

Mn2+-dependent ADP-ribose/CDP-alcohol pyrophosphatase: a novel metallophosphoesterase family preferentially expressed in rodent immune cells

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Running title: Mn²⁺-dependent ADP-ribose/CDP-alcohol pyrophosphatase

Abbreviations used: ADPRibase, ADP-ribose pyrophosphatase (used only for rat enzymes); GST, glutathione S-transferase; MALDI, matrix assisted laser desorption/ionisation; NDP, nucleoside 5'-diphosphate; NDP-X, nucleoside 5'-diphosphate-X; NMP, nucleoside 5'-monophosphate; ORF, open reading frame; TRPM2, transient receptor potential melastatin channel-2.

The DNA sequence reported in this paper has been deposited in GenBank under accession number EU037900.

Figures S1-S6 and Table S1 are included in the online supplement.

* Dedicated to Antonio Sillero, our teacher, on his 70th anniversary

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SYNOPSIS

The Mn²⁺-dependent ADP-ribose/CDP-alcohol pyrophosphatase (ADPRibase-Mn) is isolated from rat liver supernatants after separation from Mg²⁺-activated ADP-ribose pyrophosphatases devoid of CDP-alcohol pyrophosphatase activity (ADPRibase-I and ADPRibase-II). The latter are putative Nudix hydrolases, while the molecular identity of ADPRibase-Mn is unknown. MALDI mass spectrometry data from rat ADPRibase-Mn pointed to a hypothetical protein that was cloned and expressed, and showed the expected specificity. It is encoded by the RGD1309906 rat gene, so far annotated just as 'hydrolase'. ADPRibase-Mn is not a Nudix hydrolase, but it shows the sequence and structural features typical of the metallophosphoesterase superfamily. It may constitute a protein family of their own, which appears to be specific to vertebrates, plants and algae. ADP-ribose was successfully docked to a model of rat ADPRibase-Mn revealing its putative active centre. Microarray data from the GEO database indicated the mouse gen 2310004I24Rik, orthologous of RGD1309906, is preferentially expressed in immune cells. This was confirmed by northern and activity assay of ADPRibase-Mn in rat tissues. A possible role of ADPRibase-Mn in immune cell signaling is suggested by the second messenger role of ADP-ribose, that activates TRPM2 ion channels as a mediator of oxidative/nitrosative stress, and by the signaling function assigned to many of the microarray profile neighbours of 2310004I24Rik. Furthermore, an influence of ADPRibase-Mn in the CDP-choline or CDP-ethanolamine pathways of phospholipid biosynthesis cannot be discarded.

Key words: ADP-ribose, CDP-choline, metallophosphoesterase, Nudix, immune system, signalling.

INTRODUCTION

ADP-ribose is part of the NAD(P)-derived network of regulators. It is a primary product of NAD, formed by hydrolysis of the N-glycosidic linkage between nicotinamide and ribose, and also a secondary product formed by turnover of NAD primary metabolites [1, 2]. Its reducing ribose moiety is prone to react nonenzymically, which explains its protein-glycating character and its reputation of being cytotoxic [3-5]. On the other hand, intracellular ADP-ribose is also a regulator of ion channels. Very interesting is the role of ADP-ribose as an activator of transient receptor potential melastatin channel-2 (TRPM2) that participates in Ca²⁺-mediated cell death. The channel opens in response to ADP-ribose as a mediator of oxidative/nitrosative stress in the immune system, a major site for TRPM2 expression [6-11].

ADP-ribose levels in mammals are likely controlled by a low- K_m , Mg²⁺dependent, specific ADP-ribose pyrophosphatase [6, 12-15], that belongs to the Nudix superfamily of NDP-X hydrolases [16, 17]. However, there are other mammalian ADPribose hydrolases with diverse preferences for NDP-X substrates and for Mg²⁺ or Mn²⁺ as activators. In rat liver, ADP-ribose can be hydrolysed by the broad-specificity nucleotide pyrophosphatase or phosphodiesterase I ectoenzymes [18] and four intracellular hydrolases [19, 20]. ADPRibase-I (cytosolic) and ADPRibase-m (mitochondrial) are specific ADP-ribose pyrophosphatases that hydrolyse only ADPribose and the non-physiological IDP-ribose in the presence of Mg²⁺. Their specificity is less strict in the presence of Mn²⁺, but Mg²⁺ is likely to be the physiological activator. ADPRibase-II (cytosolic) is a Mg²⁺-dependent ADP- ugar pyrophosphatase, also less strict in its NDP-X specificity when assayed with Mn²⁺. Finally, ADPRibase-Mn is not activated by Mg²⁺, but by low micromolar Mn²⁺. Its specificity is unusual, as it hydrolyses ADP-ribose, IDP-ribose, CDP-glycerol, CDP-choline and CDPethanolamine, but not other non-reducing ADP-sugars or CDP-glucose [19].

None of these rat ADP-ribose hydrolases have been cloned, but ADPRibase-I and ADPRibase-m are near identical to human Nudix enzymes NUDT9 β and NUDT9 α [6, 13-15, 19-21], both encoded by the *NUDT*9 gene, while ADPRibase-II is related to human NUDT5 [21, 22], also a Nudix enzyme encoded by the *NUDT5* gene. In contrast, ADPRibase-Mn is not molecularly identified. Here, we are showing it forms a novel metallophosphoesterase family preferentially expressed in rodent immune tissues and cells.

EXPERIMENTAL

Chemicals and biochemicals

1,10-Phenanthroline monohydrate, and sodium salts of ADP-ribose, ADP-glucose, CDP-choline, CDP-ethanolamine, CDP-glycerol, CDP-glucose, UDP-glucose, CDP, AMP and CMP were from Sigma. ADP (potassium salt), Tris base (2-amino-2-[hydroxymethy1]-1,3-propanediol) and alkaline phosphatase (grade I from calf intestine) were from Roche. MnCl₂, MgCl₂, KCl, NaCl, and EDTA (Titriplex[™] III) were high-purity preparations from Merck.

Purification of rat liver ADPRibase-Mn and MALDI mass spectrometry of trypsin digests

Rat ADPRibase-Mn was purified from 100,000 g liver supernatants by a six-step chromatographic procedure. The first five steps were as previously described [19] and, for the final step, the enzyme obtained from 30 g of liver was adsorbed to a 2.5 cm x 1.5 cm Reactive Red 120-agarose column (Sigma) equilibrated in 20 mM Tris-HCl, pH 7.5, 0.5 mM EDTA, washed with 5 ml of KCl 0.5 M in the same buffer, followed by 10 ml of equilibrating buffer. The enzyme (3.5 ml) was recovered with a wash with water and concentrated to 0.1 ml in a centrifugal filter with a 3K-cutoff polyethersulfone membrane (Filtron Technology Corp, Northborough, MA). This preparation contained a single protein band of \approx 38 kDa. After staining for 1 h with Coomassie blue G-250 (1 g/l in 50% methanol) and destaining in 40% methanol, the band was cut out and processed for trypsin digestion and MALDI mass spectrometry as described [23] (Servicio de Proteómica, UAM, Madrid, Spain). The peptide-mass fingerprint (MS mode), and the ion spectra (MS/MS mode) of two fragments of 1746.88 Da and 2126.98 Da were used for database querying.

PCR cloning and sequencing of rat ADPRibase-Mn cDNA

The 1014-nt ORF coding for rat ADPRibase-Mn (mRNA accession number BC088174) was amplified by PCR from a rat liver Marathon Ready cDNA preparation with the AdvantageTM cDNA polymerase mix (both from Clontech). The forward primer GTCTG<u>GAATTC</u>CATGGCTGATAAACCGGAC contained an *Eco*RI site (underlined) followed by C and the 18 first nucleotides of the ORF, with a designed G12A substitution; the reverse primer GCTCC<u>GTCGAC</u>TTATTCCCTTTTGTAATTC contained a *Sal*I site (underlined) followed by the reverse complement of the 19 last nucleotides of the ORF. A product of the expected size was obtained, cut with *Eco*RI and *Sal*I, and inserted into the corresponding sites of the pGEX-6P-3 plasmid (Amersham) to obtain plasmid pGEX-6P-3-rADPRMn. Both strands of the insert were sequenced in the Servicio de Secuenciación Automática (IIB, CSIC-UAM, Madrid, Spain).

Recombinant ADPRibase-Mn expression and purification

One hundred-ml cultures of BL21 cells transformed with pGEX-6P-3-rADPRMn, were grown at 30°C and induced with IPTG. The GST/ADPRibase-Mn fusion recovered in post-sonication supernatants (in 10 ml of 20 mM Tris-HCl, pH 7.5, 0.5 mM EDTA, 1 mM DTT, 50 mM KCl, containing one dissolved tablet of protease inhibitor cocktail CompleteTM Mini; from Roche) was adsorbed to a GSH-Sepharose column (1.5 ml; Amersham) equilibrated in the same buffer without protease inhibitors. The unbound proteins were washed out with 15 ml of 50 mM Tris-HCl, pH 7.5, 1 mM EDTA, 1 mM DTT, 150 mM NaCl. ADPRibase-Mn was separated from the GST tag by in-column proteolysis with PreScissionTM protease followed by elution with wash buffer. The eluted enzyme (1.8 ml) was frozen at -80°C in 50- μ l aliquots. This was a high-purity enzyme preparation (Figure S1, lane 6). Upon thawing, the enzyme was stable at least for one week. The expression and purification procedure was carried out three times with the same results.

Native molecular weight of ADPRibase-Mn by gel filtration chromatography

Samples of ADPRibase-Mn and molecular weight markers were chromatographed in a Protein-Pak 125 column (7.8 mm x 300 mm) from Waters, equilibrated in 50 mM Tris, pH 7.5, 1 mM EDTA, 150 mM NaCl, and connected to a Hewlett-Packard HP1090 chromatograph. The elutions were performed with equilibrating buffer at 1 ml/min and the chromatographic profiles were recorded at 280 nm. For ADPRibase-Mn activity assay, 0.25 ml fractions were collected.

Enzyme activity assays

The hydrolytic activities of ADPRibase-Mn on NDP-X substrates were assayed at 37°C by coupling to alkaline phosphatase and measuring colorimetrically the amount of P_i formed with a dodecyl sulphate-ascorbate-molybdate reagent [21]. The assay mixtures contained, except when otherwise stated, 50 mM Tris-HCl (pH 7.5 at 25°C), 500 µM NDP-X, 100 μ M MnCl₂, 1.5 μ g/ml (5.5 units/ml) alkaline phosphatase, and the appropriate amount of enzyme sample. For the assay of the chromatographic profiles of ADPRibase-I, ADPRibase-Mn and ADPRibase-II in fractionated rat tissue extracts, 1 mg bovine serum albumin/ml was included in the reaction mixtures. Whenever the substrate was NDP or NMP, alkaline phosphatase was omitted. For the pH-activity profiles, different buffers were used, MnCl₂ was increased to 1 mM, and the concentration of alkaline phosphatase was doubled. In every case, the reaction was stopped and colour developed by addition of either 0.7 ml of standard P_i reagent to 0.05-0.1 ml or, when higher sensitivity was needed, 0.5 ml of a concentrated reagent to 0.4 ml of reaction mixture. For studies of enzyme saturation, initial rates were measured at diverse substrate concentrations and the data were adjusted by non-linear regression to the Michaelis-Menten equation [24]. For inactivation studies with o-phenanthroline, the activity was assayed in standard reaction mixtures without alkaline phosphatase, by measuring the conversion of ADP-ribose to AMP by HPLC after 15 min of reaction. Enzyme activities were linear with time and amount of enzyme. Blanks without enzyme and/or substrate were run in parallel. One unit enzyme activity is defined as 1 μ mol substrate hydrolysed/min.

HPLC detection of NMP as reaction product of ADPRibase-Mn

Reaction mixtures with all the substrates hydrolysed by ADPRibase-Mn were analysed by ion-pair reverse-phase HPLC in a 15 cm x 0.4 cm octadecylsilica column (Kromasil 100, Teknokroma, Spain) with a 1 cm x 0.4 cm pre-column of the same material, using a Hewlett-Packard HP 100 chromatograph. Two elution buffers were used: A, 5 mM sodium phosphate, pH 7.0, 20 mM tetrabutylammonium and 10 % (vol/vol) methanol; B, same as A with 100 mM sodium phosphate. Before each run, the column was equilibrated in buffer A. Samples of 20 μ l (reaction mixture) were injected, and the elution was accomplished at 1 ml/min with a linear, 10-min 5–100 mM phosphate gradient, followed by a 10-min isocratic wash with buffer B. The chromatograms were recorded at 260 nm. The retention times of compounds relevant to the reactions catalysed by ADPRibase-Mn were (min): adenosine, 2.6; AMP, 6.6; ADP, 11.7; ADPribose, 8.5; cytidine, 1.5; CMP, 4.4; CDP, 7.7; CDP-ethanolamine, 2.3; CDP-choline, 2.2; CDP-glycerol, 5.0. This allowed the identification of either AMP or CMP as the nucleotide product of hydrolysis for every substrate of ADPRibase-Mn.

Multiple tissue northern blot

A multiple rat tissue northern blot containing 2 μ g mRNA from each tissue, and prestained with Blot Stain Blue (Figure S2; Sigma), was used. The blot was probed with a digoxigenin-labeled cDNA containing the full ORF of rat ADPRibase-Mn. The labelling was performed with the DIG High Prime DNA Labeling kit (Roche) according to manufacturer instructions. Hybridisation and detection were performed by the optimized protocol described elsewhere [25] except that probe concentration was 10 ng cDNA/ml, the hybridisation steps were carried out at 55°C, and ready-to-use CSPD was the substrate for the chemiluminescent reaction catalysed by alkaline phosphataselabeled anti-digoxigenin (Roche). The results were recorded first by luminescent quantitative detection (Kodak Image Station 2000R) under conditions in which signal accumulation was linear with time and with the amount of labelled probe and, second, by exposure of X-ray film to the chemiluminescent blot.

Rat tissue or cell extract fractionation to assay ADPRibase-I, ADPRibase-Mn and ADPRibase-II

Female Wistar rats weighing 200-220 g, fed ad libitum, were provided by the Servicio de Animalario of the Universidad de Extremadura, Badajoz. The animals were killed by rapid decapitation and the organs of interest (liver, skeletal muscle, spleen and thymus) were immediately dissected. After washing in cold saline, the tissue was blotted dry and weighed. Depending on the organ, 0.35-1.3 g of tissue were processed.

The protocol to obtain rat splenocytes and to remove red blood cells was as described [26], except that spleens were crushed in 10 ml of cold saline, the red cell lysing buffer was purchased from Roche, and the final washes of splenocytes were performed with 14.5 mM Tris, pH 7.6, 5 μ M CaCl₂, 98 μ M MgCl₂, 0.54 mM KCl, 126 mM NaCl, and 0.1 g/l of glucose.

The separation of ADPRibase-I, ADPRibase-Mn and ADPRibase-II was carried out, with some simplification, as described [19]. The tissue samples were homogenised on ice, in buffer containing 50 mM Tris, pH 7.5, 0.5 mM EDTA, 1 mM PMSF, using 6 ml/g tissue (liver and muscle), 9 ml/g tissue (spleen and thymus), or 4.5 ml per washed splenocyte pellet obtained from one spleen. A hand-driven Potter type was used, except for muscle that was homogenized first in a motor-driven blendor and then finished in the Potter. With every tissue, 4-5 ml of homogenate was centrifuged for 60 min at 4°C and 33,000 rpm in a Beckman SW45 rotor. The supernatant was chromatographed in a Sephadex G-100 column (1 cm x 90 cm) equilibrated and eluted with 20 mM Tris, pH 7.5, 0.5 mM EDTA at 20 ml/h. Fractions of about 2.5 ml were collected and the Mg²⁺and the Mn²⁺-dependent ADP-ribose hydrolase activities were assayed. A small peak of activity eluting in the void volume was discarded, and the fractions of a large, included peak of ADP-ribose hydrolase, active with both Mg^{2+} or Mn^{2+} , were pooled. This peak contained ADPRibase-I, ADPRibase-Mn and ADPRibase-II, and it was applied to a DEAE-cellulose column (1.5 cm x 12 cm) equilibrated in 20 mM Tris, pH 7.5, and washed with 2-3 volumes of 20 mM Tris, pH 7.5, 0.5 mM EDTA, 50 mM KCl, followed by a 130-ml linear gradient of 50-400 mM KCl in the same buffer. The three enzymes were resolved within the gradient, allowing their individual quantitation. A careful accounting was kept such that activities quantitated after the DEAE-cellulose step could be extrapolated back and referred to total tissue protein measured in the homogenate with bovine serum albumin as a standard [27].

Docking of ADP-ribose to the homology model of rat ADPRibase-Mn

The theoretical coordinates of rat ADPRibase-Mn were taken from the SwissProt Repository [28]. This homology model is based on the crystal structure of the zebrafish protein (pdb entry: 2NXF), which contains two Zn^{2+} ions in a typical metallophosphoesterase dinuclear centre. Since there was an excellent structural similarity between the zebrafish and the rat proteins concerning the residues located within 5 Å of the Zn^{2+} ions (RMSD 0.082 Å), the metal coordinates of the crystal structure were used to place them in the homology model. The protonation state of His side chains was assigned with the program Reduce [29]. Atom types and Kollman partial charges were assigned with AutoDockTools (ADT). The zinc ions were modelled with charge 2+ and default AutoDock 4 radius and well depth. ADP-ribose was prepared with Marvin 4.1.13, 2007, ChemAxon (http://www.chemaxon.com), its Gasteiger charges were calculated with ADT (total charge: 2-), and all its torsionable groups were made flexible except 6-NH₂, and 2', 3', 2'' and 3''-OH. Docking of ADPribose was run with AutoDock 4 [30] within a scoring grid (sized 33.8 × 29.2 × 29.2 Å with a resolution of 0.375 Å) centred around the putative active-site. AutoDock was run with the Lamarckian genetic algorithm with default parameters except that: 100 independent runs were performed; the maximum number of energy evaluations per run was set to 5×10^7 ; and the step sizes were adjusted to 0.2 Å for translations and 5° for orientations and torsions.

RESULTS

Tandem-mass spectrometry of rat liver ADPRibase-Mn pointed to a hypothetical rat protein that was cloned and expressed

For identification, a peptide-mass fingerprint of the 38-kDa band of rat liver ADPRibase-Mn was obtained. In a Mascot search [31] against the NCBInr database, it did not give any significant candidate. Therefore, two peptides of 1746.88 Da and 2126.98 Da were analysed by tandem mass spectrometry. Searches with these data pointed to two tryptic peptides formed by amino acids 47-61 (HSLVHLQGAIEDWNK) and 102-118 (VPVHHTWGNHEFYNFSR) of a 337 amino-acid hypothetical rat protein (accession No. AAH88 174). Its theoretical mass of 38710 Da agreed with the 38-kDa protein band. It was recorded as a product of the rat gene *RGD1309906* (GeneID 287406) which was annotated just as 'hydrolase'. The coding sequence (accession No. BC088174) of the hypothetical protein was used to design the PCR cloning described under Experimental.

Plasmid pGEX-6P-3-rADPRMn contained, in frame with a GST affinity label, the PCR-amplified, 1014-nt ORF coding for ADPRibase-Mn. Relative to the ORF used as reference, it displayed 3 nucleotide changes: G12A (due to the design of the forward PCR primer), A131G and A636G, only one causing an amino acid change (Y44C). This sequence was deposited in GenBank (accession No. EU037900).

The GST/ADPRibase-Mn fusion of theoretical 65080 Da was expressed from pGEX-6P-3-rADPRMn as a \approx 65-kDa protein (Figure S1, lane 2), and it was adsorbed to GSH-Sepharose. The ADPRibase-Mn moiety was recovered by in-column proteolysis as a \approx 40-kDa protein (Figure S1, lane 6). This agrees with an expected mass of 39360 Da, sum of ADPRibase-Mn and the N-terminal extension (GPLGSPNS) left by proteolysis.

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Characterisation of the recombinant protein as a Mn²⁺-dependent ADP-ribose/CDPalcohol pyrophosphatase identical to ADPRibase-Mn

Like rat liver ADPRibase-Mn [19], the recombinant enzyme hydrolysed ADP-ribose and CDP-alcohols. At 500 μ M substrate and 100 μ M MnCl₂, CDP-choline gave the highest rate, followed by CDP-glycerol, CDP-ethanolamine, ADP-ribose and ADP (Table 1). However, kinetic parameters derived from hyperbolic saturation curves (Figure S3) showed ADP-ribose as the best substrate ($k_{cat}/K_m = 328000 \text{ M}^{-1}\text{s}^{-1}$), whereas CDP-alcohols and ADP were hydrolysed with 3.5-15 fold lower efficiencies (Table 1). ADP-glucose, UDP-glucose, CDP-glucose, CDP, CMP and AMP were not hydrolysed.

All the reactions involved the hydrolysis of the phosphoanhydride linkage, as: (i) P_i was formed by alkaline phosphatase from the NDP-X reaction products but not from the substrates; (ii) P_i was directly formed from ADP; (iii) AMP or CMP were identified as products by HPLC (results not shown).

ADPRibase-Mn was inactive without added Mn²⁺ and showed high affinity for it, since saturation was observed at low micromolar Mn²⁺, well below the 500 μ M substrate concentration (Figure S3). The concentration of MnCl₂ giving the half-maximal rate was 1–4 μ M. In no case, Mg²⁺ substituted for Mn²⁺.

Incubation with 2.5 mM EDTA did not affect ADPRibase-Mn activity. This was in agreement with the stability of the enzyme frozen for months in a buffer with 1 mM EDTA. However, *o*-phenanthroline caused near full inactivation in 1.5 h (Figure 1).

The effect of pH on ADPRibase-Mn activities was studied in the range of pH 6– 10. The profile of ADP-ribose hydrolase showed an optimum at pH 8.0–8.5, whereas the activities on ADP and CDP-alcohols were highest at pH 6–8, and decreased abruptly at higher pH (Figure 2).

ADPRibase-Mn behaved as a monomer, with an apparent mass of ≈ 30 kDa by gel filtration (Figure S4), in good agreement with the native mass of 32 kDa observed for the enzyme purified from rat liver [19].

Characterisation of ADPRibase-Mn as a protein belonging to the metallophosphoesterase, not to the Nudix superfamily

In contrast to other ADP-ribose pyrophophatases, ADPRibase-Mn lacked the amino acid signature of the Nudix superfamily. Searches in the InterPro site (http://www.ebi.ac.uk/InterProScan) and Superfamily database (http://supfam.org/SUFERFAMILY) identified a metallophosphoesterase domain (IPR004843), indicating that ADPRibase-Mn belongs to the Pfam metallophos family (accession ID PF00149), and is a member of the Scop metallo-dependent phosphatase superfamily (accession ID SSF56300). These proteins contain (1) a dimetal centre with diverse ion pairs or combinations, (2) a $\beta\alpha\beta\alpha\beta$ secondary structure signature within (3) a four-layered fold with two β -sheets flanked by α -helices ($\alpha/\beta/\beta/\alpha$ fold), and (4) a disperse sequence signature that includes, in five conserved regions, the amino acids coordinated with the metal ions: DX[H/X]-(X)n-GDXX[D/X]-(X)n-GNH[D/E]-(X)n-[G/X]H-(X)n-GHX[H/X] [32-35]. The aligment of ADPRibase-Mn orthologues found by BlastP searches (Table 2) revealed these five regions (Figure 3, I-V).

The crystal structure of the zebrafish (*Danio rerio*) orthologue complexed with P_i and two Zn²⁺ ions [36], and a homology model of the rat protein [28] were found in public databases. They display the $\alpha/\beta/\beta/\alpha$ metallophosphoesterase fold, with the five regions of the sequence signature brought together each at the C-end of a different β -strand: regions I-III belong to one of the β -sheets, and regions IV-V to the other. Seven of the conserved amino acids, which show an excellent structural similarity between the zebrafish and the rat proteins, form a dimetal centre located at the bottom of a groove.

ADP-ribose was succesfully docked to the rat protein model into this site, indicating that it can be the active centre (Figure 4A,B). The amino acids that form the dimetallic centre and/or might interact with the substrate are shown in Figure 4C.

The gene encoding ADPRibase-Mn in Rattus norvegicus and mammalian orthologues

A BlastN query to the *R. norvegicus* genome using the ORF sequence of rat ADPRibase-Mn revealed that the gene *RGD1309906* (GeneID 287406; also known as *MDS006* and *MGC108803*) is the only significantly related locus. It is a predicted gene located in rat chromosome 10 (10q24) that contains five exons, of which only the three last ones are translated and were hit by the BlastN search. The ADPRibase-Mn ORF spans from the 5'-end of exon 3 to within exon 5 (Figure 5). The reference rat genome contains the coding sequence of hypothetical protein LOC287406 (accession No. BC088174), identical to that coding for recombinant ADPRibase-Mn except for the nucleotide differences mentioned above. Figure 5 shows also the organisation of the mouse and human genes and the ORFs coding for ADPRibase-Mn orthologues in these organisms. TBlastN and BlastN searches pointed to an inverse 427-pb repetition including most of exon 5 and located 130 kb downstream of the rat gene, a feature not observed in the mouse or human genomes.

In human, mouse and rat, in addition to ADPRibase-Mn orthologues of 337-342 amino acids (Figure 3), BlastP searches found shorter protein variants due to an alternative 5'-splice site of the intron following the first translated exon. They were either conceptual translations from alternatively spliced transcripts (e.g. accession Nos BC001294 and BC070155 for human, and AK009142 for mouse mRNAs) or genomic predictions (accession number EDL10423 for mouse and EDM04778 for rat proteins). The alternative 5'-splice leaves 86-nt of the intron 5'-end attached to the preceding exon after splicing (Figure 5). The alternative transcription pattern contains an in-frame stop codon either within the retained intron segment (human and mouse) or in the next exon (rat). By the time of submitting this article, some databases point to the shorter version of the protein as the gene product.

Restricted taxonomic distribution of ADPRibase-Mn orthologues

Orthologous genes of *RGD1309906* are recorded only among vertebrates, higher plants and algae. As this seemed an important phylogenetic feature, we run BlastP and TBlastN searches that found ADPRibase-Mn orthologous in those phyla, but not in invertebrates and fungi (Table 2). In the latter cases, TBlastN searches of complete genomes, confirmed that the absence of orthologous proteins was not due just to their not being recorded yet.

ADPRibase-Mn gene expression: differential expression in immune and non-immune rodent tissues

Upon inspection of the Gene Expression Omnibus (GEO) database for ADPRibase-Mnrelated genes, a genomics-based investigation aimed to identify mouse immune-genes came to our attention (GEO dataset GDS2068 [37]). Out of the 8734 genomic features explored in this microarray experiment, 360 show preferential expression in thymus, spleen, peripheral blood mononuclear cells, lymph nodes (unstimulated or stimulated), or *in vitro* activated T-cells. These features were linked to 333 unique genes, one of them being 2310004I24Rik, the orthologue of ADPRibase-Mn. The probe representing this gene in the microarray is a cDNA of the alternatively spliced mRNA (AK009142) that encodes the short protein variant (see above and Figure 5). It is likely to hybridize

with both types of transcripts as they differ only in a 86-nt insert within a length of ≈ 1500 nt. The expression profile of 2310004124Rik is shown in Figure S5, together with the heat plot of its profile neighbours (listed in Table 3 with their assigned functions [37]). To confirm the preferential expression of ADPRibase-Mn in rodent immune tissues, the UniGene database was inspected, and a northern blot analysis and a fractionation study of the ADP-ribose pyrophosphatases in soluble extracts of selected rat tissues were run.

Using the UniGene's EST Profile Viewer to analyse mouse ESTs, one can see that ADPRibase-Mn is most strongly expressed in thymus (Table S1). Expression in spleen is also observed but not as prominently as in the microarray.

The northern blot was hybridised with a rat cDNA probe that revealed a major transcript of ≈ 1.4 kb in all tissues analysed (Figure 6), in agreement with the structure of *RGD1309906* gene (Figure 5). Longer transcripts of ≈ 3 kb and ≈ 5 kb were also observed in some tissues. Quantitation of the RNA signals indicated higher expression in thymus and spleen than in the other rat tissues.

As stated in the Introduction, besides ADPRibase-Mn, rat liver supernatants contain other two ADP-ribose pyrophosphatases, named ADRibase-I and ADPRibase-II, which work with either Mg^{2+} or Mn^{2+} as the activating cation [19]. To distinguish the ADP-ribose pyrophosphatase activity of ADPRibase-Mn from the other hydrolases, fractionation of tissue supernatants was necessary (Figure S6). This allowed the quantitation of the three enzymes in five different rat sources: thymus, spleen, splenocytes, liver and skeletal muscle (Table 4). ADPRibase-Mn activity in thymus and spleen was 2.5-5-fold higher than in liver and muscle. The ratio was even higher when ADPRibase-Mn was quantitated in splenocytes, where the activity was 4-8-fold higher than in the non-immune tissues (Table 4).

DISCUSSION

Our results provide a molecular identity for ADPRibase-Mn and a biochemical basis for the functional annotation of the rat gene *RGD1309906*, likely extrapolable to other mammalian genes coding for strongly similar hypothetical metallophosphoesterases. This includes the mouse gene 2310004124Rik and the human gene c17orf48, which respectively encode proteins 95% and 85% identical to rat ADPRibase-Mn.

Not every known NDP-X hydrolase is a member of the Nudix superfamily [38, 39] but, in fact, all those with a clear or relative preference for ADP-ribose [13-15, 22, 40, 41] or CDP-alcohol [41, 42] are. It has been pointed out that the Mn²⁺-dependent Nudix hydrolase YZGD, from *Paenibacillus thiaminolyticus*, displays a specificity similar to ADPRibase-Mn [41]. Therefore, that the latter is a metallophosphoesterase was relatively unexpected. In the Scop classification, the metallo-dependent phosphatase superfamily is currently divided in several families, including: purple acid phosphatase; DNA double-strand break repair nuclease; 5′-nucleotidase (syn. UDP-sugar hydrolase), N-terminal domain; protein serine/threonine phosphatase; and others. Searches in the Superfamily server (http://supfam.org/SUPERFAMILY) failed to asign ADPRibase-Mn to any established Scop family (no E-value < 0.01 was obtained), suggesting that it may constitute a novel protein family that can be provisionally described as ADP-ribose/CDP-alcohol pyrophosphatase.

ADPRibase-Mn is inactive in the absence of added metals, and is activated by low micromolar concentrations of Mn²⁺, but not of Mg²⁺. Activation by Mn²⁺ reflects metal binding to the enzyme, not just to the free substrate, as half saturation was obtained at 1-4 μ M MnCl₂, >100 fold lower than the substrate. This is compatible with Mn²⁺ being a

natural activator of the enzyme, as free Mn^{2+} is estimated to be 0.7 μM in hepatocytes from fed rats and 0.25 μM in hepatocytes from fasted rats [43].

The metal centre of metallophosphoesterases can contain a couple of the same or different metals in varying combinations. However, it is unclear whether activation by Mn^{2+} reflects binding to the dinuclear centre or otherwise. In fact, the structure of the zebrafish orthologue of ADPRibase-Mn was determined with two Zn²⁺ ions in the dinuclear centre, but it also contained two Zn²⁺ ions at other locations [36]. The time-dependent inactivation of rat ADPRibase-Mn by phenanthroline, not by EDTA, could indicate that a metal different from Mn^{2+} could be part of the enzyme, perhaps Fe or Zn, that form more stable complexes with the former chelator. However, phenanthroline could also be acting as an inhibitor by binding to the active centre. In summary, further evidence is needed to elucidate the nature of the metals bound to ADPRibase-Mn and their structural and/or functional role.

The putative active centre of ADPRibase-Mn is in the groove where P_i lies bound to the Zn^{2+} ions of the crystal structure of the zebrafish protein [36]. In the homology model of the rat protein, a docked ADP-ribose fits into that groove, with the β phosphate of ADP occupying a position similar to P_i in the X-ray structure. All the ADP-ribose interactions with the Zn^{2+} ions, and most of those established with the protein, involve the pyrophosphate group (Figure 4 and legend), and it is likely that some of them are important for catalysis. For instance, His111 (part of the conserved GNHE region III in Figure 3) interacts with the oxygen of the phosphoanhydride linkage hydrolysed by ADPRibase-Mn, as in other metallophosphoesterases where this interaction is catalytically relevant [44]. On the other hand, the specificity of ADPRibase-Mn for NDP(-X) compounds (Table 1) should be accounted for by interactions of the N and the X portions of the substrate. In the ADP-ribose docking model, while the pyrophosphate group is in a rather closed environment, adenine and the distal ribose protrude towards more open areas of the protein surface (Figure 4B). On the N side of ADP-ribose, a hydrophobic contact of adenine with Leu196 is observed, but not any hydrogen bond. On the X side, two possible interactions of the 1"-OH group with backbone atoms of His278 and His280, and a favorable contact of the ribose ring with Cys253 are observed. It is unclear whether these interactions alone account for the preferences of ADPRibase-Mn for ADP versus CDP, for NDP-X versus the corresponding NDPs, and for ADP-ribose versus ADP-glucose as summarized in Table 1. Anyhow, it must be stressed that the theoretical structure of rat ADPRibase-Mn is derived by homology to a zebrafish protein complexed with P_i, a ligand much smaller than ADP-ribose. In a true enzyme-substrate complex, specific interactions may depend on an induced fit of the enzyme which P_i would not possibly evoke.

ADPRibase-Mn may have a function in vertebrate immune systems. Mining of expression databases identified a mouse microarray experiment in which gene 2310004124Rik, that codes for the mouse orthologue of ADPRibase-Mn, is labeled as an 'immune gene', preferentially expressed in immune versus non-immune cells and tissues [37]. As a complement to this, rat ADPRibase-Mn mRNAs were found more abundant (Figure 6) and enzyme activity was higher (Table 4) in thymus and spleen than in non-immune rat tissues. ADPRibase-Mn activity was even higher in splenocytes, that include spleen immunocytes (Table 4). Such a pattern was not observed for the Nudix-type ADPRibase-I (Nudt9) and ADPRibase-II (Nudt5), either when assaying enzyme activities in rat tissues (Table 4) or in the RNA expression of the mouse orthologue of the rat *Nudt5* gene that codes for ADPRibase-II (a probe for the *Nudt9* orthologue, that codes for ADPRibase-I, was not included in the microarray [37]). Also in agreement with a role in the vertebrate immune system, a restriction was found in the taxonomic distribution of ADPRibase-Mn orthologues. BlastP and TblastN searches revealed they occur in all vertebrates with the possible exception of birds, but not in invertebrates and

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most lower eukaryots (Table 2). This contrasts, e.g., with genes orthologous to Nudt9 (coding for ADPRibase-I) which, besides in vertebrates, is represented in Drosophila and C. elegans [14], or to Nudt5 (coding for ADPRibase-II), which is represented in C. elegans and Saccharomyces [22]. The immune role of ADPRibase-Mn is unknown. In the above-mentioned microarray study [37], a cellular function was assigned to 298 of the 333 mouse immune genes identified, grouped in about 20 functional sets labeled as e.g. defense, apoptosis, immune, signaling, metabolism, etc (see e.g. Table 3). Immunegene 2310004I24Rik, here shown to code for the mouse orthologue of ADPRibase-Mn, was annotated just as metallophosphoesterase, and could not be assigned to either of those functional sets [37]. The role of ADP-ribose as a second messenger of stress in immune cells [9, 10, 45-47] points to a signaling role for ADPRibase-Mn. This agrees with the roles of the immune genes that are profile neighbours of 2310004I24Rik in the microarray study. While 16% of the 298 immune genes with a known role are marked 'signaling' [37], this increases to 43% among the profile neighbours of 2310004I24Rik (Table 3). ADPRibase-Mn in the immune system could be related to ADP-ribose turnover and to the termination of its effects on TRPM2 channels. Since the immunegene microarray does not contain a TRPM2 probe, to compare 2310004I24Rik and TRPM2 expression in mouse tissues, we inspected the UniGene database using the EST Profile Viewer. Both genes are expressed at low levels (e.g. as compared with β -actin), what makes comparisons difficult in terms of numbers of ESTs counted. Anyhow, considering only body sites for which a high number of total ESTs is on record (> 40000), one can see that TRPM2 expression is accompannied by ADPRibase-Mn. This represents only a partial correlation, because several sites express ADPRibase-Mn but not TRPM2 (Table S1).

CDP-choline and CDP-ethanolamine are key intermediates in biosynthetic pathways of, respectively, phosphatidylcholine and phosphatidylethanolamine [48, 49]. The formation of CDP-choline by CTP-phosphocholine cytidylyltransferase is ratelimiting and important for the regulation of phosphatidylcholine biosynthesis. ADPRibase-Mn is the only mammalian NDP-X hydrolase showing a relative preference for CDP-choline (Table 1). Working simultaneously with the cytidylyltransferase, ADPRibase-Mn would form a futile substrate cycle that could contribute also to regulation:

(cytidylyltransferase) CTP + phosphocholine \rightarrow CDP-choline + PP_i

(ADPRibase-Mn) CDP-choline + $H_2O \rightarrow CMP$ + phosphocholine

A similar reasoning can be applied to the CDP-etanolamine pathway. In conclusion, a role of ADPRibase-Mn related to its CDP-alcohol substrates, cannot be discarded.

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	Enzyme purified from rat liver	Enzym	ne expressed t (this w	from cloned D vork)	NA
Substrate ^a	Activity at 500 μ M substrate ^b	Activity at 500 μ M substrate ^c	$k_{\rm cat}{}^{\rm d}$	$K_{ m m}{}^{ m d}$	$k_{\rm cat}$ / $K_{\rm m}$
	(%)	(%)	s^{-1}	$\mu \mathbf{M}$	$\mathbf{M}^{-1} \mathbf{s}^{-1}$
ADP-ribose	100	100 ± 8	12.8 ± 0.2	39 ± 2	328200
CDP-choline	140	156 ± 23	28.5 ± 0.6	300 ± 22	95000
CDP-ethanolamine	89	84 ± 18	41.3 ± 1.1	1422 ± 99	29000
CDP-glycerol	89	103 ± 10	28.1 ± 1.4	304 ± 59	92400
ADP	27	26 ± 4	4.4 ± 0.2	201 ± 30	21400

Table 1 Substrate specificity and kinetic parameters of ADP-ribose/CDP-alcohol pyrophosphatase

^a Neither the purified nor the expressed enzymes displayed significant activity over

ADP-glucose, UDP-glucose, CDP-glucose, CDP, CMP and AMP (< 3%).

^b Taken from [19].

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^c Means \pm S.D. of 3 experiments; 100% was 23 μ mol/min/mg; protein assayed according to [50].

^d Kinetic parameters are expressed \pm S.E. of the pooled data of 2-3 experiments [24].

EU037900) as the query ag program failed to find a put genomes of relevant taxono Species Homo sapiens Mus musculus Gallus gallus Xenopus laevis Danio rerio Insects	The NCB1 site using the concerning the NCB1mr protein data ative orthologous protein, TB1 mic groups. The results showr Orthologous protein Marne Name Name Name Similar to putative protein product of Nbla03831 x 0006 protein Hypothetical protein LOC393393	eptual transls base. Only re lastN searcho lastN searcho BlastP E-value 6×10^{-176} 7×10^{-30} 7×10^{-30} 6×10^{-84} 6×10^{-84} 6×10^{-84}	tion of the presentatives s were run ned on Sept (%) (%) (%) (%) (%) 52 53 53 53 53 50 50 50 50 50 50 50 50 50 50 50 50 50	ORF cloned e hits are sho against the N against the N Ember 8, 200 (%) (%) 93 97 97 65 65 65 68 68 71 71 71 71 71	from rat liver cD wn for vertebrat VCBInr nucleotid 77. Query coverage (%) 100 100 38 38 95 94	NA (Gen es and pla le databas Amino acids 342 342 342 356 356 322	Bank accession unts. When the Blastl e and the assembled Accession number NP_064618 AAH04029 XP_001231601 NP_001080428 NP_001080428 NP_956715	c Gene 56985 66358 768578 380120 393393
Nematodes Echinoderms Fungi		>0.1 (best] >0.1 (best] >0.1 (best]	FBlastN E- FBlastN E- FBlastN E-	value = 0.35) value = 4.3) value = 0.033		4		
Oryza sativa	Os07g0688000 Os08g0557200	2 x 10 ⁻⁶³ 3 x 10 ⁻⁵⁸	41 40	59 54	96 94	321 339	NP_001060698 NP_001062482	4344352 4346295
Arabidopsis thaliana Ostreococcus lucimarinus	Calcineurin-like phospho- esterase family protein Predicted protein	6 x 10 ⁻⁵² 7 x 10 ⁻³²	37 31	<i>57</i> 51	92 93	311 331	NP_194204 XP_001419532	828575 5003684
(green algae)								

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Table 2 ADPRibase-Mn orthologues in representative vertebrates and plants, and absence of orthologues in invertebrates and fungi

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Table 3 ADPRibase-Mn profile neighbours of expression among the genes preferentially expressed in mouse immune cells and tissues

The list was found by automatically seaching for *2310004I24Rik* gene profile neighbours in the GEO dataset GDS2068 within the GEO Profiles database at the NCBI site. Out of the 31 records found, only those classified as 'immune genes' are listed together with their assigned functions [37]. Expression profiles are shown in Figure S5.

Gene symbol	Full name	Assigned function
Cd53	CD53 antigen	signaling
Coro1a	coronin, actin binding protein 1A	signaling
1200013B08Rik	RIKEN cDNA 1200013B08 gene	signaling
(Sly1)		
Adcy7	adenylate cyclase 7	signaling
Ptprcap	protein tyrosine phosphatase, receptor type, C polypeptide-associated protein	signaling
Syk	spleen tyrosine kinase	signaling
Cd37	CD37 antigen	signaling
Ptprc	protein tyrosine phosphatase, receptor type, C	signaling
Il2rg	interleukin 2 receptor, gamma chain	signaling
H2-Eb1	histocompatibility 2, class II antigen E beta	immune
H2-Oa	histocompatibility 2, O region alpha locus	immune
H2-Ob	histocompatibility 2, O region beta locus	immune
Cd79b	CD79B antigen	immune
Ms4a1	membrane-spanning 4-domains, subfamily A, member 1	immune
Ms4a4b	membrane-spanning 4-domains, subfamily A, member 4B	immune
Ncf4	neutrophil cytosolic factor 4	defense
Sp100	nuclear antigen Sp100	cell cycle
Ifi203	interferon activated gene 203	apoptosis
Itgb2	integrin beta 2	adhesion
Pou2af1	POU domain, class 2, associating factor 1	transcription
Tap1	transporter 1, ATP-binding cassette, sub-family B (MDR/TAP)	transport
Nup210	nucleoporin 210	nuclear pore

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Table 4 Preferential expression of ADPRibase-Mn activity in rat immune tissues

The activities were quantitated after separation of the three ADPRibases by gelfiltration of tissue supernatants followed by ion-exchange chromatography (Figure S6). The results are referred to an amount of tissue containing 1 g protein (assayed in the crude homogenates), and are means \pm S.D. of three independent experiments, each corresponding to one animal.

	ADPRibase-I	ADPRibase-Mn	ADPRibase-II
	mil	liunits / g tissue protei	n
Thymus	470 ± 140	330 ± 80	170 ± 30
Spleen	500 ± 80	300 ± 50	160 ± 60
Splenocytes	410 ± 40	550 ± 190	190 ± 30
Liver	590 ± 140	130 ± 30	270 ± 70
Skeletal muscle	180 ± 20	70 ± 20	60 ± 10

20 LicencedcopCopingisnotpermittedeseptithpriorpermissionandasalloodblaw 2008TheAuthorsJournalcompilation2008BiochemicalSociety Figure 1 Inactivation of recombinant ADPRibase-Mn by o-phenanthroline

Enzyme samples were diluted 1:1 in 5 mM (\bullet) *o*-phenanthroline or (O) EDTA each dissolved in 50 mM Tris, pH 7.5, and were incubated for the indicated lengths at 37°C. Enzyme aliquots were taken at the end of the incubation, diluted 100-fold in 50 mM Tris-HCl, pH 7.5, and immediately used for ADP-ribose hydrolase activity assay by HPLC.

Figure 2 pH profiles of the activities of recombinant ADPRibase-Mn

Initial rates of hydrolysis were measured with the indicated substrate, in the presence of 1 mM MnCl₂. The buffers used were: 100 mM Tris-acetate (pH 6-7), 100 mM Tris-HCl (pH 7-9) and 100 mM CAPS/NaOH (pH 9-10.1). Reaction mixtures of 0.4 ml were prepared and, at the end of the incubation period, 0.1 ml was used for the assay of activity, and 0.3 ml for pH mesurement with a glass electrode.

Figure 3 Protein alignment of rat ADPRibase-Mn orthologues showing the conservation of a metallophosphoesterase sequence signature

The sequences were aligned using the Clustal-W program. Orthologous proteins (with Genbank accession numbers): Rnor, *Rattus norvegicus* (conceptual translation of EU037900); Hsap, *Homo sapiens* (NP_064618); Mmu, *Mus musculus* (AAH04029); Xlae, *Xenopus laevis* (NP_001080428); Drer, *Danio rerio* (NP_956715); Atha, *Arabidopsis thaliana* (NP_194204); Osat1 and Osat2, *Oryza sativa* (NP_001060698 and NP_001062482); Oluc, *Ostreococcus lucimarinus* (green algae) (XP_001419532). The asterisks and the points below the sequences mark residues conserved as identities or similarities, respectively. The white letters on dark background represent conserved amino acids that are part of the metallophosphoesterase sequence signature divided in five regions (I-V): DX[H/X]-(X)n-GDXX[D/X]-(X)n-GNH[D/E]-(X)n-[G/X]H-(X)n-GHX[H/X] [32-35]. The vertical marks at the top of the sequences are positioned at intervals of ten amino acids relative to the *Rattus norvegicus* sequence.

Figure 4 Theoretical model of rat ADPRibase-Mn with docked ADP-ribose

The figure shows different views of the lowest-energy pose found after docking as described in the Experimental section. The two Zn²⁺ ions are shown as yellow spheres and ADP-ribose is shown in blue (adenine), red (pyrophosphate) and green (ribose-1 and ribose-2, respectively proximal and distal to the adenine). Oxygen atoms of the β -phosphate group interact with Zn1 and Zn2. (A) Full homology model showing the $\alpha/\beta/\beta/\alpha$ fold, typical of metallophosphoesterases, and the docked ADP-ribose. (B) Molecular surface of the active centre with ADP-ribose, as viewed from above in A. (C) Stereo view of a selection of amino acids that form the dimetallic centre and/or might interact with the substrate, as viewed from the right-hand side in A. Possible interactions of amino acid side chains in C are as follows. (i) Coordination with Zn1: Asp25, Gln 27, Asp74 and His280. (ii) Coordination with Zn2: Asp74, Asn110, His241 and His278. (iii) H bridge with O-P β : Gln 27, His278 and His280. (iv) H bridge with O-P α : Arg43 and His111. (v) H bridge with the phosphoanhydride O: Asn110 and His111. (vi) Hydrophobic contact with the adenine rings: Leu196. (vii) Contacts with ribose-1 or ribose-2 rings: Phe210 and Cys253, respectively. All the mentioned amino acids are

sequentially conserved (Figure 3) and superimposable to those of the zebrafish protein (not shown), except Leu196 that is conserved but not superimposable in both proteins.

Figure 5 The rat ADPRibase-Mn gene and orthologues in mouse and human genomes

The exon(black boxes)-intron(thin lines) structures are shown as they currently appear in the NCBI Gene database, including the ORFs coding for rat ADPRibase-Mn and its mouse and human orthologues after splicing. The exons of the rat gene are currently supported by a 1.3 kb mRNA (BC088174) which agrees with the size of the most abundant transcript seen in a northern blot (Figure 6). The white prolongations of the first translated exon show how the mRNAs would remain after operation of the alternative 5'-splice in the next intron (see the text). The alternative splicing would give rise to proteins shorter than ADPRibase-Mn and its orthologues, due to an in-frame stop codon either in the portion of intron retained as exon (mouse and human) or in the next exon (rat).

Figure 6 Expression of RGD1309906 transcripts in rat tissues

A multiple rat-tissue northern blot is shown after hybridisation with digoxineninlabelled ADPRibase-Mn cDNA. The figure shows a 3-h exposure of X-ray film to the chemiluminescent membrane, made 19-h after starting the detection reaction. Numbers below the lanes represent the quantitation (as percentages of the thymus signal) of the 1.4-kb band by image analysis during the first 3 h of detection.

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Figure 2

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Figure 3

Pnor	1		63
Hsan	1	MDRFDF35FADGFDF15FGVIADIQIADLEDGINFQK3KRKCIKNSVNLQGALEDWINES	63
Mmus	1		63
Xlae	1	MFNLCYKGQKLQRIVQSKRMEEPYFTFGIIADIQYADKDNRLNYLKTSMRYYRNSLTQLKVAIKEWATES	70
Drer	1	MEDPVFTFGLIADVQYADIEDGENYLRTRRRYYRGSADLLRDAVLQWRRE	50
Atha	1	MGSAA-RQPLFSFGVIA <mark>D</mark> VQYADISDGRSFLGVP-RYYRNSILVLQRAVETWNQH	53
Osat1	1	MMAVTNGVIHASS-REPLFSFGVIA <mark>D</mark> VQYADIPDGRSFLGVP-RYYRHSISVLQRAVSTWNKQ	61
Osat2	1	MLVQPPNAVAVRATAAGKEPLFSFGVIADVQYADIPDGRSFLGVP-RYYRHSLTILQRAVRSWNAHNHKHPVAG	73
Oluc	Ţ	SREPHFRFGLLADIQUCUGDDDRKNITGTQMRRYRNTLNVAERAVEYFNTH	50
Rnor	64	SMPCCVLQL CD IIDGYNAQYKVSEKSLELVMNTFQMLK-VPVHHTW <mark>GNHE</mark> FYNFSRDYLANSKLNSKF	130
Hsap	64	SMPCCVLQLGDIIDGYNAQYNASKKSLELVMDMFKRLK-VPVHHTWGNHEFYNFSREYLTHSKLNTKF	130
Mmus	64	SMPCCVLQLCDIIDGYNAQYKVSEKSLELVMNTFQMLK-VPVHHTWCNHEFYNFSRDYLASSKLNSKF	130
Xlae	71	IRFFILQLCDI IQCVNTKDNSSTIALERVLEENDKLP-IQFHHVWGN. FYNFSREYLNGSKLNSRA	137
Drer	51	-RVQCVVQLGDIIDGHNKRKDASDRALDTVMAELDACS-VDVHHVWCNHBFYNFSRPSLLSSRLNSAQ	110 110
ALNA Osat1	54	GNLRFVINMEDIVDGFCFR-DQSLAAIAALVHEERAFN-GFVIHMIGNE LINLFRE-ELLFLLAFG	126
Osat2	74	GGVERCVNFGDTVDGFCPK-DRSLAAVOAVAEFDEFH-GPAYHMLGNFCLYNLPRS-KLVSLLRMP	137
Oluc	51	-ECDFLVHNGDITDHQCAFDFASDDFKPIERGREDLGAVMRILSRAKCAEWLFTVCNHELYNFTREELRQGVASPGN	126
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Rnor	131	LEDQIPQHPETTPSENYYAYHFVPFPKFRFILLDSYDLSVLGIDQFSPKYEQCMKILREHNPN-VELNSPQG	201
Hsap	131	LEDQIVHHPETMPSEDYYAYHFVPFPKFRFILLDAYDLSV_GVDQSSPKYEQCMKILREHNPN-TELNSPQG	201
Mmus	131	LEDQIAQHPETTPSENYAYHFVPFPKFRILIDSVLGIDPSSPKYEQCMKLKENPPN-VELNSPQG	201
Dror	117	KENKI DQVVGISESGEINDESFIAINTSFFARKF STOGISSTIGKEAISLAIDISTNLAKENNEN-EDLNSFIG	186
Atha	119	RDGNAYYDFSPTPEYRVVLDGYDI SAVGWPOEHPNTIAALKILEEKNPN-TDKNSPAG	176
Osat1	127	DSDRAYYDFSPCPEYRFVVLDAYDFSALGWPRDHPVTAEAMKFLEEKNPN-SDKNSPDG	184
Osat2	138	SGRAYYDFSPFPGYRFVVLDAYDFSAVGWPRGHPVAAAARRFLDERNPN-ADKNSPSG	194
Oluc	127	S LNLKCSN ADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN	192
Oluc	127	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN	192
Oluc	127	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN	192
Oluc	127	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN	192
Oluc Rnor	202	SLNLKCSNADGEFYYSFKAGFGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270
Oluc Rnor Hsap	127 202 202	SLNLKCSNADGEFYYSFKAGFGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270
Oluc Rnor Hsap Mmus	127 202 202 202	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270
Oluc Rnor Hsap Mmus Xlae	127 202 202 202 214	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282
Oluc Rnor Hsap Mmus Xlae Drer	127 202 202 202 214 187	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282 257
Oluc Rnor Hsap Mmus Xlae Drer Atha	127 202 202 214 187 177	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282 257 247
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1	127 202 202 214 187 177 185	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282 257 247 255
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	127 202 202 214 187 177 185 195	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282 257 247 255 265 265
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	127 202 202 214 187 177 185 195 193	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	127 202 202 214 187 177 185 195 193	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282 257 247 255 265 269
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	127 202 202 214 187 177 185 195 193	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 270 282 257 247 255 265 269
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	127 202 202 214 187 177 185 195 193	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269
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Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap	127 202 202 214 187 177 185 195 193 271 271	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269 337 342
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlao	1277 2022 2022 2144 1877 1955 1933 2711 2711 2711 2712 2713	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269 337 342 340
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer	1277 2022 2022 2144 1877 1955 1933 2711 2711 2711 2711 2713 2783 258	SADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 342 340 356 322
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha	1277 2022 2022 2144 1877 1955 1933 2711 2711 2711 2711 2732 258 248	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN *** *********************************	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1	1277 2022 2022 2144 1877 1955 1933 2711 2711 2711 2713 258 2588 2586	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311 321
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2	1277 2022 2022 2144 1877 1955 1933 2711 2711 2711 2711 2713 258 258 258 258 2566	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN * * * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311 321 339
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 2144 1877 1855 1933 2711 2711 2711 2732 258 248 2566 2566 270	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN *** * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 2144 1877 1855 1933 2711 2711 2711 2732 258 248 2566 2566 2700	SLNLKCSNADGEFYYSFKAGPGWRVMVLDPYAVSIYRKGRQQGLCEAVELLRKHNPN-VDLFVKEN *** * * * * * * * * * * * * * * * * *	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 2144 1877 1855 1933 2711 2711 2731 2731 2732 258 2582 2582 2582 2582 2582 2582 258	SLNLKCSNADGEFYYSFKAGPGWPVWUDPYAVSIYRKGRQQGLCEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 2144 1877 1855 1933 2711 2711 2711 2711 2713 2588 2488 2566 2566 2700	SLNLKCSNADGEFYYSFKAGIGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 342 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 2144 1877 1855 1933 2711 2711 2711 2711 2711 2712 258 248 2566 2566 270	SLNLKCSNADGEFYYSFKAGIGWRVMVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 214 1877 1855 1933 2711 2711 2711 2711 2713 258 248 2566 2566 270	SLNLKCSNADGEFYYSFKAGPGWPUVLDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 214 1877 1955 1933 2711 2711 2711 2711 2833 2588 2488 2566 2666 270	SLNLKCSNADGEFYYSFKAGPGWP WWUDPYAVSIYRKGRQQGLCEEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 340 356 322 311 321 339 331
Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc Rnor Hsap Mmus Xlae Drer Atha Osat1 Osat2 Oluc	1277 2022 2022 214 1877 1955 1933 2711 2711 2711 2711 2833 2588 2488 2566 2666 270	SLNLKCSNADGEFYYSFKAGPGWRVJVUDPYAVSIYRKGRQQGLCEAVELLRKHNPN-VDLFVKEN ************************************	192 270 270 282 257 247 255 265 269 337 340 356 322 311 321 339 331

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Figure 6

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