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EXTRACELLULAR CYSTEINES DEFINE ECTOPEPTIDASE (APN, CD13) EXPRESSION AND FUNCTION

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Abstract—Alanyl aminopeptidase (APN) is a surface-bound metallopeptidase that processes the N-terminals of biologically active peptides such as enkephalins, angiotensins, neurokinins, and cytokines. It exerts profound activity on vital processes such as immune response, cellular growth, and blood pressure control. Inhibition of either APN gene expression or its enzymatic activity severely affects leukocyte growth and function. We show here that oxidoreductase-mediated modulations of the cell surface thiol status affect the enzymatic activity of APN. Additional evidence for the pivotal role of extracellular cysteines in the APN molecule was obtained when substitution of any of these six cysteines caused complete loss of surface expression and enzymatic activity. In contrast, the transmembrane Cys24 appears to have no similar function. Enzymatically inactive cysteine mutants were retained in the endoplasmic reticulum as shown by high-resolution imaging and Endoglycosidase H digestion. In the absence of any crystal-structure data, the demonstration that individual extracellular cysteines contribute to APN expression and function appears to be of particular importance. The data are the first to show thiol-dependent modulation of the activity of a typical surface-bound peptidase at the cell surface, probably reflecting a general regulating mechanism. This may relate to various disease processes such as inflammation or malignant transformation. © 2002 Elsevier Science Inc.

Keywords—Aminopeptidase N, Proteindisulfide isomerase, Thiol status, Disulfide bond, Site-directed mutagenesis, Free radicals

INTRODUCTION

Human alanyl aminopeptidase (aminopeptidase N, APN, CD13; EC 3.4.11.2) is a 150 kDa metalloprotease (M1 family, clan MA, gluzincins) [1,2] that shows preferential cleavage of neutral amino acids from the N-terminus of oligopeptides. This type II membrane protein with its zinc-dependent catalytic activity is constitutively expressed on the surface of a wide variety of cells and it has been identified as the leukocyte surface differentiation antigen CD13 [3]. The human APN gene, the coding sequence of which is spread over 20 exons [4], was cloned and mapped to chromosome 15 (q25–q26) [5,6].

Alanyl aminopeptidase is expressed greatly in the small intestine where it participates in the final hydrolysis of ingested nutrients. It is also expressed in the kidney and to a lesser extent in other tissues [7,8]. However, the physiological role of alanyl aminopeptidase on the surface of immune cells has not been fully established. It is believed to be involved in the degradation of neuropeptides [9–11] and cytokines [12,13], and, furthermore, it may well contribute to matrix degradation, angiogenesis, and antigen processing [14–17]. Alanyl aminopeptidase also functions as a receptor for corona viruses [18,19] and CMV [20].

With respect to the hematopoietic system, APN is predominantly expressed on cells of myelo-monocytic lineage [21]. While mature B cells and resting T cells lack detectable APN expression, acute and chronic B cell leukemia show an abnormal surface expression of alanyl aminopeptidase, which is associated with a poor prognosis in the case of adult B-ALL (acute B cell leukemia) [22,23]. A marked increase in the expression of CD13 is found on the surface of T cells activated by mitogens in
vitro [24–26] or derived from local sites of inflammation in vivo [27,28]. Furthermore, T lymphocytes show an activation-dependent increase in APN gene expression [29,30]. Antiproliferative effects result from inhibition of both alanyl aminopeptidase activity and expression [30,31]. This suggests a link between APN expression and leukocyte proliferation as recently reviewed by Lendeckel et al. [32].

The crystal structures of APN or related surface proteases have not yet been resolved. Since a number of these related enzymes share the feature of conserved extracellular cysteine residues, knowledge of the function of these residues may improve the proposed structure models for these enzymes. Cysteine residues are generally of particular importance for formation and maintenance of protein structure and, consequently, function.

To establish the role of cysteine residues in the maintenance of the tertiary structure of a membrane-bound surface protease, modulations of the cell surface thiol status are carried out. Thiols located on the cell surface are mainly represented by proteinogenic cysteiny1 groups. On B-CLL cells, the regulation of this surface thiol status is tightly controlled by membrane-bound oxidoreductases, such as protein-disulfide isomerase (PDI), which catalyze sulfhydryl-disulfide interchange reactions at the cell surface [33]. The inhibition of PDI on viable cells leads to a dramatic rise in total surface thiol expression, which is at least partly due to an increase in cysteine-derived SH groups. Furthermore, γ-glutamyl transpeptidase (GGT) is known to be involved in redox modulations of cell surface protein thiols by its participation in the metabolism of glutathione [34]. Similar to PDI inhibition, inhibition of γ-glutamyl transpeptidase leads to an increased amount of surface thiols. Therefore, modulations in the surface thiol status of U937 cells are carried out by inhibiting PDI by bacitracin and inhibitory antibodies or by inhibiting GGT by acivicin.

To evaluate the role of each individual cysteine in the APN, site-directed mutagenesis was applied. As deduced from its coding sequence [35], human alanyl aminopeptidase, a 967-amino acid protein, contains seven cysteine residues at positions 24, 113, 223, 761, 768, 798, and 834, which are all highly conserved between APNs from different species [36]. The first cysteine is located in the transmembrane region of the enzyme, while the latter six cysteines are located in the extracellular part, which also contains the active site with the HEXXH motif (amino acids 388 to 392). In earlier studies, it was shown that correct folding of the extracellular part of APN occurs independently of the residual protein [37], which suggests that substitution of cysteine 24 will not affect structure and function of alanyl aminopeptidase. In contrast, the effects of substituting cysteine residues in the extracellular part are not as predictable since alanyl aminopeptidase is known to be expressed as a homodimer of two noncovalently linked subunits [38]. However, it does seem reasonable to suppose that disulfide bridges are involved in the formation of the tertiary structure of alanyl aminopeptidase, especially within its C-terminal domain [39].

Our data indicate that (i) oxidoreductase-mediated modulations of the cell surface thiol status significantly affect the enzymatic activity of APN and (ii) the substitution of distinct cysteine residues in the extracellular portion of alanyl aminopeptidase leads to an ER retardation of enzymatically inactive mutant proteins. These findings suggest that all extracellular cysteine residues are involved in the correct formation of the tertiary structure of alanyl aminopeptidase and they support the concept that intramolecular disulfide bridges are formed within the extracellular part of APN. Moreover, these results may imply a general regulatory mechanism for ectopeptidase activity at the cell surface that is based on thio-disulfide interchange reactions.

### MATERIALS AND METHODS

#### Materials

CombiPol DNA Polymerase Mix was obtained from Invitrogen (Berlin, Germany); DNA oligonucleotides (purified by reversed-phase chromatography, 5′-phosphorylated for site-directed mutagenesis) were from BioTez Berlin-Buch GmbH (Berlin, Germany); T4 DNA ligase, all restriction enzymes, Endo Hf, and PNGase F were from New England Biolabs (Beverly, MA, USA); and Transfection reagent Effectene was from QIAGEN (Santa Clarita, CA, USA). The fluorescent probes ER-Tracker Blue-White DPX and BODIPY TR ceramide were purchased from Molecular Probes (Eugene, OR, USA). Cells were routinely grown at 37°C, 5% CO₂, and 94% humidity. Ecr-293 cells were obtained from Invitrogen (Groningen, The Netherlands) and grown in DMEM with 10% FCS (Gibco BRL, Grand Island, NY, USA) and 400 μg/ml Zeocin (Invitrogen). U937 cells (ATCC) were cultured in RPMI 1640 (Gibco BRL)/10% FCS.

#### Modulations of the cell surface thiol status

For inhibition of PDI, U937 cells were cultured in the presence of bacitracin (Sigma Chemical Co., St. Louis, MO, USA) or anti-PDI mAb clone C1D11 [33], both of which specifically inhibit the oxidoreductase activity of PDI. Cultures without bacitracin and with murine IgG (Sigma Chemical Co.) served as controls. Furthermore, U937 cells were cultured in the presence of different
thiol effectors or in thiol-deficient medium for the indicated periods of time. The following were used as thiol effectors: acivicin (Sigma Chemical Co.), inhibiting GGT; BSO (Sigma Chemical Co.), inhibiting \( \gamma \)-glutamylcysteine synthetase; NEM or diamide.

Washed samples were analyzed as follows: flow cytometrically to assess APN immunoreactivity to the anti-APN mAbs clone WM15 and Leu-M7 and for their Ala-pNA (alanylparanitroanilide) hydrolyzing activity [40].

**Construction of a wild-type APN-GFP fusion protein**

APN cDNA was amplified by PCR from U937 cDNA using following primers: primer 1, d(AGC CGG CTA GCC ACC ATC ACC ATG GCC AAG GGC); primer 2, d(GAT CCG CGG TTT GCT GTT TTC TGT GAA).

The fragment was cloned into pEGFP-N1 vector (Clontech, Palo Alto, CA, USA), which resulted in the expression vector pEGFP-APN-C24G as a template, yielding the expression vector pEGFP-APN-C24G/C834G. The resulting constructs containing the different mutations were analyzed by restriction analysis, immunoblotting, and fluorescence of the expressed mutant GFP fusion proteins.

**Transfection of EcR-293 cells**

The constructed vectors were transiently transfected into EcR-293 cells using the Effectene transfection reagent (QIAGEN). In 6-well format, \( 3 \times 10^5 \) cells per well were seeded the day prior to transfection. For complex formation, 0.4 \( \mu \)g DNA, 3.2 \( \mu \)l Enhancer, and 5 \( \mu \)l reagent were used. Cells were cultured for 2 d before assayed.

**Fluorescence microscopy**

Transfected cells grown on glass coverslips were analyzed using an Axiovert 135 TV (Zeiss, Göttingen, Germany) equipped with the Spot RT Slider camera, and Meta Series and Auto Deblur software (Visitron Systems, Puchheim, Germany) at 100× magnification. GFP fluorescence was detected using the filter set 10 (Zeiss).

Colocalization studies with organelle-specific fluorescent probes were performed. Cells grown on coverslips for 2 d were washed once with HBSS/10 mM HEPES. To stain the ER, cells were incubated with 200 nM ER-Tracker Blue White DPX in HBSS/10 mM HEPES for 30 min at 37°C and washed once with PBS. Blue fluorescence was detected using filter set 01 (Zeiss). To stain the Golgi, cells were incubated with 5 \( \mu \)M BODIPY TR ceramide-BSA-complex in HBSS/10 mM HEPES for 10 min at 37°C and washed once with PBS. Red fluorescence was detected using filter set 00 (Zeiss).

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**Table 1. Mutagenic Oligonucleotides for Site-Directed Mutagenesis of APN**

| Name       | Sequence                       |
|------------|--------------------------------|
| C24G-s     | 5′-GGCGTGGCCAG CGCGTGCCAC AATCATTGCA CTTGCA |
| C24G-as    | 5′-CGCACTTGGCC ATGTGCGCTC GACGCC |
| C113F-s    | 5′-ACGCGGGCTT TCACGCTAA AAGGGCGGT |
| C113F-as   | 5′-GACGCGGGCT TCACGCTAA AAGGGCGGT |
| C223W-s    | 5′-GGGAAGCCCT TCGCTACTG CAGGAGTTC |
| C223W-as   | 5′-GCAGCGGCC AGCCGGCTCA TGGCC |
| C761G-s    | 5′-GCGATGCAAG CCGCCGCTC GAACCGAGTT |
| C761G-as   | 5′-CAGACCATC TCCTCAC AACGGAGTTC |
| C768G-s    | 5′-AGAGAGCTCC GTGAACTCCGT TTGGAGC |
| C768G-as   | 5′-TGGAACTCCG TTGGAGC GCCATCAGCA CCGCC |
| C798W-s    | 5′-GGGACTCC GTGAACTCCGT TTGGAGC |
| C798W-as   | 5′-CTGGGCGGATA CCGCTTACTG CAGAG |
| C834G-s    | 5′-GCAGCGGCC AGCCGGCTCA TGGCC |
| C834G-as   | 5′-GTACGCAAGAC TCCTTGCTGC eGGCCAGGGC TGGGGG |

Mismatches with the template are indicated by bold lowercase letters.
Cytofluorimetric analyses

Surface staining of cells was performed at 4°C for 30 min using anti-CD13 (clone WM15, Pharmingen [Heidelberg, Germany]) and WM15 PE-conjugate, Dianova (Hamburg, Germany); clone Leu-M7, PE-conjugate, Becton Dickinson [Mountain View, CA, USA]); and anti-GFP antibodies (Living Colors Peptide antibody, Clontech). After two washes in cold PBS, cells were incubated with goat-antimouse PE-conjugate (Dianova) for WM15 staining or donkey-antirabbit PE-conjugate (Dianova) for anti-GFP staining, both at 4°C for 30 min. Cells were washed twice in cold PBS prior to analysis. For staining of cytoplasmic epitopes, permeabilization of cells was performed with a cell permeabilization kit (Fix & Perm, An der Grub Bio Research GmbH, Kaumberg, Austria) before staining. Cytofluorimetric analysis was performed using FACS Calibur (Becton Dickinson) on readings from 5000 cells per sample. Data are given as mean fluorescence intensities (mfi) ± SEM.

Preparation of the particulate fraction

Cells (2 × 10⁶) were resuspended in PBS and lyzed by sonication on ice for 30 min. The suspension was centrifuged at 60,000 × g for 30 min and then resuspended in either 500 μl PBS (for activity measurements) or 100 μl denaturing buffer (New England Biolabs) containing 1% β-mercaptoethanol and 0.5% SDS (for digestion and blotting analyses) using sonication.

Digestion with Endo Hf and PNGase F

The particulate fraction was boiled for 10 min. For Endo Hf digestion, 10 μl of the particulate fraction were incubated with 1 μl 10 × G5 buffer and 2000 U Endo Hf for 1 h at 37°C. The particulate fraction treated the same way, except for the addition of Endo Hf, served as a control. PNGase F digestion was carried out in 10 μl of the particulate fraction using 1 μl 10 × G7 buffer, 1 μl NP40 (10%), and 500 U PNGase F for 1 h at 37°C.

Immunoblotting

The particulate fractions resuspended in denaturing buffer were supplemented with two volumes Laemmli buffer (Bio-Rad, Hercules, CA, USA) and boiled for 2 min. Probes were separated on 4–12% gradient NuPage gels using MOPS-SDS running buffer (Invitrogen) at 15°C. Two different protein markers, SeeBluePlus2 Prestained Standards (Novex) and Prestained Precision Protein Standards broad range (Bio-Rad), were used to estimate protein size. After blotting on to a Protran BA-85 membrane (Schleicher & Schüll, Keene, NH, USA), APN-GFP fusion proteins were detected by means of Living Colors Peptide antibody (Clontech), goat-antirabbit-HRP (horseradish peroxidase, New England Biolabs), and SuperSignal West Dura Extended Duration chemoluminescence substrate (Pierce, Bonn, Germany).

Enzymatic assay

The particulate fractions resuspended in PBS were used to determine the enzymatic activity. To normalize the activities, the GFP fluorescence of each probe was measured using a luminescence spectrophotometer LS 50 (Perkin Elmer, Norwalk, CT, USA) with excitation at 488 nm and emission at 507 nm. Neutral aminopeptidase activity was determined by measuring the hydrolysis of Ala-pNA [40]. The specific activity was calculated as follows:

\[ \text{a}_{\text{specific}} = \frac{\text{a}_{\text{measured}} \times I_{\text{GFP}} \text{ (nontransfectedcells)}}{I_{\text{GFP}}} \]

RESULTS

Effects of PDI inhibition on the enzymatic activity of surface-bound APN

The myelo-monocytic cell line U937, coexpressing APN and the oxidoreductase PDI at the cell surface, was used to elucidate the influence of oxidoreductase-mediated changes of the cell surface thiol status on APN activity. Treatment of U937 cells with the PDI inhibitor bacitracin resulted in a concentration- and time-dependent increase in their immunoreactivity to the anti-APN clone Leu-M7 (Table 2). In contrast, the immunoreactivity of the anti-APN clone WM15 was found to be significantly decreased. After 48 h, and more pronounced after 72 h, the addition of 100 μM bacitracin led to a 1.6-fold decrease in WM15 reactivity compared to the control. In parallel, the Ala-pNA hydrolyzing activity was reduced to 78% after 48 h and to 68% after 72 h (Table 2) in comparison to untreated controls.

The possibility that these changes resulted from effects other than inhibition of oxidoreductase activity was excluded by showing that the anti-PDI mAb clone C1D11, known to inhibit PDI catalytic activity [33], affected APN activity equally. After 48 h in presence of anti-PDI mAbs, a concentration-dependent increase in Leu-M7 immunoreactivity was observed, whereas the WM15 reactivity was significantly decreased. As shown in Fig. 1, the opposite kinetics of the two anti-APN mAb clones became evident at concentrations as low as 0.2 μg/ml anti-PDI mAb C1D11 (mean fluorescence intensities of 573.7 ± 12.4 and 402.3 ± 3.8 for Leu-M7 and WM15, respectively) and enhanced at 2 μg/ml C1D11.
(mfi of $686.0 \pm 3.6$ for Leu-M7 and $272.0 \pm 5.8$ for WM15). Murine IgG, supplemented at concentrations equal to anti-PDI mAb, failed to show any effect on APN immunoreactivity.

**Effects of other thiol effectors on the enzymatic activity of surface-bound APN**

To compare the observed effects of the PDI inhibition at the surface of U937 cells with effects caused by other thiol effectors, U937 cells were cultured in the presence of 150 $\mu$M acivicin, 500 $\mu$M BSO, 100 $\mu$M NEM, or 50 $\mu$M diamide (Table 3). In contrast to PDI inhibition, these thiol effectors did not cause significant alterations of the immunoreactivity against both anti-APN mAb clones. The diminished reactivity against both mAb clones after BSO treatment may represent a decreased amount of APN-molecules at the cell surface.

**Cellular and functional characteristics of the APN-GFP fusion protein**

To determine the subcellular localization of wild-type and mutated alanyl aminopeptidase proteins, fusion proteins, with enhanced GFP (EGFP) fused in frame to the C-terminus of APN, were constructed. The C-terminal attachment of GFP to wild-type APN did not change either its subcellular localization or its function: the typical surface expression of wild-type alanyl aminopeptidase was completely retained (Fig. 2A). This implies that protein folding, glycosylation pattern, and intracellular transport were not markedly influenced. Results of cytofluorimetric analyses confirmed the intense surface expression of APN-GFP when using either anti-GFP or anti-CD13 antibodies. The mean fluorescence intensities were increased significantly when compared with non-transfected cells (Table 4). Furthermore, the neutral aminopeptidase activity of the wild-type enzyme was fully preserved. The Ala-pNA hydrolyzing activity of the particulate fraction of EcR-293 cells was significantly increased upon transient transfection with pEGFP-APN in comparison to nontransfected cells that showed only 47.7% of wild-type activity (Fig. 3, black columns).

Further evidence for the structural integrity of the wild-type APN-GFP fusion protein was obtained from SDS-PAGE and immunoblot analysis of the particulate fraction of EcR-293 cells. Using an anti-GFP antibody, two bands of approximately 170 and 190 kDa were detected. The band with the higher molecular weight corresponds to the 166 kDa monomeric form of mature APN enlarged by 27 kDa of monomeric GFP (Fig. 4, lane 2). The double band that was detected represents different glycosylation forms, as confirmed by PNGase F treatment, which leads to a single band of approximately 140 kDa. Endoglycosidase H digestion resulted in degradation of the lower molecular weight species, indicating that this was a mannose-rich glycosylation form of APN (Fig. 5).

**Integrity of the mutant proteins**

To examine the integrity of the mutants, APN-GFP fusion proteins were expressed in EcR-293 cells and analyzed for GFP-fluorescence by means of fluorescence microscopy. Protein size was determined by using reduc-
ing SDS-PAGE followed by Western blotting and detection with an anti-GFP antibody.

All constructs yielded intense GFP fluorescence upon transfection into EcR-293 cells. Furthermore, all mutant proteins showed immunoreactivity to an anti-GFP-Ab. Compared to the wild-type APN-GFP, no apparent alterations in the electrophoretic mobility in SDS-PAGE were detected for the C24G mutant. In contrast, all other mutants showed a single band only, which probably corresponded to the Endoglycosidase H-sensitive form of the wild-type protein (Fig. 4). These findings indicate that the primary structure of the protein was not severely impaired. In contrast, the glycosylation pattern seemed to be affected by substitution of extracellular cysteine residues.

The expression levels of the individual APN-GFP fusion proteins showed significant variations (Fig. 4). To compensate for these differences, which were due to variations in the transfection efficiency and expression levels, the intensity of GFP-specific fluorescence was used to normalize the enzymatic activity data.

### Table 3. Immunoreactivity of Anti-APN mAbs on U937 Cells After Modulation of the Thiol Status

| Effector  | Concentration (µM) | Time (h) | n  | Leu-M7 Ratio | SEM | WM15 Ratio | SEM |
|-----------|--------------------|----------|----|--------------|-----|------------|-----|
| Acivicin  | 150                | 1        | 3  | 1.02         | 0.04| 1.06       | 0.13|
| Acivicin  | 150                | 4        | 4  | 0.92         | 0.10| 0.94       | 0.41|
| BSO       | 500                | 4        | 4  | 0.92         | 0.14| 0.71       | 0.11|
| NEM       | 100                | 1        | 4  | 0.86         | 0.22| 1.10       | 0.31|
| Diamide   | 50                 | 1        | 4  | 0.96         | 0.04| 0.88       | 0.09|

All data are given as a mean ratio between mfi values for effector-treated and untreated cells ± SEM.

### Enzymatic activity of mutant alanyl aminopeptidase

EcR-293 cells, transiently transfected with mutant APN-GFP, were analyzed for neutral aminopeptidase activity (Fig. 3). It was confirmed that all mutants showed high levels of GFP fluorescence as well as immunoreactivity to an anti-GFP-mab, indicating a proper folding of the GFP chromophore. Therefore, GFP flou-
Table 4. Cytofluorimetric Analyses of APN-GFP Fusion Proteins

|            | Leu-M7 (%) | WM15 (%) |
|------------|------------|----------|
| Control    | 0.0        | 0.0      |
| Wild-type  | 100.0      | 100.0    |
| C24G       | 87.6       | 86.1     |
| C113F      | 15.5       | 2.3      |
| C223W      | 17.3       | 1.3      |
| C761G      | 2.2        | 1.9      |
| C768G      | 11.1       | 3.4      |
| C798W      | 1.9        | 2.1      |
| C834G      | 3.4        | 2.1      |
| C24/834G   | 2.9        | 2.2      |

Data given as relative mean fluorescence intensities; staining with mAbs after permeabilization of the cells.

Surface expression of mutant APN-GFP fusion proteins was analyzed by fluorescence microscopy and cytofluorimetric analysis in EcR-293 cells transiently transfected with the appropriate expression vectors. The substitution of cysteine residue 24 had no effect on the surface expression of APN (Fig. 2B). This is in full agreement with the corresponding enzymatic activity, which also was not affected by the substitution of this residue. All others provoked changes in the expression pattern of the APN-GFP fusion proteins, with apparently predominant intracellular localization (Figs. 2C–2I). This was confirmed by cytofluorimetric analysis using an anti-GFP antibody. A clear-cut surface expression was observed only in cells transfected with either wild-type APN-GFP or the C24G mutant (27% positive cells, mfi of 200.2 and 224.1, respectively, compared to less than 4% positive cells and mfi < 45 for all other mutants).

Therefore, detailed colocalization studies were used to investigate the subcellular localization. We used different reference markers for Golgi and ER structures. These studies were performed with all the mutants that did not contain C24G substitution. Mutants containing the C24G substitution were not included since this mutation failed to show any changes in the expression pattern of APN.

For selective staining of ER, ER-Tracker Blue-White DPX was used. The resulting fluorescence showed a typical ER-staining pattern around nonfluorescent cell nuclei (Fig. 6B). Using BODIPY TR ceramide as a trans-Golgi probe, a staining pattern that was evidently different from the ER staining was observed (Fig. 6E). As illustrated in Fig. 6 using the C113F mutant, a clear overlap of the GFP-derived fluorescence with an organelle-specific marker was only observed with the ER staining. Some partial overlap in the case of Golgi staining may have been due to high background fluorescence. Similar results were obtained for the other mutants, indicating that all enzymatically inactive mutants are retained in the ER.

![Fig. 3. Neutral aminopeptidase activity of EcR-293 cells expressing different APN-GFP fusion proteins. Specific activities of the particulate fractions of transiently transfected EcR-293 cells were measured. Nontransfected cells served as a control. Every experiment was normalized using the mean of the wild-type APN-GFP fusion protein. Results are given as percentages of wild-type activity (46.6 pkat/10⁹ cells). *A significant increase compared with the control; #a significant decrease compared with the wild-type protein (one-way ANOVA, p < .05).](image-url)
To confirm these results, analysis of Endoglycosidase H sensitivity and PNGase F treatment were carried out for all APN-GFP fusion proteins. Treatment with Endo H distinguishes high mannose (sensitive) from complex (insensitive) sugars, and, thus, indicates whether a protein is able to leave the ER. Digestion with PNGase F enables one to determine whether different protein forms are due to variations in the glycosylation pattern.

In all cases, the PNGase F digestion resulted in the formation of one single band of lower molecular weight (Fig. 6). A band of approximately 140 kDa was detected (unglycosylated APN (115 kDa) enlarged by the monomeric GFP). Corresponding to the results for wild-type protein, the double band detected in the case of the C24G mutant was reduced to a single band after PNGase F treatment, which suggests that two different glycosylation forms are present in the particulate fraction.

Treatment with Endo H revealed the presence of Endo H-sensitive proteins in all particulate fractions. One additional insensitive band of higher molecular weight, corresponding to the complex glycosylated mature form of APN that is transported to the cell surface, was found only for the wild-type APN and the C24G mutant. In contrast, in all other cases the GFP fusion protein was completely sensitive to Endo H, confirming the results of the colocalization studies, which showed a clear ER retardation of these proteins.
Immunoreactivity against anti-CD13 monoclonal antibodies

Cytofluorimetric analysis of permeabilized cells unexpectedly revealed that the substitution of cysteine residues 761, 798, and 834 was obviously associated with a complete loss of the epitope recognized by the anti-CD13 monoclonal antibody clone Leu-M7. Thus, both the single mutants C761G, C798W, C834G, and the double mutant C834G/C24G showed Leu-M7 immunoreactivity of less than 4% of the wild-type mfi, while in all other cases mfi values of more than 10% were retained (Table 4). Notably, none of the enzymatically inactive mutants bound to the anti-CD13 monoclonal antibody clone WM15 (Table 4).

DISCUSSION

Cysteine residues are known to be of particular importance in the formation and maintenance of tertiary protein structures. The aims of the present study were (i) to elucidate the influence of oxidoreductase-mediated modulations of the cell-surface thiol status on the enzymatic activity of APN on the surface of viable cells, and (ii) to use site-directed mutagenesis to study the functional role of each individual cysteine residue of APN on surface expression/subcellular localization, enzymatic activity, and binding to anti-CD13 mAbs.

Utilizing glutathione, proteindisulfide isomerase catalyzes the formation of inter- and intramolecular disulfide bridges [41]. The inhibition of PDI by either bacitracin or inhibitory monoclonal antibodies on the cell surface results in reduced enzymatic activity of APN. Thus, it is evident that the cysteine-linked sulfhydryl-disulfide turnover at the cell surface can modulate APN enzymatic activity. As expected from studies that characterized the anti-APN mAb clone WM15 as being inhibitory [42], the immunoreactivity of this clone was equally diminished. This effect is not due to a loss of APN on the cell surface, since, under the same conditions, the reactivity to the anti-APN mAb clone Leu-M7, which does not affect APN activity, was found to be increased. This could be due to the larger distance of the relevant epitope from the catalytic center, which was concluded from the finding that substitution of cysteine residues 761, 798, and 834 did cause a complete loss of Leu-M7 binding. Thus, the increase of Leu-M7 immunoreactivity could result from better access to the epitope when the protein is unfolded due to impaired disulfide bridge formation.

In contrast, treatment of U937 cells with the GGT inhibitor acivicin as well as treatment with other thiol effectors failed to give similar effects on anti-APN immunoreactivity. These findings strongly suggest that effects found after inhibition of PDI are not only due to the detected increase in cell surface thiols but also are directly linked to the enzymatic activity of PDI. It seems possible that PDI primarily regulates which disulfide links are formed, thus controlling the tertiary structure of APN as well as of other ectoenzymes.

Furthermore, the function of individual cysteine residues in the APN molecule for establishing correct structure and, consequently, enzymatic activity and surface expression, has not been previously studied. However, APN shares considerable sequence similarity with pyro-

Fig. 6. Colocalization studies of EcR-293 cells expressing mutant APN-GFP fusion protein. Cells were stained with ER-Tracker Blue-White DPX or BODIPY TR ceramide. Colocalization is shown for the C113F mutant. The GFP-derived fluorescence is shown in green and the fluorescence of the ER or Golgi marker is shown in red. (A) GFP signal of C113F; (B) ER staining of the same cells; (C) colocalization of C113F with ER; (D) GFP signal of C113F; (E) Golgi staining of the same cells; (F) colocalization of C113F with Golgi; bar = 10 μm.
glutamyl peptidase II (thyrotropin-releasing hormone degrading ectoenzyme, degrading enzyme, TRH-DE), which also belongs to the M1 family of Zn-dependent metalloproteases (Fig. 7) [43–45]. This is strikingly evident for the four cysteine residues clustered in the C-terminal region. A recent study identified TRH-DE as a disulfide-linked (via Cys68) homodimeric ectopeptidase [46]. However, covalent dimerization does not appear to be essential for its membrane expression or its enzymatic activity. Similar results were obtained in previous biochemical studies on the dimeric assembly and surface expression of APN [47]; dimerization does not seem to involve covalent bonds [38]. APN is synthesized in the RER, where it is cotranslationally inserted into the membrane and glycosylated [48,49]. It has long been assumed that high mannose content, as distinct from complex glycans characteristic of proteins that did not reach the Golgi apparatus, confirmed this ER retardation. Taken together, these data indicate that individual cysteine residues in the extracellular part of alanyl aminopeptidase are important for tertiary structure and function of APN. However, all mutants lacking one extracellular cysteine residue could not reach the cell surface. Therefore, it was not possible to directly prove that all of these residues are critical for the surface activity once the protein has reached the cell surface. Whether the dimeric assembly of APN is also impaired by the substitution of cysteine residues remains a challenging task for future studies.

To the best of our knowledge, the data provide the first evidence of a modulation of the expression and activity of a membrane-bound exopeptidase, alanyl aminopeptidase, by oxidoreductase-mediated modulation of cell-surface thiols. It is tempting to speculate that this represents a general mechanism that could apply to other surface-bound enzymes. In support of this view, studies on thiol-disulfide interchange-mediated modulations of angiotensin-converting enzyme (ACE, CD143) on activated monocytes revealed similar results. After inhibition of PDI, the ACE enzymatic activity was diminished, whereas no quantitative changes in the surface expression of ACE molecules were observed (data not shown).

Various ectopeptidases are thought to be involved in a number of disease processes, such as inflammation and malignant transformation, which are also known to be associated with severe impairment of the thiol-disulfide status. Thus, thiol-dependent regulation of membrane-bound peptidases could be a contributing cofactor in the initiation and progression of disease. APN and other surface peptidases (e.g., dipeptidyl peptidase IV, CD26) reportedly function as regulators of leukocyte growth and function [32,54], and, therefore, inhibitors of their enzymatic activities are potent immunosuppressive, anti-inflammatory, and anti-proliferative compounds [15,55,56]. The novel concept of a thiol-dependent regulation of surface activities introduces a new regulatory mechanism, which may contribute to the modulation of both the normal and pathological immune responses, and, consequently, to development or manifestation of autoimmune disease, allograft rejection, or malignant growth.

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ABBREVIATIONS

Ala-pNA—alanylparanitroanilide

APN—aminopeptidase N

B-ALL—acute B cell leukemia

BSA—bovine serum albumin

BSO—buthionine sulfoximine

CD—cluster of differentiation

CML—chronic myeloid leukemia

CMV—cytomegalovirus

ER—endoplasmic reticulum

FACS—fluorescence-activated cell sorting

FCS—fetal calf serum

GFP—green fluorescent protein

HRP—horseradish peroxidase

mi—mean fluorescence intensity

NEM—N-ethylmaleimide

NHL—non-Hodgkin lymphoma

PAGE—polyacrylamide gel electrophoresis

PBS—phosphate-buffered saline

PE—phycocerythrin

PNGase F—peptide N-glycosidase F

SDS—sodiumdodecylsulfate

TRH-DE—thyrotropin-releasing hormone-degrading ectoenzyme