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A P2X receptor from the tardigrade species *Hypsibius dujardini* with fast kinetics and sensitivity to zinc and copper

Selvan Bavan¹, Volko A Straub¹, Mark L Blaxter² and Steven J Ennion*¹

Address: ¹Department of Cell Physiology and Pharmacology, University of Leicester, PO Box 138, Leicester, LE1 9HN, UK and ²Institute of Evolutionary Biology, The Ashworth Laboratories, University of Edinburgh, Edinburgh, UK

Email: Selvan Bavan - sb314@le.ac.uk; Volko A Straub - vs64@le.ac.uk; Mark L Blaxter - mark.blaxter@ed.ac.uk; Steven J Ennion* - se15@le.ac.uk

* Corresponding author

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**Abstract**

**Background:** Orthologs of the vertebrate ATP gated P2X channels have been identified in *Dictyostelium* and green algae, demonstrating that the emergence of ionotropic purinergic signalling was an early event in eukaryotic evolution. However, the genomes of a number of animals including *Drosophila melanogaster* and *Caenorhabditis elegans*, both members of the Ecdysozoa superphylum, lack P2X-like proteins, whilst other species such as the flatworm *Schistosoma mansoni* have P2X proteins making it unclear as to what stages in evolution P2X receptors were lost. Here we describe the functional characterisation of a P2X receptor (*HdP2X*) from the tardigrade *Hypsibius dujardini* demonstrating that purinergic signalling is preserved in some ecdysozoa.

**Results:** ATP (EC₅₀ ~ 44.5 μM) evoked transient inward currents in *HdP2X* with millisecond rates of activation and desensitisation. *HdP2X* is antagonised by pyridoxal-phosphate-6-azophenyl-2',4'disulfonic acid (IC₅₀ 15.0 μM) and suramin (IC₅₀ 22.6 μM) and zinc and copper inhibit ATP-evoked currents with IC₅₀ values of 62.8 μM and 19.9 μM respectively. Site-directed mutagenesis showed that unlike vertebrate P2X receptors, extracellular histidines do not play a major role in coordinating metal binding in *HdP2X*. However, H306 was identified as playing a minor role in the actions of copper but not zinc. Ivermectin potentiated responses to ATP with no effect on the rates of current activation or decay.

**Conclusion:** The presence of a P2X receptor in a tardigrade species suggests that both nematodes and arthropods lost their P2X genes independently, as both traditional and molecular phylogenies place the divergence between Nematoda and Arthropoda before their divergence from Tardigrada. The phylogenetic analysis performed in our study also clearly demonstrates that the emergence of the family of seven P2X channels in human and other mammalian species was a relatively recent evolutionary event that occurred subsequent to the split between vertebrates and invertebrates. Furthermore, several characteristics of *HdP2X* including fast kinetics with low ATP sensitivity, potentiation by ivermectin in a channel with fast kinetics and distinct copper and zinc binding sites not dependent on histidines make *HdP2X* a useful model for comparative structure-function studies allowing a better understanding of P2X receptors in higher organisms.
Background

Vertebrate P2X receptors comprise a family of ligand gated ion channels activated by extracellular ATP [1]. They form homo or heteromeric trimers with each monomer consisting of intracellular amino and carboxy termini, two transmembrane domains and a large glycosylated extracellular region containing five disulphide bonds [2] and the agonist binding site [3]. Mammalian species possess seven distinct P2X subtypes (P2X1–7) that play important roles in a wide range of physiological processes including neurotransmission, platelet aggregation, smooth muscle contractility and bone formation [4,5]. Many studies have also described potential roles for ATP as an extracellular signalling molecule in a range of invertebrate phyla [6] and plants [7,8] leading to the assumption that ATP is a primitive signalling molecule and that the emergence of purinergic receptors occurred relatively early in evolution [9]. This assumption is supported by the definitive molecular and functional identification of P2X receptors in the slime mould Dictyostelium discoideum [10], the green alga Ostreococcus tauri [11] and the choanoflagellate Monosiga brevicollis [11]. However, several non-vertebrate organisms for which full genome data are available lack P2X-like genes, including Anopheles gambiae, Caenorhabditis elegans, Drosophila melanogaster, and Apis mellifera. Arthropods and nematodes are members of the protostome superphylum Ecdysozoa. Given that P2X receptors are present in choanoflagellates, believed to be the sister group to the Metazoa, and in some representatives of the protostome superphylum Lophotrochozoa including Schistosoma mansoni [12] and Lymnaea stagnalis [13], the absence of P2X receptors from the above fully sequenced nematode and arthropod genomes suggests a loss of this class of gene in an ancestor of the Ecdysozoa.

We identified a partial P2X-like sequence in expressed sequence tag (EST) data from the tardigrade Hypsibius dujardini. Tardigrades are microscopic animals around 200 to 500 μm in length that inhabit both marine and fresh water habitats [14,15]. They possess a fascinating ability to desiccate into a reversible state of metabolic suspension called cryptobiosis allowing them to survive for many decades in harsh environments, such as lack of oxygen, extremes in temperature, and high pressure, before rehydration to an active state within minutes [16]. Tardigrades are placed in their own phylum, Tardigrada and share features with both arthropods and nematodes. Like arthropods they possess legs and a distinctly segmented body. However they also have a triradiate pharynx more reminiscent of the nematodes. Indeed, molecular phylogenetic analyses support tardigrades as part of the Ecdysozoa [17,18]. Functional confirmation that tardigrades possess P2X receptors would expand our emerging knowledge of P2X phylogeny and better estimate the pattern of loss of P2X in nematodes and arthropods. Furthermore, a better understanding of P2X receptor function in simple invertebrate organisms may help shed new light on structure-function aspects of human P2X receptors in health and disease by virtue of conservation of functionally important amino acid residues between evolutionary remote P2X receptors and the identification of novel P2X mediated signalling pathways. The aim of this present work therefore was to determine whether the Hypsibius dujardini EST sequence corresponds to a gene encoding a functional P2X receptor and to determine its pharmacological properties. This was achieved by expression of cRNA encoding the Hypsibius dujardini P2X gene in Xenopus oocytes to enable two electrode voltage clamp recordings of ATP evoked membrane currents. Using this approach we show that the Hypsibius dujardini EST sequence does indeed code a functional P2X receptor with fast activation and desensitisation kinetics and similar to some vertebrate P2X channels, is potentiated by ivermectin and inhibited by zinc and copper.

Results

Sequence analysis of HdP2X

Existing EST data for clone Hd_mx23_13F10 consisted of 330 bp from the 5’ end of a cDNA showing similarity to vertebrate P2X receptors. Sequencing of the full insert of this clone showed that it contained an insert of 1743 bp with an open reading frame of 1440 bp. The nucleotide sequence of this cDNA has been submitted to the GenBank database [EU1979525]. The coding sequence of HdP2X is 480 amino acids in length and is predicted to contain intracellular amino and carboxy termini and two transmembrane helices (residues 47 – 67 and 345 – 365) by the TopPred algorithm [19] (Fig. 1). From the human P2X1–7 family, HdP2X was most similar to P2X1, P2X3 and P2X4 with sequence identities of 36.1, 36.5 and 38.4% respectively. However, phylogenetic analysis of the HdP2X protein alongside vertebrate and other P2X sequences suggests that, like other invertebrate P2X receptor proteins, HdP2X does not appear to be a member of any of the seven vertebrate P2X receptor subtypes, but instead represents an ortholog to the ancestor of the vertebrate paralog groups (Fig. 2). Sequence identity with the S. mansoni, D. discoideum (dP2XA), and O. tauri P2X channels was 34.2%, 15.6% and 26.0% respectively. HdP2X contains a number of conserved features typical to P2X channels including ten cysteine residues in the extracellular domain [2], a consensus protein kinase C phosphorylation site in the amino terminal [20], and the lysine residues and NFR/FT motifs shown to be involved in agonist binding (Fig. 1) [21].

ATP evoked currents in HdP2X with fast activation and desensitisation kinetics

In HdP2X expressing Xenopus oocytes clamped at -60 mV, ATP evoked transient inward currents that rapidly desen-
**Figure 1**

Amino acid alignment of Hdp2X. The predicted amino acid alignment of Hdp2X was aligned with human P2X1, and rat P2X2. P2X1 and non vertebrate P2X channels from S. mansoni (smP2X) [12], O. tauri (OtP2X) [11], M. brevicollis (MbP2X) [11] and D. discoideum (dpP2X) [10]. Predicted transmembrane regions in Hdp2X (TopPred algorithm [19]) are depicted by a black horizontal line (top). Details of the predicted secondary structure are shown in Figure 1B. Figure 1C shows the P2X (TopPred algorithm [19]) are depicted by black horizon-
Phylogenetic relationships of *Hypsibius dujardini* P2X.

Aligned protein sequences from P2X homologues from *H. dujardini* (Hd) (Tardigrada), vertebrates (represented by human (h), mouse (m) and rat (r); these taxa have seven paralogous P2X genes (P2X1–7)), *S. mansoni* (Sm) (Platyhelminthes), *L. stagnalis* (Lym) (Mollusca), *D. discoideum* (d) (Mycetozoa; Dictyostelida; five paralogues) and two *Ostreococcus* species, *O. tauri* (Ot) and *O. lucimarinus* (Ol) (Chlorophyta; Prasinophyceae) were analysed using Neighbour Joining and maximum parsimony methods. Both methods agreed on the well supported nodes figured in this phylogram. Nodes were supported with bootstrapstraps of 100% unless otherwise indicated; all nodes with less than 75% bootstrap support were collapsed.

At 100 μM ATP (~EC\(_{50}\)) the rise time from 10% to 90% peak amplitude was 71.4 ± 2.5 ms and the current decay from peak to 50% peak was 289.9 ± 16.1 ms (n = 39). The decay of the current in the continued presence of 100 μM ATP was best fit with two exponentials consisting of a fast component with a time constant of 122 ± 26 ms and a relative amplitude of 0.68 ± 0.04 and a slower component with a time constant of 627 ± 148 ms and a relative amplitude of 0.32 ± 0.04 (n = 6). As a direct comparison using the same experimental apparatus application of 100 μM ATP at the fast human P2X\(_1\) receptor gave rise and decay times of 134.5 ± 11.0 ms and 727.0 ± 88.6 (n = 15) ms respectively. Both hP2X\(_1\) (Fig. 3A) and *S. mansoni* P2X[12] display a marked reduction in current amplitude between the very first and second applications of 100 μM ATP five minutes apart (34.39 ± 5.46% and 45.76 ± 5.10% reductions respectively (n = 15)) before producing stable responses with subsequent applications of agonist at five minute intervals. In contrast, with the same experimental protocol, *Hd*P2X showed a much lower reduction in current amplitude (8.96 ± 5.46% (n = 11)) between first and second applications of ATP (first application mean = -1.77 ± 0.30 μA; second application mean = -1.62 ± 0.29 μA (Fig. 3A) demonstrating that *Hd*P2X recovers more rapidly from desensitisation than hP2X\(_1\) and *S. mansoni* P2X. The current-voltage relationship for *Hd*P2X was obtained by applying 100 μM ATP at a range of holding potentials between -60 mV to and +40 mV. Typical of a non selective cation channel the reversal potential was -3.8 mV (n = 6) and the current voltage relationship showed a slight outward rectification over the range of potentials measured (Fig. 3B and 3C).

**Agonists**

ATP induced responses at *Hd*P2X in a concentration dependent manner with an EC\(_{50}\) of 44.5 μM (pEC\(_{50}\) 4.5 ± 0.1, n = 7) and a Hill slope of 1.12 ± 0.14 (Fig. 3D and 3E). The ATP analogue Bz-ATP also evoked responses at *Hd*P2X with an EC\(_{50}\) of 12.2 μM (pEC\(_{50}\) 5.9 ± 0.2, n = 5) and a Hill slope of 0.7 ± 0.2. The efficacy of Bz-ATP was lower (p < 0.01) than ATP, with the maximum Bz-ATP response 64.6 ± 5.7% that of the maximum response to ATP (Fig. 3D). Currents evoked by Bz-ATP had a slower rise time and rate of desensitization than equivalent ATP evoked currents (p < 0.01 in both cases). With 100 μM BzATP (maximal response) the time from 10%–90% peak current was 142.3 ± 10.2 ms and the rate of current decay from peak to 50% was 446.4 ± 38.9 ms (n = 7) (Fig. 3F).

**Antagonists**

PPADS antagonised ATP evoked currents at *Hd*P2X with an IC\(_{50}\) of 15.0 μM (pIC\(_{50}\) 4.6 ± 0.1, n = 5) and a Hill slope of -1.0 ± 0.3 for responses evoked by 100 μM ATP (Fig. 4). Suramin also antagonised *Hd*P2X currents in a concentration dependent manner but not as strongly as PPADS. Suramin inhibited 100 μM ATP responses with an IC\(_{50}\) of 22.6 μM (pIC\(_{50}\) 4.7 ± 0.3, n = 5–7) and a Hill slope of -0.8 ± 0.3. A component of the *Hd*P2X current was resistant to suramin and at the highest concentration tested (300 μM), only ~75% of the 100 μM ATP evoked response was inhibited (Fig. 4).
Figure 3 (see legend on next page)
**Figure 3** (see previous page)

**Properties of ATP evoked currents.** Two-electrode voltage clamp recordings at a holding membrane potential of -60 mV were made from oocytes expressing HdP2X. A. Comparison of HdP2X and human P2X1 receptors. Currents were recorded in response to 100 μM ATP, indicated by bar. HdP2X displays faster current rise and decay times than the human P2X1, and displays faster recovery from desensitisation between the first and second applications of agonist (applications 5 minutes apart). B. Current voltage relationship of HdP2X. The reversal potential of ATP mediated currents was determined by recording ATP (100 μM, indicated by bar) induced currents at holding potentials ranging from -100 mV to +40 mV with a 5 minute interval between applications. Currents obtained in different oocytes were normalised to the current obtained at -60 mV for each individual cell (n = 6). C. Example currents for the plot depicted in B. D. Concentration response curves for ATP, Bz-ATP and αβ-me-ATP. Mean currents were normalised to the response given by 100 μM ATP (n = 5–7). E and F. Example currents recorded in response to ATP (E) and Bz-ATP (F) (concentrations in μM, agonist application indicated by bar).

**HdP2X is inhibited by zinc and copper**

Vertebrate P2X receptors can be subject to allosteric modulation by both metal ions and pH [22]. Unlike some vertebrate P2X receptors [23], neither the ATP concentration response curve nor the amplitude of ATP evoked currents for HdP2X were affected by acidic pH (pH 6.5) (n = 5, data not shown). The divalent cations zinc and copper however both caused concentration dependent inhibition of ATP-evoked currents (Fig. 5). Zinc inhibited 100 μM ATP currents, with an IC50 of 62.8 μM (pIC50, 4.2 ± 0.14, n = 5–6) and a Hill slope of -0.7 ± 0.1. Copper also acted as an inhibitor of 100 μM ATP currents with an IC50 of 19.9 μM (pIC50 4.7 ± 0.06, n = 5–6) and a Hill slope of -0.8 ± 0.1. (Fig. 5A). Histidine residues have previously been reported to be involved in zinc and copper inhibition of the P2X7 receptor [24,25] and also in the potentiation of P2X receptor currents by zinc [26]. Three histidine residues are present in the extracellular region of HdP2X and we therefore mutated these residues to alanine both individually and in combination with each other in order to probe their potential involvement in metal ion binding. Using 100 μM zinc as a test concentration, wild type currents evoked by 100 μM ATP (∼EC50) were inhibited by 59.3 ± 2.2%. The inhibition observed for the single histidine mutants H232A, H252A and H306A and the double histidine mutants H232A/H252A, H252A/H306A and H232A/H306A was not significantly different from wild type (Fig. 5B). The triple histidine mutant H232A/H252A/ H306A produced a non functional channel and therefore could not be studied. With copper (100 μM) as the inhibitor, wild type currents were inhibited by 81.0 ± 2.3%. Similar to zinc, the inhibition observed in the single histidine and the H232A/H252A double mutant with copper did not significantly differ from wild type (Fig. 5B). The inhibition observed with the double mutants H232A/ H306A and H252A/H306A however was less than wild type (p < 0.05) with H232A/H306A currents inhibited by 60.0 ± 3.9% and H252A/H306A currents inhibited by 50.7 ± 8.9%. In order to assess whether mutation had resulted in a gross change in channel function, the potency of ATP for each histidine mutation was assessed (Fig. 5C) and none of the single or double mutations produced a concentration response curve that differed significantly from wild type. Lysine, aspartic acid, and glutamic acid residues are also potential candidates for coordinating metal binding and a glutamic acid residue in rat P2X7 has been shown to be involved in zinc and copper binding [25]. We therefore mutated lysine and negatively charged residues in HdP2X that were specific between HdP2X and S. mansoni P2X since this channel is also sensitive to metal ions [27]. Five alanine substitution mutations were created (K297A, E249A, D183A, K130A and E110A) and tested for sensitivity to zinc and copper inhibition (Fig. 5B). No significant difference from the wild type inhibition for either zinc or copper was observed in any of these charged amino acid mutations.

**HdP2X currents are potentiated by Ivermectin**

Ivermectin, a broad spectrum anti-parasitic agent from the bacterium *Streptomyces avermitilis* is known to potentiate ATP-evoked currents at human and mouse P2X1 [28,29] and the *S. mansoni* [12] P2X receptors. We therefore studied the effects of this allosteric modulator on the HdP2X receptor. Ivermectin alone was unable to activate the receptor (data not shown). However, 3 μM ivermectin potentiated the response to a maximal concentration of ATP (300 μM) by 223.8 ± 14.1% of control responses, with no notable effect on the rates of current activation or decay (Fig. 6). Currents were significantly (p < 0.01) potentiated in the presence of 3 μM ivermectin at all concentrations of ATP investigated and the concentration response curve for ATP in the presence of 3 μM ivermectin had an EC50 of 32.1 μM (pEC50 4.5 ± 0.1, n = 5–10) and a Hill slope of 0.8 ± 0.2.

**Discussion**

Despite their absence in several key model organisms such as *Saccharomyces cerevisiae*, *Drosophila melanogaster* and *Caenorhabditis elegans*, functional P2X receptors have now been described in a wide range of non-vertebrate metazoan phyla including platyhelminthes [12,27], and molluscs [41], and in representatives of other eukaryotic kingdoms such as dictyostelida [10], prasinophyceae [11] and choanoflagellates [11]. Additional partial and pre-
HdP2X is antagonised by suramin and PPADS. The effects of the P2 receptor antagonists suramin and PPADS were determined in Xenopus oocytes expressing HdP2X at a holding membrane potential of -60 mV. A. Example currents in response to 100 μM ATP in the presence and absence of 100 μM suramin (five minutes between sequential applications). B. Example currents in response to 100 μM ATP in the presence and absence of 100 μM PPADS (five minutes between sequential applications). C. Inhibition curves for mean responses to 100 μM ATP in the presence of suramin (closed squares) (n = 5) and PPADS (open circles) (n = 5).

Figure 5

Copper

Zinc

% Control response

WT

H232A

H252A

H306A

H232A/H306

H252A/H306

H232A/H252A

H232A/H306

H252A/H306

H232A/H252A

% maximum response

Log [ATP]
dicted P2X-like proteins can be identified in EST and partial genome data from other animals, including additional molluscs, echinoderms [30] and cnidaria. These data provide an emerging phylogenetic understanding of the evolution of ionotropic purinergic signalling. Alignment of P2X proteins from a diverse range of taxa with the mammalian P2X (1–7) families (Fig. 2) clearly shows that the mammalian P2X receptors are a lineage-specific gene expansion, with ortholog triplets evident in mouse, human and rat. A lineage-specific expansion is also evident in D. discoideum. In the other taxa, the single P2X receptors identified appear to be orthologous to the entire vertebrate expansion. This strongly suggests that the emergence of a family of seven mammalian P2X receptors was a relatively recent evolutionary event that occurred subsequent to the split between vertebrates and invertebrates. The functional confirmation that P2X receptors exist in the tardigrade H. dujardini strongly suggests that the loss of P2X in some nematode and arthropod species occurred independently, as both traditional and molecular phylogenies place the divergence between Nematoda and Arthropoda before their divergence from Tardigrada. Pharmacological evidence for purinergic signalling in some nematode and arthropod species [6] suggests that P2X-like receptors may have been maintained in some members of these phyla. Indeed a partial transcript for a P2X gene from the nematode Xiphinema index [31] suggests that the loss of P2X in C. elegans may have occurred relatively late in nematode evolution, and P2X-like ESTs from the chelicerate Boophilus microplus similarly suggest that the loss of P2X receptors in arthropods may be restricted to Hexapoda.

Based on their kinetic parameters and ligand sensitivity, mammalian P2X receptors can be divided into three groups [32]. Group 1 consists of P2X1 and P2X3 receptors that have high sensitivity for ATP (EC50 ~1–3 μM) and show rapid channel activation and desensitisation during the continued presence of agonist. Group 2 includes P2X2, P2X4, P2X5, and P2X6 receptors with a lower sensitivity for ATP (EC50 ~10 μM) and much slower desensitization whilst group 3 is represented by P2X7 which has a very low sensitivity for ATP (EC50 ~300 μM) and shows little desensitization. Hdp2X displays an unusual phenotype in
that it has a relatively low sensitivity for ATP (EC$_{50} \sim 44.5$ μM) but has very fast activation and desensitisation kinetics. This is interesting from a mechanistic point of view regarding the relationship between channel gating and agonist binding since it demonstrates that that fast channel kinetics do not necessarily have to be associated with a high agonist sensitivity. Similar to Hdp2X, a P2X$_3$-like receptor from Zebrafish has also been shown to combine fast current kinetics with low sensitivity to ATP [33]. However, unlike Hdp2X (Fig. 3A) current amplitudes at the Zebrafish channel never fully recover from the first stimulation. Other lower organism P2X receptors that have been characterised to date have all displayed slow rates of desensitisation making Hdp2X the first example of a non vertebrate P2X channel with fast kinetics. Given that it is likely that a fast phenotype evolved independently in P2X$_{1-3}$ and Hdp2X channels, it will be interesting to determine whether the same factors determine the rates of current activation and desensitisation in these evolutionary remote channels.

Hdp2X contains several amino acid residues that are highly conserved throughout the vertebrate P2X receptor family. The ten conserved cysteine residues present in the extracellular loop of all vertebrate P2X receptors are thought to form five disulphide bonds [2] and these residues are also present in Hdp2X suggesting conservation of gross extracellular structure constrained by disulphide bond location. It is interesting to note that the P2X receptors from O. tauri and M. breviceps [11] possess only eight of these ten conserved cysteine residues in their putative extracellular regions and that in each case the two absent cysteines correspond to a proposed disulphide bond pairing in the human P2X$_1$ receptor [2] (the equivalents of hP2X$_1$C217 and C227 are absent in O. tauri and C117 and C165 are absent in M. breviceps). The agonist binding site of P2X receptors is distinct from other ATP-binding proteins and is thought to be formed from the interaction between adjacent subunits [34]. Based on site directed mutagenesis combined with methanethiosulfonate reactivity a core common mode of action of ATP binding at P2X receptors has been proposed consisting of lysine residues adjacent to the transmembrane domains coordinating phosphate binding, an NFR motif binding the adenine/ribose moiety and an FT motif also involved in agonist action [21]. The lysine residues, NFR and FT motifs are also present in Hdp2X (Fig. 1) suggesting that this mechanism of ATP binding arose relatively early in evolution and is not restricted to mammalian receptors. However, the NFR motif and the equivalent of hP2X$_1$K68 are absent in the D. discoideum P2X receptor [10], the FT motif is Y1 in S. mansoni P2X [12] and the NFR motif is NIR in the O. tauri [11] receptor suggesting that conservation of the residues involved in ATP binding is not universal to all P2X receptors. Residues at positions D280 and R278 in the rat P2X$_4$ receptor have also been shown to play an important role in the action of ATP [35]. These residues are also conserved in Hdp2X, S. mansoni P2X and M. breviceps P2X suggesting that, despite their incomplete conservation amongst the seven mammalian P2X subtypes, these two residues may also play an important role in the formation of the ATP binding site in some P2X receptor subtypes. Interestingly, the only feature with complete conservation across all vertebrate and lower organism P2X receptors characterised to date is a consen sus protein kinase C phosphorylation site in the N-terminal tail of the receptor (Fig. 1). This putative phosphorylation site has been shown to regulate desensitisation in both P2X$_1$ [36] and P2X$_2$ [20] receptors however, surprisingly direct phosphorylation of this site has been shown not to occur [37,38].

Unlike the D. discoideum [10] and O. tauri [11] P2X receptors, the general P2 receptor antagonists suramin and PPADS both inhibited Hdp2X receptor currents in a concentration dependent manner. PPADS at 100 μM effectively blocked ATP evoked currents. Suramin however was less potent and a component of the current was resistant to antagonism by suramin even at high concentrations. At the highest concentration tested (300 μM) only ~75% of the 100 μM ATP-evoked current was blocked. These properties are opposite to those of the S. mansoni P2X, which was more potently inhibited by suramin and incompletely blocked by PPADS [12]. The presence of a lysine residue at position 246 in rat P2X$_2$ and at equivalent positions in other mammalian subunits has been proposed to be involved in the inhibition of P2X receptors by PPADS, possibly forming a Schiff base with the aldehyde group of PPADS [39]. Similarly a lysine residue at position 138 in human P2X$_1$ has recently been shown to play a key role in the binding of suramin [40]. Neither of these lysine residues shown to be involved in PPADS and suramin action at mammalian receptors are conserved in either Hdp2X or S. mansoni P2X suggesting that additional residues in the ectodomain also play key roles in the mechanism of antagonist binding.

Hdp2X is inhibited by both copper and zinc in a concentration dependent manner (Fig. 5). The effects of these metal ions on vertebrate P2X receptors differ markedly with P2X$_1$ being inhibited by zinc, P2X$_2$ inhibited by both zinc and copper, P2X$_3$ potentiated by both zinc and copper and P2X$_{1-3}$ potentiated by zinc but inhibited by copper (reviewed by [41]). The action of these metal ions at P2X receptors has physiological significance since both zinc and copper are stored in presynaptic terminals and released after nerve stimulation [41,42]. In addition to Hdp2X, the modulation of P2X receptors by metal ions is also present in other lower organisms with Dictostelium P2X inhibited by copper [10] and S. mansoni P2X inhibi-
P2X channels may enhance our understanding of the process of gating and motions of the channel pore.

**Conclusion**

The identification and pharmacological characterisation of a P2X receptor from *H. dujardini* provides further evidence that P2X receptors for ATP emerged early in eukaryotic evolution. Several unusual characteristics of *Hd*P2X including fast current kinetics with low ATP sensitivity, ivermectin potentiation in a channel with fast kinetics and the likelihood of distinct copper and zinc metal binding sites make this channel a useful model for comparative structure-function studies allowing a better understanding of P2X receptors in higher organisms.

**Methods**

Identification of the *Hypsibius dujardini* P2X receptor

BLAST searches of the GenBank EST database identified a partial *Hypsibius dujardini* EST sequence [GenBank:CO741227] encoding a peptide with significant sequence similarity to the amino terminus of vertebrate P2X receptors. The insert of the corresponding clone for this EST sequence ([Hd_mz23_13F10] was fully sequenced on both strands (Automated ABI sequencing service, Leicester University, U.K.) using vector (pSPORT1) and insert specific primers and was shown to contain a full length coding sequence which was named *Hd*P2X.

**Phylogenetic analysis**

P2X protein sequences were obtained from SwissProt/UniProt and aligned with ClustalW. The alignment was adjusted by eye to minimise unique indels and subjected to phylogenetic analysis in PAUP* version 4.10b using neighbour Joining (BioNJ) and maximum parsimony methods. The alignment was 696 characters long, and had 147 invariant and 433 parsimony-informative characters. As the N-terminal and C-terminal portions of the proteins were poorly aligned, the analysis considered only the central 445 characters (of which 342 were parsimony informative, and 208 included gaps). The alignment is available from the authors (MLB). Support for the derived phylogenies was assessed by 1000 bootstrap iterations.

**Site directed mutagenesis**

Point mutations in the *Hd*P2X plasmid construct were introduced using the QuikChange™ Mutagenesis Kit (Stratagene, U.S.A.) according to the manufacturer’s instructions. Histidines at positions 232, 252 and 306, glutamic acid residues at positions 110 and 249, an aspartic acid residue at position 183 and lysine residues at positions 130 and 297 were mutated to alanine. Double mutations where two of the three histidines were both mutated to alanine were created by conducting serial rounds of site directed mutagenesis as follows: H252A/
H232A, H252A/H306A, H232A/H306A. A triple mutant where all three histidines were mutated to alanine was constructed by introducing the H306A mutation into the H252A/H232A double mutant. In all mutants, introduction of the correct mutation(s) and the absence of spontaneous mutations were confirmed by DNA sequencing (Automated ABI sequencing service, Leicester University).

Oocyte preparation

Mutant and wild type plasmids were digested with NotI to linearise and sense strand cRNA was generated using the T7 mMessage mMachine™ kit (Ambion, U.S.A.) according to the manufacturer's instructions. Manually defolliculated stage V-VI Xenopus oocytes were injected with 5 ng of cRNA (50 nl at a concentration of 0.1 ng/μl) using an Inject +Matic micro injector (J. Alejandro Gaby, Genève) and were stored at 18°C in ND96 buffer (96 mM NaCl, 2 mM KCl, 1.8 mM CaCl₂, 1 mM MgCl₂, 5 mM sodium pyruvate, and 5 mM HEPES, pH 7.5) before electrophysiological recordings were carried out 3–6 days later.

Two-electrode voltage clamp

Two-electrode voltage clamp recordings were made from oocytes using a Turbo TEC 10C amplifier (NPI Electronic Instruments, Germany) with a Digidata 1200 analogue to digital converter (Axon Instruments U.S.A.) and WinWCP acquisition software (Dr J. Dempster University of Strathclyde, Scotland). Microelectrodes were filled with 3 M KCl, and the external solution consisted of ND96 buffer with 1.8 mM CaCl₂ replaced with 1.8 mM BaCl₂ to prevent the activation of endogenous calcium-activated chloride channels. Membrane currents were recorded at a holding potential of -60 mV. Agonists, ATP (Mg²⁺ salt), 2',3'-O-4-Benzoylbenzoyl ATP (Bz-ATP) and αβ-methylene-adenosine 5'-triphosphate (αβmeATP) (Sigma, Poole, U.K.) were applied from a U-tube perfusion system, whereas ivermectin, pyridoxal-phosphate-6-azophenyl-2',4'-disulfonic acid; PPADS: Pyridoxal-phosphate-6-azophenyl-2',4'-disulfonic acid; αβmeATP: αβ-methylene-adenosine 5'-triphosphate; ADP₆hex: Hexokinase treated ADP.

Data analysis

Data are presented as means ± S.E. Differences between means were assessed by either the Student's t-test for simple comparison between two data sets or by ANOVA followed by Dunnett's post test when multiple data sets were compared (GraphPad Prism software (La Jolla, USA)). Concentration-response data were fitted with the equation $Y = \left(\frac{\text{X}^{H \cdot M}}{\text{EC}_{50}}\right) + \left(\text{X}^{H \cdot M}\right)$, where $Y$ is response, $X$ is agonist concentration, $H$ is the Hill coefficient, $M$ is maximum response, and $\text{EC}_{50}$ is the concentration of agonist evoking 50% of the maximum response. pEC₅₀ is the-log₁₀ of the EC₅₀ value. All concentration-response curves, EC₅₀ values and Hill coefficients were obtained using GraphPad Prism software.

Abbreviations

PPADS: Pyridoxal-phosphate-6-azophenyl-2',4'-disulfonic acid; αβmeATP: αβ-methylene-adenosine 5'-triphosphate; ADP₆hex: Hexokinase treated ADP.

Authors' contributions

SB carried out electrophysiological recordings, wrote the first draft of the manuscript and analysed the data. VAS participated in the design of the study and helped draft the manuscript. MLB conducted the phylogenetic analysis, helped draft the manuscript and generated the HidP2X EST clone. SJE conceived and designed the study, performed the molecular biological aspects of the study and wrote the manuscript. All authors read and approved the final manuscript.

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