Activation and selectivity of OTUB-1 and OTUB-2 deubiquitylases

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Abstract

The ovarian tumor domain (OTU) deubiquitinylating cysteine proteases OTU ubiquitin aldehyde binding 1 (OTUB1) and OTUB2 are representative members of the OTU subfamily of deubiquitinylases. Deubiquitinylation critically regulates a multitude of important cellular processes, such as apoptosis, cell signaling, and growth. Moreover, elevated OTUB expression has been observed in various cancers, including glioma, endometrial cancer, ovarian cancer, and breast cancer. Here, using molecular dynamics simulation approaches, we found that both OTUB1 and OTUB2 display a catalytic triad characteristic of proteases, but differ in their configuration and protonation states. The OTUB1 protein had a pre-arranged catalytic site, with strong electrostatic interactions between the active-site residues His-265 and Asp-267. In OTUB2, however, the arrangement of the catalytic triad was different. In the absence of ubiquitin, the neutral states of the catalytic-site residues in OTUB2 were more stable, resulting in larger distances between these residues. Only upon ubiquitin binding, did the catalytic triad in OTUB2 re-arrange and bring the active site into a catalytically feasible state. An analysis of water access channels revealed only a few diffusion trajectories for the catalytically active form of OTUB1, whereas in OTUB2 the catalytic site was solvent accessible, and a larger number of water molecules reached and left the binding pocket. Interestingly, in OTUB2, the catalytic residues His-224 and Asn-226 formed a stable hydrogen bond. We propose that the observed differences in activation kinetics, protonation states, water channels, and active-site accessibility between OTUB1 and OTUB2 may be relevant for the selective design of OTU inhibitors.

Introduction

Ubiquitinylination is a posttranslational modification and involves one or more covalent additions to the lysine residues of the target protein which is mediated by sequential action of ubiquitin ligases E1 (ubiquitin-activating enzymes), E2 (ubiquitin-conjugating enzymes) and E3 (ubiquitin ligases). It can be reversed by removal of ubiquitin (deubiquitinylation) catalyzed by ubiquitin hydrolases or deubiquitinylating enzymes (DUBs) (1). DUBs play an essential role in the regulation of cellular processes like apoptosis, cell growth, cell cycle control and DNA repair (2,3).102 DUBs were identified and can be grouped into five subfamilies (4,5). The 18 OTUs identified in human can further be grouped into four categories as OTUBs, OTUDs, A20s and OTULINs (6,7). Cysteine proteases, such as ovarian tumor deubiquitinylases Otubain-1 (OTUB1) and Otubain-2 (OTUB2) belong to the Ovarian Tumor protease (OTU) subfamily of DUB proteins (8) and were the first two OTU proteins to show DUB activity (8,9).

Elevated OTUB1 expression is associated with glioma (10) esophageal squamous cell carcinoma metastasis (11), endometrial cancer (12), hepatocellular carcinoma (HCC) (13), ovarian cancer (14), colorectal cancer (15,16), breast cancer (17), liver cancer (13) and gastric cancer (14). A recent study by Li et al. (18) have identified the role of OTUB2 and its overexpression in non-small cell lung cancer (NSCLC) tissues. Several studies have identified the therapeutic potential of DUBs for cancer treatment.

OTUB1 and OTUB2 belong to the cysteine protease class with a sequence identity and similarity of 48% and 70%, respectively. A 37 residue long N-terminal region in OTUB1 is not present in OTUB2. Despite the considerable sequence conservation and structural overlap, there is perceivable substrate specificity, OTUB1 has a slower cleavage kinetics and favor Lys48-linked poly-ubiquitin (19) whereas OTUB2 cleaves differently linked poly-ubiquitin chains.

Cysteine proteases have a Cys-His-Asn/Asp catalytic triad as the active site. When the cysteine sulfur atom is deprotonated, the histidine residue acts a base and becomes positively charged. The thiolate’s high nucleophilicity leads to an attack of the carbon of the substrate peptide bond to produce a tetrahedral intermediate; then the active site histidine acts as a proton donor to release the amino terminal fragment of the substrate (20). The intermediate is further stabilized by hydrogen bonding between the substrate oxanion and a conserved glutamine residue (21). However, this catalytic triad is performing the proteolytic activity, the protonation states (neutral or zwitterionic form) of cysteine and histidine is highly debated (22-25). The third residue (Asp/Asn) of the catalytic triad is required for a correct alignment and polarization of
catalytic histidine. (5) The Asp in the catalytic triad plays an role in the charge relay systems and is involved in the stabilization of the imidazolium intermediate and the positioning of the catalytic site histidine (26). The role of Asn in the catalytic triad is to orient the side chain of the catalytic histidine to its optimum position for various steps of the catalytic mechanism (27).

OTUB1 in absence of ubiquitin (crystal structure Pdb id: 2ZFY) is catalytically incompetent due to long distances between the catalytic triad residues histidine (His265) and cysteine (Cys91) and aspartic acid (Asp267) of 5.5Å and 4.6Å, respectively (19). Access to the active site of OTUB1 is blocked by Glu214 of the α9-α10 loop and forms backbone hydrogen bond with the catalytic Cys91. The surface charge property of the ubiquitin binding site (P’ side) of OTUB1 is different from OTUB2 located within the close proximity of Cys91 sterically restricts active site access. Conformational changes and specificity binding studies (19) show that OTUBs 1 and 2 may adopt two conformational states and display different lysine linkage specificities probably due to structural differences in their catalytic centers. Though OTUB1 and OTUB2 are close homologs (8), OTUB1 is highly specific for K48 cleavage whereas OTUB2 has broad cleavage profile (28). OTUB1 is auto-inhibited in the ubiquitin-free (apo) state and is activated upon ubiquitin binding, whereas the OTUB2 is catalytically active even in absence of ubiquitin (29). Structural analysis of OTUB1 shows the preference of ubiquitin (ub)-charged E2 acting as an N-terminal extension of OTU domain (30,31) and also binding to free ubiquitin, both mimicking the configuration of the canonical product of its DUB activity, the K48-linked di-Ub chain (32).

A long N-terminal helix of OTUB1 is critical for E2-ubiquitin recognition and inhibition; OTUB2 thus fails to recognize and inhibit E2 since it lacks N-terminal helix (32). The proximal ubiquitin binds with OTUB2 in a different manner due to the striking structural differences in the N-terminal region (absence of N-terminal helix in OTUB2) and several reports (7,32,33) show that the absence of N-terminal helix may be the reason for the non-selective deubiquitinylase activity of OTUB2. The short loop (residue 44-48) preceding the active-site helix different from all other cysteine proteases and in the crystal structure is shifted towards the catalytic residues to generate a spatially restricted active site and stabilize the oxyanion intermediate (29). Interestingly, in OTUB2 only subtle changes were observed in the catalytic residues but larger conformational changes can be seen in the disordered region 197-204 (turned into ordered region upon ubiquitin binding) and helices 1 and 2 and beta-sheets 3 and 4 (28).

Deubiquitinating enzymes have recently been identified as relevant cancer drug targets (34-36). Molecular simulations have become essential tools in identifying protein conformational dynamics and rationalizing drug selectivity. For USP7 (37) two druggable binding sites were identified of which one was essential for the inactive to active state conformational transition. MD studies elucidated the open to close conformation transition in viral DUB (Turnip yellow mosaic virus) (38) in wild type and mutants. The drug selectivity of Zn-dependent human DUBs of the JAMM family Rpn11 and CSN5 towards capzim in and CSN5i-3 could be explained due to formation of heterodimeric protein-protein complexes (39).

In this work, we focus on OTUB1 and OTUB2 DUBs to understand their differences in structural stability in the apo and holo forms and the role of water molecules in stabilizing the catalytic triad conformations using molecular dynamics simulation. Analyzing the conformational stability of the catalytic triad residues and the access and passage and pathways of solvent molecules helps us to understand the role of water molecules in stabilizing the active site. Our findings suggest that OTUB1 achieves its productive conformation in its charged state only upon ubiquitin binding, whereas in OTUB2 the productive conformation already prevails in its holo form conformation in the neutral protonation state. A very low number of water trajectories towards the catalytic site from the surface were observed in the productive conformation of OTUB1 in contrast to the very large number observed in the productive conformation of OTUB2. A number of conserved water molecules are located close to His265 and Asp267 in OTUB1 and in vicinity of Cys51 in OTUB2.
Results and Discussion

The comparison of OTUB1 and OTUB2 sequences shows 48% identity and 70% similarity. (Supplementary Figure S1). Larger structural differences are apparent in the catalytic site and in the ubiquitin binding pockets (19). A long N-terminal α-helix is present in OTUB1 but missing in OTUB2. In our nomenclature, we refer to the apo OTUB1 structure in charged and neutral states as O1U0C and O1U0N respectively; likewise the holo OTUB1 structures in charged and neutral states is O1U1C and O1U1N. Similarly, apo and holo OTUB2 charged and neutral states will be addressed as O2U0C/O2U0N and O2U1C/O2U1N. For representational purpose, we show a zwitterionic catalytic triad (in O1U1C) and a neutral charge catalytic site (in O2U0N) in Figure 1.

Absence of Ub: apo OTUB1 (O1U0C, O1U0N) and OTUB2 (O2U0C, O2U0N)

Molecular dynamics simulations of OTUB1 (charged and neutral) and OTUB2 (charged and neutral) were replicated three times. The RMSD (Ca) plots of all apo OTUB1 and OTUB2 simulations were analyzed and show that the structures are stable throughout the simulations (see Supplementary Figures S2 and S3). In addition, the global protein RMSFs reveal no large fluctuations. Only the loop region connecting sheets 2 and 3 in OTUB2 (O2U0C, O2U0N) shows a high degree of flexibility (see Supplementary Figures S4 and S5).

Structural characteristics of the catalytic triad in ubiquitin-free OTUB1 and OTUB2

Catalytic protease activity requires an accurate positioning of the Cys/His residues to perform their catalytic function. The interatomic distances of the catalytic residues of both OTUB1 and 2 were monitored. The average distances and their standard deviations are given in Table 1: in the charged state of OTUB1 the distances are significantly smaller than in the neutral state (His223ND1…Cys91SG were 5.8±1.8Å compared to 6.4±0.9Å; His223NE2…Asp287OD1 4.8±1.4Å and 7.0±1.2Å). This is in good agreement with the catalytic geometry of the experimental structure and shows that the charged state is dominant in the X-ray structures.

For the zwitterionic state O2U0C average contact distances for His223ND1…Cys91SG of 5.5±0.9Å and His223NE2…Asn226OD1 4.7±1.9Å were obtained. Unexpectedly, the neutral O2U0N state shows shorter interatomic distances for His223ND1…Cys91SG of 3.9±0.8Å and His223NE2…Asn226OD1 of 4.5±1.8Å (see Figure 2B and Figure 3C and D). The His223 dihedral angle X(NE1-Ca-CB-CG) is 174±5°-154±44° degree in O2U0N, but in O2U0C it is 149±44°-138±52° and in O1U0C, O1U0N it is 84±52°-156±41° and 145±45°-67±19°, respectively (Table 1).

Apparently, there is no conformational change of the catalytic residues associated upon activation of OTUB2. The differences in structural parameters such as interatomic distances and dihedral angle are only minor and show that this is in agreement with experimental results which demonstrate the prevalence of the neutral state of apo OTUB2 to be catalytically active.

The Ub-bound protein conformation of OTUB1 and OTUB2

Simulations of the charged and neutral states of the catalytic triad residues of OTUB1 and OTUB2 in the ubiquitin-bound conformation were performed in triplicates. The RMSD (Ca) plots of all holo OTUB1 and OTUB2 show the stability of the charged and neutral states of OTUB2 but not for OTUB1. Whereas the OTU-domain of OTUB1 is rather rigid with a RMSD of 2 Å, it is the flexible N-terminus which is stabilized by interactions with the E2-linked ubiquitin binding and responsible for the very large RMSD of 6-8 Å in OTUB2 (see Supplementary Figure S6). The RMSF analysis further confirms that the large fluctuations originate from the N-terminal helical region (first 21 amino acids of the protein) and the rest of the protein is less flexible. (see Supplementary Figure S7). The analysis of the RMSF further also shows local residue fluctuations in the loop connecting the helices 2 and 3 in both charge states. Less pronounced flexible regions are helices 8 and 9 in the charged state (O1U1C) and the loop connecting the sheets 3 and 4 in the neutral state (O1U1N) (see Supplementary Figure S7).
OTUB2, which does not possess the N-terminal helix, reveals a smaller RMSD in both charge states (O2U1C, O2U1N) during the entire simulations (see Supplementary Figure S8). The analysis of the RMSFs of holo OTUB2 shows that the neutral state O2U1N is structurally more stable than the O2U1C except that the latter one shows sharp fluctuations in the short N-terminal helix and the small loop connecting the sheet 5 and 6 (see Supplementary Figure S9).

**Analysis of catalytic triad conformation in ubiquitin-bound OTUB1 vs. OTUB2**

In O1U1C, the average contact distance in the MD trajectories between His265ND1...Cys91SG is 3.3±0.2Å and the His265NE2 maintains a close contact distance of 2.7±0.1Å with Asp267OD1 (Figure. 2A; Figure 3A and B). The dihedral angle $\chi_1$(N-CA-CB-CG) of His265 is very well preserved throughout the simulation with the range of 173.7±4.6° and -175.4±3.6°. In O1U1N, the average distance of His265ND1 with Cys91SG in the simulations is significantly larger with 4.7±1.5Å and the His265NE2...Asp267OD1 distance of 4.6±2.1Å is likewise large and not arranged for catalysis. The dihedral of the His265 $(\chi_1)$ falls in the range of 170±17°/-147±45°. In the case of OTUB2, the different charge states O2U1C and O2U1N display average inter-residue distances between His265ND1 and Cys91SG of 5.9±0.7Å and 5.4±1.2Å, respectively. The average His265NE2...Asp267OD1 distances are 7.5±1.8Å and 6.5±1.7Å, respectively. The histidine dihedral $(\chi_1)$ for O2U1C is 74±44°/-59±23° and for O2U1N it is 126±62°/-79±46° (see Table 1). This shows that structural parameters for the holo form in the charged state of OTUB1 (O1U1C) are in good agreement with the crystal structure. The other feasible charge states for holo conformations of OTUB1 and OTUB2 are significantly deviating from the experimental structural structures.

In OTUB1, throughout the total simulation time of the O1U1C state, the strong salt bridge interaction between the catalytic site His265$^+$ and Asp267$^-$ is persistent. Conformational restrictions of the active site residues in OTUB1 are also due to Pro87 that is located in close proximity of Cys91 and also the small loop before the His265 has an effect on the Cys91 orientation. The role of α9-α10 loop to influence the conformational change on Cys91 and His265 upon the ubiquitin binding was only inferred from a static picture in the X-ray structure (19). A hydrogen bonding interaction between the C-terminal end of α3 and the backbone of the catalytic Cys91 was speculated to inhibit substrate access to the active site and be the reason for an OTUB1 ligand-induced activation step.

**Water access of active site in Ub-free in apo OTUB1 and OTUB2**

Water molecules in the active site play an important role in regulating the activity, stability and selectivity of enzymes (40-42). The buried active sites in OTUB1 and OTUB2 are connected with the bulk solvents by water tunnels. The molecular properties of the amino acids of the water tunnels control the access of the water molecules. Studying the water tunnels is important, as it is the major route of the substrate delivery to the active site center. The analysis of water molecules diffusing from the surface of the protein to the active site is a versatile tool to discriminate structurally similar proteins. Despite our special focus on dissecting the active states’ conformations O1U1C and O2U1N, we have also analyzed the complementary conformations of the OTUB1 and OTUB2 proteins in the different charge states.

In O1U1C, the number of water inlets is 1125 of which 471 are incoming and 654 outgoing paths. The localization of water molecules in vicinity of the active site residues His265 and Cys91 is shown in Figure 4A. A high density of water molecules is also observed close to these residues with longer times of residence due to strong hydrogen bonding and electrostatic interactions (Figure 4C). In O1U1N, the number of inlets is smaller with 776, of which 387 are incoming and 389 outgoing paths. The water inlets are mostly located close to the region of His265 and Asp267 (Figure 4B). The local distribution of water molecules is high near the side chain oxygen of Asp267, and low close to His265 and not present in vicinity of Cys91 (see Figure 4D). This shows that the catalytic aspartate residue plays a vital role in trapping water molecules close to the active site region of OTUB1.

In O2U1C, 260 water inlets can be identified and the numbers of incoming and outgoing channels are almost evenly distributed with 135 and 123, respectively. In this case, the number of water molecules which enter or reside close to the active
site His224 is high compared to other two active site residues cysteine and aspartate (see Figure 5A). The local distribution of water molecules is high near the His224 sidechain compared to the other active site residue Asn226 (Figure 5C). In O2U1N, on the other hand, 1499 inlets are observed out of which 833 are incoming and 666 outgoing paths (see Figure 5B). O2U0N has numerous water trajectory paths with more of incoming and outgoing paths compared to a small number of water molecules residing trapped in the active site region (Figure 5D). In O2U0N, the absence of the long N-terminal helix allows the opening of two large cavities located near the entrance of the catalytic triad, which act as the main route of access for the water molecules in this truncated structure. The water molecules are not densely located close to one catalytic residue but sparsely distributed around the whole catalytic triad. More specifically, analysis of the local water distribution density shows that the water molecules reside longer in vicinity of the catalytic cysteine residue and can less frequently be found surrounding the catalytic histidine and asparagine residues.

**Water access in the Ub-bound conformation of OTUB1 and OTUB2**

In the catalytically active state of OTUB1 (O1U1C), the water access to the active site is limited with a total of just 19 inlets comprising of 10 outgoing and 9 incoming paths (see Figure 6A). In O1U1C, the highly fluctuating extended N-terminal helix may acts as a swinging gate and not enable many water molecules to diffuse from the surface to the active site region and vice versa. In O1U1C, very few molecules enter and leave the active site; compared to the number of incoming and outgoing water molecules’ trajectories more water molecules seem to be trapped inside the catalytic triad. This can be attributed to the strong salt bridge and stable interactions observed in the charged catalytic site state and also the presence of the long N-terminal helix. The local distribution density of water molecules enables the identification of essential residues capable of trapping water molecules and it can also be used for conserved water identification or regulation of ligand access (43). In O1U1C, a very low water distribution density is found. In O1U1C, the water molecules are localized around the catalytic residues His265 and Asp267, especially close the His265 (Figure 6C). In contrast to the charge state of OTUB1, in the neutral state (O1U1N), the total number of inlets is larger with 361, of which almost equal half of them are incoming and outgoing (see Figure 6B). Analysis of the water residence time distribution shows that the water molecules are trapped far from the catalytic triad and remain in that pocket during the entire simulation. Although O1U1N has a larger number of water sites compared to O1U1C, they are not located in vicinities of the catalytic sites and hence the access to the catalytic site is restricted. The water distribution density also shows that during the simulations the water molecules are found distant from the catalytic sites and only very few are approach the catalytic cysteine residue (see Figure 6D).

In the charged state of the Ub-bound conformation of OTUB2 (O2U1C), 586 water inlets were identified out of which 307 are incoming and 279 are outgoing (Figure 7A). The local distribution density of water molecules is higher for the catalytic residue histidine (His224) than for the other two residues (Figure 7C). In the neutral state of OTUB2 (O2U1N), out of 896 water inlets identified the number of incoming and outgoing paths is equally distributed with 450 and 446, respectively (Figure 7B). The local water distribution density is not high close to the catalytic residues but at a certain distance from the active site histidine and asparagine residues and remote from the active site cysteine (see Figure 7D).

**Comparative analysis of water trajectories of OTUB1 in Ub-free vs. Ub-bound protein conformations**

The comparison of the OTUB1 zwitterionic state in the ubiquitin-free (apo) and ubiquitin-bound (holo) protein conformations shows that there are thousands of water inlets and the outgoing paths are larger in number in the free protein. The holo state shows less water inlets with just around 20, which equally distributed among the incoming and outgoing. If you compare the local water distribution density, both apo and holo conformations show a preferred water distribution density close to the catalytic histidine and cysteine
91 in the apo form and aspartate 265 in the holo form.

As an aside, the speculative neutral protonation state of OTUB1 in the apo and holo protein conformations shows very different results. In the apo state around 780 inlets are equally distributed between incoming and outgoing paths. In the holo state, there is a larger number of water inlets compared to the charged state; there are around 360 which are equally distributed as in the apo state. The major site of localization is remote from the catalytic region and not significant for the stabilization of the catalytic triad residues.

Comparative analysis of water trajectories of OTUB2 in Ub-free vs. Ub-bound protein conformations

The comparison of charged states of apo and holo structures of OTUB2 shows a significant difference between Ub-free and Ub-bound protein conformations. In the apo conformation, only half of the water inlets are present compared to the holo conformation. In the apo state (O2U0C), 260 water inlets are almost equally distributed among incoming and outgoing channels (only slightly less outgoing paths) and the large number of 586 water inlets in the holo form (O2U1C) are also almost equally distributed among the incoming and outgoing (slightly less in outgoing). In both cases, the local water distribution density is high in the catalytic site.

The same observation can also be made for the neutral states. In the OTUB2 apo conformation (O2U0N), of the total 1500 water inlets there are more (833) incoming and 666 outgoing (666) identified. However, in the holo conformation (O2U1N), around 900 inlets are almost equally distributed between incoming and outgoing. Interestingly in the Ub-free conformation, the local water distribution density is high in vicinity of the side chain atoms of all three catalytic residues, whereas as in the holo conformation state it is remote from the active site.

Catalytically active forms OTUB1 (O1U1C) vs OTUB2 (O2U0N)

OTUB1 is catalytically active only when ubiquitin is bound which points to a substrate-induced activation process. The active site in OTUB1 is not competent for catalysis in its apo state since the short loop connecting β4-β5 and the loop between helices 9 and 10 have an influence on the catalytic cysteine orientation and mediate the conformational change upon ubiquitin binding. Based on the careful analysis of catalytic site geometry from our molecular dynamics simulations, we infer that the zwitterionic state of OTUB1 in the Ub-bound conformation (O1U1C) represents the catalytically active state.

In contrast, OTUB2 is already active in the Ub-free apo state and the active site residues are in a pre-arranged, catalytically active conformation. However, according to our results, we stress that the neutral charge state of OTUB2 (Cys-SH, His0, Asn0, Asp0) is in a catalytically active conformation (O2U0N) and only minor proton transfer events are required to initiate the protease activity.

The comparison of the water trajectories between the catalytically active OTUB states, however, shows that there are significant differences between the OTUs. OTUB2 (O2U0N) possesses a huge number of water inlets compared to OTUB1 (O1U1C) and the clustering of water molecules is close to the catalytic site region in OTUB2 but remote from the catalytic site in OTUB1. The preferred sites of localization are all three catalytic residues in OTUB2 but only histidine and aspartate in OTUB1. The inner pocket volumes calculated based on the water molecule distribution densities reveal a smaller pocket of just 62 Å³ in OTUB1, whereas in OTUB2 it is nearly six fold larger with a volume of 362 Å³ (see Figure 8).

Water-mediated interactions in the catalytic triad residues

After the completion of pathway analysis of water molecules and its distribution in the active site region, we also analyzed the hydrogen bonding interactions of the conserved water molecules with the key catalytic residues. We only mention the water-mediated hydrogen bond interactions between the catalytic residues for the productive conformations of OTUB1 and OTUB2.

In O1U1C, in only 1.4% of the simulation time direct water-mediated interactions between the ND1 of His265 and the sulfur atom (SG) of cysteine residue (Cys91) can be observed (see Figure 9). No extended water-mediated
stabilizations of two or more water molecules are found. The NE2 of His265 and OD1 of Asp267 show water-mediated interactions for around 11% of the simulation time. For only 1% of the simulation time, extended water-mediated contacts are observed. In O1U1C, the salt bridge interaction between the positively charged His265 and the negatively charged Asp267 is stable and likewise a hydrogen bond interaction between the proton at the NE2 atom of His265 and OD1 atom of Asp267. The water molecule also makes a hydrogen bond interaction with OD2 atom of Asp267 and backbone oxygen atom of the His265 (Figure 10A).

In the case of O2U0N, ND1 of His224 and SG of Cys51 show direct water-mediated interactions for 11.4% of the simulation time and furthermore extended water-mediated contacts for nearly 2% of the simulation time. Similarly, the NE2 of His 224 has direct and extended water mediated interaction for nearly 9% and 4.2%, respectively of the simulation time with OD1 of Asn226 (see Figure 9).

In case of O2U0N, the water molecule mediates the interaction between histidine and asparagine only when the distance between these residues increases sufficiently during the simulation. Then, the water molecule bridges the OD1 atom of Asn226 and NE2 atom of His224 (see Figure 10B). If the distance further increases, an extended water-mediated interactions can be observed in which more than one water molecule form an interaction bridging the OD1 atom of Asn226 and NE2 atom of His224 (Figure 10C).

The results show that the persistent and strong electrostatic interactions between the zwitterionic catalytic residues in O1U1C and the presence of a long flexible N-terminal helix restrict and control the entry and egress of water molecules to the catalytic triad.

CONCLUSIONS

The detailed structural comparison of the zwitterionic and neutral states of Ub-free and Ub-bound protein conformations of OTUB1 and OTUB2 shows that the apo OTUB1 structure with a charged catalytic site (O1U1C) is in a stable but catalytically incompetent conformation. OTUB1 requires substrate-induced protease activation. A possible explanation is that OTUB1 is in inactive conformation in absence of ubiquitin and a conformational change may induce a movement of His265 in closer proximity of Cys91 upon ubiquitin substrate binding. Also the DUBs USP7 (44) and USP12 crystallize in an auto-inhibited state; USP12 requires an extra allosteric activation upon UAF1 and WDR20 binding (45). The allosteric binding to remote binding sites increases the catalytic rate $k_{cat}$ by a factor of 20 whereas the substrate affinity ($K_m$) remains almost unchanged. OTUB1 is highly selective for K48-linked polyubiquitin chains, but not cleaving K63-, K29-, K6-, or K11-linked polyubiquitins (31). The selectivity could be rationalized by a bidentate ubiquitin binding model with distinct sites for catalytic activity (Cys95) and a second distal binding site. Also, the role of the N-terminal extension of its core OTU-domain is not fully resolved yet (32). According to our work, the long extended N-terminal helix resolved in the O1U1C is flexible throughout the simulation and acts as a gate of specificity regulation to control substrate and solvent access to the catalytic site.

In OTUB2, the catalytic triad is stabilized by unique hydrogen bonding network configuration (29). The similarity in the side-chain orientations of Cys51 and His224 in OTUB2 to those of other cysteine protease structures implies that they are in a catalytically productive geometry even in absence of ubiquitin binding. On the basis of sequence analysis of OTUs, the active site catalytic triad is commonly annotated as Cys51/His224 and Asp48, the latter stabilizing the orientation of the protonated histidine. In the apo and holo crystal structures and in our simulations, we can show that this aspartate is at too far distance to directly interact with His224, but Asn226 forms hydrogen bonds with His224 and nearby other residues. This configuration is stable in both Ub-free and Ub-bound states and requires only a minor proton transfer event from Cys51 to His224 without any configurational re-orientation to initiate the protease activity. The role of Asn226 in the isopeptidase reactions is corroborated by the absence of catalytic activity upon an Asn226Ala mutation. Based on the water trajectory analysis, OTUB2 is more prone to enable water diffusion and passage to and from the active site compared to OTUB1.
DUBs and OTUs have recently emerged and promising new drug targets for treating various types of cancer. The role of an asparagine residue in a stable configuration to replace an aspartate in cysteine proteases, confirms that there are clearly distinct functional subclasses within the OTU family. The difference in protonation states of the residues of the catalytic triad (diad) plus selective ubiquitin binding sites proximal and distal to the active site will allow the design of selective inhibitors.

**EXPERIMENTAL DETAILS**

**Structure preparation**

Crystal structures of OTUB1 and OTUB2 in the apo and co-crystallized ubiquitin-bound states are available from the PDB (www.pdb.org); for OTUB1 structures in absence of ubiquitin (2ZFY), and in the ubiquitin-bound state (4DDG); and for OTUB2 (ubiquitin-free structure 1TFF) and co-crystallized with the ubiquitin (4FJV) (Table 2). Downloaded structures were preprocessed to add missing loops, amino acid side chains and revert introduced mutations (in the crystal structure 4DDG-S1091C) using Prime module (Schrodinger Inc, USA); then further processed to assign bond orders, add hydrogen atoms and identify hydrogen bond interactions using the protein preparation wizard. In the resulting structures, only hydrogen atoms were minimized using the Epik module (Schrodinger Inc, USA). To carry out an unbiased study, the ubiquitin molecule, the E2 ligase and solvent molecules other structural units were removed from both OTUB1 and 2 structures. The cysteine and histidine residues in the catalytic triad were prepared in neutral (protonated Cys-SH/His\(^{\text{a}}\) N\(_{\text{e}}\) and charged (deprotonated Cys-S'/His\(^{+}\) imidazolium cation) using the 3D builder (Schrodinger Inc, USA).

**Molecular dynamics simulation**

Every simulation was performed in triplicates of all-atom 100 ns simulations using Gromacs 2018.3 (46) and the CHARMM36 force field (47). The TIP3P water model was used and 0.15 M of salt (Na\(^{+}\)Cl\(^{-}\)) was added to neutralize the system. Energy minimization was carried out using the steepest descent algorithm for 5000 steps.

Equilibrations were carried out in NVT and NPT ensembles for 1 ns and 2 ns, respectively. Temperature (310 K) and pressure (1 atm) were controlled by the velocity rescaling thermostat and a Parrinello-Rahman barostat (48), respectively. For the treatment of long-range electrostatic interactions, the Particle-mesh Ewald summation (PME) was used. The three independent production runs for 100 ns each were carried out without any position restraints in an NPT ensemble. A time step of 2 fs was used and the coordinates were saved for every 2 ps.

**Water pathway and analysis**

The paths of the solvent diffusion to and from the active site area (within a radius of 4Å from the catalytic triad) to the surface of the protein were analyzed using AQUA-DUCT software (49). Aqua-duct is a tool to perform a detailed analysis of solvent access and channels in proteins. Each water molecule, which diffuses between a defined active site sphere to the surface of the protein, is individually tracked. The raw paths connecting the surface of the protein and the active site are determined by looping over all frames of the simulation trajectories. Separate paths for each water molecule, which enters and leaves the surface of the protein and active site sphere during the simulations are calculated. A clustering of the inlets (points where the traceable water molecules enters or leave the surface of the protein) is performed using different clustering algorithms. Finally, the results can be visualized using Pymol (50). It also allows to classify pockets as ‘inner’ and ‘outer’ when analyzing the distribution of local water molecules on a grid spanning all paths with default size of 1Å. The maximal space explored by traced water molecules represents the ‘outer’ pocket whereas the ‘inner’ pocket is the area, which is easily accessible by water molecules. Along the entire trajectory, the number of paths crossing the grid cells divided by number of frames gives the average density of traced residues. It can also detect high-density water points (aka ‘water hotspots’) from the distribution of local densities on the grid, traced water molecules trapped in hydrophobic cages or attracted by favorable interactions with nearby amino acids and long residence times. The results were visualized using PYMOL software (50).
Computation of direct and water-mediated interactions in the active site

The HBonds plugin in VMD was used to calculate the hydrogen bonds along the molecular dynamics simulations (51) and the geometric criteria of donor-acceptor distance should be <3.5Å and a donor-hydrogen-acceptor angle >100°. A water-mediated interaction (WB) between two residues is calculated when one water molecule is bridging a hydrogen bond donor and acceptor. An extended water-mediated interaction (WB2) is defined as a water-mediated hydrogen bond formed between two residues by two bridging water molecules. Though we have analyzed the entire structures, the focus in this work is on the water-mediated interaction between the key atoms of the catalytic triad residues.

Data availability All prepared protein structural files in pdb format and topology files (top) to reproduce the Molecular Dynamics simulations with GROMACS are freely available at https://doi.org/10.5281/zenodo.3741678.
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REFERENCES

1. Reyes-Turcu, F. E., Ventii, K. H., and Wilkinson, K. D. (2009) Regulation and cellular roles of ubiquitin-specific deubiquitinating enzymes. Annual Review of Biochemistry 78, 363-397
2. Zhang, D., Zaugg, K., Mak, T. W., and Elledge, S. J. (2006) A role for the deubiquitinating enzyme USP28 in control of the DNA-damage response. Cell 126, 529-542
3. Adhikari, A., Xu, M., and Chen, Z. J. (2007) Ubiquitin-mediated activation of TAK1 and IKK. Oncogene 26, 3214-3226
4. Abdul Rahman, S. A., Kristariyanto, Y. A., Choi, S. Y., Nkosi, P. J., Weidlich, S., Labib, K., Hofmann, K., and Kulathu, Y. (2016) MINDY-1 Is a Member of an Evolutionarily Conserved and Structurally Distinct New Family of Deubiquitinating Enzymes. Molecular Cell 63, 146-155
5. Clague, M. J., Barsukov, I., Coulson, J. M., Liu, H., Rigden, D. J., and Urbe, S. (2013) Deubiquitylases from genes to organism. Physiological Reviews 93, 1289-1315
6. Sun, J., Shi, X., Mamun, M. A. A., and Gao, Y. (2020) The role of deubiquitinating enzymes in gastric cancer. Oncology Letters 19, 30-44
7. Mevissen, T. E., Hospenthal, M. K., Geurink, P. P., Elliott, P. R., Akutsu, M., Arnaudo, N., Ekkebus, R., Kulathu, Y., Wauer, T., El Oualid, F., Freund, S. M., Ovaa, H., and Komander, D. (2013) OTU deubiquitinases reveal mechanisms of linkage specificity and enable ubiquitin chain restriction analysis. Cell 154, 169-184
8. Balakirev, M. Y., Tcherniuk, S. O., Jaquinod, M., and Chroboczek, J. (2003) Otubains: a new family of cysteine proteases in the ubiquitin pathway. EMBO Reports 4, 517-522
9. Borodovsky, A., Ovaa, H., Kolli, N., Gan-Erdene, T., Wilkinson, K. D., Ploegh, H. L., and Kessler, B. M. (2002) Chemistry-based functional proteomics reveals novel members of the deubiquitinating enzyme family. Chemistry & Biology 9, 1149-1159
10. Xu, L., Li, J., Bao, Z., Xu, P., Chang, H., Wu, J., Bei, Y., Xia, L., Wu, P., Yan, K., Lu, B., and Cui, G. (2017) Silencing of OTUB1 inhibits migration of human glioma cells in vitro. Neuropathology : Official Journal of the Japanese Society of Neuropathology 37, 217-226
11. Zhou, H., Liu, Y., Zhu, R., Ding, F., Cao, X., Lin, D., and Liu, Z. (2018) OTUB1 promotes esophageal squamous cell carcinoma metastasis through modulating Snail stability. Oncogene 37, 3356-3368
12. Stanisic, V., Malovannaya, A., Qin, J., Lonard, D. M., and O’Malley, B. W. (2009) OTU Domain-containing ubiquitin aldehyde-binding protein 1 (OTUB1) deubiquitinates estrogen receptor (ER) alpha and affects ERalpha transcriptional activity. The Journal of Biological Chemistry 284, 16135-16145
13. Ni, Q., Chen, J., Li, X., Xu, X., Zhang, N., Zhou, A., Zhou, B., Lu, Q., and Chen, Z. (2017) Expression of OTUB1 in hepatocellular carcinoma and its effects on HCC cell migration and invasion, Acta Biochimica et Biophysica Sinica 49, 680-688
14. Weng, W., Zhang, Q., Xu, M., Wu, Y., Zhang, M., Shen, C., Chen, X., Wang, Y., and Sheng, W. (2016) OTUB1 promotes tumor invasion and predicts a poor prognosis in gastric adenocarcinoma. American Journal of Translational Research 8, 2234-2244
Zhao, L., Wang, X., Yu, Y., Deng, L., Chen, L., Peng, X., Jiao, C., Gao, G., Tan, X., Pan, W., Ge, X., and Wang, P. (2018) OTUB1 protein suppresses mTOR complex 1 (mTORC1) activity by deubiquitinating the mTORC1 inhibitor DEPTOR. *The Journal of Biological Chemistry* **293**, 4883-4892

Liu, X., Jiang, W. N., Wang, J. G., and Chen, H. (2014) Colon cancer bears overexpression of OTUB1. *Pathology. Research and Practice* **210**, 770-773

Karunarathna, U., Kongsema, M., Zona, S., Gong, C., Cabrera, E., Gomes, A. R., Man, E. P., Khongkow, P., Tsang, J. W., Khoo, U. S., Medema, R. H., Freire, R., and Lam, E. W. (2016) OTUB1 inhibits the ubiquitination and degradation of FOXM1 in breast cancer and epirubicin resistance. *Oncogene* **35**, 1433-1444

Li, J., Cheng, D., Zhu, M., Yu, H., Pan, Z., Liu, L., Geng, Q., Pan, H., Yan, M., and Yao, M. (2019) OTUB2 stabilizes U2AF2 to promote the Warburg effect and tumorigenesis via the AKT/mTOR signaling pathway in non-small cell lung cancer. *Theranostics* **9**, 179-195

Edelmann, M. J., Iphofer, A., Akutsu, M., Altun, M., di Gleria, K., Kramer, H. B., Fiebiger, E., Dhe-Paganon, S., and Kessler, B. M. (2009) Structural basis and specificity of human otubain 1-mediated deubiquitination. *The Biochemical Journal* **418**, 379-390

Coulombe, R., Grochulski, P., Sivaraman, J., Menard, R., Mort, J. S., and Cygler, M. (1996) Structure of human procathepsin L reveals the molecular basis of inhibition by the prostate. *The EMBO Journal* **15**, 5492-5503

Blow, D. M., Birktoft, J. J., and Hartley, B. S. (1969) Role of a buried acid group in the mechanism of action of chymotrypsin. *Nature* **221**, 337-340

Elsasser, B., Zauner, F. B., Messner, J., Soh, W. T., Dall, E., and Brandstetter, H. (2017) Distinct Roles of Catalytic Cysteine and Histidine in the Protease and Ligase Mechanisms of Human Legumain As Revealed by DFT-Based QM/MM Simulations. *ACS Catalysis* **7**, 5585-5593

Wei, D., Huang, X., Liu, J., Tang, M., and Zhan, C. G. (2013) Reaction pathway and free energy profile for papain-catalyzed hydrolysis of N-acetyl-Phe-Gly 4-nitroanilide. *Biochemistry* **52**, 5145-5154

Shokhen, M., Khazanov, N., and Albeck, A. (2011) The mechanism of papain inhibition by peptidyl aldehydes. *Proteins* **79**, 975-985

Shokhen, M., Khazanov, N., and Albeck, A. (2009) Challenging a paradigm: theoretical calculations of the protonation state of the Cys25-His159 catalytic diad in free papain. *Proteins* **77**, 916-926

Craik, C. S., Roczniai, S., Largman, C., and Rutter, W. J. (1987) The catalytic role of the active site aspartic acid in serine proteases. *Science* **237**, 909-913

Vernet, T., Tessier, D. C., Chatellier, J., Plouffe, C., Lee, T. S., Thomas, D. Y., Storer, A. C., and Menard, R. (1995) Structural and functional roles of asparagine 175 in the cysteine protease papain. *The Journal of Biological Chemistry* **270**, 16645-16652

Altun, M., Walter, T. S., Kramer, H. B., Herr, P., Iphofer, A., Bostrom, J., David, Y., Komsany, A., Ternette, N., Navon, A., Stuart, D. I., Ren, J., and Kessler, B. M. (2015) The human otubain2-ubiquitin structure provides insights into the cleavage specificity of poly-ubiquitin-linkages. *PloS One* **10**, e0115344

Nanao, M. H., Tcherniuk, S. O., Chroboczek, J., Dideberg, O., Dessen, A., and Balakirev, M. Y. (2004) Crystal structure of human otubain 2. *EMBO Reports* **5**, 783-788

Nakada, S., Tai, I., Panier, S., Al-Hakim, A., Iemura, S., Juang, Y. C., O'Donnell, L., Kumakubo, A., Munro, M., Sicheri, F., Gingras, A. C., Natsume, T., Suda, T., and Durocher, D. (2010) Non-canonical inhibition of DNA damage-dependent ubiquitination by OTUB1. *Nature* **466**, 941-946

Wang, T., Yin, L., Cooper, E. M., Lai, M. Y., Dickey, S., Pickart, C. M., Fushman, D., Wilkinson, K. D., Cohen, R. E., and Wolberger, C. (2009) Evidence for bidentate substrate binding as the basis for the K48 linkage specificity of otubain 1. *Journal of Molecular Biology* **386**, 1011-1023

Juang, Y. C., Landry, M. C., Sanches, M., Vittal, V., Leung, C. C., Ceccarelli, D. F., Mateo, A. R., Pruneda, J. N., Mao, D. Y., Szilard, R. K., Orlicky, S., Munro, M., Brzovic, P. S., Klevit, R. E.,
Sicheri, F., and Durocher, D. (2012) OTUB1 co-opts Lys48-linked ubiquitin recognition to suppress E2 enzyme function. *Molecular Cell* **45**, 384-397

33. Wiener, R., DiBello, A. T., Lombardi, P. M., Guzzo, C. M., Zhang, X., Matunis, M. J., and Wolberger, C. (2013) E2 ubiquitin-conjugating enzymes regulate the deubiquitinating activity of OTUB1. *Nature Structural & Molecular Biology* **20**, 1033-1039

34. D'Arcy, P., Wang, X., and Linder, S. (2015) Deubiquitinase inhibition as a cancer therapeutic strategy. *Pharmacology & Therapeutics* **147**, 32-54

35. Fraile, J. M., Quesada, V., Rodriguez, D., Freije, J. M. P., and Lopez-Otin, C. (2012) Deubiquitinases in cancer: new functions and therapeutic options. *Oncogene* **31**, 2373-2388

36. Shen, M., Schmitt, S., Buac, D., and Dou, Q. P. (2013) Targeting the ubiquitin-proteasome system for cancer therapy. *Expert Opinion on Therapeutic Targets* **17**, 1091-1108

37. Srivastava, M., Suri, C., Singh, M., Mathur, R., and Asthana, S. (2018) Molecular dynamics simulation reveals the possible druggable hot-spots of USP7. *Oncotarget* **9**, 34289-34305

38. Jupin, I., Ayach, M., Jomat, L., Fieulaine, S., and Bressanelli, S. (2017) A mobile loop near the active site acts as a switch between the dual activities of a viral protease/deubiquitinase. *PLoS Pathogens* **13**, e1006714

39. Stepankova, V., Khabiri, M., Brezovsky, J., Pavelka, A., Sykora, J., Amaro, M., Minofar, B., Prokop, Z., Hof, M., Ettrich, R., Chaloupkova, R., and Damborsky, J. (2013) Expansion of access tunnels and active-site cavities influence activity of haloalkane dehalogenases in organic cosolvents. *ChemBioChem: a European Journal of Chemical Biology* **14**, 890-897

40. Yang, L., Dordick, J. S., and Garde, S. (2004) Hydration of enzyme in nonaqueous media is consistent with solvent dependence of its activity. *Biophysical Journal* **87**, 812-821

41. Mitusinska, K., Magdziarz, T., Bzowka, M., Stanczak, A., and Gora, A. (2018) Exploring Solanum tuberosum Epoxide Hydrolase Internal Architecture by Water Molecules Tracking. *Biomolecules* **8**, 143

42. Li, H., Lim, K. S., Kim, H., Hinds, T. R., Jo, U., Mao, H., Weller, C. E., Sun, J., Chatterjee, C., D'Andrea, A. D., and Zheng, N. (2016) Allosteric Activation of Ubiquitin-Specific Proteases by beta-Propeller Proteins UAF1 and WDR20. *Molecular Cell* **63**, 249-260

43. Huang, J., and MacKerell, A. D., Jr. (2013) CHARMm36 all-atom additive protein force field: validation based on comparison to NMR data. *Journal of computational chemistry* **34**, 2135-2145

44. Nosé, S., and Klein, M. L. (1983) Constant pressure molecular dynamics for molecular systems. *Molecular Physics* **50**, 1055-1076

45. Magdziarz, T., Mitusinska, K., Bzowka, M., Raczynska, A., Stanczak, A., Banas, M., Bagrowska, W., and Gora, A. (2019) AQUA-DUCT 1.0: structural and functional analysis of macromolecules from an intramolecular voids perspective. *Bioinformatics*

46. DeLano, W. L., and Lam, J. W. (2005) PyMOL: A communications tool for computational models. *Abstr Pap Am Chem S* **230**, U1371-U1372

47. Humphrey, W., Dalke, A., and Schulten, K. (1996) VMD: visual molecular dynamics. *Journal of Molecular Graphics* **14**, 33-38
Table 1. Comparison of characteristic catalytic inter-residue distances from available X-ray structures and calculated averages +/- standard deviation from MD simulations of OTUB1 and OTUB2 in Å.

| Protein | Protein conformation | Catalytic site protonation state | His-Cys (Å) | Asp-His(Å) | His (N-CA-CB-CG) dihedral angle (deg) |
|---------|----------------------|---------------------------------|-------------|------------|-------------------------------------|
| OTUB1   | Ub-bound             | 4DDG                            | 3.5         | 3.3        | 165                                 |
|         | O1U1C                | Charged                         | 3.3±0.2     | 2.7±0.1    | 173±05/-175±04                      |
|         | O1U1N                | Neutral                         | 4.7±1.5     | 4.6±2.1    | 170±17/-147±45                      |
|         | Ub-free              | 2ZFY                            | 5.5         | 6.7        | -65                                 |
|         | O1U1C                | Charged                         | 5.8±1.8     | 4.8±1.4    | 084±52/-156±41                      |
|         | O1U1N                | Neutral                         | 6.4±0.9     | 7.0±1.2    | 145±45/-067±19                      |
| OTUB2   | Ub-bound             | 4FJV                            | 4.0         | 2.9        | -179                                |
|         | O2U1C                | Charged                         | 5.9±0.7     | 7.5±1.8    | 074±44/-059±23                      |
|         | O2U1N                | Neutral                         | 5.4±1.2     | 6.5±1.7    | 126±62/-079±46                      |
|         | Ub-free              | 1TFF                            | 4.0         | 2.9        | -176                                |
|         | O2U1C                | Charged                         | 5.5±0.9     | 4.7±1.9    | 149±44/-138±52                      |
|         | O2U1N                | Neutral                         | 3.9±0.8     | 4.5±1.8    | 174±05/-150±44                      |

Table 2. Protein structures with pdb codes used in the study

| S.No | Pdb id | Protein | Complex | Active site                   |
|------|--------|---------|---------|-------------------------------|
| 1.   | 4DDG   | OTUB1   | Ubiquitin | Ser1091*-His1265-Asp1267     |
| 2.   | 2ZFY   | OTUB1   | n.a.     | Cys91-His265-Asp267          |
| 3.   | 4FJV   | OTUB2   | Ubiquitin | Cys51-His224-Asn226        |
| 4.   | 1TFF   | OTUB2   | n.a.     | Cys51-His224-Asn226        |

*Active site cysteine was mutated to serine
Figure 1. Structures of (A) OTUB1 (O1U₁C) in the ubiquitin-bound protein conformation and (B) OTUB2 (O2U₀N) in the ubiquitin-free protein conformation. Residues of the catalytic triad are labelled and their charged states are given in the first.

Figure 2. Interatomic distances with standard deviations (in Å) of the key atoms of the catalytic residues monitored throughout simulations (A) OTUB1 (O1U₁C) and (B) OTUB2 (O2U₀N).

Figure 3. Monitoring of interatomic distances of key catalytic residues (His-Cys-Asp/Asn) in MD simulations. (A) His⁺…Cys⁺, and (B) His⁺…Asp⁻ residues in OTUB1 (O1U₁C); (C) His⁰…Cys⁰, and (D) His⁰…Asn⁰ residues of OTUB2 (O2U₀N).
Figure 4. Analysis of water access in ubiquitin-free OTUB1 structures. **Top panel:** Water inlets to the catalytic site for incoming - outgoing water molecules. **Bottom panel:** Local distribution density of water molecules to identify stable water molecules **(A)** and **(C)** charged state of active site O1U0C; **(B)** and **(D)** neutral charge state of active site O1U0N.

Figure 5. Analysis of water access to the active site in ubiquitin-free OTUB2 structures. **Top panel:** Water inlets to the catalytic site for incoming - outgoing water molecules. **Bottom panel:** Local distribution density of water molecules to identify stable water molecules. **(A)** and **(C)** charged states of catalytic triad residues O2U0C; **(B)** and **(D)** neutral charge state of catalytic triad residues O2U0N.
Figure 6. Water trajectory analysis of ubiquitin-bound OTUB1 structures. **Top panel:** Water inlets to the catalytic site for incoming - outgoing water molecules. **Bottom panel:** Local distribution density of water molecules to identify stable water molecules. (A) and (C) charged state of catalytic residues O1U\textsubscript{C}; (B) and (D) neutral state of active site O1U\textsubscript{N}.

Figure 7. Water trajectory analysis of ubiquitin-bound OTUB2 structures. **Top panel:** Water inlets to the catalytic site for incoming - outgoing water molecules. **Bottom panel:** Local distribution density of water molecules to identify stable water molecules. (A) and (C) O2U\textsubscript{C} and (B) and (D) O2U\textsubscript{N}.
Figure 8. Solvent accessibility of active sites. Top panel is O1U1C and bottom panel is O2U0N. (A) and (D) localization of water molecules (represented as small balls) with passages to and from the active site of OTUB1 and OTUB2. (B) and (E) water paths cluster density was absent in O1U1C and spread across the entire catalytic triad in O2U0N. (C) and (F) Local water distribution densities represented as spheres and inner pocket volume represented as mesh.

Figure 9. Comparison of hydrogen bonding interactions in different states and protein conformations of OTUB1 and OTUB2. Direct water-mediated interactions (WB) by one water molecule and extended water-mediated interactions by more than one water molecule (WB2).
Figure 10. Water-mediated hydrogen bond (WB) interactions by one water molecule between the key catalytic residues in (A) O1U<C and (B) O2U<N. (C) Extended water-mediated hydrogen bond (WB2) by two water molecules in O2U<N. Hydrogen bonding interactions are given as black and salt bridge interactions as blue dashed lines.
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