High-resolution structure of mammalian PI31–20S proteasome complex reveals mechanism of proteasome inhibition

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Proteasome-catalyzed protein degradation mediates and regulates critical aspects of many cellular functions and is an important element of proteostasis in health and disease. Proteasome function is determined in part by the types of proteasome holoenzymes formed between the 20S core particle that catalyzes peptide bond hydrolysis and any of multiple regulatory proteins to which it binds. One of these regulators, PI31, was previously identified as an in vitro 20S proteasome inhibitor, but neither the molecular mechanism nor the possible physiologic significance of PI31-mediated proteasome inhibition has been clear. Here we report a high-resolution cryo-EM structure of the mammalian 20S proteasome in complex with PI31. The structure shows that two copies of the intrinsically disordered carboxyl terminus of PI31 are present in the central cavity of the closed-gate conformation of the proteasome and interact with proteasome catalytic sites in a manner that blocks proteolysis of substrates but resists their own degradation. The two inhibitory polypeptide chains appear to originate from PI31 monomers that enter the catalytic chamber from opposite ends of the 20S cylinder. We present evidence that PI31 can inhibit proteasome activity in mammalian cells and may serve regulatory functions for the control of cellular proteostasis.

Intracellular protein degradation plays a critical role in mediating and regulating many aspects of normal cellular function. Accordingly, abnormal or dysregulated protein degradation is a feature of many human diseases including cancer and those associated with muscle wasting and neurodegeneration (1-4). Most protein degradation in eukaryotic cells is catalyzed by the proteasome, a modular protease system consisting of multiple holoenzymes composed of a common protease, the 20S proteasome, and any of various regulatory proteins including 19S/PA700, PA28αβ, PA28γ, PA200, p97, and PI31 (5, 6). The 20S proteasome (aka Core Particle) is a 28-subunit protease complex composed of four axially stacked heteroheptameric rings (7, 8). Each of the two identical outer rings contains seven unique α-type subunits,

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whereas each of the two identical inner rings contains seven unique β -type subunits, resulting in an $\alpha 1-7/\beta 1-7/\beta 1-7/\alpha 1-7$ cylindrical architecture. The β 1, β 2, and β 5 subunits of each inner ring harbor N-terminal threonine residues that line the central interior chamber of the cylinder and function as catalytic nucleophiles for peptide bond hydrolysis with, respectively, different substrate specificities (7, 9). Substrate access to these structurally sequestered catalytic sites occurs via gated pores in the center of the α -subunit rings. Gating of the substrate access pore is an important function of regulators that compose proteasome holoenzymes. Thus, in addition to conferring specific catalytic and regulatory properties on their respective holoenzymes, such as ATP-dependent degradation of ubiquitin-modified protein in the case of 19S/PA700 (10), most regulators license the complex for substrate hydrolysis by allosterically relieving constitutive pore occlusion upon binding to the apical surface of the 20S α -subunit rings (11–14).

PI31 (Proteasome Inhibitor of 31,000 Da), the product of the PSMF1 gene, was originally identified as an in vitro inhibitor of artificially activated 20S proteasome (15). PI31 has a globular N terminus (FP domain, residues 1-151) and a natively disordered, proline/glycine-rich C terminus (residues 152-271) that is necessary and sufficient for proteasome inhibition (Figs. 1A and S1) (16). In contrast to its welldocumented in vitro function, the physiologic role of PI31 has been uncertain and the subject of seemingly conflicting results. For example, neither overexpression nor genetic reduction of PI31 in mammalian cells affects overall proteasome activity in a manner expected of a general proteasome inhibitor (17, 18). Such results, as well as uncertainty about cellular roles of regulator-free 20S proteasome, suggest that PI31's in vitro proteasome inhibition may lack physiologic significance. Moreover, other studies indicate that PI31 is a positive regulator of cellular proteasome function via various mechanisms whose generality and relationship to one another are not readily apparent (18-21).

To gain further insight into physiologic roles of PI31, we solved a high-resolution cryo-EM structure of a mammalian PI31-20S complex. The structure shows that the natively unstructured C terminus of PI31 enters the central 20S catalytic chamber through the substrate access pores and interacts

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Figure 1. Cryo-EM structure of bovine 20S proteasome complexed with human PI31. *A*, domain structure of human PI31, showing residues that interact with β 5 and β 2 catalytic sites (described in text). *B*, EM map of the human PI31-bound bovine proteasome at average resolution of 2.54 Å. The well-resolved densities belonging to the two PI31 C-terminal domains (CTDs) are shown in orange and green, respectively. *C*, superposition of the PI31 atomic model with the EM density. *D* and *E*, the EM density around the β 2 (*D*) and the β 5 active site (*E*). *F*, cartoon view of the two modeled PI31 CTDs. One PI31 CTD runs from the β s site of one β -ring to the β 2 and β 1 active sites of the other β -ring. *G*, *B*-factor plot showing that PI31 CTDs around the β 2 and β 5 sites are more stable. The highest and lowest B-factors are in *red* and *blue*, respectively.

with catalytic sites in a manner consistent with inhibition of peptide bond hydrolysis. While our work was in progress, a comparable structure was reported for a complex between a mutant, open-gate yeast 20S proteasome and Fub1, a yeast PI31 ortholog (22). More recently, the structure of a complex between 20S proteasome and a truncated PI31-like protein has been described from dormant spores of *Microsporidia* (23). Although these structures are in agreement regarding a general mechanism by which PI31 interacts with and inhibits the proteasome, they have a number of specific significant differences including the paths by which the PI31 polypeptide chain interacts with individual catalytic sites and the identities of PI31 residues involved in these interactions. We have also identified conditions that manifest PI31 inhibition of proteasome function in mammalian cells and provide new context and explanations for results of earlier reports. Our work reveals new insights about the molecular and cellular roles of PI31 in regulation of proteasome function.

Results

The C-terminal domain of human PI31 enters the central chamber of mammalian 20S proteasome and interacts directly with catalytic sites

We determined the cryo-EM structure of a mammalian PI31-20S proteasome complex after incubating recombinant human PI31 (hPI31) with the purified latent bovine 20S proteasome featuring a predominately closed-gate conformation (24). Human and bovine PI31 have 91% sequence identity and function indistinguishably in inhibition assays (Fig. S1 and refs (15, 16)) The maps of PI31-free and PI31-bound proteasome were refined to the resolutions of 2.23 Å and 2.54 Å, respectively (Figs. 1B, S2 and S3; Table S1). After model building, extra densities were identified as C-terminal domain residues (aa 189-253) of hPI31 located in the interior of the 20S particle (Fig. 1, B and C). The hPI31 densities near the β 2 and β 5 active sites within the central chamber of 20S were especially well resolved (Fig. 1, D and E). Portions of two hPI31 C-terminal domains were identified within a given 20S particle and featured the same C2 symmetry characteristic of the β -subunit rings. However, the C1 map was used for PI31 model building to avoid average deviations. The modeled structure is shown as a linearized hPI31 chain extending from the vicinity of the β 5 active site of one β -ring and then crossing the interface of the two β -rings toward the β 2 and β 1 active sites of the opposing β -ring (Fig. 1F). Although some hPI31 densities are present near the $\beta 1$ site, we were unable to build a model at this location due to the disconnection of the densities. Because the end of the modeled hPI31 is near the β 1 site, we predict an interaction between this site and the C terminus of hPI31. Biochemical analyses described below support this prediction. The C-terminal domain of hPI31 (aa 152-271) is composed of 35% proline and glycine residues and is structurally disordered (16). Accordingly, there is no discernable PI31 secondary structure in the resolved complex. The lack of hPI31 secondary structure indicates that the C terminus of hPI31 is flexible and needs to be stabilized by the surrounding residues of the proteasome. The b-factor plot demonstrates that the hPI31 sequences near the $\beta 2$ and $\beta 5$ active sites are the most stable parts in the modeled hPI31 structure (Fig. 1G), suggesting that the interactions between hPI31 and the proteasome active sites constitute the basis for the inhibitory activity of PI31.

The structure of this mammalian PI31–20S complex is generally similar to that of a recently reported complex between Fub1, the yeast PI31 ortholog, and an α 3 Δ yeast mutant 20S proteasome (22). In each structure, two copies of the hPI31/Fub1 C-terminal domain reside within the central chamber of 20S where they appear to interact with each of the three distinct catalytic sites. Notably, the regions of hPI31/ Fub1 involved in these interactions represent the most highly conserved regions of the two proteins (Fig. 2, *A* and *B*). Despite these similarities, there are also a number of differences. For example, in the yeast structure, Fub1 starts (N to C terminus) from the β 1 active site and then extends over the center of the β -ring to the β 5 active site where it dimerizes with the second copy of Fub1. Fub1 then contacts the β 5 active site of the

opposing β -ring and returns to the β 2 active site of original β ring. In contrast, hPI31 does not traverse the center of proteasome in the mammalian complex. Instead, it begins from the β 5 active site of one β -ring and curls toward the β 2 active site of the opposing β -ring (Fig. 2, C and D). Thus, although PI31 and Fub1 have conserved active-site interacting regions, the structural arrangement of these interactions appears to be distinct (Fig. 2, B-D). Fub1 blocks the β 5 activity through a unique conformation between two separate Fub1 molecules, whereas hPI31 achieves this inhibition using two conserved sequences (aa 194-200 and aa 206-212) within the same chain near the $\beta5$ active site. Specifically, the antiparallel hydrogen bonding between Gly194-Gly195-Glu196 and Gly206-Gly207 creates a conformation that fits into the \$\beta5\$ binding site (Fig. 2E). Human PI31 mimics the substrate binding, P4-P3-P2-P1-P1'-P2'-P3'-P4' to S4-S3-S2-S1-S1'-S2'-S3'-S4', to inhibit the proteolytic activity of proteasome. At the $\beta 2$ active site, residues 242 to 246 of hPI31 occupy the β 2 pocket in an antiparallel direction with Asp244 (P1) and Arg242 (P3) inserting into the S1 and S3 binding pockets, respectively (Fig. 3A). The hPI31 residues Arg242, Asp244, Pro245, and Phe246 are hydrogen bonded with the surrounding residues Thr1, Gly47, Ala49, Gly128, and Ser129 of B2 subunit to stabilize the binding sequence (Fig. 3*B*). At the β 5 active site, the dipeptide Asp197-Leu198 and residues 208 to 214 of hPI31 fit into the \$65 binding pocket in a parallel direction with the Met208(P1) and Ile209(P3) inserting into the S1 and S3 binding pockets (Fig. 3C). The hPI31 residues Asp197, Met208, and Ile209 are hydrogen bonded with the surrounding residues Thr1, Thr21, Gly47, Ala49, and Ser130 of \$5 subunit to sustain the unique binding conformation (Fig. 3D). These results show that hPI31 interacts with each of the proteasome's catalytic sites in a manner likely to directly inhibit substrate catalysis.

PI31 interacts with catalytic sites of the proteasome in a manner that prevents its own degradation

The ability of PI31 to stably inhibit the proteasome requires that PI31 itself is resistant to degradation. Several features of the hPI31 bound to the β 2 active site may contribute to this resistance. First, Lys33 of the B2 subunit was proposed to promote the deprotonation of the catalytic Thr1 and to generate the oxyanion for nucleophilic attack (25, 26). The hydrogen bond between functional carboxyl oxygen (OD1) of Asp244 of hPI31 and the hydrogen of the hydroxyl group (HG1) of Thr1 may impede the deprotonation of HG1 from the gamma oxygen (OG1) of Thr1 by Lys33 and prevent the formation of the threonine nucleophile. Second, the trypsinlike activity of the $\beta 2$ active site has a preference for basic residues at the P1 site. In the structure of the PI31-B2 site, P1 is occupied by Asp244, which is unfavorable for hydrolysis. According to the Keil rule, trypsin can cleave after Arg or Lys but not at those sites before Pro (27). Therefore, Pro245 at the P1' site likely hampers the trypsin-like activity toward PI31 at the $\beta 2$ active site. Third, the distance between the cleavage site (P1'-P1) and the catalytic Thr1 is 6 Å (Fig. 3B), which further



Figure 2. Similarities and differences between PI31 and Fub1 in 20S binding. *A*, sequence alignment of the C-terminal regions of human PI31 (aa 159–271) and yeast Fub1 (aa 134–250). Structurally conserved sequences are show in red brackets. *B*, structural alignment of the C-terminal regions of human PI31 and yeast Fub1. The β 5 subunit of the human PI31–bovine proteasome structure was aligned with that of Fub1– α 3 Δ yeast proteasome mutant structure. Only the regions near the β 2 and β 5 active site are structurally conserved. The structurally conserved regions are shown in *red squares. C*, zoomed view of the aligned structures of PI31 and Fub1 near the β 2 site. *D*, zoomed view of the aligned structures of PI31 and Fub1 near the β 5 site. Instead, PI31 forms a unique conformation within the same molecule to block the β 5 site. *E*, the special conformation of PI31 near β 5 is constructed by the antiparallel hydrogen bonding between the main chains of the two conserved sequences, Gly194-Gly195-Glu196 and Gly206-Gly207.

restrains the proteolytic activity of the β 2 active site. At the β 5 site, Asp197 of PI31 forms a hydrogen bond with the β 5 catalytic Thr1 (Fig. 3*D*). Like the hydrogen bonding between Asp244 and Thr1 of β 2, the hydrogen bond between Asp197 and Thr1 of β 5 may prevent the formation of threonine nucleophile at the β 5 site. In addition, the unique antiparallel conformation formed by Gly194-Gly195-Glu196 and Gly206-Gly207 of PI31 forces Asp197 to sit at the S1' site and residues 208 to 214 to bind to β 5 pocket in a parallel direction, which is in the reverse direction of normal substrate binding. This binding profile leaves no peptide bond between P1 (Met208) and P1' (Asp197) and renders PI31 unfavorable for hydrolysis at the β 5 site.

The C-terminal domain of hPI31 breaches the closed-gate conformation of wildtype mammalian 20S proteasome

The mammalian PI31–20S complex structure reported here was obtained with latent wildtype 20S proteasome featuring a constitutively closed-gate conformation. This form of the proteasome exhibits low protease activity in the absence of endogenous gate-opening activators, raising questions about how hPI31 gains access to the structurally sequestered 20S catalytic sites. Although PI31 was present in 5-fold molar excess of 20S in samples prepared for cryo-EM analysis, only 34% of 20S particles contained extra densities after extensive 3D classification (Fig. S3). This finding suggests that structural and/or functional features of these



Figure 3. Pl31 mimics substrate binding to the proteasome active sites. *A*, binding of hPl31 to the β 2 active site. The side chains of Asp244 (P1) and Arg242 (P3) are inserted into the S1 and S3 binding pockets, respectively. The Pro245 is at the P1' position. *B*, Pl31 in the β 2 active site was stabilized by the surrounding amino acids of the β 2 site. The distance between Thr1 and the cleavage site is 6 Å and may contribute to the proteolytic resistance to the β 2 site. *C*, binding of hPl31 to the β 5 active site. The side chains of M208 (P1) and I209 (P3) are inserted into the S1 and S3 binding pockets, respectively. Asp197 is at the P1' position. *D*, hydrogen bonds between Pl31 and the surrounding amino acids of β 5 help to stabilize the bound special conformation of Pl31 in β 5. The special conformation leaves no peptide bond between the P1 (Met208) and P1' (Asp197) for the β 5 activity.

proteins limit their interaction. It is possible that the C terminus of PI31 enters the proteasome during transient openings of the predominately closed-gate conformation during the 20S-PI31 preincubation (28, 29). Alternatively, our preparations of bovine 20S proteasome might contain low levels of constitutively activated (i.e., open-gate) 20S known to be promoted by certain purification conditions (30). Either possibility would be consistent with the relatively low efficiency of complex formation revealed by 3D classification even after prolonged incubation of 20S with a molar excess of PI31. In either case, we predicted that hPI31 should be a much more efficient inhibitor of open-gated 20S. To test this prediction, we compared PI31 inhibition of 20S in the presence and absence of low concentrations of SDS, commonly used to enhance proteasome activity by opening the substrate access gates. In fact, when 20S was activated by low concentrations of SDS, PI31 inhibited proteasome activity to a much greater extent than with latent 20S (Fig. 4). These results suggest that PI31 readily gains access to the proteasome's catalytic sites when its C terminus passes through the central pore of the apical α -subunit rings.

Individual catalytic subunit types are inhibited by specific sequences of hPI31

The structure of the hPI31-20S complex described above predicts hPI31 residues likely to be involved in inhibition of specific catalytic sites. To test these predictions, we first expressed and purified hPI31 truncation mutants that contain or lacked various combinations of these residues and tested these mutants in activity assays specific for each of the proteasome's three catalytic subunit types (Fig. 5A). As expected, both full-length wildtype hPI31 and a truncation mutant lacking the FP domain (residues 1-151) but containing the entire C-terminal domain (residues 152-271) inhibited the catalytic activity of each subunit type by >95% (Fig. 5B). Similarly, a mutant consisting of residues 1 to 261 inhibited the activity of each catalytic subunit type. In contrast, a mutant consisting of residues 1 to 181 (i.e., residues N-terminal to any observed catalytic site-interacting residues of PI31) had no inhibitory activity. These results confirm that PI31 amino acids responsible for inhibition of proteasome active sites reside between residues 182 and 262. Accordingly, PI31 variants 1 to 244 and 152 to 244 each inhibited β 5 activity but not β 1



Figure 4. Open-gated 20S proteasome is inhibited more effectively than closed-gate proteasome by PI31. Purified wildtype hPI31 was incubated with purified latent 20S proteasome in the absence or presence of 0.03% SDS prior to assay of proteasome activity with peptide substrates specific for indicated catalytic subunits, as described in Experimental Procedures. Proteasome activity in the absence of SDS or PI31 was set to 100, and all other activities are expressed as a percentage of that value. Bars represent mean values of indicated triplicate assays. Differences in activity were analyzed by two-way ANOVA, and multiple comparisons within each catalytic subunit dataset were tested by Sidak's multiple comparisons test. β 1: control versus PI31 (-SDS) p = 0.0263; PI31 (-SDS) versus PI31 (+SDS) p < 0.0001; β 5: control versus PI31 (-SDS) p = 0.651; PI31 (-SDS) versus PI31 (+SDS) p < 0.0001; β 5: control versus PI31 (-SDS) p = 0.651; PI31 (-SDS) versus PI31 (+SDS) p < 0.0001. Similar data were obtained in at least four independent experiments.

activity, as predicted. However, neither of these latter mutants inhibited $\beta 2$ activity, as expected, but instead stimulated $\beta 2$ catalyzed hydrolysis. This surprising result suggests that these truncated proteins lack residues required for the stable orientation of the amino acid(s) that contact(s) the active site β2 threonine. The differential effects of these PI31 mutants on specific types of proteasome activities were also demonstrated in an orthogonal assay using the activity probe, Me₄BodipyFL-Ahx₃Leu₃-VS (Fig. 5C). Me₄BodipyFL-Ahx₃Leu₃-VS covalently binds to each of the proteasome's catalytically active sites, and the resulting fluorescently tagged subunits can be detected and identified after SDS-PAGE (31). Thus, inhibitors of any catalytic site will reduce labeling of that respective subunit. As expected, both full-length, wildtype and the Cterminal domain (residues 152-271) PI31s greatly reduced labeling of each proteasome subunit type, whereas PI31¹⁻¹⁸¹ had no effect on labeling of any subunit. In contrast, a mutant consisting of residues 1 to 261 inhibited labeling of each catalytic subunit type (Fig. 5C). These functional results are consistent with the structural predictions for identification of interaction sites between PI31 and individual catalytic subunits that cause proteasome inhibition and indicate that despite a clear structural resolution of PI31 residues in contact with the β1 catalytic site the latter interaction occurs between PI31 residues 245 and 261 (Fig. 5A).

To further refine the functional specificity of interactions between individual 20S catalytic sites and corresponding PI31 residues involved in inhibition, we expressed and purified hPI31 with point mutations (D to A) at residues 197 and 244. Our structure shows that D197 and D244 interact directly with the catalytic threonine residues of β 5 and β 2 proteasome subunits, respectively (Fig. 3). Because of the unresolved nature of the interaction between hPI31 and the β 1 site, we

allow an analogous test for this site. Consistent with our predictions, hPI31^{D197A} inhibited β5-specific peptide hydrolysis 3 times less well than wildtype hPI31 but had little or no effect on the inhibition of hydrolysis of β1- or β2-specific substrates compared with wildtype hPI31 activities (Fig. 6A). A similar β5-specific diminution of PI31 inhibition by the hPI31^{D197A} mutant was observed in an assay using the activesite probe, Me₄BodipyFL-Ahx₃Leu₃-VS (Fig. 6B). Likewise, hPI31^{D244A} had significantly weaker inhibitory activity against β2-specific peptidase activity but did not affect inhibition of the β 5-specific substrate hydrolysis (Fig. 6A). This latter mutant also blocked normal PI31 inhibition of \u03b31-specific substrate hydrolysis to a lesser extent. Although the basis of this effect is unclear, we note that the proteasome's catalytic sites are known to have interdependent allosteric effects (32). Therefore, selective inhibition at one or more catalytic sites may affect activity at other sites.

cannot confidently predict a specific mutation that would

Monomeric PI31 is sufficient for proteasome inhibition

Previous studies established that PI31 forms homodimers mediated by its N-terminal FP domain (16, 33). This finding raises questions about whether the two C-terminal PI31 polypeptide chains observed in the catalytic chamber of 20S originate from PI31 dimers entering through a single a-subunit ring, from monomers entering through opposite α -subunit rings, or from selective entrance of only one of a dimer's C termini from opposite α -subunit rings. To distinguish among these possibilities, we compared the inhibitory activity of PI31 variants with differing abilities to dimerize. First, we used mass photometry to confirm conclusions from previously reported centrifugation analysis that wildtype hPI31 exists in a monomer-dimer equilibrium (ref. (16) and Fig. 7A). At a concentration estimated to approximate that found in cells, hPI31^{WT} featured nearly equal amounts of monomer and dimer (2183 and 2064 mass photometry counts, respectively). Next, we expressed and purified hPI31 containing a point mutation (V6R) in the FP domain predicted to disrupt PI31 dimerization (33). The V6R mutation significantly shifted the monomer-dimer equilibrium in favor of monomeric PI31 (1766 monomer counts versus 751 dimer counts) (Fig. 7B). Finally, we expressed hPI31¹⁵²⁻²⁷¹, which lacks the FP dimerization domain and is assumed to be monomeric. Unfortunately, mass photometry cannot be used to confirm the monomeric state of hPI31¹⁵²⁻²⁷¹ (a predicted 13-kDa protein) due to limitations of lower mass detection by this method. Therefore, we used chemical cross-linking followed by SDS-PAGE to further evaluate and compare the relative oligomeric states of hPI31^{WT}, hPI31^{V6R}, and hPI31^{152–271} (Fig. 7*C*). As expected, cross-linking of hPI31^{WT} produced detectable protein at a size (~60 kDa) consistent with dimeric PI31. A significantly reduced amount of crosslinking-dependent protein was detected for the hPI31^{V6R} mutant, consistent with the predominately monomeric nature of the protein shown by mass photometry. Importantly, crosslinking of hPI31^{152–271} produced no detectable products, a



Figure 5. Identification of PI31 interaction sites with different 20S proteasome catalytic subunit types. *A*, recombinant wildtype and indicated truncation mutants of hPI31 were expressed and purified as described in Experimental Procedures. Mutants are depicted schematically and show proposed interaction sites for β 5, β 2, and β 1 catalytic sites (see Fig. 1*A*). The proposed interaction site for the β 1 subunit is indicated in blue. *B*, 20S proteasome (50 nM) activity for each catalytic subunit type was measured in the absence (–) or presence of wildtype and indicated hPI31 mutants at 10- to 20-fold molar excess. Proteasome activity for each subunit type in the absence of PI31 (–) was set to 100, and other activities were expressed as a percentage of that value. Bars represent mean values (±SD) of indicated triplicate assays. Similar results were obtained in three independent experiments. Differences in proteasome activity among PI31 treatments were analyzed prior to normalization by ANOVA and Tukey's HSD. For β 1 activity, pairwise differences between all treatments were observed at *p* < 0.0001; for β 2 activity, pairwise differences between all treatments were observed at *p* < 0.001; for β 2 activity, pairwise differences between all treatments were observed at *p* < 0.001 except for (–) PI31 *versus* 1 to 181 (*p* = 0.0456) and WT *versus* 152 to 244 (*p* = 0.7878). C, activated 20S proteasome (115 nM) was preincubated in the absence (–) or presence of indicated hPI31 variants (10 μ M) and then labeled with Me₄BodipyFL-Ahx₃Leu₃-VS probe (500 nM) for 15 min to avoid signal saturation. Assays were quenched with SDS-PAGE sample buffer, and samples were subjected to SDS-PAGE and imaging as described in Experimental Procedures.

result consistent with the conclusion that this mutant is completely monomeric. Despite their differing oligomeric states, each of these hPI31 variants was competent for proteasome inhibition. Surprisingly, however, inhibitory activities were indistinguishable when assayed at comparable $[M]_T$ (where $[M]_T = [monomer] + 2 \times [dimer]$) (Fig. 7*D*). Nevertheless, a direct comparison of the relative inhibitory effects of these proteins is complicated by the nonhomogenous oligomerization state of PI31^{WT} and PI31^{V6R}. Assuming [monomer]/[M]_T ratios of 0.32, 0.54, and 1.0 for hPI31^{WT}, hPI31^{V6R}, and hPI31^{152–271}, respectively, hPI31^{WT} achieves 50% inhibitory activity at a lower amount of functional monomers than hPI31^{V6R} and hPI31^{152–271} ([monomer] = 64 nM, 108 nM, and 200 nM, respectively). Thus, inhibitory activity may be a function of total subunit number ([M]_T) rather than oligomerization state. Although we favor a model in which the 20S–PI31 complex structure determined here results from two PI31 monomers that enter the 20S catalytic sites from opposite ends of the 20S cylinder, we cannot exclude related models including those that feature the inhibitory action of PI31 dimers (Fig. 8*D*; see Discussion). Additional work will be required to resolve this issue.



Figure 6. Identification of individual PI31 residues involved in sitespecific inhibition of proteasome activity. A, 20S proteasome (143 nM) activity for each indicated catalytic sites was measured as described in Experimental Procedures following incubation with 10-fold molar excess of PI31 (1.4 μ M). Activity in the absence of PI31 was set to 100, and other activities are expressed as a percentage of that value. Bars represent mean values (±SD) of indicated triplicate assays. Similar results were obtained in three independent experiments. Differences in proteasome activity were analyzed by one-way ANOVA and Tukey's HSD posthoc test. p-Values are shown for significant differences between PI31 treatments. B, 20S proteasome (143 nM) was activated by SDS and incubated in the absence or presence of indicated variants of PI31 (1.4 µM). Samples were then labeled with Me₄BodipyFL-Ahx₃Leu₃-VS activity-based probe (1 µM) for 40 min. Assays were guenched with SDS-PAGE sample buffer, and samples were subjected to SDS-PAGE and imaging as described in Experimental Procedures.

PI31 overexpression inhibits proteasome activity in mammalian cells

The structure of the PI31-20S complex and consequent proteasome inhibition described above raises questions about their physiologic significance. Previous work by us and others has failed to demonstrate an expected inhibitory effect of PI31 overexpression on proteasome activity in mammalian cells (17, 18). However, global quantitative proteomic analysis of multiple mammalian cell lines documents that PI31 is normally expressed at low levels in most cells and is exceeded on a molar basis by 20S proteasome by 50- to 100-fold (34-37). Retrospective analysis of our previous experiments reveals that PI31 overexpression in those studies was only 3- to 10-fold over endogenous levels, making it unlikely that any effect on proteasome activity could be detected since PI31 would remain significantly substoichiometric to the proteasome. We have repeated those experiments under conditions that express PI31 by >700 fold over endogenous levels and in >20fold molar excess of proteasomes (Fig. S4). Under these conditions, overall proteasome activity in cell extracts was reduced by 25 to 30% as monitored by proteasome-dependent hydrolysis of peptide substrates (Fig. 9). Although additional experiments will be required to determine the exact basis for and specificity of this effect, these data are consistent with a model in which PI31 can bind to and inhibit cellular proteasome.

Discussion

The high-resolution structure of the mammalian PI31-20S proteasome complex reported here provides a structural basis for our original discovery and description of PI31 as an in vitro inhibitor of 20S proteasome activity (15). The structure is in accord with mutational analysis showing that the C-terminal domain of PI31 is necessary and sufficient for inhibitory activity (Fig. 5 and (16)). This mammalian structure and two recently published structures of yeast and Microsporidia PI31like proteins in complex with 20S proteasome reveal a surprising mechanism of inhibition via direct interaction of PI31 with the proteasome's catalytic sites (22). This mechanism differs significantly from a previously anticipated model of inhibition in which the C terminus of PI31 binds to the exterior of the proteasome near the substrate access pore and sterically blocks the entrance of substrates to the catalytic chamber (16, 38). Rationale for this model was reinforced by recognition that PI31 contains a C-terminal HbYX motif found in various proteasome regulators that bind to HbYX-cognate sites between outer ring α -subunits (17, 39, 40). However, as discussed below, the functional relationship between the hPI31's HbYX motif (residues 269-271) and the structure of the hPI31-20S complex remains uncertain.

Unlike the unstructured proline/glycine-rich C-terminal domain, the N-terminal FP domain of hPI31 is well folded (33, 41) and therefore cannot transit the narrow substrate access pore regardless of its gating status. Our previous data showed that this domain neither binds to nor inhibits the 20S



Figure 7. PI31 monomers are sufficient to inhibit 20S proteasome activity. Mass photometry analysis of PI31 dimerization using purified, recombinant (*A*) hPI31^{WT} and (*B*) PI31^{VGR} dimerization mutant proteins. Proteins were diluted to 40 nM total monomer (1.2 ng μ L⁻¹) in PBS during recording and analyzed as described in Experimental Procedures. *C*, analysis of PI31 (WT, V6R, 152–271) dimerization by chemical cross-linking. Protein, 23 μ M, was cross-linked with 0.05% glutaraldehyde or buffer controls as described in Experimental Procedures. One microgram of denatured proteins was separated by SDS-PAGE and blotted against PI31 antibodies. *D*, inhibition of proteasome activity by hPI31^{WT}, hPI31^{VGR}, and hPI31^{52–271}. 20S proteasome (50 nM) was activated by 0.03% SDS prior to incubation with indicated amounts of total PI31 subunits, [M]_T (where [M]_T = [monomer] = 2 x [dimer]), up to 10-fold molar excess. Proteasome activities were measured in duplicate as described in Experimental Procedures and expressed as percentages relative to controls lacking PI31.

proteasome (16). No density for the FP domain was detected in our high-resolution electron microscopy (EM) map of the hPI31–20S complex, likely due to its flexibility on the exterior of the proteasome. However, a low-pass filter of the map shown at a lower display threshold reveals densities on the apical surface of 20S α -rings of hPI31-bound proteasome (Fig. 8, *A* and *B*). We also examined the difference map between the PI31-bound and PI31-free EM maps and observed hPI31 densities in the 20S antechambers as well as in the catalytic chambers. These findings indicate that the C-terminal regions of PI31 in the catalytic chamber are threaded from the antechambers (Fig. 8*C*). This observation is consistent with the proposed orientation of active site engagement along the primary structure of the hPI31 C terminus, as described above and discussed further below.

Although the mammalian and yeast PI31/Fub1–20S complexes share general structural features, they have a number of significant distinctions. Importantly, the mammalian complex described here was achieved using native 20S proteasome as opposed to a mutant 20S proteasome with a constitutively open gate. This result demonstrates the capacity of PI31 to inhibit a physiologic form of 20S proteasome normally found in cells and

rules out effects that might result from an artificial α -ring. The manner by which the polypeptide chains of hPI31/Fub1 interact with the proteasome's multiple catalytic sites also differ appreciably. The Fub1 chain traverses a path (in the N-to-C direction) from $\beta 1$ to $\beta 5$ in the opposing β ring and back to $\beta 2$ in the original β ring. Moreover, Fub1's interaction with and proposed inhibition of β5 employs the cooperative action of a second Fub1 molecule. In contrast, our data show a hPI31 interaction path (in the N-to-C direction) from $\beta 5$ to $\beta 2$ to $\beta 1$. As with Fub1, hPI31 interacts with $\beta 5$ on the opposing ring from that of its interactions with β 1 and β 2. However, unlike Fub1, hPI31 utilizes two different segments of the same polypeptide chain for its $\beta 5$ interaction. This feature indicates that a single molecule of hPI31 is sufficient to engage one copy of each β -type catalytic subunit. The functional consequence of inhibition of only one of two copies of each catalytic subunit type is unknown. Interestingly, an interaction path similar to that of the mammalian proteins has been observed for a PI31-like peptide and 20S proteasome from Microsporidia (23). Nevertheless, the finding that PI31 orthologs from evolutionally diverse organisms directly contact proteasome catalytic sites suggests a highly conserved in vivo inhibitory function for this interaction.



Figure 8. Proposed model for hPI31 interaction with 20S proteasome. *A*, the low-pass filtered EM map of 20S proteasome surface rendered at a low threshold of $\sigma = 2$. *B*, the low-pass filtered EM map of PI31-bound 20S proteasome surface rendered at a low threshold with $\sigma = 2$. *C*, difference map (*pink* surface) between the two above-mentioned low-pass filtered EM maps revealing extra PI31 densities inside both catalytic and antechambers. The 20S atomic model is superimposed to indicate the chamber location of the difference densities. *D*, alternative models of the PI31–20S proteasome complex involving either monomeric (*left*) or dimeric (*right*) PI31 showing the interaction of the PI31 C-terminal domain with proteasome catalytic sites (see text for details).

hPI31 can form homodimers mediated by the N-terminal FP domain (16, 33, 41) but exists in a monomer-dimer equilibrium at PI31 concentrations likely found in cells (Fig. 7 and (16)). We favor a model in which the hPI31-20S complex results from 20S interaction with PI31 monomers (Fig. 8D, left). This feature might contribute to the observation that high molar ratios of PI31 to 20S are required for inhibition in vitro. It is also consistent with the conclusion that the two C termini residing in the catalytic chamber enter from opposite ends of the proteasome. Nevertheless, we cannot exclude models of inhibition involving PI31 dimers (Fig. 8D, right). In one variant of this model, transit through the substrate access pore may be limited to one of the two C termini while the HbYX motif of the other subunit promotes gate opening via α-subunit binding. Although such a mechanism is consistent with previously shown gate-opening properties of hPI31 C-terminal peptides, it cannot be obligatory because PI31 lacking HbYX residues retains inhibitory activity (Fig. 5 and (16, 17)). Alternatively, both C termini of a PI31 dimer may enter the antechamber

through the same pore while only one gains access to the central catalytic chamber. Interestingly, previous work has shown that the open proteasome pore can accommodate two unstructured polypeptide chains (42). Additional work will be required to distinguish among these and related models and to determine their functional implications.

In its position on the apical surface of the α -rings, the FP domain of PI31 may serve as a binding partner of other proteins. In fact, FbxO7, an F-box protein whose mutant forms are responsible for early-onset autosomal recessive Parkinson's disease, has been identified as a PI31 binding partner and this interaction is mediated by the FP domains of both proteins (33, 43, 44). Likewise, the FP domain is likely involved in a proposed role for PI31 as an adaptor between the proteasome and motor proteins for axonal transport of proteasomes to intracellular sites of action (21). The relationship between PI31-mediated proteasome inhibition and such functions is unclear and would appear to require release of inhibition upon delivery. Although we have not identified *in vitro* conditions that reverse



Figure 9. Overexpression of hPI31 inhibits proteasome activity in HepG2 cells. HepG2 cells were transfected with pCMV3-FlagPI31 or vehicle controls as described in Experimental Procedures. Extracts from control and PI31 overexpression conditions were prepared and assayed for proteasome activity using peptide substrates for each of the indicated catalytic subunits, as described in Experimental Procedures. Bars are mean (\pm SD) activities from n = 4 lysates of independently transfected cells. Activity in control (-FlagPI31) extracts was set as 100, and other activities within the same experiment are expressed as a percentage of that value. Differences in activity were analyzed by two-way ANOVA, and multiple comparisons test. Similar results were obtained in three independent experiments.

PI31–proteasome binding and inhibition, such a process has been proposed as a mechanism for termination of PI31-likemediated proteasome inhibition that allows assembly of active 20S–19S/PA700 holoenzyme complexes after germination of dormant *Microsporidia* spores (23). The generality of the latter process is unknown and appears to be inconsistent with previous evidence that PI31 blocks the *in vitro* assembly and activation of proteasome holoenzymes with 19S/PA700 and PA28 regulators (16, 38). Alternatively, irreversible PI31 binding could result in clearance of aberrant or unneeded proteasomes, as was hypothesized for Fub1. Such a process might require binding of the FP domain with proteins involved in proteasome disposal and is consistent with a general role for the exposed FP domain in protein–protein interactions.

Because hPI31 normally is present in cells at much lower levels than the proteasome, it can affect only a small proportion of total cellular proteasome. Therefore, any physiologic effect of even full-engaged PI31 would escape detection in assays that measure global proteasome activity. In contrast, we have shown that proteasome activity was detectably inhibited when PI31 was ectopically expressed to levels exceeding those of the proteasome (Fig. 9). Although additional work is required to determine the exact mechanism of this effect, it is consistent with a direct inhibitory role for cellular PI31. We are unaware of any physiologic or pathologic condition in which PI31 levels are increased to such levels. Nevertheless, endogenous levels of PI31 may serve to terminate the activity of a subpopulation of proteasomes that function normally in a regulator-free fashion (45-48) or to prevent dysregulation of cellular proteostasis by inappropriately activated 20S proteasomes (22, 46).

Experimental procedures

Protein purification

20S proteasome was purified from bovine red blood cells as described (30). Recombinant wildtype human PI31 and PI31

truncation mutants were expressed in E.coli and purified as described (16). A pET28a(+) vector containing N-terminally tagged, wildtype hPI31 was utilized for site-directed mutagenesis. Primers and annealing temperatures for site-directed mutagenesis were predicted using the NEBaseChanger online tool (version 1.3.3, https://nebasechanger.neb.com/). Sitedirected mutagenesis was performed on 10 ng pET28a(+)-HisPI31^{wt} template with the Q5 Site-Directed Mutagenesis Kit (New England BioLabs) according to manufacturer's instructions. Based on the length of the plasmid, an extension time of 2.5 min was used for each PCR cycle. The mutagenesis PCR product was ligated using the Kinase Ligase DpnI (KLD) mix provided in the kit. Ligated plasmid $(1 \mu l)$ was then transformed into One Shot BL21 Star (DE3) competent cells (Invitrogen) with outgrowth in SOC media for 1 h at 37 °C before plating on LB agar plates containing kanamycin (50 µg/ ml). Sequences were confirmed by Sanger sequencing at the UT Southwestern McDermott Center Sequencing Core with Big Dye Terminator 3.1 and capillary instruments (Applied Biosystems Inc).

Recombinant His-PI31 mutant proteins were purified by metal affinity chromatography. BL21 Star (DE3) cells containing pET28a(+) HisPI31 mutants from monoclonal stab cultures were used to inoculate 150 ml LB broth containing 50 μg/ml kanamycin. Cells were grown with shaking overnight at 37 °C. The following morning, 100 ml of LB/Kan broth was added to cultures with 250 to 500 µM IPTG to induce expression. Cells were grown at 37 °C for an additional 3 to 6 h. Following expression, cells were pelleted by centrifugation at 10,000g for 10 min at 4 °C. Cell pellets were resuspended in 8 ml Binding Buffer (50 mM sodium phosphate, pH 7.6, 500 mM NaCl, 10 mM imidazole, pH 8.0) containing 0.1 mg/ ml lysozyme and 1x cOmplete, EDTA-free protease inhibitor cocktail (Roche). Cells were incubated on ice for 30 min before sonication. Lysed cells were then centrifuged at 14,000 rpm, 20 min, 4 °C. The 8 ml supernatant was applied to 1 ml packed volume of Ni-NTA His-Bind Resin (Millipore Sigma). Binding was performed with gentle mixing for 1 h at 4 °C and loaded onto a Poly Prep Chromatography Column (Bio-Rad). The loaded resin was then washed 3 to 4 times with 6 ml Wash Buffer (50 mM sodium phosphate, pH 7.6, 500 mM NaCl, 20 mM imidazole, pH 8.0) until protein was undetectable with Bradford Reagent (Pierce). His-PI31 was eluted in four 0.5-ml fractions with 250 mM imidazole, 50 mM sodium phosphate, pH 7.6, 500 mM NaCl. Fractions containing peak protein (as assessed by Bradford assay) were pooled and dialyzed against Buffer H (20 mM Tris-HCl, pH 7.6 at 4 °C, 20 mM NaCl, 1 mM EDTA, 5 mM β-mercaptoethanol) containing 10% glycerol. Purification was confirmed by SDS-PAGE Coomassie staining and Western blotting with anti-His antibody.

Proteasome activity assays

Inhibitory activity of hPI31 proteins against bovine and human 20S proteasome was analyzed using 7-amino-4methylcoumarin (amc)-linked fluorogenic peptide substrates specific to each of the proteasome catalytic sites: caspase-like (β1, Z-LLE-amc), trypsin-like (β2, Ac-RLR-amc or Z-VVRamc), and chymotrypsin-like (β5, Suc-LLVY-amc). In some experiments, latent 20S proteasome was first activated with preincubation in 0.03% SDS at 37 °C for 5 min. PI31 (wildtype, D197A, or D244A) was then incubated for an additional 15 min at 37 °C. Proteasome activity was assayed by addition of peptide substrate (50 µM final concentration). Free amc produced by proteasome hydrolysis was monitored (Ex. 360 nm, Em. 460 nm) at 45-s intervals over 21 min at 37 °C (Synergy Mx Multi-Mode Microplate reader, BioTek). Proteasome activity was expressed as rates of amc production, and all were linear with respect to time and proteasome concentration in these assays. All assays were performed in triplicate and repeated independently at least three times. Differences between the effects of PI31 mutants on proteasome activity were analyzed by one-way ANOVA and Tukey's honestly significant difference (HSD) test using GraphPad Prism 9 (v9.5.0). PI31 mutant inhibitory activity was also assessed using direct fluorescent labeling of proteasome active sites with a $Me_4BodipyFL-Ahx_3Leu_3-VS$ activity-based probe (31). This probe covalently binds each of the β 1, β 2, and β 5 active sites and can be visualized following separation by SDS-PAGE. Proteasomes were first SDS-activated and incubated with PI31 as described above. Activity-based probe (0.5-1 µM final concentration) was then added for an additional 15 to 40 min incubation at 37 °C. Because the probe labels each subunit with different affinity, concentrations and incubation times were adjusted to avoid signal saturation of some subunits while maintaining detectability of others. Labeling was quenched by denaturing proteasomes in SDS-PAGE sample loading dye, and proteins were separated on a SDS-PAGE gel. Gels were either imaged directly using an Odyssey M Imager (LI-COR, 520 nm fluorescence channel) or transferred to nitrocellulose membranes and visualized on ChemiDoc MP Imager (Bio-Rad, fluorescein channel). Images were imported into Image Studio (LI-COR, v5.0.21) for quantification.

Western blotting and antibodies

Western blotting was conducted as described (17). Transferred membranes were imaged using a LiCor Odyssey CLx. Rabbit polyclonal antibody against human PI31 was purchased from Enzo (PW9710) and was used at 1:2000 dilution. Rabbit polyclonal antibody against human β 5 subunit of the proteasome was produced and characterized in the DeMartino lab as described (49, 50). Rabbit polyclonal antibody against human β tubulin was purchased from Cell Signaling Technology (2146) and used at 1:1000 dilution. Goat anti-rabbit IRDye680RD was purchased from Li-COR and was used at 1:10,000 dilution.

Overexpression of PI31 in mammalian cells

A mammalian expression vector, pCMV3-N-FlagPI31^{wt}, was obtained from Sino Biological Inc (catalog #HG17079-NF). HepG2 cells were seeded in six-well plates at 3×10^5 cells well⁻¹ and grown overnight at 37 °C, 5% CO₂ in Dulbecco's modified Eagle's medium containing 10% fetal bovine serum, penicillin, and streptomycin. On the following day, 750 ng of pCMV3-N-FlagPI31 vector was mixed with 5 μ l P3000 reagent diluted in OptiMEM medium and added to 5 μ l of diluted Lipofectamine3000 according to manufacturer's instructions (Life Technologies). Vectors or vehicle-only controls were added to replicate wells and transfected for 24 h. Following transfection, cells were washed three times with PBS and harvested in a hypotonic lysis buffer in the presence of ATP (20 mM Tris, pH 7.6 at 37 °C, 20 mM NaCl, 5 mM MgCl₂, 100 μ M ATP, 1 mM DTT, 0.1% NP-40). Cells were then passed through a 27½ G needle 15 times and centrifuged at 14,000 rpm, 20 min, 4 °C. Supernatants were divided into three sets of triplicate wells in a 96-well plate and assayed for proteasome activity using fluorogenic peptide substrates (β 1, Z-LLE-amc; β 2, Z-VVR-amc; and β 5, Suc-LLVY-amc) as described above for pure proteins.

Proteasome and PI31 amounts in HepG2 cells transfected with vehicle controls or pCMV3-N-FlagPI31 were analyzed by semiquantitative Western blotting using polyclonal antibodies against the proteasome β 5 subunit and the PI31 C terminus. Within the same gel, a range of pure proteins (bovine 20S proteasome or recombinant His-PI31) was used as standards for quantification. Secondary antibodies with IRDye (LI-COR) were used and near infrared signals were quantified on Odyssey CLx imager (LI-COR). Signals were converted to protein masses based on the standard curve of pure proteins and then converted to moles assuming values of 700 kDa for proteasome and 31 kDa for PI31 (Fig. S4). Amounts of each protein were then normalized to total protein content (determined by BCA protein assay, Pierce) to compare molar ratios of PI31 and proteasome in HepG2 cells.

Analysis of PI31 oligomerization by mass photometry

Purified His-PI31 (wildtype or V6R mutant) proteins were dialyzed against 20 mM Tris HCl (pH 7.6 at 5 °C), 20 mM NaCl for analysis of oligomerization by mass photometry (TwoMP, Refeyn). Protein concentrations were determined by BCA assay (Pierce) and converted to molar concentrations assuming 31 kDa molecular weights. PI31 was then diluted in 0.2 µm filtered PBS into a total monomer concentration of 400 nM. Cover slips were cleaned with isopropanol and MilliQ water and fitted with silicon gaskets. The mass photometer optics were focused on 16.2 µl buffer prior to the addition and mixture of 1.8 µl sample. Sample mixture into the prefocused droplet resulted in an additional one-tenth dilution (40 nM final), and recordings were initiated immediately after mixing. Bovine serum albumin and thyroglobulin oligomers were used as molecular weight standards to calibrate ratiometric contrast measurements to molecular weight. Automatic Gaussian curve fitting for peaks was performed by DiscoverMP software (Refeyn). which identified peaks centered at 36 to 37 and 56 to 59 kDa for monomers and dimers, respectively.

Cross-linking analysis of PI31 oligomerization

Purified hPI31^{WT}, hPI31^{V6R}, and hPI31^{152–271} proteins were dialyzed against 20 mM Hepes (pH 7.8), 20 mM NaCl to avoid interference of Tris buffers in cross-linking. Protein

concentrations were determined by Bradford assay, and samples were diluted to 23 μ M based on molecular weights of 31, 31, and 13 kDa for WT, V6R, and 152 to 271, respectively. Glutaralde-hyde or a buffer control was added to a final concentration of 0.05% and incubated for 15 min on ice. Cross-linking was terminated by the addition of 10 mM Tris-HCl, and samples were denatured in SDS sample buffer. Proteins were then separated by 12% SDS-PAGE and blotted against PI31 as described above.

Cryo-EM sample preparation and data collection

Cryo-EM was conducted at the Cryo-EM facility of Van Andel Institute. To obtain the human PI31-bovine proteasome complex, we mixed purified hPI31 and 20S at a molar ratio of 5:1 and incubated the mixture at 37 °C for 1 h in 20 mM Tris, pH 7.5, 5 mM MgCl₂, 100 mM KCl, and 1 mM DTT. The sample concentration was adjusted to 3 mg/ml for EM grid preparation. Cryo-EM grids were prepared in Vitrobot Mark IV (FEI) with the chamber set to 5 °C and 100% humidity. We first glow-discharged Quantifoil R 1.2/1.3 300 mesh gold holey carbon grids with O2-air for 30 s at 30 W in a Gatan Model 950 Advanced Plasma System, applied a droplet of 3 µl sample to the EM grid, waited for 1 s, blotted the grid for 4 s using a piece of 595 filter paper with a blotting force set to 3, then vitrified the sample by plunging the grid into liquid ethane precooled with liquid nitrogen. The cryo-EM dataset was recorded in a Titan Krios (FEI) equipped with a Quantum 967 energy Filter plus a post GIF K3 summit direct electron detector (A.K.A. BioQuantum). Movies were recorded using SerialEM software in super-resolution counting mode with a defocus range of -1.3 to -1.8 µm and a pixel size of 0.414 Å/pixel. The dose rate was 46 electrons/Å²/s with 1.3-s exposure time for a total dose of 60 electrons/Å². A total of 19,264 movies were collected during a 2-day session.

Cryo-EM data processing

Image processing was performed in Relion 3.1.1 and Cryosparc 3.3.2 (51, 52). We first corrected the beam-induced motion of each movie micrograph using MotionCor2 with binning factor of 2 in Relion (53). The corrected micrographs were imported into Cryosparc for further processing. We used CTFFIND4 to estimate the defocus value for each micrograph and used the detected value to correct for the contrast transfer effect. Based on the CTF detection limit, low-resolution micrographs were removed and a total of 18,528 micrographs were retained for particle picking. We first used blob picking to generate several templates for the next stage template-based particle picking. We used the proteasome particles from template picking and with good 2D class averages to train Topaz and extracted 1,573,299 particles by the trained Topaz. After reference-free 2D classification and subsequent 3D ab initio 3D reconstruction, a total of 1,467,640 particles were selected for 3D multireference heterogenous refinement. The 3D class with extra density inside the catalytic β -ring of the 20S proteasome was chosen for further 3D heterogenous refinement using the classified maps with and without extra density as the references. The particles belonging to the two 3D classes (with or without extra density) were segregated and refined separately, first by homogeneous refinement and followed by nonuniform refinement (51). By applying C1 symmetry during refinement, we obtained a 3D EM map of the 20S proteasome with extra chamber density (PI31) at 2.54 Å average resolution. We applied C2 symmetry to the 20S proteasome alone particles and refined a 3D EM map at 2.23 Å average resolution. The resolutions were estimated by applying a soft mask around the map of proteasome and used the gold standard Fourier shell correlation = 0.143 as the threshold.

Model building

Both 20S alone and the PI31–20S complex were modeled using the coordinates of bovine 20S proteasome (Protein Data Bank ID 1IRU) as the initial model. The PI31 structure was built manually in Coot after completion of the 20S proteasome modeling (54). Real space refinements were performed in Phenix to correct geometric errors, and the model quality was estimated using MolProbity in Phenix (55).

Data availability

The structures of the 20S proteasome and the 20S proteasome–PI31 complex were deposited in the Protein Data Base with accession codes 8FZ5 and 8FZ6, respectively. Any data not in the article can be shared upon request.

Supporting information—This article contains supporting information (16, 22, 23).

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Abbreviation—The abbreviations used are: EM, electron microscopy.

References

Hanna, J., Guerra-Moreno, A., Ang, J., and Micoogullari, Y. (2019) Protein degradation and the pathologic basis of disease. *Am. J. Pathol.* 189, 94–103

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- Lecker, S. H., Goldberg, A. L., and Mitch, W. E. (2006) Protein degradation by the ubiquitin-proteasome pathway in normal and disease states. *J. Am. Soc. Nephrol.* 17, 1807–1819
- **3.** Tanaka, K., and Matsuda, N. (2014) Proteostasis and neurodegeneration: the roles of proteasomal degradation and autophagy. *Biochim. Biophys. Acta* **1843**, 197–204
- 4. Türker, F., Cook, E. K., and Margolis, S. S. (2021) The proteasome and its role in the nervous system. *Cell Chem. Biol.* **28**, 903–917
- Rock, K. L., Gramm, C., Rothstein, L., Clark, K., Stein, R., Dick, L., et al. (1994) Inhibitors of the proteasome block the degradation of most cell proteins and the generation of peptides presented on MHC class I molecules. Cell 78, 761–771
- DeMartino, G. N., and Gillette, T. G. (2007) Proteasomes: machines for all reasons. *Cell* 129, 659–662
- Groll, M., Ditzel, L., Lowe, J., Stock, D., Bochtler, M., Bartunik, H. D., et al. (1997) Structure of the 20S proteasome from yeast at 2.4A resolution. *Nature* 386, 463–471
- Baumeister, W., Walz, J., Zühl, F., and Seemüller, E. (1998) The proteasome: paradigm of a self-compartmentalizing protease. *Cell* 92, 367–380
- 9. Bochtler, M., Ditzel, L., Groll, M., Hartmann, C., and Huber, R. (1999) The proteasome. *Ann. Rev. Biophys. Biomol. Struct.* **28**, 295–317
- Collins, G. A., and Goldberg, A. L. (2017) The Logic of the 26S proteasome. *Cell* 169, 792–806
- Finley, D., Chen, X., and Walters, K. J. (2016) Gates, channels, and switches: elements of the proteasome machine. *Trends Biochem. Sci.* 41, 77–93
- Whitby, F. G., Masters, E. I., Kramer, L., Knowlton, J. R., Yao, Y., Wang, C. C., *et al.* (2001) Structural basis for the activation of 20S proteasomes by 11S regulators. *Nature* 408, 115–120
- Smith, D. M., Chang, S. C., Park, S., Finley, D., Cheng, Y., and Goldberg, A. L. (2007) Docking of the proteasomal ATPases' carboxyl termini in the 20S proteasome's alpha ring opens the gate for substrate entry. *Mol. Cell* 27, 731–744
- Sakata, E., Eisele, M. R., and Baumeister, W. (2021) Molecular and cellular dynamics of the 26S proteasome. *Biochim. Biophys. Acta Proteins Proteom.* 1869, 140583
- Ma, C.-P., Slaughter, C. A., and DeMartino, G. N. (1992) Purification and characterization of a protein inhibitor of the 20S proteasome (macropain). *Biochim. Biophys. Acta* 1119, 303–311
- McCutchen-Maloney, S. L., Matsuda, K., Shimbara, N., Binns, D. D., Tanaka, K., Slaughter, C. A., *et al.* (2000) cDNA cloning, expression, and functional characterization of PI31, a proline-rich inhibitor of the proteasome. *J. Biol. Chem.* 275, 18557–18565
- Li, X., Thompson, D., Kumar, B., and DeMartino, G. N. (2014) Molecular and cellular roles of PI31 (PSMF1) protein in regulation of proteasome function. *J. Biol. Chem.* 289, 17392–17405
- Zaiss, D. M., Standera, S., Kloetzel, P. M., and Sijts, A. J. (2002) PI31 is a modulator of proteasome formation and antigen processing. *Proc. Natl. Acad. Sci. U. S. A.* 99, 14344–14349
- Bader, M., Benjamin, S., Wapinski, O. L., Smith, D. M., Goldberg, A. L., and Steller, H. (2011) A conserved F-box-regulatory complex controls proteasome activity in Drosophila. *Cell* 145, 371–382
- Cho-Park, P. F., and Steller, H. (2013) Proteasome regulation by ADPribosylation. *Cell* 153, 614–627
- Liu, K., Jones, S., Minis, A., Rodriguez, J., Molina, H., and Steller, H. (2019) PI31 is an adaptor protein for proteasome transport in axons and required for Synaptic development. *Dev. Cell* 50, 509–524.e10
- Rawson, S., Walsh, R. M., Jr, Velez, B., Schnell, H. M., Jiao, F., Blickling, M., et al. (2022) Yeast PI31 inhibits the proteasome by a direct multisite mechanism. Nat. Struct. Mol. Biol 29, 791–800
- Jespersen, N., Ehrenbolger, K., Winiger, R. R., Svedberg, D., Vossbrinck, C. R., and Barandun, J. (2022) Structure of the reduced microsporidian proteasome bound by PI31-like peptides in dormant spores. *Nat. Commun* 13, 6962
- 24. McGuire, M. J., and DeMartino, G. N. (1989) The latent form of macropain (high molecular weight multicatalytic protease) restores ATP-dependent proteolysis to cell-free extracts of BHK fibroblasts pretreated

with antimacropain antibodies. *Biochem. Biophys. Res. Commun.* 160, 911–916

- 25. Löwe, J., Stock, D., Jap, B., Zwickl, P., Baumeister, W., and Huber, R. (1995) Crystal structure of the 20S proteasome from the archaeon T. acidophilum at 3.4 Ä resolution,. *Science* 268, 533–539
- Groll, M., Heinemeyer, W., Jäger, S., Ullrich, T., Bochtler, M., Wolf, D. H., *et al.* (1999) The catalytic sites of 20S proteasomes and their role in subunit maturation: a mutational and crystallographic study. *Proc. Natl. Acad. Sci. U. S. A.* **96**, 10976–10983
- Keil, B. (1992) Essential substrate residues for action of endopeptidases. In Specificity of Proteolysis, Springer, Berlin, Heidelberg: 43–228
- Religa, T. L., Sprangers, R., and Kay, L. E. (2010) Dynamic regulation of archaeal proteasome gate opening as studied by TROSY NMR. *Science* 328, 98–102
- 29. Osmulski, P. A., Hochstrasser, M., and Gaczynska, M. (2009) A tetrahedral transition state at the active sites of the 20S proteasome is coupled to opening of the alpha-ring channel. *Structure* 17, 1137–1147
- 30. McGuire, M. J., McCullough, M. L., Croall, D. E., and DeMartino, G. N. (1989) The high molecular weight multicatalytic proteinase, macropain, exists in a latent form in human erythrocytes. *Biochim. Biophys. Acta* 995, 181–186
- Schipper-Krom, S., Sanz, A. S., van Bodegraven, E. J., Speijer, D., Florea, B. I., Ovaa, H., *et al.* (2019) Visualizing proteasome activity and intracellular localization using fluorescent proteins and activity-based probes. *Front. Mol. Biosci.* 6, 56
- 32. Kisselev, A. F., Akopian, T. N., Castillo, V., and Goldberg, A. L. (1999) Proteasome active sites allosterically regulate each other, suggesting a cyclical bite-chew mechanism for protein breakdown. *Mol. Cell* 4, 395–402
- 33. Kirk, R., Laman, H., Knowles, P. P., Murray-Rust, J., Lomonosov, M., Meziane, e. K., *et al.* (2008) Structure of a conserved dimerization domain within the F-box protein Fbxo7 and the PI31 proteasome inhibitor. *J. Biol. Chem.* 283, 22325–22335
- 34. Schwanhäusser, B., Busse, D., Li, N., Dittmar, G., Schuchhardt, J., Wolf, J., et al. (2011) Global quantification of mammalian gene expression control corrigendum: global quantification of mammalian gene expression control. Nature 473, 337–342
- Beck, M., Schmidt, A., Malmstroem, J., Claassen, M., Ori, A., Szymborska, A., *et al.* (2011) The quantitative proteome of a human cell line. *Mol. Syst. Biol.* 7, 549
- 36. Itzhak, D. N., Tyanova, S., Cox, J., and Borner, G. H. (2016) Global, quantitative and dynamic mapping of protein subcellular localization. *Elife* 5, e16950
- 37. Kulak, N. A., Pichler, G., Paron, I., Nagaraj, N., and Mann, M. (2014) Minimal, encapsulated proteomic-sample processing applied to copynumber estimation in eukaryotic cells. *Nat. Methods* 11, 319–324
- 38. Zaiss, D. M., Standera, S., Holzhutter, H., Kloetzel, P., and Sijts, A. J. (1999) The proteasome inhibitor PI31 competes with PA28 for binding to 20S proteasomes. *FEBS Lett.* **457**, 333–338
- **39.** Kim, Y. C., and DeMartino, G. N. (2011) C termini of proteasomal ATPases play nonequivalent roles in cellular assembly of mammalian 26 S proteasome. *J. Biol. Chem.* **286**, 26652–26666
- Rabl, J., Smith, D. M., Yu, Y., Chang, S. C., Goldberg, A. L., and Cheng, Y. (2008) Mechanism of gate opening in the 20S proteasome by the proteasomal ATPases. *Mol. Cell* 30, 360–368
- 41. Shang, J., Huang, X., and Du, Z. (2015) The FP domains of PI31 and Fbxo7 have the same protein fold but very different modes of proteinprotein interaction. J. Biomol. Struct. Dyn. 33, 1528–1538
- Liu, C.-W., Corboy, M. J., DeMartino, G. N., and Thomas, P. J. (2003) Endoproteolytic activity of the proteasome. *Science* 299, 408–411
- 43. Di Fonzo, A., Dekker, M. C., Montagna, P., Baruzzi, A., Yonova, E. H., Correia Guedes, L., *et al.* (2009) FBXO7 mutations cause autosomal recessive, early-onset parkinsonian-pyramidal syndrome. *Neurology* 72, 240–245
- Vingill, S., Brockelt, D., Lancelin, C., Tatenhorst, L., Dontcheva, G., Preisinger, C., et al. (2016) Loss of FBXO7 (PARK15) results in reduced



proteasome activity and models a parkinsonism-like phenotype in mice. *EMBO J.* **35**, 2008–2025

- Asher, G., Reuven, N., and Shaul, Y. (2006) 20S proteasomes and protein degradation "by default". *Bioessays* 28, 844–849
- 46. Baugh, J. M., Viktorova, E. G., and Pilipenko, E. V. (2009) Proteasomes can degrade a significant proportion of cellular proteins independent of ubiquitination. J. Mol. Biol. 386, 814–827
- Kumar Deshmukh, F., Yaffe, D., Olshina, M. A., Ben-Nissan, G., and Sharon, M. (2019) The contribution of the 20S proteasome to proteostasis. *Biomolecules* 9, 190
- Ramachandran, K. V., and Margolis, S. S. (2017) A mammalian nervous system-specific plasma membrane proteasome complex that modulates neuronal function. *Nat. Struct. Mol. Biol.* 24, 419–430
- 49. Fabunmi, R. P., Wigley, W. C., Thomas, P. J., and DeMartino, G. N. (2001) Interferon γ regulates accumulation of the proteasome activator PA28 and immunoproteasomes at nuclear PML bodies. *J. Cell Sci.* 114, 29–36
- 50. Kim, Y. C., Li, X., Thompson, D., and DeMartino, G. N. (2013) ATP binding by proteasomal ATPases regulates cellular assembly and

substrate-induced functions of the 26 S proteasome. *J. Biol. Chem.* **288**, 3334–3345

- Punjani, A., Rubinstein, J. L., Fleet, D. J., and Brubaker, M. A. (2017) cryoSPARC: algorithms for rapid unsupervised cryo-EM structure determination. *Nat. Methods* 14, 290–296
- 52. Zivanov, J., Nakane, T., Forsberg, B. O., Kimanius, D., Hagen, W. J., Lindahl, E., *et al.* (2018) New tools for automated high-resolution cryo-EM structure determination in RELION-3. *Elife* 7, e42166
- 53. Zheng, S. Q., Palovcak, E., Armache, J. P., Verba, K. A., Cheng, Y., and Agard, D. A. (2017) MotionCor2: anisotropic correction of beam-induced motion for improved cryo-electron microscopy. *Nat. Methods* 14, 331–332
- Emsley, P., Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Features and development of Coot. *Acta Crystallogr. D Biol. Crystallogr.* 66 (Pt 4), 2010, 486–501.
- 55. Liebschner, D., Afonine, P. V., Baker, M. L., Bunkóczi, G., Chen, V. B., Croll, T. I., *et al.* (2019) Macromolecular structure determination using X-rays, neutrons and electrons: recent developments in Phenix. *Acta Crystallogr. D Struct. Biol.* **75**(Pt 10), 861–877