Enhanced Purification of Ubiquitinated Proteins by Engineered Tandem Hybrid Ubiquitin-binding Domains (ThUBDs)*

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Ubiquitination is one of the most common post-translational modifications, regulating protein stability and function. However, the proteome-wide profiling of ubiquitinated proteins remains challenging due to their low abundance in cells. In this study, we systematically evaluated the affinity of ubiquitin-binding domains (UBDs) to different types of ubiquitin chains. By selecting UBDs with high affinity and evaluating various UBD combinations with different lengths and types, we constructed two artificial tandem hybrid UBDs (ThUBDs), including four UBDs made of DSK2p-derived ubiquitin-associated (UBA) and ubiquitin 2-derived UBA (ThUDQ2) and of DSK2p-derived UBA and RABGEF1-derived A20-ZnF (ThUDA20). ThUBD binds to ubiquitinated proteins, with markedly higher affinity than naturally occurring UBDs. Furthermore, it displays almost unbiased high affinity to all seven lysine-linked chains. Using ThUBD-based profiling with mass spectrometry, we identified 1092 and 7487 putative ubiquitinated proteins from yeast and mammalian cells, respectively, of which 362 and 1125 proteins had ubiquitin-modified sites. These results demonstrate that ThUBD is a refined and promising approach for enriching the ubiquitinated proteome while circumventing the need to overexpress tagged ubiquitin variants and use antibodies to recognize ubiquitin remnants, thus providing a readily accessible tool for the protein ubiquitination research community. Molecular & Cellular Proteomics 15: 10.1074/mcp.O115.051839, 1381–1396, 2016.

Ubiquitination, a universal post-translational modification, refers to the covalent attachment of ubiquitin to lysine residues or the N terminus of proteins (1). It is a cascade process catalyzed by ubiquitin-activating enzyme (E1), ubiquitin-conjugating enzyme (E2), and ubiquitin ligase (E3) (2). Ubiquitination is reversible by the action of deubiquitinating enzymes, which cleave ubiquitin moieties from the substrates (3, 4). Proteins with ubiquitin-binding domains (UBDs) can bind to ubiquitin, thereby expanding the functional diversity of the ubiquitination modification signaling network (5, 6). In addition, ubiquitination plays pivotal regulatory roles in a broad spectrum of cellular processes such as protein stability, gene transcription, cell cycle progression, DNA damage, and the immune response (7–10).

Ubiquitin can be conjugated to substrate proteins in different forms, including monoubiquitin, multiple monoubiquitin, and polyubiquitin. All seven lysine residues and the N terminus of ubiquitin can be involved in the formation of polyubiquitin chains (11). Different lengths and linkages of ubiquitin modifications are linked to distinct physiological functions in cells. For example, monoubiquitin regulates DNA repair and receptor endocytosis (12–14), and polyubiquitin chains, such as the well studied Lys-48-linked and Lys-63-linked polyubiquitin.

1 The abbreviations used are: UBD, ubiquitin-binding domain; hUBD, hybrid UBD; ThUBD, tandem hybrid UBD; UBA, ubiquitin-associated domain; LQT, linear trap quadrupole; SPR, surface plasmon resonance; GSH, glutathione; NHS, N-hydroxysuccinimide; ACN, acetonitrile; SILAC, stable isotope labeling with amino acids in cell culture; AQUA, absolute quantification; Ub, ubiquitin; RP-LC, reversed-phase liquid chromatography; qUBA, the four tandem ubiquitin-1 UBD domain; Ni-NTA, nickel-nitrilotriacetic acid; FA, formic acid; HCC, human hepatocellular carcinoma.

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ubiquitin chains, function in proteasomal degradation (15), endosomal trafficking to the lysosome, intracellular signaling, and DNA repair (16). The specific N-terminal linear polyubiquitin chain activates NF-κB kinase (17). Recent studies also suggest that Lys-11 linkage plays important roles in the endoplasmic reticulum-associated protein degradation pathway (18) and cell cycle (19), whereas Lys-33 linkage regulates cell surface receptor-mediated signal transduction (20) and post-Golgi transport (21). However, the biological functions of other atypical ubiquitinated chains still remain poorly understood (22).

Large scale profiling of ubiquitin conjugates and mapping of ubiquitination sites are instrumental to understanding the ubiquitin regulatory system. However, profiling is still technically challenging due to the low abundance and rapid degradation of the ubiquitin conjugates. To this end, several different strategies have been used to enrich the ubiquitinated proteome or ubiquitinome (23). One well-established method is purifying tagged ubiquitin from yeast (24). Later, this approach has been employed for ubiquitin profiling in mammalian cells (25–27). However, this method is not applicable in animal tissues or pathological specimens, because of the technical challenge of expressing tagged ubiquitin in those samples. In addition, overexpressing ubiquitin in mammalian cells may interfere with normal cellular functions.

High affinity antibodies are a successful alternative approach to enriching ubiquitin conjugates (28–30). However, off-target antibody-interacting proteins or the high background of the antibody itself in liquid chromatography-tandem mass spectrometry (LC-MS/MS) analysis can be difficult to exclude. Xu et al. (31) developed a di-Gly-lysine-specific antibody against the signature di-glycine residues that are the remnants of ubiquitin after trypsin digestion, to directly enrich the ubiquitinated peptides in a human cell line. This method has been improved and is the primary strategy in ubiquitinome research (32–34). However, proteins modified by neural precursor cell-expressed developmentally down-regulated 8 (NEDD8) or interferon-stimulated gene 15 (ISG15) can have identical di-Gly remnants on the modified Lys residues, making it impossible for LC-MS/MS to distinguish among these modifications. In addition, the di-Gly antibody has varied affinity for different epitopes, causing biases for different ubiquitinated peptides during purification (35, 36).

UBDs are small protein module families that can recognize and bind to ubiquitin modifications and are used as alternative tools to enrich ubiquitinated proteomes. To date, more than 20 different UBD families have been identified (5, 37). The affinity of different UBDs to ubiquitin spans a wide range, from 2 to 500 μM (6), with various preferences for different types of ubiquitin linkages (38). For example, the ubiquitin-associated (UBA) domains of hHR23A and MUD1 selectively bind to Lys-48-linked chains (39, 40), whereas the Npl4 zinc finger domain of TGFβ-activated kinase 1 (TAK1)-binding protein 2 prefers Lys-63-linked chains (41), and the UBAN domain of the NF-κB essential modulator (NEMO) specifically binds to linear ubiquitin chains (42, 43). Although single and tandem UBDs have been used to purify ubiquitinated proteins from mammalian cells (44–46), they are usually restricted to well-studied UBA domains (such as UBA from ubiquilin 1 and hHR23A) with ubiquitin chain preference, rather than reflecting the global ubiquitination landscape.

To address the bias in UBDs and efficiently recover the ubiquitinome during purification, we systematically evaluated the affinity of a number of UBDs, and we constructed an artificial tandem hybrid UBD (ThUBD) that exhibited no strong bias toward any of the seven tested polyubiquitin chains. By analyzing complex samples of yeast and mammalian cells with ThUBD, we found a dramatic improvement in binding affinity compared with previously reported UBD methods. Our study demonstrates that ThUBDs are promising tools for the efficient and unbiased analysis of the ubiquitinated proteome, especially for animal tissues or pathological specimens. This approach may aid in the discovery of novel biomarkers or therapeutic strategies against human diseases.

**EXPERIMENTAL PROCEDURES**

**Cloning, Protein Purification, and Immobilization**—The UBDs used in this study were DSK2p-derived UBA (DSK2, 327–371 amino acids), ubiquin 1-derived UBA (UQ1, 541–586 amino acids), ubiquilin 2-derived UBA (UQ2, 575–624 amino acids), RABGEF1-derived A20-ZnF (A20, 9–73 amino acids), and HDAC6-derived ZnF-UBP (HDAC6, 985–1152 amino acids). Five different UBDs were cloned by PCR amplification and inserted into the pGEX-4T-2 vector (GE Healthcare, Chalfont St Giles, UK) using BglII and EcoRI sites. Tandem repeats of UBDs were constructed using BglII and BamHI restriction sites as designed by Hjerpe et al. (47). ThUBDs (DSK2 and UQ2 (termed as UDQ2) and DSK2 and UQ2 (termed as UDQ2)) and ThUBDs (ThUDQ2 and ThUDQ2) were constructed in the same way. All of the proteins were overexpressed in Escherichia coli BL21 (DE3) cells, which were induced with 0.5 mM isopropyl β-D-galactopyranoside for 4 h at 30 °C. The harvested cells were lysed by sonication in lysis buffer (1 mM DTT, 1% Triton X-100 in PBS), GST-UBD fusion proteins were purified from cell lysates using glutathione-Sepharose (GSH) 4B beads (Qiagen, Valencia, CA) according to the manufacturer’s instructions. The purified GST-UBDs were competitively eluted by reduced glutathione and then resuspended in coupling buffer (0.2 M NaHCO3, 0.5 M NaCl, pH 8.3) before being coupled to NHS-activated Sepharose (GE Healthcare, Munich, Germany) following the manufacturer’s instructions. The UBD-conjugated agarose was stored at 4 °C in PBS supplemented with 30% glycerol.

**Yeast Strains and Affinity Purification**—The yeast strain SUB592 (48) was grown at 30 °C in yeast extract peptone dextrose (YPD) medium (1% yeast extract, 2% Bacto-peptone, 2% dextrose) and harvested in the early log phase (A600 = 1.0). His-Myc-tagged ubiquitin conjugates were purified from yeast under denaturing conditions with Ni-NTA agarose (24) or under native conditions by UBDs (18). For native conditions, cells were lysed with glass beads in 200 μl of buffer A (50 mM Na2HPO4, pH 8.0, 500 mM NaCl, 0.01% SDS, 5% glycerol). Protein extracts were centrifuged at 70,000 × g for 30 min and then incubated with immobilized GST-UBD beads at 4 °C for 30 min. After incubation, the beads were washed with buffer A and then buffer B (50 mM NH4HCO3 and 5 mM iodoacetamide), followed by 50 mM NH4HCO3 to remove iodoacetamide. Bound ubiquitin conjugates were eluted by boiling in 1× SDS-PAGE loading buffer (50 mM Tris-
Cell Culture and Affinity Purification—Human hepatocellular carcinoma (HCC) MHCC97-H cells (49) were cultured to confluence before harvest. Ubiquitin conjugates were purified from cells under native conditions by ThUBDs (ThUDA20) as described above (45). Finally, the purified ubiquitin conjugates were released by boiling in 1× SDS-PAGE loading buffer.

SDS-PAGE and Western Blot Analysis—Affinity-purified proteins from yeast cells were separated on 10% SDS-polyacrylamide gels, transferred onto nitrocellulose membranes (Bio-Rad), and then probed with antibodies against Myc (Abcam, Cambridge, UK) or visualized by silver staining.

Sample Preparation for MS Analysis—Yeast or human ubiquitin conjugates were separated on SDS-polyacrylamide gels or underwent off-line high pH reverse phase LC separation (24, 50). The ubiquitin conjugates separated by SDS-PAGE were sliced into different gel pieces based on molecular weight markers and digested with trypsin overnight. The tryptic peptides were extracted with extraction buffer (5% total cell lysates (FA) + 45% acetonitrile (ACN)) and then ACN and were finally dried using a vacuum dryer (Labconco CentriVap, Kansas City, MO).

Peptides were analyzed using an ultra-performance LC-MS/MS platform of hybrid LTQ-Orbitrap Velos mass spectrometer (Thermo Fisher Scientific, San Jose, CA) equipped with a Waters nano-platform of hybrid LTQ-Orbitrap Velos mass spectrometer (Thermo Fisher). The LC separation was performed on an in-house packed capillary column (75 μm inner diameter × 15 cm) with 3-μm C18 reverse-phase fused silica (Michrom Bioresources, Inc., Auburn, CA). Then the sample was eluted with a 60–140-min nonlinear gradient ramped from 8 to 40% of mobile phase B (phase B, 0.1% FA in ACN; phase A, 0.1% FA + 2% ACN in water) at a flow rate of 0.3 μl/min. Eluting peptides were analyzed using an LTQ-Orbitrap Velos mass spectrometer. The MS1 was analyzed with a mass range of 300–1600 at a resolution of 30,000 at m/z 400. The automatic gain control was set as 1×106, and the maximum injection time was 150 ms. The MS2 was analyzed in data-dependent mode for the 20 most intense ions subjected to fragmentation in the linear ion trap (LTQ). For each scan, the dynamic range was set at 45–60 s to suppress repeated detection of the same ion peaks.

Data Analysis—The raw data files were searched by the Sorcerer™-SEQUEST™ (version 4.0.4 build, Sage-N Research, Inc.) and SEQUEST HT algorithms embedded in Proteome Discovery (version 1.4.1.14, Thermo Fisher) against the Swiss-Prot reviewed database (version released in 2013.10 for yeast, contains 6652 entires; version 2014.04 for human, contains 20,266 entries). MS search parameters consisted of full tryptic restriction, fixed modification of cysteine carbamidomethylation, methionine oxidation, and di-Gly-lysine as variable modifications. Peptides were allowed up to three cleavage sites. Precursor mass tolerance was set at 20 ppm and that of the MS2 fragments was set at 0.5 Da. Peptide matches were filtered by a minimum length of seven amino acids. For ubiquitination identification, a re-searching strategy was used for protein identification; di-Gly-lysine was added in the second search. The C-terminal di-Gly-lysine site identifications were removed. The peptides and proteins were filtered until a false discovery rate lower than 1% was estimated using a target-decoy search strategy.

Targeted Quantitative MS of Polyubiquitin Linkages Using the Stable Isotope Labeling by Amino Acids in Cell Culture-Absolute Quantification (SILAC-AQUA) Approach—To accurately quantify the abundance of the seven polyubiquitin chains from each ubiquitin conjugate purified with different UBDs, ubiquitin conjugates labeled with heavy isotope 13C6-lysine (Lys-6) and 15N4-lysine (Arg-10) were purified with nickel beads as a quantification standard to mix equally with each sample (18). The ubiquitin conjugates were digested as described above, and the resulting peptides were dissolved in a sample buffer (1% FA, 1% ACN) and analyzed by LC-MS/MS. Eluted peptides were detected on the Orbitrap mass spectrometer in a survey scan (300–1600 m/z, resolution 30,000) followed by selective reaction monitoring scans for seven ubiquitin chains ions in the LTQ. The peptide intensity for quantification was manually analyzed by ion chromatograms using Xcalibur version 2.0 software (Thermo Finnigan, San Jose, CA).

Determination of the Amount of Immobilized UBDs Using the SILAC-AQUA Approach—To accurately compare the binding affinity to ubiquitin of different UBDs, the same amount of each UBD was used. We selected two typical peptides (supplemental Fig. S1) from glutathione S-transferase (GST) protein as internal standards for quantifying the amount of GST-UBDs coupled to agarose beads. The GST protein was overexpressed and completely labeled with [13C6]lysine (+6.0201 Da) using the SILAC-labeled E. coli BL21 (DE3) cells and was then purified using GSH 4B beads. The amount of purified heavy-labeled GST was measured using the BCA protein assay kit (Thermo Scientific, Rockford, IL) and Coomassie staining (51). A total of 10 μg of heavy-labeled GST was digested with Lys-C, and the resulting peptides were used as quantitative standards. To quantify the GST-UBD proteins, on-bead digestion with Lys-C was performed to release peptides from GST-UBDs coupled to agarose beads. The resulting peptides were desalted by passing through a C18 microcolumn as reported previously (52), and then mixed with the heavy-labeled and Lys-C-digested GST peptides followed by nano-LC-MS/MS analysis. The detailed quantification method was described as reported previously (53). According to the light and heavy intensities of the same peptides from GST proteins, we calculated the amount of light GST in samples, which represented the amount of GST-UBD proteins.

Surface Plasmon Resonance Imaging Measurements and Analysis—The surface plasmon resonance imaging binding measurements were performed on the PlexArray® HT system (Plexera LLC, Washington, WA). Experiments were performed in PBS buffer at 25 °C. Purified GST-UBD proteins were resuspended in PBS buffer and then immobilized onto NanoCapture Gold™ SPR-active slides following the standard amine coupling chemistry, as described previously (54). Then different concentrations (10, 5, 2.5, 1.25, and 0.625 μg/μl) of Lys-48-linked tetra-ubiquitin (Boston Biochem, Cambridge, MA) were injected for a flow rate of 2 μl/s for 300 s. All of the injections were performed as randomized duplicates, and the entire experiment was repeated three times. The binding level was tracked by measuring the change in response units, according to the different injection concentrations. Kd values were calculated using BIAevaluation software.

Virtual Western Blot—With the extracted ion chromatograms of each protein in different bands and the experimental molecular weight of each band, we reconstructed the experimental lanes using an in-house-developed MATLAB script (55). The reconstructed images revealed the different purification methods with regard to molecular weight and abundance.

RESULTS

hUBDs Showed Improved Binding Affinity to Ubiquitinated Proteins—UBDs, which directly interact with ubiquitin, have been utilized for purifying ubiquitinated proteins (44–47). Certain UBA domains from DSK2 and UQ1 are preferred to enrich ubiquitinated proteins, because they appear to bind equally to different types of ubiquitin linkages (39). However, these results were obtained by in vitro pulldown assays with radiolabeled Ub4 linked through different lysine residues, lacking
Novel Approach for Ubiquitinome Research Using ThUBD

| Name        | Source                        | Type     | Binding affinity $K_D$ μM |
|-------------|-------------------------------|----------|-------------------------|
| DSK2        | *Saccharomyces cerevisiae* Dsk2p (DSK2) | UBA      | 14.8 ± 5.3             |
| UQ1         | *Homo sapiens* ubiquitin 1 (UBQLN1) | UBA      | 20          |
| UQ2         | *H. sapiens* ubiquitin 2 (UBQLN2) | UBA      | ND         |
| A20         | *H. sapiens* RAB guanine nucleotide exchange factor (GEF 1 (RABGEF1)) | ZnF-A20  | 12–22       |
| HDAC6       | Mouse (HDAC6)                 | ZnF-UBP  | 0.06       |
| UDQ2        | This study                    |          |             |
| UDA20       | This study                    |          |             |

* See Ref. 61.  
+ See Ref. 62.  
& See Ref. 63.  
* See Ref. 60.  
& See Ref. 64.  
+ See Ref. 65.

Further systematic verification. In this study, we quantitatively compared the binding characteristics of seven UBDs (Table I and supplemental Table 1) to endogenous ubiquitinated proteins in the cell lysates by Western blotting and LC-MS/MS.

We first compared the relative binding efficiency of seven UBDs to ubiquitinated proteins (Fig. 1A). Purified GST-UBDs (Fig. 1B) were coupled to NHS beads as the bait. The amount of GST-UBDs coupled to NHS beads was analyzed by SILAC-AQUA (supplemental Fig. S1). Equal amounts of cell lysates from SUB592 were incubated with the same molar concentrations of GST-UBDs for affinity experiments. Purified ubiquitin conjugates by different UBDs were split into two portions, and 5% (v/v) ubiquitin conjugates were resolved by SDS-PAGE for Western blot analysis. The rest of the ubiquitin conjugates were mixed with equal amounts of the respective heavy isotope-labeled ubiquitin conjugates purified with Ni-NTA under denaturing conditions, followed by separation on SDS-polyacrylamide gels, digestion with trypsin, and analysis by LC-MS/MS. Compared with the A20_mut, which does not bind to ubiquitin (56), all seven UBDs pulled down ubiquitinated proteins from total cell lysates. However, different amounts of ubiquitinated proteins were captured by the UBDs (Fig. 1D), suggesting that different UBDs have diverse binding efficiencies to ubiquitinated proteins. Among these, A20, DSK2, UQ2, and two hUBDs (UDQ2 and UDA20) enriched more ubiquitinated proteins than the other UBDs. Two hUBDs, especially UDA20, enriched the most ubiquitinated proteins, suggesting that a combination of different UBDs can significantly increase the binding efficiency to ubiquitinated proteins.

Next, we investigated the binding affinity of the UBDs to seven ubiquitin linkages. The heavy isotope-labeled ubiquitinated proteins purified by Ni-NTA-agarose were used as a reference (Fig. 1A), representing the proportion of different polyubiquitin linkages in yeast cells. The relative amounts of seven polyubiquitin chains purified by different UBDs were quantitatively measured with Ub-AQUA (18, 55). As a result, Lys-6-, Lys-11-, Lys-27-, Lys-29-, Lys-33-, Lys-48-, and Lys-63-linked polyubiquitin chains were all detected from the proteins pulled down by the UBDs (Fig. 1E), demonstrating the efficient enrichment of ubiquitinated proteins. In addition, LC-MS/MS analyses further confirmed that different UBDs exhibit diverse interaction properties with polyubiquitin chains. For instance, the A20 domain preferred Lys-63-linked chains and UQ1 purified more Lys-11-linked chains, whereas the hUBD (especially UDA20) apparently had a higher binding affinity to all seven polyubiquitin chains. These results are not completely consistent with previous studies using either SPR analyses or pulldown assays (39). These differences may be due to binding affinity that was normalized based on the proportion of different polyubiquitin linkages purified by Ni-NTA-agarose under denaturing conditions. It also provides a new way to systematically analyze the binding efficiency of UBDs to polyubiquitin chains. These data suggest that hUBDs not only show increased binding affinity to endogenous ubiquitinated proteins but also have no bias toward different polyubiquitin chains. Additional studies are needed to fully understand the binding mechanism of hUBDs to ubiquitin and different linkages of ubiquitin chains.

ThUBDs Have Superior Binding to Ubiquitinated Proteins—To achieve higher binding affinity to ubiquitin (45, 47), ThUBDs were also constructed (Fig. 2A). Five UBDs, including A20, DSK2, UQ2, and two hUBDs (UDQ2 and UDA20), were selected based on their higher binding affinities to ubiquitin (Fig. 1D). Purified ThUBDs (Fig. 2B) were coupled to NHS beads and quantified by LC-MS/MS as described previously. All five TUBDs efficiently purified ubiquitinated proteins from total cell lysates (Fig. 2C). The different ubiquitination protein patterns of TUBDs suggest that these UBDs may have specific preferences for ubiquitinated substrates. Furthermore, two ThUBDs (4_UDQ2 and 4_UDA20) also enriched more ubiquitinated proteins than the other three TUBDs, as dem-
onstrated in our pulldown experiments. This was especially the case for proteins with higher molecular masses (>50 kDa) (Fig. 2, C and D).

We also compared the binding affinity of TUBDs to seven polyubiquitin chains. By AQUA analysis, we found that the relative binding affinity of TUBDs to different polyubiquitin chains was significantly higher than that of single UBDs (Fig. 2E), in accordance with data from previous reports (45, 47). In addition, most TUBDs possess diverse binding selectivity to different ubiquitin linkages. Particularly, two ThUBDs (4_UDQ2 and 4_UDA20) showed no obvious bias to seven polyubiquitin chains. Hence, they were selected for further characterization.

**Binding Affinity of the Dimer of ThUBDs Reaches to Saturation**—To optimize UBDs for the ubiquitinated proteomics study, we investigated the effects of the tandem units on ubiquitin binding affinity. We constructed the respective mono-, di-, and tetra-ThUBDs (Fig. 3A), and we found that the di-ThUBDs purified more ubiquitinated proteins than the mono-ThUBDs (Fig. 3, B and C), suggesting that repeats of...
the ThUBD monomer can increase ubiquitin binding affinity. However, the purified tetra-ThUBDs had lower solubility and stability, due to their high molecular weight, which might explain why there was no apparent increase in ubiquitin binding affinity to tetra-ThUBDs compared with di-ThUBDs (Fig 3, B and C). In addition, the di-ThUBDs, especially 2_UDA20 repeats, showed no apparent bias to all seven polyubiquitin chains (Fig. 3D), further demonstrating the superiority of this hybrid UBD on the ubiquitinome study.

To further validate the pulldown results, quantitative SPR analysis was applied on mono-, di-, and tetra-UDA20. The di-ThUBD showed the highest binding affinity to Lys-48-Ub4 (Table II), in agreement with the Ub-AQUA results (Fig. 3D). The binding affinity of 2_UDA20 to Lys-48-Ub4 was also better than the other three tandem UBDs (Table II), including ubiquitin 1 TUBE (47), HR23A TUBE (47), and qUBA (45), which have been successfully used to purify ubiquitinated proteins in many cell lines. These results suggest that 2_UDA20 might be an optimal tool for the global ubiquitinated proteome research.

**ThUBD (2_UDA20) Shows Superior Binding to Ubiquitinated Proteins than qUBA**—To evaluate the effectiveness of ThUBD, we compared the relative binding efficiency of 2_UDA20 and qUBA to ubiquitinated proteins from yeast cell lysates (Fig. 2E). The binding affinity of ubiquitin chains to the hybrid UBD was quantified using the SILAC-AQUA strategy (33). The results showed that 2_UDA20 bound to ubiquitinated proteins with higher efficiency than qUBA, indicating that 2_UDA20 is a more effective tool for the global ubiquitinated proteome research.
lysates (Fig. 4A). After being resolved on SDS-polyacrylamide gels, proteins captured by ThUBD and qUBA exhibited similar patterns (Fig. 4B), suggesting that both UBDs are capable of efficiently pulling down ubiquitinated proteins from yeast cells. Interestingly, we also noticed that the amount of ubiquitinated proteins enriched by the ThUBD was more than the amount for qUBA, even for proteins in the high molecular weight range. These results strongly support the fact that the ThUBD binds more efficiently to ubiquitinated proteins than the qUBA.

These results were also corroborated by LC-MS/MS analysis (Fig. 4C) and quantitative SPR (Table II). In the high molecular mass region (>50 kDa), the superiority of ThUBD binding to ubiquitin was more evident (Fig. 4C). Protein ubiquitination, especially polyubiquitination, can shift a protein’s molecular weight to a higher molecular mass region, indicating that ThUBD prefers binding to polyubiquitinated proteins. In fraction 18, which was under 20 kDa and mainly represented the monomer and dimer ubiquitin regions, we found extremely large amounts of ubiquitin purified by ThUBD rather than qUBA.

**TABLE II**

| Name       | Binding affinity Kd (μM) | Source |
|------------|-------------------------|--------|
| UDA20      | 8.09                    | This study |
| 2_UDA20    | 4.46                    | This study |
| 4_UDA20    | 6.41                    | This study |
| qUBA       | 6.8                     | This study |
| Ubiquilin 1 TUBE | 8.94              | 47     |
| HR23A TUBE | 6.86                    | 47     |

*Normalized based on heavy isotope-labeled Ub conjugates purified by Ni-NTA in denature condition; Top two in each column were highlighted.*
than by qUBA. Within this fraction, we extracted the spectra of ubiquitin and found that almost 50% of the spectra purified by ThUBD contained ubiquitin-modified sites, suggesting that most of these ubiquitins were derived from polyubiquitinated rather than monoubiquitinated proteins. However, the opposite results were shown in fraction 18, which was purified by qUBA (supplemental Fig. S2A). Together, these results indicate that ThUBD captures more ubiquitinated proteins, especially polyubiquitinated proteins, than qUBA from yeast cell lysates.

Through LC-MS/MS analysis, we identified 3837 and 3594 proteins, including 362 and 260 ubiquitinated proteins with ubiquitin-modified sites purified by ThUBD and qUBA (Fig. 4D), respectively. Based on the virtual Western blot strategy for validating ubiquitinated proteins as described previously (55), we further screened out 905 and 538 proteins with a 10-kDa increase in molecular mass (supplemental Fig. S2, B and C). Combining proteins with ubiquitin-modified sites and molecular weight increases, we identified 1092 and 690 proteins purified by ThUBD and qUBA as reliable ubiquitinated proteins (supplemental Tables 2 and 3). The number of overlapped ubiquitinated proteins was 509, representing 46.5% and 73.8% of the identified ubiquitinated proteins purified by ThUBD and qUBA, respectively (supplemental Fig. S2D). Not
only was the number of ubiquitinated proteins purified by ThUBD higher than that purified by qUBA, the relative amounts of these proteins enriched by ThUBD were also significantly enhanced (Fig. 4E). The same trend was also shown for the ubiquitinated proteins identified from samples enriched by ThUBD and qUBA (supplemental Fig. S2E), consistent with our gel imaging data.

Compared with qUBA, a known purification material that is also based on the binding affinity of UBD to ubiquitin, ThUBD purified more ubiquitinated proteins in the large scale purification from yeast cells, demonstrating that ThUBD is a promising purification reagent for global ubiquitinated proteomic research.

Comparison of Enrichment Efficiency between the Optimized ThUBD (2_UDA20) and Ni-NTA—To further evaluate the efficiency of ThUBD as a reagent for global ubiquitinated proteomic research, we also compared it with the Ni-NTA affinity purification method. Ubiquitinated proteins were purified by ThUBD under native conditions or by Ni-NTA under denaturing conditions from the JMP024 strain, which expresses His-Myc-tagged ubiquitin (24), and were then digested by trypsin after separation by SDS-PAGE, followed by LC-MS/MS analysis. Both reagents successfully purified the ubiquitinated proteins and displayed an apparent smear pattern in the high molecular mass region, compared with total cell lysates (Fig 5A). ThUBD purified even more ubiquitinated proteins than Ni-NTA, indicating its higher efficiency. Quantification of peptides representing ubiquitin ions (Ser-57, Lys-48, and Lys-63) in gel fraction 2 (Fig. 5A) also showed that significantly more ubiquitin was captured by ThUBD than by Ni-NTA, consistent with our gel imaging results. The same trend was shown in the gel fractions in the high molecular mass region (>50 kDa) (Fig. 5C), suggesting that ThUBD prefers binding to polyubiquitinated proteins. However, in the low molecular mass region, especially in the 10-kDa region (which often represents the ubiquitin pool), more ubiquitin was captured by Ni-NTA than by ThUBD. The overall higher enrichment of the larger molecular weight ubiquitinated proteins could be attributed to the protection of ThUBDs for polyubiquitinated proteins from deubiquitinase or proteasomal degradation (47), showing the superiority of ThUBD in ubiquitinated protein purification.

Based on the same criteria used in ThUBD, we finally identified 710 ubiquitinated proteins, including 268 ubiquitinated proteins with ubiquitin-modified sites purified by Ni-NTA under denaturing conditions (supplemental Tables 2 and 3). The shared number of ubiquitinated proteins with ubiquitin-modified sites was 118, representing 32.6 and 44% of the identified ubiquitinated proteins purified by ThUBD and Ni-NTA, respectively. By reconstructing virtual Western blots for ZEO1 (YOL109W), which was identified by ThUBD and Ni-NTA as ubiquitinated proteins (Fig. 5E), we found that ZEO1 was modified by ubiquitin, indicating the reliability of identification. However, the profile of ZEO1 purified by ThUBD and Ni-NTA was quite different. The ZEO1 protein purified by ThUBD showed a distinct molecular weight increase to the high molecular mass region, suggesting polyubiquitination. These results further confirmed the preference of ThUBD to polyubiquitinated proteins.

The total of 1092 candidate ubiquitinated proteins identified by ThUBD represents one of the largest datasets publicly available for yeast ubiquitinome study in a single experiment, compared with the 710 proteins identified by Ni-NTA (this study), 1075 proteins identified by Ni-NTA without filtration (24), and 1108 proteins identified by di-Gly antibody in yeast (57). The overlap of ubiquitinated proteins identified in these four datasets increased reliability of the identified ubiquitinated proteins (supplemental Fig. S3A). Among the overlapped proteins, about 39.7% (434/1092) of the ubiquitinated proteins were uniquely identified by ThUBD, which suggests the high binding efficiency of ThUBD in the purification of ubiquitinated proteins. Among the identified ubiquitinated proteins with ubiquitin-modified sites, there were shared proteins across all four datasets (Fig. 5D). More than 64.4% (233/362) of identified proteins by ThUBD are novel ubiquitinated proteins that are not identified by Ni-NTA. These results imply that a potentially large pool of ubiquitinated proteins remains to be explored.

The relative amount of ubiquitinated proteins enriched by ThUBD was also significantly enhanced, compared with that purified by Ni-NTA (supplemental Fig. S3B). Together, these results demonstrate that ThUBD can efficiently capture ubiquitinated proteins, especially polyubiquitinated proteins, under native conditions.

Large Scale Identification of the Endogenous Ubiquitinated Proteins in MHCC97-H Cells Using ThUBD (2_UDA20)—MHCC97-H cells are a highly metastatic HCC cell line that has high invasion ability (49). To investigate possible misregulation in the ubiquitination system of MHCC97-H cells, we performed global ubiquitinome analysis in these cells based on ThUBD. As expected, ubiquitinated proteins from MHCC97-H cells were successfully purified by ThUBD (Fig. 6A) and displayed the characteristic smear pattern on the SDS gel. Quantification of peptides representing ubiquitin ions (Ser-57, Lys-48, and Lys-63) in gel fraction 3 showed an apparent increase in the samples purified by ThUBD, compared with that in total cell lysates (Fig. 6B). Additionally, a significant portion of proteins had a clear shift toward higher molecular weight regions in purified ubiquitinated proteins, compared with that in total cell lysates (Fig. 6C), both of which demonstrated the effectiveness of using ThUBD to purify ubiquitinated proteins.

We identified 1663 credible ubiquitinated proteins from MHCC97-H cells based on the same criteria used in yeast, which was a greater number than the ubiquitinated proteins identified by other UBDs (Fig. 6D) (supplemental Tables 4 and 5) (45, 46). However, only 19.2% (319/1663) of the identified proteins had ubiquitin-modified sites, which was
FIG. 5. Comparison of the enriched ubiquitinated proteins from yeast total cell lysates by ThUBD and Ni-NTA beads under denaturing condition (Ni_denature).

A, comparison of ubiquitinated proteins purified with ThUBD and Ni-NTA beads under denaturing condition by SDS-PAGE. Ubiquitin conjugates were purified from the same amount of yeast total cell lysates, resolved on a 10% SDS-polyacrylamide gel, stained with Coomassie Blue, excised into 24 gel bands, digested by trypsin, and analyzed by LC-MS/MS. B, base-peak chromatogram of three MS runs (upper) and elution profiles of peptides Ser-57, Lys-48 (LIFAGK QLEDGR), and Lys-63 (TLSDYNIQK ESTLHLVLR) (bottom) resulting from trypsin digestion of ubiquitin or ubiquitin chains for the samples of the gel band 2 on A for total cell lysates (TCL), Ni_denature, and ThUBD, respectively. Peptides derived from the gel band 2 were loaded on the column and then eluted with a 2–30% gradient of buffer B over 50 min. The peak area (AA) corresponds to the relative ion intensity of the analyzed peptide. C, comparison of ubiquitin in total cell lysates (TCL) or purified by ThUBD and Ni-NTA under denaturing condition in each gel band, reflecting quantitative data in all gel bands in the total cell lysates or ubiquitin conjugates. The ubiquitin abundance was represented by the intensity signal on MS of Ser-57 peptide extracted.
lower than we expected. The current gel separation method might be responsible for the low identification of ubiquitin-modified sites, because the proteins were separated on SDS-polyacrylamide gel by molecular weight, and polyubiquitinated proteins may disperse in different molecular weight regions (Fig. 7A), resulting in a lower abundance in each gel fraction.

To further improve the identification of ubiquitinated proteins by mass spectra, we introduced an off-line high pH RP-LC separation strategy. Purified ubiquitinated proteins by ThUBD from MHCC97-H cells were digested with trypsin, and separated by high pH RP-LC (supplemental Fig. S4A). We collected 20 fractions by high pH RP-LC in a 60-min gradient (supplemental Fig. S4B), after which each fraction was analyzed by MS. One advantage of off-line RP-LC separation is that peptides are separated based on physicochemical properties, and hence the peptides from high abundance proteins (such as ubiquitin) will be aggregated in the same fraction (Fig. 7A). This prevents signals of the targeted GG peptides from being diluted and suppressed by adjacent highly abundant ions in LC-MS/MS analysis. Indeed, we found that proteins showed different distribution patterns in high pH RP-LC fractions, compared with that observed by gel separation. Taking ubiquitin as an example, it dispersed with moderate intensity in all gel fractions, while mainly being concentrated in one fraction with extremely high intensity in high pH RP-LC fractions (Fig. 7B). This also held true for the ubiquitin-modified peptide (ITITNDK ↓ GR) from HSPA8 (Fig. 7C), demonstrating the advantage of the high pH RP-LC separation strategy in protein identification, especially for low abundance proteins.

A total number of 7345 candidate ubiquitinated proteins were identified by the high pH RP-LC separation method, including 939 ubiquitinated proteins with ubiquitin-modified sites, which is more than that identified by gel separation (supplemental Tables 4 and 5). Up to 91.5% (1521/1663) of the identified ubiquitinated proteins by gel separation were also identified by high pH RP-LC separation (Fig. 7D), indicating the reliability of the identified candidate ubiquitinated proteins. A total of 133 identified ubiquitinated proteins with ubiquitin-modified sites was shared by the two methods, which means that more than 85.8% (806/939) of the identified proteins with ubiquitin-modified sites by the high pH RP-LC separation method were uniquely identified (Fig. 7E). These results further demonstrate the advantage of the high pH RP-LC separation method for identifying ubiquitinated proteins and ubiquitination sites. By comparing our dataset with previously published datasets that used di-Gly antibodies (32, 33), the number of shared proteins for all three datasets was 2294 (supplemental Fig. S4C), which only represents 31.2% of those identified from this study, 51.9% of identified proteins by Kim et al. (32), and 55.8% of identified proteins by Wagner et al. (33), suggesting the high quality of identification performed in the high pH RP-LC separation method. Several previously validated ubiquitinated proteins were also identified in our study, including RH0A (58) and EPN1 (supplemental Fig. S5, A and B) (59).

A functional annotation of ubiquitinated proteins with ubiquitin-modified sites, identified from the ubiquitinated proteins enriched with ThUBD (combined gel separation and high pH RP-LC separation methods), was assigned by Ingenuity pathway analysis. The identified proteins involved a large variety of biological processes including but not limited to protein translation, ubiquitination, localization, cell cycle, and apoptosis as reflected in biological processes (supplemental Fig. SSC) and signaling pathway analyses (Fig. 7F). Ingenuity pathway analysis integrates all of the identified proteins into 63 canonical pathways (p < 0.01), of which the top 15 pathways are shown in Fig. 7F. The canonical pathways describing the translational process as well as post-translational modification such as EIF2 signaling, mechanistic target of rapamycin signaling, and ubiquitination pathway are significantly enriched. In agreement with the high metastatic ability characteristic of MHCC97-H cells (49), a significant proportion of captured proteins was involved in adherents junctions and actin cytoskeleton signaling, suggesting that ubiquitination may play important roles in a metastasis-related process.

Ubiquitination is a conserved post-translational modification from yeast to mammalian cells. To examine the properties of amino acids surrounding ubiquitination sites, the frequencies of neighboring amino acid residues were analyzed based on the enriched ubiquitination peptides from 97H and yeast cells, respectively. We observed an obvious preference for Leu and Glu adjacent to ubiquitination sites both in mammalian and yeast cells (Fig. 7G). However, in contrast to phosphorylation motifs, we found no apparent sequence motif (supplemental Fig. S6), suggesting that the ThUBD enrichment technique used here has no bias to particular sequences surrounding the ubiquitination sites. In addition, based on the homologous search, we found 305 pairs of conserved proteins enriched from mammalian cells and yeast cells (supplemental Table 6). GO annotation (Fig. 7H) showed that these conserved proteins are involved in almost identical pathways in both mammalian and yeast cells, suggest-
Fig. 6. Large scale profiling of ubiquitinated proteins enriched by ThUBD from liver cancer cell MHCC97-H. A, comparison of MHCC97-H total cell lysates (TCL) and ubiquitinated proteins purified by ThUBD displayed on SDS-PAGE. Both samples are resolved on a 10% SDS-polyacrylamide gel, stained with Coomassie Blue, excised into 18 gel bands, digested by trypsin, and analyzed by LC-MS/MS. B, base-peak chromatogram of two MS runs (upper) and elution profiles of peptides Ser-57, Lys-48, and Lys-63 (bottom) resulting from trypsin digestion of ubiquitin or ubiquitin chains for the samples of the gel band 3 on A for total cell lysates and ubiquitinated proteins (ubc), respectively. Peptides derived from the gel band 3 were loaded on the column and then eluted in a 2–30% gradient of buffer B over 50 min. The peak area (AA) corresponds to the relative ion intensity of the analyzed peptide. C, \( \Delta MW \) (the molecular weight difference between the experimental and theoretical one) histogram of the proteins identified in the total cell lysates (n = 4518) and ubiquitinated proteins (ubc) (n = 2551). The dataset was produced through SDS-PAGE separation and then analyzed by LC-MS/MS. D, comparison of ubiquitinated proteins purified by ThUBD, qUBA, and TUBE from mammalian cells.

|                         | Purified ubiquitinated proteins candidate | Ubiquitination sites | Ubiquitinated proteins with ubiquitin modified sites | Ubiquitinated proteins with \( \Delta MW = 10 \text{kDa} \) | Combined ubiquitinated proteins | Reference |
|-------------------------|------------------------------------------|----------------------|----------------------------------------------------|-------------------------------------------------|---------------------------------|-----------|
| ThUBD                  | 3411                                     | 458                  | 319                                                | 1522                                            | 1663                            | This study |
| qUBA                   | –                                        | 294                  | 223                                                | –                                               | –                               | Ref [42]  |
| TUBE                   | 643                                      | –                    | –                                                  | –                                               | –                               | Ref [43]  |
ing that these pathways are highly conserved and ubiquitinated across different species. These results further demonstrated the conservation of ubiquitination in different species, from essential amino acid sequences to intricate pathway levels.

DISCUSSION
One of the challenges of ubiquitinome research is the enrichment of ubiquitinated proteins, which are in low abundance in cells and prone to rapid degradation by the 26S proteasome. In this study, we developed a recombination of...
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hybrid UBDs to effectively purify ubiquitinated proteins with no apparent biases under native conditions. Using ThUBD (2_UDA20), we identified one of the largest ubiquitinated protein datasets from yeast and mammalian cells in a single experiment to date.

In this study, we systematically compared the binding characteristics of different UBDs to ubiquitin and seven polyubiquitin chains. We found that different UBDs showed diverse binding affinity to ubiquitin and linkages, such as A20 appears to prefer Lys-63-linked chains, and UQ1 specificity binds to Lys-11-linked chains. Among the seven evaluated UBDs, DSK2 and A20 showed the highest binding affinity to ubiquitin (Fig. 1C). This is also corroborated by the calculated equilibrium dissociation constants (K_D) (Table I). To the best of our knowledge, this is the first study to identify the linkage specificity of UBDs by LC-MS/MS. This approach provides a new method to systematically evaluate the linkage specificity of UBDs under natural conditions.

In cells, many proteins carry multiple copies of UBDs or several types of UBDs. For example, protein RAD23 in yeast contains UBL (ubiquitin-like) and UBA domains at the N and C termini, respectively. It is thought that the increase of UBD copies and types in one protein might contribute to the high affinity of ubiquitin. Based on this assumption, we designed and artificially constructed hybrid UBDs, comprising two different UBDs, to increase the binding affinity to ubiquitin. The top three UBDs that showed high binding affinity to ubiquitin were chosen to construct two hybrid UBDs (i.e. UDA20 and UQD2) (Fig. 1C). We found that hUBDs not only increased binding affinity to ubiquitin but also to the seven polyubiquitin chains by quantitative comparison. This might due to the different binding mechanism of UBDs to ubiquitin and linkages. For example, most UBAs contact the Ile-44-containing surface of ubiquitin (6), although A20 in Rabex-5 binds to a polar region centered on Asp-58 of ubiquitin (60). Therefore, the recombined hUBDs may utilize both binding sites to increase the binding affinity to ubiquitinated proteins. More importantly, this structure may reduce the bias to polyubiquitin chains, showing the effective binding to all seven polyubiquitin chains. With the increasing discovery of novel UBDs, the combination of these UBDs may provide better hUBDs with higher binding affinity to ubiquitin and polyubiquitin chains.

Compared with qUBA and Ni-NTA, which are two powerful approaches for the purification of ubiquitinated proteins, the ThUBD (2_UDA20) captured the largest number of candidate ubiquitinated proteins from yeast, especially polyubiquitinated proteins, demonstrating that ThUBD is a promising reagent for the global ubiquitinated proteome research. A20 in Rabex-5 binds to a polar region centered on Asp-58 of ubiquitin (60). Therefore, the recombined hUBDs may utilize both binding sites to increase the binding affinity to ubiquitinated proteins. More importantly, this structure may reduce the bias to polyubiquitin chains, showing the effective binding to all seven polyubiquitin chains. With the increasing discovery of novel UBDs, the combination of these UBDs may provide better hUBDs with higher binding affinity to ubiquitin and polyubiquitin chains.

In conclusion, using ThUBD, we identified the largest ubiquitinated protein dataset of mammalian cells, demonstrating that ThUBD is a promising reagent for the global ubiquitinated proteomic research.

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