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Evolution of Acetylcholinesterase and Butyrylcholinesterase in the Vertebrates: An Atypical Butyrylcholinesterase from the Medaka Oryzias latipes

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Abstract

Acetylcholinesterase (AChE) and butyrylcholinesterase (BChE) are thought to be the result of a gene duplication event early in vertebrate evolution. To learn more about the evolution of these enzymes, we expressed in vitro, characterized, and modeled a recombinant cholinesterase (ChE) from a teleost, the medaka Oryzias latipes. In addition to AChE, O. latipes has a ChE that is different from either vertebrate AChE or BChE, which we are classifying as an atypical BChE, and which may resemble a transitional form between the two. Of the fourteen aromatic amino acids in the catalytic gorge of vertebrate AChE, ten are conserved in the atypical BChE of O. latipes; by contrast, only eight are conserved in vertebrate BChE. Notably, the atypical BChE has one phenylalanine in its acyl pocket, while AChE has two and BChE none. These substitutions could account for the intermediate nature of this atypical BChE. Molecular modeling supports this proposal. The atypical BChE hydrolyzes acetylthiocholine (ATCh) and propionylthiocholine (PTCh) preferentially but butyrylthiocholine (BTCh) to a considerable extent, which is different from the substrate specificity of AChE or BChE. The enzyme shows substrate inhibition with the two smaller substrates but not with the larger substrate BTCh. In comparison, AChE exhibits substrate inhibition, while BChE does not, but may instead show substrate activation. The atypical BChE from O. latipes also shows a mixed pattern of inhibition. It is effectively inhibited by physostigmine, typical of all ChEs. However, although the atypical BChE is efficiently inhibited by the BChE-specific inhibitor ethopropazine, it is not by another BChE inhibitor, iso-OMPA, nor by the AChE-specific inhibitor BW284c51. The atypical BChE is found as a glycophosphatidylinositol-anchored (GPI-anchored) amphiphilic dimer (G2+), which is unusual for any BChE. We classify the enzyme as an atypical BChE and discuss its implications for the evolution of AChE and BChE and for ecotoxicology.


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Introduction

Acetylcholinesterase (AChE; EC 3.1.1.7) hydrolyzes acetylcholine at the neuromuscular junction of vertebrates. Higher vertebrates also contain an evolutionarily related cholinesterase (ChE), butyrylcholinesterase (BChE; EC 3.1.1.8). The function of BChE is unknown but is suggested to play a role in growth and development and to act as a scavenger of cholinergic toxins as well as having an auxiliary role in synaptic transmission [1,2]. The two ChEs may be distinguished functionally both kinetically and pharmacologically: AChE hydrolyzes acetylcholine (Ach) and is virtually inactive on the larger substrate butyrylcholine (BCh). BChE is less selective, hydrolyzing both substrates comparably. AChE exhibits inhibition at high substrate concentrations, while BChE shows substrate activation instead [3]. The two enzymes may also be distinguished by their susceptibility to diagnostic inhibitors [4].

Within species, AChE and BChE have ~50% amino acid identity, and the overall tertiary structures of the two enzymes are similar [5,6]. Individual amino acid residues involved in determining the molecular basis of the differences in substrate and inhibitor specificity of AChE and BChE have been identified in the acyl pocket, located at the bottom of a deep catalytic gorge; the peripheral site, located at the lip of the gorge; the oxyanion hole; and the choline-binding site of the hydrophobic patch, also located within the gorge [7–14]. Although the dichotomy between AChE and BChE is generally clear in birds and mammals [1,15,16], the two enzymes often more closely resemble one another functionally in fish. In the cartilaginous fish, the electric ray Torpedo marmorata [17], and the bony fishes, the plaice Pleuronectes platessa [18], the flounder Platichthys flesus [19], and perhaps the surgeonfish Acanthurus dussumieri [20,21], ChEs with properties intermediate to and atypical of AChE and BChE are found along with AChE. These enzymes have alternatively been considered atypical ChEs [18,19] or atypical pseudo-cholinesterases (pseudo-ChEs) [17,20]; we are designating them as atypical BChEs, as suggested by Whitaker [22]. Although a number of cDNAs have been cloned for AChEs from these organisms, molecular information about the atypical BChEs present is...
unavailable. Moreover, only a single ChE, AChE, has been identified functionally and molecularly in the jawless fish, the lamprey *Petromyzon marinus* [23] and the hagfish *Myxine glutinosa* [24]. These observations suggest that AChE is the ancestral ChE in the vertebrates and that an early gene duplication event and subsequent divergent structural and functional evolution produced the AChE and BChE of higher vertebrates [23,25].

AChE and BChE also exist in a variety of homomeric and heteromeric molecular forms. The catalytic subunit of AChE is found in different variants as a result of alternative splicing of the C-terminus, producing R, H, and T (or AChET, AChEH, or AChET2) subunits [26,27]. The R, or read-through, transcript is rare and produces soluble non-amphiphilic monomers, G₂₄₈, [29]. AChET has a hydrophobic C-terminus, which is replaced by a glycosphatidylinositol-phospholipid (GPI) anchor and produces amphiphilic dimers, G₂₄₈, [29]. AChEH is capable of forming G₂₄₈, G₂₄₈, and G₂₄₈₄ [29], as well as “tailed” forms (thus the T subunit) by associating with a transmembrane protein, the Proline-Rich Membrane Attachment (PRiMA) [30] and the triple helical collagen Q (ColQ; Q for queue, tail in French) [31,32]. In brain and at the neuromuscular junction, PRiMA localizes AChE to the cell membrane of synapses, forming G₂₄₈ (or G₂₄₈). ColQ anchors AChE to the junctional basal lamina of the neuromuscular junction, producing A₄₄₄, A₄₄₄, and A₄₄₄₄, which represent one, two or three tetramers attached to the ColQ triple helix. While AChET is found in all classes of vertebrates, AChET₄ exists in cartilaginous fish (*Torpedo spp.*) [33], perhaps amphibians (*Xenopus laevis*) [34], and mammals [35], but has not been reported in jawless or bony fish, reptiles, or birds, raising questions about the evolution of this splice variant [26].

BChE does not exhibit alternative splicing and is considered to be found solely as a T variant (*BChET*) [36,37] that also associates with PRiMA and ColQ [30,36]. R and H variants of BChE have not been reported. However, according to the *Xenopus tropicalis* genome project [38] and other evidence [34,39–42], an H variant of BChE appears to be present in amphibian *Xenopus species*. The atypical BChEs of *T. marmorata* and *A. dussumieri* are T variants (*BChET*), assembling a collection of globular and asymmetric forms [17,20]. In remarkable contrast, the atypical BChE of *P. japonicus* is BChEH, assembling only into GPI-anchored G₂₄₈ forms [19].

The medaka *Oryzias latipes* is a teleost fish that is of interest as a vertebrate model system for developmental, genomic, and evolutionary biology [43–45]. It was previously reported that *O. latipes* possesses an AChE [46]. Here we report the cloning and expression of a ChE from *O. latipes*, and briefly discuss the implications of the structure and function of this enzyme for the evolution of the ChEs. Additionally, the presence of ChEs with anomalous inhibitor specificities has ecotoxicological implications for *O. latipes* [47,48] and other fish [49,50].

**Results**

**Sequence Analysis Reveals Two ChEs in *O. latipes***

Two expressed sequences for ChEs are present in the *O. latipes* genome: AChE (GenBank EST DK110600) and an enzyme that we are classifying as an atypical BChE [45] (GenBank cDNAs AV668390 and GU792751). The sequence for the AChE is truncated near the carboxyl terminus and contains 561 amino acids. The sequence of the mature polypeptide for the atypical BChE from *O. latipes* contains 564 amino acids (Fig. 1). Pair-wise BLAST alignments of sequences from the catalytic region of ChEs show that the AChE from *O. latipes* clearly resembles *T. californica* AChE rather than *Homo sapiens* BChE (68/80% identity/similarity to AChE compared to 54/70% for BChE), while the atypical BChE resembles both AChE and BChE more or less equally (46/68% for AChE and 49/67% for BChE). A phylogenetic tree of vertebrate and deuterostome invertebrate ChEs is shown in Fig. 2; the AChE of *O. latipes* is found in the AChE clade, while the atypical BChE of *O. latipes* is found in the BChE clade.

The members of the catalytic triad of ChEs are found as Ser203, Glu330, and His461 in the AChE, and Ser203, Glu330, and His463 in the atypical BChE. Of the fourteen aromatic amino acids that line the catalytic gorge of vertebrate AChE, all are conserved in the *O. latipes* AChE, and ten are conserved in the atypical BChE; in contrast, eight are conserved in vertebrate BChE (Fig. 1; Table 1). The *O. latipes* atypical BChE is missing two of the three aromatic residues of the peripheral site of AChE, while BChE lacks all three. Additionally, while AChE has two Phe residues in the acyl pocket and BChE none, the *O. latipes* atypical BChE has one Phe (Fig. 1; Tables 1, 2). As the *O. latipes* AChE conserves all ten aromatic residues, it has two Phe residues in its acyl pocket.

The three pairs of conserved cysteine residues involved in intra-chain disulfide bonding are also found as Cys69-Cys96, Cys257-Cys268, and Cys405-Cys543 in the AChE; Cys69-Cys96, Cys257-Cys268, and Cys405-Cys543 in the atypical BChE of *O. latipes*. Another cysteine (Cys540), near the carboxyl terminal that normally mediates inter-chain disulfide bonding, is also conserved in the atypical BChE (Fig. 1). The carboxyl terminal of the enzyme is of the H-type (Fig. 3), characterized by a loosely defined GPI anchor signal, including an ω cleavage/attachment site followed by a stretch of hydrophobic amino acids [51]. The H-peptides show very little sequence homology to one another. We did not find evidence for the existence of T-type carboxyl terminus for the atypical BChE either as a T-exon in the genome or a T-type carboxyl terminus in the ESTS of *O. latipes*. The truncated AChE is missing its C-terminal sequence.

**Diagnostic Inhibitors Show the Presence of Two ChE Activities in Adult *O. latipes***

To demonstrate the presence of two ChE activities in *O. latipes*, extracts from adult medaka were incubated with the inhibitors physostigmine, which inhibits all ChEs; BW294c51, which inhibits AChE selectively; and ethopropazine, which inhibits BChE preferentially, and assayed for activity with ATCh and PTCh [16,52,53]. Different dose-response curves were observed with the two substrates for each inhibitor, suggesting the presence of at least two ChE activities (Fig. 4). The dose-response curves for inhibition of PTCh hydrolysis by BW294c51 and ethopropazine are biphasic, clearly indicating the presence of two ChE activities, presumably the AChE and atypical BChE identified in the *O. latipes* genome project.

**Kinetic Analysis of the Atypical BChE Indicates Its Anomalous Properties**

As we were interested in the functional characteristics of the atypical BChE in *O. latipes*, we cloned and expressed in *vitro* a cDNA for the enzyme in COS-7 cells. To determine the substrate specificity of the enzyme, we assayed the hydrolysis of the substrates acetylthiocholine (ATCh), propionylthiocholine (PTCh), and butyrylthiocholine (BTCh) by the recombinant enzyme (Fig. 5). The smaller substrates ATCh and PTCh are hydrolyzed more or less equally, as indicated by the similar values of kcat/Substrate/ kcat/ATCh (Table 3); the larger BTCh is hydrolyzed at about a quarter of the rate of the other two substrates. The K_a is inversely proportional to the length of the acyl group, with BTCh having the lowest K_a. The highest catalytic efficiency (kcat/K_a) is seen with PTCh. Additionally, ATCh and PTCh produce

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**An Atypical Cholinesterase from Medaka**

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Anomalous Properties
substrate inhibition (i.e., lower enzyme activity at high substrate concentrations and b parameter values of <1), although the inhibition by PTCh is weak (Fig. 5; Table 3). BTCh does not produce substrate inhibition. Overall, this pattern of substrate hydrolysis is not typical of either AChE or BChE.

Pharmacological Analysis of the Atypical BChE Confirms Its Anomalous Properties

Since the recombinant enzyme from O. latipes exhibited anomalous kinetic properties, to characterize further this atypical BChE activity, we determined the half maximal inhibitory concentration (IC_{50}) values of the enzyme for the inhibitors physostigmine, which inhibits all ChEs; BW284c51, which inhibits AChE preferentially; and iso-OMPA and ethopropazine, which inhibit BChE preferentially. Phystostigmine and ethopropazine inhibit the enzyme at sub-μM concentrations; by contrast, much higher concentrations of BW284c51 and Iso-OMPA are required for inhibition under the conditions tested (Fig. 6; Table 4). This pattern of inhibition is also not characteristic of either AChE or BChE.

Analysis of Molecular Forms of the Atypical BChE further Demonstrates Its Unusual Nature

ChEs exist in various homomeric and heteromeric molecular forms depending, in part, on the nature of their carboxyl termini. Since the amino acid sequence of the atypical BChE indicates an H-type C-terminus, we performed velocity sedimentation on sucrose gradients in the presence and absence of the non-ionic detergent Triton X-100 to determine the molecular forms of the recombinant enzyme produced in vitro by COS-7 cells. The extract contains G_2^a forms on the basis of the sedimentation coefficient (3.33±0.10 S; Mean±SE, N = 6) and its shift to higher values in the absence of detergent due to aggregation of the enzyme (0.9±0.05; Mean±SE, N = 6) (Fig. 7). Digestion of intact COS-7 cells with phosphatidylinositol-specific phospholipase C (PIPLC) releases ~80% of the surface enzyme activity. Spontaneous release of activity into the supernatant during incubation in the absence of PIPLC was ~10% (Fig. 8). These data indicate that the G_2^a produced is a glyrophosphatidylinositol-anchored (GPI-anchored) form. It is unusual for any BChE to be found as a GPI-anchored form.

Molecular Modeling of the Atypical BChE Illustrates Its Differences with AChE and BChE

We built a homology model of the atypical BChE from O. latipes based on the X-ray structures of H. sapiens BChE, and T. californica and Drosophila melanogaster AChEs, in order to get a structural understanding of the special pharmacological and enzymatic properties of the enzyme (Fig. S1). A comparison of the active site grooves is presented in Figure 9. The volume of the O. latipes atypical BChE catalytic gorge (630 Å³) is much closer to that of H. sapiens BChE (690 Å³) than that of T. californica AChE (410 Å³). The difference in volume is related to the lack of aromatic residues in three gorge subsites: the peripheral site, the acyl binding pocket, and the choline binding pocket. T. californica AChE possesses three conserved residues (Tyr70, Tyr121, and Trp279) that form an aromatic peripheral binding site and restrict access to the gorge. None of these aromatic residues is conserved in H. sapiens BChE and only one aromatic residue is present in the O. latipes atypical BChE (Tyr282), thus enlarging the gorge entrance (Table 1).

The acyl binding pocket of T. californica AChE is also smaller than that of the atypical BChE from O. latipes, which is in turn smaller than that of H. sapiens BChE, due to the respective decreasing number of the phenylalanines shaping the pocket. These differences are expected to change the selectivity of the enzymes for substrates and inhibitors bearing large chains oriented toward this pocket.

Another major difference between the O. latipes atypical BChE and H. sapiens BChE and T. californica AChE is located in the choline binding pocket. Phe330 (a tyrosine in mammalian AChE) is substituted by a non-aromatic residue in both BChEs (Ala328 in the H. sapiens BChE and Cys333 in the O. latipes atypical BChE). Phe330 can adopt different conformations in the presence of different ligands thus providing an efficient way to modulate the shape and size of the choline binding pocket, even acting as a molecular lid. The absence of a gating aromatic residue at this position in BChE leaves the gorge wide open at all times. Notably, just next to this position, the substitution of Met442 in H. sapiens BChE by Ala440 in O. latipes BChE results in a significantly widened choline binding pocket.

All things considered, the active site gorge of the atypical BChE from O. latipes shares more structural features with that of H. sapiens BChE than that of T. californica AChE, and it appears legitimate to consider this enzyme as an atypical BChE from a structural point of view.

Discussion

Overview

In addition to possessing an AChE, the medaka O. latipes has an atypical BChE that is in many respects different from either vertebrate AChE or BChE (Table 5). Of the fourteen aromatic amino acids in the catalytic gorge of vertebrate AChE, ten are conserved in the atypical BChE from O. latipes; by contrast, eight are conserved in vertebrate BChE. These substitutions may account for the intermediate nature of the atypical BChE. Molecular modeling supports this interpretation. The enzyme hydrolyzes ATCh and PTCh preferentially, but BTCh to a considerable extent. In contrast, AChE is highly specific for ATCh compared to BTCh and even PTCh, while BChE hydrolyzes the larger substrates preferentially. The K_m for substrate hydrolysis by the atypical BChE are inversely proportional to the length of the acyl group, which is more a characteristic of BChE than AChE. The enzyme shows substrate inhibition with the two smaller substrates but not the larger substrate BTCh. By comparison, AChE exhibits substrate inhibition, while BChE does not, but may instead show substrate activation. The O. latipes enzyme also shows an atypical pattern of inhibition by diagnostic inhibitors. It is effectively inhibited by the ChE inhibitor physostigmine, typical of all ChEs. However, although the atypical BChE is efficiently inhibited by the BChE-specific inhibitor ethopropazine, it is not inhibited by another BChE inhibitor, iso-OMPA, nor by the AChE-specific bis-quaternary inhibitor BW284c51. The atypical BChE is found as a GPI-anchored G_2^a membrane-bound dimeric form, also unusual for a BChE. We consider the enzyme an
atypical BChE that has implications for the evolution of AChE and BChE in the vertebrates.

Comparison of Kinetic and Pharmacological Parameters of the Atypical BChE from *O. latipes* with AChE, BChE, and other Atypical BChEs in Vertebrates

Fluck [46] characterized the acylcholine hydrolyzing activity of early embryos of *O. latipes* and found that ATCh, PTCh, and BTCh were hydrolyzed at the relative rates of 1.0, 0.4, and 0.14, respectively; that ATCh, at least, produced substrate inhibition; and that 10 μM BW284c51 inhibited 90% of the activity, while 10 μM iso-OMPA inhibited only 10% of the activity. He concluded that the enzyme present was AChE. Given the results from the *O. latipes* genome project, which indicate two ChEs in the genome; our results for BW284c51 and ethopropazine inhibition of the ChE activity of adult medaka, which also suggest two ChEs; and our data for the recombinant atypical BChE from *O. latipes*, it seems likely that both AChE and BChE were present in the embryos studied by Fluck [46].

The *Kₘ* for substrate hydrolysis by the atypical BChE from *O. latipes* is inversely proportional to the length of the acyl chain of the substrate (Table 5). This relationship is seen in *H. sapiens* BChE, which is representative of the typical vertebrate BChE; but not *T. marmorata*, which is representative of vertebrate AChE. This pattern is the most conserved characteristic of the atypical BChEs from the flounder *P. flesus*, the placid *P. platessa*, the surgeonfish *A. dussumieri*, and the ray *T. marmorata*, suggesting that the non-covalent stabilization of binding of substrate is determined similarly in these enzymes. The acyl pocket, Phe288 and Phe290 in *Torpedo* spp. AChE, is implicated in this binding [8,12-14]. A detailed discussion of the molecular basis of *Kₘ* and other parameters of substrate and inhibitor specificity is given in the next section; since sequence information is only available for the atypical BChE from *O. latipes*, it alone, among the atypical BChEs, can be compared to AChE and BChE at the molecular level.

In contrast to the clear series seen for *Kₘ* the substrate hydrolysis specificities of the enzymes, as defined by relative *Vₘₐₓ* or *kₐₙ* values, do not show as consistent a pattern. Most of the atypical BChEs hydrolyze ATCh preferentially and BTCh the least. An exception is the *O. latipes* atypical BChE, which hydrolyzes PTCh maximally, albeit only slightly faster than ATCh. However, there are two additional exceptions to this generalization: the *P. platessa* atypical BChE hydrolyzes BTCh faster than PTCh, and the *A. dussumieri* atypical BChE hydrolyzes BTCh only marginally better than does AChE. The variation in relative *Vₘₐₓ* or *kₐₙ* values suggests that there are differences in the stabilization in the covalent Michaelis complexes of substrates in the various atypical BChEs. Such differences could be due to differences in the nature of the conformation of the histidine of the catalytic triad, which appears to be determined by different sets of structural interactions in AChE and BChE [9,54].

There are also four different patterns of substrate inhibition among the five atypical BChEs, indicating another heterogeneity among the enzymes. The physiological relevance and molecular mechanism of substrate inhibition are unclear [55–57]. Nevertheless, these differences in the enzymes are probably due to alterations of amino acids, probably non-aromatic substitutions, comprising the peripheral site and/or other regions of the catalytic gorge in the various enzymes that are important for substrate inhibition [9,13,58,59]. It is notable that the substrate hydrolysis curve for PTCh hydrolysis by the *O. latipes* atypical BChE, particularly in the region of substrate inhibition resembles the curves for AChT and PTCh hydrolysis by the atypical BChE from *A. dussumieri*, as the substrate inhibition levels off in both of the enzymes [21].

The atypical BChEs also show different patterns of sensitivity to inhibitors; however, given the range of inhibitors used, and focusing on the atypical BChE from *O. latipes*, it is possible to compare their inhibition only for the AChE and BChE diagnostic inhibitors BW284c51 and ethopropazine, and even for these inhibitors the data are incomplete and a consistent pattern elusive.

Table 1. Aromatic Amino Acids in the Catalytic Gorge of Vertebrate ChEs.

<table>
<thead>
<tr>
<th>Subsite</th>
<th>Torpedo AChE</th>
<th>Oryzias AChE</th>
<th>Oryzias BChE</th>
<th>Homo BChE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peripheral Site</td>
<td>Tyr70</td>
<td>Phe72</td>
<td>Met69</td>
<td>Asn68</td>
</tr>
<tr>
<td></td>
<td>Tyr121</td>
<td>Tyr124</td>
<td>Val124</td>
<td>Gly119</td>
</tr>
<tr>
<td></td>
<td>Trp279</td>
<td>Trp282</td>
<td>Tyr282</td>
<td>Ala277</td>
</tr>
<tr>
<td>Hydrophobic Patch¹</td>
<td>Trp84</td>
<td>Trp86</td>
<td>Trp83</td>
<td>Trp82</td>
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<tr>
<td></td>
<td>Tyr130</td>
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<td>Trp133</td>
<td>Tyr128</td>
</tr>
<tr>
<td></td>
<td>Phe330</td>
<td>Phe333</td>
<td>Cys333</td>
<td>Ala128</td>
</tr>
<tr>
<td></td>
<td>Phe331</td>
<td>Phe334</td>
<td>Phe334</td>
<td>Phe329</td>
</tr>
<tr>
<td>Acyl Pocket</td>
<td>Phe288</td>
<td>Phe291</td>
<td>Leu291</td>
<td>Leu286</td>
</tr>
<tr>
<td></td>
<td>Phe290</td>
<td>Phe293</td>
<td>Phe293</td>
<td>Val288</td>
</tr>
<tr>
<td>Wall of Gorge</td>
<td>Trp114</td>
<td>Trp117</td>
<td>Trp119</td>
<td>Trp114</td>
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<tr>
<td></td>
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<td></td>
<td>Tyr442</td>
<td>Tyr463</td>
<td>Tyr445</td>
<td>Tyr440</td>
</tr>
</tbody>
</table>

Conserved aromatic residues are shown in bold. Torpedo AChE is representative of all vertebrate AChEs and Homo BChE is representative of all vertebrate BChEs.

¹Includes the choline-binding site.

doi:10.1371/journal.pone.0017396.t001
Of the atypical BChEs, those from *O. latipes* and *P. platessa* resemble most closely ChE, being sensitive to ethopropazine, but not BW283c51; the BChE from *T. marmorata* is equally sensitive to both inhibitors, while the BChEs from *P. platessa* and *A. dussumieri* appear to be sensitive and resistant, respectively. These inhibitors are sensitive to the presence or absence of aromatic amino acids in the catalytic gorge [12,14,60,61], which could be and are sensitive to the presence or absence of aromatic amino acids [12,14,60,61]. Overall, the fact that the atypical BChE from *O. latipes* has little effect on the Km for ATCh, but decreases Km for BTCh 10-fold [12]. In *T. marmorata* BChE, the F295L (F288L) mutation is consistent with this explanation.

### Molecular Basis of Substrate and Inhibitor Specificity in the Atypical BChE from *O. latipes* in Comparison to AChE and BChE

The atypical BChE from *O. latipes* maximally hydrolyzes ATCh and PTCh almost equally and the larger substrate BTCh at about a quarter of the rate of the two other smaller substrates. This substrate specificity is not typical of either AChE, which hydrolyzes ATCh maximally, PTCh adequately, but not BTCh at all; nor BChE, which hydrolyzes the three substrates more or less equally [15]. An important molecular determinant of ChE specificity is the acyl pocket, which in vertebrate AChE is characterized by two Phe residues (Phe295, Phe290, and Val288 (Phe290), seems sufficient to explain the intermediate substrate specificity of the enzyme. Our molecular modeling is consistent with this explanation.

In the mouse *Mus musculus* AChE, the F295L (F288L) mutation has little effect on the Km for ATCh but lowers the Km for BTCh 30-fold [14]. In *H. sapiens* AChE, F295L has little effect on the Km for ATCh or PTCh but decreases Km for BTCh 10-fold [12]. In contrast, F295A also spares the Km for ATCh, but decreases Km for PTCh and BTCh 4-fold and 33-fold, respectively. With respect to
The F295L, A mutants do not affect appreciably the $k_{cat}$ (stickleback), F. rubripes $K_m$ H. sapiens ATCh and increases it 14-fold for BTCh. In the results that we have obtained for the factor of three. These data are consistent with, if not identical to, found for BTCh, with the efficiency for all three substrates within a ATCh; whereas, for the F295L, A mutants, the highest efficiency is hydrolyzed appreciably, the which lacks the homologous Phe: all three substrates are not conserved in BChE is well stabilized by interactions with an adjacent residue corresponding to Phe288 affects both the binding and substrates are within a factor of two. Thus, it appears that the shape the acyl binding pocket of AChE, along with the aromatic side chain of the residue and the diethylamino-2-isopropyl amino acids of the choline-binding site at the bottom of the gorge and the peripheral site at its rim. In contrast, BChE has a number of these residues replaced by aliphatics. In M. musculus AChE, the peripheral site double mutant Y72N/Y124Q (Y70N/Y121Q) increases $K_i$ for BW284c51 69-fold [13]. Comparably, in H. sapiens AChE the same double mutation increases $K_i$ for the inhibitor 35-fold. Moreover, the choline-binding site mutation Y337A (Phe330A) increases the $K_i$ 5-fold [60]. Making the reasonable assumption of a synergistic effect for the mutations [60], a triple Y72N/Y124Q/Y337A mutation encompassing both the peripheral and choline-binding sites could increase $K_i$ by over two orders of magnitude. Thus in the O. latipes atypical BChE, which has the aliphatic substitutions Met69, Val124, and Cys333 at the homologous sites, preventing the necessary cation-π and π-π interactions to aromatic amino acids of the choline-binding site at the bottom of the gorge and the peripheral site at its rim. In contrast, BChE has a number of these residues replaced by aliphatics. In M. musculus AChE, the peripheral site double mutant Y72N/Y124Q (Y70N/Y121Q) increases $K_i$ for BW284c51 69-fold [13]. Comparably, in H. sapiens AChE the same double mutation increases $K_i$ for the inhibitor 35-fold. Moreover, the choline-binding site mutation Y337A (Phe330A) increases the $K_i$ 5-fold [60]. Making the reasonable assumption of a synergistic effect for the mutations [60], a triple Y72N/Y124Q/Y337A mutation encompassing both the peripheral and choline-binding sites could increase $K_i$ by over two orders of magnitude. Thus in the O. latipes atypical BChE, which has the aliphatic substitutions Met69, Val124, and Cys333 at the homologous sites, preventing the necessary cation-π and π-π interactions to aromatic amino acids, one might expect inefficient inhibition by BW284c51, which is exactly what is observed.

Notably, the structure of the acyl pocket does not appear to influence the binding of BW284c51 [13,14,60], consistent with the situation in the atypical BChE of O. latipes.

The atypical BChE of O. latipes is inhibited by a low concentration of the tricyclic amine-containing phenothiazine, ethopropazine, which contains a bulky diethylamino-2-isopropyl exocyclic group – inhibition typical of vertebrate BChE. Inhibition of AChE and BChE by ethopropazine appears to be sensitive to the nature of the residue corresponding to Phe330 of the aromatic patch in Torpedo spp. AChE and to the volume of the active center in BChE [13,61]. In M. musculus AChE, the Y337A (F330A) mutation decreased $K_i$ almost 2700-fold, making it comparable to BChE [13], presumably because the side chain of Tyr337 sterically hinders the binding of ethopropazine via an interaction between the aromatic side chain of the residue and the diethylamino-2-isopropyl moiety of the inhibitor. By contrast, the converse mutations A328F and A328Y in H. sapiens BChE do not substantially affect the $K_i$ for the inhibitor [61]. There is, however, a large difference in the gorge dimensions between AChE and BChE, and this difference was used to explain the lack of effect. The volume of the lower portion of the gorge in Torpedo spp. AChE was reported to be 302 Å³, and in a modeled H. sapiens BChE it was measured as 302 Å³. The van der Waals volume of ethopropazine is 318 Å³, explaining why the drug does not bind.
The F330A mutation of *Torpedo* spp. AChE increases the lower gorge volume to 338 Å³ and allows ethopropazine to bind. In contrast, the A328Y and A328F mutations in the *H. sapiens* BChE model only decrease the volume of the lower gorge to 410 and 406 Å³, respectively, which allow the gorge to still be large enough to bind ethopropazine easily [61].

Our own volume calculations using the narrower part of the bottleneck to define the gorge entrance and taking into account the conserved structural water molecule, gives an overall volume of 690 Å³ for *H. sapiens* BChE, 630 Å³ for *O. latipes* atypical BChE and 410 Å³ for *Torpedo* spp. AChE, in good agreement with the observed trend for ethopropazine inhibition.

Iso-OMPA is an effective organophosphate inhibitor of BChE but not AChE [52], where it is over 10,000 times less reactive [14], and this selectivity appears dependent on the dimensions of the active center of the enzyme, particularly the acyl pocket, affecting the affinity of the enzymes for inhibitor. In *M. musculus* AChE, the acyl pocket mutation F295L (F288L) increases $k_i$ 90-fold, F297I (F290I) 200-fold, and the double mutant, over 500-fold for iso-OMPA [14]. In *H. sapiens* AChE, similar, although more complex, results were seen for the less bulky organophosphates, diisopropyl phosphorofluoridate (DFP), diethyl phosphorofluoridate (DEFP), and paraoxon [64]. Replacement of aromatic amino acids in the acyl pocket with aliphatic residues increases $k_i$ up to 130-fold, with substitutions at Phe295 (Phe288) having the greater effect. The differences in $k_i$ were primarily due to decreases in $K_d$ with $k_2$ relatively unaffected, suggesting that the substitutions relieved steric interference in the binding of the inhibitors and enhanced enzyme activity.

Figure 4. Concentration dependencies for inhibition of ATCh and PTCh hydrolysis by extract from adult *O. latipes*. Adult fish were extracted in HIS buffer and assayed with ATCh and PTCh in the

Figure 5. Substrate concentration dependencies for hydrolysis of ATCh, PTCh, and BTCh by recombinant BChE from *O. latipes*. Transfected COS-7 cells producing BChE were extracted in HIS buffer and assayed with ATCh (●), PTCh (●), or BTCh (●) and fit as described in Materials and Methods.

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doi:10.1371/journal.pone.0017396.g005
Table 3. Kinetic Parameters for Recombinant ChE from O. latipes. a

<table>
<thead>
<tr>
<th>Substrate</th>
<th>$K_m$ μM</th>
<th>$V_{max}$ mM</th>
<th>$b$</th>
<th>$k_{cat}/k_{m}$ min$^{-1}$</th>
<th>$k_{cat}$ Substrate/ATCh</th>
<th>$k_{cat}/K_m$ M$^{-1}$ min$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATCh</td>
<td>270±18</td>
<td>516±52</td>
<td>0</td>
<td>2.76±0.11×10^4</td>
<td>1.00</td>
<td>1.03±0.05×10^6</td>
</tr>
<tr>
<td>PTCh</td>
<td>156±15</td>
<td>20±12</td>
<td>0.67±0.02</td>
<td>3.07±0.24×10^4</td>
<td>1.11</td>
<td>2.00±0.05×10^6</td>
</tr>
<tr>
<td>BTCh</td>
<td>72±6</td>
<td>39±29</td>
<td>0.92±0.07</td>
<td>0.70±0.03×10^4</td>
<td>0.25</td>
<td>0.99±0.07×10^6</td>
</tr>
</tbody>
</table>

aData are the mean ± SE of 6-8 determinations.

Table 4. IC$_{50}$ Values for Inhibition of Recombinant ChE from O. latipes. a

<table>
<thead>
<tr>
<th>Inhibitor</th>
<th>IC$_{50}$ M</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physostigmine</td>
<td>5.85±0.79×10$^{-8}$</td>
</tr>
<tr>
<td>Ethopropazine</td>
<td>8.98±0.75×10$^{-7}$</td>
</tr>
<tr>
<td>BW284c51</td>
<td>4.80±0.82×10$^{-4}$</td>
</tr>
<tr>
<td>Iso-OMPA</td>
<td>4.58±1.52×10$^{-3}$</td>
</tr>
</tbody>
</table>

aData are the mean ± SE of 3-5 determinations.

doI:10.1371/journal.pone.0017396.t003

doI:10.1371/journal.pone.0017396.t004

doI:10.1371/journal.pone.0017396.g006

affinity, but did not alter the rate of phosphorylation. Unlike
ethopropazine, converse site-directed mutagenesis of the acyl pocket
has not been reported for BChE and iso-OMPA. The atypical BChE
from O. latipes is relatively insensitive to, but is inhibited by iso-
OMPA. This result, as well as the substrate specificity of the enzyme,
is in good agreement with the reduction in size of the acyl pocket,
comparing to BChE, associated with the presence of only one of the
two aromatic residues in the subsite in our molecular modeling. This
explanation, as well as the others offered on the molecular basis of
substrate and inhibitor specificity could be tested by site-directed
mutagenesis. In any event, it should be kept in mind that vertebrate
ChEs cannot always be characterized as AChE or BChE simply on
the basis of diagnostic inhibitor specificities [53], and that molecular
analysis can provide additional valuable information about the
nature of ChE activity. Since molecular data are not available for the
atypical BChEs from T. marmorata, P. flesus, P. platessa, and A.
dussummeri, we cannot compare at the molecular level the atypical
BChE from O. latipes with these enzymes.

Comparison of the Molecular Form of O. latipes Atypical
BChE with AChE, BChE, and other Atypical BChEs in
Vertebrates

The atypical BChE from O. latipes is a GPI-anchored G$_2^a$
membrane-bound dimer and thus resembles a BChE$_{H}$, possessing
an H-type C-terminus, which is removed upon the addition of the
GPI anchor. The atypical BChE of P. flesus is also a GPI-anchored
G$_2^a$ [19]. However, the atypical BChEs from T. marmorata and A.
dussummeri are BChE$_{T}$ types since they are capable of assembling
into G$_2$, G$_3$, and G$_4$ molecular forms, and are able to interact with
the collagenic tail protein ColQ [17,20]. The forms of the P.
platessa enzyme are not known. While many vertebrate AChEs,
including Torpedo spp. and mammalian AChE are found as
AChE$_{T}$ and AChE$_{H}$ as the result of alternative splicing, all
vertebrate BChEs are considered to be of the T-type [26]; the
atypical BChEs from O. latipes and P. flesus are the only certain
exceptions to this rule. The nature of the ChE activity in Xenopus
spp. is perplexing. A PIPLC-sensitive G$_2$ AChE$_{H}$ has been
reported in X. laevis muscle [34,41], even though its substrate
specificity is characteristic of BChE [41], but the enzyme has been
classified as an AChE on the basis of diagnostic inhibition [39,40].
However, the ChE in X. laevis tadpoles was found to be resistant to
various carbamate and organophosphate inhibitors and to not
show inhibition by excess substrate [42]. Moreover, the genome
project for X. tropicalis indicates that only AChE$_{T}$ sequences and
not AChE$_{H}$ sequences are present. In striking contrast though, a
BChE$_{H}$ sequence is found. The H-peptide is widespread in the
AChEs of invertebrates, although these sequences are not
homologous among the invertebrates nor with the vertebrates,
BChE sequence is found. The H-peptide is widespread in the
AChEs of invertebrates, although these sequences are not
homologous among the invertebrates nor with the vertebrates,
containing only a few functional similarities: a cysteine near the
carboxyl terminal that mediates inter-chain disulfide bonding in
the dimer, and a GPI anchor signal consisting of an
attachment site followed by a non-conserved stretch of hydropho-
bic amino acids [51]. These variant kinetic, pharmacologic, and
molecular form data among the atypical BChEs raise questions
about their evolution and the evolution of AChE and BChE in the
vertebrates.

The Evolution of AChE and BChE

In terms of substrate and inhibitor specificity and inhibition, it is
tempting to speculate that the atypical BChE of O. latipes and the
other atypical BChEs discussed represent a transitional form of

Figure 6. Concentration dependencies for inhibition of ATCh
hydrolysis by recombinant BChE from O. latipes. Transfected COS-7
cells producing BChE were extracted in His buffer and assayed with
ATCh in the presence of various concentrations of inhibitors. Extracts
were incubated with inhibitor for 20 minutes prior to being assayed for
activity: physostigmine ( ), ethopropazine ( )
BW284c51 ( ), iso-
OMPA ( ). Data were fit to a three-parameter logistic function as
described in Materials and Methods.

doI:10.1371/journal.pone.0017396.g006

An Atypical Cholinesterase from Medaka
ChE as BChE gradually evolved from an ancestral AChE in the vertebrates subsequent to a gene duplication event early in vertebrate evolution. Assuming Darwinian gradualism, one would expect that if AChE were the ancestral vertebrate ChE, with two Phe residues in its acyl pocket, that upon gene duplication the two residues would be replaced sequentially with an intermediate enzyme having one of the Phe residues and decreased substrate specificity, a property that is consistent with the putative role of BChE in detoxification mechanisms and our data [1,2,65–67]. Subsequently, under selection pressure, the enzyme would lose its second acyl pocket Phe residue to obtain the substrate specificity of BChE in the higher vertebrates. Likewise, the atypical BChEs show different intermediate patterns of substrate inhibition or lack thereof, which could be interpreted as a transition to the complete loss of substrate inhibition with all substrates, and the acquisition of substrate activation, also a property consistent with the detoxifying role of BChE. While substrate inhibition may be physiologically relevant for the role of AChE in synaptic transmission, it would not be adaptive toxicologically. The issue of inhibitor sensitivity is more complicated. One might expect selective pressures on a detoxifying enzyme to be directed towards increased resistance to inhibitors. While non-aromatic substitutions decrease the sensitivity of the enzyme to some inhibitors, they increase the volume of the catalytic gorge in BChE, literally opening it up to inhibition by bulkier inhibitors that cannot gain access to AChE. Thus, there may be a tradeoff between decreased substrate specificity and substrate inhibition, and decreased inhibitor inhibition with the maximization of all three impossible. By contrast, the larger gorge and increased sensitivity to some inhibitors allows BChE to act as a stoichiometric scavenger of natural and man-made carbamate and organophosphate inhibitors [2]. Clearly, other non-aromatic substitutions in the acyl pocket, in other subsites in the catalytic gorge, and in the peripheral site, also producing kinetic and pharmacological differences between the two enzymes have occurred in this transition as the two enzymes diverged structurally and functionally subsequent to gene duplication [9,54,68,69].

Because of the possibility of three extensive (even genome-wide) gene duplication events early in vertebrate evolution at (1) the origin of the vertebrates, (2) the emergence of the jawed fish, and (3) the appearance of the ray-finned fishes [70–72], the timing of the putative gene duplication event producing vertebrate BChE from AChE is uncertain. The jawless vertebrates, the lamprey *P. marinus* and the hagfish *M. glutinosa*, appear to have only one ChE, AChE [24,73], and there is evidence for two ChEs in the cartilaginous jawed fish *T. marmorata*, AChE, and an atypical BChE [17]. Therefore, it is possible that a duplication of an ancestral AChE gene accompanied the wide-spread gene duplication event that coincided with the emergence of the jawed fish, with the atypical and typical BChEs of bony fish and other derived vertebrates descendents of this gene. However, given the genome duplication event in the ray-finned fish lineage, it is possible that these atypical BChEs are unique to the ray-finned fish and not ancestral to the BChE of higher vertebrates, as the land vertebrates descended from the lobe-finned fishes, represented today by the lungfish and the coelacanth, which diverged from the ray-finned fish prior to the duplication event [74]. Currently there is not any information about ChEs from the coelacanth genome project, but such information could shed light on the timing of the gene duplication event and the evolutionary origins of BChE in tetrapods. In any case, for the time being, the atypical BChE of *O. latipes* can serve as a model for the evolutionarily intermediate ChE between AChE and BChE.

Given that the poorly conserved H-transcripts are widespread in the AChEs of invertebrates, while in vertebrates they are present only in the AChE of the elasmobranch *Torpedo* spp., and mammals, and apparently in a BChE in the amphibian *Xenopus* spp., but not reported in reptiles or birds, Massoulé et al. [26] stated that sequences characteristic of a GPI-addition signal were “invented” on several occasions during evolution. We can now add the presence of the H-peptide in the atypical BChEs of the teleosts *P. flesus* and *O. latipes*. Combes et al. [73] speculated that the conservation of splice sites at the C-terminus of AChE in insects, nematodes, and vertebrates suggests that exon shuffling has occurred at the 3’end of ChE genes at various times throughout evolution. Such shuffling is one mechanism for appearance of an alternatively spliced exon [76]. Another explanation is exonization [77,78]; indeed, the presence of read-through or R-transcripts [79–
in AChE could be considered a nascent or abortive exonization event, and the low abundance of such R-transcripts is consistent with an exonization process [78]. Thus, it is possible that the evolution of the C-termini of ChEs is independent of the evolution of the catalytic subunit. Further discussion of the evolution of ChEs can be found in Pezzementi and Chatonnet [82].

Ecotoxicological Implications
Pesticide use is a major concern in aquatic environments, where runoff from agricultural and urban ecosystems impacts their ecology. The major pesticides in use today are organophosphate and carbamate acetylcholinesterase inhibitors, and pyrethroid ion-channel agents [83,84]. These toxins are transported from terrestrial to aquatic ecosystems, placing the latter at risk; thus, it is important to understand the effects of these compounds on the resident vertebrate and invertebrate fauna [85]. Inhibition of ChE activity in fish generally correlates with mortality. There may also be sub-lethal behavioral and physiological effects, including reduced swimming ability, and altered feeding and social behavior. However, there are marked species differences in these effects [86]. These differences could be due to different levels of AChE and BChE in the nervous system.

Figure 9. Active site gorges of H. sapiens BChE (hBChE; pdb code 1p0i), T. californica AChE (TcAChE; pdb code 1ea5) and a homology model of O. latipes atypical BChE (OlBChE). The side chains of key residues lining the gorges are represented as sticks (carbon in orange, oxygen in red, and nitrogen in blue). The top of the gorge and the entrance to and exit from the enzyme are shown at the top of the figures. The acyl pocket, which accommodates the acyl group of substrates, is comprised of residues F288 and F290 in T. californica AChE, L286 and V288 in H. sapiens BChE, and L291 and F293 in O. latipes AChE. The central residue of the choline binding site at the bottom of the gorge is W84 in T. californica AChE, W82 in H. sapiens BChE, and W83 in O. latipes atypical BChE. Substrates are bound between these two subsites. The solvent accessibility surface of the gorge was calculated by taking into account highly conserved structural water molecules, using the software "HOLLOW" and represented as grey dots [99].

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An Atypical Cholinesterase from Medaka

Table 5. Comparison of Available Kinetic and Pharmacological Parameters and Splice Variants of Vertebrate AChE, BChE, and Atypical BChEs.

<table>
<thead>
<tr>
<th>Parameter/Species</th>
<th>T.marmorata AChE*</th>
<th>O.latipes BChE</th>
<th>P.fresus BChE*</th>
<th>P.platessa BChE*</th>
<th>A.dussumieri BChE*</th>
<th>T.marmorata BChE*</th>
<th>H.sapiens BChE</th>
</tr>
</thead>
<tbody>
<tr>
<td>$K_m$ ATCh mM</td>
<td>0.05</td>
<td>0.27</td>
<td>2.20</td>
<td>0.90</td>
<td>1.24</td>
<td>0.4</td>
<td>1.4*</td>
</tr>
<tr>
<td>$K_m$ BTCh mM</td>
<td>0.2</td>
<td>0.16</td>
<td>1.03</td>
<td>0.15</td>
<td>0.24</td>
<td>0.15</td>
<td>0.97*</td>
</tr>
<tr>
<td>$V_{max}$ ATCh/µM</td>
<td>N.M.†</td>
<td>0.07</td>
<td>0.32</td>
<td>0.14</td>
<td>0.06</td>
<td>0.05</td>
<td>0.91*</td>
</tr>
<tr>
<td>Substrate inhibition†</td>
<td>+A, −P</td>
<td>+AP, −B</td>
<td>+APB, −AP, +B</td>
<td>−AP, +B</td>
<td>+APB</td>
<td>−APB</td>
<td>−APB</td>
</tr>
<tr>
<td>IC₅₀ Ethopropazine µM</td>
<td>158</td>
<td>0.90</td>
<td>12.6</td>
<td>N.D.‡</td>
<td>N.D.‡</td>
<td>100</td>
<td>15.3†</td>
</tr>
<tr>
<td>IC₅₀ BW284C51 µM</td>
<td>0.04</td>
<td>480</td>
<td>63</td>
<td>0.79</td>
<td>Weak</td>
<td>100</td>
<td>651‡</td>
</tr>
<tr>
<td>Splice Variant†</td>
<td>H, T-type</td>
<td>H-type</td>
<td>H-type</td>
<td>N.D.‡</td>
<td>T-type</td>
<td>T-type</td>
<td>T-type</td>
</tr>
</tbody>
</table>

*Data are from Toutant et al., [17]; the IC₅₀ values are estimates.
†Data are from Davies et al. [19].
‡Data are from Ludin and represent titrometric measurement of the hydrolysis of the oysterse rather than the titoester of substrates [18].
§Data are from Leibel and inhibition by BW284C51 is based on quantitative observations of the inhibition of ATCh staining of enzyme following non-denaturing gel electrophoresis [21]. Splice variant data are from Leibel [20].
∥Data are from Davies et al. [100].
¶Data are from Ucar et al. [101].
Data are from Loewenstein-Lichtenstein et al.; IC₅₀ was back calculated from a calculated $K_i$ [69].
N.M. not measureable.
N.D. not determined.
Splice variant determined by molecular forms present and sequence data when available.

doi:10.1371/journal.pone.0017396.t005

Materials and Methods

Ethics Statement

All animal procedures were conducted in strict adherence to the European Council Directive of November 24, 1986 (86-609). Approval for this study was provided by Comité Regional d’Etique Languedoc Roussillon C34-172-10.

Materials

Dulbecco’s modified Eagle medium, fetal bovine serum, Opti-MEM medium, and phosphatidylinositol-specific phospholipase C (PIPLC) were purchased from Invitrogen, Carlsbad, California. FuGene was obtained from Roche, Indianapolis, Indiana. Acetylthiocholine (ATCh), butyrylthiocholine (BTCh), propionylthiocholine (PTCh), [4-5-[4-(dimethyl-prop-2-enyl-ammonio)-phenyl]-3-oxo-pentyl]phenyl]-dimethyl-prop-2-enyl-azanium di-bromide (BW284c51), 5-(3-carboxy-4nitro-phenyl)disulfanyl-2-phenyl-[3-oxo-pentyl]phenyl]-dimethyl-prop-2-amine (iso-OMPA), 10-(2-diethylaminopropyl) phenothiazine hydrochloride (ethopropazine), and (3aS-cis)-1,2,3,3a,8,8a-hexahydro-1,3a,8-trimethyl-phenyloxy-[3,3-b-jindol-3-ol methylcarbamate (physostigmine) were purchased from Sigma, St. Louis, Missouri. The high-affinity ligand 7-[(dithioethoxy)phosphoryloxy]-1-methylquinolinium iodide (DEPQ) was a gift from Yaacov Ashani.

O. latipes adults and embryos were from facilities at INRA. Fish were maintained at 28°C on a 13-h light/11-h dark cycle.

cDNA cloning and sequence analyses

The cDNA clone Ola.23452 (Genbank AV668390) from an O. latipes library was obtained from UniGene. The clone was extended by PCR on the basis of the gene structure on scaffold23502 in UCSC genome project, and the full cDNA sequence (Genbank GU797251) was cloned into the expression vector pCMV SPORT 6.1 (Invitrogen).

Sequences were aligned with Clustal W or Clustal X for molecular modeling or phylogenetic analysis by the neighboring method [89]. Putative cleavage/attachment sites and downstream stretches of hydrophobic amino acids of H-type C-termini of ChEs were predicted with PredGPI [89] and ProtScale [90], respectively.

In vitro expression and extraction of enzyme

COS-7 monkey cells (American Type Culture Collection) were grown in Dulbecco’s modified Eagle medium containing 10% fetal calf serum. Cells were plated at a density of 2.5×10⁵ cells/75 cm² culture flask, incubated overnight, and transferred to OptiMEM medium. FuGene was then used to transfect the cells with 7.8 µg of DNA. The cells were then incubated for 48 h at 37°C before the medium was removed and the cells extracted in high ionic strength (HIS) buffer: 10 mM NaHPO₄, pH 7, 1 M NaCl, 1% Triton X-100, 1 mM EDTA. Extracts were centrifuged at 20,000 g for 20 min, and the supernatants were assayed for ChE activity.

Measurement and analysis of BChE activity and inhibition

AChE activity was measured according to the method of Ellman et al. [91] as modified by Doctor et al. [92] in 100 mM NaHPO₄,
Menten kinetics is observed. The kinetic parameters $K_m$, $V_{max}$, and $b$, and $V_{max}$, were determined by using SigmaPlot to fit the data to the equation below as described by Radić et al. [13] and Kaplan et al. [9]. $K_c$ is the dissociation constant for the binding of substrate to a second site on the enzyme, and the parameter $b$ indicates the relative catalytic efficiency of the SES complex compared to SE. If $b < 1$, the enzyme shows substrate inhibition; if $b > 1$, the enzyme shows substrate activation, and if $b = 1$, Michaelis-Menten kinetics is observed.

$$v = \frac{1+b[S]/K_c}{1+S/K_c} \cdot \frac{V_{max}}{1+K_m/S}$$

The turnover number $k_{cat} (V_{max}/[Enzyme])$ was determined by enzyme titration with DEPQ [93] as described previously [63]. Values of IC$_{50}$ for the inhibitors were used by determining IC$_{50}$ with various concentrations of drug for 20 min and then assaying for enzyme activity in the presence of AChe. SigmaPlot was then used to fit the data to a three-parameter logistic function, yielding IC$_{50}$. Biphasic inhibition curves were fit with GOSAfit. Since we were just looking for classical diagnostic differential inhibition, it was not necessary to determine $k_c$ or $K_V$ values for the inhibitors [4,24,52].

**Velocity Sedimentation on Sucrose Gradients; PIPLC Digestion**

The molecular forms of ChE were analyzed by velocity sedimentation in 5–25% isokinetic sucrose gradients prepared in HIS buffer (with or without Triton X-100) containing 1 mg/ml bovine serum albumin. Sedimentation was in an SW 41 rotor at 30,000–37,000 rpm for times satisfying the equation $[(rpm)^2 \times t \ (h)] = 2.5 \times 10^{10}$ as described previously [24]. Apparent sedimentation coefficients were calculated relative to the sedimentation of catalase (11.3 S). Data were plotted as fractional activity of total ChE activity on the gradient as a function of sedimentation coefficient: fractional activity on gradient $= \text{activity in a given fraction/total activity on gradient}$; sedimentation coefficient $= \text{fraction number} / (11.3 \text{S/fraction number of catalase peak})$.

PIPLC digestion was performed by a modification of the method of Gibney et al. [94]. These enzyme assays were performed in Ellman’s solution prepared in phosphate-buffered saline.

**Molecular Modeling**

Sequence alignment was performed with ClustalX 1.04 and homology modeling was carried out using MODELLER 9v3 program [95]. MODELLER can implement comparative protein structure modeling by satisfying spatial restraints in terms of probability density functions. A 3D structural model of medaka AChE was built by using the crystal structures of H. sapiens BChE (pdb code 1p0i), T. californica AChE (pdb code 1e5) and D. melanogaster AChE (pdb code 1dx4) as structural templates. Structural water molecules that are conserved among the three templates were also retained in the modeling procedure. A series of 200 runs of MODELLER were carried out using standard parameters, and the outcomes were ranked on the basis of the internal DOPE scoring function. The model with the highest score was chosen as the candidate model. Then, energy minimization was performed using GROMACS 4.05 according to the software protocol [96]. The final energy-minimized model and the templates were aligned using Theseus [97] and analyzed in PyMOL [98]. Active site gorge surfaces and volumes were calculated using the software HOLLOW 1.1 [99] and taking into account highly conserved structural water molecules. For each structure, the active site entrance forms a bottleneck that was used to delimit the volume of the gorge in the calculations.

**Supporting Information**

**Figure S1** The pdb dataset for the model of *O. latipes* BChE.

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**Author Contributions**

Conceived and designed the experiments: AC FN LP. Performed the experiments: AC FN LP. Analyzed the data: AC FN LP. Contributed reagents/materials/analysis tools: AC FN LP. Wrote the paper: AC FN LP.

**References**


