ABSTRACT: Post-translational covalent modification by ubiquitin and ubiquitin-like proteins (UBLs) is a major eukaryotic mechanism for regulating protein function. In general, each UBL has its own E1 that serves as the entry point for a cascade. The E1 first binds the UBL and catalyzes adenylation of the UBL’s C-terminus, prior to promoting UBL transfer to a downstream E2. Ubiquitin’s Arg 72, which corresponds to Ala72 in the UBL NEDD8, is a key E1 selectivity determinant: swapping ubiquitin and NEDD8 residue 72 identity was shown previously to swap their E1 specificity. Correspondingly, Arg190 in the UBA3 subunit of NEDD8’s heterodimeric E1 (the APPBP1-UBA3 complex), which corresponds to a Gln in ubiquitin’s E1 UBA1, is a key UBL selectivity determinant. Here, we dissect this specificity with biochemical and X-ray crystallographic analysis of APPBP1-UBA3-NEDD8 complexes in which NEDD8’s residue 72 and UBA3’s residue 190 are substituted with different combinations of Ala, Arg, or Gln. APPBP1-UBA3’s preference for NEDD8’s Ala72 appears to be indirect, due to proper positioning of UBA3’s Arg190. By contrast, our data are consistent with direct positive interactions between ubiquitin’s Arg72 and an E1’s Gln. However, APPBP1-UBA3’s failure to interact with a UBL having Arg72 is not due to a lack of this favorable interaction, but rather arises from UBA3’s Arg190 acting as a negative gate. Thus, parallel residues from different UBL pathways can utilize distinct mechanisms to dictate interaction selectivity, and specificity can be amplified by barriers that prevent binding to components of different conjugation cascades.
targets, and downstream functions (9–12). In order to ultimately understand the unique functions of different UBLs, it is important to understand the molecular “code” by which UBLs are distinguished by their enzymatic and effector machineries. To date, some of the best progress in this area has been in deciphering sequences dictating E1 discrimination between ubiquitin and NEDD8. Several previous studies indicate that E1s provide critical UBL selection during initial noncovalent binding, and that this selection hinges on the identity of a UBL’s residue 72, the only C-terminal tail residue differing between ubiquitin (Arg) and NEDD8 (Ala) (Figure 1A). First, ubiquitin’s Arg 72 was identified as a critical determinant for E1 binding during adenylation (19). Second, “ubiquitinizing” NEDD8 with an Ala72Arg mutation allows binding to ubiquitin’s E1 (20), and “NEDD8ylizing” ubiquitin with Arg72Leu or Arg72Ala mutations allows ubiquitin to be activated by NEDD8’s E1 (21). Third, a “NEDD8ylized” ubiquitin can be transferred to NEDD8’s E2, albeit with ~80-fold diminished $k_{cat}/K_m$, suggesting either that the same UBL residue dictates both E1 and E2 selectivity, or E1 is fully responsible for dictating an E2’s UBL selection (22). This latter notion was supported by the finding that an E2’s UBL selectivity is dictated by side-chains involved in E1-, but not UBL- binding (23). Interestingly, ubiquitin and NEDD8’s residue 72 also plays a fundamental role in DUB specificity (24–26).

The basis for UBL selectivity has been partially characterized for NEDD8’s E1. The crystal structure of NEDD8’s heterodimeric E1 (APPBP1-UBA3) in complex with NEDD8 and ATP allowed modeling ubiquitin in place of NEDD8 in the structure (21). In the model, a conserved Arg in UBA3 (Arg190) would clash with ubiquitin’s specificity-conferring Arg72. By contrast, the corresponding residue in E1s for ubiquitin is absolutely conserved throughout evolution as a glutamine (Figure 1B), and a UBA3 Arg190Gln substitution allows adenylation of ubiquitin to some extent (21). Interestingly, in the wild-type E1-UBL complex, although UBA3’s Arg190 faces NEDD8’s Ala72, these two residues do not contact each other. There are two simplistic explanations for how the UBA3 Arg190Gln mutation allows NEDD8’s E1 to catalyze ubiquitin adenylation. First, it is possible that the polar Gln attracts ubiquitin through positive interactions with Arg72, and because NEDD8’s E1 lacks this Gln it fails to bind ubiquitin. Second, it is possible that NEDD8’s E1 actively prevents ubiquitin binding via electrostatic repulsion, with UBA3’s Arg190 repelling ubiquitin’s Arg72. Here we structurally and biochemically dissect roles of NEDD8’s residue 72 and UBA3’s residue 190, in order to gain further insights into mechanisms underlying E1-UBL selectivity.

**EXPERIMENTAL PROCEDURES**

**Protein Preparation.** Mutations were generated by PCR, and the entire coding sequence for each construct was verified by sequencing. Expression and purification of human APPBP1-UBA3, Ubc12, NEDD8, ubiquitin, and variants have been described (21, 23, 27, 28). Biochemical experiments were performed with full-length versions of APPBP1-UBA3. Surface plasmon resonance experiments examining noncovalent UBL interaction with APPBP1-UBA3 utilize UBA3 mutants harboring the catalytic Cys216Ala mutation, whereas those examining transthiolation retain the catalytic Cys. For BIACORE experiments, GST-APPBP1-UBA3 (Cys216Ala) and mutants were purified by glutathione affinity and gel filtration chromatography in 50 mM Tris-HCl, 150 mM NaCl, 0.5 mM DTT, pH 7.6. NEDD8 and variants were expressed as His-MBP-fusions from pRSF-1b with a GGS linker inserted upstream of NEDD8 to facilitate thrombin cleavage. His-MBP-NEDD8 was purified by nickel affinity, link inserted upstream of NEDD8 to facilitate thrombin cleavage. His-MBP-NEDD8 was purified by nickel affinity, link inserted upstream of NEDD8 to facilitate thrombin cleavage. His-MBP-NEDD8 was purified by nickel affinity, link inserted upstream of NEDD8 to facilitate thrombin cleavage. His-MBP-NEDD8 was purified by nickel affinity, link inserted upstream of NEDD8 to facilitate thrombin cleavage. His-MBP-NEDD8 was purified by nickel affinity, link inserted upstream of NEDD8 to facilitate thrombin cleavage.
antibodies (Biacore, Inc.) were covalently attached to a carboxymethyl dextran-coated gold surface (CM-4 Chip; Biacore, Inc.). The carboxymethyl groups of dextran were activated with N-ethyl-N’-(3-dimethylaminopropyl) carbodiimide (EDC) and N-hydroxysuccinimide (NHS), and anti-GST antibodies were attached at pH 5.0 in 10 mM sodium acetate. Any remaining reactive sites were blocked by reaction with ethanolamine. Anti-GST antibodies were immobilized at levels of ~4000 RU for each flow cell.

The kinetics of association and dissociation were monitored at a flow rate of 100 µL/min. GST-APPBP1-UBA3 constructs were captured to a level of ~1250~1450 RU for each experiment. GST was captured on the reference surface to account for any nonspecific binding to the GST tag. The UBL analytes were prepared in 10 mM Tris (pH 7.6), 150 mM NaCl, 0.1 mg/mL bovine serum albumin, and 0.005% P20 surfactant. Binding was measured for concentration ranges of 41 nM to 10 µM for all UBLs. To account for injection artifacts, a series of sensorgrams was recorded throughout the experiment after injecting only buffer (blank injections). The analytes dissociated completely from the chip surfaces, eliminating the need for a regeneration step. Data reported are the difference in SPR signal between the flow cell containing the APPBP1-UBA3 construct and the reference cell without APPBP1-UBA3. Additional instrumental contributions to the signal were removed by subtraction of the average signal of the blank injections from the reference-subtracted signal (29). Triplicate injections were made, and the data were analyzed globally by equilibrium affinity analysis using the software package Scrubber 2 (BioLogic Software).

**Ubc12–UBL Thioester Assays.** APPBP1-UBA3-catalyzed formation of the thioester-linked Ubc12–NEDD8 covalent intermediate was assayed in 20 µL volumes containing 100 nM of APPBP1-UBA3 or mutants, 3 µM Ubc12, 4 µM wild-type NEDD8 or mutants, and 50 mM Tris-HCl, 50 mM NaCl, 10 mM MgCl₂, 5 mM ATP, 1 mM DTT, pH 7.6. The reactions were started by adding the E1 enzyme to the assay mixture, incubated at room temperature for 5 min, and quenched by the addition of 10 µL 2× nonreducing SDS–PAGE sample buffer. Reaction products were resolved by 15% SDS–polyacrylamide gels, and transferred to nitrocellulose membranes. The membranes were saturated with 3% bovine serum albumin (BSA) in TBS-T (50 mM Tris-HCl, 150 mM NaCl, 0.1% Tween 20, pH 7.4) prior to incubation with anti-NEDD8 antibody. Anti-NEDD8 polyclonal antibodies (RCA) were raised in rabbits against the C216A Arg190Gln-NEDD8Ala72Arg (hereafter referred to as APPBP1-UBA3 Arg190Gln-NEDD8Ala72Arg or “wild-type-opposite”, Figure 1C) and APPBP1(Δ254–258)-UBA3(AN12 C216A) Arg190Ala-NEDD8Ala72Arg (hereafter referred to as APPBP1-UBA3Arg190Ala-NEDD8Ala72Arg or “wild-type- opposite”, Figure 1C) and APPBP1(Δ254–258)-UBA3(AN12 C216A)-NEDD8Ala72Gln (hereafter referred to as APPBP1-UBA3Arg190 wt-NEDD8Ala72Gln or “ubiquitinized-opposite”, Figure 1C) were grown with reservoir solution containing 0.1 M Tris, 0.4 M ammonium acetate, 9~10% PEG 10K, 5 mM DTT, pH 7.0~8.0. These crystals were soaked in 0.1 M Tris, 0.4 M ammonium acetate, 12~13% PEG 10K, 5 mM DTT, pH 7.0~8.0 supplemented with increasing concentrations of glycerol (5~30%, in 5% increments) prior to flash-freezing in the presence of 30% glycerol and data collection at the 8.2.2 beamline at the Advanced Light Source. All crystals formed with four complexes in the asymmetric unit in the space group P2_1_2_1, with unit cells and packing similar to the wild-type complex reported previously (21). Reflection data were indexed, integrated and scaled using HKL2000 (31). The structure of APPBP1-UBA3Arg190Ala-NEDD8Ala72Arg was determined first, by rigid body refinement using a model derived from the structure of wild-type APPBP1-UBA3-NEDD8 (21), in which UBA3’s Arg190 side-chain was truncated after the β-carbon. Rigid body refinement, simulated annealing, energy minimization and restrained B-factor refinement were carried out using the program CNS (32), iteratively with manual rebuilding using O (33). The structure of APPBP1-UBA3Arg190Ala-NEDD8Ala72Arg, but with NEDD8’s residue 72 side-chain truncated after the β-carbon, was then used as a model for the other two structures, which were refined using a similar protocol. Simulated annealing (2000 K) omit maps confirm the presence and the correct placement of mutated residues in all three structures. In one complex from the asymmetric unit of all these structures, additional vector-derived residues N-terminal of NEDD8 participate in crystal packing. The four complexes in the asymmetric unit for the final model of APPBP1-UBA3Arg190Ala-NEDD8Ala72Arg contain: APPBP1 residues 6–204, 207–534; 6–200, 202, 204–205, 208–534; 6–203, 208–253, 261–534; 6–203, 205–534; UBA3 residues −1 and 12–442; −1 and 12–442; −1 and 12–441; −1 and 12–441; and NEDD8 residues −10 to −1 and 1–76; 1–76; 1–76; 1–76, respectively. The four complexes in the asymmetric unit for the final model of APPBP1-UBA3Arg190Gln-NEDD8Ala72Arg contain: APPBP1 residues 6–200, 208–534; 6–201, 211–534; 6–199, 212–252, 262–534; 6–200, 209–534; UBA3 residues −1 and 12–442; −1 and 12–441; −1 and 12–441; and NEDD8 residues −10 to −1 and 1–76; 1–76; 2 to −1 and 1–76; −1 and 1–76, respectively. The four complexes in the asymmetric unit for the final model of APPBP1-UBA3Arg190 wt-NEDD8Ala72Gln contain: APPBP1 residues 6–204, 207–534; 6–203, 205–534; 6–200, 202, 204–205, 208–253, 260–534; 6–203, 208–534; UBA3 residues −1 and 12–442; −1 and 12–442; −1 and 12–441; −1 and 12–441; and NEDD8 residues −10 to −1 and 1–76; 1–76; 2 to −1 and 1–76; −1 and 1–76, respectively.
12–441; −1 and 12–442; −1 and 12–441; and NEDD8 residues 10 to 1 and 1–76; 1–76; 1–76; 1–76, respectively. Other residues are not modeled, presumably due to disorder. Ramachandran statistics were analyzed with Procheck (34). Structures are shown using PyMol (35).

RESULTS

U3A’s Arg190 Gates against Binding to a UBL with a Basic Side-Chain at Residue 72. In order to quantify effects of mutations on APPBP1-UBA3-NEDD8 interactions, we developed a binding assay using BIACORE (Table 1, Figure 2). In our assay, GST-APPBP1-UBA3 binds NEDD8 with a $K_d$ value of 333 ± 1 nM. Furthermore, a NEDD8 mutant harboring the “ubiquitinizing” Ala72Arg mutant does not display any binding in our assay. Thus, we examined the role of U3A’s Arg190 in this discrimination. Consistent with previous studies, “ubiquitinizing” GST-APPBP1-UBA3 with an Arg190Gln mutation allows binding to NEDD8 Ala72Arg, with a $K_d$ of 1.02 ± 0.1 µM.

We next asked whether relief from negative repulsion from the E1’s Arg is sufficient to allow binding of NEDD8 Ala72Arg, offline whether attractive electrostatic interaction with the E1’s Gln is required. A U3A Arg190Ala substitution would address this question by removing barriers from the Arg side-chain, without providing opportunities for favorable polar interactions with the UBL Arg. GST-APPBP1-UBA3 Arg190Ala binds NEDD8 Ala72Arg with a $K_d$ value of 1.66 ± 0.01 µM. The similar $K_d$ values (1.02 ± 0.1 µM v 1.66 ± 0.01 µM) for the Gln and Ala substitutions suggest that U3A’s Arg190 primarily acts as a gate that negatively selects against a UBL’s Arg72. The U3A Arg190Ala substitution also allows binding to a NEDD8 with a Lys substitution at residue 72, although in this case a Gln does appear to impart more positive interactions (3-fold better binding).

U3A’s Residue 190 and NEDD8’s Residue 72 Do Not Play Equivalent Roles in E1-Ubl Selectivity. The U3A Arg190Ala-NEDD8 Ala72Arg pair essentially swaps Arg and Ala selectivity determinants between E1 and NEDD8. Thus, we asked whether these E1 and UBL positions play equivalent roles in determining binding with a complete set of Ala, Arg, and Gln substitutions at both positions (Table 1, Figure 2). The results reveal consistent, nonequivalent, directional effects. First, the wild-type E1-NEDD8 pair binds best, ∼5-fold better than the “opposite” pair in which the identities of U3A’s residue 190 and NEDD8’s 72 have been swapped [U3A Arg190Ala and NEDD8 Ala72Arg]. Second, the “ubiquitinizing” pair (with the U3A Arg190Gln and NEDD8 Ala72Arg substitutions to residues found in ubiquitin’s E1 and ubiquitin) associate with a ∼9-fold lower $K_d$ than the “opposite ubiquitinized” pair, wild-type U3A (Arg190) and NEDD8 Ala72Gln.

Interestingly, both the APPBP1-UBA3 Arg190Gln and Arg190Ala variants preferentially bind to NEDD8 Ala72Arg (or NEDD8 Ala72Lys), rather than to wild-type NEDD8 (Ala72), or the NEDD8 Ala72Gln mutant. This raises the possibility that an Arg serves a function beyond dictating E1-UBL selectivity.

NEDD8’s E1 Contains Additional Determinants Selecting against Ubiquitin. Having established U3A’s residue 190 as gating against a UBL’s Arg72, we asked the extent to which this single interaction establishes APPBP1-UBA3’s selectivity against initial noncovalent binding to ubiquitin. Although binding is too weak for us to measure a $K_d$ within the limits of our assay, some signal is detected for interactions between wild-type GST-APPBP1-UBA3 and ubiquitinArg72Ala, but not for wild-type ubiquitin (Figure 2D). In a related vein, GST-APPBP1-UBA3 Arg190Gln and Arg190Ala do show evidence of interaction with wild-type ubiquitin. Therefore, although the same trends are observed for mutations allowing APPBP1-UBA3 interaction with “ubiquitinized” NEDD8 Ala72Arg, additional residues also contribute to NEDD8 E1’s discrimination against noncovalent binding to ubiquitin.

NEDD8’s E1-E2 Transthiolation with ULBs Harboring Arg at Position 72. Although wild-type ubiquitin is excluded from NEDD8’s E1, a previous study found that NEDD8’s E1 could transfer the “NEDD8ylized” ubiquitinArg72Leu mutant to NEDD8’s E2 (Ubc12), albeit with less efficiency overall than the NEDD8 transthiolation cycle (22). This result raised the question as to whether specificity was established entirely by noncovalent binding by the E1, or whether the ubiquitin mutation relieves discrimination against an Arg at the UBL’s position 72 by APPBP1-UBA3 and Ubc12 during transthiolation. To address this question, we assayed the different APPBP1-UBA3 mutants for NEDD8 Ala72Arg transfer to Ubc12 (Figure 3). We observe no detectable activity with wild-type APPBP1-UBA3, consistent with gating against this “ubiquitinizing” mutation. However, both the APPBP1-UBA3 Arg190Gln and Arg190Ala mutants can transfer NEDD8 Ala72Arg to Ubc12. Similar results are also observed for ubiquitin, although reactions are less efficient, consistent with the weaker noncovalent binding of ubiquitin to these mutants.

Crystallographic Dissection of Arg, Ala, and Gln at U3A’s Residue 190 and NEDD8’s Residue 72. Our biochemical data pointed toward a directional gating mechanism by which NEDD8’s E1’s Arg190 influences interaction with a UBL’s residue 72. In order to understand the structural basis underlying this specificity, we determined crystal structures of three mutant APPBP1-UBA3-NEDD8 complexes: “wild-type-opposite” [U3A Arg190Ala-NEDD8 Ala72Arg, 2.85 Å resolution]; “ubiquitinizing” [U3A Arg190Gln-NEDD8 Ala72Arg, 3.05 Å resolution]; and “ubiquitinizing-opposite” [U3A Arg190 (wt)-NEDD8 Ala72Gln, 2.90 Å resolution]. Details of data and refinement statistics are given in Table 2. Overall, for all three complexes, the electron density 2Fo-Fc maps were continuous and well-defined over all four copies in the asymmetric units (data not shown). Moreover, the side chains at both positions NEDD8’s E1 residue 190 and NEDD8 residue 72 were clearly present in simulated annealing omit maps (Figure 5), generated
from a model lacking the residues UBA3’s 190 and NEDD8’s 72. For each mutant complex, the four complexes in the asymmetric unit are similar: “wild-type-opposite”, 0.5 Å rmsd, “ubiquitinized”, 0.6 Å rmsd, and “ubiquitinized-opposite”, 0.5 Å rmsd. Therefore, the figures display only one copy for each mutant complex. The overall structures all superimpose well with the prior wild-type APPBP1-UBA3-NEDD8 structure, with rmsds between 0.4 and 0.5 Å.

**Structures of “Wild-Type-Opposite” and “Ubiquitinized” Complexes: Unlocking UBA3’s Gate against a UBL’s Arg72.** In order to understand how UBA3’s Arg190 gates against a UBL’s Arg72, we compared the structures of the previously determined wild-type APPBP1-UBA3-NEDD8 complex, in which UBA3 harbors an Arg, and the “wild-type-opposite” and “ubiquitinized” complexes, in which NEDD8 has an Arg. Strikingly, in all three complexes, the Arg guanidium group is in the same relative location (Figure 4). This explains how UBA3’s Arg would gate against a UBL’s Arg by clashing and repulsion. The results also show how simply removing UBA3’s Arg190 side-chain with an Ala mutation allows a UBL’s Arg.

In all three structures, the Arg makes many favorable contacts (Figure 5). In the wild-type complex, Arg190 forms hydrogen bonds with UBA3’s Tyr207 and Thr203, and a...
salt-bridge with UBA3’s Asp179. These latter two electrostatic interactions are preserved when the Arg comes from the mutant NEDD8. The aliphatic portion of the Arg side-chain also preserves hydrophobic interactions with Tyr207 and Tyr321. In the “ubiquitinized” complex, a hydrogen bond between NEDD8’s mutant Arg72 and UBA3’s mutant Gln190 explains the moderate preference for the “ubiquitinizing” Gln over an Ala at position 190 in UBA3.

**Insights into Directionality of the UBA3 190 and UBL 72 gate from the “Wild-Type-Opposite” Structure.** We wished to obtain a structural understanding of the directional preference toward residue identity at UBA3’s position 190.
and NEDD8’s position 72, revealed from our binding assay.

Comparison of the wild-type (UBA3Arg190, NEDD8Ala72) with the “wild-type-opposite” (UBA3 mutant Ala190, NEDD8 mutant Arg72) reveals that swapping the Arg side-chain between UBA3 and NEDD8 removes van der Waals contacts, between UBA3 Arg190s gamma and delta carbons and Asp179’s beta carbon (Figure 5A-B). Asp179 is responsible for orienting UBA3’s Arg151, which in turn aligns the backbone of UBA3’s Mg$^{2+}$-binding Asp146 and NEDD8’s C-terminal Gly-Gly motif. Although at the resolution of our structures subtle structural differences are not detectable, we speculate that slight variations in this region influences affinity for NEDD8.

Although Arg-Gln and Gln-Arg electrostatic interactions are observed in both the “ubiquitinated” (UBA3 mutant Gln190, NEDD8Ala72) and “ubiquitinated-opposite” (UBA3 Arg190, NEDD8 mutant Gln72) complexes, the “ubiquitinated-opposite” (UBA3 Arg190, NEDD8 mutant Gln72) interaction is lower affinity. The Arg190 hydrophobic contact to UBA3’s Asp179 is lost in both of these complexes, as in the “wild-type-opposite” complex (Figure 5C-D). Moreover, the UBA3 Asp179 - Arg salt-bridge is lost in the “ubiquitinated-opposite”, in which the UBA3 Arg guanidium group has moved away, in order to accommodate and form a hydrogen bond with the NEDD8 mutant Gln side-chain (Figure 4C).

The structure of the “ubiquitinated-opposite” complex suggests that the relatively higher affinity interaction between wild-type UBA3, with an Arg190, and wild-type NEDD8, with an Ala72 is an indirect effect. Despite the lack of positive electrostatic 190–72 interactions, an Ala at NEDD8’s position 72 allows proper positioning of UBA3’s Arg. By contrast, in order to accommodate a Gln at NEDD8’s position 72, other interactions are disrupted for relocation of UBA3’s Arg190 (Figure 4C, Figure 5D).

**DISCUSSION**

UBL modifications regulate a vast array of eukaryotic pathways. Thus, it is of great interest to understand the detailed mechanisms by which different UBLs are directed to their targets, and thereby modify function. Ubiquitin’s and NEDD8’s residue 72 has emerged as a key determinant distinguishing protein–protein interactions between the two cascades. Here we dissect the structural mechanisms by which the identity of a UBL’s residue 72, and the corresponding E1’s residue 190, dictates interaction selectivity. We find that ubiquitin’s Arg72 actively prevents misactivation by NEDD8’s E1 due to direct negative exclusion by UBA3’s Arg190. This contrasts with a model where NEDD8’s E1 simply lacks residues dictating positive interactions with ubiquitin’s Arg72.

Negative selectivity has emerged as a general mechanism separating the ubiquitin and NEDD8 pathways. In addition to barriers preventing NEDD8’s E1 from interacting with ubiquitin, NEDD8’s E2 (Ubc12) also displays negative
selectivity determinants that oppose mismatching with ubiquitin (23). Ubc12’s UBL specificity, from NEDD8 to ubiquitin, was switched 1010-fold simply by removing surface elements, without substituting sequences of ubiquitin E2s (23). NEDD8’s E1 also displays barriers against structures conserved among ubiquitin E2s (36). It seems likely that negative interactions will be found to play major roles in distinguishing members of other UBL cascades. Indeed, negative interactions may hinder binding between the HECT E3 E6AP and the ubiquitin E2 UbcH5 (37), and the altered specificity of a charge-swapped RING E3 CNOT4-UbcH5B pair may involve altered negative selectivity as well as designed positive interactions (38).

Subtle structural differences, which may be distributed over ubiquitin’s and NEDD8’s entire surfaces, also contribute substantial selectivity. Such differences appear to add E1-UBL specificity, on top of the predominant role of a UBL’s residue 72 (19, 22) (also Figure 2D, Figure 3B). Not surprisingly, the native UBA3Arg190-NEDD8Ala72 interaction is most favored. Our structural data suggest this may be an indirect effect of proper positioning of UBA3’s Arg190, rather than from direct positive interactions with NEDD8’s Ala72. Fine variation also appears to contribute at least a factor of 10 to the differential binding of NEDD8 and ubiquitinArg72Ala to the NEDD8-specific ubiquitin-like protease DEN1/NEDP1/SENP8 (25, 26). Insights into how slight differences in presentation of ubiquitin surface residues can lead to large differences in affinities come from the finding that hydrophobic core mutations selectively diminish binding to ubiquitin-interacting motifs, but not to ubiquitin-associated domains (39).

In contrast to apparently indirect functions of NEDD8’s residue 72, ubiquitin’s Arg72 mediates important positive interactions with many cognate partners. The most striking effect is observed for ubiquitin interactions with the ubiquitin C-terminal hydrolase Yuh1, where the NEDD8Ala72Arg mutant binds Yuh1 as well as ubiquitin (24). The structure of a Yuh1-ubiquitin aldehyde complex revealed essential positive interactions between Yuh1’s Asp35 and a UBL’s Arg72 (24). In a related vein, a ubiquitin Arg72Ala mutation decreases cleavage by the ubiquitin-specific protease HAUSP (26). It seems that ubiquitin’s Arg72 likely makes positive interactions with ubiquitin’s E1 (19, 20). Consistent with this notion, we observe hydrogen bonding between a UBL’s Arg72 and a "ubiquitinized" UBA3 Gln190 mutant. Ubiquitin’s E1 also contains an Asp in place of UBA3’s Leu206 and Tyr207, which further contact NEDD8’s residue 72. Thus, residue 72 in ubiquitin and NEDD8 appear to mediate specificity in distinct ways.

It seems likely that structures mediating key positive interactions in some other UBL pathways may also serve as negative determinants that prevent interaction with noncognate conjugation cascades. Indeed, it is possible that conceptually related, but distinct mechanisms confer specificity even among E1-UBL antecedents found in prokaryotes. The bacterial enzymes MoeB and ThiF are prokaryotic ancestors of E1s, with roles in molybdopterin and thiamin biosynthesis, respectively (3, 40). MoeB and ThiF resemble the E1 adenylation domain both structurally and functionally, catalyzing adenylation of the bacterial ubiquitin-fold proteins MoaD and ThiS. Here, specificity may be dictated by UBL insertions, between the /beta2A and /beta2B strands in MoaD and the /betaD and /betaE strands in ThiS, which make unique contacts to MoeB and ThiF, respectively (41, 42). Interestingly, the insertions in both MoaD and ThiS contain arginines, which may also gate against noncognate interactions.
Role of UBA3 Arg190 in NEDD8’s E1 UBL Specificity

Knowledge of the “code” for selective interactions will lay a foundation for manipulating UBL ligation machineries to transform protein activities as desired.

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