Dual Function of Phosphoubiquitin in E3 Activation of Parkin*

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Mutations in the gene encoding parkin, an auto-inhibited E3 ubiquitin ligase that functions in the clearance of damaged mitochondria, are the most common cause of autosomal recessive juvenile Parkinsonism. The mechanism regulating parkin activation remains poorly understood. Here we show, by using isothermal titration calorimetry, solution NMR, and fluorescence spectroscopy, that parkin can bind ubiquitin and phosphomimetic ubiquitin by recognizing the canonical hydrophobic patch and C terminus of ubiquitin. The affinity of parkin for both phosphomimetic and unmodified ubiquitin is markedly enhanced upon removal of the ubiquitin-like (UBL) domain of parkin. This suggests that the agonistic binding of ubiquitin to parkin in trans is counterbalanced by the antagonistic activity of the parkin UBL domain in cis. Intriguingly, UBL binding is enthalpy-driven, whereas ubiquitin binding is driven by an increase in the total entropy of the system. These thermodynamic differences are explained by different chemistry in the ubiquitin- and UBL-binding pockets of parkin and, as shown by molecular dynamics simulations, are not a consequence of changes in protein conformational entropy. Indeed, comparison of conformational fluctuations reveals that the RING1-IBR element becomes considerably more rigid upon complex formation. A model of parkin activation is proposed in which E2~Ub binding triggers large scale diffusional motion of the RING2 domain toward the ubiquitin-stabilized RING1-IBR assembly to complete formation of the active parkin-E2~Ub transfer complex. Thus, ubiquitin plays a dual role in parkin activation by competing with the inhibitory UBL domain and stabilizing the active form of parkin.

Mutations in the PARK2 gene that encodes the protein parkin have been found to cause a familial form of Parkinson disease termed autosomal recessive juvenile parkinsonism (AR-JP) (1–3). Parkin is an E3 ubiquitin ligase of the RBR family (Fig. 1A). RBR-type ubiquitin ligases function by first forming a covalent thioester intermediate between a key catalytic cysteine and the C terminus of ubiquitin, and then transferring the activated ubiquitin to their substrate (4). Under basal conditions, parkin resides in the cytosol (5, 6), and cytosolic parkin is thought to be catalytically inactive (5). Accordingly, recently reported crystal structures show parkin in an auto-inhibited conformation; both the interaction platform with the incoming E2 enzyme and the catalytic cysteine residue Cys431 are occluded because of intramolecular interactions (Fig. 1B) (7–9). Autoregulation of E3 activity by the formation of such inhibitory interactions is a common feature of RBR ligases and has been previously described for the ubiquitin ligases HHARI and HOIP (4, 10, 11).

Parkin functions in the ubiquitylation of mitochondrial proteins, an event that initiates the degradation of dysfunctional mitochondria by mitophagy (5, 6); this implies that the latent E3 ubiquitin ligase activity of parkin can be released in response to external factors. At least two events are required to release the latent E3 activity of parkin in cells. It has been shown that upon loss of the electrochemical potential across the inner mitochondrial membrane, the protein kinase PINK1, which is anchored on the outer mitochondrial membrane with the kinase domain facing the cytoplasm, phosphorylates parkin at residue Ser65 of its UBL domain (12–14). Furthermore, it has been reported that PINK1 also phosphorylates ubiquitin, intriguingly at the homologous residue Ser65, which is conserved between ubiquitin and the UBL domain of parkin (15–17).

How these two phosphorylation events act together to convert parkin from an inactive to a catalytically active state is not fully understood. Alone, the two events are not sufficient to induce translocation of parkin to mitochondria, but they are required for its translocation and are sufficient to release the latent E3 ubiquitin ligase activity of parkin in vivo (15, 16).

A leading hypothesis for parkin activation is that Ser65-phosphorylated ubiquitin (hereafter referred to as "phosphoubiquitin") may activate parkin in an allosteric manner (15, 18, 19). Indeed, it has been demonstrated in vitro that the presence of phosphoubiquitin in a ubiquitylation assay solution is sufficient to activate parkin without the need for phosphorylation of the parkin UBL domain (18).

The aim of this study was therefore to investigate whether ubiquitin can exert a direct influence on the multidomain structure of parkin, which ultimately leads to release of its latent ubiquitin ligase activity.

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‡ The abbreviations used are: AR-JP, autosomal recessive juvenile parkinsonism; HSQC, heteronuclear single quantum coherence; HOIP, HOIL-1L interacting protein; IBR, in-between RING; ITCl, isothermal titration calorimetry; MD, molecular dynamics; RING, really interesting new gene; r.m.s.d., root mean square deviation; RMSF, root mean square fluctuation; RBR, RING-in-between-RING; UBL, ubiquitin-like; Ub, ubiquitin; PDB, Protein Data Bank; REP, repressor element.
Results

Ubiquitin Binding to Parkin—Direct allosteric activation of parkin by ubiquitin would require a physical interaction of the two molecules in solution. Indeed, we detected such a non-covalent association by NMR spectroscopy. Addition of an equimolar amount of unlabeled parkin to 15N-labeled ubiquitin markedly changed the HSQC spectrum of ubiquitin (Fig. 2A). Several resonances of ubiquitin, such as Ile36, shifted in a fast exchange-like manner upon the addition of parkin, while displaying a small degree of line broadening. Conversely, other resonances, such as Val70, were severely broadened and were no longer detectable, suggesting that ubiquitin binds to parkin by a specific interaction. Such non-covalent binding of ubiquitin to an E3 ligase is not typical because catalytic processing of ubiquitin exclusively involves its covalent transfer via cysteine residues (20). Thus, we inferred that binding of ubiquitin may have a specific role in parkin function. The complete loss of Leu8 and Val70 signals suggested that parkin binds to the canonical hydrophobic patch on ubiquitin (Fig. 2B) (20). Moreover, loss of the amide cross-peaks of Arg72, Leu73, and Arg74 of ubiquitin indicated that parkin also interacts with the flexible C terminus of ubiquitin in solution.

Direct determination of the ubiquitin-binding site on parkin by NMR spectroscopy was obstructed by inefficient folding of parkin in isotope-labeled minimal media. While this study was in progress, a crystal structure of phosphoubiquitin bound to the parkin core was reported (19). Superimposing ubiquitin on that crystal structure brought residues showing severe line-broadening in our NMR experiments close to the surface of parkin, thereby explaining the line-broadening as interaction of ubiquitin’s hydrophobic patch and C terminus with residues on the surface of parkin (Fig. 2C).

Accordingly, these data argue that parkin and ubiquitin specifically interact in solution. Notably, ubiquitin can interact with parkin even without phosphorylation of Ser65. Because the intracellular concentration of ubiquitin is on the order of ~85 μM (21), which is much higher than that of other proteins, including parkin and phosphoubiquitin (15), our data suggest that cytosolic (inactive) parkin may be often bound to unmodified ubiquitin in vivo. Nevertheless, it has been reported that only phosphoubiquitin or phosphomimetic (S65D) ubiquitin activates parkin in cells (15). On the basis of these considerations, we hypothesized that, although unmodified ubiquitin can bind to parkin, binding of phosphoubiquitin might exert a specific effect that facilitates the release of latent E3 activity. The nature of such an effect is not evident from the currently available static structural data alone because, in the recent crystal structure of phosphoubiquitin-bound parkin, the catalytic cysteine residue is still buried inside the protein and thereby unable to participate in the ubiquitylation cascade (Fig. 1, C and D) (19). Thus, we assumed that phosphoubiquitin binding to parkin must play a more subtle role in the activation of parkin.

Thermodynamic Evaluation of Ubiquitin Binding to Full-length Parkin—To evaluate the binding of ubiquitin and parkin in solution in more detail, we employed isothermal titration calorimetry (ITC), which confirmed the binding of ubiquitin to parkin. Intriguingly, addition of ubiquitin to parkin was found to be an endothermic reaction with a dissociation constant of ~70 μM (Fig. 3A). Because it is not ubiquitin per se, but rather phosphoubiquitin that is thought to activate parkin, we pre-
FIGURE 2. Physical interaction of parkin with ubiquitin. A, transverse relaxation-optimized spectroscopy-HSQC spectrum of 100 μM 15N-labeled ubiquitin before (black) and after (red) addition of 1 mol eq of unlabeled full-length parkin, showing selective line broadening after the addition of parkin. B, residue-specific NMR signal loss in ubiquitin upon the addition of parkin. Green bars, residues not observed in the reference spectrum of free ubiquitin; red bars, residues for which near-complete signal loss occurred upon the addition of parkin. Red line, average intensity ratio. C, structural model of ubiquitin binding to parkin based on PDB code SCAW (19). Residues (one-letter code) with significant line broadening in the NMR experiment are colored red.

FIGURE 3. Ubiquitin binding to parkin is entropy-driven. A and B, isothermal titration calorimetry thermogram for ubiquitin (A) and phosphomimetic ubiquitin (B) binding to full-length parkin, showing an endothermic reaction. Upper panels show raw data; lower panels show the integrated heat values. C, dissociation constants determined from the ITC experiments. Unmodified ubiquitin and phosphomimetic ubiquitin bind to full-length parkin with similar avidity (p = 0.16; Student’s t test). D, entropy drives the binding of both ubiquitin and phosphomimetic ubiquitin to parkin. The binding process is disfavored enthalpically, but this is counteracted by a large favorable entropic term. Error bars indicate the S.E. in C and D of three independent experiments (n = 3).
dicted that the affinity of phosphoubiquitin for parkin would be
stronger than that of unmodified ubiquitin.

To examine this possibility, we employed the S65D ubiquitin
mutant, which has been successfully used as a mimic for phos-
phomimetic ubiquitin (15, 16, 22). ITC experiments with phos-
phomimetic ubiquitin resulted in a dissociation constant of
\(K_d \approx 35 \mu M\); however, statistical analysis precluded the conclusion
that phosphomimetic ubiquitin bound to full-length parkin
more avidly than wild-type ubiquitin (Fig. 3, B and C). Interest-
ingly, the entropic contribution to the binding reaction was
enhanced in the case of phosphomimetic ubiquitin. These
results suggest that parkin has a specific, entropy-driven bind-
ing mode for phosphorylated ubiquitin.

**Ubiquitin Binding to the Parkin Core**—Direct interpretation
of the thermodynamic driving force behind the binding is com-
plex, because changes in both solvent and protein conforma-
tional entropy must be taken into account. Moreover, the pres-
ence of the UBL domain and the ~70-residue linker between
the UBL and the parkin core (Fig. 1A) further complicated the
interpretation. Therefore, to analyze the effect of ubiquitin
binding on the catalytic core of parkin specifically, we deleted
the N-terminal UBL domain and its adjacent linker in parkin
and measured binding of the parkin core (hereafter parkin
UBL) to ubiquitin using ITC.

Parkin UBL showed increased affinity for unmodified ubiq-
uitin (Fig. 4A; \(K_d \approx 18 \mu M\)) as compared with full-length parkin
(Fig. 3A; \(K_d \approx 70 \mu M\)). Notably, deletion of the UBL domain did
not change the endothermic nature of the reaction, indicating
that the entropic driving force behind the reaction is indepen-
dent of the UBL domain and the adjacent linker.

Because parkin bound unmodified and phosphomimetic
ubiquitin with comparable affinity, we next determined
whether this behavior would be preserved in the absence of
the UBL domain of parkin. ITC experiments showed that phospho-
mimetic ubiquitin bound to parkin UBL with a \(K_d\) of ~1.5 \(\mu M\)
(Fig. 4, B and D). Thus, parkin UBL showed a 10-fold prefer-
ence for phosphomimetic ubiquitin over unmodified ubiquitin. Again, the reaction was entropy-driven, and the entropic contribution was larger for phosphomimetic ubiquitin than for unmodified ubiquitin (Fig. 4E).

Because binding of (phospho-) ubiquitin to the parkin core is entropy-driven, we considered the role of the UBL domain in full-length parkin from a thermodynamic point of view. Using ITC, we found that the parkin UBL domain bound parkinΔUBL with a $K_d$ of $16 \mu M$ (Fig. 4C). Accordingly, parkinΔUBL showed essentially the same affinity toward unmodified ubiquitin and its UBL domain in trans but bound phosphomimetic ubiquitin 10-fold more tightly (Fig. 4E). Unexpectedly, we found that binding of the UBL domain to parkinΔUBL is an exothermic reaction. Thus, although ubiquitin and the parkin UBL bind to parkinΔUBL with essentially the same affinity, the driving force behind the reaction is fundamentally reversed. Taken together, these observations imply that binding of phospho-ubiquitin to parkin induces subtle structural changes in the core domains of parkin (Fig. 5E).

**Changes in the Conformational Dynamics of Ubiquitin upon Complex Formation**—The comparably small red shift of the fluorescence emission spectra upon complex formation indicated that (phospho-) ubiquitin binding-induced conformational changes in parkin must be rather modest. Indeed, the static structural differences between free parkin (7–9) and its complex form with phospho-ubiquitin (19) seem to be limited. We thus considered the possibility that ubiquitin affects the structural dynamics, rather than the static structure of parkin, to modulate parkin function. To obtain detailed insight into the conformational dynamics of the free and phospho-ubiquitin-bound parkin core (parkinΔUBL), we performed molecular dynamics (MD) simulations with parkinΔUBL, (phospho-) ubiquitin, and the parkin-phospho-ubiquitin complex in explicit water.

First, we obtained MD trajectories for ubiquitin and phospho-ubiquitin. To enable a statistical analysis of the output of our simulations, we repeated each calculation 10 times, starting from the same structure, but with different randomized initial velocities. In the case of phospho-ubiquitin, two distinct conformations have been reported; the structure of the major state of phospho-ubiquitin in solution was used for our simulations because this is the conformation of phospho-ubiquitin that binds to parkin (18, 19).
Interaction of Parkin with Ubiquitin

On the nanosecond time scale, ubiquitin and phosphoubiquitin exhibited similarly rigid overall motion with residual flexibility in the loop regions (residues 7–10 and 46–48), as well as the well-known high flexibility in the C-terminal tail (Fig. 6, A and B). Second, we obtained trajectories of the parkin ΔUBL-phosphoubiquitin complex. In the complex, the backbone fluctuations of phosphoubiquitin were essentially identical to that of the free form in the regions of secondary structure. However, the backbone motion of phosphoubiquitin was even more suppressed in the loop regions (residues 7–10 and 46–48) and the C-terminal tail (Fig. 6, A and B, red trace). This suggests that phosphoubiquitin gives up some of its residual dynamic character upon formation of a complex with parkin.

Conformational Dynamics of the Parkin Core—Next, we obtained MD trajectories of the free parkin core (parkin ΔUBL). The core domains of parkin adopted a stable conformation with a total Cα r.m.s.d. value of ~4–5 Å (Fig. 7A). Among these domains, the IBR domain showed the highest RMSF values, indicating the high mobility of this domain with respect to the other domains of parkin. RING1 was rigid, with average RMSF values ranging from ~1.5 to 2 Å (Fig. 7, B and C). This result is reasonable considering that RING1 engages in non-covalent interactions with all other domains of parkin (Fig. 1B). Interestingly, the catalytic cysteine residue Cys431 was observed at a minimum of backbone flexibility (Fig. 7, B and C). Likewise, the key auto-inhibitory residue Trp403, alanine mutation of which key-activated parkin (7), was observed in another local minimum of backbone flexibility (Fig. 7B). Thus, auto-inhibition of parkin works not only by the static auto-inhibitory interactions, as described by recent x-ray crystallographic reports (7–9), but also by the intrinsically rigid motion of the protein backbone among residues flanking key auto-regulatory sites.

Changes in the Conformational Dynamics of Parkin upon Complex Formation—Many residues of parkin showed differences in conformational dynamics between the free and phosphoubiquitin-bound forms, as indicated by the respective RMSF values (Fig. 7B). Interestingly, phosphoubiquitin binding to RING1 changed the backbone fluctuations in parkin, and this effect was not confined to the immediate vicinity of the binding site. Rather, the global conformational dynamics in the whole parkin molecule were redistributed after complex formation (Fig. 7, B and D).

Phosphoubiquitin binding had no significant effect on the RING2 domain, which showed very similar dynamics between the free and phosphoubiquitin-bound form (Fig. 7, B and D). This is reasonable, considering that the RING2 domain is located away from the ubiquitin-binding site. The conformational fluctuations in all other domains of parkin were significantly affected by phosphoubiquitin binding. The RING1 domain engaged in non-covalent interactions with phosphoubiquitin, rendering it even more rigid as indicated by average RMSF values of ~1 Å (Fig. 7, B and D). Several residues in the RING0 domain also showed increased rigidity, whereas regions more remote to the ubiquitin-binding site seemed to be less affected (Fig. 7, B and D). Importantly, the highly dynamic IBR domain showed increased rigidity after complex formation.

Taken together, these data suggest that the conformational fluctuations in the parkin core are redistributed by phosphoubiquitin binding leading to significantly enhanced rigidity in the RING1-IBR element, while exerting no apparent influence on RING2 and large parts of the RING0 domain.

Discussion

Thermodynamic Change upon Ubiquitin Binding—We have shown that both ubiquitin and phosphomimetic ubiquitin can bind to full-length parkin and the isolated parkin core (parkin ΔUBL) in solution (Figs. 2–4). Binding of (phosphomimetic) ubiquitin to parkin is enthalpically disfavored, but it is driven by an increase in the total entropy of the system (Figs. 3 and 4). Because the total entropy change of binding ($-T\Delta S_{\text{total}}$) can be dominated by changes in protein conformational entropy ($-T\Delta S_{\text{conf}}$) (23–28), we considered changes in protein dynam-
ics upon complex formation. However, MD analysis indicated that both phosphoubiquitin and the parkin core become less dynamic after complex formation. Accordingly, the entropy-driven nature of ubiquitin binding to parkin must be explained by increases in solvent entropy.

The entropic nature of ubiquitin binding to parkin can be explained by the hydrophobic effect; the phosphoubiquitin-binding site on the RING1 domain of parkin contains several hydrophobic residues whose exposed side chains engage in interactions with phosphoubiquitin (Ala152, Val226, Val303, Ala321, Phe325, Ile326, Ile343, Leu344, and Ile345 in PDB code 5CAW in Fig. 8A) (19). In the absence of phosphoubiquitin, these hydrophobic side chains cannot participate in hydrogen bonding with the solvent, leading to the formation of highly ordered water molecules caging the binding site. Phosphoubiquitin binding displaces many of these highly ordered water molecules, resulting in less ordered water (higher solvent entropy), which drives the reaction.

Conversely, only two hydrophobic side chains (Leu266 and Val269 in PDB code 4K95, see Fig. 8B (7)) are exposed to the solvent in the case of the UBL-binding site. Thus, the solvent at this site is unlikely to form highly ordered water clusters, and UBL binding does not result in a significant entropy gain. These considerations are in good agreement with our observation that binding of the UBL domain to parkin is driven by enthalpy (Fig. 4E).

**Effect of AR-JP-associated Mutations on Parkin**—Pathogenic mutations can affect parkin by a variety of mechanisms (Fig. 9). Some of the mutations appear to perturb the stability of structural elements of parkin or directly alter the enzymatic active site. Importantly, two pathogenic mutations L283P and G284R are located in a phosphoubiquitin-binding element of the RING1 domain, and the AR-JP mutation A240R interferes with E2 binding (7, 19). Other Parkinson disease-associated mutations are located at domain-domain interfaces. This highlights that both intra- and intermolecular interactions have to be mediated by the hydrophobic effect.

**FIGURE 7.** Conformational fluctuations in the parkin core before and after binding to phosphoubiquitin. A, time traces of the r.m.s.d. calculated for Cα atoms after rotational and translational fitting to the Cα coordinates of the first time frame during molecular dynamics simulations of parkin ΔUBL (black) and the parkin ΔUBL-phosphoubiquitin complex (red). All simulations were performed 10 times with different randomized initial velocities. Average r.m.s.d. values are shown. The shaded area represents the S.E. of the respective r.m.s.d. B, RMSF of Cα atoms of parkin ΔUBL (black) and the parkin ΔUBL-phosphoubiquitin complex (red) averaged over the time interval (5 ns; 50 ns). The average RMSF value of 10 simulations, each of 50 ns, is shown. Error bars indicate the S.E. Residue numbers refer to parkin from R. norvegicus and the alignment was performed by Clustal Omega (53). The respective domains are indicated. C, conformational fluctuations in the free parkin core (B) as visualized on the structure of the first time frame. Residues are color-coded as follows: red, RMSF > RMSFaverage; blue, RMSF < RMSFaverage − 1σ; gray, all other residues. Magenta sphere, catalytic cysteine. D, changes in conformational fluctuations in the parkin core upon binding to phosphoubiquitin. The ratio of the RMSF values of B is visualized on the structure of free parkin. The phosphoubiquitin-binding site is indicated. Color code: blue, ρi > Paverage + 1σ; light blue, ρi > Paverage; gray, all other residues; ρi, RMSF (B) ratio for residue i of the free to the phosphoubiquitin-bound form of the parkin core. Magenta sphere, catalytic cysteine.
Intriguingly, the role of a large number of pathogenic mutations remains largely unexplained (Fig. 9, gray). Because auto-inhibition of parkin activity is critical for proper regulation of its activity, this group of point mutations may exert a subtle influence on the conformational dynamics of the autoinhibited enzyme (Fig. 7), resulting in a slight imbalance of its autoregulatory mechanism. However, a detailed thermodynamic and kinetic analysis will be required to elucidate the exact impact of these point mutations on parkin activity.

Role of PINK1 in Parkin Activation—On the basis of our data and recently published reports, it is possible to construct a model of parkin activation. First, the mitochondrial protein kinase PINK1 phosphorylates ubiquitin at residue Ser65 (12–14). Although unmodified ubiquitin can bind to parkin (Figs. 2–5), phosphorylation by PINK1 increases the affinity of ubiquitin for the parkin core (Fig. 4D). Importantly, the absence of the parkin UBL domain enhances ubiquitin binding to parkin (Figs. 3D and 4D), suggesting that ubiquitin and the parkin UBL domain compete for binding to parkin, even though they bind at distinct sites (19, 29). Phosphoubiquitin binding displaces (19, 29) the auto-inhibitory UBL domain from the parkin core, leading to solvent exposure of Ser65 of the UBL domain, which is buried in the structure of free parkin.

This exposure renders Ser65 accessible to PINK1, which then phosphorylates the UBL domain of parkin. This phosphorylation event prevents the UBL from rebinding back onto the parkin core (19, 29). Thus, PINK1 plays a dual regulatory role in this pathway by (i) enhancing the affinity of ubiquitin for the parkin core (Fig. 4D) and (ii) decreasing the affinity of the UBL domain for the parkin core. Here, PINK1 intriguingly uses the same mechanism, phosphorylation of Ser65, but on different molecules (ubiquitin and the UBL domain of parkin) with opposite effects (respectively, enhanced and decreased affinity for the parkin core) to shift the UBL/ubiquitin competition equilibrium toward a phosphoubiquitin-bound and UBL-released state of parkin, comprising an elegant positive feedback mechanism (Fig. 10A).

Structural Rearrangements in the Parkin Core—We observed structural changes in the core RING domains of parkin by monitoring the fluorescence red shift that accompanies the solvent exposure of tryptophan residues upon complex formation. The absolute magnitude of red shift is comparably small, indicating that the tryptophan residues do not become completely exposed to the solvent as would be the case in protein unfolding ($\lambda_{max} \approx 355$ nm). Rather, the observed red shift may suggest that, upon phosphoubiquitin binding, subtle rearrangements in the conformational ensemble of parkin take place in solution.

Conformational rearrangements on a much larger scale are required to convert the conformation of auto-inhibited parkin into a fully activated form, because superposition of the E2~Ub conjugate onto the E2-binding site in any of the known crystal structures of parkin results in models in which the active sites of the E2 and E3 enzymes are placed ~50 Å apart from each other (7–9, 19). Thus, currently available crystal structures of parkin cannot explain the transthiolation reaction.

Role of Ubiquitin in Parkin Activation—Phosphoubiquitin binding to the parkin core results in modest local changes to the static structure of the parkin core such as the straightening of an $\alpha$-helix in the RING1 domain, although most other regions seem to be rather unaffected (Fig. 10A, right) (19). By contrast, our MD analysis indicated that the structural rigidity of parkin is increased in the phosphoubiquitin-bound state, and importantly, this effect is most pronounced in the RING1-IBR element (Fig. 7, B and D).

Comparison of parkin to a recently published structure of an active RBR/E2~Ub transfer complex of HOIP suggests a possible mechanism for the conversion of parkin to an activated state. In that structure (31), the RING1-IBR element makes non-covalent contacts with an allosteric ubiquitin molecule, which according to the authors was not expected to be found in the electron density, but it proved absolutely necessary for crystal formation (Fig. 10B). Thus, we infer that ubiquitin exerts a stabilizing effect on the RING1-IBR element in the crystal of the HOIP RBR/E2~Ub, in analogy to the stabilizing effect that phosphoubiquitin exerts on the RING1-IBR element in parkin (Fig. 7D).

Fixation of RING1 and the dynamic (Fig. 7C) IBR domain relative to one other (Fig. 7D) would permit the RING2 domain
to undergo diffusion with respect to the phosphoubiquitin/RING1-IBR arrangement, facilitated by the linker between the IBR and RING2 domains (Fig. 10C). Importantly, the final conformation of the RING2 domain is unlikely to show marked structural changes (Fig. 10D), suggesting that simple rotational and translational diffusion of the RING2 domain toward the phosphoubiquitin-fixed RING1-IBR element is sufficient to bring the catalytic cysteine within close proximity of the C-terminal glycine residue (Gly76) of the activated ubiquitin, thereby yielding an active state of parkin.

**Conclusion—AR-JP mutations may unbalance the conformational dynamics of parkin by a variety of mechanisms.** Pathogenic mutations linked to autosomal recessive juvenile parkinsonism (8, 30, 54–62) are mapped onto the crystal structure of parkin (spheres). Several mutants appear to affect the structure of parkin by interfering with Zn$^{2+}$ binding (blue) or the correct formation of secondary structure elements (magenta). The mutants G430D and C431F directly affect the active site (yellow). Other PD mutants may alter domain-domain interactions (green). Ala240 (orange) is critical for binding to the E2, and Leu283 and Gly284 (red) participate in phosphoubiquitin binding (19). Many of the remaining PD mutants are less well understood (gray) but may modulate the overall conformational dynamics of the enzyme. The structure of parkin was drawn from PDB code 4K95 (7).

**Experimental Procedures**

**Proteins—**Ubiquitin was expressed and purified as described previously (33). Parkin from *Rattus norvegicus* was expressed from a pGEX-6P1 expression vector that was obtained from Addgene with kind permission from Dr. Kalle Gehring (7). Full-length parkin (residues 1–465) and its fragment lacking the N-terminal UBL domain (parkin ∆UBL; residues 141–465) were expressed and purified as reported previously (7). In brief, BL21(DE3) *Escherichia coli* cells harboring the parkin expression vector were grown to an $A_{600}$ of 0.5 at 37 °C. The temperature was decreased to 16 °C, and expression was induced by the addition of 25 μM isopropyl 1-thio-β-D-galactopyranoside overnight. GST-parkin was purified by glutathione-Sepharose 4FF (GE Healthcare) column chromatography. After cleavage of the GST tag by PreScission protease, parkin was further purified by size-exclusion chromatography on a HiLoad 16/60 Superdex 75pg column (GE Healthcare). The parkin UBL domain (residues 1–76) was expressed and purified as described previously (30).

**NMR Spectroscopy—**NMR spectra were acquired on an Avance II 700 MHz spectrometer equipped with a 5-mm 1H/13C/15N z-gradient triple resonance cryoprobe (Bruker). The free induction decays were apodized, zero-filled, and Fourier-transformed using NMRPipe (34). Resolution in the indirect dimension was improved by linear prediction. Spectra were analyzed in CCPN (35). Ubiquitin was measured at a concentration of 100 μM at 310 K, and its heteronuclear single quantum coherence (HSQC) spectrum was acquired as transverse relaxation-optimized spectroscopy-type experiment. Figures of spectra were created by the Python packages Nmrglue and Matplotlib (36, 37).

**Isothermal Titration Calorimetry—**Isothermal titration calorimetry (ITC) experiments were conducted at 298 K on an ITC200 system (MicroCal). All protein samples were dialyzed against ITC buffer (20 mM HEPES, pH 7.5, 100 mM sodium chloride, 0.1 mM tris(2-carboxyethyl)phosphate) overnight and thoroughly degassed before each experiment. Samples were incubated at room temperature (298 K) before each experi-
FIGURE 10. Model of parkin activation by phosphoubiquitin. A, left, in the initial state, parkin adopts an auto-inhibited conformation with the UBL domain bound to the parkin core (7). PINK1 phosphorylates Ser65 of ubiquitin, which competes with the UBL for binding to the parkin core. Release (19, 29) of the UBL domain from the parkin core to the solvent enables PINK1 to phosphorylate Ser65 of the UBL, which prevents the UBL from rebinding back on the parkin core. Accordingly, the action of PINK1 drives the UBL/phosphoubiquitin competition equilibrium toward a phosphoubiquitin-bound/UBL-exposed state of parkin (right, schematic).

B, structure of an active RBR/E2/H11011 Ub transfer complex of the RBR-type E3 ubiquitin ligase HOIP (31). The linear ubiquitin chain-determining domain is omitted for clarity. An allosteric ubiquitin molecule (Uballo) orients the IBR and RING1 domains with respect to one other. C, as compared with the inactive conformation of parkin (A), the RING2 domain in active HOIP (B) is rotated and translated, bringing it in close proximity to the RING1-IBR element. Such a large scale structural change is permitted by the long linker between the IBR and RING2 domains, but it requires release of the REP element from RING1. This displacement of the REP element is probably driven by competitive binding of E2—Ub to the E2-binding site on RING1 (blue arrow). D, comparison of the inactive and active states of the RING2 domain of parkin. Left, in the inactive conformation, the catalytic cysteine is occluded by the RING0 domain. Right, structural model of the RING2 domain of parkin in the active state as generated by SWISS-MODEL (63) on the basis of the active state of HOIP (B). The sequence identity of the RING2 domain between HOIP and parkin, as judged by SWISS-MODEL, is 43%. Apart from formation of a new α-helix, the overall structure of the RING2 domain is not markedly changed, indicating that simple rotation and translation of the RING2 domain with respect to the RING1-IBR element is sufficient to bring the catalytic cysteine within close proximity of the activated Gly76 of the incoming ubiquitin (Ubact), leading to formation of an active parkin-E2—Ub transfer complex. Domains other than RING0 and RING2 are omitted for clarity. Structures are drawn from PDB codes 4K95, 5CAW, and 5EDV, respectively.
ment. The concentration of the syringe protein (ubiquitin, phosphomimetic ubiquitin, or parkin UBL) ranged from 0.8 to 1.5 mM, whereas the cell contained protein (parkin or parkin \( \Delta \)UBL) at 87–100 \( \mu \)M. The syringe protein was injected into the cell at 3-min intervals. The resulting data were processed using Origin 7 (MicroCal Software, Inc.). Errors in the derived data represent the standard error of the mean of three independent experiments.

Molecular Dynamics—All molecular dynamics simulations were performed by GROMACS 5.0.6 (38–43) in explicit solvent using the AMBER99SB-ILDN force field (43) and the following crystal structures: \textit{Pediculus humanus} parkin \( \Delta \)UBL (residues 143–461; PDB code 5CAW (19)) in complex with phosphoubiquitin (residues 1–75); \textit{R. norvegicus} parkin \( \Delta \)UBL (residues 140–465; PDB code 4K7D (7)); human ubiquitin (residues 1–75; PDB code 1UBQ (44)); Ser\(^{65}\)-phosphorylated human ubiquitin (major conformation; residues 1–75; PDB code 4WZP (18)). Force field parameters for phosphoserine were obtained from Ref. 45. Residues missing in the crystal structures due to disorder were rebuilt using the ModLoop interface to Modeller (46). The respective protein was solvated in a triclinic box with SPC/E water. In structures involving phosphoubiquitin, the system charge was neutralized by addition of two \( \text{Na}^+ \) ions. \( \text{Zn}^{2+} \) ions of the RING domains of parkin were distance restrained to the respective cysteine S\(_\alpha\) atoms (47, 48). Energy minimization was performed by the steepest-descent algorithm with a maximum force \( F_{\text{max}} \) of 1000 kJ mol\(^{-1}\) nm\(^{-1}\). Equilibration of protein and solvent was conducted in two steps. First, the system was simulated under the canonical (NVT) ensemble for 100 ps with position restraints on all protein atoms. Second, 100 ps of position-restrained NPT simulation was performed to stabilize the density of the system. The Parrinello-Rahman algorithm (49) was applied for pressure coupling (1 bar, 2.0-ps coupling constant), and the temperature was coupled using the modified Berendsen (40) thermostat (310 K, 0.1 ps coupling constant). The integration step time was set to 2 fs. Bonds were constrained by the LINCS algorithm (42). The parameters for fast particle mesh Ewald electrostatics (50) were as follows: Lennard-Jones cutoff distance, 0.8 nm; Coulomb cutoff distance, 0.8 nm; Fast Fourier Transform grid spacing, 0.08 nm; interpolation order, cubic. For each simulation, 10 independent calculations were performed starting from the same structure with different initial velocities randomly assigned from the Maxwell-Boltzmann distribution at 310 K. All simulations were carried out on Linux workstations running Fedora 21 with CUDA-compatible NVIDIA graphics processing units. Analysis was done by routines integrated in GROMACS, and figures were created using the Python package Matplotlib (37). r.m.s.d. and RMSF values were calculated with reference to the structure of first time frame of the simulation by rotational and translational fitting of the \( C_{\alpha} \) coordinates. RMSF values were calculated for time frames in the range of 5–50 ns.

Fluorescence Spectroscopy—Fluorescence was measured using a FluoroMax4 fluorescence spectrometer (HORIBA). Tryptophan fluorescence was selectively excited at 300 nm, and emission spectra were acquired over wavelengths of 310–400 nm with the slit width set to 5 nm at 298 K. Ubiquitin has no tryptophan residues and contributes only baseline noise to the spectra. In parkin, tryptophan residues cluster in the catalytic center with the single exception of the linker residue Trp\(^{57}\), which is not resolved in any of the reported crystal structures.

Before the experiments, proteins were dialyzed into 20 mM HEPES, pH 7.5, 100 mM sodium chloride, 0.1 mM tris(2-carboxyethyl)phosphine. Parkin was diluted to a final protein concentration of 7 \( \mu \)M and measured at a volume of 500 \( \mu \)L. Each acquisition was repeated 10 times, and the averaged spectrum was subjected to analysis. The spectral contribution of the buffer was subtracted from the acquired spectra. The red shift of the fluorescence spectrum upon ligand addition was evaluated by calculating the baricentric mean of the spectrum. The baricentric mean was calculated as shown in Equation 1,

\[
\lambda_{\text{bcm}} = \frac{\sum F(\lambda) \lambda}{\sum F(\lambda)}
\]

(Eq. 1)

in which \( F(\lambda) \) is the tryptophan fluorescence emission intensity at \( \lambda \) nm, and the sum is over the entire spectrum (310–400 nm). Fluorescence red shifts are reported as differences from the initial baricentric mean: \( \Delta \lambda_{\text{bcm}} = \lambda_i - \lambda_{\theta} \), in which \( \lambda_i \) is the baricentric mean of the spectrum after the \( i \)th addition of ligand, and \( \lambda_{\theta} \) is the baricentric mean of the fluorescence spectrum before the addition of ligand. A red shift of the tryptophan emission spectrum was interpreted as an increased solvent exposure of tryptophan residues (51, 52).

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Note Added in Proof—After our manuscript was accepted, we became aware of results previously reported by Kumar \textit{et al.} (64). Their results partially overlap with the conclusions of this study but were not discussed in the original version of the manuscript. The authors sincerely apologize for this oversight.

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