Cloning and characterization of the ecto-nucleotidase NTPDase3 from rat brain: Predicted secondary structure and relation to other members of the E-NTPDase family and actin

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Abstract

The protein family of ecto-nucleoside triphosphate diphosphohydrolases (E-NTPDase family) contains multiple members that hydrolyze nucleoside 5'-triphosphates and nucleoside 5'-diphosphates with varying preference for the individual type of nucleotide. We report the cloning and functional expression of rat NTPDase3. The rat brain-derived cDNA has an open reading frame of 1590 bp encoding 529 amino acid residues, a calculated molecular mass of 59.1 kDa and predicted N- and C-terminal hydrophobic sequences. It shares 94.3% and 81.7% amino acid identity with the mouse and human NTPDase3, respectively, and is more closely related to cell surface-located than to the intracellularly located members of the enzyme family. The NTPDase3 gene is allocated to chromosome 8q32 and organized into 11 exons. Rat NTPDase3 expressed in CHO cells hydrolyzed both nucleoside triphosphates and nucleoside diphosphates with hydrolysis ratios of ATP:ADP of 5:1 and UTP:UDP of 8:1. After addition of ATP, ADP is formed as an intermediate product that is further hydrolyzed to AMP. The enzyme is preferentially activated by Ca2+ over Mg2+ and reveals an alkaline pH optimum. Immunocytochemistry confirmed expression of heterologously expressed NTPDase3 to the surface of CHO cells. PC12 cells express endogenous surface-located NTPDase3. An immunoblot analysis detects NTPDase3 in all rat brain regions investigated. An alignment of the secondary structure domains of actin conserved within the actin/HSP70/sugar kinase superfamily to those of all members of the NTPDase family reveals apparent similarity. It infers that NTPDases share the two-domain structure with members of this enzyme superfamily.

Abbreviations: ACR – apyrase conserved region; CHO – Chinese hamster ovary; ConA – concanavalin A; Cy3 – indocarbocyanin-3; DAPI – 2-(4-aminophenyl)-6-indolecarbamidine dihydrochloride; DMEM – Dulbecco’s modified Eagle’s medium; EDTA – ethylenediaminetetraacetic acid; E-NTPDase – ecto-nucleoside triphosphate diphosphohydrolase; EST – expressed sequence tag; NGF – nerve growth factor; PC12 – pheochromocytoma 12; PCR – polymerase chain reaction; ORF – open reading frame

Introduction

The protein family of ecto-nucleoside triphosphate diphosphohydrolases (E-NTPDase family) contains multiple members that differ regarding tissue distribution, cellular location and substrate specificity. They hydrolyze nucleoside 5'-triphosphates and nucleoside 5'-diphosphates with varying preference for the individual type of nucleotide. Depending on subtype their catalytic site may be in ecto-position facing the extracellular medium or it may be located in the lumen of intracellular organelles. Whereas the surface-located members of the protein family are thought to be involved mainly in the control of ligand availability for P2 receptors, the functional role of the intracellular members is less defined [1, 2].

The gene family has members within vertebrates, invertebrates, plants, yeast and protozoans but has not been identified in bacteria (references in [1, 3–5]). Hallmarks of all E-NTPDases are five highly conserved sequence domains (‘apyrase conserved regions,’ [3, 4, 6]) that presumably are involved in the catalytic cycle. Interestingly, E-NTPDases share two common sequence motifs with members of the actin/HSP70/sugar kinase superfamily, the actin-HSP 70-hexokinase β- and γ-phosphate binding motif [(I/L/V)X(I/L/V/C)DXG(T/S/G)(T/S/G)XX(R/K/C)] [3, 7–9], with the DXG sequence strictly conserved.
In contrast to the members of the actin/HSP70/sugar kinase superfamily, mammalian E-NTPases are membrane-anchored proteins with one or two transmembrane domains. NTPDase5 and NTPDase6 contain a single predicted N-terminal hydrophobic domain. They are located to the endoplasmic reticulum or Golgi apparatus respectively but they can also be released in soluble form from transfected cells [10–12]. They preferentially hydrolyze nucleoside diphosphates. The two forms of NTPDase4, that differ by alternative splicing, have predicted N- and C-terminal transmembrane domains and were allocated to the Golgi apparatus (UDPase) [13] and to lysosomal/autophagic vacuoles (LALP70) [14, 15], respectively. The Golgi enzyme hydrolyzes a number of nucleoside 5'-di and triphosphates but not ATP and ADP. NTPDase7 [16] is localized to intracellular organelles and hydrolyses a variety of nucleoside triphosphates with the exception of ATP. In mammals, four different surface-located subtypes of E-NTPases have been cloned and characterized. They share a membrane topography with N- and C-terminal transmembrane domains: NTPDase1 [17–20], NTPDase2 [8, 21], NTPDase3 [9, 22, 23], and most recently NTPDase8 [24]. They all hydrolyze nucleoside tri- and diphosphates but differ significantly in catalytic properties [25].

Surface-located mammalian E-NTPases have been cloned and characterized mainly from rat, human and mouse tissues. Since these enzymes vary regarding substrate preference and product pattern formation it is necessary to compare the catalytic properties of individual enzymes within the same species. We have previously cloned and characterized rat NTPDase1 and rat NTPDase2 [8, 20]. Here we report the cloning and characterization of rat NTPDase3. We analyze its functional properties and tissue distribution in brain and compare some of the key properties of primary and secondary structure to those of other members of the gene family.

**Materials and methods**

**Materials**

Rat brain cDNA library SuperScript™, reverse transcriptase SuperScript™II, Trizol™ reagent, oligo(dt)-cellulose, fetal calf serum, horse serum, DMEM (Dulbecco’s modified Eagle’s medium), penicillin and streptomycin, and trypsin/EDTA solution was obtained from Invitrogen (Karlsruhe, Germany). The cell culture medium Ham’s F12 (www.ncbi.nlm.nih.gov/BLAST/) was used for obtaining the complete full length sequence of the cDNA clone 3.1.1.1. The Oligo 2.0 sequence analysis program (Oxford Molecular Ltd., Oxford, UK) was used for assembling sequence fragments, translating DNA into amino acid sequences, generating hydrophobicity blots and amino acid alignment (CLUSTAL W algorithm). To align the amino acid sequences for the denguevirus, ClustalX 1.81 was employed. For prediction of transmembrane domains, the software TMHMM 2.0 (www.ncbi.nlm.nih.gov/BLAST/) was used. Protein motif search was performed with the 288-bp cDNA fragment labeled with [α-32P]dCTP by PCR. Positive signal areas were amplified and rescreened for single positive colonies.

**cDNA library screening**

Total RNA from rat brain was isolated with Trizol reagent. For isolation of polyadenylated RNA from total RNA, oligo(dt)-cellulose was used according to the manufacturer’s instructions. A cDNA library was synthesized with SuperScript™II from 0.5 μg of mRNA with an oligo(dT)18 primer in accordance with the manufacturer’s instructions. Two mouse sequences from expressed sequence tag (EST) databases (GenBank accession numbers bf302156 and w46136) were used for primer design. As a probe, a 288-bp PCR fragment was amplified using forward primer 5'-CCGTCCCTGCTCCCAAGATT-3' and reverse primer 5'-CAGGACACGAGGCGGATTACG-3' and rat brain cDNA library as a template. For library screening, electrocompetent Escherichia coli DH5α were transformed with a rat brain pCMV-SPORT 2 cDNA library amplified and plated on Luria–Bertani/ampicillin agar plates. The resulting transformants were screened by colony hybridization with the 288-bp cDNA fragment labeled with [α-32P]dCTP by PCR. Positive signal areas were amplified and rescreened for single positive colonies.

**cDNA sequencing and computational sequence analysis**

DNA sequencing was performed by Scientific Research and Development GmbH (Oberursel, Germany). Primer walking in both directions was employed for obtaining the complete full length sequence of the cDNA clone 3.1.1.1. The Oligo 2.0 sequence analysis program (Oxford Molecular Ltd., Oxford, UK) was used for assembling sequence fragments, translating DNA into amino acid sequences, generating hydrophobicity blots and amino acid alignment (CLUSTAL W algorithm). To align the amino acid sequences for the denguevirus, ClustalX 1.81 was employed. For prediction of transmembrane domains, the software TMHMM 2.0 (www.ncbi.nlm.nih.gov/BLAST/) was used. For signal peptide and sorting analysis, SignalP 3.0 (www.cbs.dtu.dk/services/SingalP/) and PSORT II (http://psort.nibb.ac.jp/form2.html) were used. The DNA and deduced amino acid sequences were analyzed for similarity to known sequences with the NCBI Blast Network service (www.ncbi.nlm.nih.gov/BLAST/). Protein motif search was
performed using the prosite database (www.expasy.org/prosite/). Secondary structure prediction of the amino acid sequences was performed using the SSPro tool (www.igb.uci.edu/tools/scratch/). The genomic library was screened using BLAST and the splice analysis of the genomic sequence was performed using the splice site analysis tool www.fruitfly.org/seq-tools/splice.html.

Expression of recombinant proteins
For recombinant expression, the EcoRI/NotI cDNA fragment of clone 3.1.1.1 was cloned into the EcoRI/NotI sites of pcDNA3. Chinese hamster ovary (CHO) cells were cultured as previously described [8]. Cells were transfected by electroporation with the rat NTPDase3 pcDNA3 plasmid or with plasmids expressing rat NTPDase1 [20] or NTPDase2 [8] in electroporation buffer (in mM: 137 NaCl, 5 KCl, 0.7 Na2HPO4, 6 dextrose, 20 Hepes, pH 7.0) using a BTX Electrocell manipulator 600. In control experiments cells were transfected with empty vector alone. Twenty-four hours after electroporation the culture medium of transfected CHO cells was exchanged to remove dead cells and debris.

Preparation of membrane fractions
Forty-eight hours after electroporation, the culture medium was removed, cells were washed twice with isotonic buffer (in mM: 140 NaCl, 5 KCl, 0.5 EDTA, 2 MOPS, pH 7.4) and scraped from the plates in ice cold homogenization buffer (in mM: 250 sucrose, 2 EDTA, 2 iodoacetamide, 30 MOPS, pH 7.4) containing a mixture of protease inhibitors (in µg/ml: 2 chymostatin, 2 aprotinin, 1 peptastin, 150 benzamidine, 2 antipain, 2 leupeptin). After centrifugation at 300 gav, cells were resuspended in homogenization buffer, homogenized, and centrifuged for 10 min at 300 gav at 4 °C. The resulting supernatant was sonicated and subsequently centrifuged at 100,000 gav, for 60 min at 4 °C, pellets were resuspended in storage buffer (in mM: 2 iodoacetamide, 25 Hepes, pH 7.4) containing the protease inhibitor mixture and 50% (v/v) glycerol, and stored at −20 °C until further processing. For preparation of membrane fractions from various brain tissues, Wistar rats obtained from Charles River Wiga (Sulzfeld, Germany) were used. Animals were anaesthetized with CO2, decapitated, the brain was dissected, homogenized in five volumes of homogenization buffer containing the protease inhibitor mixture and further processed as described above.

Measurement of nucleotidase activities
Nucleotidase activity was determined by measuring the formation of Pi liberated from nucleotides [26]. Membrane fractions were incubated in phosphate-free solution containing 500 µM CaCl2 and 25 mM Hepes (pH 7.4), and 500 µM nucleoside tri- or diphosphates, respectively, at 37 °C. Samples were heat-inactivated for 2 min at 95 °C prior to determination of inorganic phosphate. To investigate the dependence of enzyme activity on metal ions, 50–400 µM CaCl2 or MgCl2 was added, or CaCl2 and MgCl2 were replaced by 1 mM EDTA. pH-dependency was determined using a combined buffer (25 mM Hepes and 50 mM glycine) ranging from pH 3 to 10, containing 500 µM ATP and 500 µM CaCl2 or 500 µM MgCl2. Catalytic activity of membrane fractions derived from cells transfected with the empty plasmid was subtracted from that obtained with cDNA-transfected cells. It was verified for each experimental condition applied that hydrolysis rates were constant (10–30 min). At the end of the reaction with nucleoside triphosphate it was ensured that less than 10% of the initial substrate was hydrolysed.

For determination of product formation following ATP hydrolysis (250 µM ATP, 250 µM CaCl2), aliquots were collected at various time points and subjected to HPLC analysis. Following heat inactivation, samples were centrifuged at 20,000 gav for 15 min (4 °C). ATP, ADP, AMP, and adenosine were separated by a Sepsil C18 reverse phase column (Jasco, Groß-Umstadt, Germany) and eluted with the mobile phase, consisting of 50 mM potassium-phosphate buffer (pH 6.5), 6% methanol and 5 mM tetrabutylammonium hydrogen sulfate [20]. The absorbance at 260 nm was continuously monitored and nucleotide concentration was determined from the area under each absorbance peak.

Immunoblotting
The anti-rat NTPDase3 antibody (N3-3i4) was raised in rabbits by direct injection of the encoding NTPDase3-pcDNA3 plasmid [27]. Previous to Western blot analysis of membrane fractions from transfected CHO cells or mouse brain tissues, NTPDase3 was enriched using ConA-Sepharose. The 100,000 gav, pellet was solubilized in ConA-buffer (in mM: 150 NaCl, 2 MgCl2, 2 CaCl2, 2 MnCl2, 20 Tris/HCl, pH 7.4) containing 0.1% Triton X-100. After an overnight incubation with ConA-Sepharose at 4 °C, the ConA-Sepharose was washed several times with ConA-buffer containing 0.1% Triton X-100, and protein was eluted with the same buffer containing 200 mM methyl-D-mannopyranoside. For immunoblotting the ConA eluate was precipitated with 10% trichloroacetic acid and the pellet was resuspended in sample buffer without reducing agents. Polyacrylamide gel electrophoresis was carried out on minigels (10% acrylamide). Immunoblotting using the polyclonal NTPDase3 antibody (dilution 1:4000) was performed using the Amersham enhanced chemiluminescence system according to the manufacturer’s instructions.

Immunofluorescence
Transfected CHO cells (15,000 cells per well) or non-transfected PC12 cells (15,000 cells per well) were seeded onto poly-D-lysine-coated (10 µg/ml) glass cover slips (10 mm diameter) and cultured for 2 days (CHO cells) or up to 14 days (PC12 cells) [28]. An aliquot of 50 ng/ml of NGF was added to PC12 cells every 3–4 days. For surface
Figure 1. DNA sequence and predicted protein sequence of rat NTPDase3. The five 'apprase conserved regions' (ACR) are indicated by boxes and numbered. Cysteine residues and potential N-glycosylation sites are indicated by arrow heads and filled circles, respectively. Predicted N- and C-terminal hydrophobic sequences are shaded. Amino acid residues conserved from NTPDase1 and NTPDase8 are in bold. In ACR 1 and ACR 4 these include the DXG-containing phosphate-binding motif and a conserved glycine in ACR 5. (GenBank accession number AJ437217.)
labeling, the anti-rat NTPDase3 antibody (1:1000) was applied to viable cells for 15 min at 37 °C followed by repeated washing with phosphate-buffered saline (PBS, in mM: 137 NaCl, 3 KCl, 15 Na+/K+ phosphate buffer, pH 7.4) at room temperature, methanol fixation at −20 °C and application of Cy3-conjugated anti-rabbit IgG antibody together with 2-(4-aminomophenyl)-6-indolecarbamidine dihydrochloride (DAPI, 1 µg/ml). After immunolabeling, cells were mounted and investigated with an epifluorescence microscope equipped with an MCID 4 imaging analysis system (Imaging Research, St. Catherines, Canada).

Results

Cloning and sequencing of rat NTPDase3

Two mouse EST sequences (GenBank accession numbers bf302156 and w46136) homologous to human NTPDase3 (GenBank accession number AF034840) were used for primer design. With these primers a 288-bp fragment was amplified from rat brain cDNA by RT-PCR. Using colony hybridization, six positive cDNA clones were isolated from a rat brain pCMV-SPORT 2 cDNA library using the 288-bp radiolabeled PCR fragment as a probe. One of the clones (3.1.1.1) was completely sequenced in both directions by primer walking. The cDNA of clone 3.1.1.1 had a length of 3087 bp and revealed a deduced open reading frame (ORF) of 1590 bp from nt 214 to 1803. The ORF encodes 529 amino acid residues with a calculated protein molecular mass of 59.1 kDa, and an isoelectric point of 5.75. A polyadenylation signal (AAUAAA) was located 24 nucleotides upstream of the poly(A) tail. The deduced amino acid sequence (Figure 1) contains seven potential N-glycosylation sites located to the extracellular domain. It reveals 13 cysteine residues of which 10 are located to the extracellular domain, five potential protein kinase C phosphorylation sites and six potential casein kinase II phosphorylation sites. The hydrophobicity analysis predicts two strong hydrophobic stretches in the polypeptide chain, close to the N- and C-terminal ends (Figure 1). These regions represent predicted transmembrane domains with the N- and C-terminus at the cytosolic side, separated by a large extracellular loop. This corresponds to the membrane topography of the known plasma membrane-located NTPDases [1].

A homology search using the rat genomic sequence database localized the NTPDase3 gene (Entpd3) to chromosome 8q32 (Genebank Acc.-No. NW_047803 RN8_2327: 3551369-3651370). Alignment of the rat NTPDase3 cDNA against the rat genomic sequence revealed that Entpd3 covers 31012 bp and is organized into 11 exons and 10 introns of which exons 2 to 11 encode the open reading frame (Figure 2). Exon 1 contains most of the 5'-untranslated region of the NTPDase3 cDNA. The ACRs 1–4 are contained in exons 4, 5, 6 and 7, respectively, whereas ACR5 is contained in exon 11. The N-terminal transmembrane domain is situated in exon 3 whereas the C-terminal transmembrane domain and the stop codon are localized in exon 11.

![Figure 1](image1.png)

**Figure 1.** The amino acid sequence of rat NTPDase3.

![Figure 2](image2.png)

**Figure 2.** Chromosomal localization of rat NTPDase3 and intron-exon structure. The upper part of the figure depicts the position of the NTPDase3 gene in chromosome 8q32. The lower part enlarges the genomic sequence of NTPDase3. Upper line: Exons are numbered and indicated by black boxes in relation to the length of the encoded sequences (gDNA fragment = 31 kb). The lower line indicates the length of individual exons. The ORF is shaded. The position of ACRs 1 to 5 is indicated by white boxes, that of the two putative transmembrane domains by black boxes.

![Figure 3](image3.png)

**Figure 3.** Hypothetical phylogenetic tree derived for 22 selected members of the E-NTPDase family (NTPDase1 to NTPDase8) from rat (r), human (h) and mouse (m). Amino acid sequences were aligned using ClustalX 1.81 and the dendrogram was prepared using TreeView 1.6.6. The length of the lines indicates the differences between amino acid sequences. The graph depicts a clear separation between intracellular (top) and surface-located NTPDases (bottom). Protein sequences are derived from mRNA sequences. The GenBank accession numbers of the sequences are as follows: (NTPDase1, Homo sapiens (S73183), Mus musculus (AF037366), Rattus norvegicus (U81295); NTPDase2, Homo sapiens (U91510), Mus musculus (AF042811), Rattus norvegicus (Y11835); NTPDase3, Homo sapiens (AF034840), Mus musculus (AY376710), Rattus norvegicus (AJ437217); NTPDase4, Homo sapiens (AF016032), Mus musculus (AK004761); NTPDase5, Homo sapiens (AF039918), Mus musculus (AF006482), Rattus norvegicus (BC62044); NTPDase6, Homo sapiens (AF039916), Mus musculus (NM_127117), Rattus norvegicus (AJ277748); NTPDase7, Homo sapiens (AK055540), Mus musculus (AF288221); NTPDase8, Homo sapiens (AY430414), Mus musculus (AY364442), Rattus norvegicus (AY536920).
Relation to other members of the E-NTPDase family

The deduced amino acid sequence of the cloned rat NTPDase3 shares 94.3% and 81.7% amino acid identity, respectively, with the mouse [23] and human [22] orthologs. Compared to other rat members of the E-NTPDase family, NTPDase3 is most closely related to cell surface-located NTPDase2, NTPDase8, and NTPDase1 (36.9%, 36.3%, 35.1% identity, respectively). It is more distantly related to the intracellularly located rat NTPDase5 and rat NTPDase6 (17.8% and 15.2% identity, respectively). The graphic depiction of a multiple sequence alignment of 22 mammalian members of the E-NTPDase family illustrates the subdivision of the family into intracellular and surface-located enzymes (Figure 3). Whereas the origins of the branches for the four surface-located members (NTPDase1, NTPDase2, NTPDase3, NTPDase8) are located at close distance to each other, the intracellular members are divided into two sub-branches. One sub-branch consists of NTPDase4 and NTPDase7 that share their membrane topology with the surface-located enzymes. The other sub-branch consisting of NTPDase5 and NTPDase6 is characterized by a single N-terminal hydrophobic domain, which can be cleaved resulting in the formation of a soluble enzyme [2].

The surface-located members of the E-NTPDase family share 10 cysteine residues in comparable location within the sequence (Figure 4). Of these, two residues are situated between ACR1 and ACR2 and eight residues between ACR4 and ACR5. The closely related NTPDase1 and NTPDase2 possess an additional cysteine residue located at the N-terminal transmembrane domain. In contrast, rat NTPDase3 has one additional cysteine residue at the N-terminal intracellular domain and two cysteine residues at...
Around ACRI of E-NTPDases would thus correspond to the C-terminal transmembrane domain. The plasma membrane-located E-NTPDases possess seven to eight predicted N-linked glycosylation sites. Their distribution within the protein sequence is similar but not identical. Only the N-glycosylation site between ACRI and ACRI2 (N81 in rat NTPDase3) is conserved between NTPDase1, NTPDase2, and NTPDase3 (rat, mouse, and human sequences). This site is essential for full enzymatic activity of human NTPDase3 [29] but it does not exist in NTPDase8 (mouse, human). The majority of the potential N-glycosylation sites is situated between ACRI4 and ACRI5.

**Secondary structure prediction and relation to the actin/HSP70/sugar kinase superfamily**

The similarities in the distribution of the ACRs between the members of the E-NTPDase family imply structural conservation relevant for catalytic activity. It has previously been emphasized that NTPDases share two nucleotide binding motifs with the actin/HSP70/sugar kinase superfamily [3, 8, 9]. Members of this superfamily share little overall sequence identity except for the strictly conserved DXG motifs in the consensus sequence. These motifs can be identified in ACRI and ACRI4 of all E-NTPDases.

There are, however, apparent similarities in the secondary structure of members of the actin/HSP70/sugar kinase superfamily. Its members possess two major domains with similar topology (β1β2α1β4α2β5α3) that fold into a pocket for substrate binding [30–32]. The two domains presumably are the result of gene duplication. Each of these domains is composed of two subdomains (a and b). The opposing subdomains Ia and Iib share the same basic topology (β1β2α1β4α2β5α3) that is involved in nucleotide binding. The nucleotide binding site bridges both domains. The two subdomains Ib and Iib (formed by sequences in between β3 and α1) are different from one subfamily to another [30]. Figure 5 shows an alignment of the secondary structure conserved between members of the actin/HSP70/sugar kinase superfamily, as depicted for actin, with the predicted secondary structure of NTPDase1 to NTPDase8. The atomic structure of actin has previously been revealed [33, 34]. At the N-terminus, the alignment is oriented around the DXG motif in ACRI1 of NTPDases and the corresponding DXG motif in actin. The two β strands around ACRI1 of E-NTPDases would thus correspond to β1 and β2 of actin. The overall pattern of conserved actin secondary structure (α1β1α2β2α3) is very similar to that of NTPDases. This β1β2α1β4α2β5α3 topology is repeated in the second domain of actin. Similar to actin, the conserved ACRI4 of the NTPDases (second DXG motif) is embedded between two β strands. Taking further into consideration the distance between β3 and α1 in the second domain of actin and assuming that the glycine residue conserved between E-NTPDases in ACRI5 (Gly462 in NTPDase3) corresponds to the conserved glycine residue in α3 of the actin/HSP70/sugar kinase superfamily (Gly342 in actin), the secondary structure of the C-terminal half of E-NTPDases may also be aligned.

The degree of similarity in secondary structure mirrors the phylogenetic distance between sequences (comp. Figure 3). In all NTPDases, the sequences ACRI1 and ACRI4 are situated between two β strands. The alignment not only implies considerable structural conservation between E-NTPDases and members of the actin/HSP70/sugar kinase superfamily. It also suggests that NTPDases – in spite of their membrane anchorage – may consist of two major domains that repeat basic topology and key conserved sequence domains.

**Catalytic properties of heterologously expressed rat NTPDase3**

Preliminary experiments revealed that the ATPase activity at the surface of viable [20] mock-transfected CHO cells was 8.9 ± 0.4% (n = 2, triplicate determinations in each) of the activity obtained after transfection with the NTPDase3-encoding plasmid (500 μM ATP). In the isolated mem-

![Figure 6](image)

**Figure 6.** Ion dependence of NTPDase3 catalytic rate. ATPase (top) or ADPase (bottom) activity of membrane fractions derived from transfected CHO cells was analyzed in the presence of increasing concentrations of Ca²⁺ (closed symbol) or Mg²⁺ (open symbol) (500 μM ATP or ADP). The 100% values (± S.D., n = 3) correspond to 452 ± 23 and 89 ± 2 nmol/min/mg protein for ATPase and ADPase activity, respectively.
brane fraction, ATPase activity after mock-transfection was 3.1 ± 2.2% (n = 7) of the activity obtained after NTPDase3 transfection. No significant ADP hydrolysis could be determined in membrane fractions from mock-transfected cells. In the following, catalytic properties of rat NTPDase3 were analyzed in membrane fractions, whereby the catalytic activity derived from cells transfected with the empty plasmid was subtracted from that obtained with cDNA-transfected cells. Activity of rat NTPDase3 strongly depended on the presence of Ca<sup>2+</sup> or Mg<sup>2+</sup>. ATP hydrolysis was stimulated by Ca<sup>2+</sup> to a larger extent than by Mg<sup>2+</sup>, particularly at low concentrations (Figure 6). The difference is less prominent for ADP hydrolysis. In the presence of EDTA (1 mM) no significant nucleotidase activity could be measured (not shown).

ATPase activity revealed an activity optimum between pH 7.5 and 8.5 for both Ca<sup>2+</sup> and Mg<sup>2+</sup> activation (Figure 7). At pH 5 the rates were approximately 45% of maximal activity. The following experiments were performed at pH 7.4 in the presence of 500 μM Mg<sup>2+</sup> and 500 μM nucleotide substrate. An HPLC analysis of product formation using ATP as a substrate showed that ADP is formed in a time-dependent manner and accumulates in the medium. It is further hydrolyzed to AMP. No adenosine is formed (Figure 8).

In addition we compared the hydrolysis rates for ATP, GTP, CTP, and UTP as well as the respective diphosphates. NTPDase3 hydrolyzed nucleoside triphosphates–uridine or pyrimidine nucleotides – at similar rates (Table 1). Nucleoside diphosphates were hydrolyzed at considerably lower rates. In these conditions, the ratio of hydrolysis rates (triphospho- to diphosphonucleotides) varied between 4.1 and 9.5. ATP was hydrolyzed five times faster than ADP and UTP eight times faster than UDP.

### Antibody production and enzyme distribution

A rabbit polyclonal antibody was produced by genetic immunization. The specificity of the antibody was first investigated in CHO cells transfected with the cDNA encoding NTPDase1, NTPDase2, or NTPDase3 (Figure 9).

![Figure 7. pH dependence of NTPDase3 catalytic rate. ATPase activity (500 μM ATP) of membrane fractions derived from transfected CHO cells was analyzed at varying pH in the presence of either Ca<sup>2+</sup> (closed symbol) or Mg<sup>2+</sup> (open symbol) (500 μM each). The 100% values (± S.D., n = 3) correspond to 641 ± 602 and 370 ± 339 nmol/min/mg protein for Ca<sup>2+</sup>- and Mg<sup>2+</sup>-ATPase activity, respectively.](image)

When applied to the surface of viable cells the anti-NTPDase3 antibody revealed a significant surface staining of NTPDase3-transfected cells, demonstrating surface localization of the expressed protein. No immunofluorescence was obtained after application of the anti-NTPDase3 antibody to NTPDase1- and NTPDase2-expressing cells. Control experiments verified that antibodies against NTPDase1 or NTPDase2 yielded a cell surface immunosignal with CHO cells transfected with the respective cDNA (not shown).

Surface staining revealed endogenous expression of NTPDase3 by the rat-derived PC12 cells (Figure 9e). Immunofluorescence was observed over the entire cell surface. The formation of small immunofluorescent dots implies a partial clustering of the protein. In addition, immunostaining was enhanced at growth cones. Application of preimmune serum either to CHO or PC12 cells yielded negative results (not shown).

NTPDase3 immunoreactivity could be detected by immunoblotting in all brain regions analyzed. In membrane

![Figure 8. ATP hydrolysis and product formation. ATPase activity (250 μM ATP) of membrane fractions derived from transfected CHO cells was analyzed at varying time points in the presence of Ca<sup>2+</sup> (250 μM). ATP hydrolysis and product formation was monitored by HPLC. The contribution of each nucleotide is expressed as percentage of total nucleotides present in the sample. Values of 100% correspond to the initial amount of ATP substrate. Values represent means ± S.D of three experiments. Equal amounts of starting ATPase activity (7.9 nmol Pi/min) were employed for each assay.](image)

### Table 1. Substrate specificity of recombinant rat NTPDase3 in membrane fractions of transfected CHO cells.

| Substrate | Activity (% of ATPase activity) | NTP:NDP-ratio |
|-----------|--------------------------------|---------------|
| ATP       | 100.0                          | 5.0:1         |
| ADP       | 20.1 ± 1.3                     |               |
| CTP       | 126.2 ± 21.9                   | 9.5:1         |
| CDP       | 13.3 ± 2.4                     |               |
| GTP       | 103.2 ± 11.1                   | 4.1:1         |
| GDP       | 25.0 ± 3.2                     |               |
| GTP       | 92.1 ± 8.8                     | 4.6:1         |
| GDP       | 20.2 ± 0.9                     |               |
| UTP       | 109.5 ± 20.6                   | 7.7:1         |
| UDP       | 14.3 ± 6.0                     |               |

Nucleotidase activity (500 μM ATP) of membrane fractions derived from transfected CHO cells was analyzed in the presence of Ca<sup>2+</sup> (500 μM). Values represent means ± S.D of three experiments. For ATP 100% values correspond to 120 ± 67 nmol/min/mg protein.
fractions obtained from tissue extracts NTPDase3 was hardly detectable (not shown). However, when the protein was enriched by binding to concanavalin A, clear bands could be visualized at 80 kDa (Figure 10). These protein bands corresponded to that obtained from membrane fractions of CHO cells heterologously expressing NTPDase3. No immunosignal was obtained with CHO cells transfected with the empty vector or with transfected CHO cells expressing rat NTPDase1 or rat NTPDase2 (not shown).

Discussion

Rat NTPDase3 differs significantly in its functional properties from the two other surface-located NTPDases previously isolated from rat, namely NTPDase1 and NTPDase2. Whereas rat NTPDase1 expressed in CHO cells exhibits the typical catalytic properties of an apyrase with a ratio of hydrolysis rates for ATP and ADP of 1:0.8, rat NTPDase2 has dominant ATPase activity with a ratio of 1:0.05 or 1:0.03 [8, 20, 35]. When expressed in CHO cells, rat NTPDase3 reveals an ATP to ADP hydrolysis ratio of 1:0.2 and thus represents a functional intermediate between the two other NTPDases. This difference is further underlined when the product formation following hydrolysis of ATP is analyzed. Whereas NTPDase1 hydrolyzes ATP directly to AMP with minimal formation of free ADP, ADP accumulates in the reaction medium following hydrolysis of ATP by NTPDase2 [20]. NTPDase3 reveals intermediate properties. In the presence of ATP, NTPDase3 accumulates extracellular ADP which is finally converted to AMP. Rat NTPDase3 differs significantly from rat NTPDase1 and NTPDase2 regarding cation dependence. Whereas rat NTPDase1 and NTPDase2 are equally activated by Ca\(^{2+}\) or Mg\(^{2+}\) [20], rat NTPDase3 shows a clear preference for activation by Ca\(^{2+}\). In addition, ATPase and ADPase activities are differentially affected by Ca\(^{2+}\) or Mg\(^{2+}\). This differential activation remains unexplained and may depend on the difference in phosphate chain length between ATP and ADP and thus the potential coordination with the respective metal cation.
within the protein. Removal of divalent cations abolishes catalytic activity of all three enzymes. None of these enzymes hydrolyzes AMP. NTPDase3 shares, however, with NTPDase1 and NTPDase2 its broad substrate specificity towards purine and pyrimidine nucleoside triphosphates. It can be expected that the differences in catalytic properties between individual subtypes of NTPDases differentially affect P2 receptor signaling either by activating or inactivating P2 receptors [36].

The principal functional properties of rat NTPDase3 are similar to those of human NTPDase3 (HB6, [22]) and mouse NTPDase3 [23]. Interestingly, the pH-dependence of ATP hydrolysis by rat NTPDase3 clearly differs from that of mouse NTPDase3. Whereas mouse NTPDase3 expresses a considerably higher activity at pH 5 than at pH 7–8 [23], rat NTPDase3 reveals its maximal activity at alkaline pH. Interestingly, the recently cloned NTPDase8 [24] shares principal functional properties with NTPDase3 rather than with NTPDase1 or NTPDase2. Mouse NTPDase8 is preferentially activated by Ca2+ over Mg2+, has an ATP to ADP hydrolysis ratio of approximately 1:0.5 and accumulates ADP that is effectively further hydrolyzed to AMP [24]. Human NTPDase3 forms a dimer [37] and glycosylation is essential for functional expression [38]. The length of the ORF of rat, mouse [23] and human [22] NTPDase3 is identical (529 aa). The three enzymes share 13 cysteine residues and reveal 7 (rat, human) or 8 (mouse) potential N-glycosylation sites. The rat gene (31.0 kb) is slightly longer than the mouse gene (26.9 kb) but it shares its general organization into 11 exons of which exons 2 to 11 contain the ORF.

Rat NTPDase1 and 2 have previously been immuno-localized in the brain. NTPDase1 is associated with the endothelium of blood vessels and smooth muscle as well as with microglia [39, 40]. NTPDase2 is expressed by neural stem cells in the subventricular zone of the lateral ventricles [41]. The immunoblot analysis identified NTPDase3 in all brain regions investigated. NTPDase3 could be detected by immunocytochemistry at the surface of viable PC12 cells. The corresponding RNA has previously been identified in PC12 cells by RT-PCR together with that of NTPDase1 and NTPDase2 [42]. Interestingly, the hydrolysis ratio of ATP:ADP (1:0.28) and the product formation following application of ATP to viable PC12 cells is much closer to that of NTPDase3 than of NTPDase2 for which also a weak immunostaining was obtained [42]. This suggests that NTPDase3 is the predominant ecto-nucleotidase of PC12 cells. In addition we observed an enhancement of NTPDase3 immunoreactivity at growth cones of PC12 cells suggesting that the enzyme may be preferentially associated with sites of active membrane incorporation.

The availability of all NTPDase isoforms expected from genomic analysis opens the possibility of structural and functional comparison. To date structural data for this enzyme family are not available. However, the atomic structure of a considerable number of enzymes belonging to the actin/HSP70/sugar kinase superfamily, including glycerol kinase has been derived. All these proteins are soluble, have ATP phosphotransferase or hydrolyase activity, depend on divalent metal ion and tend to form oligomeric structures [32]. Individual enzyme families lack global sequence identity. They share, however, the principal structure of two major domains (I and II) of similar folds on either side of a large cleft with an ATP binding site at the bottom of the cleft [30]. These two domains are expected to undergo conformational changes involving movement relative to each other. Both, domain I and II are divided into subdomains (Ia, Ib, Ila, Ilb). Subdomains Ia and Ila have the same basic fold with conserved secondary structure elements that share considerable similarity with those of NTPDases.

A comparison of the conserved secondary structure (Figure 5) reveals duplicate conservation of DXG motifs between β strands (ACR1 and ACR4) of NTPDases that correspond to the β- and γ-phosphate binding motifs in subdomains Ia and Ila of actin, as well as a conserved glycin in ACR5 that can be identified among all members of the actin/HSP70/sugar kinase superfamily (α3, [32]). This further supports the notion that E-NTPDases are members of the actin/HSP70/sugar kinase superfamily. It implies in addition that E-NTPDases, like the other members of this superfamily, consist of two major domains with one phosphate binding motif in each domain and the binding of the nucleotide in a cleft between the two opposing domains [31]. Interestingly, some members of the E-NTPDase superfamily are entirely soluble (e.g., potato apyrase or the nucleoside triphosphatases of the protozoan parasite Toxoplasma gondii, references in [35]), others have one transmembrane domain and can be cleaved to form catalytically active soluble enzymes (NTPDase5, NTPDase6), and yet others are (NTPDase 1, 2, 3, 4, 8) are firmly anchored to the membrane via two transmembrane domains. The transmembrane domains of NTPDase1 were found to be important for maintaining catalytic activity and substrate specificity [6, 43], presumably by affecting tertiary and/or quaternary structure. The two transmembrane domains of NTPDase1 interact both within and between monomers and may undergo coordinated motions during the process of nucleotide binding and hydrolysis [44].

NTPDases differ from members of the actin/HSP70/sugar kinase superfamily by additional conserved short sequence domains (ACR2, ACR3). Differences in sequence, secondary and tertiary structure are believed to account for differences in catalytic properties between related NTPDases [35]. The essential role of the ACRs for catalytic activity has been underpinned by a considerable number of studies using point mutations in the ACRs or ACR deletions [2, 29, 45–48].

Our present study shows that rat NTPDase3 displays catalytic properties distinctly different from those of rat NTPDase1 and rat NTPDase2. Rat NTPDase3 differs from mouse NTPDase3 in its pH dependence. The enzyme is expressed in multiple brain regions and at the surface of PC12 cells. A comparison of the conserved secondary structure of actin and of NTPDases reveals apparent similarities, inferring that also basic tertiary structure may
be conserved between members of the actin/HSP70/sugar kinase superfamily and NTPDases.

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