crystal structures. NMR dynamics experiments show significant mobility for the IBR domain and segments on either side of a short helix within the tether region where poor electron density is frequently observed in crystal data (see fig S4 in Kumar et al., 2015). Details of the partial E3 ligase activation of parkin have been shown in complexes with phosphorylated ubiquitin (pUb; Kumar et al., 2015, 2017; Wauer et al., 2015), where pUb binds to a broad crevasse between the RING0 and RING1 domains (Fig 1A). The association of pUb causes rearrangement of the IBR domain and results in the formation of a large gap between the IBR and RING1 domains. This results in decreased mobility of the IBR domain due to its juxtaposition with the pUb molecule as shown by heteronuclear nOe experiments (Fig EV1) and increased mobility in the tether region, especially residues V380–A390 that immediately follow the IBR domain. In this optimized state, PINK1 is able to phosphorylate the Ubl domain more efficiently (Ordureau et al., 2014; Aguirre et al., 2018) resulting in nearly 10-fold decreased affinity of the pUb domain and increased affinity for pUb at the RING0/RING1 interface. This allosterically releases the pUb domain from its interaction with the RING1 domain (Kumar et al., 2015; Sauvé et al., 2015; Aguirre et al., 2017) although no structures of this complex for full-length parkin exist. To assess this event, the structure and dynamics of autoinhibited parkin and those of phosphorylated parkin in complex with pUb (pParkin:pUb) were probed by HDX experiments measured by mass spectrometry (Fig IB–D and Appendix Fig S1). These experiments were used to provide a measure of the accessibility of the backbone amides to solvent and
the strength of hydrogen bonding in parkin. Based on previous structural data, these experiments should probe at least three aspects of parkin activation: (i) the binding site for the pUb molecule, (ii) changes in the position of the Ubl domain upon phosphorylation and (iii) indirect structural changes that result from either of these events. The HDX data, which show 91% coverage for parkin (Appendix Fig S2), highlight that many regions of the RING0 and RING1 domains exhibit the largest differences in their HDX properties (Fig 1C). For example, a series of peptides covering regions of the RING0 (Y147-Q155, F209-A225) and RING1 (F277-L283, I298-E309, Y312-E322, V324-L331) domains show slower exchange in the pParkin:pUb state due to protection of these regions by pUb binding at the RING0/RING1 interface (Fig 1D). Coincident with this, peptides at the extreme C-terminus of the IBR domain (C365-A379), the tether region (Y391-Q400) and nearly the entire Ubl domain show increased deuterium exchange (Fig 1C and D). One region of the Ubl domain (R42-E49) shows slower exchange upon phosphorylation. This is likely a direct effect of phosphorylation only since several residues in this region (I44, A46, G47) have decreased amide exchange in the isolated Ubl domain (Aguirre et al., 2017). These observations show the pUb domain is no longer bound at the IBR/RING1 interface and, consistent with previous NMR, sedimentation velocity and computational experiments, indicates the pUb domain adopts a range of bound/free conformations in the pParkin:pUb state (Fig 1D; Caulfield et al., 2014; Aguirre et al., 2017). In this scenario, release of the pUb domain exposes the predicted E2–Ub binding site, based on other RING/E2 complexes (Lechtenberg et al., 2016; Dove et al., 2017; Yuan et al., 2017).

The Parkin/UbcH7–Ub complex reveals an E2–Ub conformation poised for Ub transfer

Phosphorylation of Ub, its recruitment to parkin and subsequent phosphorylation of the Ubl domain result in an increased affinity for the UbcH7–Ub conjugate (Kumar et al., 2015) used by parkin to efficiently transfer ubiquitin to the RING2(Rcat) domain and propagate the ubiquitination reaction. As a first step to identify how UbcH7–Ub stimulates the transfer of its Ub cargo to the catalytic cysteine within the RING2(Rcat) domain, we used NMR spectroscopy to examine how the E2–Ub complex is recruited to the R0RBR C-terminal region of parkin. This work required multiple modes of isotopic labelling, chemical shift assignment of the selected proteins and in some cases purification of the assembled complexes. For example, the chemical shift assignment of R0RBR parkin was completed using TROSY-based triple-resonance experiments using 1H,13C,15N R0RBR parkin (Kumar et al., 2015). A non-hydrolysable E2–Ub conjugate was assembled using a triple-substituted UbcH7 protein (UbcH71H15N/C17S/C668K/C137S) to allow formation of the isopeptide linkage between the UbcH7 and Ub (UbcH7–Ub) and prevent oxidation of the E2 enzyme during NMR experiments. Due to the size of the complexes formed, which compromises the quality of NMR spectra, we utilized mixtures containing a triple-labelled (2H,13C,15N) protein titrated with a deuterated partner. For example, 1H13C15N-labelled R0RBR parkin was complexed with invisible 2H-labelled pUb and purified to homogeneity by size-exclusion chromatography (R0RBR:pUb) prior to the addition of 2H-labelled UbcH7–Ub conjugate.

Initial chemical shift perturbation experiments monitored the 1H–15N TROSY spectra of the 1H,13C,15N-labelled R0RBR–2H-labelled pUb complex with 2H-labelled UbcH7–Ub. These data show that many resonances in R0RBR parkin shift in the presence of E2–Ub and undergo line broadening indicative of a nearly 70-kDa complex being formed (Fig 2A). Formation of the UbcH7–Ub complex with R0RBR:pUb occurs with slow exchange binding kinetics indicative of a KD ≲ 1 μM measured previously by isothermal titration calorimetry (Kumar et al., 2015, 2017). Surprisingly, spectral changes were obvious in two distinct regions of R0RBR—one region where residues are mostly surface exposed and a second region where residues are largely buried (Figs 2B and EV2). The first area consists of resonances belonging to surface residues within the RING1, IBR and tether region indicative of a canonical E2/RING E3 binding interface as well as the Ub interaction site. Resonances affected by E2 binding were located in L1 (C241, T242), helix H1 (L266, V269-L272), L2 (C293) and the tether (Q389-R392, D394-R396) regions of R0RBR while residues in the IBR domain (V330-R334) and tether (A379-F381) corresponded to the Ub binding region (Figs 2B and EV2A). The E2 binding region was verified through separate NMR experiments that monitored the individual binding of 2H-labelled UbcH7 with 1H,13C,15N-labelled R0RBR (Fig EV2B). This experiment showed more limited chemical shift changes but included distinctive line broadening and loss of signals for the L1, helix H1 and L2 regions of the RING1 domain confirming the E2 interaction site with little effect on resonances from residues in the IBR domain.

Reciprocal NMR experiments were conducted to identify the binding surfaces for the UbcH7 and Ub components within the UbcH7–Ub conjugate upon binding to the R0RBR:pUb complex. In these experiments, we titrated size-exclusion purified 2H–R0RBR:2H-pUb complex into a solution of 1H,13C,15N-labelled UbcH7–Ub conjugate and monitored chemical shift changes using 1H–15N TROSY NMR spectroscopy (Appendix Fig S3). These experiments were complicated because UbcH7–Ub predominantly forms a closed state in the absence of a binding partner, but reverts to an open state upon binding to an RBR E3 ligase (Dove et al., 2016). Thus, we expected to see chemical shift changes in UbcH7–Ub that reflected both conversion to the open state and binding to R0RBR:pUb. The closed state of UbcH7–Ub was easy to identify based on previous experiments (Dove et al., 2016). Initial 1H–15N HSQC spectra of the UbcH7–Ub conjugate showed numerous resonances in UbcH7 (F22, V40, N43, N56, K100-N113) and Ub (G47-L50) that reflect close proximity between helix H2 in UbcH7 and Ub (Appendix Fig S4). Upon binding to R0RBR:pUb, most of these signals return to a similar position as found in the unconjugated (free) UbcH7 protein indicating Ub is not occupying the closed position. These observations indicate that parkin converts the UbcH7–Ub conjugate from the predominantly closed state to a more open conformation, a similar observation as made for the RBR E3 ligase HHARI (Dove et al., 2016). Indirectly this also shows the helix H2 region (K100-N113) in UbcH7 and the K48 loop in Ub are not at the R0RBR interface. Thus these chemical shift changes were not considered to map the binding interface in the complex. In contrast to the changes observed for conversion of the closed to the open states of UbcH7–Ub, titration of the R0RBR:pUb complex into the UbcH7–Ub conjugate resulted in broadening of many resonances. Notably, several signals in both UbcH7 and
Ub underwent significant chemical shift changes or shifted such that they could not be identified in spectra, where the UbcH7-Ub conjugate was saturated with the R0RBR:pUb complex, due to slow or slow–intermediate chemical shift exchange. These resonances correspond to those in helix 1 (R6, M8 and L11), loop 4 (A59, F63) and loop 7 (N94, K96, A98) of UbcH7 and the β1–β2 loop (V5, T7, L8, T9 and I13), the linker following helix α1 (E34, I36) and C-terminus (V70, G75) of Ub (Appendix Fig S3).

We used the results from chemical shift perturbation experiments to determine a model of the UbcH7-Ub conjugate bound to R0RBR:pUb using HADDOCK (Dominguez et al., 2003). This was done using the crystal structures of R0RBR:pUb (PDB 5N2W), UbcH7 (PDB 4Q5E) and Ub (PDB 1UBQ) as starting points and imposing distance restraints between the proteins to conduct three-molecule docking of UbcH7 and Ub to the R0RBR:pUb complex. Distance restraints were selected for residues that became unobservable or shifted more...
than one standard deviation above the average shift (Fig EV2 and Appendix Fig S3), and had a side chain surface exposure > 20%. We also included neighbouring residues to those with large chemical shift changes that were solvent exposed (“passive” residues according to the HADDOCK protocol). This resulted in a total of 23 ambiguous distance restraints between R0RBR and the UbcH7 moiety and 25 ambiguous restraints between the R0RBR and Ub proteins. A single unambiguous restraint was used to mimic the isopeptide bond between the carboxylate in G76 of Ub and the side chain amine from K86 in UbcH7 during calculations. The resulting models showed the location of the UbcH7 and Ub proteins with respect to R0RBR:pUb is similar in all 100 water-refined complexes. The best 20 complexes have a backbone RMSD of 0.71 ± 0.10 for R0RBR:pUb:UbcH7-Ub. Comparison of the structures shows that some variation in the orientation of the Ub molecule is observed that was not evident for R0RBR, pUb or UbcH7 proteins in the models. This may indicate the Ub protein in the UbcH7-Ub complex is more dynamic (Fig EV3) than the remainder of the complex. In the lowest energy structure (Fig 2C), the UbcH7-Ub conjugate takes on an open conformation in the complex where the UbcH7 moiety interacts mostly with the RING1 domain and the tether region while the Ub molecule interacts with the RING1/IBR pocket and the N-terminus of the tether region. In the complex, helix H1 (R6, K9) and loop L4 (E60, F63-K64) in UbcH7 contact the R0RBR loop L1 (T240, T242) and helix H1 (L266, T270, R271, D274) respectively in RING1 (Fig 2C). As with other E2/RBR E3 ligase complexes (Lechtenberg et al., 2016; Dove et al., 2017; Yuan et al., 2017), the L7 loop (N94, K96, P97, A98) in UbcH7 sits adjacent to loop L2 (V290-G292) in RING1, but has additional contacts with Y391-D394 just prior to the short helix in the tether region. We noted that some of the largest

Figure 2.
chemical shift changes were in this region of the tether (Y391, R392, D394), which undergoes multiple rearrangements in crystal structures of parkin (Kumar et al., 2017). NMR dynamics experiments (Fig EV1; Kumar et al., 2015) and a structure of the isolated IBR-tether–RING2(Rcat) region (Spratt et al., 2013) indicate this region is very flexible and likely adopts multiple conformations in solution. We interpreted the chemical shift changes within the tether region to result from both direct E2 binding and an altering of the tether position to accommodate the E2 enzyme. The open arrangement of the E2-Ub bound to R0RBR parkin more closely resembles the interaction of Ubch5b-Ub with HOIP (Lechtenberg et al., 2016) than either of the structures for Ubch7-Ub with HHARI (Dove et al., 2017; Yuan et al., 2017; Fig EV4). Ub binding is governed predominantly by contacts from β1-L1-L2 (K6, L8, K11, I3-T14), the linker following helix α1 (Q31-D32) and C-terminus (L73, R74) to an R0RBR surface including β1 (P333, P335) and the C-terminus of the IBR domain (E370) and adjacent tether (V380, F381, S384, T386), RING1 helix H1 (N273) and the straightened RING1 helix H3 (R314, Y318). The Ub binding site of the IBR domain from Ubch7-Ub binding agrees well with potential ubiquitin-binding regions (UBR2, UBR3) inferred from crystallographic studies and supported through ubiquitination activity assays (Kumar et al., 2017). Further, the location of the Ub molecule provides insight into the next step of the ubiquitination process, the transfer to the RING2(Rcat) domain. The Ub conjugate is positioned such that two hydrophobic regions including the I44 patch and the C-terminus (V70, L71) of Ub are pointed away from the RING1 domain and exposed to solvent. Although not identical, the activated Ub in the HOIP/Ubch5b complex and donor Ub in the RING2L transfer complex are positioned similarly (Stiegitz et al., 2013; Lechtenberg et al., 2016). These hydrophobic regions (I44, V70, L71) are used to recruit helix β2 from the catalytic RING2L domain in the domain-swapped dimer structure (Lechtenberg et al., 2016). This suggests that a similar mechanism might exist for parkin whereby the RING2(Rcat) domain is repositioned adjacent to the hydrophobic sites in Ub. The orientation of the Ubch7-Ub conjugate in our structure poises the C-terminus of Ub for translocation by exposing the G76 carboxyl in the isopeptide linkage towards the tether side of parkin. This arrangement suggests that nucleophilic attack by the catalytic C431 in the RING2(Rcat) domain would come from this direction (backside as shown in Fig 2C). The structure also shows that helix H2 in Ubch7, previously used to interact with Ub in the closed E2-Ub conformation, is exposed on the same side as the C86K-G76 linkage. Overall, the current structure shows how Ubch7 might position its Ub cargo for transfer to the catalytic cysteine (C431) of the RING2(Rcat) domain and provides clues that suggest the RING2(Rcat) domain is eventually repositioned near the Ubch7-Ub conjugate to promote Ub transfer.

In order to further test the parkin recruitment site for Ubch7-Ub observed in our models, a series of ubiquitination assays were performed. We first analysed the observed ubiquitin surface involved at the Parkin:Ubch7-Ub interface using distinct assays that monitor transfer of ubiquitin from the E2 onto parkin and subsequently onto the substrate. To assess Ubch7 mediated ubiquitin loading of parkin, we generated a pParkin RING2(Rcat) mutant (C431S H433A, referred to as pParkinCH) that is able to trap an E3–Ub oxyester intermediate (Spratt et al., 2013; Kumar et al., 2017). We rationalized that ubiquitin loading to pParkinCH would depend on the interaction of Ub in the Ubch7–Ub conjugate with pParkin. In the presence of ATP, E1, Ubch7 and wild-type ubiquitin, we observe the formation of a pParkin CH–CbUb intermediate (Fig 2D, top). In contrast, alanine mutants of ubiquitin β1-L1-L2 residues (K6A, L8A, K11A) observed at the Ub interface with the parkin IBR domain result in defective ubiquitin transfer onto parkin even in the presence of pUb. Consequently, these ubiquitin mutants were also compromised in pParkin-mediated substrate (HA-tagged Miro1) ubiquitination (Fig 2D, bottom). The ubiquitin mutant E34A shows minimal effects suggesting the C-terminus of the long helix in Ub has a lesser role in directing the Parkin:Ubch7–Ub interface. A similar observation was made for the HOIP:Ubch5b-Ub complex (Lechtenberg et al., 2016). In a complementary approach, we tested the Ubch7-Ub interface with parkin by using ARJ and non-ARJ parkin variants to assess E3 ligase activity (Fig 2E). All assays were done in the absence and presence of parkin mediated substrate (HA-tagged Miro1) ubiquitination (Fig 2D, bottom). The ubiquitin mutant E34A shows minimal effects suggesting the C-terminus of the long helix in Ub has a lesser role in directing the Parkin:Ubch7–Ub interface. A similar observation was made for the HOIP:Ubch5b-Ub complex (Lechtenberg et al., 2016). In a complementary approach, we tested the Ubch7-Ub interface with parkin by using ARJ and non-ARJ parkin variants to assess E3 ligase activity (Fig 2E). All assays were done in the absence and

Figure 2. Model of the E2-Ub conjugate bound to pUb-activated parkin.

A Portion of the 1H,13C,15N TROSY NMR spectrum showing signals from R0RBR parkin within the R0RBR:pUb complex (black) and in the presence of one equivalent of the unlabelled Ubch7-Ub conjugate (blue). Signals that shift in the presence of Ubch7-Ub are indicated by arrows. Some signals shift and broaden and cannot be identified in the bound state. Boxes indicate signals that have shifted and are visible at very low contour level.

B Two distinct surfaces are revealed on parkin upon binding of the Ubch7-Ub conjugate. Chemical shift perturbations were measured from 1H,13C,15N-labelled R0RBR parkin bound to unlabelled pUb in the absence and presence of one equivalent unlabelled Ubch7-Ub. Chemical shift perturbations (absence of signal, average shift + 1 SD) are modelled onto the surface of R0RBR bound to pUb (PDBe5N2W). The Ubch7-Ub binding site comprises the RING1 and IBR regions (magenta). An adjacent site composed of many buried residues was also observed (cyan) that results from E2-Ub binding but does not include the E2 binding site. Sections of the tether and linker regions not visible in the crystal structure were added using Modeller (Eswar et al., 2006) so that chemical shift changes for residues not observed in crystal structures could be mapped.

C Model of R0RBR-pUb in complex with Ubch7-Ub derived from NMR chemical shift data and HADDOCK docking (Dominguez et al., 2003). The lowest energy structure is shown, and the top 20 structures all showed excellent agreement (RMSD 0.71 ± 0.1 Å) although some minor differences were noted for the orientation of the Ub molecule. The structure shows that Ubch7-Ub binds in an open conformation. Ubch7 uses canonical E2–RING3 interactions that include residues from two loops in the RING1 domain (L1, L2) and two loops in the E2 enzyme (L1, L2) to stabilize the interaction. Ub binds to a RING1/IBR pocket. In this model, no attempt was made to move or re-orient any of the domains in parkin.

D Effects of ubiquitin surface mutants based on the Ubch7-Ub:R0RBR-pUb model for the ubiquitin loading and off-loading potential of pParkinCH (pParkinCH–C431S H433A). Coomassie-stained gels (top) depict the formation of a pParkinCH–Ub oxyester intermediate after 60 min in the absence and presence of pUb using the indicated ubiquitin species. Anti-HA blots (bottom) depict ubiquitination of HA-tagged Miro1181H–Ub derived from NMR chemical shift data and HADDOCK docking (Dominguez et al., 2003). The lowest energy structure is shown, and the top 20 structures all showed excellent agreement (RMSD 0.71 ± 0.1 Å) although some minor differences were noted for the orientation of the Ub molecule. The structure shows that Ubch7-Ub binds in an open conformation. Ubch7 uses canonical E2–RING3 interactions that include residues from two loops in the RING1 domain (L1, L2) and two loops in the E2 enzyme (L1, L2) to stabilize the interaction. Ub binds to a RING1/IBR pocket. In this model, no attempt was made to move or re-orient any of the domains in parkin.

E Autoubiquitination assay for parkin using ARJ and non-ARJ substitutions in parkin observed near the interface with Ubch7-Ub. Assays were done in the absence and presence of pUb. Experiments with PINK1 were done by treating parkin:pUb with PINK1 for 30 min prior to adding other reagents needed for ubiquitination. Assays were monitored using a Dylight-labelled Ub protein and measuring fluorescence at 800 nm.
The presence of pUb or with PINK1 preincubated with parkin and pUb to enable phosphorylation of the Ubl domain. As expected, significant increases in ubiquitination were observed in the presence of both phosphorylation steps. ARJP variant T240R in the RING1 helix H1 shows significant decreases in activity due to disruption of interactions with F63 and P97 in UbcH7. The P335C/C337G substitution in parkin expected to disrupt one of the Zn-binding sites in the E2 domain that interacts with K11, T13 and H14 of ubiquitin also had diminished activity. The R271S substitution that is near the UbcH7 L4 loop in our structure had a minor decrease consistent with modification of that interaction. The activity of two other substitutions, N273S and R314A, had similar activities to that of the wild-type E3 ligase, likely a reflection of a weakened interaction with the Ubl facilitating its phosphorylation and stimulating ubiquitination as previously observed (Sauvé et al., 2015).

**UbcH7-Ub binding leads to a re-modelled RING0/RING2 interface in parkin**

In addition to the UbcH7-Ub binding site on parkin, analysis of chemical shift perturbation experiments of 2H,13C,15N-labelled R0RBR:pUb and unconjugated UbcH7 revealed many changes localized at the interface between the C-terminus of the tether region (A398-T414), RING0 (S145, F146), RING1 (Q252, R256, H257) and RING2(Rcat); T415, E426, K427, N428, D464; Figs 2B, and 3A and B). In contrast, NMR experiments performed between 2H,13C,15N-labelled R0RBR:pUb and unconjugated UbcH7 revealed many of these chemical shift perturbations are absent, or present to a lesser extent (Fig EV2), suggesting the intact UbcH7-Ub conjugate is necessary to induce these additional changes. Three-dimensional structures show that most of the affected residues form a cluster anchored by W403 that follows the short helix in the tether region and is essential for packing the tether against the RING0/RING1/RING2(Rcat) core (Fig 3A and B). In the absence of Ub or Ubl phosphorylation by PINK1, a W403A parkin variant dramatically increases ubiquitination activity (Trempe et al., 2013), likely a result of a structural rearrangement near W403 and exposure of the subsequent catalytic C431 in the RING2(Rcat) domain. Therefore, we hypothesized that UbcH7-Ub binding to R0RBR:pUb might cause a similar rearrangement near the RING0/RING1/RING2(Rcat) core in the wild-type protein as in the W403A substituted version (R0RBRW403A). To test this, we compared the positions of resonances from a 15N-labelled R0RBRW403A HSQC spectrum with those from a 15N-labelled R0RBR spectrum (Appendix Fig S5). Remarkably, in the R0RBRW403A data many signals from residues that neighbour W403 are either undetectable or undergo significant chemical shift changes similar to those observed upon UbcH7-Ub binding to R0RBR:pUb but not directly at the E2 binding site. The similarity of the chemical shift changes and the buried nature of many of these residues suggest the RING0/RING1/RING2(Rcat) interface is re-modelled during the E2–Ubl interaction with parkin.

**The UbcH7-Ub conjugate enhances pUbl domain re-binding to parkin**

Upon presentation of pUb and phosphorylation of the Ubl domain of parkin, both HDX and NMR dynamics experiments show the pUbl domain is dislodged from the autoinhibitory site against the IBR and RING1 domains. This is supported by the nearly 10-fold poorer affinity of the Ubl domain for R0RBR parkin upon phosphorylation (Kumar et al., 2015). In order to test how the UbcH7-Ub conjugate might alter the interaction of the pUbl domain with the remainder of the E3 ligase, we again used HDX mass spectrometry experiments of phosphorylated parkin bound to pUb (pParkin:pUb), this time in the presence of UbcH7-Ub. The dissociation constant for this complex is near 0.5 μM, so these experiments were done using 10 μM pParkin:pUb in the presence of non-hydrolysable UbcH7-Ub (10 μM) to achieve more than 80% saturation of pParkin:pUb with the E2-Ub conjugate during HDX measurements.

In the presence of the E2-Ub complex, the majority of parkin shows greater exchange than the phosphorylated, activated state (Fig 4A and Appendix Fig S6). In general, this indicates that a global rearrangement has occurred in the protein, stimulated by UbcH7-Ub conjugate binding. For example, the HDX data show significant increases in exchange for the RING0 and RING2(Rcat) domains.

This observation is incongruous with current structures of parkin, and our HDX data for pParkin:pUb, that show most of these domains are protected from solvent and not exposed. In contrast, these regions correspond closely to the RING0/RING1/RING2(Rcat) interface with the tether region suggested to be re-modelled based on NMR chemical shift perturbation experiments (Fig 3A and B). The increase in deuteration upon UbcH7-Ub binding indicates this interface is becoming much looser with a loss of both hydrophobic and hydrogen bonding interactions. One possible interpretation of this result is that the RING2(Rcat) domain is displaced from the interface exposing its surface, the RING0 interface with RING2(Rcat) and the C-terminus of the tether (Fig 4B).

Coincident with the changes in HDX for the RING0 and RING2(Rcat) domains, numerous increases are noted near the pUb interface with the RING0/RING1 cleft and extending towards the IBR domain. These regions undergo multiple rearrangements in structure and orientation upon pUb and Ubl binding (Kumar et al., 2015; Wauer et al., 2015; Kumar et al., 2017). The increase in HDX could indicate further rearrangement occurs upon UbcH7-Ub binding leading to a more extended structure. The HDX experiments also show that the pUb domain remains exposed to solvent. One exception is the C-terminus of the pUb domain (D62-M80) that is significantly more protected. This region contains the phosphorylated S65 residue suggested to interact with three basic residues (K161, R163 and K211) in the RING0 domain that show attenuated parkin autoubiquitination when substituted (Wauer et al., 2015). Two of these sites are locations of ARJP substitutions (K161N, K211R, K211N). A minority of parkin crystal structures also show a bound sulphate ion in this area. Although we did not observe a decrease in HDX in this region of the RING0 domain, the decreased exchange in the N-terminus of the pUbl domain could be consistent with increased binding to the RING0 domain stimulated by the UbcH7-Ub conjugate.

To test how UbcH7-Ub binding might influence the potential relocation of the pUb domain, we examined a series of 1H-15N HSQC spectra for 15N-labelled phosphorylated parkin in complex with unlabelled pUb and compared this to the isolated pUb domain. We then compared spectra of 15N-labelled pUb in the absence and presence of R0RBR:pUb and UbcH7-Ub. In full-length
phosphorylated parkin, the $^{1}H$-$^{15}N$ HSQC spectrum is very complicated due to a large number of signals that are visible from the highly flexible linker and tether regions in the protein (Fig 4C). However, nearly all signals are visible from the pUbl domain in the full-length protein when a short relaxation delay is used in the HSQC experiment to suppress broad signals arising from the compact, folded R0RBR region (see also fig S9 in Aguirre et al., 2017). When compared to the $^{1}H$-$^{15}N$ HSQC spectrum of the isolated pUbl domain (Fig 4C), it is clear that very small chemical shift changes and decreases in intensities occur for residues pSer65, D62, Q63 and Q64, consistent with a weak interaction of the pUbl with the remainder of the protein. However, many other signals from the pUbl are hardly attenuated (Aguirre et al., 2017). These observations are consistent with our HDX experiments, which indicate the pUbl domain spends most of its time dissociated from the remainder of the protein in the absence of an E2-Ub conjugate (Fig 1D).

When the $^{1}H$-$^{15}N$ HSQC spectrum of $^{15}N$ pUb is compared to that with added R0RBR:pUb, we observe no significant chemical shift changes (Fig 4D). This supports previous observations of a weak interaction for the pUbl domain with R0RBR parkin in trans (Kumar et al., 2015; Sauvè et al., 2015). Upon addition of UbcH7-Ub, we observe small but measurable changes for the positions of several signals including pSer65, D62, Q63 and Q64 (Fig 4D). These chemical shift changes occur on the fast exchange timescale consistent with weak binding of the pUbl domain to R0RBR parkin that is accentuated upon UbcH7-Ub recruitment by the E3 ligase. This indicates that the pUbl domain undergoes a weak interaction with the remainder of the parkin protein that is stimulated through binding of the UbcH7-Ub conjugate.

**Ubch7-Ub and pUb work together to modulate C431 reactivity**

Our NMR experiments using R0RBR parkin allowed us to determine the UbcH7-Ub binding site within parkin as one requirement for ubiquitination and have identified how UbcH7-Ub binding might re-model the RING0/RING1/RING2(Rcat) interface with the tether. However, in the absence of the pUbl domain these experiments are deficient in establishing how the pUbl might work with an E2-Ub conjugate to achieve optimal activity. In particular, all three-dimensional structures to date have been unable to show how the reactivity of the catalytic C431 residue in the RING2(Rcat) domain might be altered upon phosphorylation of the Ub domain and addition of the E2-Ub conjugate. In order to assess this, we examined the ability of full-length parkin and R0RBR parkin to form a non-hydrolysable ubiquitin adduct using ubiquitin-vinyl sulphone (UbVS). Although smaller probes to test E3 ligase activity are available (Pao et al., 2016), we used UbVS to mimic the parkin-Ub product that would be expected following Ub transfer from the UbcH7 enzyme. Using different combinations of parkin, phosphorylated parkin, phospho-ubiquitin and UbcH7-Ub, this approach tested the reactivity, and hence accessibility, of the catalytic C431 in the RING2(Rcat) domain during each step of the parkin activation cycle (Fig 5).

As expected, autoinhibited and pUb-bound parkin or R0RBR showed minimal reaction with UbVS (Fig 5 and Appendix Fig S7) in agreement with previous three-dimensional structures and reactivity profiles that indicate the catalytic C431 is mostly occluded by neighbouring RING0 domain interactions (Riley et al., 2013; Wauer & Komander, 2013). Remarkably, phosphorylated parkin activated by pUb shows rapid product formation with UbVS (Fig 5A and B),
visible even after 1 min (not shown). A similar reaction with R0RBR:pUb when the parkin pUbl domain is added in trans shows little modification with UbVS even after 60 min (Fig 5C and D). The most logical explanation for these observations is that the pUbl domain is facilitating access of the UbVS probe to the catalytic C431 site in the RING2(Rcat) domain by binding to another region in parkin. Since the pUbl interaction exhibits a weaker affinity in trans, the effect of the UbVS probe is much lower than observed for the intact protein. Interestingly, introduction of UbcH7-Ub to either the pParkin:pUb sample (Fig 5A and B) or the pUbl:R0RBR:pUb complex (Fig 5C and D) leads to opposite results. In the full-length protein, we observe a reproducible lower conversion rate in the presence of the E2-Ub conjugate than in its absence (Fig 5A and B). In the pUbl:R0RBR:pUb sample, the reactivity to UbVS is enhanced in the presence of UbcH7-Ub (Fig 5C and D). At a minimum, this shows that the pUbl domain and E2-Ub act synergistically to alter the reactivity of the catalytic C431 residue in the RING2(Rcat) domain. Introducing the “activating” W403A mutation did not
significantly increase reactivity with the UbVS probe, suggesting the re-modelling observed in our NMR experiments may not directly expose parkin’s catalytic Cys431 (Appendix Fig S7). In agreement with our NMR data (Fig 4), the E2-Ub conjugate has the ability to re-model the RING0/RING2(Rcat) interface and increase binding of the pUbl domain in trans that results in increased reactivity of the catalytic C431 residue. Meanwhile, the small decrease in C431 accessibility in the pParkin:pUb complex in the presence UbcH7-Ub suggests a further conformational change occurs due to E2 binding that decreases the availability of the catalytic site. This is in agreement with our HDX experiments that suggest a reorganization of the RING2(Rcat) domain (Fig 4).

Discussion

Although parkin was originally thought to be a constitutively active enzyme, it is now known to regulate its activity through intramolecular domain–domain interactions and binding to effectors (Chaugule et al., 2011; Kumar et al., 2015; Sauvé et al., 2015). In particular, PINK1 regulates parkin activity through phosphorylation of the Ubl domain, and of ubiquitin itself (Kumar et al., 2015; Sauvé et al., 2015; Wauer et al., 2015; Kumar et al., 2017), which then acts as an effector. Multiple structures of autoinhibited parkin reveal that the E2 binding site is blocked; however, static crystal structures have shown that binding to pUb does not render the proposed E2 binding site, nor the phosphorylation site in the Ubl domain accessible (Fig 6A). Comparison of crystal structures in the absence/presence of the Ubl domain reveals reorganization of residues between the RING0/RING1 interface and multiple arrangements of the IBR domain, shown to be flexible by previous NMR dynamics experiments (Kumar et al., 2015). Though pUb binding to parkin decreases the affinity of R0RBR for the Ubl domain, it is not sufficient to dislodge the Ubl domain in the crystal (Kumar et al., 2015, 2017). Rather, this step optimizes parkin for Ubl phosphorylation and E2 bind-

The pUbl domain and UbcH7-Ub synergistically modulate catalytic C431 reactivity.

A Exposure of the catalytic Cys431 is parkin as assessed by reaction with a UbVS probe. The different stages of activation using parkin, parkin:pUb, pParkin:pUb and pParkin:pUb in combination with an isopeptide-linked UbcH7-Ub conjugate are indicated above each gel. Following addition of UbVS, samples were taken at the times indicated (0–10 min) and visualized by SDS–PAGE.

B Relative percentages of pParkin and the pParkin-Ub adduct as a function of time. Intensity percentages were calculated as a function of total intensity of pParkin-Ub, parkin and UbVS/pUb bands. Error bars represent standard deviation from the average for duplicate measurements.

C Exposure of the catalytic Cys431 is R0RBR parkin as assessed by reaction with a UbVS probe. The different stages of activation using R0RBR, R0RBR:pUb, R0RBR:pUb: pUbl and R0RBR:pUbl:pUbl in combination with an isopeptide-linked UbcH7-Ub conjugate are indicated above each gel. Following addition of UbVS, samples were taken at the times indicated (0–60 min) and visualized by SDS–PAGE.

D Relative percentages of R0RBR and the R0RBR-Ub adduct as a function of time. Intensity percentages were calculated as a function of total intensity of the R0RBR-Ub, R0RBR and UbVS/pUb/pUbl bands.
(Gladkova et al., 2018; Sauvé et al., 2018) show that the pUbl domain is nearly completely exposed to solvent in this state compared to its position in autoinhibited parkin where it is surrounded by the RING1, IBR and tether regions. This interpretation agrees with NMR relaxation data, analytical centrifugation experiments and computational work that conclude the pUbl domain samples a large conformational space that includes weak interaction with other regions of parkin (Caulfield et al., 2014; Aguirre et al., 2017; Fig 6C). While this manuscript was under review, two crystal structures of partially activated parkin appeared, which showed the pUbl domain has the ability to bind to the RING0 domain (Gladkova et al., 2018; Sauvé et al., 2018) comprised of a previously identified (Wauer & Komander, 2013) basic patch (K161, R163, K211) that included two ARJP substitutions. Mutation of these residues renders parkin unreactive with an E2-based activity probe, consistent with a requirement for pUbl interaction (Pao et al., 2016). Yet other experiments show parkin retains appreciable ubiquitination ability in the absence of its Ubl domain (Chaugule et al., 2011; Kazlauskaite et al., 2014; Kumar et al., 2015), suggesting phosphorylation and the Ubl domain itself are less important. Our NMR data of full-length pParkin:pUb (Fig 5) indicate the native interaction of the pUbl domain with the RING0 domain is weak in this state in agreement with previous affinity experiments (Kumar et al., 2015; Sauvé et al., 2015). Consistent with this, it was necessary to remove the entire RING2(Rcat) domain and the tether region from parkin in recent structures in order to capture the bound pUb state (Gladkova et al., 2018; Sauvé et al., 2018), suggesting removal of the RING2(Rcat) domain may enhance the pUb interaction through removal of steric hindrance.

One of the main outcomes of pUb recruitment and subsequent phosphorylation of the Ubl domain is to unmask the RING1 binding site for the E2 conjugating enzyme and rearrangement of the IBR domain to engage the conjugated Ub molecule (Kumar et al., 2015; Kumar et al., 2017). This interaction with UbcH7-Ub shows that the donor Ub is in the open conformation favoured by RBRs (Dove et al., 2016). The interaction site of UbcH7 with the RING1 domain in our structure is in agreement with previous crystal structures of HOIP with UbcH5b-Ub (Lechtenberg et al., 2016) and HHARI with UbcH7-Ub (Dove et al., 2017; Yuan et al., 2017) although minor orientation differences occur. This likely arises due to differences in the L2 loop regions of the RBR E3 ligases as previously noted (Spratt et al., 2014; Dove et al., 2016), and in particular differences in the linchpin residue that directs the E2–Ub conjugate to its open state during interaction (Dove et al., 2017). The position of the UbcH7 enzyme (in UbcH7-Ub) with parkin is also nearly identical to a recent structure of truncated parkin in complex with an unconjugated E2 enzyme (Sauvé et al., 2018). This indicates the donor Ub does not play a major role in directing the E2 binding in agreement with its poor binding affinity for parkin on its own.

Our NMR data indicate that UbcH7-Ub binding to partially activated parkin re-models the hydrophobic cluster involving W403 in the tether region at the junction of the RING0/RING1/RING2(Rcat) domains. This is supported by substitution of W403 that produces NMR chemical shift changes analogous to those for binding of UbcH7-Ub. Further, experiments using an E2-based activity probe show a W403A substitution can partially recapitulate catalytic cysteine labelling of parkin even in the absence of Ubl domain phosphorylation (Pao et al., 2016).

Together these observations suggest that it is the E2–Ub binding step that induces a conformational change in the W403 cluster (Fig 6D), rather than W403 being re-modelled to allow E2–Ub engagement, that facilitates the interaction of the pUb domain with the RING0 domain of parkin. Although this step could not be shown in recent crystal structures that lacked the RING2(Rcat) domain and tether regions, our HDX data show that both phosphorylation and E2–Ub binding together lead to nearly complete exposure of the RING2(Rcat) domain. The interplay between pUb and E2–Ub binding to modulate ubiquitination efficiency is also borne out from the reactivity of the catalytic cysteine (C431) in RING2(Rcat) to react with a ubiquitin probe. In R0RBR parkin, reactivity required both

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**Figure 6. Model of parkin activation by combined pUb and E2–Ub interactions.**

A Autoinhibited state whereby the Ubl domain masks the E2 binding site on the RING1 (R3) domain as described by Chaugule et al (2011) and supported through crystallographic studies.

B Optimization step controlled by PINK1 phosphorylation of ubiquitin (Ub) and subsequent binding of pUb to the RING1/IBR interface resulting reorganization of the RING0/RING1 interface and movement of the IBR domain. Steps (A) and (B) have been previously described.

C PINK1 phosphorylation of the Ubl domain causes its dissociation from the RING1 binding site allowing it to sample a large conformational space in solution including weak binding to the RING0 domain favouring an uncovering of the catalytic cysteine C431.

D E2–Ub binding to the RING1/IBR domains making use of a cryptic ubiquitin binding site uncovered through dissociation of the pUb domain. Re-modelling of the RING0/RING1/RING2(Rcat) interface with the tether region occurs based on NMR chemical shift data and HDX experiments.

E Fully activated state of parkin utilizes synergistic binding of pUb domain to the RING0 domain observed in crystal structures (Gladkova et al., 2018; Sauvé et al., 2018) and E2–Ub binding that re-models the RING0/RING1/RING2(Rcat) interface to maximize accessibility to the catalytic cysteine.
pUb and E2-Ub addition for adduct formation. In similar NMR experiments, addition of E2-Ub was required to observe binding of pUb to R0RBR parkin. The fast exchange observed in current NMR experiments between the pUb domain and the RING0 site observed in crystal structures (Gladkova et al., 2018; Sauvé et al., 2018) suggests that this interaction is short-lived. We suggest that upon E2–Ub recruitment, the RING2(Rcat) domain is dislodged to provide a more optimal binding site for the pUb domain. Indeed, one recent structure shows that an additional segment of the linker that precedes the RING0 domain takes the place of the RING2(Rcat) domain (Gladkova et al., 2018), perhaps providing an extra level of regulation. Taken together, we suggest that pUb binding alone to the RING0 domain is not sufficient to drive the conformational changes required and that E2–Ub binding is necessary to re-model the W403 region for full activation (Fig 6D and E). These events show that both E2–Ub binding and pUb recruitment to the RING0 domain act synergistically to propagate the ubiquitination reaction.

Recruitment of the UbcH7-Ub conjugate to pParkin:pUb appears to cause a significant conformational change based on HDX and NMR chemical shift experiments. An unresolved question then is: What is the conformational change that occurs that allows ubiquitin transfer from the E2 conjugating enzyme to the RING2(Rcat) domain? It is tempting to consider that the RING2(Rcat) domain repositions itself nearby the open E2–Ub conjugate to accept the donor ubiquitin. In the absence of data to directly show this, several lines of evidence suggest this is a possibility. For example, our HDX data show that although the RING2(Rcat) domain is exposed in the presence of UbcH7-Ub, its extreme C-terminus (E452-W462) is more protected compared to that in the absence of E2–Ub. This portion of parkin has been suggested to form important interactions with the E2–Ub conjugate based on the similarity of hydrophobic residues with HOIP and the position of the swapped RING2L domain in HOIP that neighbours the UbcH5b-Ub conjugate (Lechtenberg et al., 2016). UbVS experiments show that the reactivity of C431 in RING2(Rcat) is slightly diminished in the presence of UbcH7-Ub (compared to its absence in pParkin:pUb), suggesting possible rearrangement and protection might impede its reactivity in this assay. Further, the region immediately preceding the RING2(Rcat) domain contains a ubiquitin-binding motif (Chaugule et al., 2011) that harbours at least one ARJP substitution (T415N) that impairs all ubiquitination activity (Matsuda et al., 2010). A similar ubiquitin-binding motif has been observed for HHR1L, and substitutions in this region show significant decreases in Ub affinity and ubiquitination activity (Dove et al., 2016). Although unstructured in parkin, the corresponding region in HOIP forms a short α-helix (helix h12) that interacts with the 144 patch of the donor Ub in the UbcH5b-Ub complex. These observations hint at a catalytic complex where interactions between both components of the UbcH7–Ub conjugate are important for recruitment of the tether and RING2(Rcat) regions. Alternatively, multiple studies have suggested that the ubiquitin-transfer complex is more complicated and requires co-operation between multiple parkin and E2–Ub molecules (Lazarou et al., 2013; Kumar et al., 2017) to facilitate ubiquitin transfer. An attractive feature of this model, which also utilizes alternate ubiquitin binding sites, is that it provides a framework for the processivity of autoubiquitination observed for parkin. With this in mind, it is worth noting that recent crystal structures lack structural resolution of several linkers in parkin that preclude identification of intra- vs. intermolecular interactions of the pUb or E2 enzyme with the RING0 or RING1 domains, respectively (Gladkova et al., 2018; Sauvé et al., 2018). By the same note, our NMR-based model of UbcH7-Ub in complex with R0RBR:pUb is unable to identify how the tether and RING2(Rcat) might be rearranged upon UbcH7-Ub and pUb engagement. Nevertheless, our study shows that synergy exists between these two effectors to fully control parkin activity. Understanding the next steps in detail will be essential to target this important enzyme for modulation during the pathogenesis of Parkinson’s disease.

Materials and Methods

Protein constructs and purification

Full-length human parkin (1–465), Ubl (1–76), R0RBR (141–465), Drosophila melanogaster RING2 (410–482) and other parkin variants were expressed and purified as described previously (Chaugule et al., 2011; Spratt et al., 2013). Briefly, His-smt3-parkin constructs were expressed in BL21(DE3) cells at 37°C to an OD600 of 0.8. Expression was induced at 16°C with 25 μM IPTG for parkin, 0.1 mM IPTG for R0RBR and 0.5 mM IPTG for Ubl or RING2 for 18 h. All growths, except the Ubl domain, were supplemented with 0.5 mM ZnCl2. Purification utilized an initial HisTrap FF column followed by Ulpl cleavage at 4°C, a second HisTrap FF column and final Superdex 75 10/300 size-exclusion chromatography. Selectively 2H,13C,15N-labelled R0RBR:pUb or R0RBR and selectively 2H,13C,15N or 2H,15N-labelled Ubl, UbcH7 or UbcH7-Ub were expressed and purified as previously described (Kumar et al., 2015).

His-tagged Uba1 was expressed in BL21(DE3)CodonPlus-RL cells at 37°C to an OD600 of 0.8. Expression was induced with 0.5 mM IPTG at 18°C for 12 h. His-tagged Uba1 was purified on a HisTrap FF column by washing with a buffer containing 50 mM Tris, 200 mM NaCl, 250 μM TCEP and 25 mM imidazole (pH 8.0) and then washing with 14% of elution buffer that contained 250 mM imidazole. The His-tagged Uba1 was then eluted with 100% elution buffer and stored in aliquots at −80°C for ubiquitination assays.

His-TEV-tagged human UbcH7C175,C368,K,C172 - was expressed in BL21(DE3)CodonPlus-RL cells at 37°C to an OD600 of 0.8, and expression was induced with 1 mM IPTG at 30°C for 18 h. UbcH7 was purified on a HisTrap FF column, cleaved at 4°C overnight and purified on a second HisTrap FF column. His-tagged UbcH7 was expressed in BL21(DE3)CodonPlus-RL cells. Complexes that contained 1:1 R0RBR:pUb were formed using a 1:5-fold excess pUb compared to R0RBR in a buffer containing 20 mM Tris, 75 mM NaCl and 250 μM TCEP (pH 8). The 1:1 complex mixture was purified on a Superdex 75 10/300 size-exclusion column to ensure excess pUb was not present in samples for NMR studies.

Glutathione S-transferase (GST)-HA-tagged human Miro1 (residues 181–592) was expressed in BL21(DE3) cells purified using standard protocols (Kumar et al., 2015). The GST tag was removed using GST-3C protease, and the cleaved material was further purified on a Superdex 200 Increase 10/300 size-exclusion column.

Protein phosphorylation

Phosphorylation of Ub, Ubl and parkin was done using purified Pediculus humanus PINK1 (126–575) as described previously...
(Kumar et al., 2015; Aguirre et al., 2017). For pUb and pUbl (1–76), typically 10 μM PINK1 was sufficient to stoichiometrically phosphorylate either 100 μM Ub or Ubl in 1 h at 24°C. For parkin, typically 75 μM PINK1 was sufficient to phosphorylate 150 μM parkin for 3.5 h at room temperature. Reactions were visualized by Phos-tag gel. PINK1 was removed using a GSTrap FF column. Phosphorylated proteins were purified using a Superdex 75 10/300 size-exclusion column and confirmed by mass spectrometry.

**Synthesis of UbcH7-Ub isopeptide-linked conjugate**

UbcH7-Ub isopeptide-linked conjugate was synthesized using an optimized version of the protocol of Plechanovová et al. (2012). Briefly, 200 μM His-tagged Ub, 400 μM UbcH7 C17S/C66K/C137S, 25 μM non-cleavable His-tagged Uba1 and 10 mM Mg2+ /ATP were incubated together in a buffer containing 50 mM CHES and 150 mM NaCl (pH 9.0) at 37°C for 6–16 h to form approximately 60% UbcH7-Ub isopeptide-linked conjugate based on SDS–PAGE analysis. The resulting mixture was passed through a HisTrap FF column to eliminate unconjugated UbcH7. The eluted His-tagged proteins were TEV-cleaved overnight at 4°C. The mixture was purified on a second HisTrap FF column to eliminate non-cleavable His-tagged Uba1. The remaining UbcH7-Ub was separated from unreacted Ub using a HiLoad Superdex 16/60 size-exclusion column.

**NMR experiments**

All NMR data were collected at 25°C on a Varian Inova 600-MHz NMR spectrometer equipped with a triple-resonance cryogenic probe and z-field gradients. Samples were prepared in a buffer containing 25 mM HEPES, 50 mM NaCl and 500 μM TCEP (pH 7.0) with 10% D2O (v/v) using DSS as an internal reference and imidazole to monitor pH. Backbone assignments of R0RBR parkin (Kumar et al., 2015) and the pUbl domain (Aguirre et al., 2017) were completed using standard triple-resonance methods as previously reported. 1H-15N HSQC spectra were collected in TROSY mode (Pervushin et al., 1997) to follow backbone chemical shift perturbations. 1H-15C HMQC spectra (Tugarinov et al., 2004) were collected to monitor chemical shifts of Ub side chain methyl groups. 1H-13C and 1H-15N TROSY spectra were collected using different combinations of 1H-13C, 15N-labelled and 2H-15N-labelled R0RBR:pUb or R0RBR with 2H-13C, 15N or 2H-15N-labelled pUb, Ub, UbcH7 or UbcH7-Ub. For the 13N-labelled pUb binding experiments, 1H-15N HSQC experiments were collected to monitor pUb chemical shift changes with addition of equimolar amounts of unlabelled R0RBR:pUb and UbcH7-Ub. Chemical shift perturbation measurements for amide backbone resonances were calculated using δ = (δH 2 + δN/5)1/2 and for side chain methyl groups using δ = (δH 2 + (δC/3.3) 2)1/2. All data were processed using 60°-shifted cosine bell-weighting functions using NMRPipe and NMRDraw (Delaglio et al., 1995) and were analysed using NMRViewJ (Johnson & Blevins, 1994).

**Ubiquitin–Vinyl sulphone reactions**

Individually purified proteins were acquired as described above. The final concentration of each component was 10 μM in a final volume of 45 μl [R0RBR and RING2(Rcat)] or 55 μl (parkin) in 50 mM HEPES and 50 mM NaCl (pH 8.0). Time started when the UbVS (Boston Biochem) was added to the reaction at 37°C. 10 μl was removed at each time point, and the reaction was quenched with 3× SDS sample buffer. 16.5% SDS–PAGE gels were run and stained with Coomassie Blue. Gels were imaged on a Bio-Rad ChemiDoc XR5–. Band intensities of parkin-Ub (R0RBR-Ub), parkin (R0RBR) and UbVS/pUb/pUbl were measured using ImageJ software (Schneider et al., 2012). The per cent contribution per band was calculated from the normalized intensity in each lane.

**Ubiquitination assays**

These assays were monitored by fluorescence using Ub containing an N-terminal cysteine residue linked to DyLight 800 Maleimide (Ub800; Thermo Fisher Scientific) as previously described (Kumar et al., 2015; Aguirre et al., 2018). All reactions were performed at 37°C and contained purified 1 μM wild type or substituted parkin, 0.5 μM UbcH7, 0.1 μM Uba1, 4 μM Ub and 0.5 μM Ub800 in 5 mM MgATP, 50 mM HEPES (pH 7.5). pUb was added to 0.5 μM when needed. To induce in situ phosphorylation, 0.01 μM of purified GST-PINK1 was added to the parkin/pUb/ATP samples 30 min before initiating ubiquitination. The ubiquitination reactions were quenched with 3× SDS sample buffer and 1 M DTT. 4–12% Bis-Tris gradient gels (Thermo Fisher Scientific) were used with MES running buffer (250 mM MES, 250 mM Tris, 0.5% SDS and 5 mM EDTA, pH 7.3). Fluorescence intensity at 700 and 800 nm was measured using an Odyssey Imaging System (LI-COR).

HA-Miro1 ubiquitination reactions were performed at 30°C for 10 min in a buffer containing 50 mM Tris (pH 7.5), 100 mM NaCl, 5 mM MgCl2, 1 mM TCEP and 5% (v/v) glycerol. Reactions contained 25 nm E1, 250 nM UbcH7, 250 nM pParkin, 10 μM Ub (wild type or mutant), 5 μM HA-Miro1 and 5 mM ATP in final reaction volume of 10 μl. Non-activatable pUb-6His (2.5 μM) was used as an allosteric activator where indicated. Reactions were terminated using NuPAGE LDS Sample Buffer (Invitrogen), resolved on 4–12% Bis-Tris gradient gels (Thermo Fisher Scientific) and transferred onto nitrocellulose membranes using iBlot Gel Transfer Device (Invitrogen). Membranes were subjected to immunoblotting using anti-HA mouse monoclonal primary antibody (901515, Bio Legend, 1/5,000 dilution) and fluorescent-labelled secondary antibody (926-32213, Li-COR, 1/10,000 dilution). Blots were visualized using Li-COR Odyssey Infrared Imaging System.

**Parkin ubiquitin loading**

Reactions monitoring pParkin-ubiquitin oxyester formation were performed at 30°C for 60 min in a buffer containing 50 mM Tris (pH 7.5), 100 mM NaCl, 5 mM MgCl2, 1 mM TCEP and 5% (v/v) glycerol. Reactions contained 100 nM E1, 2.5 μM UbcH7, 2.5 μM pParkin C431S+H433A, 10 μM Ub (wild type or mutant) and 5 mM ATP in final reaction volume of 10 μl. Non-activatable pUb-6His (2.5 μM) was included as an allosteric activator where indicated. Reactions were stopped using NuPAGE LDS Sample Buffer (Invitrogen) that contained reducing agents and boiled for 5 min. The samples were resolved on 4–12% Bis-Tris gradient gels (Thermo Fisher Scientific) and analysed by Coomassie staining.
Hydrogen–deuterium exchange mass spectrometry

Deuteration of proteins occurred at 20 ± 1°C in 90% D2O and 10% H2O with 50 mM HEPES, 100 mM NaCl and 250 μM TCEP at pH 7.0. Parkin and pParkin:pUb sample concentrations were 1 μM. UbcH7-Ub conjugate and pParkin:pUb concentrations were both 10 μM and were diluted to 1 μM in the above H2O buffer at pH 2.3. 100-μl aliquots were removed at time points between 15 s and 10 min following deuteration. Aliquots were quenched with ice-chilled 10% HCl in H2O to reach a pH of 2.3 and then flash-frozen in liquid nitrogen. Zero time point (m0) controls were created by adding ice-chilled D2O to an ice-chilled protein sample under quench conditions (pH 2.3) and flash-frozen. Fully exchanged controls (m100) were also created by exposing the proteins to D2O at pH 2.3 and heated to 70°C with a water bath for 8 h. These samples were then flash-frozen in liquid nitrogen. Aliquots were thawed to approximately 0°C and injected into a Waters HDX nanoACQUITY HPLC system. Online digestion of the proteins was performed with a POROS pepsin column at 15°C. The resulting peptides were trapped and analysed on a Waters BEH C18 column at 0°C using a water/acetonitrile with 0.1% formic acid gradient at 40 μl/min. The peptide masses were measured using a Waters Synapt G2 Q-TOF mass spectrometer. Peptides were identified through MS/MS. The resulting peptides were analysed with Waters DynamX 3.0. Deuteration is expressed here as per cent deuteration uptake where m is the centroid mass at time t and m0 and m100 are described above according to the following equation:

\[
\% \text{ D uptake} = \frac{m_t - m_0}{m_{100} - m_0} \times 100
\]

Model determinations for UbcH7-Ub binding to R0RBR:pUb

Interacting residues were identified from NMR experiments (described above) and defined as those amides that shifted greater than the average + one standard deviation and residues that broadened/shifted and could not be identified in the bound state (Fig EV2 and Appendix Fig S3). These residues were filtered for those that had > 20% side chain accessible surface area in each starting set of coordinates (below). Passive residues were defined according to the HADDOCK protocol (Dominguez et al, 2003; Bonvin et al, 2018) as residues that neighboured active residues having > 20% side chain accessible surface area and had chemical shift changes greater than average. This approach led to a set of ambiguous restraints between the UbcH7 (K9, A59, E60, F63, K64, E93, N94, K96, A98) and R0RBR parkin (T242, L266, T270, Q276, A291, G292, Q389-D394, R396) and between Ub (F4, T7-G10, I11, E34, I36, V70, L71, L73, G75, G76) and R0RBR parkin (Q276, V330, L331, R366, A379-S384, G385, T386).

The UbcH7-Ub conjugate was docked to R0RBR:pUb using HADDOCK (Dominguez et al, 2003) using the residues described above. Starting coordinates from the crystal structure of pUb:UbIRORBR (PDB code 5N2W) were used following removal of the UbI domain and adjoining linker coordinates (residues 1–83). Several linker sections absent in crystal structures of parkin were incorporated using the Modeller (Eswar et al, 2006) plug-in for UCSF Chimera (Pettersen et al, 2004). The tether (387–405) that partly occludes the RING1 binding site in the starting coordinates was allowed to move. Coordinates for ubiquitin (PDB code 1UBQ; Vijay-Kumar et al, 1987) and UbcH7 (PDB code 4Q5E; Grishin et al, 2014) were used to sequentially dock the UbcH7 and Ub moieties untethered according to restraints and using a single unambiguous restraint between the C-terminal G76 of ubiquitin and the catalytic C86K of UbcH7 to create the UbcH7-Ub conjugate in the complex. Upper distance limits of 4.0 Å were set for ambiguous distance restraints, while the unambiguous distance restraint was set to 6.8 Å. Standard HADDOCK parameters were used except inter_rigid (0.1) which was set to allow tighter packing of the proteins, and the unambiguous force constant was set fivefold higher compared to those used the ambiguous constants. A total of 1,000 initial complexes were calculated, and the best 100 structures were water-refined.

Data availability

Coordinates for models of the UbcH7-Ub conjugate bound to R0RBR:pUb reported here have been deposited to the Protein Data Bank under accession number 6N13.

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Author contributions

TECC collected and assigned all NMR experiments, completed and analysed HADDOCK modelling for the E2-Ub complex with pParkin:pUb, and wrote the manuscript. KMD conducted ubiquitination assays and NMR-based interaction experiments, completed UbVS reactivity experiments, and analysed and wrote the manuscript. EAF did HDX experiments and NMR interaction experiments, analysed the data and wrote the manuscript. KRB expressed and purified proteins for ubiquitination assays, NMR experiments and HDX experiments and wrote the manuscript. JDA helped with NMR-based interaction experiments. VKC completed ubiquitin loading and off-loading experiments and wrote the manuscript. YX and LK helped with the design and analysis of HDX experiments. HW and GSS conceived the study, designed experiments, analysed the data and wrote the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

References


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