ADAMDEC1 Is a Metzincin Metalloprotease with Dampered Proteolytic Activity

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Background: ADAMDEC1 is a putative ADAM-like metalloprotease with a short domain structure and a noncanonical active site.

Results: Recombinant ADAMDEC1 cleaves α2-macroglobulin and casein. The activity is significantly enhanced upon reconstituting the consensus zinc-binding site.

Conclusion: ADAMDEC1 is secreted as a proteolytic active, glycosylated metalloprotease.

Significance: The ADAMDEC1 has adapted a reduced catalytic activity, possibly compensating the loss of auxiliary specificity determining domains.

ADAMDEC1 (Decysin-1) is a putative ADAM (a disintegrin and metalloprotease)-like metalloprotease with an unknown physiological role, selectively expressed in mature dendritic cells and macrophages. When compared with other members of the ADAM family, ADAMDEC1 displays some unusual features. It lacks the auxiliary cysteine-rich, EGF, and transmembrane domains, as well as the cytoplasmic tail. The active site of ADAMDEC1 is unique by being the only mammalian ADAM protease with a non-histidine zinc ligand, having an aspartic acid residue instead. Here we demonstrate that ADAMDEC1, despite these unique features, functions as an active metalloprotease. Thus, ADAMDEC1 is secreted as a mature, glycosylated, and proteolytically active metalloprotease, capable of cleaving macromolecular substrates. In the recombinant form, three of the four potential N-linked glycosylation sites are modified by carbohydrate attachment. Substitution of basic residues at the predicted proprotein convertase cleavage site blocks proprotein processing, revealing both specific ADAMDEC1-dependent and specific ADAMDEC1-independent cleavage of the prodomain. The pro-form of ADAMDEC1 does not have proteolytic activity, demonstrating that the prodomain of ADAMDEC1, like in other members of the ADAM family, confers catalytic latency. Interestingly, the proteolytic activity of mature ADAMDEC1 can be significantly enhanced when a canonical ADAM active site with three zinc-coordinating histidine residues is introduced.

ADAMDEC1 (Decysin-1) mRNA was first identified in 1997 by Mueller et al. (1) and predicted to encode a novel ADAM2 (a disintegrin and metalloprotease)-like protease based on sequence homology with members of the ADAM family of metzincin metalloproteases. ADAMDEC1 bears closest resemblance to ADAM-28 and ADAM-7, with sequence identities of 47 and 36%, respectively, whose genes also cluster together with ADAMDEC1 on chromosome 8p12 in the human genome (2). Phylogenetic analysis suggests that the partial gene duplication event giving rise to ADAMDEC1 and ADAM-7 and -28 from a common ancestor followed the mammalian divergence from amphibians, indicating that ADAMDEC1 is of relative late descendant when compared with other ADAMs (2, 3).

The ADAM family constitutes type-1 transmembrane metzincin metalloproteases, which in addition to the metzincin/reprolysin-type metalloprotease domain have multiple C-terminal auxiliary domains including a disintegrin-like domain, a cysteine-rich domain, an EGF-like domain, a transmembrane domain, and a cytoplasmic tail (4, 5). The domains positioned C-terminal to the metalloprotease domain are involved in cell adhesion, integrin binding, signaling events, and substrate recognition (5–7). In metzincin superfamily members, the metalloprotease domain contains the active site consisting of an elongated zinc-binding motif (HEXXHXXG/NXXHH/DD) with three zinc-coordinating ligands (underlined) immediately followed by a family-specific residue (5, 8). For the catalytic active ADAM subfamily members, the consensus sequence conforms to HEXXHXXGXXHD with three histidine residues coordinating the zinc ion, which together with a glutamic acid residue polarizes the catalytic solvent molecule promoting the nucleophile attack on the substrate, a catalytic mechanism similar to that of thermolysin (9). The conserved glycine within the active site allows for a sharp turn positioning the third histidine for zinc binding (4). In addition, ADAMs contain a conserved methionine (placed in the so-called Met-turn) positioned beneath the three zinc-binding histidines acting as a hydrophobic base (4). Catalytically active ADAM proteases are synthesized as zymogens with a large N-terminal prodomain shown to maintain catalytic latency (5, 10).

Like other ADAMs, ADAMDEC1 mRNA encodes an N-terminal signal peptide for directing secretion and a relatively
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The ADAMDEC1 cDNA, optimized for mammalian cell expression, has a large (173-residue) prodomain, predicted to be cleaved off during maturation. Indeed, furin has been shown in vitro to be capable of processing the prodomain in the murine ortholog (11). The mature protein only comprises a metzincin metalloprotease domain and a short disintegrin-like domain. Consequently, ADAMDEC1 lacks most of the auxiliary domains otherwise present in the ADAMs and is predicted to be secreted as a soluble protein (1, 2). The metzincin metalloprotease domain harbors a putative zinc-binding active site consensus sequence (HEXXHXXGXXD). However, the third zinc-binding ligand in ADAMDEC1 is an aspartic acid residue, as found in some bacterial metzincins, instead of the histidine residue found in all other proteolytically active ADAMs. Due to the differences in primary structure, ADAMDEC1 is regarded as the first (and only) member of a novel subgroup of mammalian ADAMs (1).

Little is known about the physiological role of this putative metalloprotease, but ADAMDEC1 is expressed in monocytes at low levels and at increased levels during 1α,25-dihydroxyvitamin D₃-induced differentiation into mature macrophages (12). Moreover, expression is strongly induced by spontaneous as well as by CD4₀- or LPS-stimulated maturation of immature dendritic cells (1, 12). ADAMDEC1 expression was found to be up-regulated in atherosclerotic plaques (13), pulmonary sarcoidosis (14), intestinal inflammation (15), intracranial tumor (craniopharyngioma) (16), and infection of lymphoblastoid dendritic cells by Epstein-Barr virus (17), whereas it is down-regulated in colorectal cancer (18). In addition, ADAMDEC1 is up-regulated during pregnancy (19). Collectively, the expression pattern of ADAMDEC1 suggests that it plays a role in the immune response (2). However, ADAMDEC1 may have additional physiological roles as well.

Given the lack of functional and structural characterization of this unusual ADAM-like protein, we investigated the putative proteolytic activity of human ADAMDEC1 expressed in a mammalian expression system. We demonstrate that ADAMDEC1 is a secreted, active enzyme capable of cleaving α₂-macroglobulin (α2M) and casin. In addition, we show that the prodomain confers catalytic latency. The same prodomain undergoes specific ADAMDEC1-dependent cleavage, which can occur in trans. The proteolytic activity of mature ADAMDEC1 is enhanced by reconstructing the canonical ADAM active site consensus sequence with three histidine residues coordinating the zinc ion. Finally, we demonstrate that three of the four potential sites are modified by N-linked glycosylation.

**EXPERIMENTAL PROCEDURES**

**Materials**—The FreeStyle 293 expression system and all additives, SDS-PAGE, and Western blot materials (Novex/iBlot System were from Invitrogen. Azocasein and (1,10)-phenanthroline were from Sigma-Aldrich and VWR International, respectively. PNGase F endoglycosidase was from New England Biolabs. Plasma-derived human 25(OH)-vitamin D₃ was from MP Biomedicals. HepG2 cells were from ATCC and cultured in Iscove’s modified Dulbecco’s medium containing 10% calf serum and 1% penicillin/streptomycin (Invitrogen) at 37 °C, 5% CO₂, and 125 rpm. Subcloning and mutagenesis primers were purchased from MWG Eurofins (Ebersberg, Germany), and primer sequences are available in Table 1.

**Protein Expression and Western Blot Analysis—**HEK293-6E cells were grown in FreeStyle 293 expression medium at 37 °C, 8% CO₂, and 125 rpm; for HEK293-6E cells, the medium was supplemented with 25 μg/ml G-418 Geneticin and 0.1% (v/v) Pluronic F-68. Cells were transfected with 1 μg of DNA/10⁶ cells using 293Fectin and Opti-MEM according to the manufacturer’s protocol. Media were harvested 3–4 days after transfection, sterile-filtered, and stored at −80 °C. For SDS-PAGE and Western blot analysis, proteins were denatured by boiling in LDS sample buffer. DTT was added to reduced samples to 50 mM prior to boiling. Samples were separated in 4−12% Bis-Tris gels using MES or MOPS running buffer. Proteins were transferred to nitrocellulose membranes using the iBlot dry blotting system and blocked in 5%(W/V) skim milk in wash buffer (20 mm Tris (pH 7.4), 140 mm NaCl, 5 mm CaCl₂, 0.05%(v/v) Tween 20). 0.43 μg/ml murine anti-human ADAMDEC1 mAb (Abcam ab57224) and 1.3 μg/ml HRP-labeled rabbit anti-mouse IgG (Dako) were used for detection of ADAMDEC1 expression. N-Linked glycans were removed under denaturing conditions using PNGase F according to the manufacturer’s protocol. For N-terminal sequencing, proteins

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**Table 1**

**DNA primer list**

| Subcloning | Reverse | Sequence |
|---|---|---|
| ATACATATGGTCACATTTCATCACGCTCTGGCUTGG | TGTCCGACCTCTGCTTCGTTG |

| Insertion of N-terminal HPC4 tag | Forward | Internal forward | Internal reverse |
|---|---|---|---|
| TATGCTACATGTCACATTTCATGGCAAT | CAACTGACCCGCAAGTACATTTCATCACGCTCTGGCUTGGCAAT | CAACTGACCCGCAAGTACATTTCATCACGCTCTGGCUTGGCAAT |

**Mutagenesis**

| Mutagenesis | Sequence |
|---|---|
| E353A | GGAATGATGTCACATTTCATCACGCTCTGGCUTGG |
| D362H | GTGCTGGGCATGCCTCATGACGTGCCTTTCAACACC |
| H373A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| K55A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| R56A | GCAACATGCTCAGACGCGGACCGGGAGATCC |
| E353A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N61Q | GCGGAGATCAAGAACCAACAGACCGAGAAGCACGGC |
| K160A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| P161A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| H54A | GCACATCCTGCACAAGCGGGAGATCGCAAGACCG |
| R200K/R2003A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N184Q | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N55A | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N61Q | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N184Q | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N270Q | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |
| N466Q | GCCATCAGAGGACCCTGCCCAACACCTGCGGCTG |

**Note:** Mutagenesis primers were stated as the forward primer sequence. For each forward primer, a reverse complementary primer was added to each mutagenesis reaction.
were separated by SDS-PAGE and transferred to PVDF membranes. Individual bands were cut out and subjected to Edman amino acid sequence analysis using an Applied Biosystems Precise HT protein sequencer with on-line identification of phe- nythiohydantoin derivatives.

**Proteolytic Activity Assays**—Cell supernatants were concentrated ~60 times, diluted 30 times in reaction buffer (50 mM Hepes (pH 7.5), 100 mM NaCl, 1 mM CaCl₂, 10 μM ZnCl₂), and concentrated again to a final concentration ~60 times that of the supernatant using Amicon Ultra 10,000 molecular weight cut-off centrifugal filters (Millipore). The relative concentrations of ADAMDEC1 variants were evaluated by densitometric analysis of Coomassie Brilliant Blue-stained SDS-PAGE using the ImageJ software (20). An added amount of cell supernatant was adjusted in an iterative fashion until sample normalization was obtained. Supernatants from cells with added transfection reagents without DNA vector (Mock) were used as background control.

Identical amounts of concentrated ADAMDEC1 variant proteins, corresponding to ~400 μl of ADAMDEC1 WT in cell supernatant, and an excess amount of Mock cell supernatant were added to azocasein solubilized in reaction buffer to a final concentration of 2 mg/ml (~94 μM). Reaction mixtures were incubated at 37 °C for 3 days. The proteolysis was terminated by the addition of TCA to a final concentration of 6% (w/v) followed by a 30-min incubation on ice and removal of undigested protein by centrifugation at 10,000 × g and 4 °C for 10 min. NaOH was added to the supernatants to a final concentration of 0.26 M, and the absorbance was measured at 440 nm using a SpectraMax 190 (Molecular Devices). Absorption data were subtracted from buffer contribution and compared with the data of Mock using one-way analysis of variance analysis with Bonferroni’s adjustment. Individual sample pairs were additionally compared using an unpaired, two-tailed t test.

Proteolysis of human plasma-derived α2M was carried out by incubating ADAMDEC1 variant proteins (corresponding to 150 μl of cell supernatant) with α2M (final concentration: 0.9 mg/ml (1.25 μM)) in reaction buffer for ~65 h at 37 °C. The result was visualized by reducing SDS-PAGE, and cleavage of α2M was observed by appearance of an 85-kDa fragment (21, 22). Proteolysis of α2M in human plasma was followed by adding identical amounts of ADAMDEC1 variants (equal to 200 μl of cell supernatant) to human plasma stabilized with 200 nM tick anticoagulant peptide and 1 μM hirudin. After ~40 h of incubation at 37 °C, α2M fragments were visualized by Western blot analysis using 1 μg/ml mouse anti-human α2M antibody (2D9, Abcam ab36995) and 1.3 μg/ml HRP-labeled rabbit anti-mouse IgG (Dako). Cross-linking of ADAMDEC1 to plasma-derived α2M was visualized by Western blot analysis using 1.31 μg/ml anti-HPC4 tag antibody and 1 μg/ml HRP-labeled goat anti-human IgG.

**RESULTS**

**ADAMDEC1 Is Expressed and Secreted as a Mature and Glycosylated Protein**—ADAMDEC1 protein was detected in the cell supernatant of HEK293 cells transfected with pCI-ADAMDEC1, encoding full-length human wild-type ADAMDEC1 (WT) (resides 1–470, Fig. 1A). SDS-PAGE and Western blot analysis both showed expression by transfected cells, and no endogenous ADAMDEC1 expression was detected in the supernatant of Mock-transfected cells (Fig. 1, B and C). The observed molecular mass of ADAMDEC1 is ~33 kDa, corresponding to the proADAMDEC1 being processed and secreted as the mature protein. A decreased mobility of reduced ADAMDEC1 indicates the presence of disulfides in the secreted protein (Fig. 1C).

ADAMDEC1 has two predicted N-linked glycosylation sites in the prodomain and two in the mature protein. Deglycosylation using PNGase F resulted in an increased electrophoretic mobility and an apparent molecular mass of 30 kDa, which correlates with the predicted molecular mass of 29.4 kDa for the mature protein without glycan and shows that mature ADAMDEC1 is modified with at least one N-linked glycan. We individually removed the glycosylation sites in the metalloprotease domain (Asn-237 and Asn-466) in the truncated disintegrin-like domain (Asn-237 and Asn-466) analyzed in anti-ADAMDEC1 Western blot analysis both showed expression by transfected cells, and no endogenous ADAMDEC1 expression was detected in the supernatant of Mock-transfected cells (Fig. 1, B and C). The observed molecular mass of ADAMDEC1 is ~33 kDa, corresponding to the proADAMDEC1 being processed and secreted as the mature protein. A decreased mobility of reduced ADAMDEC1 indicates the presence of disulfides in the secreted protein (Fig. 1C).

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**Expression and glycosylation status of mature ADAMDEC1.** A, domain structure of full-length ADAMDEC1 consisting of a signal peptide (SP, res. 1–30), a prodomain (PRO, res. 31–203) predicted to be processed at a proprotein convertase recognition site (PC), a metzincin metalloprotease domain (MP, res. 204–411) containing the putative zinc-binding active site (AS, res. 352–362), and a short disintegrin-like domain (DIS, res. 412–470) roughly two-thirds the length of the disintegrin-like domain in other ADAMs (1, 2). The mAb used for Western blot analysis is raised against a C-terminal fragment (res. 361–470, indicated on the figure). Confirmed N-linked glycosylation sites (Asn-61, Asn-184, and Asn-237) are indicated by black lollipops. B, secretion of recombinant human ADAMDEC1 WT to mammalian cell supernatant shown in nonreducing (NR) SDS-PAGE loaded with ~19 μl of cell supernatant. MW, molecular weight markers. C, anti-ADAMDEC1 Western blot analysis of nonreduced (NR), reduced (R), and reduced + PNGase F-treated (R+PF) ADAMDEC1 WT. D, electrophoretic mobility of ADAMDEC1 variants containing substitutions in the two predicted sequons in the mature protein (Asn-237 and Asn-466) analyzed in anti-ADAMDEC1 Western blot and compared with reduced (R) and reduced + PNGase F-treated (R+PF) ADAMDEC1 WT.
whether ADAMDEC1 was able to cleave α2M, an important regulator of a broad spectrum of proteases including several ADAMs (23–25). We incubated ADAMDEC1 with plasma-derived α2M (Fig. 2A) and stabilized human plasma (Fig. 2B). α2M proteolysis was analyzed by SDS-PAGE followed by Coomassie Brilliant Blue staining and anti-α2M Western blot analysis, respectively. We found that ADAMDEC1 cleaves α2M, demonstrated by generation of an immunoreactive 85-kDa fragment (marked by arrow in Fig. 2, A and B). In addition, SDS-stable ADAMDEC1-α2M complexes were observed by Western blot analysis specific for the recombinant ADAMDEC1, demonstrating thioester-mediated cross-linking (Fig. 2C). Further, cleavage of α2M by ADAMDEC1 was inhibited by 5 mM (1,10)-phenanthroline, confirming metal ion dependence (Fig. 2, A and C). Substitution of the predicted catalytic glutamic acid residue (E353A, Table 2) completely abrogated the proteolytic activity, confirming the identity of the ADAMDEC1 active site (Fig. 2, A–C). Incubation of α2M with supernatant from Mock-transfected cells did not induce α2M proteolysis.

When applied to the broadly used substrate azo-labeled casein (azocasein), HEK293 cell supernatant containing recombinant ADAMDEC1 WT exhibited caseinolytic activity, as observed by a significant increase in release of azo-labeled peptides, whereas supernatant from Mock-transfected cells showed no activity (Fig. 2D). Again, the E353A substitution completely abrogated the proteolytic activity.

Immediately downstream of the long zinc-binding consensus sequence, metzincins have a so-called family-specific residue (5), which for adamalysins/ADAMs in general comprises an aspartic acid. Interestingly, it is a valine in ADAMDEC1 (Table 2). In addition, the third zinc-binding residue of the consensus sequence is an aspartic acid, whereas it is generally a histidine in other adamalysins/ADAMs and other metzincins with the exception of snapalysins and, possibly, thurinylsins (5). This feature could have resulted from an evolutionary substitution event or alternatively from a deletion event. To reconstruct an intact zinc binding motif, two new variants were constructed: D362H, substituting the predicted zinc-coordinating aspartic acid with histidine, and His-362insert, inserting a histidine residue at position 362, thereby creating the complete ADAM zinc-coordinating sphere (HEXXXHXXGXXHD) followed by the ADAM family-specific aspartic acid residue (Table 2). When tested as described above, the D362H variant demonstrated significantly enhanced proteolytic activity toward both α2M and azocasein, whereas no proteolytic activ-

**FIGURE 2.** **Proteolytic activity of ADAMDEC1.** A, incubation of plasma-derived α2M with supernatants from Mock, WT, and HPC4-tagged E353A-transfected cells, respectively, in the absence or presence of 5 mM (1,10)-phenanthroline (Phen). Proteolysis of α2M is demonstrated by generation of an 85-kDa α2M fragment indicated by the arrow in reducing SDS-PAGE. Intact 180-kDa α2M monomeric subunit, 120- and 60-kDa α2M heat fragments (47), and the added ADAMDEC1 variants (~33 kDa) are indicated, as well as a 50-kDa unidentified impurity. MW, molecular weight markers. B, anti-α2M Western blot analysis of stabilized human plasma incubated with supernatants from Mock-, WT-, E353A-, and D362H-transfected cells, respectively. The arrow marks the immunoreactive active 85-kDa α2M proteolytic fragment. C, covalent cross-linking between HPC4-tagged ADAMDEC1 WT and plasma-derived α2M, in the absence or presence of 5 mM (1,10)-phenanthroline, visualized by anti-HPC4 Western blot analysis. D, proteolytic activity assay of ADAMDEC1 WT and variants using azocasein as a substrate. The sample means are shown by a horizontal line. ***, p < 0.0001; ns., not significant relative to Mock.

**TABLE 2**

| PC recognition site | Active site |
|---------------------|------------|
| Ille99-Ser204        | His352-Asn366 (Phe365) |
| ADAM consensus       | xHEBGBxSBGBxHJD x x x |
| WT                  | IRISRFS    |
| E353A               | IRISRFS    |
| D362H               | IRISRFS    |
| H362insert          | IRISRFS    |
| R200K/R203A          | IKISA S    |
| R200K/R203A/E353A    | IKISA S    |

**Notes:** The predicted catalytic residue (Glu-353) is in italic font, predicted zinc-coordinating ligands are underlined, and mutations are in bold. The predicted active site sequence is compared with the ADAM family consensus sequence (gray box) where X is a less conserved residues and B is a bulky hydrophobic residues. Promodain processing among the ADAMs is subject to great variation and is therefore not included in this comparison.
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**Processing of proADAMDEC1—N-terminal sequencing** of the recombinant ADAMDEC1 WT protein by Edman degradation confirmed processing at the furin-like proprotein convertase recognition site: RISSRLSLKS (Fig. 3A). To enable studying of the ADAMDEC1 pro-form in regard to post-translational modification, we generated a variant with two mutations substituting residues Arg-200 and Arg-203 with lysine and alanine, respectively (R200K/R203A, Table 2), rendering the protein impervious to proprotein convertase processing. This variant was expected to be secreted as the full-length proADAMDEC1 form after signal peptide cleavage, starting with residue Ile-31 (Fig. 1A). Surprisingly, R200K/R203A was found present in the medium as three distinct species with apparent molecular masses of 55, 50, and 42 kDa, respectively, rather than as a single unprocessed pro-form (Fig. 3A). Cumulative molecular masses of 55, 50, and 42 kDa, respectively, rather than the untreated protein, consistent with removal of the three N-linked glycans at Asn-61, Asn-184, and Asn-237, respectively (Fig. 3B).

**Sequential Proteolytic Events Remove N-terminal Parts of the ADAMDEC1 Prodomain**—An alanine-scanning mutagenesis of the P3-P3' residues constituting the Arg-56 cleavage site (HKR56-EIK), using the R200K/R203A variant as template, demonstrated that the P1 Arg-56 to alanine substitution (R56A) inhibited all observed proteolytic processing of the prodomain, resulting in the secretion of a single species pro-form. Reduced proteolysis at all cleavage sites in the medium was observed with the K55A P2 variant (Fig. 4A, left). This surprisingly demonstrates that the two cleavage events resulting in the 50- and 42-kDa species depend on initial cleavage after Arg-56. None of the other substitutions had any visible effect on the distribution between proADAMDEC1 species. N-terminal sequencing of the R56A/R200K/R203A secreted protein (Fig. 4A, lane R56) resulted in the sequence IAIKQT, demonstrating that the cleavage after residue 56 was inhibited and confirming the predicted signal peptide cleavage site in proADAMDEC1 at Ala-30 (TQA30-Iai) (Fig. 1A).

**Alanine-scanning mutagenesis of P2-P2’ residues around the Pro-161 cleavage site did not inhibit cleavage at Pro-161**. However, the distribution of the individual processing fragments seems to vary. All four variants give rise to the ~42-kDa species, produced from cleavage at Pro-161, but mutagenesis of the P1' Leu residue appears to result in a slight accumulation of the 50-kDa species (Fig. 4A, right).

**Processing of the prodomain at an unidentified position between Arg-56 and Pro-161 and at Pro-161, giving rise to the 50- and 42-kDa species, respectively, is dependent on the cleavage at residue Arg-56** and ADAMDEC1 proteolytic activity could be detected for the His-362 insert variant (Fig. 2, B and D). Combination of the D362H substitution and the inactivating E353A mutation completely abrogated the proteolytic activity (data not shown), thus confirming the reconstitution of the zinc-binding site in ADAMDEC1.

**Prodomain Glycosylation**—The prodomain of ADAMDEC1 contains two predicted N-linked glycosylation sites (Asn-61 and Asn-184, Fig. 1A). Availability of a single species pro-form containing all predicted glycosylation sites (R200K/R203A/E353A) allowed us to evaluate these by site-directed mutagenesis (Fig. 3B). Substitution of either Asn-61 or Asn-184 with glutamine resulted in increased electrophoretic mobility when compared with R200K/R203A/E353A, demonstrating that both sites are modified by N-linked glycosylation. The apparent molecular masses of R200K/R203A/E353A, N61Q/R200K/R203A/E353A, and N184A/R200K/R203A/E353A were determined to be 55.0, 52.0, and 52.4 kDa, respectively, showing that the effect of one N-linked glycans is a decrease in mobility corresponding to a difference in molecular mass of 2.6–3 kDa. Enzymatically deglycosylated R200K/R203A/E353A migrated with an apparent molecular mass of 47.5 kDa, ~7.5 kDa less than the untreated protein, consistent with removal of the three N-linked glycans at Asn-61, Asn-184, and Asn-237, respectively (Fig. 3B).

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(Figs. 3A and 4A). To investigate whether the ADAMDEC1-dependent processing can take place in trans, we co-incubated proteolytically inactive R200K/R203A/E353A with supernatants from Mock-, ADAMDEC WT-, and D362H-transfected cells, respectively, in the absence or presence of (1,10)-phenanthroline (Fig. 4B). ADAMDEC1 WT and, more efficiently, D362H were able to generate an ~50-kDa proADAMDEC1 fragment (Fig. 3A). This proteolysis was completely inhibited by the addition of (1,10)-phenanthroline, and no cleavage was observed after incubation with Mock cell supernatant. These observations indicate that ADAMDEC1 can perform auto-cleavage of its own prodomain in trans, giving rise to an ~50-kDa species.

ADAMDEC1 Prodomain Confers Partial Catalytic Latency—Several ADAM proteases have been shown to be synthesized aszymogens, and removal of the prodomain is required for proteolytic activity (26–33). To determine whether a similar regulatory mechanism applies to ADAMDEC1, we investigated the proteolytic activity of three ADAMDEC1 variants with different prodomain lengths, but with intact active sites: mature ADAMDEC1 WT (33 kDa, residues 204–470), the R200K/R203A variant predominately found as the 42-kDa species (residues 162–470) cleaved after Pro-161, and the R56A/R200K/R203A variant, exclusively found in the full-length pro-form (57 kDa, residues 31–470) (Fig. 5A). We used azocasein as substrate and included the inactive R200K/R203A/E353A variant as a control (Fig. 5B). Interestingly, the full-length R56A/R200K/R203A pro-form shows significantly reduced activity when compared with mature ADAMDEC1 WT, indicating substantial inhibition of ADAMDEC1 by the intact prodomain (Fig. 5B). Even the 42-kDa R200K/R203A variant exhibited significantly reduced proteolytic activity when compared with the mature WT enzyme, although to a much lower degree than the full-length proenzyme, demonstrating that the 42 remaining residues of the prodomain (res. 162–203) are partially inhibitory to ADAMDEC1. As expected, the R200K/R203A/E353A variant did not cleave azocasein (Fig. 5B).

FIGURE 4. ADAMDEC1 prodomain auto-catalysis and mutational analysis. A, secretion pattern of cells transfected with ADAMDEC1 R200K/R203A-based variants subjected to alanine-scanning mutagenesis around the identified prodomain cleavage sites at position Arg-56 (left) and Pro-161 (right). The alanine-scanned residues (res) and their individual positions (pos) relative to the scissile bond are indicated above the lanes. R56A/R200K/R203A is identified as full-length proADAMDEC1 by Edman sequencing. MW, molecular weight markers. B, co-incubation of R200K/R203A/E353A (RRE) with Mock, ADAMDEC1 WT, or D362H in the absence or presence of 5 mM (1,10)-phenanthroline (Phen). Proteolytic cleavage resulting in an ~50-kDa species is indicated with an arrow.

FIGURE 5. Proteolytic activity of proADAMDEC1. A, reducing anti-ADAMDEC1 Western blot of ADAMDEC1 variants including mature wild-type (WT, 33 kDa,-res. 204–470), R200K/R203A (RR, predominately 42-kDa species, res. 162–470), R56A/R200K/R203A (R56A-RR, 57 kDa, res. 31–470), and the inactivated R200K/R203A/E353A (RRE, 55 kDa, res. 57–470), MW, molecular weight markers. B, activity assay of ADAMDEC1 variants with different prodomain lengths using azocasein as a substrate. Sample means are shown by horizontal lines. *** , p < 0.0001; ns, not significant relative to Mock.
ADAMDEC1 is a putative ADAM-like metzincin metalloprotease with a rare zinc-binding consensus sequence and an unusually short domain structure. It is expressed during differentiation of dendritic cells and is constitutively expressed in macrophages (1, 12). Until now, ADAMDEC1 has primarily been studied on a transcriptional level, where it has been found differently regulated in a number of pathologies (13–18).

We expressed recombinant, human ADAMDEC1 in HEK293 cells and detected the mature and glycosylated protein secreted into the cell supernatant. Other ADAM family members are membrane proteins due to their type-1 transmembrane domain, but secretion of ADAMDEC1 into the cell medium was expected because the short domain structure does not contain a transmembrane domain (1).

Mature ADAMDEC1 produced in HEK293 cells cleaves α2M, resulting in the appearance of a characteristic 85-kDa protein fragment consistent with bait region cleavage (21). We were able to demonstrate covalent trapping of ADAMDEC1 by α2M, as is seen for e.g. ADAM-12 (28), resulting from activation of the α2M cross-linking thioester. Both covalent cross-linking of ADAMDEC1 to α2M and the observed molecular mass of the α2M proteolytic fragment are supportive of cleavage of α2M inside the bait region (34). This also indicates that α2M could be a physiological inhibitor of ADAMDEC1. ADAMDEC1 also cleaves azocasein, resulting in release of azo-labeled peptides.

The amino acid sequence of human ADAMDEC1 contains four predicted N-linked glycosylation sites, which are conserved for higher primates; however, more distant orthologs lack one or more modification site(s). Mutational analysis of the four predicted sites demonstrated modification by N-linked glycosylation at Asn-61, Asn-184, and Asn-237. N184Q and N237Q both exhibited reduced expression levels when compared with mature and pro-form ADAMDEC1, respectively, whereas the expression level of N61Q was comparable with mature and pro-form ADAMDEC1, respectively, as is seen for ADAM-12 (28), resulting from activation of the α2M cross-linking thioester. Both covalent cross-linking of ADAMDEC1 to α2M and the observed molecular mass of the α2M proteolytic fragment are supportive of cleavage of α2M inside the bait region (34). This also indicates that α2M could be a physiological inhibitor of ADAMDEC1. ADAMDEC1 also cleaves azocasein, resulting in release of azo-labeled peptides.

The sequence of human ADAMDEC1 contains a proprotein convertase site (RISR203) (1). Our findings confirm processing of human ADAMDEC1 at RISR203, SLKS consistent with furin-like proprotein convertase activity. In comparison, murine ADAMDEC1 contains two proprotein convertase-like recognition sites, RDQR156 and RTSR203, where in vitro furin processing at the latter site was demonstrated to generate the mature protein (11). Moreover, we demonstrate that full-length proADAMDEC1 has reduced proteolytic activity, and thus the prodomain maintains partial catalytic latency of the protease. A number of ADAMs have been shown to be synthesized as zymogens; however, prodomain processing varies among the members. ADAM-9 (26), -10 (27), -12 (28), -15 (29), -17 (30), and -19 (31) are all activated by a subtilisin/Kex2p-like proprotein convertase (e.g. furin) at a paired basic recognition site (e.g. RXRR), whereas ADAM-8 (32) and -28 (33) are activated by auto-catalytic events, inhibited by substitution of the catalytic glutamic acid residue.

Interestingly, removal of the proprotein convertase recognition site in ADAMDEC1 revealed extensive catalytic processing of the prodomain at three positions. First, proADAMDEC1 is cleaved after Arg-56 by an unidentified protease. This cleavage site has positively charged residues at the P1, P2, and P6 positions relative to the scissile bond and could therefore likely be a substrate for a proprotein convertase family member (38). Prevention of this cleavage, by introducing the P1 variant R56A, inhibited catalytic processing of the prodomain at the two additional sites and resulted in secretion of full-length proADAMDEC1, with the predicted N-terminal sequence after signal peptidase cleavage at Ala-30. However, if the Arg-56 cleavage event is allowed to occur, proADAMDEC1 is further cleaved at two positions. One cleavage occurs after Pro-161, resulting in a 42-kDa species, and the other occurs at an unidentified site N-terminal to Pro-161, resulting in a ~50-kDa ADAMDEC1 species. These events both depend on ADAMDEC1 proteolytic activity because the inactivated pro-form R200K/R203A/E353A does not undergo proteolysis at these sites. Further, this ADAMDEC1-dependent catalysis can occur in trans, demonstrated by co-incubating the inactivated R200K/R203A/E353A with mature ADAMDEC1 WT. This in trans cleavage may be a partial activating event for ADAMDEC1, as is the case for ADAM-8 (32) and ADAM-28 (33). However, in trans cleavage is only observed at the unknown site, giving rise to the 50-kDa species, potentially reflecting differences in the kinetics between cis and trans cleavage of the two ADAMDEC1-dependent proteolysis events.

ADAMDEC1 displays an unusual active site with a non-histidine zinc coordination ligand, which is conserved within all sequenced ADAMDEC1 orthologs available in the Merops database (39). Despite this rare feature, we were able to demonstrate proteolytic activity of the wild-type protein. Interestingly, substituting the aspartic acid constituting the third zinc-binding ligand with histidine (D362H), thereby reconstructing the ADAM active site consensus sequence (HEXXHXXGXXH), increases the proteolytic activity significantly. These data indicate that the change of the zinc ligand chemical properties, from a “medium/soft,” polarizable imidazole nitrogen to a “hard” negatively charged carboxyl oxygen, significantly influences the effectiveness of the zinc ion as a catalyst of hydrolysis. We speculate that this could be due to a reduced potency to activate the coordinated water molecule for nucleophilic attack or a lowered ability to polarize the carbonyl of the scissile bond. Also, the presence of a negatively charged ligand would be expected to reduce the stabilization of the negatively charged transition state (40). Introduction of an aspartic acid residue as the third zinc-coordinating residue also potentially changes the zinc binding geometry of ADAMDEC1, leading to a reduced zinc binding affinity (41, 42). The significant attenuation of the
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proteolytic activity may have evolved to compensate for the short disintegrin domain and loss of C-terminal auxiliary domains, which confers substrate specificity through exo-site interactions in other ADAMS, e.g. ADAM-17 (43) and ADAM-10 (7). Interestingly, inserting a histidine at position 362 (His-362insert), thereby creating an ADAM-like elongated zinc-binding motif followed by an ADAM family-specific aspartic acid residue (HEXXHXGXXHD), completely abrogates the proteolytic activity. We hypothesize that the His-362 insertion may perturb the structure immediately C-terminal to the active site, thereby inhibiting catalytic activity, and suggest that the appearance of the aspartic acid is most likely a result of an evolutionary substitution event. Two distantly related bacterial metzincin families, the snapalysins (ScNP, Merops M7 family) and the thuringilysins (Merops M6 family) (1, 2, 8, 39), share the HEXXXHXGXXD active site sequence with ADAMDEC1. The low overall sequence identity between ADAMDEC1 and members of the M6 and M7 families, however, suggests convergent evolution of this particular zinc-binding sequence (2). Snapalysin from Streptomyces has been shown to be an active protease (44, 45), and a high resolution crystal structure of snapalysin (Protein Data Bank (PDB) 1C7K) shows that the aspartic acid residue functions as the third zinc-coordinating ligand, in an identical manner to what we expect in ADAMDEC1 (46).

In conclusion, ADAMDEC1 is a proteolytic active metzincin metalloprotease despite the rare zinc-coordinating site and C-terminal truncated domain structure. It appears to have evolved with a dampened proteolytic activity because substituting the zinc-ligating aspartic acid with a histidine increases the activity significantly. Decreasing ADAMDEC1 proteolytic activity may have been evolutionarily important to reduce unspecific proteolysis, which we suggest would be the consequence of lacking the C-terminal auxiliary domains.

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