Activity-Based Profiling Reveals Reactivity of the Murine Thymoproteasome-Specific Subunit β5t

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SUMMARY

Epithelial cells of the thymus cortex express a unique proteasome particle involved in positive T cell selection. This thymoproteasome contains the recently discovered $\beta5t$ subunit that has an uncharted activity, if any. We synthesized fluorescent epoxomicin probes that were used in a chemical proteomics approach, entailing activity-based profiling, affinity purification, and LC-MS identification, to demonstrate that the β 5t subunit is catalytically active in the murine thymus. A panel of established proteasome inhibitors showed that the broad-spectrum inhibitor epoxomicin blocks the ß5t activity and that the subunit-specific antagonists bortezomib and NC005 do not inhibit §5t. We show that §5t has a substrate preference distinct from $\beta 5/\beta 5i$ that might explain how the thymoproteasome generates the MHC class I peptide repertoire needed for positive T cell selection.

INTRODUCTION

The ability to recognize nonself oligopeptides is a key feature of mammalian immunity. T cells that recognize antigenic oligopeptides elicit a directed adaptive immune response aimed at the identification and eventual eradication of the invading pathogen that is the source of the nonself protein from which the antigenic oligopeptide is derived (Janeway and Bottomly, 1994; Medzhi-tov, 2007). T cell recognition is effected by binding of specific T cell receptors to the antigenic peptides that are complexed to either major hisocompatibility complex (MHC) class I or MHC class II molecules (Huseby et al., 2005; Takahama et al., 2008). MHC I molecules present oligopeptides derived from cytosolic and nuclear proteins to CD8⁺ cytotoxic T lymphocytes (CTL) and by this virtue report on the presence of virally encoded proteins (Kloetzel and Ossendorp, 2004). T cells specific for nonself peptides are produced by thymic selection. The generation in the thymus of nonself peptide-selective CTL proceeds in two discreet events (Nitta et al., 2008). Positive selection is mediated by cortical thymic epithelial cells. In this process, thymocytes expressing T cell receptors are confronted with tissues expressing MHC I molecules loaded with oligopeptides. Current understanding is that the MHC I/peptide antigen complexes produced by cortical thymic epithelial cells are low-affinity T cell receptor binders. Thymocytes passing through the thymic cortex that bind to MHC I molecules carrying a peptide load are selected from thymocytes expressing nonbinding receptors. In the ensuing negative selection step, mediated by medullary thymic epithelial cells, thymocytes from the positively selected pool that are responsive to MHC I molecules exposing selfpeptides are eliminated.

Recently, Tanaka and co-workers made a major breakthrough toward understanding how positive selection proceeds (Murata et al., 2007). They found that epithelial cells at the thymic cortex express, next to the constitutive proteasome and the immunoproteasome, a third 20S proteasome particle which was dubbed the thymoproteasome. The 20S core particle of the proteasome is assembled from α and β subunits in a pattern of four, stacked, heptameric rings (α 1-7, β 1-7, α 1-7) generating a barrelshaped structure that contains two copies of the catalytically active β subunits: β 1 (post acidic), β 2 (tryptic-like), β 5 (chymotriptic-like) peptidase activities (Baumeister et al., 1998). The thymoproteasome contains the β 1i and β 2i subunits just like the immunoproteasome, with the important exception that the unique subunit β 5t replaces the immunoproteasome-specific subunit, β 5i.

The thymoproteasome is the most abundant proteasome species in cortical thymic epithelial cells (cTEC). Thymoproteasome expression may have implications for the repertoire of oligopeptides presented by MHC I molecules on the surface of cTECs that might significantly differ from to the repertoire produced by medullary thymic epithelial cells. Closer inspection of the thymoproteasome 20S particle revealed that, in contrast to the constitutive and the immunoproteasome, it possessed little chymotryptic activity, a finding that seems to correlate



activity based probesproteasome inhibitors $f_{i}^{+}f_{i}^$

Figure 1. Activity-Based Probes and Proteasome Inhibitors Used in This Study

In addition to the enzyme reactive group (warhead) and targeting sequence of the inhibitors, activity-based probes are equipped with a fluorophore for in-gel detection, a biotin tag for affinity purification or with both.

with the hydrophilic nature of the putative substrate-binding site of β 5t compared with β 5/ β 5i (Murata et al., 2007). In theory, β 5t can contribute in two ways to the generation of specific MHC I peptides used in positive T cell selection (Murata et al., 2008). It could act as an impassive, catalytically inactive bystander, in which case β 1i/ β 2i produces the majority of MHC I peptides with a bias toward their substrate preferences. Alternatively, it could actively participate in protein degradation and assist in producing "nonself" peptides thanks to its intrinsic substrate preference, which then must be distinct from that of β 5/ β 5i.

Activity-based probes are synthetic compounds bearing a reporter or affinity tag and an enzyme reactive group that can covalently bind to the active site of an enzyme (Cravatt et al., 2008). The tagged enzymatic activities can than be visualized by fluorescence or affinity purified, digested with trypsin, and identified by LC/MS analysis. We here demonstrate, by making use of activity-based proteasome probes (Verdoes et al., 2009), that β 5t is in fact a catalytically active subunit and show that its preference toward established proteasome inhibitors differs substantially from those of β 5/ β 5i.

RESULTS AND DISCUSSION

Activity-Based Profiling Reveals **B5t** Activity

As the first experiment, we incubated whole tissue thymus homogenate from 3-week-old mice with the fluorescent broadspectrum ABPs 1, (Verdoes et al., 2008) 2, 4, and MV151 (Verdoes et al., 2006) shown in Figure 1 (for the synthesis of probes 2 and 4, see Supplemental Experimental Procedures available online). Proteins were resolved by SDS-PAGE under reducing conditions and fluorescently labeled proteasome subunits were visualized by in-gel fluorescence scanning. In Figure 2A, MV151 shows the typical band pattern of staining that is similar to that of the EL4 cell line expressing the constitutive and the immunoproteasome (Kessler et al., 2001) (Figure S1) indicating that both particles are expressed in the thymus. Peptide vinyl sulphone 1, the biotinylated derivative of MV151, shows a similar pattern as MV151. Interestingly, the peptide epoxyketones 2 and 4 show two new bands that run below and above the constitutive and immunoproteasome subunits. Of these, the lower band corresponds to β 1i.



Figure 2. Activity-Based Profiling, Affinity Purification, and LC-MS Identification of Proteasome β Subunits in Murine Tissues Lysates (A) In-gel fluorescence detection of active proteasome β subunits in 3-weekold wild-type murine thymus homogenate after labeling with MV151, ABP 1, 2, and 4 (see also Figure S1). M indicates the molecular marker band of 25 kDa. (B) In-gel fluorescence and silver stain detection of active proteasome β subunits in young and adult thymus after labeling with ABP 1, 2, 3, and affinity purification. Protein identification by LC-MS analysis of in-gel digested silver-stained bands (see Tables S1 and S2 for details).

(C) In-gel fluorescence detection with ABP 4 of β 5t activity in wild-type and absence of activity in the (-/-) β 5t knockdown thymus from 3-weeks-old mice.

(D) Activity-based proteasome profiling using ABP 4 shows β 5t activity in murine thymus (young and adult) but not in heart, lung, liver, spleen, brain, testes, or kidney.

To ascertain whether the new, epoxyketone-sensitive protein, with a gel mobility corresponding to the predicted molecular weight of β 5t (Murata et al., 2007), is indeed β 5t and not a thymus-specific gene product unrelated to the thymoproteasome, we performed a pull-down experiment by making use of the biotin moiety present in ABPs 1 and 2. Biotinylated proteins from thymus homogenate were captured by streptavidin-coated magnetic beads, resolved by SDS-PAGE and detected both by fluorescence and silver staining. Figure 2B shows the specific purification of several proteins that run in a pattern similar with that of Figure 2A. Protein ID, indicated by arrows in Figure 2B, was determined by on-bead (Table S2) and in-gel tryptic digestion followed by LC-MS/MS analysis. Oligopeptides corresponding to the expected constitutive proteasome ($\beta 1/\beta 2/\beta 5$) and immunoproteasome (\beta1i/\beta2i/\beta5i) were captured by ABP 1 but no evidence for $\beta 5t$ was found. Peptides derived from $\beta 5t$ were found by affinity purification with ABP 2, indeed in the band running higher than the other active β subunits. However, the protein yield achieved by pull-down with ABP 1 and 2 was low, and we hypothesized that the short biotinylated epoxomicin ABP 3 might increase the pull-down efficiency (for the synthesis of probe 3, see Supplemental Experimental Procedures). ABP 3 performed as expected, showing bands of similar pattern as ABP 2, stronger signal in silver-stained gels and reliable LC-MS identification of proteins (Table S1). Thymus from adult animals treated in the same fashion shows β 5t activity as well, which suggests that the murine thymoproteasome remains active for at least 6 months. Next to the active proteasome β subunits, only four endogenously biotinylated background proteins were recovered with this method, a result that reflects the selectivity of ABPs 1, 2, 3, and 4 toward proteasomes. Thymus lysates of 2-week-old mice in which the β5t protein expression was genetically knocked down show normal activity of immuno- and constitutive proteasome compared with the wild-type, but complete absence of β 5t activity (Figure 2C). To characterize the expression of $\beta 5t$ in murine tissues, we performed a tissue scan with ABP probe 4. Figure 2D shows that β5t activity is exclusively present in the young thymus and at lower activity in thymus of 6-month-old mice. Integration of the fluorescent signal from young thymus indicated that β 5t contributes to some 4% of the total active signal in this full thymus lysate. Heart, lung, liver, spleen, brain, testes, and kidney do not show ß5t activity. The presence of immunoproteasome bands in the heart, lung, liver, and spleen tissues is explained by the presence of lymphocytes in these organs.

LC-MS³ Analysis of the β 5t Active-Site Peptide

Isolation and analysis of the active-site peptide covalently bound to ABP probe **3** would be the ultimate proof for the β 5t acitivity. Biotin-epoxomicin binds to the catalytic N-terminal threonine via an irreversible morpholino ring formation shown in Figure 3A. The β5t active-site peptide (Figure 3B) is generated after denaturation and tryptic digest of the thymoproteasome. Given that biotin-epoxomicin binds to all active β subunits, we expect to find six different active-site peptides because the tryptic peptides derived from $\beta 5$ and $\beta 5i$ are identical (see Table S3). After LC-MS analysis, the active-site peptides were identified from the high resolution full MS scans by their exact mass and charge (Figure 3C). Further evidence was provided by the MS/MS (MS²) fragmentation that revealed the presence of the biotin-epoxomicin signature ions b_1 , b_2 , b_3 , and b_4 from Figure 3D. In fact, the favored fragmentation of the morpholino ring due to push-pull radical stabilization of the ions (Carey and Sundberg, 2008) yields mainly two major ions b_4 and y_7 , where y_7 contains the peptide sequence of the β subunit active-site. By electrostatic trapping and further MS³ fragmentation of the y_7 ion, the LAFR sequence of the β 5t active-site peptide was identified (Figure 3E). Taken together, this data set demonstrates that β5t is, indeed, a reactive proteasome subunit.

Competitive Activity-Based Profiling Reveals β 5t Substrate Specificity

The finding that β 5t reacts with epoxyketones **2**, **3**, and **4**, but not with peptide vinyl sulphones **1** and MV151 gives a first indication of an altered substrate specificity compared to β 5/ β 5i. With probe **4** in hand as readout, we set out to investigate the β 5t substrate preference by competitive activity-based studies

Chemistry & Biology Activity-Based Profiling Shows β5t Reactivity





(A) Reaction mechanism of biotin-epoxomicin **3** with the catalytically active N-terminal Thr residue of active proteasome β subunits. The morpholino ring formation results in a covalent and irreversible binding.

(B) Schematic representation of the biotin-epoxomicin modified, N-terminal active-site tryptic peptide of β5t. Amino acid residues are represented in a three-letter code.

(C) LC-MS elution profile of the six unique biotinylated tryptic peptides derived from the active sites. Notice that $\beta 5$ and $\beta 5i$ active-site peptides are identical (see also Table S3).

(D) LC-MS² determination of the β 5t active-site fragmentation pattern. The parent ion $[m/z (M+2H)^{2+} = 767.44]$ was fragmented. The b_1 , b_2 , b_3 , and b_4 ions are signature ions of the biotin-epoxomicin N-terminal part. The abundant y_7 ion containing the β 5t active-site peptide sequence was selected for further (MS³) fragmentation (see E).

(E) LC-MS³ determination of the y_7 ion (MH⁺ = 821.32) revealing the β 5t activesite peptide amino acid sequence.



Figure 4. Analysis of β 5t Substrate Specificity in Juvenile Murine Thymus Lysates by Competitive Activity-Based Profiling with ABP 4 (A) Lysates were exposed to increasing concentrations Lactacystin or MG132; residual β 5t activity was stained with ABP 4 and visualized by in-gel fluorescence detection. The inhibitors are not reactive toward the β 5t activity.

(B) Bortezomib efficiently inhibits β 5, β 5i, β 1, and β 1i activity but not β 5t. NC005 specifically targets β 5 (indicated by the gray arrow) but not β 5t. The mixed inhibitor containing the boronic ester warhead equipped with the epoxomicin tail efficiently blocks β 5t.

(C) In analogy to (B), installation of the epoxomicin llelleThrLeu peptide targeting motif to the vinyl sulphone warhead affords a potent inhibitor of the β 5t activity.

with established proteasome inhibitors of diverse chemical characteristics. Figure 4A shows the results of the most commonly used proteasome inhibitors lactacystin (Fenteany et al., 1995) and MG132 (Rock and Goldberg, 1999). Both require concentrations higher that 10 µM for broad-spectrum proteasome inhibition with marked affinity for β 5 and β 2 but do not inhibit the β 5t activity. Figure 4B shows that the Bortezomib boronic ester 5 effectively blocks \(\beta1\), \(\beta1\)i, \(\beta5\), and \(\beta5\)i as previously described (Adams et al., 1998; Verdoes et al., 2007), while the subunitspecific inhibitor NC005 (Britton et al., 2009) selectively inhibits the *β5/β5i* subunits. Neither bortezomib nor NC005 appears to interact with the β5t. However, mixing of the potent boronic ester warhead with the Acl₂TL peptide sequence inherent to epoxomicin as in compound 13 abolished the subunit preference of bortezomib and efficiently inhibited ß5t. A similar effect is revealed in Figure 4C where the vinyl sulphone warhead (Bogyo et al., 1997) was equipped with the AdaAhx₃ extended I₂TL motif. Apparently, the presence of the hydrophilic threonine side chain at P2 in an inhibitor or ABP probe is favorable for affinity to the ß5t subunit. From the results, some interesting trends pointing toward a substrate preference of β 5t that is rather distinct to that of $\beta 5/\beta 5i$ appear. Whereas $\beta 5t$ is sensitive toward the broad-spectrum proteasome epoxomicin 6, it is quite unreactive toward the \u03b35/\u03b35i-biased compounds. Bortezomib boronic ester 5 (which at the concentrations used disables $\beta 1/\beta$ β 1i/ β 5/ β 5i) and lactacystin **8** are unreactive toward β 5t, as is the case with vinyl sulfone 7, peptide aldehyde 9, and epoxyketone 12

As a final research objective, we set out to establish whether β5t activity could be detected in fluorogenic substrate assays. In a first experiment, lysates of 3-week-old wild-type and β 5t-/murine thymi were preincubated with 1 μ M 10, 6, 9, or the β 5/ β 5ispecific inhibitor NC005 (12) prior to treatment with the reported β5-specific fluorogenic substrate, Ac-LLVY-amc (Kisselev and Goldberg, 2005). No differential signal could be detected in the wt versus the β 5t-/- lysates (data not shown), indicating that this substrate does not react with β5t. We next synthesized two novel fluorogenic substrates, Ac-IITL-amc and Ac-SSTL-amc (Supplemental Experimental Procedures) that were expected to be cleaved by all catalytically active β subunits, including β 5t. Preincubation of wild-type and β 5t-/- lysates with 1 μ M **10**, **6**, **9**, or **12** followed by β 5t read-out with Ac-IITL-amc or Ac-SSTL-amc again did not result in a discernible ß5t-specific signal (data not shown). We next fractionated the lysates by 10%-40% sucrose gradient ultracentrifugation and used the enriched (thymo)proteasome fractions as determined by ABP profiling with 4 (Figure S2). The tryptic- and caspase-like activities were inhibited with 1 or 10 μ M 6 or 7 and the β 5t activity was assayed with Ac-IITL-amc or Ac-SSTL-amc (Figure S3). Again, no differences could be found between wild-type and β 5t-/- (thymo)proteasomes.

In conclusion, we have demonstrated that β 5t has an intrinsic reactivity resembling that of the other proteasome catalytic ß5t subunits and have obtained evidence suggesting that this thymoproteasome-specific subunit plays an active role in positive T cell selection. Our discovery that $\beta 5t$ is an active species is based on activity-based probes, and we acknowledge the fact that the reactive groups differ in nature from the natural substrates (vinyl sulphone/epoxyketone as opposed to amide bonds). We note, however, that all known proteasome catalytic subunits are active toward these electrophilic traps and moreover that all nonactives are not. In fact, we argue that activity-based enzyme profiling is especially attractive for probing putative proteasome active sites, which are active only in the context of the 20S conglomerate as a whole and by activity-based profiling, can still be reported on individually. We indeed were unable to detect $\beta 5t$ activity by using fluorogenic substrates, due to the fact that the substrates we were able to obtain or design based on our data were either specific for $\beta 5/\beta 5i$ (in which case no $\beta 5t$ activity is to be expected based on our inhibition studies) or are pan-reactive (in which case we could not discriminate between the different activities). More research is required to identify a fluorogenic substrate specific for β 5t, and we argue this may be accomplished by screening (libraries of) designed inhibitors against thymoproteasomes in an activity-based profiling assay such as disclosed here. Altogether, our data, revealing that β 5t is catalytically active toward inhibitors that contain a hydrophilic residue (Thr) in a hydrophobic stretch, point toward the involvement of β5t in the generation of a unique set of oligopeptides complexed to MHC I molecules for optimal positive T cell selection.

SIGNIFICANCE

We demonstrate for the first time to the best of our knowledge that thymoproteasome-specific β 5t subunit is catalytically active. Interestingly, active β 5t is also found in adult thymi but not in other murine organs. We provide the first insight into the nature of the substrate preference of $\beta 5t$ that is probably biased toward peptides with hydrophilic side chains. This body of evidence was made possible by the direct action of activity-based probes (ABP) with emphasis on the bifunctional ABPs that facilitate both readout and affinity purification. Moreover, the Bodipy-epoxomicin probe shows superior labeling and SDS-PAGE resolution for the active β subunits. The biotin-epoxomicin probe facilitated the LC-MS³ identification of the β 5t active-site peptide that is the ultimate proof for the reactivity of this subunit. This work illustrates the versatility of fluorescent activity-based probes and opens new avenues of detection as fluorescence microscopy or flow cytometry in living cells. Therefore, the probes presented here should be considered as a powerful addition to the toolbox for proteasome activity determination and guantification besides the classic fluorogenic substrate assays.

EXPERIMENTAL PROCEDURES

Animals and Tissues

Thymus and other organs were isolated from young (3 week old) or adult mice and kindly provided by Ine Tijdens, Chantal Pont, and Prof. Dr. Bob van de Water. Thymus from β 5t knockout mice was kindly provided by Dr. Keiji Tanaka. Organ isolation was approved by the animal experimentation ethical committee of the Leiden University and Tokyo Metropolitan Institute of Medical Science.

Compounds

Design, synthesis, and mechanism of action of the activity-based probes **1–4**, MV151, and the proteasome inhibitors **5–11**, **13** is reviewed in Verdoes et al. (2009). NC005 is described in Britton et al. (2009). Lactacystin, MG132, and all other compounds of analytical grade were purchased from Sigma-Aldrich.

Activity-Based Profiling

Tissues were homogenized in 3 volumes of ice-cold lysis buffer (50 mM Tris-HCI [pH 7.5], 250 mM sucrose, 5 mM MgCl₂, 1 mM DTT, 2 mM ATP, 0.025% digitonin, 0.2% NP40; Kisselev and Goldberg, 2005) with a tissue homogenizer and further disrupted by 2 × 30 s sonication. Lysates were cleared by cold centrifugation at 13,000 × g, protein concentrations determined by Bradford assay and kept at -80° C until use. For comparative activity-based profiling, equal amounts of protein were incubated with ABPs for 1 hr at 37°C and resolved by 12.5% SDS-PAGE, and the wet gel slab was scanned on a Thyphoon scanner (GE Healthcare) with the TAMRA settings ($\lambda_{ex} = 530$ nm, $\lambda_{em} = 560$ nm). Competitive activity-based profiling was done by first incubating thymus lysates with increasing concentrations of various proteasome inhibitors for 1 hr at 37°C, followed by 1 hr incubation with 0.5 μ M ABP 4 for the in-gel detection of the residual proteasome activity. Images were acquired, processed, and quantified with Image Quant (GE Healthcare).

Affinity Purification

Approximately 1 or 2 mg of protein was incubated with 10 μ M biotinylated ABPs **1**, **2** or **3** for 1 hr at 37°C, denatured by boiling for 5 min with 1% SDS and precipitated with chloroform/ methanol (C/M, Wessel and Flügge, 1984). The protein pellet was rehydrated in 180 μ l 8 M urea/100 mM NH₄HCO₃, reduced with 10 μ l 90 mM DTT for 30 min at 37°C, alkylated with 15 μ l 200 mM iodoacetamide at room temperature in the dark, cleared by centrifugation at 13,000 × g, and desalted by C/M. The pellet was dispersed in 25 μ l PD buffer (50 mM Tris-HCl [pH 7.5], 150 mM NaCl) with 2% SDS in a heated (37°C) sonic bath. Stepwise (3 × 25 μ l, 4 × 100 μ l, 1 × 500 μ l) addition of PD buffer afforded a clear solution that was incubated with 50 μ l MyOne T1 Streptavidin-grafted beads (Invitrogen) at room temperature with vigorous shaking for 2 hr. The beads were stringently washed with 2 × 300 μ l PD buffer I (4 M urea/50 mM NH₄HCO₃), 2 × 300 μ l wash buffer II (50 mM Tris-HCl [pH 7.5], 10 mM NaCl),

and 2 × 300 µl water. For in-gel analysis, two-thirds of the beads were eluted with 100 µl 1× sample buffer containing 10 µM biotin by boiling for 5 min at 90°C and resolved by 12.5% SDS-PAGE. Proteins were visualized by fluorescence and silver stain, in-gel digested (Shevchenko et al., 2006), and desalted (Rappsilber et al., 2007). For on/bead digest, one-third of the beads were digested with 300 ng trypsin in 100 µl digest buffer (100 mM Tris-HCI [pH 7.8], 100 mM NaCl, 1 mM CaCl₂, 2% ACN) overnight at 37°C. Peptides were eluted with 2 × 80 µl 10 µM biotin in 5% formic acid/25% ACN/70% H₂O for 30 min at 37°C and desalted after ACN evaporation.

LC-MS Analysis

Trytic peptides were analyzed on a Surveyor nanoLC system (Thermo) hyphenated to a LTQ-Orbitrap mass spectrometer (Thermo). Gold and carbon coated emitters (OD/ID = 360/25 μ m tip ID = 5 μ m), trap column (OD/ID = 360/25 μ m top ID = 5 μ m), trap column (OD/ID = 360/25 μ m 120 Å) and analytical columns (OD/ID = 360/75 μ m packed with 25 mm robust Poros10R2/ 15 mm BioSphere C18 5 μ m 120 Å) and analytical columns (OD/ID = 360/75 μ m packed with 20 cm BioSphere C18 5 μ m 120 Å) were from Nanoseparations (Nieuwkoop, The Netherlands). The mobile phases (A: 0.1% FA/H2O, B: 0.1% FA/ACN) were made with ULC/MS grade solvents (Biosolve). The emitter tip was coupled end-to-end with the analytical column via a 15 mm long TFE Teflon tubing sleeve (OD/ID 0.3 × 1.58 mm, Supelco, USA) and installed in a stainless steel holder mounted in a nano-source base (Upchurch scientific, Idex, USA).

General mass spectrometric conditions were as follows: an electrospray voltage of 1.8 kV was applied to the emitter, no sheath and auxiliary gas flow, ion transfer tube temperature 150°C, capillary voltage 41 V, tube lens voltage 150 V. Internal mass calibration was performed with air-borne protonated polydimethylcyclosiloxane (m/z = 445.12002) and the plasticizer protonated dioctyl phthalate ions (m/z = 391.28429) as lock mass (Olsen et al., 2005).

For shotgun proteomics analysis, 10 µl of the samples was pressure loaded on the trap column with a 10 µl/min flow for 5 min followed by peptide separation with a gradient of 35 min 5%–30% B, 15 min 30%–60% B, 5 min A at a flow of 300 µl/min split to 250 nl/min by the LTQ divert valve. For each data-dependent cycle, one full MS scan (300–2000 *m/z*) acquired at high mass resolution (60,000 at 400 *m/z*, AGC target 1 × 10⁶, maximum injection time 1000 ms) in the Orbitrap was followed by three MS/MS fragmentations in the LTQ linear ion trap (AGC target 5 × 10³, maximum injection time 120 ms) from the three most abundant ions (Baek et al., 2008). MS² settings were as follows: collision gas pressure 1.3 mT, normalized collision energy 35%, ion selection threshold of 500 counts, activation *q* = 0.25 and activation time of 30 ms. Fragmented precursor ions that were measured twice within 10 s were dynamically excluded for 60 s and ions with *z* < 2 or unassigned were not analyzed.

A parent ion list of the *m/z* ratios of the active-site peptides was compiled and used for LC-MS³ analysis in a data-dependent protocol. The parent ion was electrostatically isolated in the ion trap of the LTQ and fragmented by MS², and the most intense peak was isolated and further fragmented in MS³ to reveal the amino acid sequence of the active-site peptide. Data from MS² and MS³ were validated manually.

SUPPLEMENTAL INFORMATION

Supplemental Material includes Supplemental Experimental Procedures, three figures, and three tables and can be found with this article online at doi:10.1016/j.chembiol.2010.05.027.

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