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Crystal Structure of Mouse Acetylcholinesterase

A PERIPHERAL SITE-OCCLUDING LOOP IN A TETRAMERIC ASSEMBLY

Yves Bourne‡, Palmer Taylor§, Pierre E. Bougis¶, and Pascale Marchot\n
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The crystal structure of mouse acetylcholinesterase at 2.9-Å resolution reveals a tetrameric assembly of subunits with an antiparallel alignment of two canonical homodimers assembled through four-helix bundles. In the tetramer, a short O loop, composed of a cluster of hydrophobic residues conserved in mammalian acetylcholinesterases along with flanking a-helices, associates with the peripheral anionic site of the facing subunit and sterically occludes the entrance of the gorge leading to the active center. The inverse loop-peripheral site interaction occurs within the second pair of subunits, but the peripheral sites on the two loop-donor subunits remain freely accessible to the solvent. The position and complementarity of the peripheral site-occluding loop mimic the characteristics of the central loop of the peptidic inhibitor fasciculin bound to mouse acetylcholinesterase. Tetrameric forms of cholinesterases are widely distributed in nature and predominate in mammalian brain. This structure reveals a likely mode of subunit arrangement and suggests that the peripheral site, located near the rim of the gorge, is a site for association of neighboring subunits or heterologous proteins with interactive surface loops.

Acetylcholinesterase (AChE),\(^1\) a member of the family of proteins with an \(\alpha/\beta\)-hydrolase fold (1, 2), rapidly terminates cholinergic neurotransmission by hydrolysis of acetylcholine (3, 4). In mammals, AChE is encoded by a single gene, yet a multiplicity of molecular forms arise through alternative mRNA processing and association of the catalytic subunits with structural subunits. Alternative mRNA processing gives rise to three splicing options: (a) a soluble monomer without the capacity for disulfide linkage; (b) a glycoprophospholipid-linked form that is membrane-associated and exists as monomers and dimers; and (c) an amphiphilic form found as monomers, dimers and tetramers (in the tetramer, amphiphilic character may be lost presumably through occlusion of the hydrophobic surfaces). Further structural complexity is achieved in the tetrameric assemblies through disulfide association of the catalytic subunits with structural subunits that are either amphiphatic or collagen-like, giving rise to predominant forms in brain and skeletal muscle. The tetrameric forms are dimers of dimers where one set of disulfides links with a polyproline-containing structural subunit (5) and the other set is formed between monomers (3, 4). However, there is currently little structural information about the subunit orientation in the tetramer and the association of tetramers with structural subunits.

Differences in the molecular forms of the cholinesterases are the primary determinants of their cellular disposition (3). Abnormal associations of AChE are apparent in Alzheimer’s dementia, where a selective loss of the amphiphilic tetramers is observed (6, 7). AChE was suggested to associate with or to nucleate the neuritic plaques found in the disease (8–12); however, the mode of this association is unknown.

The active center of AChE, which consists of the triad Ser\(^{203}\), Glu\(^{334}\), His\(^{447}\) in mammals (13), is nearly centrosymmetric to the subunit and is located at the bottom of a narrow gorge (14). Inhibitors may bind at the active center or at a distant allosteric site, the peripheral anionic site, located at the gorge rim. The structure of recombinant monomeric mouse AChE (mAChE) in a complex with the peptidic inhibitor fasciculin 2 (Fas2), bound to the peripheral anionic site, provided the first mammalian cholinesterase template (15). Only slight differences were observed in the conformation of Fas2-associated mAChE compared with uncomplexed Torpedo californica AChE (TeAChE) (14). However, it was not possible to distinguish species-related differences from changes in conformation accompanying Fas2 binding. Indeed, little difference in TeAChE conformation was observed between structures of the apoenzyme (14) and the Fas2 complex (16). In addition, currently available crystal structures of AChE reveal the entrance of the active site gorge to be either occluded by a symmetry-related molecule (14, 17, 18) or sealed with high surface complementarity by Fas2 (15, 16), therefore precluding structural analysis of an unprotected peripheral anionic site region.

The crystal structure of mAChE refined to 2.9-Å resolution has good stereochemistry, with an \(R\)-factor of 21.5\% for data in the 15 to 2.9-Å resolution range, and contains four mAChE...
monomers assembled as a tetramer, three GlcNAc moieties and one GlcNAc-β1,4-GlcNAc-α1,6-Fuc trisaccharide moiety linked to Asn residues, four decamethonium (DECA) molecules, four phosphate groups, four glycerol molecules, and 205 water molecules in the asymmetric unit. The structure of mAChE provides significantly improved accuracy in the positions of the main chain and side chains of the molecule over the Fas2mAChE structure, reveals distinctive features of the mouse enzyme, and permits direct comparison of solvent-exposed and occluded peripheral anionic sites within the same crystal unit. More importantly, this structure highlights surface determinants that could participate in formation of oligomeric structures that could participate in formation of oligomeric units A and D) and residues Glu1–Cys257 and Asn265–Ala547 (subunits B and Thr 436 oxygen atoms and the carbon atoms in van der Waals interactions with the side chain of Met 85. A phosphate group arising from the crystallization liquor was positioned between the Lys 332 and Arg395 side chains, with its oxygen atoms bound to the Trp 442 carbonyl site (see “Results”). A glycerol molecule arising from the cryoprotection solution was positioned into a density found in the vicinity of Glu81, side chains were adjusted, and the Pro 258–Gly264 portion, which was missing in the search model, was built into 2Fo – Fa difference electron density maps. In each of the subunits in the tetramer, a DECA molecule was fitted into a residual density observed within the active site (see “Results”). A glycerol molecule arising from the cryoprotection solution was positioned into a density found in the vicinity of Glu41, with the glycerol oxygen atoms hydrogen-bonded to the Glu652, Glu41, and Thr455 oxygen atoms and the carbon atoms in van der Waals interactions with the side chain of Met55. A phosphate group arising from the crystallization liquor was positioned between the Lys372 and Arg905 side chains, with its oxygen atoms bound to the Trp442 carbonyl oxygen and Lys385 nitrogen atoms. Since the non-crystallographic symmetry (NCS) was not restrained throughout these early stages of model building and refinement, subunits were then superimposed within each dimer, and different NCS constraints were applied along the molecule and used during all subsequent refinement steps. This model was refined with REFMAC (27), including all low resolution data. A conservative number of solvent molecules were manually added into the model; most of them were found to be located at nearly identical positions in the 2.5-Å resolution structure of TcAChE (Protein Data Bank 1MAH) (15) as a search model with the AMoRe program package (24). Four mAChE subunits were positioned within the asymmetric unit (correlation = 75.2%, R-factor = 29.9% in the 15 to 4.5-Å resolution range) and found to assemble as a dimer of the same dimers as seen in the Fas2mAChE structure. Rigid-body refinement applied to the whole tetramer, each of the two dimers, and the individual subunits was then performed with X-PLOR (25) using synchrotron data between 8 and 3 Å and gave an R-factor of 33%. For 2% of the reflections against which the model was not refined, Rfree was 33%. Several cycles of Powell conjugate-gradient minimization were then performed, and electron density maps calculated with the subsequent model were inspected with TURBO-FRODO (26).

The positions of misplaced amino and carboxyl termini and of a few side chains were adjusted, and the Pro352–Gly364 portion, which was missing in the search model, was built into 2Fo – Fa, Fo – Fa, electron density maps. In each of the subunits in the tetramer, a DECA molecule was fitted into a residual density observed within the active site (see “Results”). A glycerol molecule arising from the cryoprotection solution was positioned into a density found in the vicinity of Glu41, with the glycerol oxygen atoms hydrogen-bonded to the Glu652, Glu41, and Thr455 oxygen atoms and the carbon atoms in van der Waals interactions with the side chain of Met55. A phosphate group arising from the crystallization liquor was positioned between the Lys372 and Arg905 side chains, with its oxygen atoms bound to the Trp442 carbonyl oxygen and Lys385 nitrogen atoms. Since the non-crystallographic symmetry (NCS) was not restrained throughout these early stages of model building and refinement, subunits were then superimposed within each dimer, and different NCS constraints were applied along the molecule and used during all subsequent refinement steps. This model was refined with REFMAC (27), including all low resolution data. A conservative number of solvent molecules were manually added into the model; most of them were found to be located at nearly identical positions in the 2.5-Å resolution structure of TcAChE (Protein Data Bank 1MAH) (15) as a search model with the AMoRe program package (24). Four mAChE subunits were positioned within the asymmetric unit (correlation = 75.2%, R-factor = 29.9% in the 15 to 4.5-Å resolution range) and found to assemble as a dimer of the same dimers as seen in the Fas2mAChE structure. Rigid-body refinement applied to the whole tetramer, each of the two dimers, and the individual subunits was then performed with X-PLOR (25) using synchrotron data between 8 and 3 Å and gave an R-factor of 33%. For 2% of the reflections against which the model was not refined, Rfree was 33%. Several cycles of Powell conjugate-gradient minimization were then performed, and electron density maps calculated with the subsequent model were inspected with TURBO-FRODO (26).

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and C) (see Figs. 1 and 2). High temperature factors and weak electron density include mAChE residues Glu\(^{-}\)–Gln\(^{-}\) and Arg\(^{245}\)–Pro\(^{286}\) in all four subunits; Pro\(^{272}\)–Gly\(^{284}\) is not visible in electron density maps of subunits B and C. The r.m.s. deviation between the two mAChE subunits of a canonical dimer is 0.06 Å for tight-constraint NCS atoms, 0.16 Å for medium-constraint NCS atoms, and 0.64 Å for loose-constraint NCS atoms. Structures B and C were solved using the refined structure A (inhibitor and solvent molecules removed) as an initial model and refined to 3.1 Å. The r.m.s. deviation values for the backbone atoms and all atoms are 0.21 and 0.3 Å between structures A and B, 0.13 and 0.2 Å between structures A and C, and 0.19 and 0.26 Å between structures B and C, respectively. The stereochemistry of all three structures was analyzed with PROCHECK (29) and WHATIF (30) (Table I). Figures were generated with the programs RIBBON (31), GRASP (32), and TURBO-FRODO (26).

RESULTS AND DISCUSSION

Overall Description of the Structure

The mAChE molecule, which consists of a 12-stranded central-mixed \(\beta\)-sheet surrounded by 14 \(\alpha\)-helices, has the same overall conformation as found for the Fas2–mAChE complex (15): the r.m.s. deviations between the free and complexed mChEs are 0.5 Å for the backbone atoms and 0.6 Å for all atoms, with the largest deviations occurring in surface loops (Ala\(^{24}\)–Gly\(^{25}\), Pro\(^{106}\)–Ser\(^{110}\), Glu\(^{225}\)–Leu\(^{224}\), Asp\(^{272}\)–Ala\(^{274}\), Leu\(^{338}\)–Pro\(^{341}\), and Asp\(^{381}\)–Ser\(^{387}\)), in the position of domain Tyr\(^{341}\)–Glu\(^{399}\), and in the orientation of Tyr\(^{337}\).

mAChE, which is devoid of the carboxyl-terminal amphiphatic helix and intersubunit disulfide-linking Cys, is monomeric in dilute solution (19), but dimeric in the Fas2–mAChE crystals, where a homodimer assemble through a tightly packed four-helix bundle (15). In the mAChE crystal, two identical homodimers assemble as a tetramer in which the two four-helix bundles are aligned antiparallel (Fig. 1). The main axes of the two dimers are tilted by \(\sim 35^\circ\) from each other to form a compact, pseudo-square planar tetramer with overall dimensions of 90 \(\times\) 90 \(\times\) 100 Å.

In the tetramer, a short surface loop and flanking \(\alpha\)-helices from subunit A of the first dimer tightly associate with the peripheral site region at the gorge entrance of the facing subunit, subunit C of the second dimer (Fig. 1). The inverse loop-peripheral site interaction involves the loop and \(\alpha\)-helices of subunit D in the second dimer and the peripheral site region of subunit B in the first dimer. As a result, peripheral sites of the two loop-acceptor subunits, B and C, are occluded, whereas peripheral sites of the two loop-donor subunits, A and D, are accessible to the solvent, a feature unique to this structure. At the tetramer center, the carboxyl-terminal \(\alpha\)-helices in the four-helix bundles converge at the interface. The dimer-dimer interface, which extends over 90 Å in a direction roughly perpendicular to the four-helix bundle axis, buries to a 1.6-Å probe radius, a 2500-Å\(^2\) surface area for the two pairwise dimer-dimer interfaces, an area falling in the highest range for functionally relevant crystal packing interfaces (33). Including the area buried at the four-helix bundle interface within each dimer (15), the total mChE surface area buried within the tetramer encompasses 6600 Å\(^2\).

Detailed Description of the Structure (Structure A)

At the dimer-dimer interface, mAChE loop 257–272, a short \(\Omega\) loop (34) that is bridged by two Cys residues and contains the Pro–Pro–Gly–Gly–Ala–Gly–Gly sequence conserved in mammalian AChEs along with flanking \(\alpha\)-helices \(\alpha_6^6\) and \(\alpha_7^7\) (Fig. 2), is associated with the peripheral anionic site at the gorge entrance of the facing mAChE subunit. The helix-loop-helix domain sterically occludes access to the active center gorge (Figs. 1–3). This part of the interface involves hydrophobic and polar contacts of 12 residues from each subunit (Fig. 3). The Gly\(^{260}\)–Gly\(^{264}\) cluster of hydrophobic residues at the loop tip packs against peripheral site residues Tyr\(^{72}\), Trp\(^{286}\), and Tyr\(^{441}\), with predominant interactions between dipeptide Gly\(^{260}\) Gly\(^{261}\) and Trp\(^{286}\) and between dipeptide Gly\(^{261}\) Ala\(^{262}\) and Tyr\(^{441}\). Two discrete patches of polar interactions involve the loop extremities and flanking helices with the boundary of the peripheral anionic site and appear to form three hydrogen bonds. The guanidinyl moieties of Arg\(^{245}\) and Arg\(^{253}\), located near the amino and carboxyl termini of helix \(\alpha_6^6\), respectively, form key hydrogen bonds with the side chain carboxylate and the amide backbone nitrogen atom of Gly\(^{263}\), and the Gly\(^{263}\) nitrogen forms a hydrogen bond with the Tyr\(^{341}\) carboxyl oxygen. The loop shows high complementarity to the peripheral anionic site, a feature arising from the flexibility imparted by the vicinal Gly residues; limited internal stabilization of conformation is achieved from the adjacent Pro residues, with Pro\(^{258}\) in the cis-conformation and in van der Waals contacts with Ala\(^{262}\). Flexibility of the loop is evident since it cannot be
seen in the absence of external stabilization, as found for subunits B and C in the tetramer (Fig. 1) and in the Fas2-mAChE complex at 3.2-Å resolution (15); only the Pro doublet is seen in the Fas2-mAChE complex at 2.7-Å resolution.2

The second region of the dimer-four-helix interface is located centrosymmetric to the tetramer where the two four-helix bundles converge (Fig. 1). The carboxy-terminal region, not seen in the Fas2-mAChE structure, is ordered sufficiently to be seen in subunits B and C, with only the carboxy-terminal Pro548 being disordered. The two Thr543 residues at and near the carboxy-terminal ends of helices a10 of subunits A and C, respectively, face each other and are separated by only 7.5 Å. The segment from Ala544 to Ala547 is not part of helix a10; instead, it deviates from the helix axis and exits the plane of the tetramer, perhaps because of charge repulsion between Glu546 residues of neighboring subunits in the tetramer. Accordingly, the carboxy-terminal helix a10 ends at Thr543, the last residue encoded by exon 4 before the region of alternative splicing (13, 19).

GlcNAc moieties are observed in two of the three consensus sequences for N-linked glycosylation, Asn150 in subunit A, also seen in the Fas2-mAChE complex (15), and Asn265 in subunits A and D, although the density is weaker. In addition, a GlcNAc-β1,4-GlcNAc-α1,6-Fuc trisaccharide moiety is linked to Asn350 in subunit D (Figs. 1 and 2) (35, 36).

Comparison of the mAChE and Fas2-mAChE Structures

The structure of mAChE at 2.9-Å resolution provides greater delineation of the positions of the α-carbon and side chains than the Fas2-mAChE complex (3.2 Å) (15) and establishes several distinctive features of mAChE compared with TcAChE (Protein Data Bank 2ACE) (28). Thus, the conformations of the amino- and carboxyl-terminal segments were resolved, and the mAChE-specific positioning of the Met85 side chain in its stacking interaction with the indole of Trp36 was confirmed. The Tyr419-Ser399 rigid-body motion that was found when comparing the Fas2-mAChE and TcAChE structures is not apparent in mAChE; hence, movement of this domain is induced by Fas2 association (15).

The most prominent feature arising upon superimposition of the structures is the close resemblance of the two peripheral site-occluding loops, the short Ω loop of mAChE subunit A and the central loop, loop II, of Fas2 bound to mAChE, in their positions and surface complementarity (Fig. 4). The mAChE loop and the tip of Fas2 loop II, although positioned roughly perpendicular with their respective Pro doublets non-superimposable, overlap at the gorge entrance. Several side chains and backbone carbons of the mAChE loop mimic the side chains of Fas2 loop II in their interactions with residues in the peripheral anionic site. In particular, interaction of Gly263 in mAChE subunit A with Trp266 at the gorge entry of mAChE subunit C mimics the key interaction of Met225 in Fas2 with Trp256 in Fas2-associated mAChE; backbone atoms of Gly263-Gly264 in mAChE align with the side chain of Lys32 in Fas2; and Gly260
in mAChE establishes the same van der Waals contacts with Tyr72 as does Fas2 Leu48. Several residues in helix α6,7 of mAChE subunit A also adopt positions similar to residues in bound Fas2: the side chain of Arg255 in mAChE mimics, in both position and orientation, the side chain of Arg247 in Fas2, and mAChE Gly256 forms the same van der Waals contact in the Fas2-mAChE complex. The mAChE molecular surfaces buried at the loop-peripheral site and Fas2-mAChE interfaces are displayed in gold and brown, respectively. Panel B, close-up view of the similarity between the mAChE short Ω loop and the tip of Fas2 loop II (same color codes as in panel A). Residues belonging to the peripheral site of mAChE subunit C and the side chains arising from the loop region of subunit A are displayed as green bonds with colored spheres, and the side chains arising from Fas2 are displayed as orange bonds with colored spheres. Residues are indicated in black and blue for mAChE subunits A and C, respectively, and in orange for Fas2.

**Fig. 4. Comparison of the mAChE loop-peripheral site interface with the Fas2-mAChE interface.** Panel A, superimposition of the short Ω loop and flanking α-helices in mAChE subunit A (dark blue ribbon, green helices, and red loop) with Fas2 (yellow ribbon with orange tip of loop II) bound to mAChE (light blue molecular surface) according to all C-α atoms of mAChE subunit C in the tetramer and of mAChE in the Fas2-mAChE complex. The mAChE molecular surfaces buried at the loop-peripheral site and Fas2-mAChE interfaces are displayed in gold and brown, respectively. Panel B, close-up view of the similarity between the mAChE short Ω loop and the tip of Fas2 loop II (same color codes as in panel A). Residues belonging to the peripheral site of mAChE subunit C and the side chains arising from the loop region of subunit A are displayed as green bonds with colored spheres, and the side chains arising from Fas2 are displayed as orange bonds with colored spheres. Residues are indicated in black and blue for mAChE subunits A and C, respectively, and in orange for Fas2.

**The mAChE Gorge and Peripheral Anionic Site**

Despite extensive dialysis (19), a DECA molecule is still present, approaching full occupancy in the active site gorges of all four mAChE subunits (Fig. 5). Superimposition of subunit A, the peripheral site of which is free of an occluding loop and totally accessible to the solvent, with TcAChE in the DECA-TcAChE complex (Protein Data Bank 1ACL) (17) shows that the proximal (N-1) and distal (N-2) quaternary ends of DECA are in van der Waals contact with the indole rings of Trp286 in the peripheral site and Trp346 in the active center and superimpose in the two structures. In mAChE, however, the 10-carbon methylene chain adopts an inverted curvature and follows the opposite wall of the gorge, a difference that involves a rotated C-5–C-6 bond and a positioning of the C-7–C-8 bond nearly 5 Å away from its position in TcAChE; hence, the C-7–C-8 portion of the methylene chain winds behind the Tyr341 ring in mAChE. The conformation of DECA in mAChE likely results from the 90° rotation of the Tyr337 phenol ring, which lies perpendicular to the gorge axis, whereas the homologous Phe330 ring in TcAChE lies parallel to the gorge axis (17, 28). The tyrosine hydroxyl, unique to mammalian AChEs, forces a different pathway for the methylene chain by its steric and

**Fig. 5. Stereo views of the 2.9-Å resolution electron density maps of the DECA molecules found in the active site gorges of subunits A and C.** Panel A, superimposition of DECA in subunit A (blue DECA molecule and mAChE side chains) and in the DECA-TcAChE complex (red DECA molecule and TcAChE side chains) showing the omit Fo − Fc map (contoured at 2.5σ) of DECA and surrounding residues in mAChE subunit A. The coordinates of this region (2.5% of total number of atoms) were omitted, and the protein coordinates were refined by simulated annealing before the phase calculation for the map. The peripheral anionic site of subunit A is free of an interacting loop and totally accessible to the solvent. The 90° rotation of mAChE Tyr337 compared with the orientation of TcAChE Phe330 and the change in the position of Tyr341 compared with TcAChE Tyr337 are evident. Panel B, superimposition of DECA in subunit C (orange DECA molecule, yellow mAChE side chains, and red loop from subunit A) and in subunit A (blue DECA molecule and mAChE side chains) showing the αs weighting 2Fo − Fc map (contoured at 1.2σ) of DECA and surrounding residues in subunit C with the loop arising from subunit A (180° from panel A). The 20° tilt of Trp346 is evident. The side chain of Tyr341, which would occlude the figure, is not displayed.
Polar contributions. Also, the Tyr\(^{341}\) ring is shifted toward the center of the gorge, \(-1.5\) Å removed from its position in TcAChE; hence, the hydroxyl moieties on Tyr\(^{341}\) and Tyr\(^{337}\) are separated by 3.8 Å, compared with 3 Å in TcAChE. The orientation of Tyr\(^{337}\) also observed in the Fas2-mmAChE complex (15), could either be intrinsic to the mouse enzyme or arise from complexation with DECA or Fas2; it is, however, identical to that of Phe\(^{330}\) in TcAChE in complexes with various inhibitors other than DECA (17, 28). Hence, fluctuations of configuration of the active center gorge of AChE permit accommodation of a large set of active center ligands.

The mAChE structure provides an initial opportunity to compare exposed and occluded peripheral sites within an asymmetric unit. However, superimposition of loop-donor subunit A with loop-acceptor subunit C does not reveal large differences in conformations of the two sites. r.m.s. deviations are 0.44 and 0.77 Å for the main chain and all atoms, respectively; only the side chain of Trp\(^{286}\) at the gorge entrance of subunit C undergoes a 20° tilt, which may arise from the steric constraint induced by the interacting loop. However, the presence of DECA in the catalytic site of non-occluded subunit A might mask structural rearrangements expected from the unliganded peripheral site. Yet in subunit C, the bound loop partially dissociates DECA, which, although disordered, seems to adopt two discrete conformations with a hinge point near the C-4–C-5 bond. In one conformation, the proximal quaternary end of DECA interacts weakly with the indole ring of Trp\(^{286}\). In the second conformation, this quaternary end is displaced and flips toward the Phe\(^{332}\) and Ile\(^{294}\) side chains, some 5 Å away (Fig. 5). Residual inhibitor was observed in structures of native TcAChE, perhaps even with a tetramethylammonium-eluted enzyme (14, 17, 28). In contrast, DECA is not found in the gorge of the Fas2-mmAChE complex (15), consistent with the inhibitor molecule being displaced by bound Fas2 and eliminated during dialysis of the complex prior to crystallization (19).

**Other Crystal Forms of mAChE**

Structures B and C are essentially identical to structure A. However, structure B contains residual EDR instead of DECA in the active site of all four subunits (data not shown). Comparison of mAChE in this structure with TcAChE in the EDR/TcAChE complex (Protein Data Bank IACK) (17) shows similar overall positioning of the EDR molecule, located between His\(^{447}\), Ser\(^{203}\), Tyr\(^{337}\), and Trp\(^{86}\) in mAChE; however, a 30° tilt of the EDR ring is observed, which may arise from accommodation to the Tyr\(^{337}\) orientation found in mAChE. There is no difference between subunits A and C, indicating that the occluding loop, which is too short to sterically destabilize the associated EDR, does not perturb it allosterically. Also, bound Fas2 does not dissociate residual EDR from the active site of the complexed enzyme.

Structure C originates from a crystal grown in conditions different from those used for the structure A crystal and contains no bound phosphate. In subunit A, a sialic acid moiety, Sia\(_\alpha_{2,6}\)-Gal, was fitted into an unaccounted density proximal to His\(^{287}\) and appears to be terminal to the oligosaccharide chain GlcNAc-\(\beta_{1,4}\)-GlcNAc-\(\beta_{1,4}\)-Man-\(\alpha_{1,3}\)-Man, linked to Asn\(^{350}\) (data not shown) (36); the junction between the last Man and Gal moieties was not resolved, but Asn\(^{350}\) is located 25 Å away from His\(^{287}\), a value consistent with the mean distance between the two ends of an N-linked carbohydrate chain. A similar carbohydrate chain may be linked to Asn\(^{350}\) in subunit D, although a weaker density is observed. mAChE is composed of two species of roughly equal populations proposed to differ in their carbohydrate composition or some other post-translational modification (19). Combined gel-filtration, polyacrylamide gel electrophoresis, and isoelectric focusing experiments showed that the population of a higher mass is of a lower pl, an observation consistent with sialic acid in only two subunits in the mammalian tetramer (35).

**Significance of the Structure**

**Quaternary Structure of mAChE—**A square planar model for association of catalytic and structural subunits was recently proposed based on sequence alignments, computational predictions, and mutagenesis and biochemical data (38). The tetrameric structure of mAChE (Fig. 1) largely agrees with this model. Recombinant mAChE, however, is truncated after residue 548; hence, it lacks the carboxyl-terminal sequences encoded by the alternatively spliced exons and is expressed as a monomer (19), a feature that poses the question of whether physiological forms of the enzyme would form a similar tetramer. In mouse, the alternatively spliced sequences are (a) a 30-amino acid polar sequence encoded by the retained intron, (b) a glycosphospholipid signal sequence that is cleaved at selected positions upon addition of the glycosphospholipid, or (c) a 40-amino acid peptide containing an amphiphilic helix and a free cysteine. Only the last splicing option allows for tetrameric assembly in situ (4, 39); assembly may be facilitated by the amphiphilic helix- and polyproline-containing peptides (5).

Nevertheless, truncated mAChE associates as dimers in concentrated solution and as dimers and tetramers in the crystals. Upon gel filtration at protein concentrations used for crystallization and low ionic strength (10–12 mg/ml; \(\mu = 50\) mM), mAChE elutes as a dimer, whereas at concentrations <100 \(\mu\)g/ml, it elutes as a monomer; tetramers were not detected under these conditions or at high ionic strengths (\(\mu = 3\) M) approaching the crystallization conditions. Since protein concentrations exceeding those for dimer formation are apparently required to detect tetramers, it seems unlikely that occlusion of internal sites as seen in the structure would influence kinetic studies, typically conducted at low enzyme concentrations.

Deviation of the mAChE carboxyl termini out from the tetramer plane (Fig. 1) suggests that the twisted square planar structure could accommodate, on each side and with minimal perturbation, alternatively spliced sequences serving either as attachment sites for structural subunits or as membrane-anchoring domains. Since both amphiphilic and non-amphiphilic tetramers appear to form from a single splice variant (40), discrete tetrameric species are likely where the amphiphatic carboxyl-terminal sequences are either buried or exposed. Related tetrameric arrangements of catalytic subunits were recently observed in two 4.5-Å resolution crystal structures determined from two distinct crystal forms of trypsin-released tetrameric AChE from *Electrophorus electricus*: (a) a loose pseudo-square planar arrangement with free space in the center where the carboxyl-terminal sequences might be buried and (b) a compact square non-planar arrangement in which a folding of the tetramer along the dimer-dimer interface axis exposes the four sequences on the same side. Comparison of these two arrangements with that of mAChE indicates significant flexibility of the tetramer about the four-helix bundle axis and along the dimer-dimer interface axis. Hence, more than a single tetrameric arrangement may exist to accommodate the discrete carboxyl-terminal sequences.

**Conformational Mobility of mAChE—**\(\Omega\) loops are often important for protein stability and function and are so positioned for molecular recognition; \(\Omega\) loop lids, which are flexible until substrate or inhibitor is bound, are thought to play a role in enzymatic catalysis (34). Conformational gating provides a
mechanism to recruit substrates to the active site, to exclude solvent, to sequester reactive intermediates, or to enhance substrate specificity. Conformational mobility of loop Cys<sup>692</sup>-Cys<sup>696</sup> of the large Ω loop that is structurally homologous to the lid that sequesters the substrate in certain neutral lipases, which also show the α2-hydrolase fold (1, 2), has been proposed in gating accessibility of small molecules to the active center and in allosteric modulation of AChE catalysis (41–43). In the lipases, lid opening uncovering the catalytic site is a key feature of interfacial activation; however, the lid domain may be formed by one or more helices or surface loops that differ not only in length and sequence, but also in location relative to the α2-hydrolase core and the active site (44–49). In several α2-hydrolase proteins, the domain located between strands β<sub>s</sub> and β<sub>c</sub> is quite independent of the α2-hydrolase core and can accommodate large insertions such as a regulatory module (50), a putative helical lid (45), or a capping domain for the active site (51), without perturbing the core structure. The flexible short Ω loop Cys<sup>257</sup>-Cys<sup>272</sup>, which is also located between strands β<sub>s</sub> and β<sub>c</sub> of mAChE and protrudes to the surface of the molecule not far from the gorge entrance (Fig. 2), and helix α<sub>1</sub>–α<sub>7</sub>, where several residues interact with the facing subunit in the tetramer (Figs. 1 and 3), are virtually conserved in the entire α2-hydrolase family (cf. alignment in Refs. 2, 50, and 52–55). Yet a conformational rearrangement of helix α<sub>1</sub>–α<sub>5</sub>, or α<sub>2</sub>–α<sub>6</sub> (or both) would be required for the mAChE helix-loop-helix region to fold back and interact with the peripheral anionic site of the same mAChE subunit. Partial melting of α-helices into β-sheets participates in structural reorganization of the human pancreatic lipase (49). Unwinding of a helical turn, refolding of a flexible connection, and cis,trans-isomerization of a Pro residue acting as a hinge point accompany positional shift of the flap in Candida rugosa lipase (47). The mobile surface loop of cytochrome c peroxidase mutant W191G contains the Pro<sup>190</sup>-Gly-Gly-Ala-Ala-Asn-Asn<sup>196</sup> sequence, which shows striking similarity to the mAChE short Ω loop, and is followed by an α-helix starting at Asn<sup>196</sup> similar to mAChE helix α<sub>5</sub>–α<sub>7</sub>. Movement of the peroxidase loop involves a double-hinged rotation about Pro<sup>190</sup> and Asn<sup>196</sup> with a cis,trans-isomerization of Pro<sup>190</sup> and an interchange in the positions of the Asn<sup>195</sup> side and main chain atoms (56, 57). Hence, the sequence of the short Ω loop and the cis-conformation of Pro<sup>258</sup> at its base should confer conformational flexibility to the helix-loop-helix domain of mAChE.

**Heterologous Associations on AChE Surfaces**—Certain brain regions contain large amounts of AChE, but are devoid of acetylcholine and its biosynthesis capacity. This disparity in localization and the appreciable secretion of AChE into extracellular space suggest that AChE may have a signaling function independent of substrate catalysis (58–60). Non-catalytic properties of AChE may also play a role in neuropathology. Colocalization of AChE and amyloid β-peptide deposits in the brain of Alzheimer’s disease patients indicates a presence, if not an active role, in amyloid plaque formation (8–11). AChE was demonstrated to accelerate the assembly of amyloid β-peptides into Alzheimer’s fibrils, with a possible involvement of the peripheral anionic site (12, 61–63).

The solution structure of amyloid β-peptide-(1–40) shows a helix-loop-helix motif with a disordered central loop (64). Superimposition of this structure with that of the helix-loop-helix domain of mAChE (subunit A) brings several side chains in the β-peptide and mAChE in the same position (Fig. 6). In particular, Lys<sup>16</sup>, Ala<sup>21</sup>, and Val<sup>24</sup> in the β-peptide superimpose with Arg<sup>247</sup>, Ala<sup>252</sup>, and Val<sup>255</sup> in mAChE, respectively. Moreover, the side chains of His<sup>16</sup> in the β-peptide and Arg<sup>245</sup> in mAChE align, a position suitable for hydrogen bonding with Glu<sup>292</sup> in mAChE (cf. subunit C). Finally, the side chains of Glu<sup>22</sup> in the β-peptide and Arg<sup>253</sup> in mAChE also align (as does the side chain of Arg<sup>27</sup> in bound Fas2), a substitution that likely disrupts polar interactions, but not van der Waals contacts, with Pro<sup>250</sup> and His<sup>287</sup> in the peripheral site of mAChE. In solution, the high mobility of the two loops may bring dipetides Val<sup>24</sup>-Gly<sup>25</sup> and Gly<sup>291</sup>-Ala<sup>292</sup> in the β-peptide in the same positions as Val<sup>255</sup>-Gly<sup>256</sup> and Gly<sup>262</sup>-Gly<sup>263</sup> in mAChE. Hence, the amyloid β-peptide and the mAChE helix-loop-helix domain may interact in a similar manner with the AChE peripheral anionic site, which could serve as a nucleation site to promote aggregation.

Conformational plasticity of the amyloid β-peptides permits not only variable positioning of α-helices, but also local β-strand structures (64–66). In the amyloid β-peptides, as in the prion proteins, an α-helix to β-sheet conversion was proposed to accompany aggregation (67). Sequence analysis revealed similarity of the amyloid protein region Gln<sup>15</sup>–Gly<sup>37</sup> to the prion protein region Met<sup>120</sup>–Gly<sup>142</sup> (64), a region suggested to undergo α-helix to β-sheet transition in the pathogenic states (68–70). In addition to its structural similarity to the amyloid β-peptide, the mAChE helix-loop-helix domain with its glycine cluster shows intriguing sequence similarity to the Met<sup>120</sup>–Gly<sup>142</sup> region of the prion proteins (Fig. 6). This feature also makes rearrangement of the mAChE domain an attractive consideration.

In summary, the crystal structure of mAChE shows a tetrameric assembly of subunits; points to flexibility within the active center gorge; and reveals the peripheral anionic site, at the rim of the gorge, to be the site of association of surface loops not only from inhibitory peptides, but also from adjacent subunits. Hence, this may also be a site for heterologous protein association with AChE.

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**Fig. 6. Comparison of the mAChE helix-loop-helix domain with the human amyloid β-peptide and the rat prion protein.** Panel A, superimposition of the structures of the mAChE domain (residues 232–280) and the amyloid β-peptide-(1–40) (1 conformer out of 20) (64) according to the Cα atoms of mAChE helix α<sub>5</sub>–α<sub>7</sub> and β-peptide helix α<sub>5</sub>–α<sub>7</sub>, with blue coils, green α-helices, and red (mAChE) and magenta (β-peptide) loops. The side chains of selected residues in the mAChE domain and in the β-peptide are displayed as green and gray bonds with colored spheres, respectively. Panel B, comparison of the mAChE domain sequence with alignment of the amyloid β-peptide sequence (Aβp) and the prion protein sequence (Prp; sequence 85–146, with the asterisk representing a 23-residue insertion) (64) (same color codes as in panel A). Black dots above the mAChE sequence indicate the mAChE residues that bury 10% or more of their side chain surface area at the loop-peripheral site interface. Numbering above and below the mAChE and amyloid β-peptide sequences refers to residues whose side chains are displayed in panel A.
Peripheral Site-occluding Loop in Mouse Acetylcholinesterase

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