Involvement of Glutamine-238 in the Substrate Specificity of Human Laeverin/Aminopeptidase Q

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Human laeverin/aminopeptidase Q (APQ) is a novel member of the M1 family of zinc aminopeptidases and is specifically expressed on the cell surface of extravillous trophoblasts. In this study, we examined the significance of Gln-238 of laeverin/APQ, a putative S1 site residue, by site-directed mutagenesis for its enzymatic activity and substrate specificity. Replacement of Gln-238 with Ala caused a significant change in substrate specificity rather than a decrease in enzymatic activity. These results indicate that Gln-238 is important for the substrate specificity of laeverin/APQ. In addition, our data suggest that direct electrostatic interaction between substrate and S1 site of the enzyme is not involved in the mutant enzyme’s preference for basic amino acids.

Key words aminopeptidase; laeverin; endoplasmic reticulum aminopeptidase 1; site-directed mutagenesis; substrate specificity

Aminopeptidases hydrolyze the N-terminal amino acid of proteins or peptide substrates. Among them, the M1 family of zinc aminopeptidases (gluzincins) shares the consensus GAMEN and HEXXH(X)18E motifs essential for enzymatic activity. This family, which consists of 11 enzymes in humans, 1–3 plays important roles in several pathophysiological processes, such as angiogenesis, cell cycle regulation, reproduction, memory retention, blood pressure control, and antigen presentation to major histocompatibility complex (MHC) class I molecules. 4–12

Laeverin was originally identified as a cell-surface protein specifically expressed on human extravillous trophoblasts. 13 cDNA cloning of human laeverin revealed that it contains both consensus motifs and thus is a novel member of the M1 family. Another group also predicted the existence of the ‘leaverin’ gene in the human genome through a genomic search and named it aminopeptidase Q (APQ). 14 Recently, we established a large-scale production system of the enzyme and characterized its enzymatic properties in detail. 15 We found that the enzyme degraded several placenta-derived peptide hormones, such as angiotensin III, endokinin C, and kisspeptin-10. Furthermore because its exopeptidase motif GAMEN is uniquely composed of the HAMEN sequence, we examined the enzymatic significance of His-379 of human laeverin/APQ and characterized its enzymatic properties in detail. 15 Considering the susceptibility of these peptides and their specific expression in the placenta, we speculated that laeverin/APQ plays important roles in the maintenance of normal pregnancy in humans.

In our previous work, we identified a residue affecting enzymatic activity and the substrate specificity of endoplasmic reticulum aminopeptidase (ERAP)1, a member of the M1 aminopeptidase family. 16 In particular, replacement of Gln-181 with Asp caused a marked change in the substrate specificity of ERAP1. On the other hand, replacement with Ala led to almost complete loss of enzymatic activity. To further analyze the residue affecting enzymatic activity and substrate specificity of the M1 family of aminopeptidases, we examined in the present study the effect of replacing Gln-238 of human laeverin/APQ with Ala. We found that Q238A laeverin/APQ retained substantial enzymatic activities with marked change of its substrate specificity, indicating the significance of this residue for the enzymatic characteristics of laeverin/APQ. Data presented in this study imply that direct electrostatic interaction is not involved in the mutant enzyme’s preference for basic amino acids, which was suggested in the previous work. 17

MATERIALS AND METHODS

Molecular Modeling of Laeverin/APQ The published X-ray crystallographic structure of Thermoplasma acidophilum Tricorn-interacting factor F3 (TIF3) (Protein Data Bank code 1Z5H) 18 was used as a template for modeling the catalytic site of ERAP1 with the three-dimensional Jigsaw Protein Server (http://www.bmm.icnet.uk/~3djigsaw/). The structure was displayed using the CueMol program (Ishitani, R., CueMol: Molecular Visualization Framework, http://cuel.
mol.sourceforge.jp).

Site-Directed Mutagenesis To obtain a large amount of recombinant human laeverin/APQ, we slightly modified its cDNA by polymerase chain reaction (PCR). Briefly, to express the enzyme as a soluble protein, the coding sequences for the cytosolic and transmembrane region of the enzyme (Met1–Gln64) were replaced with that for human trypsin II signal peptide (i.e. MNLLILTFVAAVAA). 15 Hexahistidine tag was added at its C-terminal end. Amplified cDNA was cloned into the BssHIII–XhoI site of the baculovirus transfer vector pFastbac-1 (Invitrogen, CA, U.S.A.). The cDNA encoding mutant Q238A laeverin/APQ was generated using a QuickChange site-directed mutagenesis kit (Stratagene, La Jolla, CA, U.S.A.). The primers used to replace Gln-238 with Ala in human laeverin/APQ cDNA were as follows: sense primer, 5‘-GCCCTGTAGCGTCCGGTCTGGAACCAAC- ATTTG-3’, and antisense primer 5‘-CAAATGTTGGTTCC- AGACGGACGCTAACAGGGC-3‘.

The sequences of the products were confirmed by automated sequencing on an Applied Biosystems model 3730.

Expression and Purification of Recombinant Wild-
Type and Mutant Laeverin/APQs in the Baculovirus System

The pFastbac-1 vectors containing laeverin/APQ cDNAs were transformed to competent DH10bac *Escherichia coli* cells harboring the baculovirus genome (bacmid) and a transposition helper vector (Invitrogen). Subsequently, insect SF9 cells were transfected with recombinant bacmids using the Cellfectin reagent (Invitrogen). After a 3-d incubation period, recombinant baculoviruses were isolated and used to infect SF9 cells at a multiplicity of infection of 0.1. Three days after infection, the amplified viruses were harvested.

For the production of laeverin/APQ, SF9 cells were grown at 27 °C in 100 ml SF-900III medium (Gibco, CA, U.S.A.) and 1.5×10⁶ cells/ml were infected at a multiplicity of infection of 1—3. After 3 d, the culture medium was collected after centrifugation at 5000 g for 15 min.

The culture medium was loaded onto a hydroxyapatite (Nacalai Tesque, Kyoto, Japan) column (bed volume: 10 ml) pre-equilibrated with 5 mM phosphate buffer (pH 7.5). After extensive washing with the same buffer, laeverin/APQ was eluted from the column with 100 mM phosphate buffer (pH 7.5). The eluate was then loaded onto a Ni²⁺-chelating Sepharose (Pharmacia, Uppsala, Sweden) column (bed volume: 1 ml) pre-equilibrated with 10 mM phosphate buffer (pH 7.5) containing 0.1 M NaCl. After extensive washing with 10 mM phosphate buffer (pH 7.5) containing 0.1 M NaCl and 5 mM imidazole, laeverin/APQ was eluted with 10 mM phosphate buffer (pH 7.5) containing 0.1 M NaCl and 150 mM imidazole. The laeverin/APQ-containing fractions were extensively dialyzed against 25 mM Tris–HCl buffer (pH 7.5) containing 0.125 M NaCl, concentrated with an ultrafiltration membrane and stored at -20 °C prior to use.

Measurement of Aminopeptidase Activity of Laeverin/APQs

Aminopeptidase activities of wild-type and mutant laeverin/APQs were routinely determined by endpoint assay with various fluorogenic aminoacyl-4-methylcoumaryl-7-amides (aminoacyl-MCAs) as substrates. The reaction mixture containing 100 μM aminoacyl-MCA and 1 μg/ml enzyme in 0.5 ml of 25 mM Tris/HCl buffer (pH 7.5) was incubated at 37 °C for 15 min. The reaction was terminated by adding 2.5 ml of 0.1 M sodium acetate buffer (pH 4.3) containing 0.1 M sodium monochloroacetate. The amount of 7-amino-4-methylcoumarin released was measured by spectrofluorophotometry (F-2000; Hitachi) at an excitation wavelength (360 nm) and an emission wavelength (460 nm), and is given in arbitrary units. To determine the kinetic parameters, the reaction mixture containing various concentrations of substrates and either wild-type or mutant laeverin/APQ in 500 μl of 25 mM Tris/HCl buffer (pH 7.5) was incubated at 37 °C for 5 min. The amount of 7-amino-4-methylcoumarin released was then measured. The kinetic parameters (K_m and V_max) were calculated using Lineweaver–Burk plots.

Materials

Asp-, benzyl-Cys-, and Glu-MCAs were purchased from BACHEM AG (Bubendorf, Switzerland). Ala-, Arg-, Leu-, Lys-, Met, and Phe-MCAs were obtained from the Peptide Institute (Osaka, Japan). Bestatin was purchased from Sigma.

RESULTS AND DISCUSSION

Molecular Modeling of the Catalytic Pocket of Laeverin/APQ

In the previous work, we showed that Gln-181 of ERAP1 was critical for the enzymatic activity and substrate specificity of ERAP1. Alignment of ERAP1 and laeverin/APQ indicates that Gln-238 of laeverin/APQ occupies the corresponding site of Gln-181 of ERAP1. Using TIF3 as a template, we modeled the structure of the catalytic pocket of human laeverin/APQ (Fig. 1). As in the case of ERAP1, Gln-238 of laeverin/APQ is expected to be located near the catalytic HEXXH(X)₄E motif and at a possible S1 site of the enzyme, suggesting the importance of this residue for enzymatic activity. To elucidate the role of Gln-238 of human laeverin/APQ in its enzymatic properties further, this residue was substituted with Ala to generate Q238A laeverin/APQ, and its enzymatic property was compared with wild-type enzyme. We chose an Ala mutant to check the significance of Gln-238 residue, because the substitution of Gln-181 residue with Ala caused almost complete loss of the enzymatic activity of ERAP1, thus indicating the essential role of this residue in the enzymatic activity.

Characterization of the Hydrolytic Activities of Wild-Type and Q238A Laeverin/APQs

Figure 2 shows the enzymatic activities of wild-type and Q238A laeverin/APQs toward various substrates. As shown previously, the wild-type enzyme cleaved Leu-MCA most efficiently, followed by Met-, Arg- and Lys-MCA. Hydrolysis of Ala-, Phe-, and benzyl-Cys-MCA was also observed. On the other hand, in sharp contrast to Q181A ERAP1, which showed little hydrolytic activity toward substrates tested, Q238A laeverin/APQ retained substantial activity toward basic amino acid substrates, Arg and Lys-MCA. Little activity was detected toward other substrates tested. These results indicate that while wild-type enzyme shows broad substrate specificity, enzymatic activity of the mutant enzyme is rather restricted to basic amino acids.

Table 1 shows the kinetic parameters of wild-type and Q238A laeverin/APQs using Leu-MCA and Arg-MCA as substrates. The calculated kcat/Km value from a Lineweaver–Burk plot of wild-type enzyme for the most efficient substrate Leu-MCA was 451±16 μM⁻¹ s⁻¹. The catalytic efficiency (kcat/Km) of the wild-type enzyme for Arg was 205±33 μM⁻¹ s⁻¹, about 2.3-fold less than for Leu-MCA. On the other hand, when comparing the hydrolytic activity of mutant enzyme with that of the wild-type toward Arg-MCA, mutant enzyme showed comparable efficiency (232±49 μM⁻¹ s⁻¹). Both K_m and kcat values of mutant laeverin/APQ were about 2-fold less than those of wild-type enzyme, suggesting that replacement of Gln-238 with Ala slightly decreased the enzyme recognition mechanism toward basic amino acids. It showed little activity toward Leu-MCA with K_m value of 2.79±0.12 μM⁻¹ s⁻¹, because of a marked decrease in the kcat value. These results indicate that mutant enzyme retained substantial enzymatic activity toward Arg-MCA and its substrate specificity was different from that of the wild-type enzyme and showed rather a restricted pattern. Taken together, we concluded that the substitution of Gln-238 with Ala caused a change of substrate specificity rather than total decrease of the enzymatic activity toward substrates, which is quite different from the case of ERAP1.
Effect of Bestatin on the Enzymatic Activities of Wild-Type and Q238A Laeverin/APQs

Because laeverin/APQ is unique in that bestatin inhibits its enzymatic activity more efficiently than amastatin, we next compared the effect of bestatin on the enzymatic activity of the wild-type and Q238A laeverin/APQs. As shown in Fig. 3, bestatin inhibited the enzymatic activity of the wild-type enzyme toward Leu-MCA comparably with that toward Arg-MCA. On the other hand, it inhibited the mutant enzyme about 50-fold less efficiently than the wild-type enzyme when measured using Arg-MCA as a substrate. Considering that bestatin is a small peptide having an aromatic amino acid-like residue at its N-terminus, consistent with the notion that wild-type laeverin/APQ binds to aromatic amino acids with higher affinity than the Q238A mutant enzyme.

Table 1. Kinetic Parameters of Wild-Type and Q238A Laeverin/APQs toward Leu-MCA and Arg-MCA

<table>
<thead>
<tr>
<th>Substrate</th>
<th>( K_m ) (( \mu )M)</th>
<th>( k_{cat} ) (s(^{-1}))</th>
<th>( k_{cat}/K_m \times 10^3 ) (s(^{-1})·( \mu )M(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leu-MCA</td>
<td>Wild-type 119±21.9</td>
<td>53±8.06</td>
<td>451±15.7</td>
</tr>
<tr>
<td></td>
<td>Q238A 83.2±6.70</td>
<td>0.23±0.01</td>
<td>2.79±0.12</td>
</tr>
<tr>
<td>Arg-MCA</td>
<td>Wild-type 27.9±7.82</td>
<td>5.56±0.81</td>
<td>205±32.9</td>
</tr>
<tr>
<td></td>
<td>Q238A 11.5±4.33</td>
<td>2.53±0.52</td>
<td>232±49.4</td>
</tr>
</tbody>
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\( a \) Values are mean±S.D. (n=3).

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In this study, we identified Gln-238 as a residue crucial in the substrate specificity of human laeverin/APQ. Replacement of this residue with Ala caused the enzyme preference for basic amino acids and rather restricted substrate specificity. In our previous work, we reported that replacement of Gln-181 of human ERAP1 with Asp increased the enzyme preference for basic amino acids. We speculated at present that both Gln-238 of human laeverin/APQ with neutral amino acid Ala and replacement of Gln-181 with acidic amino acids might cause a local conformational change of the substrate pocket, resulting in rather restricted substrate specificity through electrostatic interaction between the residue and N-terminal basic amino acid of the substrate. However, replacement of Gln-238 of human laeverin/APQ with neutral amino acid Ala caused the enzyme preference for basic amino acids, suggesting that the direct electrostatic interaction between Ala-238 and N-terminal basic amino acid of the substrates may have little role in the basic amino acid preference of the enzyme. Instead, we speculate at present that both Gln-238 of wild-type and Ala-238 of Q238A mutant laeverin/APQs maintain their catalytic pocket structures by interacting with or interfering with another unidentified residues, and thus form unique S1 sites, as discussed elsewhere. It is possible that unique S1 site structure of the mutant enzyme but not that of the wild-type may allow the electrostatic interaction between neighboring acidic amino acid and basic amino acid substrate, causing the basic amino acid preference.

The corresponding site of the M1 family of aminopeptidases is occupied either Gln or Asp (Fig. 1B). When the site is occupied by Asp, this site may interact with either Ca$^{2+}$ (APA) or substrates (L-RAP/ERAP2) through electrostatic interaction and thus contribute to the unique properties of the enzymes. To elucidate the mode of the interaction between S1 site and substrates definitively, it is essential to determine the crystal structure of the enzymes.

In summary, through substitution of Gln-238 with Ala of human laeverin/APQ, we have shown that this residue is important for its substrate specificity; however, as direct electrostatic interaction is not likely to be involved in the interaction between N-terminal amino acid of the substrate and S1 site of the enzyme, it is conceivable that another acidic residue might participate in their interaction.

REFERENCES