Conserved lysine 79 is important for activity of ecto-nucleoside triphosphate diphosphohydrolase 3 (NTPDase3)

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Abstract

Cell membrane-bound ecto-nucleoside triphosphate diphosphohydrolases (NTPDases) are homooligomeric, with native quaternary structure required for maximal enzyme activity. In this study, we mutated lysine 79 in human ecto-nucleoside triphosphate diphosphohydrolase 3 (NTPDase3). The residue corresponding to lysine 79 in NTPDase3 is conserved in all known cell surface membrane NTPDases (NTPDase1, 2, 3, and 8), but not in the soluble, monomeric NTPDases (NTPDase5 and 6), or in the intracellular, two transmembrane NTPDases (NTPDase4 and 7). This conserved lysine is located between apyrase conserved region 1 (ACR1) and an invariant glycosylation site (N81), in a region previously hypothesized to be important for NTPDase3 oligomeric structure. This lysine residue was mutated to several different amino acids, and all mutants displayed substantially decreased nucleotidase activities. A basic amino acid at this position was found to be important for the increase of nucleotidase activity observed after treatment with the lectin, concanavalin A. After solubilization with Triton X-100, mutants showed little or no decrease in activity, unlike the wild-type enzyme, suggesting that the lysine at this position may be important for maintaining proper folding and for stabilizing the quaternary structure. However, mutation at this site did not result in global changes in tertiary or quaternary structure as measured by Cibacron blue binding, chemical cross linking, and native gel electrophoretic analysis, leaving open the possibility of other mechanisms by which mutation of this conserved lysine residue might decrease enzyme activity.

Abbreviations: ACR – apyrase conserved region; BSA – bovine serum albumin; Con A – concanavalin A; DMEM – dulbecco’s modified eagle medium; DMSO – dimethyl sulfoxide; DSS – disuccinimido suberate; DTT – dithiothreitol; MOPS – 3-[N-morpholino] propane sulfonic acid; NTPDases – nucleoside triphosphate diphosphohydrolases; Pi – inorganic phosphate; PVDF – polyvinylidene fluoride; TBS – tris buffered saline

Introduction

The eNTPDases are a family of enzymes that hydrolyze extracellular nucleoside 5’ di- and triphosphates [1]. The nucleotidase activity of these enzymes is dependent on the presence of divalent cations (usually Ca2+ or Mg2+). Four of the six members of the human eNTPDases characterized to date (NTPDase1–4) are integral membrane glycoproteins, with large extracellular domains and two transmembrane domains located near the N- and C-termini [1]. Three of these four integral membrane eNTPDases (NTPDase1–3) are cell-surface-associated, while NTPDase4 is associated with Golgi membranes [2]. Recently, another cell-surface associated member of this family was reported, mouse NTPDase8, which is expressed at the highest level in liver [3].

Glycosylation of asparagine 81 (N81), an invariant putative glycosylation site located near apyrase conserved region 1 in the NTPDases, has been shown to be essential for full enzymatic activity of NTPDase3 [4]. Enzymatic deglycosylation of this site (by the deglycosylation enzyme, peptide N-glycosidase-F) was shown to be responsible for the inactivation of the wild-type enzyme [4]. Removal of the glycosylation site at this position was hypothesized to have its effects by decreasing the stability of the native oligomeric structure [4].

A lysine residue has been hypothesized to be involved in the coordination of a metal ion involved in the catalytic pathway of CD39 (NTPDase1 [5]). However, multiple sequence alignments of NTPDases reveal very few conserved lysine residues, and no conserved lysine residues in the putative phosphate binding domains of the NTPDases, apyrase conserved regions 1 and 4 (ACR1 and ACR4 [6]).
Nevertheless, there are two fairly well conserved lysine residues nearby ACR1, Lys 56 and Lys 79 in NTPDase3. In the present study, we analyzed mutations of Lys 79, a conserved lysine residue located nearby the conserved N81 glycosylation site. Lys 79 is conserved in all known sequences of the oligomeric, cell surface NTPDases 1, 2, 3, and 8, but not in the soluble, monomeric enzymes, NTPDase5 and NTPDase6 (see Table 1). Analyses of the mutants included solubilization (and monomerization) by Triton X-100, native gel electrophoresis, cross-linking by DSS, Cibacron blue binding, and Concanavalin A binding, in addition to nucleotide assays in the presence of Ca^{2+} or Mg^{2+}. The results indicate that lysine 79 is essential for full nucleotide activity of NTPDase3, and we speculate this residue might be important for maintaining some subtle aspect of native folding and/or native oligomeric structure necessary for maximal enzymatic activity.

Materials and methods

Materials

The QuikChange™ site-directed mutagenesis kit was purchased from Stratagene. Oligonucleotides were synthesized by the DNA Core Facility at the University of Cincinnati. Lipofectamine Plus Reagent, Dulbecco’s modified eagle medium (DMEM), calf serum, and antibiotics/antimycotics were obtained from Gibco/Life Technologies. The mammalian expression vector pcDNA3 was obtained from Invitrogen. The chemical cross-linking reagent disuccinimidy1 suberate (DSS) and the SuperSignal chemiluminescence reagents were purchased from Pierce Chemical. Cibacron Blue Gel (Affi-Gel Blue), pre-cast SDS-PAGE 4%–15% gradient mini-gels, and goat anti-rabbit horseradish peroxidase conjugated secondary antibody were obtained from Bio-Rad Laboratories. Ampicillin, nucleotides and other reagents were from Sigma.

Site-directed mutagenesis of NTPDase3

Mutagenesis of NTPDase3 in pcDNA3 vector was performed using the QuikChange™ site-directed mutagenesis kit (Stratagene) as described previously [6–11]. The sense nucleotides used for mutagenesis are as follows: K79A, 5’-CAATGGCCACGACAGAAAGCAGAGATAATACCGGAGTGG-3’; K79E, 5’-CAATGGCCACGAGAGAAAAGCAGAGATAATACCGGAGTGG-3’; K79G, 5’-CAATGGCCACGAGAGAAAAGCAGAGATAATACCGGAGTGG-3’; K79R, 5’-CAATGGCCACGAGAGAAAAGCAGAGATAATACCGGAGTGG-3’; Altered codons are underlined, but the complementary antisense oligonucleotide also necessary for mutagenesis are not shown. DNA sequencing of 600–800 bases surrounding the mutated bases both confirmed the desired mutation and demonstrated that no unintended mutations occurred. The entire coding sequence was not sequenced because our experience in generating more than 100 mutants using this cDNA and methodology have led us to the conclusion that unwanted mutations typically occur within 20–30 bases of the desired mutation. The mutated NTPDase3 cDNA was used to transform competent cells as described by the manufacturer (Stratagene).

Transient transfection

COS-1 cells were grown and transfected with wild-type and mutant NTPDase3 cDNA, as previously described [6–11]. An empty pcDNA3 vector was transfected into
COS cells and used as a background control for nucleotidase assays. Cells were harvested 48 h post-transfection and crude total membrane preparations were obtained as described [6–11].

**Protein assay**

Protein concentrations were determined using the Bio-Rad Coomassie blue dye binding assay, using bovine serum albumin as the standard, with the modifications of Stoscheck [12].

**Nucleotidase assays**

Nucleotidase activities were determined by measuring the concentration of inorganic phosphate (P$_i$) released from the ATP or ADP substrates in the presence of Mg$^{2+}$ or Ca$^{2+}$. Assays were performed at 37 °C as previously described [9–11], modified from Fiske and Subbarow [13]. Nucleotidase activities were corrected for COS-1/pcDNA3 background (provided by the empty pcDNA3 vector transfected into COS-1 cells) and for differences in expression levels relative to those of wild-type, as determined by Western blotting.

**Western blot analysis**

For Western blot, proteins were resolved in a 4%–15% linear gradient SDS-PAGE gradient gel (BioRad 4%–15%) and transferred to a polyvinylidene fluoride (PVDF) membrane. Blots were probed with an anti-C-terminal peptide NTPDase3 antibody, as previously described [6].

**Chemical cross-linking**

COS cell membrane preparations (0.1 mg/ml) were diluted in 20 mM MOPS, 5 mM MgCl$_2$ buffer, pH 7.4, and incubated for 20 min at 22 °C with 200 μM disuccinimido suberate (DSS) freshly dissolved in DMSO. This lysine specific cross-linking agent is able to cross-link NTPDase3 monomers in contact in the native oligomer, and has been used as a probe for oligomeric structure [14]. Non-cross-linked samples were treated with the same volume of DMSO, and the final DMSO concentration of all samples was less than 2% of the total sample volume. The cross-linking reaction was stopped by incubation with 10 mM lysine for 5 min at 22 °C. After reducing sample buffer was added, samples were boiled for 5 min, loaded on a SDS-PAGE gradient gel, and Western blotted, as described above.

**Analysis of Cibacron blue binding**

Cibacron blue binding assays have been described previously [7, 9, 10]. Basically, this triazine dye binds to many nucleotide-handling enzymes, presumably at the nucleotide binding site. Therefore it has been shown to be a useful probe of the native tertiary structure by demonstrating if various mutations have affected the gross tertiary structure of NTPDase3, thereby affecting the ability of the mutated enzyme to bind the triazine dye. Native gel electrophoresis of wild-type and mutant NTPDases

NTPDase3 membrane preparations were solubilized for 10 min at room temperature in digitonin (1% final concentration), a detergent known to preserve the native quaternary structure and the activity of the NTPDases [15, 16]. Solubilized proteins were isolated by centrifugation and electrophoresed on a 6% Laemmli native gel containing 0.1% digitonin as described previously [10].

**Treatment of wild-type and mutant NTPDase3 with concanavalin A**

Concanavalin A (Con A) was prepared at a concentration of 5 mg/ml in 20 mM MOPS buffer containing 100 mM NaCl, 1 mM MnCl$_2$ and 1 mM CaCl$_2$, pH 7.4. Wild-type, K79A, K79E, K79G and K79R NTPDase3 total membrane preparations (2 μg) were incubated with 5 μl of 5.0 mg/ml Con A at 37 °C for 15 min before the addition of substrate. Nucleotidase assays were performed in the presence of 5 mM Ca$^{2+}$ or Mg$^{2+}$, at a final nucleotide concentration of 2.5 mM.

**Time dependence of Mg-ATPase activity**

Kinetics of Mg$^{2+}$-dependent ATPase hydrolysis before and after treatment with the lysine-specific cross linker, DSS, were measured in a Beckman DU-800 spectrophotometer, using a NADH-linked enzyme spectrophotometric assay [17], as described previously for NTPDase2 [18] and NTPDase3 [4]. This assay measures the oxidation of NADH (via the decrease in absorbance at 340 nm), which is coupled enzymatically to the hydrolysis of ATP in the presence of Mg$^{2+}$, allowing continuous measurement of Mg$^{2+}$-ATPase activity. After pre-incubation to bring the cuvettes and samples to 37 °C, the reactions were initiated by addition of a small volume of NTPDase3 enzyme.

**Solubilization and nucleotidase assays**

Cell membranes (0.1 mg/ml) were solubilized in 1% Triton X-100, 5 mM MgCl$_2$ and 20 mM MOPS buffer pH 7.4 at 22 °C for 10 min with occasional mixing, followed by centrifugation at 150,000 g for 30 min at 22 °C. Mg$^{2+}$ and Ca$^{2+}$ ATPase and ADPase activities were measured in presence of 0.1% Triton X-100, after adding nucleotide to a final nucleotide concentration of 0.435 mM, using a malachite green phosphate assay [19] to measure nucleotidase activities, because Triton X-100 causes turbidity in the modified Fiske and Subbarow assay [13].

**Results**

**Expression and characterization of NTPDases**

Mutation of the fairly conserved Lys 56 to Ala did not result in diminished nucleotidase activities (data not shown) and
Table 2. Nucleotidase activities of wild-type (wt) and mutant NTPDase3 enzymes.

| NTPDase3 enzyme | % wt level of expression | Normalized Mg$^{2+}$-ATPase activity (% wt) | Normalized Ca$^{2+}$-ATPase activity (% wt) | Normalized Mg$^{2+}$-ADPase activity (% wt) | Normalized Ca$^{2+}$-ADPase activity (% wt) |
|-----------------|--------------------------|--------------------------------------------|--------------------------------------------|--------------------------------------------|--------------------------------------------|
| Wild-type       | 100                      | 144 ± 10 (100)                             | 413 ± 35 (100)                            | 57 ± 5 (100)                               | 83 ± 15 (100)                             |
| K79A            | 68 ± 5                   | 22 ± 2 (15*)                               | 92 ± 12 (22)                              | 16 ± 4 (28*)                               | 36 ± 4 (43)                               |
| K79E            | 50 ± 7                   | 25 ± 4 (17*)                               | 97 ± 18 (23)                              | 15 ± 2 (26*)                               | 36 ± 4 (43)                               |
| K79G            | 60 ± 4                   | 18 ± 4 (12*)                               | 88 ± 22 (21)                              | 7 ± 2 (12*)                                | 30 ± 5 (36)                               |
| K79R            | 74 ± 8                   | 36 ± 5 (25)                                | 113 ± 16 (27)                             | 12 ± 3 (21*)                               | 34 ± 6 (40)                               |

Values given represent the means ± standard deviations of three separate transfections (which were matched to their own wild-type and empty pcDNA3 vector controls). Activities were measured in presence of 5 mM MgCl$_2$ or CaCl$_2$ at a final concentration of 2.5 mM ATP or ADP. Values were normalized for different NTPDase3 protein expression levels by dividing the nucleotidase activity (expressed as μmol Pi/mg protein/h) by the expression level relative to the wild-type enzyme. Note the differences in the degree of loss of Mg$^{2+}$-stimulated activity compared to the loss of Ca$^{2+}$-stimulated activity for most of the K79 mutants, especially regarding the ADPase activities. Values for the normalized % wild-type Mg-nucleotidase activities (in parentheses) that are statistically different ($P \leq 0.05$) from the corresponding Ca-nucleotidase activities (also in parentheses) are indicated by an asterisk (*).

Therefore, was not investigated further. Wild-type and K79 mutant NTPDase3 proteins were transiently expressed in COS cells and expression levels were calculated by quantification of Western blots as previously described [10]. The values presented in Table 2 are the means of three separate transfections, and the specific activities for ATP and ADP of the NTPDase3 mutants were corrected for variations in expression levels relative to those of the wild-type (see Table 2). ATPase and ADPase nucleotidase activities of all mutants were determined in presence of either Mg$^{2+}$ or Ca$^{2+}$. For wild-type NTPDase3, activities for Mg$^{2+}$ and Ca$^{2+}$-ATPase were 144 and 413 μmol/mg/h, respectively, while activities for Mg$^{2+}$ and Ca$^{2+}$-ADPase were 57 and 83 μmol/mg/h, respectively (Table 2). Interestingly, K79 mutants exhibited different amounts of residual Ca$^{2+}$ versus Mg$^{2+}$-mediated nucleotidase activities. For example, in Mg$^{2+}$ buffer, K79 NTPDase3 mutants show a 4- to 8-fold decrease in ATPase activity and a 3.5- to 8-fold decrease in ADPase activity, while in the presence of Ca$^{2+}$, K79 mutants show smaller (3.5–5) fold decreases in ATPase activity and approximately threefold decreases in ADPase activity (Table 2). Thus, the K79 mutants showed a larger proportional decrease in Mg$^{2+}$-stimulated activity compared to Ca$^{2+}$-stimulated activity. Those differences that are statistically significant ($P \leq 0.05$) when comparing the % wild-type activity remaining for Ca$^{2+}$ versus Mg$^{2+}$-activities of each mutant are indicated in Table 2 with an asterisk (*).

$Mg^{2+}$-ATPase kinetics of wild-type and K79A mutant NTPDase3 after cross-linking

To investigate the effect of the lysine specific cross-linker DSS on Mg$^{2+}$-ATPase activity of wild-type NTPDase3 and the K79A lysine mutant, the activity was monitored as a function of time utilizing the linked enzyme assay. Interestingly, DSS treatment increased Mg$^{2+}$-ATPase activity of the wild-type enzyme (consistent with earlier results [4]), but decreased Mg$^{2+}$-ATPase activity of the K79A mutant (see Figure 1). Previously, such increases in wild-type NTPDase enzyme activity upon chemical cross-linking have been attributed to stabilization of the native quaternary structure [4, 15].

Analysis of the global quaternary structure of the mutants by native gel electrophoresis and DSS cross-linking

The native gel electrophoretic mobilities of wild-type NTPDase3 and K79 mutants are indistinguishable, suggesting that the mutants have the same gross oligomeric structure as wild-type NTPDase3 (Figure 2, top panel). In addition, a lysine-specific cross-linking agent, disuccinimido substrate (DSS), was also used to investigate quaternary structure. After reducing SDS-PAGE and Western blotting, both wild-type and mutant NTPDases exist as non-cross-linked monomers in the absence of DSS (data not shown).

Figure 1. Time dependence of Mg$^{2+}$-ATPase activity of wild-type and K79A NTPDase3 after DSS cross-linking. Wild-type and mutant COS cell membranes (0.1 mg/ml) were treated with DSS (200 μM) for 10 min at room temperature, the reaction was stopped with lysine, and Mg$^{2+}$-stimulated activity was measured by the NADH-linked enzyme assay [17]. Due to the lower amount of activity of the K79A mutant, 3 μg was used for each wild-type sample, while 9 μg was used for each K79A NTPDase3 sample, to obtain curves of approximately the same shape, thus allowing easier visual comparison. For clarity of presentation, the data for wild-type NTPDase3 (both control and DSS cross-linked) have been shifted down by subtracting 0.2 absorbance units from each data point.
Following treatment with 200 μM DSS, wild-type and all NTPDase3 mutants made in this study were similarly cross-linked into dimers, suggesting that the gross quaternary structures of the mutants are similar to those of the wild-type (see Figure 2, middle panel).

Analysis of the global tertiary structure of the mutants by Cibacron blue binding

To explore the possibility of global misfolding induced by point mutations leading to the inability to bind the nucleotide analogue triazine dye, Cibacron blue, Cibacron blue binding assays were performed as previously described [10]. Grossly misfolded mutants are unable to bind Cibacron blue affinity matrix, as is denatured NTPDase3 (exemplified by the boiled, wild-type enzyme [10]). All the mutants described in this study bound to the Cibacron blue matrix like the wild-type enzyme (see Figure 2, bottom panel), suggesting no global changes in the tertiary conformations of the mutants affecting the ability to bind the nucleotide analogue, Cibacron blue.

Effect of concanavalin A on NTPDase3 wild-type and lysine mutants

Concanavalin A is a tetravalent protein that binds to glycans and can stabilize the oligomeric structure of glycoproteins by induction of the formation of protein oligomeric lattices [20]. Concanavalin A was previously shown to increase the nucleotidase activity of several cell surface, oligomeric NTPDases [4, 18, 21–23]. Unlike wild-type NTPDase3 and the K79R mutant, which show approximately a 1.9-fold increase in activity after incubation with Con A, the K79A, K79E, and K79G mutant ATPase activities were not substantially stimulated by Con A (Figure 3). Thus, the nature of the amino acid at this position plays an important role. A basic, positively charged residue at this position is necessary for the stimulatory effect of Con A, while acidic (K79E) or uncharged (K79G and K79A) substitutions at this position greatly diminish this effect. In contrast, the K79R mutant NTPDase3 Mg2+-ATPase activity was decreased by DSS cross-linking (data not shown), similar to the K79A mutant, but dissimilar to the wild-type enzyme shown in Figure 1. Thus, the K79R mutant that responded like wild-type NTPDase3 to Con A treatment (mediated via glycan binding) did not respond like wild-type enzyme to chemical cross-linking by DSS (mediated via reaction at lysine residues).

Do mutations at position 79 affect the NTPDase3 oligomer?

To investigate whether the effects of mutation of Lys 79 might be dependent on the oligomeric structure, wild-type and K79 mutants were solubilized with Triton X-100. Ca2+ and Mg2+ ATPase activities as well as ADPase activities were determined (Figure 4). Solubilization decreases both Mg2+ and Ca2+-ATPase for wild-type NTPDase3 (Figure 4A). For the wild-type enzyme, Triton X-100 treatment decreased the Mg2+-ATPase activity 75%, while only decreasing the Ca2+-ATPase activity by 37%. Mg2+-ADPase activity of wild-type was decreased by 67% while Ca2+-ADPase activity was unchanged or increased slightly (Figure 4B). In contrast, for the lysine mutants, the activity...
remaining after Triton X-100 solubilization is 80% for Mg\(^{2+}\)-ATPase, while there is a 30% increase in the Ca\(^{2+}\)-ATPase activity for all the mutants, except the K79G mutant, whose Ca\(^{2+}\)-ATPase activity remains unchanged (Figure 4A). Both the Mg\(^{2+}\)-ADPase and Ca\(^{2+}\)-ADPase activities of the K79 mutants were increased by 25%–30%, except for the K79G mutant, which remained unchanged (Figure 4B).

Discussion

In this study we used site-directed mutagenesis to mutate the conserved lysine residue at position 79 to several different amino acids to determine the role of lysine at this position in NTPDase3. This lysine residue is conserved in all cell-membrane associated NTPDases [1–3, 8] sequenced to date, but is not conserved in the soluble NTPDase5 and NTPDase6 enzymes, or in the intracellular membrane associated NTPDase4 and NTPDase7 enzymes (see Table 1). This led us to consider the possibility that K79 is important for native oligomeric protein interactions of the cell surface membrane-bound NTPDases.

Continuous measurement of Mg\(^{2+}\)-ATPase activity indicates that wild-type NTPDase3 activity increases after cross-linking, in contrast to the Mg\(^{2+}\)-ATPase activity of the K79A mutant, which is inhibited after cross-linking (Figure 1). The increase in wild-type activity after cross-linking has been proposed to be the result of inter-molecular cross-linking and the resultant stabilization of the native oligomeric structure for both NTPDase3 [4], and other [15, 18] membrane-bound NTPDases. Therefore, this result is consistent with the possibility that K79 is important for, or directly involved in, the formation of an inter-molecular cross-link that stabilizes the native quaternary structure, thereby increasing enzyme activity. The observed decrease in K79A activity after cross-linking could be due to (non-cross-linking) modification of essential lysine residues, or to intra-molecular cross-linking, or both. These inhibitory effects of DSS treatment would presumably be masked in the wild-type enzyme by the larger stimulatory effect of inter-molecular cross-linking, possibly involving K79.

Concanavalin A, a tetravalent lectin known to stabilize the oligomeric structure of glycoproteins by induction of high-order protein oligomer lattice formation [20], increases the nucleotidase activity of several eNTPDases [21–27], presumably by modulating the oligomeric structure/stability of the enzyme. Unlike the wild-type enzyme, the K79A, K79E, and K79G NTPDase3 mutants are not stimulated substantially by Con A. However, when lysine is replaced by another basic amino acid (arginine, i.e., the K79R mutant), the residual activity behaves like wild-type with regard to its stimulation by interaction with Con A (see Figure 3), suggesting that a positively charged amino acid at this position is needed for the increase in activity induced by the lectin. However, this K79R mutant does not behave like the wild-type enzyme in that it has significantly lower nucleotidase activity than wild-type enzyme (Table 2), and its activity is not increased by DSS cross-linking (data not shown).

A Cibacron Blue binding assay, which is a measure of gross tertiary structure changes, suggests that all these mutants are not globally misfolded (Figure 2). Chemical cross-linking and native gel electrophoresis results suggest that these mutants also do not undergo global changes in quaternary structure (Figure 2). Thus, the results presented in Figure 2 suggest that the decreases in nucleotidase activity observed for the K79 mutants are not due to global protein misfolding or to gross changes in oligomeric structure.

Triton X-100 solubilization is known to disrupt the oligomeric structure of the cell-surface NTPDases, resulting in monomeric enzymes [11, 16]. After Triton X-100 solubilization, wild-type NTPDase3 loses most of its Mg-ATPase activity and there is a substantial decrease in Ca\(^{2+}\)-ATPase activity (see Figure 4A). However, solubilization has little effect on the Mg\(^{2+}\)-ATPase activity of the K79
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mutants, and Ca\(^{2+}\)-ATPase activities of these mutants are slightly increased after solubilization (Figure 4A). One interpretation of these results is that the lysine residue at this position may, in some subtle way, not detectable by DSS cross-linking/Western blot or native gel electrophoresis (Figure 2), stabilizes the quaternary structure of wild-type NTPDase3. The conserved glycosylation site, which is located two residues removed from K79 (at N81), was previously shown to be important for enzyme activity, apparently mediated by its affect on the native quaternary structure [4]. Thus, it is possible that mutation of K79 might affect the local protein conformation and therefore the nearby N81 glycosylation site. Consistent with this hypothesis is the inability of Concanavalin A to increase the enzymatic activity of most of the K79 mutants (Figure 3), as well as the lack of inhibition of the K79 mutants by monomerization with Triton X-100 (Figure 4). Therefore, these results lead us to hypothesize that K79, and this region of the protein in general, may be important in some subtle way for the stability or fine-tuning of the native quaternary structure necessary for maximal enzyme activity. It follows that since the quaternary structures of the K79 mutants are already non-optimal, further disruption of the quaternary structure by Triton X-100 monomerization would have a reduced effect on the residual Mg\(^{2+}\)-ATPase activity of these mutants, which is what is observed (see Figure 4). Alternatively, the K79 residue could be directly important for enzyme catalysis. However, this seems not as likely, since this residue is only two amino acids from a known, conserved glycosylation site (N81 in NTPDase3), which should be solvent-accessible, and not in a relatively shielded active site catalytic cleft typical of enzymes. In addition, K79 is not likely to be directly involved in a generic NTPDase active site, since it is not conserved in either the soluble or the intracellular NTPDases (Table 1). Also, mutation of an active site residue would likely diminish Mg\(^{2+}\)- and Ca\(^{2+}\)-supported nucleotidase activities to roughly the same extent, which is not the case for the K79 mutants (Table 2).

Although the membrane-bound activities of the wild-type NTPDase3 and many of the mutant NTPDase3 constructs (made in this and previous studies) are higher in Ca\(^{2+}\) than in Mg\(^{2+}\), the reason for this is unclear. Based on the differences observed in the Ca\(^{2+}\) versus Mg\(^{2+}\) nucleotidase activities (Figure 4), we speculate that Ca\(^{2+}\) stabilizes the tertiary and quaternary structures of the NTPDase3 relative to Mg\(^{2+}\), giving rise to a more stable native oligomeric structure and therefore a higher nucleotidase activity. This effect is seen with the enzyme expressed in COS cell membranes, and is even more apparent after solubilization of wild-type NTPDase3 with Triton X-100 (see Figure 4), a detergent that disrupts the oligomeric structure of NTPDases, including NTPDase1/CD39 [16] and NTPDase3 [11]. Thus, the % decrease of the ATPase activity after solubilization/monomerization of NTPDase3 is greater when assayed using Mg\(^{2+}\)-ATP as substrate as opposed to Ca\(^{2+}\)-ATP as substrate. After Triton X-100 solubilization, the remaining Mg\(^{2+}\)-ATPase activities of the K79 NTPDase3 mutants are virtually the same as the wild-type enzyme, unlike the results obtained in Ca\(^{2+}\) buffer (see Figure 4). This suggests that Ca\(^{2+}\) may have a stabilizing effect on the monomeric tertiary structure, or that Ca\(^{2+}\)-ATP is a better substrate than Mg\(^{2+}\)-ATP for the solubilized, monomeric NTPDase3.

In summary, conserved lysine 79, located near ACR1, is essential for maximal NTPDase3 nucleotidase activity. Due to its location very near a known glycosylation site (N81), it seems unlikely that it is directly involved in the active site. Con A and DSS effects on K79 NTPDase3 mutant nucleotidase activities, as well as the effects of Triton X-100 on nucleotidase activities, are consistent with K79 being important for native quaternary structure. However, Cibacron blue binding, native gel electrophoresis, and DSS cross-linking of NTPDase3 protein indicate no gross changes in tertiary or quaternary structure for the K79 mutants. This suggests that the changes in tertiary or quaternary structure in these mutants must be subtle, and not able to be detected by these techniques. Future application of more sensitive and/or more discriminating techniques to adequately measure small changes in tertiary and quaternary structures may definitively answer the question of the mechanism of loss of activity induced by mutation of lysine 79.

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