Ubiquitin-Activated Interaction Traps (UBAITs) identify E3 ligase binding partners

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Abstract

We describe a new class of reagents for identifying substrates, adaptors, and regulators of HECT and RING E3s. UBAITs (Ubiquitin-Activated Interaction Traps) are E3-ubiquitin fusion proteins and, in an E2- and E2-dependent manner, the C-terminal ubiquitin moiety forms an amide linkage to proteins that interact with the E3, enabling covalent co-purification of the E3 with partner proteins. We designed UBAITs for both HECT (Rsp5, Itch) and RING (Psh1, RNF126, RNF168) E3s. For HECT E3s, trapping of interacting proteins occurred in vitro either through an E3 thioester-linked lariat intermediate or through an E2 thioester intermediate, and both WT and active-site mutant UBAITs trapped known interacting proteins in yeast and human cells. Yeast Psh1 and human RNF126 and RNF168 UBAITs also trapped known interacting proteins when expressed in cells. Human RNF168 is a key mediator of ubiquitin signaling that promotes DNA double-strand break repair. Using the RNF168 UBAIT, we identify H2AZ—a histone protein involved in DNA repair—as a new target of this E3 ligase. These results demonstrate that UBAITs represent powerful tools for profiling a wide range of ubiquitin ligases.

Keywords  HECT E3s; RING E3s; Ubiquitin; Ubiquitin ligases
Subject Categories Methods & Resources; Post-translational Modifications, Proteolysis & Proteomics
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Introduction

Ubiquitin is a post-translational modifier that determines the fate of a large fraction of all proteins in eukaryotic cells, and the specificity of ubiquitylation is controlled by hundreds of ubiquitin ligases (> 600 in human cells) [1]. The complex network of enzyme-substrate interactions that mediate ubiquitylation is only partially characterized, with the vast majority of E3s remaining essentially uncharacterized with respect to substrates and functions. Unraveling this network has the potential to yield many insights into cellular regulation and disease pathways and provide innumerable opportunities for developing clinically useful modulators of specific E3s or classes of E3s [2].

E3 enzymes interact either directly or indirectly, via adaptor proteins, with their substrates. In human cells, E3s are supported by two E1 enzymes (Uba1 and Uba6) and approximately 60 E2 enzymes [3,4]. The E1 enzymes activate ubiquitin in an ATP-dependent reaction and transfer ubiquitin to E2 proteins, catalyzing formation of a thioester bond between the terminal carboxyl group of ubiquitin and the active-site cysteine of the E2s [5]. RING E3s are the most numerous type of ubiquitin ligases, representing approximately 95% of all E3s encoded by human cells [1]. The RING domain forms a platform for recruiting a ubiquitin-charged E2 protein. The E3 facilitates transfer of ubiquitin from the E2 to one or more lysines of an E3-bound substrate [6,7] and can activate the E2 for discharge of ubiquitin [8–11]. For simple RING E3s, substrate recruitment is mediated by other domains within the E3 protein, while for more complex E3s, such as CRLs, other proteins within the multi-protein complex mediate substrate recognition (e.g., F-box proteins). RING E3s control a wide range of cellular processes and play a key role in a wide range of human disease states, including cancer and the innate immune response to viral infections (Mdm2 and TRIM5, respectively) [12,13].

HECT E3s are relatively few in number, with five in Saccharomyces cerevisiae and 28 human HECT E3s. They are defined by a approximately 350 amino acid C-terminal catalytic domain (the HECT domain) and range in size from approximately 90 kDa to over 500 kDa. The regions N-terminal to the HECT domain are involved in substrate recognition, localization, and regulation [14]. Structures of isolated HECT domains have revealed that the catalytic domain consists of an approximately 250 amino acid N-terminal lobe, containing the E2 binding site, and a approximately 100 amino acid C-terminal lobe, containing the active-site cysteine [15–17]. The HECT domain binds ubiquitin-charged E2s, ubiquitin is transferred to the active-site cysteine of the E3, and the E3 directly catalyzes substrate ubiquitylation.

In some cases, substrate recognition by HECT E3s is mediated by obvious protein–protein interaction modules, such as WW domains. WW domains define the Nedd4 subfamily of HECT E3s, which includes S. cerevisiae Rsp5 and human Nedd4, Nedd4L, Itch, and others [18]. WW domains are 30–45 amino acids in length and recognize proline-containing “PY” motifs (PPXY consensus), with most Nedd4 family substrates, substrate adaptors, or regulatory
proteins containing one or more PY motifs [19–22]. At least half of all human HECT E3s do not have obvious protein–protein interaction modules, and a basis for substrate recognition is not readily apparent. Such an example is human E6AP/Ube3A, which is involved in HPV-associated cervical cancer and Angelman syndrome, a severe neurologic disease [23–25]. Other HECT E3s involved in important aspects of human biology include Huw1/Arf-BP1, Herc1, Herc2, and Herc5 [26]. A third class of E3s, the RBR E3s, have biochemical characteristics of both RING and HECT E3s, in that they contain a RING domain that recruits an E2 enzyme, yet they also contain an active-site cysteine that accepts ubiquitin from the E2 and transfers it to substrates [27–29]. Important members of this class of E3s include Parkin, HOIP, and HHARI [30].

Many genetic and biochemical approaches have been employed to identify substrates and regulatory proteins of E3s, including yeast two-hybrid assays, co-immunoprecipitation approaches, and protein–protein interaction arrays [31–35]. While each of these has its own advantages and disadvantages, we sought a method that would be applicable to a wide range of E3s and would overcome the challenges posed by potentially weak or transient enzyme–substrate interactions. We describe a method, based on E3-ubiquitin fusion proteins, to covalently trap E3s to their substrates and other interacting proteins.

**Results**

**HECT E3 UBAITs**

The C-terminus of ubiquitin is critical for all chemistry involved in ubiquitin activation and conjugation. Ubiquitin molecules with small N-terminal epitopes are generally very good substrates in conjugation reactions (e.g., 6×His-Ub [36]), and we envisioned that a very large N-terminal epitope—consisting of a approximately 100-kDa HECT E3 with a flexible linker connecting it to ubiquitin—might also be competent for activation by the E1 ubiquitin-activating enzyme and subsequent transfer to an E2 enzyme (Fig 1A; species i and ii, respectively). If so, we predicted, based on the structure of a HECT domain in complex with an E2-ubiquitin complex [37], that the active-site thiolate of the HECT domain might be able to attack the Ub–E2 thioester bond in an intramolecular reaction, forming a thioester-linked protein lariat (Fig 1A, species iii). The reaction of a substrate lysine with the lariat protein would then yield an amide-linked E3-Ub-target protein complex (Fig 1A, species iv), thus covalently “trapping” the substrate protein to its E3. Alternatively, even if the lariat structure did not form, the direct reaction of a substrate lysine with the Ub–E2 thioester would be predicted to yield an identical product (species v, via species iv). In both cases, affinity purification of the HECT-Ub fusion protein would co-purify covalently trapped target proteins, which could then be identified by standard LC-MS/MS techniques. We refer to these tools for identification of substrates, and potentially regulatory or other interacting proteins, as UBAITs, for Ubiquitin-Activated Interaction Traps.

**Characterization of Rsp5 UBAITs in vitro**

Saccharomyces cerevisiae Rsp5 is one of the most extensively characterized HECT E3s, with many identified substrates, adaptors, and regulatory proteins [38,39]. We purified bacterially expressed Rsp5 UBAITs with a 12-amino acid linker (consisting of GA dipeptide repeat) between the E3 and ubiquitin (Fig 1B). UBAITs consisted of either wild-type Rsp5 fused to wild-type ubiquitin (WT UBAIT), the Rsp5 active-site C-to-A mutant fused to wild-type ubiquitin (C-A UBAIT), or wild-type Rsp5 fused to a non-conjugatable form of ubiquitin (ΔGG UBAIT, deleted of the terminal GG residues). The UBAITs were 32P-labeled at an N-terminal kinase recognition epitope and assayed for their ability to form E1 and E2 thioester complexes. As shown in Fig 2A, the WT UBAIT formed an E1 thioester in the presence of E1 alone (lane 2), and the addition of both E1 and E2 (lane 4), but not E2 alone (lane 3), resulted in the appearance of a band of the predicted size for a UBAIT–E2 adduct. The E1 and E2 adducts were greatly diminished in the presence of DTT (lanes 6 and 8), although residual E2 adducts remained (lane 8), likely a reflection of E2 auto-conjugation (i.e., auto-UBAITylation). These results indicate that the ubiquitin moiety of the wild-type Rsp5 UBAIT is competent for activation by E1 and transfer to E2. The Rsp5 C-A UBAIT behaved identically to the WT UBAIT in the E1 and E2 thioester assays, while the AGG UBAIT, as expected, did not form any E1 or E2 adducts (H.F. O’Connor and J.M. Huibregtse, unpublished observation). No extra bands with the WT UBAIT, corresponding to a potential lariat thioester, were detected in the
presence of E1 and E2, although the lariat would not necessarily migrate differently than the UBAIT protein, itself.

To determine whether the Rsp5 UBAIT was capable of covalently trapping targets in vitro, the WT, C-A, and ΔGG UBAITs were combined with E1 and E2 enzymes and products were analyzed by SDS–PAGE in the absence or presence of DTT in the loading buffer. The migration points of the UBAIT–E1 and UBAIT–E2 thioesters are indicated. The asterisk marks a contaminant of the purification process (uncut GST-UBAIT).

**Figure 2. Rsp5 UBAITs are functional in vitro.**

A Rsp5 can be activated by E1 and transferred to an E2. Purified 32P-labeled Rsp5 UBAIT (WT) was incubated with or without E1 and E2 enzymes, and products were analyzed by SDS–PAGE in the absence or presence of DTT in the loading buffer. The migration points of the UBAIT–E1 and UBAIT–E2 thioesters are indicated. The asterisk marks a contaminant of the purification process (uncut GST-UBAIT).

B Rsp5 UBAITs were incubated with E1 and E2 proteins, as indicated, and 32P-labeled Wbp2. The migration points of the major products are indicated with arrows.

C Rsp5 UBAITs were incubated with E1 and E2 proteins, as indicated, and 32P-labeled Sna3. The migration points of the major products are indicated with arrows. The asterisk marks a contaminant of the purification process (uncut GST-Sna3).

not the ΔGG, formed a stable covalent conjugate of the predicted size to both target proteins (compare lanes 4 and 6 of Fig 2B and C). Conjugation formation with the WT UBAIT was shown to be dependent on the PY motif of Sna3 (Appendix Fig S1). Higher molecular weight conjugates corresponded in size to multi-UBAITylated targets, which are likely to arise due to binding of the PY motif of a UBAIT-target conjugate to a WW domain of a second UBAIT molecule. The apparent inefficiency of the substrate UBAITylation reaction compared to the substrate ubiquitylation catalyzed by Rsp5 is due, at least in part, to the fact that the UBAIT reaction is single-turnover, whereas Rsp5 rapidly depletes free substrate by polyubiquitylating many molecules of the substrate over a relatively short time period [40]. Interestingly, the C-A mutant UBAIT also formed conjugates to both target proteins (lane 5, Fig 2B and C), although less efficiently than with WT UBAIT, particularly for Sna3. This suggests that the Rsp5 UBAIT is capable of trapping targets by both mechanisms illustrated in Fig 1A: either through nucleophilic attack of an E3-Ub lariat intermediate, or by direct attack of the E2 thioester. Based on the relative efficiency of the WT UBAIT to C-A UBAIT reactions, the lariat mechanism appears to be the predominant mechanism in vitro.

We examined the effect of the linker length on target trapping in vitro by expressing Rsp5 UBAITs containing a GGSG flexible linker sequence [33,41,42] repeated 1, 3, or 5 times, resulting in linker lengths of 4, 12, or 20 amino acids. As shown in Appendix Fig S2, all three WT UBAITs supported target trapping in vitro, with the longer linker lengths supporting more efficient conjugation to Wbp2. Therefore, a range of linker lengths appears to be permissible for the Rsp5 UBAIT, with a longer linker being generally more favorable.

**In vivo validation of Rsp5 UBAITs**

To determine whether UBAITs were able to trap substrates or interacting proteins in vivo, N-terminal TAP-tagged Rsp5 UBAITs (WT, C-A, and ΔGG, with a 12-amino acid GA-repeat linker) and a TAP-only control protein (consisting of two copies of protein A and the calmodulin-binding protein) were expressed from a galactose-regulated multicopy plasmid in a wild-type RSP5 haploid yeast strain. An anti-TAP immunoblot showed a similar distribution of conjugate formation with the WT UBAIT was shown to be dependent on the PY motif of Sna3 (Appendix Fig S3), suggesting that the Rsp5 UBAIT conjugates, expression of the TAP-UBAIT proteins was induced for three hours, cell lysates were prepared, and the proteins were affinity-purified via the protein A tag on IgG sepharose. Bound proteins were then denatured with SDS and heat and, after dilution of the SDS, purified on IgG sepharose a second time. The final products were subjected to SDS–PAGE, and gel slices were excised from above the migration point of the TAP-Rsp5 proteins (approximately 130 kDa). In-gel tryptic digestions
were performed, and peptides were identified by LC-MS/MS. Results were analyzed using Thermo Proteome Discoverer software with high stringency filters. Proteins with two or more spectral counts from the WT UBAIT sample and greater than or equal to five-fold higher spectral counts in the WT UBAIT sample compared to the TAP control were considered “positive hits” and potential Rsp5-interacting proteins. Table 1 shows the total spectral counts for known Rsp5-interacting proteins that met the filtering criteria in two biological replicates (25 total proteins), and Appendix Table S1 shows an additional 57 proteins that also met filtering criteria and have not previously been reported to interact with Rsp5. Among the known Rsp5-interacting proteins were regulatory proteins (Rup1, Ubp2), substrates (Mup1, Zrt1, Sec7, Rpb1, Mga2), and adaptor proteins (Art5, Rvs167, Lsb1, Bsd2) (Table 1). The potential novel interactors included Sul2 (a sulfur permease), two plasma membrane-associated t-SNAREs (Sso1 and Sso2), and Rvs161, a lipid raft protein that interacts with Rvs167 [44], a known Rsp5 target protein (Appendix Table S1).

Further examination of the Rsp5-interacting proteins showed that they fell into two groups with respect to UBAIT trapping. The first group contained proteins that were isolated with both the WT and C-A UBAITs, but not the ΔGG UBAIT. There were no proteins that were isolated solely with the WT UBAIT, further indicating that the predominant mechanism of trapping in vivo was the non-lariat mechanism. Both known and potentially novel Rsp5 interacting proteins were present in this group, as well as a subset of ubiquitin E2 proteins (Ubc4, Ubc6, and Ubc13), which were likely to have been isolated as a result of E2 auto-UBAITylation (as observed in vitro; Fig 2A). The second group of proteins was, surprisingly, isolated with all three UBAITs, including the non-conjugatable ΔGG UBAITs, suggesting that these were isolated via non-covalent binding to the Rsp5 proteins. Many of these proteins were well-characterized Rsp5-interacting proteins, including Bul1, several Art proteins (Art1, Art3, Art4, Art6, Art8), and Rpb1 [32,45–48]. In general, the spectral counts for these conjugation-independent hits were among the highest of any of the proteins identified, consistent

Table 1. Known Rsp5-interacting proteins isolated with Rsp5 UBAITs in Saccharomyces cerevisiae.

<table>
<thead>
<tr>
<th>ACC#</th>
<th>Protein</th>
<th>MW [kDa]</th>
<th>Experiment 1</th>
<th>Experiment 2</th>
<th>PY-like motif</th>
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Total spectral counts of known Rsp5-interacting proteins isolated with the Rsp5 WT UBAIT, C-A UBAIT, ΔGG UBAIT, and empty TAP control (TAP). Proteins shown passed the following filtering criteria in two biological replicates (Experiments 1 and 2): two or more spectral counts for the WT sample and greater than or equal to five-fold more counts in WT sample compared to TAP control. Identified proteins were analyzed for the presence of PY-like motifs [94].
with these proteins being among the most abundant and/or robust Rsp5-interacting proteins. It is likely that these proteins were able to re-bind non-covalently to the Rsp5 proteins in the second IgG pull-down, after denaturation and subsequent dilution of the SDS. To test this possibility, an additional experiment was performed with the WT and ΔGG UBAITs in parallel with TAP-Rsp5 (non-UBAIT) as a control. As shown in Appendix Table S2, the same proteins that were isolated with the ΔGG UBAIT were also isolated with Rsp5 (non-UBAIT). In contrast, those that had only been isolated with the WT and C-A UBAITs (and not the ΔGG UBAIT) were not isolated with TAP-Rsp5 (Appendix Table S3). As most of these non-covalently interacting proteins were bona fide Rsp5-interacting proteins, further attempts to eliminate them during purification were not made.

A potential caveat to the UBAIT approach is the use of the UBAIT as a source of ubiquitin in reactions catalyzed by other cellular E3s. We therefore determined whether Rsp5 (non-UBAIT) could utilize, in vitro, a RING E3 UBAIT (based on Psh1; discussed further below) as the source of ubiquitin in a ubiquitylation reaction. As shown in Appendix Fig S5, Rsp5 could indeed use Psh1-Ub as a source of ubiquitin to modify Wbp2 in vitro. This suggests that many other cellular E3s might also be capable of using a large ubiquitin fusion protein in a substrate ubiquitylation reaction. At one extreme, one might therefore have expected to isolate the complete cellular “ubiquitinome” with the Rsp5 UBAIT. This clearly was not the case, however, as 39% of all covalently trapped proteins (16 out of 41) represented known Rsp5 interactors. A possible explanation for these seemingly contradictory results is that the UBAITylated products of other cellular E3s would be distributed among a very large number of substrates, such that the signal from any given protein would be very low, whereas the signal from the covalently trapped Rsp5 substrates would be concentrated among a relatively small number of proteins. The fact that the Rsp5 WT and C-A UBAITs isolated a broad range of known interacting proteins (as did other UBAITs; below) implies that the potential use of the fusion proteins as a source of ubiquitin is not a significant impediment to the approach.

Itch UBAITs in human cells

Itch is one of nine human Nedd4 family HECT E3s. These E3s have many functions and binding partners in human cells and play a role in ubiquitin-mediated endocytosis and immune regulation [49,50]. TAP-Itch UBAITs (WT and ΔGG) were expressed by transfection in HEK293T cells to determine whether UBAITs could trap targets in human cells. Cell lysates were prepared 36 h post-transfection, and conjugates were purified on IgG sepharose. An anti-TAP immunoblot showed a distribution of high molecular weight conjugates to the WT UBAIT, but not the ΔGG UBAIT (Appendix Fig S6). As with Rsp5, the ubiquitin moiety did not destabilize Itch in human cells (Appendix Fig S4B), and a distribution of conjugates emanated upward from the Itch WT UBAIT, but not the ΔGG. Total UBAIT conjugates were analyzed by mass spectrometry as described above for Rsp5. Three biological replicates of the experiment were performed.

Table 2 shows total spectral counts for proteins that co-purified specifically with the Itch WT UBAIT in all three replicates. Out of the 18 proteins that met the filtering criteria, seven were known Nedd4 family interactors (38%). These proteins included Stam and Hrs, which function in endosomal sorting complexes required for transport (ESCRT) pathway [50,51]. Alix, also part of the ESCRT pathway, was isolated, which has been reported to interact with other Nedd4 family members [52,53]. Wbp2, which interacts with the WW domains of Nedd4 family proteins, was also isolated [54]. Unlike the Rsp5 UBAITs, there were no proteins that co-purified with both the WT and the ΔGG Itch UBAITs.

RING E3 UBAITs

The results presented above indicated that the Rsp5 UBAITs trapped targets in cells via a non-lariat mechanism, independent of the HECT catalytic cysteine. This suggested that the approach could be applied to other types of E3s, including RING E3s, which do not function via a thioester intermediate. This was first tested with Psh1, a yeast RING E3. Psh1 is responsible for ubiquitilation of Cse4/CENP-A, a histone H3 variant. Cse4 is normally localized to centromeric chromatin, and it is ubiquitylated and degraded when it is mislocalized to euchromatin [55,56]. Cse4 is so far the only known substrate of Psh1. It was also recently shown that Psh1 interacts with Sp16, a component of the FACT (Facilitates Chromatin Transcription/Transactions) complex, and that this interaction is necessary for Psh1 to ubiquitylate Cse4 in cells but not in vitro [57].

To determine whether the Psh1 UBAIT was capable of covalently trapping Cse4 in vitro, Psh1 UBAITs (WT and ΔGG) were combined with E1 and E2 (Ubch) proteins and 32P-labeled Cse4. As predicted, the WT UBAIT, but not the ΔGG, formed a covalent conjugate to Cse4 (Appendix Fig S7A).

TAP-tagged Psh1 UBAITs (WT and ΔGG) were expressed in yeast. TAP-Psh1 UBAITs were expressed at similar levels to TAP-Psh1, indicating that the ubiquitin moiety did not result in destabilization of the fusion protein (Appendix Fig S4C). Conjugates were purified and analyzed as described above. Three biological replicates were performed, and a total of 15\* proteins were common to all three replicates (Table 3). As predicted, both Cse4 and Sp16 were isolated specifically with the WT Psh1 UBAIT, but not the ΔGG UBAIT. Two other histone proteins (H3 and H4) were also isolated. While these are not thought to be ubiquitylated by Psh1, they have been shown previously to co-immunoprecipitate with Psh1, likely as a result of their indirect association with Cse4 in nucleosome complexes [56]. Their covalent “trapping” to Psh1 may therefore reflect their proximity to Cse4 in nucleosome complexes. Two proteasome proteins were also trapped (Rpn10 and Pup3), although the significance of this is not known. It should be noted that while the spectral counts for both Sp16 and Cse4 were very low, the fact that both proteins were identified in three independent experiments points to the specificity and sensitivity of the UBAIT method for profiling RING E3s.

RNF126 is a human RING ubiquitin ligase involved in degradation of mislocalized and aggregation-prone proteins. This pathway is dependent on the interaction of RNF126 with the Bag6 complex.
UBAIT was capable of covalently trapping Bag6 (Bag6/Trc35/Ubl4A) [58]. To determine whether the RNF126 conjugates to only the WT UBAIT (Fig 3B). Transfected cells were confirmed expression of all three proteins and formation of expressed by transfection in HEK293T cells (Fig 3A). Immunoblot-D TAP-RNF168 and TAP-RNF168 UBAITs (WT and characterized by immunodeficiency and radiosensitivity [62 number of quality control substrates of the Bag6 complex. Many additional proteins were also isolated with the WT UBAIT, which may reflect the large WT UBAIT trapped Bag6 in three biological replicates, and the specific targets using human RNF168. RNF168 is a key mediator of DNA repair [60,61], and RNF168 mutations are associated with Riddle syndrome, a human disorder characterized by immunodeficiency and radiosensitivity [62–64]. TAP-RNF168 and TAP-RNF168 UBAITs (WT and ΔGG) were expressed by transfection in HEK293T cells (Fig 3A). Immunoblotting confirmed expression of all three proteins and formation of conjugates to only the WT UBAIT (Fig 3B). Transfected cells were treated with ionizing radiation, and purification and mass spectrometry analysis of the UBAITs identified histone H2A only in the WT UBAIT sample (Table 4). Ubiquitylation of H2A by RNF168 is a vital signaling event in the DNA damage response and represents the most well-established target of RNF168 [65,66]. In addition, histone H2AZ, a member of the H2A family of histone variants, was identified as a potentially novel RNF168 substrate. H2AZ is recruited to sites of DSBs to facilitate repair by non-homologous end joining [67]. To validate H2AZ as a target, we first overexpressed RNF168 in HEK293T cells. This led to an additional H2AZ ubiquitylation product of H2AZ, beyond the previously characterized monoubiquitylated species generated by the RING1B ligase (Fig 3C) [68]. RNF168 also ubiquitylated H2AZ in vitro in H2AZ-containing reconstituted nucleosome core particles (NCPs) (Fig 3D). The D94A mutation in H2AZ disrupts an acidic patch that is required for repair activity. The analogous mutation in H2A blocks recruitment of RNF168 to nucleosomes [69], and this mutation also prevented in vitro ubiquitylation of H2AZ in NCPs (Fig 3D). Together, these results validate H2AZ as a target of RNF168 and demonstrate the utility of the UBAIT approach for identification of novel ubiquitin ligase targets.

### Discussion

In vitro and in vivo results presented here validate Ubiquitin-Activated Interacting Traps (UBAITs) as useful tools for identification

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Total spectral counts of targets identified with Itch WT UBAIT, ΔGG UBAIT, and empty TAP control (TAP). Experiment was performed in triplicate. Data was filtered according to the following parameters: proteins with four or more spectral counts for the WT sample and greater than or equal to five-fold more counts in WT sample compared to TAP control. Identified proteins were analyzed for the presence of PY-like motifs [54]. Known Nedd4 family binding partners are shown in bold.
of substrates and other interacting proteins for both HECT and RING ubiquitin ligases. While the approach was conceived to be specific for HECT ubiquitin ligases (e.g., Rsp5 and Itch), the lack of dependence on the active-site cysteine suggested that the same approach could be applied to RING ubiquitin ligases, and this was validated with three different RING E3s (Psh1, RNF126, and RNF168). Furthermore, the RNF168 UBAIT identified histone H2AZ as a new target of this E3.

Thirty percent of the proteins that were isolated with the Rsp5 UBAIT and passed filtering criteria were known Rsp5-interacting proteins, and these represented approximately 32% of all proteins reported to interact with Rsp5 (Appendix Table S5). Of the known interacting proteins that were covalently trapped by the Rsp5 WT and C-A UBAITs in yeast cells, many were substrate adaptor proteins, including six out of the ten Art proteins [70], rather than classical ubiquitylation substrates. For example, Art5 was trapped, but the Rsp5/Art5 target, ltr1 [71], was not. Art1 was also trapped, and while the Rsp5/Art1 target Mup1 was trapped, Can1 and Lyp1 were not [70]. A bias toward adaptor proteins might simply reflect the steady-state distribution of Rsp5 with its associated proteins in cells. Also, the yeast experiments reported here were performed under standard growth conditions, and the lack of identification of some targets might reflect the physiologic state of the cells. For example, the ubiquitylation of Fur4 by Rsp5 occurs predominantly under nutrient deprivation [72,73].

Several potential novel targets of Rsp5 were also identified. Bmh2, a 14-3-3 protein, which has previously been shown to regulate the Rsp5 adaptor Bul1 [74], was isolated, and Ino1 (inositol 3-phosphate synthase) and Pdi1 (a protein disulfide isomerase) were also previously isolated in large-scale proteomic analyses of Rsp5-associated proteins [75,76]. Sul2 (a sulfur permease), two plasma membrane-associated t-SNAREs (Sso1 and Sso2), and Rvs161, a lipid raft protein, were also isolated, consistent with the role of Rsp5 in membrane protein trafficking. Other membrane-associated interacting proteins included Vti1 and Sac1, which are Golgi membrane proteins, and Pil1, which is localized to the mitochondrial membrane [77-79].

Endocytic pathway proteins were well represented among the Itch UBAIT conjugates in human cells, including Alix and SCAMP3, which have been reported to interact with other Nedd4 family members [52,53]. Expression of Itch UBAITs in other cell types may yield target proteins related to the function of Itch in immune regulation. For example, Bcl-10, JunB, and Cbl-b, which were not isolated with the Itch UBAIT, are Itch targets that play a role in immune responsiveness [80-83].

The UBAIT approach with RING E3s was validated with yeast Psh1 and human RNF126 and RNF168. Psh1 associates with Spt16, and this interaction is required for recognition and ubiquitylation of Cse4/CENP-A [57]. The UBAIT trapped both Spt16 and Cse4, the only two known interactors of Psh1, which points to the sensitivity and specificity of the approach. The Psh1 UBAIT also trapped other histone proteins, which may reflect the proximity of those proteins to Cse4 in nucleosome complexes. This highlights the potential usefulness of UBAITs as tools to map intracellular locale or proximity to other proteins. While the RNF126 UBAIT trapped its best-characterized associated protein, Bag6, it also trapped a large number of previously uncharacterized RNF126-interacting proteins. We speculate that at least some of these may reflect quality control client proteins of the Bag6/RNF126 complex [58]. The RNF168 UBAIT trapped histone H2A, the best-characterized substrate of this ligase, as well as H2AZ, a previously unknown substrate of RNF168. The nucleosome acidic patch region of H2AZ was required for

Table 3. Proteins identified in Psh1 UBAITs in Saccharomyces cerevisae.

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Total spectral counts for proteins common to all three biological replicates of Psh1 WT UBAIT, ΔGG UBAIT, and empty TAP control (TAP). Data were filtered according to the following parameters: proteins with two or more spectral counts for the WT sample and greater than or equal to five-fold more counts in WT sample compared to TAP control. Known Psh1-interacting partners are shown in bold.
ubiquitylation events [34]. While the UBA fusion approach was designed to specifically characterize ubiquitin ligases, the UBAIT approach is not necessarily limited to characterizing ubiquitin ligases. It has the potential to be used as a general protein–protein interaction tool, where virtually any protein of interest (POI) could be fused to ubiquitin as a way of trapping its interacting partners. This assumes that the POI does not have to be a ubiquitin ligase in order to trap interacting proteins, and preliminary experiments with RING UBAITs deleted of the RING domain suggest this is the case (H.F. O’Connor and J.M. Huibregtse, unpublished observation). Similar to the UBAIT concept, the “NEDDylator” system and the BioID biotinylation-based proximity ligation approaches can potentially be adapted for any POI [33,34]. The NEDDylator is a POI-Ubc12 fusion protein, which can be added to cell lysates or expressed in cells along with a tagged form of NEDD8. A subsequent tagged NEDD8 immunoprecipitation identifies NEDDylated conjugates, including the targets of the POI. In the BioID system, the addition of exogenous biotin and a promiscuous BirA-POI fusion protein biotinylates proximal binding partners. The principal advantage of the UBAIT method over these is that the reaction product is covalently trapped to the POI, as opposed to being a reaction product that is no longer in complex with the bait protein; this may increase the confidence that a target protein was indeed trapped as a result of close proximity to the bait protein.

The UBAIT system may be further optimized by modification of either the linker peptide connecting the POI to ubiquitin or the purification epitope. While we have not yet observed significantly different outcomes based on linker length or linker sequence for HECT or RING UBAITs, it is conceivable that specific UBAITs might have unique requirements. The UBAITs described here have an N-terminal purification epitope in addition to the C-terminal linker-ubiquitin epitope. Incorporation of the affinity purification into or adjacent to the linker sequence would allow one-step C-terminal epitope tagging which would be useful for genomic tagging of individual genes in yeast or human cells. Finally, while HECT E3s share a common architecture, with the HECT domain located at the C-terminus of the protein, monomeric RING E3s are structurally very diverse. Of the three RING E3s examined here, two have their RING domain at their extreme N-terminus (Psh1 and RNF168), while the RING domain of RNF126 is located near the C-terminus of the protein. The fact that all three RING UBAITs successfully isolated interacting proteins suggests that UBAIT functionality is not limited in any obvious way by domain

RNF168-dependent ubiquitylation, and although this region has been shown to promote ubiquitylation of H2A [66,69,84,85], H2AZ contains an extended acidic patch which may confer unique structure and function to H2AZ compared to other H2A family members [86].

The potential to capture potentially weak and transient protein–protein interactions is a key feature of the UBAIT approach. Other approaches for characterizing E3s have been recently reported, including one that fuses UBA ubiquitin binding domains to F-box proteins to capture the product of F-box protein-mediated ubiquitylation events [34]. While the UBA fusion approach was designed to specifically characterize ubiquitin ligases, the UBAIT approach is not necessarily limited to characterizing ubiquitin ligases. It has the potential to be used as a general protein–protein interaction tool, where virtually any protein of interest (POI) could be fused to ubiquitin as a way of trapping its interacting partners. This assumes that the POI does not have to be a ubiquitin ligase in order to trap interacting proteins, and preliminary experiments with RING UBAITs deleted of the RING domain suggest this is the case (H.F. O’Connor and J.M. Huibregtse, unpublished observation). Similar to the UBAIT concept, the “NEDDylator” system and the BioID biotinylation-based proximity ligation approaches can potentially be adapted for any POI [33,34]. The NEDDylator is a POI-Ubc12 fusion protein, which can be added to cell lysates or expressed in cells along with a tagged form of NEDD8. A subsequent tagged NEDD8 immunoprecipitation identifies NEDDylated conjugates, including the targets of the POI. In the BioID system, the addition of exogenous biotin and a promiscuous BirA-POI fusion protein biotinylates proximal binding partners. The principal advantage of the UBAIT method over these is that the reaction product is covalently trapped to the POI, as opposed to being a reaction product that is no longer in complex with the bait protein; this may increase the confidence that a target protein was indeed trapped as a result of close proximity to the bait protein.

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### Table 4. Proteins identified in RNF168 UBAIT samples in human 293T cells.

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Total spectral counts of targets identified with RNF168 WT UBAIT, ΔGG UBAIT, and empty TAP control (TAP) following ionizing radiation treatment. Experiment was performed in triplicate. Data were filtered according to the following parameters: proteins with four or more spectral counts for the WT sample and greater than or equal to five-fold more counts in WT sample compared to TAP control (see Appendix Table S4 for complete unfiltered data). Identified proteins were analyzed for the presence of PY-like motifs [94].
Materials and Methods

DNA constructs and recombinant protein expression

*Escherichia coli* expression vectors and the production of Ubc1, Sna3–K125, Sna3–PAAA, and wild-type Rsp5 and Wbp2 proteins were previously described [40, 87, 88]. The MYC-RNF168 construct was previously described [69], and bacterial expression vectors for core histones (human H2B, H3, and H4) in *E. coli* and pPROEX HTa-RNF168 (residues 1–113) were previously described [89]. Itch was PCR-amplified from *Mus musculus* cDNA. Itch UBAITs, Rsp5 (GGS)5 UBAIT WT and Psh1 UBAITs were generated by homologous recombination in *Saccharomyces cerevisiae* strain FY56 (MATa his4-1128 rps12-1286 ura3-52) [45]. RNDF2 UBAITs were generated using the one-step cloning vector for generating E3-Ub fusion proteins (Appendix Fig S8). Further cloning information and primer sequences are provided below.

ORFs were sub-cloned into pGEX-6p-1 and pGEX-6p-K for bacterial expression of GST fusion proteins [40], and were sub-cloned into pYES2-NTAP or pcDNA3-TAP for in vivo expression in yeast and human cells, respectively [87, 90]. pYES2-NTAP constructs were transformed into *S. cerevisiae* FY56. GST fusion proteins were expressed in *E. coli* by standard methods and affinity-purified on Glutathione-Sepharose beads (GE Healthcare). Proteins expressed from pGEX-6p-1 vector were cleaved from GST by using PreScission protease (GE Healthcare) under manufacturer-recommended conditions, and these proteins were used in ubiquitin-thioester and ubiquitination reactions. The His6-tagged RNF168 (1–113) construct was expressed in Rosetta 2 (DE3) pLysS cells and purified over a Ni-NTA column (Qiagen) as described [69].

Overlapping extension PCR

Rsp5 (GGS)5 UBAIT WT and ΔGG were generated by overlapping PCR. Using Rsp5 wild-type and C-A mutant plasmids as template [91], round 1A product was generated with primers Rsp5F (GGGGCG GCCGCT CACGCA GG) and Rsp5UBaGA6R (CCGGCT CCAGCC CCGCGG CCTCTT TGACAA AACCTT ATG). Using yeast ubiquitin as template, round 1B product was generated with primers Psh1F (GGGGCG GCCGCT CATCTT AGCACA AGA), respectively. Round 2 PCR was generated using 1A and 1B as template and primers Rsp5F and yUbWTR or yUbGGR to generate Psh1 UBAIT WT and Psh1 UBAIT ΔGG, respectively.

Similarly, Psh1 UBAIT WT and ΔGG were constructed by overlapping PCR. For round 1A PCR, Psh1 was amplified from *S. cerevisiae* FY56 genomic DNA using PSH1F (GGGGAA TTCATG GGCGAC GAATTA CACACA CG) and PSH1UbLX1R (CCAGAT CCGCGG CTCATG TCACAG TCTCTT AGCTAG TCTCTT AGC) (round 1A, Psh1). Round 1B (yeast ubiquitin wild-type or ΔGG mutant) was generated with PSH1F (CCAGAT CCGCGG CTCATG TCACAG TCTCTT AGCTAG TCTCTT AGC) and yUbWTR or yUbGGR, respectively. Round 2 PCR was generated using 1A and 1B as template and primers PSH1F and yUbWTR or yUbGGR to generate Psh1 UBAIT WT and Psh1 UBAIT ΔGG, respectively.

Itch UBAIT WT and ΔGG were generated usingITCHF (CCGGAT CCTCTT AAGATG ACCAGT ACTCGC ATCTTG TTTCTG) and ITCHUBLX1R (GCTGGA GCCGGG GCAGGG GCCCAG ATTTTC GTCTCT) as template and primers PSH1F and yUbWTR or yUbGGR to generate Itch UBAIT WT and Itch UBAIT ΔGG, respectively.

Homologous recombination

Rsp5 (GGS)5 UBAIT WT and ΔGG were generated by homologous recombination in a two-round process with the pYES2-NTAP vector in *S. cerevisiae*. Firstly, wild-type or ΔGG mutant ubiquitin PCR product was amplified with GGSG-encoding oligonucleotides at the 5′ end and 30–60 bp overhangs complementary to the desired site of recombination in pYES2-NTAP at the 5′ end (RILX1F (CCCTCG CCCGCC TGAGGG ATGAGC ATGGCG GGGCGG CTATCT GACGGA GAGGAG GGTCCG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively). Restriction digestion of 100 ng pYES2-NTAP with BgIII was performed overnight and transformed, along with 600 ng of GGSG-Ub PCR product [92]. Following selection on SC-URA agar plates, DNA was extracted from yeast cells. Positive clones were verified by electroporation of the yeast plasmid DNA into *E. coli*, followed by sequence analysis. In a second round, Rps5 was amplified by PCR along with overhangs complementary to pYES2-NTAP at the 5′ end (R2RSP5LX5R (ACCAGA TCCGCC ACCAGA TCCGCC ACCAGA AG) and R2RSP5LX1R (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively). Rps5 (GGS)5 UBAIT WT and ΔGG were generated using pYES2-NTAP Rsp5 (GGS), UBAIT WT as template, using R2RSP5LSX5F (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively) and 3′ end (R1WTUBR (TGAATG TAAGCG TGACAT CGGTGG GTAGGG GTGGGG GTTCG) for (GGS)1, 3, 5, respectively).
TCCGCC ACCACT TCCACG TCCGGA ACCCGA CCGTCC TCCCGA ACC

One-step UBAIT cloning vector

A one-step cloning vector was constructed to allow a UBAIT to be constructed with one round of PCR and ligation. Wild-type and ΔGG mutant ubiquitin were amplified using UBVVX1F (AAATTG GCAGCC GCAGTC GCTCAG ATTTTC TCAAG ACT) and UBVVX1R (CTTCTT TCTAGA TCAACG ACCCTT TACGG TACAGT AAGGG) or ubGGVVX1R (CTTCTT TCTAGA TCAACG ACCCTT TACGG TACAGT AAGGG) using H2AZF (ATGGCC GAGGCG TCGCCG CA) and H2AZR (GAGCTA GCGGCC GCCGAG TTGCTT) and cloning into pcDNA3-TAP. Itch (no Ub fusion) was generated by PCR with ATGGCC GAGGCG TCGCCG CA) and RNF126R (GAGCTA GCGGCC GCCGAG TTGCTT) and cloning into pcDNA3-TAP. Psh1 (no Ub fusion) was generated by PCR from FY56 genomic DNA using PSH1F (GGGGAA TTCATG ATCCAG) and RNF168MUTR (CTGGAT GGATCT GGTCAG ATTTTC GTCAAG ACT) and cloning into pcDNA3-TAP-GCTCTA CCCAAA GACGCC) and RNF168VX1R (TTTGCG GCCGC C) to generate pGEX-6p-K-Bag6 (1–2015) harboring N-terminal 6× His tag. S'-biotin-tagged 601 nucleotide sequence to be used for reconstitution of nucleosomes was generated as previously described [69].

Mutations in H2AZ (for D94A) were generated by site-directed mutagenesis following standard protocols with H2AZD94AF (GCAAAT TGCTAT TGCTTG AGCTGA AGAATT GGATTC TCTC) and H2AZD94AR (GAGAGA ATCCAA TTCTTC AGCTCC ACGAAT) using H2AZD94AF

Biochemical assays

Ubiquitin-thioester reactions were performed as described previously [87]. Reaction mixtures were incubated for 15 min for thioester assays at room temperature. For substrate ubiquitylation assays, 32P-labeled proteins, generated as described previously [40], were incubated with 5 μg/ml of Rsp5 or Rsp5 UBAIT in the presence of 25 mM Tris–HCl (pH 7.5), 50 mM NaCl, 5 mM MgCl2, 0.1 mM DTT, 50 μg/ml of ubiquitin (Sigma), 1.25 μg/ml of human E1 (Boston Biochem), 1.25 μg/ml of UbC1 (40 μl total volume), and 10 mM ATP. All reactions were initiated by the addition of ubiquitin or UBAIT and stopped by the addition of 4× SDS–PAGE loading buffer either lacking or containing 0.4 M DTT. Reaction mixtures were incubated for 1 h at room temperature. Ubiquitylated products were analyzed using Typhoon FLA 9500 (GE Healthcare).

For RNF168 in vitro ubiquitylation assays, 2.5 μg of recombinant mononucleosomes was incubated in a 50-μl reaction buffer containing 50 μM Tris–HCl, pH 7.5, 100 mM NaCl, 10 mM MgCl2, 1 μM ZnOAc, 1 mM DTT, 30 nM ubiquitin-activating enzyme E1 (Boston Biochem), 1.5 μM ubiquitin-conjugating enzyme UbcH5a (Boston Biochem), 4 μM RNF168 (1–113), 22 μM ubiquitin (Boston Biochem), and 3.33 mM ATP at 30°C for 4 h [69].

Construction of non-UBAIT-encoding plasmids

Cse4 was amplified by PCR from S. cerevisiae genomic DNA using Cse4F (GGGAAT TCAATG CAAGTA ACAAAC AATTG) and Cse4R (GGGCAG CCGGCT AAATAA ACTGTC CCTGGG AA) to generate pGEX-6p-K-Cse4. Bag6 was amplified by PCR from HEK293T cDNA with RNF126VX1F (ATAGTC GAATTC ATGGCC GAGGCG TCGCCG CA) and RNF126VX1R (GGGCGG CCGCTT ACTCTT GTCCAA ATCCTT) and cloning into pcDNA3-TAP. Itch (no Ub fusion) was generated by amplifying RNF126 by PCR from HEK293T cDNA with RNF126VX1F (ATAGTC GAATTC ATGGCC GAGGCG TCGCCG CA) and RNF126VX1R (GAGCTA GCGGCC GCCGAG TTGCTT) and cloning into pcDNA3-TAP. Psh1 (no Ub fusion) was generated by PCR from FY56 genomic DNA using PSH1F (GGGGAA TTCATG ATCCAG) and RNF168MUTR (CTGGAT GGATCT GGTCAG ATTTTC GTCAAG ACT) and cloning into pcDNA3-TAP-GCTCTA CCCAAA GACGCC) and RNF168VX1R (TTTGCG GCCGC C) to generate pGEX-6p-K-Bag6 (1–167 amino acids).

RNF168 (no Ub fusion) was generated by amplifying RNF126 by PCR from HEK293T cDNA with RNF126VX1F (ATAGTC GAATTC ATGGCC GAGGCG TCGCCG CA) and RNF126VX1R (GAGCTA GCGGCC GCCGAG TTGCTT) and cloning into pcDNA3-TAP. Iitch (no Ub fusion) was generated by PCR with ITCHF (CGGGAT CCTCTG AACAGT GCCGCC AGCCT) andITCHR (GGGGCC CCGTCT ACTCTG GCCGAA CT) and cloning into pcDNA3-TAP. Iitch (no Ub fusion) was generated by PCR with S.cerevisiae FY56 genomic DNA using PSH1F (GGGAAT TCAATG CAAGTA ACAAAC AATTG) and cloning into pcDNA3-TAP. Psh1 (no Ub fusion) was generated by PCR from S.cerevisiae FY56 genomic DNA using PSH1F (GGGAAT TCAATG CAAGTA ACAAAC AATTG) and cloning into pcDNA3-TAP. Psh1 (no Ub fusion) was generated by PCR from S.cerevisiae FY56 genomic DNA using PSH1F (GGGAAT TCAATG CAAGTA ACAAAC AATTG) and cloning into pcDNA3-TAP. Psh1 (no Ub fusion) was generated by PCR from S.cerevisiae FY56 genomic DNA using PSH1F (GGGAAT TCAATG CAAGTA ACAAAC AATTG) and cloning into pcDNA3-TAP.

Human H2AZ cDNA was cloned into a N-terminal FLAG-tagging vector using Gateway LR reaction. Human H2AZ cDNA was cloned using H2AZF (ATGGCT GGCGGT AGGCTG) and H2AZR (TTAGAGT TCGCTG GCCGCT) into Gateway compatible entry vector (pDONR201) and subcloned into bacterial expression vector
collected by gentle centrifugation, and following three washes with RIPA buffer, the cells were pelleted, and 75 µl SDS-PAGE loading buffer was added to the beads. Beads were boiled at 95°C for 5 min. Samples were analyzed by SDS-PAGE (8% gel) and stained with Coomassie blue. In addition, a fraction of the material was reserved for an anti-TAP immunoblot. The gel section above the expected size of the UBAIT was excised and, along with the corresponding section in the TAP empty control, was subject to in-gel tryptic digestion and LC-MS/MS. This protocol was also followed for Psb1 UBAITs, except cells were grown to OD$_{600nm}$ of 0.5 prior to the addition of 2% galactose.

TAP-tagged protein purification from human cells

For large-scale UBAIT purifications, ten 10-cm plates of 293T cells (3 µg DNA per 10 cm plate) were transfected with each plasmid construct. RNF168 samples were treated with 10 grays of ionizing radiation (Faxitron X-ray) prior to lysis. Cells were harvested, lysed, and pooled 36 h post-transfection in NP-40 lysis buffer. Cleared cell lysates were collected and diluted with 4 ml RIPA buffer. Conjugates were purified on IgG sepharose beads (GE Healthcare), binding at 4°C for 4 h. Beads were washed three times with high salt RIPA buffer (400 mM NaCl), followed by three times with RIPA buffer. Proteins were eluted from beads by adding SDS-PAGE loading buffer and boiling the samples at 95°C for 5 min, and the samples were analyzed by SDS-PAGE and stained with Coomassie blue. The gel section above the unconjugated UBAIT was excised from the gel, subject to in-gel tryptic digestion and LC-MS/MS.

Mass spectrometry

Samples were subject to in-gel digestion and protein identification using LC-MS/MS at the University of Texas at Austin Proteomics Facility as previously described, with minor adjustments [93]. Briefly, LC-MS/MS was carried out by reverse phase using a Dionex Ultimate 3000 Nano UPLC interfaced with a Thermo Orbitrap Elite mass spectrometer in positive mode. The LC utilizes an Acclaim PepMap 100 Nano-Trap column (75 µM × 2 cm, C18, 3 µM, 100 angstroms) with two Acclaim PepMap RSLC columns in tandem with each other: a 15-cm column (75 µM × 15 cm, C18, 2 µM, 100 angstroms) and a 25-cm column (75 µM × 15 cm, C18, 3 µM, 100 angstroms). A packed spray tip with 15 cm of C18 was utilized, and the total column resin length is 55 cm. Peptides were eluted from a linear gradient starting with 5–40% buffer B over a period of 120 min with a flow rate of 300 nL/min. MS spectra and MS/MS spectra were acquired via data-dependent acquisition, with the full-scan MS occurring in the Orbitrap at 120,000 resolution. The top 20 most intense precursor ions were then selected for MS/MS CID fragmentation and the product ions were detected in the ion trap, with a dynamic exclusion of 60 s.

Spectra were searched against the UniProt human database (Itch, RNF126 and RNF168) and UniProt yeast database (Rsp5 and Psb1) along with decoy databases using SEQUEST HT (Proteome Discoverer 1.4, Thermo Scientific). Fully tryptic peptides with up to 2 missed cleavages were considered. Mass tolerance filters of 10 ppm (MS1) and 0.8 Da (MS2) were applied. PSMs were filtered using Percolator (implemented in Proteome Discoverer) to control false discovery rates (FDR) to < 1% as determined using a reverse sequence decoy database.

Proteins with two or more spectral counts from the WT UBAIT sample and greater than or equal to five-fold higher spectral counts in the WT UBAIT sample compared to the TAP control were considered “positive hits” and potential Rsp5-interacting proteins. Hits that were common to unrelated UBAITs in yeast (Cpr1, Ddi1, Ecm21, histone H2B, Met4, Pal1, Pgk1, Uba1, Ubc1, Ub14, Yhr097c) and human (Amot, Fam115a, Spg20) cells were not further considered as potential interactors. Raw data files are available at ProteomeXchange, accession number PXD002791.

NCP reconstitution

Recombinant human histones H2B, H3, and H4 were expressed in Escherichia coli BL21 (DE3)/RIL cells and purified as described [69]. Histone H2AZ was extracted from the soluble fraction. Octamers were refolded from purified histones by mixing the four histones in equimolar ratios (10% more of H2AZ/H2B relative to H3/H4), followed by dialysis into 2 M NaCl, and then purified on a Superdex 200 (16/60) size exclusion column (GE Healthcare). Nucleosome core particles (NCPs) were reconstituted by salt deposition and confirmed as previously described [69].

Western blot

Western blots were detected using Li-cor Odyssey Detection System (Li-cor Biosciences). For RNF168 experiments, ubiquitylated products were analyzed using ECL Prime Western blotting detection reagent (GE Healthcare).

Antibodies


Primary antibody rabbit anti-H2AZ (2718, Cell Signaling Technologies) and secondary antibody HRP-linked goat anti-rabbit IgG (7074, Cell Signaling Technologies).

Author contributions

The concept was developed by HFO and JMH, and experiments were performed by HFO, NL, JWL, PA, and CDS. The paper was written by HFO, KMM, and JMH.

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Conflict of interest
The authors declare that they have no conflict of interest.

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Correction added on 2 December 2015 after first online publication: References in the Materials and Methods section were corrected and references [91–93] were added.