Cloning and characterisation of hAps1 and hAps2, human diadenosine polyphosphate-metabolising Nudix hydrolases

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Abstract

Background: The human genome contains at least 18 genes for Nudix hydrolase enzymes. Many have similar functions to one another. In order to understand their roles in cell physiology, these proteins must be characterised.

Results: We have characterised two novel human gene products, hAps1, encoded by the NUDT11 gene, and hAps2, encoded by the NUDT10 gene. These cytoplasmic proteins are members of the DIPP subfamily of Nudix hydrolases, and differ from each other by a single amino acid. Both metabolise diadenosine-polyphosphates and, weakly, diphosphoinositol polyphosphates. An apparent polymorphism of hAps1 has also been identified, which leads to the point mutation S39N. This has also been characterised. The favoured nucleotides were diadenosine 5',5''-pentaphosphate (kcat/Km = 11, 8 and 16 x 10³ M⁻¹ s⁻¹ respectively for hAps1, hAps1-39N and hAps2) and diadenosine 5',5''-hexaphosphate (kcat/Km = 13, 14 and 11 x 10³ M⁻¹ s⁻¹ respectively for hAps1, hAps1-39N and hAps2). Both hAps1 and hAps2 had pH optima of 8.5 and an absolute requirement for divalent cations, with manganese (II) being favoured. Magnesium was not able to activate the enzymes. Therefore, these enzymes could be acutely regulated by manganese fluxes within the cell.

Conclusions: Recent gene duplication has generated the two Nudix genes, NUDT11 and NUDT10. We have characterised their gene products as the closely related Nudix hydrolases, hAps1 and hAps2. These two gene products complement the activity of previously described members of the DIPP family, and reinforce the concept that Ap5A and Ap6A act as intracellular messengers.

Background

In addition to the canonical ribonucleoside and deoxyri- bonucleoside phosphates and cofactors, cells contain a large number of minor nucleotides. Among these are the diadenosine polyphosphates (ApnA, where n = 2–7 [1]). Ap3A and Ap4A are the most intensively studied of these and are generally present in the soluble fraction of eukaryotic and prokaryotic cells at concentrations between 10 nM and 5 µM [2]. Platelet dense granules, adrenal chromaffin granules and certain synaptic vesicles have been reported to contain high concentrations of Ap5A and Ap6A in addition to Ap3A and Ap4A, all of which can be exocytosed following appropriate stimuli and bind to target cell purinoceptors causing a variety of physiological responses in the cardiovascular and central and peripheral nervous systems [1,3–5]. However, although Ap6A has been detected in erythrocytes [6], there are no substantiated measurements of Ap5A and Ap6A in the soluble fraction of nucleated cells, and it is likely that they are typically present at concentrations much lower than those of Ap3A.
and Ap$_3$A. The estimated 7 µM Ap$_3$A reported in cardiac muscle [7] may also be confined to granules.

Whilst the extracellular diadenosine polyphosphates have partially characterised signalling properties, the possible functions of the soluble, intracellular compounds remain unclear. On the one hand, they may simply be by-products of several enzymatic reactions (e.g. those catalysed by aminoacyl-tRNA synthetases and other ligases [8]). On the other, they may have one or more of a number of important regulatory roles, including involvements in DNA replication/repair, metabolic stress responses, the determination of cellular fate, and the regulation of enzyme activities and ion channels (see [1]). For example, Ap$_5$A is a potent inhibitor of adenylate kinase [9].

It is clear that cells have a variety of relatively specific enzymes able to degrade these compounds. These include symmetrically-cleaving Ap$_4$A hydrolases (in prokaryotes), members of the histidine triad protein family such as the FHT tumour suppressor protein and (in lower eukaryotes) the related Ap$_4$A phosphorylases, and most widespread of all, members of the Nudix hydrolase family (see [10,11] for reviews). Nudix hydrolases cleave predominantly the diphosphate linkage in compounds of general structure: nucleoside diphosphate linked to another moiety, $\chi$. Some family members have a relatively broad substrate specificity, while others have a much more restricted range. They all possess the Nudix sequence signature motif Gx$_3$Ex$_5$ [UA]xResEx$_3$Ex$_2$Gx (where U is an aliphatic hydrophobic amino acid), which represents the catalytic site of the enzyme. Animal and plant Ap$_4$A hydrolases (EC 3.1.6.17) degrade Ap$_4$A, Ap$_5$A and Ap$_6$A, always producing ATP as one of the products. They are most active towards Ap$_4$A; however, the related enzymes from Escherichia coli [12] and Rickettsia prowazekii [13] show a marked preference for Ap$_5$A.

More recently, distinct members of the Nudix family that prefer Ap$_6$A have been purified and characterised. The three human enzymes termed diphosphoinositol polyphosphate phosphohydrolases 1 [14,15], 2α and 2β [16] (hDIPP-1, -2α and -2β) as well as Schizosaccharomyces pombe Aps1 [17] and Saccharomyces cerevisiae Ddp1p [18] all favour Ap$_6$A over Ap$_5$A. Activity with Ap$_6$A is lower, and is in fact lacking in the case of Ddp1p. Remarkably these Ap$_6$A hydrolases are also able to utilise the structurally unrelated non-nucleotide substrates, diphosphoinositol pentakishosphate (PP$_3$InsP$_5$) and bisdiphosphoinositol tetrakisphosphate ([PP]$_2$InsP$_4$), hence they are commonly referred to as the DIPP subfamily. These substrates are generally favoured in vitro, with $k_{cat}/k_m$ ratios being 50–500-fold higher for PP-InsP$_5$ compared to Ap$_6$A, except for DIPP-2β, which has a significantly reduced relative activity with PP-InsP$_5$ compared to the others. The intracellular levels of diphosphoinositol polyphosphates appear to be lower than those of the nucleotide substrates. PP-InsP$_5$ levels are often between 1 [19] and 3% [20,21] of the levels of InsP$_6$, which has been estimated as being between 15 and 100 µM (see [22]) and [PP]$_2$-InsP$_4$ levels are an order of magnitude lower [20,21], and as such are virtually undetectable in many cell types. PP-InsP$_5$ turnover is regulated by the tumour promoter thapsigargin [19], while [PP]$_2$-InsP$_4$ turnover is regulated by cyclic AMP- and cyclic GMP-dependent processes via an undefined mechanism [23]. However, as for the Ap$_n$As, no clear functions have yet been found for the diphosphoinositol polyphosphates, although they have been implicated in vesicle trafficking [24], apoptosis [25] and DNA repair [26].

We describe here two further members of the human DIPP subfamily with novel properties. These proteins, hAps1 and hAps2, products of the NUDT11 and NUDT10 genes respectively, closely resemble the other DIPPs in primary structure, but show a selectivity towards Ap$_4$A and Ap$_6$A and reduced activity towards PP-InsP$_5$ and [PP]$_2$-InsP$_4$. They also display a novel pattern of tissue-specific expression.

**Results**

**Sequence alignment**

An initial BLAST search of the GenBank expressed-sequence tag (EST) database with the hDIPP-1 sequence identified two closely related but previously uncharacterised predicted proteins, which we have called hAps1 and hAps2 (Human ApsXh hydrolases 1 and 2). Alignment of the cDNA sequences with the human genome indicated that the genomic sequences encoding these proteins, FLJ10628 and LOC139770, lie about 150 kb apart on the X chromosome at Xp11.23 and are transcribed in opposite directions (Fig. 1). In accordance with the guidelines for the Nudix protein family, the genes for hAps1 and hAps2 have been designated NUDT11 and NUDT10 respectively by the HUGO Gene Nomenclature Committee (see [http://www.gene.ucl.ac.uk/nomenclature/gene-family/npym.html](http://www.gene.ucl.ac.uk/nomenclature/gene-family/npym.html)). These proteins are each 164 amino acids long and are most similar to hDIPP-2β, hAps1 showing 76, 90 and 91% identity at the amino acid level to hDIPP-1, 2α and 2β, respectively. In particular, they both possess the additional Gln residue (Q86 in DIPP-2β) that distinguishes hDIPP-2β from hDIPP-2α and hDIPP-1 and which is responsible for the reduced activity of hDIPP-2β towards diphosphoinositol polyphosphates compared to the other DIPPs. hAps1 and hAps2 are also identical to the hDIPP-2 enzymes throughout the Nudix motif (Fig. 2). These observations led us to anticipate that hAps1 and hAps2 would be potent dinucleoside polyphosphate hydrolases with reduced activity towards diphosphoinositol polyphosphates. The existence of two such similar ex-
pressed gene products was surprising. Comparison of the open reading frames (ORFs) of hAps1 and hAps2 revealed 6 differences at the nucleotide level, leading to five silent changes (T8, E28, E69, D88 and stop), and an Arg-Pro substitution at position 89. A comparison of the cDNAs for the two gene products showed further divergence in the 3' untranslated region. It appears that the evolution of these closely related genes occurred recently, as hAps1 and hAps2 show greater similarity with each other than with the Aps-like sequences from mouse (of which two genes, XM_135784 and XM_135786 appear to yield the same protein product) or cow (Fig 3). Further examination of all available hAps1 ESTs revealed a sequence divergence, leading to a point mutation S39N, in the coding region. This is present in two ESTs (Table 1) and was detected in 10% (3 from 30) of polymerase chain reaction (PCR) products generated from a range of normal human cDNA samples from diverse sources. In all other respects investigated, including the 3' untranslated region, it appeared to be identical to hAps1. This would suggest that this minor variant, hAps1-39N, represents a polymorphism present in the human population. Its properties were also examined in this study.

**Purification and properties of hAps1 and hAps2**

The ORFs for hAps1, hAps1-39N and hAps2 were cloned into the pGEX6P-1 expression vector and expressed in Escherichia coli (E. coli). After incubation with isopropyl β-D-thiogalactoside (IPTG) (100 µM) at 26°C for 8 hours, a major protein of the expected size was induced and expressed in a soluble form in each case. The glutathione S-transferase (GST)-tagged recombinant proteins were purified by chromatography on Glutathione Sepharose 4 Fast Flow resin (Fig. 4).

Initially, the ability of hAps1 and hAps2 to metabolise Ap5A was determined at 37°C in the presence of 50 mM HEPES, pH 7.6 and 100 µM substrate. A divalent cation was essential for activity, with Mn²⁺ by far the most effective between 2 and 6 mM (Fig. 5a). Cu²⁺ supported less than 30% and Zn²⁺ and Co²⁺ each less than 3% of the maximum activity. Ni²⁺, Ca²⁺ and, surprisingly, Mg²⁺ were unable to activate hAps1 or hAps2. When assayed in the presence of 1 mM MnCl₂, hAps1 and hAps2 showed alkaline pH optima of pH 8.5 (Fig. 5b), a feature common among Nudix hydrolases. Activity increased from undetectable at pH 6 to over 70% maximum between 7.5 and 9. Previous characterisation of hDIPP-1, 2α and 2β, as well as *S. pombe* Aps1 was performed in a buffer containing 1 mM MnCl₂ at pH 7.6. In order to compare results for hAps1 and hAps2 directly with hDIPP-1, hDIPP-2 and *S. pombe* Aps1, we have performed assays using identical conditions.

When hAps1, hAps1-39N and hAps2 were assayed with a wide range of potential substrates all three proteins were found, as expected, to metabolise Ap₅A and Ap₆A. Of the other substrates tested, only p₄A and Gp₄G showed significant rates of hydrolysis by all three, while Ap₄A and Gp₄G were also effective substrates for hAps2. All three enzymes showed a marked preference for adenine over guanine nucleotides. Table 2 shows rates of metabolism and products formed where identified. *Kₘ* and *kₐₐₙ* values for each enzyme with Ap₅A and Ap₆A are shown in Table 3. The most rapidly metabolised substrate appeared to be Ap₅A, although Ap₆A was bound with higher affinity.

Prolonged treatment of Ap₅A and Ap₆A identified the major products as adenosine 5'-monophosphate (AMP) plus...
were found to have little activity towards PP-InsP5 or hAps1, hAps2 and the DIPPs, hAps1 and hAps2. Despite the overall high degree of sequence similarity between hAps1, hAps2 and the DIPPs, hAps1 and hAps2 were found to have little activity towards PP-InsP5 or [PP]2-InsP4. hAps1 and hAps1-39N showed first order rate constants (k-1) with PP-InsP5 of 19 ± 4 and 7 ± 5 µg-1 min-1 respectively. Activity of hAps2 was similar to that of hAps1, with a k-1 = 13 ± 6 µg-1 min-1. These compare with k-1 values of 2200, 2000 and 220 µg-1 min-1 for hDIPP-1, -2α and -2β respectively [16]. Km and kcat values were determined for each of the proteins (Table 3) and these suggested that the reduction in activity towards PP-InsP5 compared with hDIPP-1 and hDIPP-2 was primarily due to a reduced affinity for this substrate. Activity towards [PP]2-InsP4 was even weaker. hAps1 and hAps2 gave k-1 = 1.0 ± 0.2 and 0.9 ± 0.2 µg-1 min-1 respectively. hAps1-39N showed no activity towards [PP]2-InsP4 under the conditions tested. In contrast, hDIPP-1, -2α and -2β gave k-1 values of 320, 90 and 32 µg-1 min-1 respectively, whereas S. pombe Aps1 and S. cerevisiae Ddp1p showed similar activity with k-1 = 8.3 and 1.7 µg-1 min-1 respectively [15].

**Tissue distribution**

Analysis of the tissue distribution of hAps1 and hAps2 gene expression was performed using PCR and human tissue cDNA samples. Using primers selective for hAps1,
Figure 3
Comparison of hAps1 with translated EST sequences from mouse and cow. No full-length bovine EST for the hAps1 homologue is available, hence the composite of two separate sequences is shown. A predicted rat protein from genomic sequencing is also shown.

Table 1: Tissue sources of known ESTs for hAps1 and hAps2

| Tissue               | hAps1                      | hAps2                      |
|----------------------|----------------------------|----------------------------|
| Brain                | 15434059*, hippocampus, hAps1 | 15936725, medulla           |
|                      | 15495588, hypothalamus, hAps1 | 13980266, hypothalamus      |
|                      |                            | 15743679, pooled brain      |
| Testis               | 14001386, hAps1             | 14083118                    |
| Breast               | 15996913, hAps1             |                            |
| Ovary                | 19353207, hAps1             |                            |
| Uterus               | 11970540, tumour cell line, hAps1 |                          |
| Eye                  | 19090579, hAps1             |                            |
| Teratocarcinoma      | 7022778, 10989803, hAps1-39N |                            |

*Numbers are GenBank gene identifiers
PCR products were readily visible from brain, pancreas and testis and less so from lung, thymus, prostate, ovary, small intestine and heart (the least) (Fig. 6). Selective primers for hAps2 produced strong signals from brain and liver, but also from heart, placenta, lung, kidney, pancreas, spleen, prostate, testis and ovary (Fig. 6). No signal for hAps1 was detected in placenta, liver, skeletal muscle, kidney, spleen, colon or peripheral blood leukocytes. Likewise, no signal for hAps2 was found in skeletal muscle, thymus, small intestine, colon or peripheral blood leukocytes. A number of tumour cell line cDNA samples were also tested; hAps1 was present in prostate PC3, and ovary GI-102 cells. The prostate PC3 cell line was also found to express hAps2. Tumour cell lines from breast GI-101, lung LX-1, colon CX-1, lung GI-117, colon GI-112 and pancreas GI-103 were found to express neither hAps1 nor hAps2. These results are in broad agreement with the sources of ESTs for these two protein families (Table 1). This list is for comparison and is not exhaustive. In contrast, hDIPP-1 was found in all the tissues and cell types tested above (S.T. Safrany, unpublished data).

Whilst an exhaustive analysis has not been performed for hDIPP-2, it was found to be highly expressed in heart, skeletal muscle, kidney and pancreas, and weakly expressed in brain, placenta lung and liver [16].

Subcellular localisation
The subcellular localisation of hAps1, hAps1-39N and hAps2 was investigated by individual expression of the three proteins tagged at the N-terminus with EGFP in HEK293 and PC12 cells, and visualisation of the expressed fusion proteins by deconvolved fluorescence microscopy. These results show that in both cell types EGFP-tagged proteins showed a cytosolic location indistinguishable from EGFP-hDIPP-2α, EGFP-hDIPP-2β and EGFP alone (Fig. 7a and 7b). Similar data were obtained with proteins tagged at the C-terminus with EGFP (data not shown). This is in agreement with localisation predicted using PSORTII.

Discussion
The human genome contains at least 18 Nudix genes, several of which encode multiple products, and 9 Nudix hydrolases have been characterised so far to varying degrees (see [http://www.gene.ucl.ac.uk/nomenclature/gene-family/npym.html]). These include hDIPP-1, hDIPP-2α and hDIPP-2β, encoded by the NUDT3 (hDIPP-1) and NUDT4 (hDIPP-2α and hDIPP-2β) genes, all of which hydrolyse both the higher order diadenosine polyphosphates and diphosphoinositol polyphosphates. In this report, we describe the cloning and functional characterisation of hAps1 and hAps2, two new members of the DIPP subfamily with a novel pattern of expression and substrate specificity. A polymorphism of hAps1 (hAps1-
39N) has also been found, its activity and localisation appearing broadly similar to those of hAps1. Unlike the other subfamily members, hAps1 and hAps2 show little activity towards diphosphoinositol polyphosphates, both gene products favouring diadenosine polyphosphates, Ap₆nA, where n = 5 or 6. In each case, Ap₆nA binds with higher affinity than Ap₅nA, whereas the kₗₐₜₜ for the latter is greater. Given that no other PP-InsP₃ hydrolase activity was observed during the purification of rat hepatic DIPP (most similar to hDIPP-1) and assuming that tissue expression and physiology are broadly conserved between human and rat, these results would suggest that Aps-like proteins do not contribute significantly to inositol phosphate metabolism.

Both hAps1 and hAps2 show typical Nudix hydrolase requirements, namely an alkaline pH optimum in vitro and a requirement for divalent cations [12]. An unusual feature of hAps1 and hAps2 is that they both require Mn²⁺ for activation, and that Mg²⁺ is without effect. Optimal [Mn²⁺] is approximately 4 mM, which is in contrast to that estimated for free intracellular [Mn²⁺] of below 0.5 μM [27]. Free [Mn²⁺] in our in vitro assays was not determined. It is well acknowledged that Mn²⁺ binds serum albumin very tightly, but despite this we would anticipate free [Mn²⁺] in the assays to be far in excess of intracellular levels. It is known that the activity of metal ion-requiring enzymes in different cellular compartments can be regulated through the controlled trafficking of these ions and therefore through the controlled access of these enzymes to their required ions. This principle has been most thoroughly studied in the case of Ca²⁺[28], although analogous observations have been made with Cu²⁺[29] and Mn²⁺[30]. Since the [Mn²⁺] throughout the cell is known to be non-uniform, the unusual requirement of the hAps enzymes suggests a possible mechanism of regulation through access to this ion.

Several biochemically distinct Ap₅/₆A hydrolases have been described previously. S. pombe Aps1, S. cerevisiae Ddp1p and hDIPP-1 and hDIPP-2 have all been shown to metabolise Ap₅A and Ap₆A. The activities of hAps1 and hAps2 towards Ap₅A and Ap₆A are comparable to hDIPP-1, but their activities towards the diphosphoinositol polyphosphates PP-InsP₅ and [PP]₂-InsP₄ are greatly reduced, thus making them some 100–300-fold more selective towards Ap₅A and Ap₆A than hDIPP-1 or hDIPP-2. S. pombe Aps1 and S. cerevisiae Ddp1p, the sole representatives of the DIPP subfamily in the two yeast genomes, appear to favour the diphosphoinositol polyphosphates. Their modes of action have been shown to differ from their mammalian counterparts. The major products of Ap₅A hydrolysis by S. pombe Aps1 and S. cerevisiae Ddp1p are ADP and p₄A [17,18]. As with hDIPP-1 [15] and hDIPP-2 (Safrany, data not shown), hAps1 and hAps2 produced AMP and p₅A as major products. With Ap₃A as substrate, S. pombe Aps1 favoured the production of ADP plus ATP, whereas S. cerevisiae Ddp1p favoured production of AMP and p₄A. In this respect hAps1 and hAps2 as well as hDIPP-1 and hDIPP-2 resemble Ddp1p.

Due to the chromosomal proximity of Aps1 and Aps2, and the greater degree of sequence identity with each other than with the Aps-like genes from mouse, rat or cow, it seems likely that this gene pair has arisen as the result of a recent gene duplication event on the short arm of the X chromosome [31]. A popular view of the evolution of catalytic motifs envisages the duplication of relatively non-specific progenitors followed by adaptation to perform special tasks, and the Nudix hydrolase motif appears to be a good illustration of this process. The existence of multiple forms of these enzymes would suggest that the diadenosine polyphosphates have significant biological functions, with the specialised task of hAps1 and/or hAps2 being the regulation of Ap₃A and Ap₅/A levels.

Conclusions

Previously, the human DIPP family comprised hDIPP-1, a ubiquitous enzyme able to hydrolyse both inositol phosphates and diadenosine polyphosphates, and hDIPP-2α and -2β, which are predominantly diphosphoinositol polyphosphate phosphohydrolases highly expressed in heart, skeletal muscle, kidney and pancreas. We have now identified two further members, hAps1 and hAps2, which have high selectivity towards the diadenosine polyphosphates. The distinction between DIPP and Ap₆A hydrolase activities reinforces the importance of Q86 in DIPP-2β and the role of Q85 in hAps1 and hAps2. The addition of this amino acid in hDIPP-2 has been previously shown to have a substantial negative effect upon its activity towards diphosphoinositol polyphosphates, whereas Ap₃A hydrolase activity was increased. The expression of hAps1 and hAps2 is also restricted. Of the tissues tested, only pancreas showed high levels of hDIPP-2 and hAps1, and only heart showed high levels of hDIPP-2 and hAps2, although ESTs were found suggesting that hDIPP-2 is expressed in testis, ovary and prostate. The presence of hAps1 and hAps2 in such a variety of tissues suggests that Ap₃A and Ap₅A may have roles other than as secreted molecules. We find that hAps1 is present in the pancreas, but there is no suggestion that it is secreted. For this reason, hAps1 and hAps2 are expected to control intracellular Ap₃A and Ap₅/A levels.

In conclusion, we have identified two human Nudix hydrolases that share a novel substrate specificity, metabolising predominantly Ap₃A and Ap₅/A. These two gene products complement the activity of previously described members of the DIPP family, and reinforce the concept of Ap₃A and Ap₅/A acting as intracellular messengers.
**Figure 5**  
**Properties of hAps1 and hAps2.**  
(a) Mn^{2+}-dependency of hAps1 (filled circles) and hAps2 (open circles) activity towards Ap5A (100 μM).  
(b) pH dependency of hAps1 (filled circles) and hAps2 (open circles) activity towards Ap5A (100 μM).
Authors' contributions
NRL performed the cloning and subcloning, determined subcellular localisation in mammalian cells, participated in sequence analysis and obtained the genomic alignment. AGMcL assisted in the writing of the manuscript. STS conceived of this study, prepared recombinant enzymes, performed enzyme assays, determined the tissue distribution, participated in sequence analysis and drafted the manuscript. All authors read and approved the final manuscript.

Materials and Methods

Materials
Ap$_4$G, Ap$_5$G and p$_4$A were purchased from Jena Bioscience, Jena, Germany. Other nucleotides were purchased from Sigma. Non-radioactive PP-InsP$_5$ was a kind gift from J.R. Falck (University of Texas Southwestern Medical Center, Dallas, TX, USA). $[^3]$H]-PP-InsP$_5$ and $[^3]$H]-[PP]$_2$-InsP$_4$ were prepared as previously described [15].

Cloning and plasmid construction
PCR products corresponding to the hAps1 sequence were amplified from a human testis library (Clontech), using the primers SSD019 (5'-gccagatatcgaattggaaccacgctgg-3') and SSD020 (5'-ggcagatatcgaattggaaccacgctgg-3') (BamHI sites are underlined). PCR was carried out using a HiFidelity Expand Kit (Roche), and the products cloned into the vector pCR2.1TOPO (Invitrogen) and fully re-sequenced.

Table 2: Nucleotide substrate utilisation by hAps1, hAps1-39N and hAps2

| Nucleotide | hAps1 | hAps1-39N | hAps2 | Major products |
|------------|-------|-----------|-------|----------------|
| Ap$_6$A    | 0.25 ± 0.04 | 0.19 ± 0.01 | 0.28 ± 0.04 | AMP + p$_3$A |
| Ap$_5$A    | 0.41 ± 0.05 | 0.37 ± 0.02 | 0.9 ± 0.1 | AMP + p$_4$A |
| Ap$_4$A    | 0.02 ± 0.01 | 0.02 ± 0.01 | 0.09 ± 0.03 | AMP + ATP |
| Ap$_3$A    | 0       | 0         | 0     |                |
| Ap$_2$A    | 0       | 0         | 0     |                |
| Gp$_5$G    | 0.04 ± 0.02 | 0.02 ± 0.01 | 0     |                |
| Gp$_4$G    | 0.04 ± 0.02 | 0.04 ± 0.02 | 0.11 ± 0.04 | GMP, GDP, GTP |
| Gp$_3$G    | 0       | 0         | 0     |                |
| Gp$_2$G    | 0       | 0         | 0     |                |
| Ap$_5$G    | 0       | 0         | 0     |                |
| Ap$_4$G    | 0       | 0         | 0     |                |
| p$_4$A     | 0.07 ± 0.03 | 0.10 ± 0.03 | 0.13 ± 0.04 | ATP |
| ATP        | 0       | 0.03 ± 0.02 | 0     |                |
| ADP-ribose | 0       | 0         | 0     |                |
| ADP-glucose| 0       | 0         | 0     |                |
| dATP       | 0       | 0.07 ± 0.04 | 0     |                |
| p$_4$G     | 0.15 ± 0.05 | 0.21 ± 0.05 | 0.21 ± 0.03 | GTP |
| GTP        | 0.02 ± 0.01 | 0.03 ± 0.01 | 0     |                |
| GDP-glucose| 0.01 ± 0 | 0         | 0     |                |
| GDP-mannose| 0       | 0         | 0     |                |
| dGTP       | 0.05 ± 0.03 | 0.05 ± 0.03 | 0     |                |
| dUTP       | 0.04 ± 0.03 | 0.05 ± 0.02 | 0     |                |
| dTTP       | 0.05 ± 0.04 | 0.05 ± 0.03 | 0     |                |
| CTP        | 0.03 ± 0.02 | 0.03 ± 0.02 | 0     |                |
Table 3: Kinetic parameters of hAps1, hAps1-39N and hAps2

| Enzyme    | Substrate | $K_m$ (µM) | $k_{cat}$ (s⁻¹) | $k_{cat}/K_m$ (10³M⁻¹s⁻¹) |
|-----------|-----------|------------|----------------|---------------------------|
| hAps1     | Ap₂A      | 13 ± 5     | 0.17 ± 0.04     | 13.1                      |
|           | Ap₃A      | 37 ± 9     | 0.4 ± 0.1       | 10.8                      |
|           | PP-InsP₅ (n = 2) | 4    | 0.20           | 50.0                      |
| hAps1-39N | Ap₂A      | 11 ± 3     | 0.16 ± 0.02     | 14.5                      |
|           | Ap₃A      | 50 ± 10    | 0.4 ± 0.1       | 8.0                       |
|           | PP-InsP₅ (n = 2) | 0.3  | 0.09           | 290.0                     |
| hAps2     | Ap₂A      | 19 ± 1     | 0.20 ± 0.01     | 10.5                      |
|           | Ap₃A      | 50 ± 10    | 0.8 ± 0.2       | 16.0                      |
|           | PP-InsP₅  | 1.3 ± 0.6  | 0.14 ± 0.02     | 110.0                     |

Figure 6
Tissue distribution of hAps1, hAps2 and β-actin. PCR using human tissue first-strand cDNA as template was performed with primers specific for β-actin and for hAps1 or hAps2. 35 cycles of amplification were used for β-actin, 40 cycles for hAps1 and hAps2.
The 3' nucleotide of SSD019 introduced a silent mutation into hAps2, and was allowed to remain. Likewise, SSD020 also introduced a silent mutation into the stop codon of hAps2. This was also allowed to remain. Vectors for the expression of Nudix hydrolase proteins fused to the C-terminus of Enhanced Green Fluorescent Protein (EGFP) were produced by ligating a cDNA encoding each protein as a BamHI digestion fragment from the GST-fusion vectors described above, into pEGFP-C1 (Clontech) previously digested with BglII and BamHI.

**Bioinformatics**

Multiple sequence alignments were performed using MAP and BoxShade, both run by EMBnet, Lausanne, Switzer-

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Figure 7

Subcellular localisation of EGFP, EGFP-hDIPP-2 (α and β) and hAps1 and hAps2. Proteins fused to the C-terminus of EGFP were expressed in (a) HEK293 or (b) PC12 cells for 24 h and visualised by fluorescence microscopy.
land. Blast searches of EST and genomic databases were performed using the facilities provided by NCBI, Bethesda, MD, USA.

**Expression and purification of hAps1, hAps1-39N and hAps2**

Full-length hAps1, hAps1-39N and hAps2 were expressed and purified from *E. coli* as GST-fusion proteins. Expression plasmids based on pGEX-6P1 were transformed into *E. coli* strain BL21, and induced at 26°C overnight with 100 µM IPTG. Cells were harvested in buffer A (20 mM Tris, 150 mM NaCl, 2 mM DTT, 0.1 mM EGTA, pH 7.5), supplemented with 5 µg/ml leupeptin and 1 µg/ml aprotinin, and disrupted by sonication (3 × 15s). Particulate matter was removed by centrifugation. The supernatant was applied to (and subsequently eluted from) a 5 mm × 5 cm Glutathione Sepharose 4 Fast Flow column (Amersham Pharmacia Biotech) at a flow rate of 1 ml/min. The column was washed for 5 min with buffer A. Bound protein was eluted with a gradient generated by mixing buffer A with buffer B (buffer A plus 25 mM glutathione, pH 7.5) as follows: 0-10 min, 0-50% buffer B; 10-13 min, 100% buffer B. Fractions containing pure hAps1, hAps1-39N or hAps2 were supplemented with glycerol (10% v/v final), and stored at -80°C.

**Tissue distribution**

The tissue distribution of hAps1 and hAps2 expression was determined by PCR from human tissue cDNA samples (human cDNA panels, Clontech) and human cDNA libraries (testis from Clontech, brain from Dr Peter Cheung of the MRC Protein Phosphorylation Unit, Wellcome Trust Biocentre, Dundee, Scotland, UK). β-Actin-specific and hAps1 and hAps2-specific primers were used. The hAps1 primers were SSD049 (5’-cgctctgcagacacagcgtc-3’), and SSD045 (5’-caaaagccacacatggtgcc-3’) and the hAps2 primers were SSD050 (5’-gctctgcagacacagcgtc-3’) and SSD046 (5’-gctgcaaacctgagaatgcttga-3’). Primers SSD045 and SSD046 were selective for the 3’ untranslated regions of hAps1 and hAps2 respectively. β-Actin cDNA was amplified using β AS (5’-acagtctgcctctgaggg-3’) and β AA (5’-cctctgcctcgtctgcaatg-3’). PCR was carried out using a HiFidelity Expand Kit (Roche), and cDNA samples from heart, brain, placenta, lung, liver, skeletal muscle, kidney, pancreas (human MTC panel I #K1420-1, Clontech), spleen, thymus, prostate, testis, ovary, small intestine, colon, peripheral blood leukocytes (human MTC Panel II #K1421-1, Clontech). The following tumour cell lines were also tested: breast GI-101, lung LX-1, colon CX-1, lung GI-117, prostate PC3, colon GI-112, ovary GI-102 and pancreas GI-103 (human MTC Tumor Panel #K1422-1, Clontech).

**Subcellular localisation**

The subcellular localisation of hAps1, hAps1-39N and hAps2 proteins fused to EGFP was investigated using transfected HEK 293 epithelial cells and PC12 pheochromocytoma cells. PC12 cells were grown in DMEM/5% foetal bovine serum/5% horse serum. HEK293 were grown in DMEM/10% foetal bovine serum. Both cell types were grown on coverslips and transfected using Fugene-6 (Roche) with expression vectors for each Nudix hydrolase fused to the C-terminus of EGFP. 24 hours after transfection, cells were observed using a Leica inverted stage fluorescence microscope and a Hamamatsu Orca charge-coupled-device camera. Images were analysed using Improvision OpenLab deconvolution software.

**Enzyme assays**

For the analysis of nucleotide metabolism, assays were performed as previously described [15], unless otherwise stated. The UV-absorbing (260 nm) nucleotide reaction products were analysed using a 4.6 × 125 mm Partisphere 5 µm SAX HPLC column. Substrates and products were eluted at 1 ml/min by the following gradient generated by mixing water with buffer C (1.3 M (NH4)2HPO4, pH 4.8 with H3PO4): 0-5 min, 0% buffer C; 5-55 min, 0-50% buffer C; 56-70 min, 0% buffer C. pH optima of recombinant enzymes were determined in a series of buffers comprising 1 mM MnCl2, 100 µg/ml BSA and 15 mM MES, 15 mM HEPES or 15 mM Tris (pH 6-9 with HCl or NaOH). Divalent ion-dependency was determined in a buffer comprising 50 mM HEPES, 100 µg/ml BSA, pH 7.6, and 50 µM-10mM (where possible) divalent metal as the chloride salt. Inositol phosphate metabolism was determined exactly as described previously [14]. Km and kat values were determined by hyperbolic regression analysis.

**Authors’ note**

During the reviewing of this manuscript a paper was submitted and accepted in the Journal of Biological Chemistry (K Hidaka, JJ Caffrey, L Hua, T Zhang, JR Falck, GC Nickel, L Carrel, LD Barnes, SB Shears: An adjacent pair of human NudT genes on chromosome X are preferentially expressed in testis and encode two new isoforms of di-phosphoinositol polyphosphate phosphohydrolase. J Biol Chem in press). DIPP-3β corresponds to hAps1, whereas DIPP-3α corresponds to hAps2.

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