X-ray structure of dopamine transporter elucidates antidepressant mechanism

Aravind Pennarsa1*, Kevin H. Wang1* & Eric Gouaux1,2

Antidepressants targeting Na⁺/Cl⁻–coupled neurotransmitter uptake define a key therapeutic strategy to treat clinical depression and neuropathic pain. However, identifying the molecular interactions that underlie the pharmacological activity of these transport inhibitors, and thus the mechanism by which the inhibitors lead to increased synaptic neurotransmitter levels, has proven elusive. Here we present the crystal structure of the Drosophila melanogaster dopamine transporter at 3.0 Å resolution bound to the tricyclic antidepressant nortriptyline. The transporter is locked in an outward-open conformation with nortriptyline wedged between transmembrane helices 1, 3, 6 and 8, blocking the transporter from binding substrate and from isomerizing to an inward-facing conformation. Although the overall structure of the dopamine transporter is similar to that of its prokaryotic relative LeuT, there are multiple distinctions, including a kink in transmembrane helix 12 halfway across the membrane bilayer, a latch-like carboxy-terminal helix that caps the cytoplasmic gate, and a cholesterol molecule wedged within a groove formed by transmembrane helices 1a, 5 and 7. Taken together, the dopamine transporter structure reveals the molecular basis for antidepressant action on sodium-coupled neurotransmitter symporters and elucidates critical elements of eukaryotic transporter structure and modulation by lipids, thus expanding our understanding of the mechanism and regulation of neurotransmitter uptake at chemical synapses.

Chemical neurotransmission is initiated by Ca²⁺-induced release of neurotransmitters into the synaptic cleft. Upon release into the synaptic cleft, neurotransmitters such as glutamate, dopamine, noradrenaline, serotonin, glycine and GABA (-aminobutyric acid) activate G-protein-coupled receptors and ligand-gated ion channels, resulting in excitatory or inhibitory postsynaptic signalling cascades and currents. The widespread and critical roles of neurotransmitters in both central and peripheral nervous systems necessitate a requirement for strict spatiotemporal control of their levels at neural synapses. The primary mode of neurotransmitter clearance from the synaptic cleft is through secondary active transporters localized in presynaptic cells and glial cells that harness ionic gradients, across the cell membrane, to drive the uphill transport of neurotransmitters. This symport process requires both Na⁺ and Cl⁻ ions, which has led to the solute carrier 6 (SLC6) family of secondary transporters being referred to as neurotransmitter sodium symporters (NSSs).

Disregulation of NSS function is associated with several debilitating disorders that include depression, attention deficit hyperactivity disorder, orthostatic intolerance, epilepsy, Parkinson’s disease and infantile parkinsonism dystonia. NSSs are also the primary targets of antidepressants, drugs to treat neuropathic pain, attention deficit hyperactivity disorder, anxiety and of habit-forming substances of abuse such as cocaine and amphetamines. Development of antidepressants had a serendipitous beginning in the 1950s, followed by the discovery that the tricyclic antidepressant (TCA) imipramine inhibits noradrenaline reuptake in tissues. Numerous variants of imipramine, and the subsequent discovery of selective serotonin reuptake inhibitors, have revolutionized antidepressant treatment. To date, inhibition of neurotransmitter uptake remains the most widely used strategy for antidepressant therapy, despite numerous side effects.

Gains in our understanding of the molecular mechanisms underlying sodium-coupled transport have been derived from the structures of multiple conformations of LeuT, a bacterial sodium-coupled amino acid transporter with ~20% sequence identity to the eukaryotic NSSs. Models of eukaryotic NSSs based on LeuT have provided valuable insights into substrate and ion specificities, pharmacology and transport mechanisms in NSS members. However, bacterial NSS models fall short of answering questions concerning the elements of NSS structure and function, including the local structure of NSSs in regions that are unrelated to LeuT in amino acid sequence, the determinants of substrate selectivity and the atomic-level details of transport inhibition by antidepressants and addictive compounds. Moreover, there is no understanding, at the level of three-dimensional structure, of the role of lipids and post-translational modifications in NSS structure and mechanism.

Here we present a 3.0 Å X-ray crystal structure of the Drosophila melanogaster dopamine transporter (DAT) in complex with the TCA nortriptyline. The Drosophila DAT has greater than 50% sequence identity with its mammalian counterparts and harbours a pharmacological profile that is a hybrid of the mammalian DATs, noradrenaline transporters (NETs) and serotonin transporters (SERTs), making it a powerful vehicle to study NSS pharmacology and substrate specificity. The DAT structure reveals atomic details of TCA recognition, novel structural elements of NSS protein architecture and suggests a role for cholesterol in the allosteric control of transport in eukaryotic NSS members.

Thermostabilization and crystallization

Wild-type Drosophila DAT is labile, loses ligand-binding activity upon detergent extraction from the cellular membranes and is refractory to crystallization. To stabilize DAT for functional characterization, antibody generation and crystallization, we screened single point mutants for ligand-binding activity at increased temperatures, ultimately combining five mutations into the construct used for crystallization and structure determination (DAT_cyst). Purified DAT_cyst binds to the high-affinity inhibitor nisoxetine with a dissociation constant...
(K_d) of 29 nM (Supplementary Fig. 2a), and the TCA nortriptyline exhibits an inhibition constant (K_i) of 156 nM (Supplementary Fig. 2b). Unfortunately we were unable to measure the binding of nortriptyline to wild-type DAT because of its instability. Nortriptyline has a K_d of 18 nM at human SERT and 4.4 nM at human NET, values that are ~9-fold and ~35-fold lower than that for Drosophila DAT_crys. In dopamine-uptake measurements with the wild-type DAT and amitriptyline, a precursor of nortriptyline, transport is inhibited with a K_i of 30 nM, whereas the DAT_crys construct is inactive in transport (Supplementary Fig. 2c, d). Crystallization was further enhanced by the use of a complex with a Fab, resulting in crystals of a DAT_crys–Fab complex that diffract X-rays to 3.0 Å resolution.

Architecture of DAT

The structure of Drosophila DAT_crys bound to nortriptyline exhibits an outward-open conformation whereby the antidepressant is bound in a cavity halfway across the membrane bilayer and accessible to solvent from only the extracellular side of the membrane (Fig. 1). The transporter displays an overall LeuT-like fold with 12 transmembrane helices (TM)s in which helices 1–5 and 6–10 are related by inherent pseudo-symmetry, akin to LeuT (Supplementary Fig. 3). Residues in TM1 and TM6 make numerous interactions with the ligand and ions via non-helical, hinge-like regions at the approximate mid-points of these TM’s, connecting the bonding networks of all three ions with the inhibitor. Residues at the bend in TM3 contribute to the hydrophobic pocket that cradles the tricyclic moiety of the ligand, which lays approximately perpendicular to the TM’s, mimicking a wedge separating the jaws of a vice. One cholesterol molecule is located in a groove between TM5 and TM7 and poised to modulate the movement of TM10 that occurs during the transport cycle (Fig. 1a).

The primary binding site accommodates nortriptyline but cannot adopt the subsequent helical movements of TM1b and 6a required to form the occluded state. Using LeuT for comparison, the occluded state of LeuT is formed in the presence of sodium and leucine substrate, but not in the presence of tryptophan, which binds to the primary site, comparable to the TCA in the context of DAT_crys. We propose that both tryptophan and TCA stabilize the outward-open conformations of LeuT and DAT_crys, respectively, by targeting the primary binding site and sterically blocking the extracellular domains of the transporter, preventing the extracellular gate from closing and thus acting by way of a foot-in-the-door mechanism (Supplementary Fig. 4a, b and Supplementary Table 2).

Whereas the core of DAT_crys closely resembles that of LeuT, the periphery of DAT_crys exhibits several features distinct from LeuT and important for neurotransmitter transport and cellular localization. In TM12, a kink in the centre at Pro 572 causes the second half of the helix to turn away from the transporter, indicating that the dimerization interface of LeuT is not the same as potential oligomerization interfaces of eukaryotic SSs (Fig. 1a, b and Supplementary Fig. 5). Although previous studies indicate that SSsss oligomerize and DAT_crys is monomeric in detergent micelles and in the crystal lattice (Supplementary Fig. 6), thus suggesting that a membrane bilayer or additional molecules may be required for SSs assembly. The variable extracellular loop 2 (EL2) region has numerous predicted N-linked glycosylation sites and one disulphide bond, modifications that have critical roles in proper trafficking of SSs to the plasma membrane. The strictly conserved disulphide linkage was observed in the structure between two conserved cysteines, Cys 118 and Cys 157 (Supplementary Fig. 7). In the crystal, EL2 has a central role in lattice contacts, packing against a neighbouring Fab with an 870 Å interface (Supplementary Fig. 6). Because 43 residues were deleted from EL2 in the DAT_crys construct, further studies are required to determine the role of the full-length EL2 in transporter structure and function (Supplementary Fig. 8a). Together with EL2, EL4 harbours a Zn^{2+}-binding site in mammalian DATs that modulates transport (27). The equivalent residues in Drosophila DAT_crys are within a Cα-Cα distance of 10 Å, but because their identities are Ghu 161, Leu 374 and Ala 395, they do not form a high-affinity Zn^{2+}-binding site in DAT_crys.

TCA-binding site

Unambiguous density for nortriptyline in DAT_crys was observed in the primary site, approximately halfway across the membrane bilayer (Fig. 2a). In accordance with previous chimaeric studies, swapping TM regions between NET and DAT (28, 29), the drug-binding site is surrounded primarily by helices 1, 3, 5 and 6 in a region equivalent to the substrate-binding pocket of LeuT (29), and in close proximity to the densities for sodium and chloride ions (Fig. 2a). The dibenzocycloheptene ring of nortriptyline is oriented as a saddle, curving around the central region of TM3 and engaging in hydrophobic interactions with Val 120, Tyr 124 and Ala 117 (Fig. 2b). Val 120 is extensively conserved and faces the cycloheptene ring (Supplementary Fig. 4a), and replacement of the corresponding Ile 122 in human SERT with larger substitutions such as methionine markedly reduce affinity towards most SS inhibitors (30). This location was previously found in human SERT.

Figure 1 | Architecture of Drosophila DAT_crys. a, Structure of DAT_crys viewed parallel to membrane. Nortriptyline, sodium ions, a chloride ion and a cholesterol molecule are shown in sphere representation in magenta, purple, green and yellow, respectively. b, View of DAT_crys from the extracellular face. c, Surface representation showing that ligand and ion binding sites are accessible from the extracellular vestibule. Nortriptyline and TMs 1, 3, 6 and 8 are scored as in a.
to be protected from crosslinking agents in the presence of inhibitor or substrate. Phe 325 in TM6b forms an edge-to-face aromatic interaction with one of the benzo groups of norryptiline. Residues Gly-125 (TM3) and Ala 479 (TM10) also interact with the tricyclic group of the drug. The N-methylpropylamine group of the drug extends across the width of the drug-binding site and prevents TM5b and 6a from closing the extracellular gate above the drug. The amine group forms a hydrogen bond with the main-chain carbonyl of Phe 43 and a carboxy-an interaction with the side chain of Phe 43 (Fig. 2b). Interestingly, residues equivalent to Val 120 and Phe 43 (Ile 172 and Tyr 95) in SERT are necessary for interactions with antidepressants.

The biogenic amine transporters harbour a crucial aspartate residue in TM1 and in the DATcrys structure we see how Asp 46 substitutes for the absence of the carboxylate group in biogenic amines as compared to amino acid substrates transported by LeuT and the GABA and glycine transporters (Supplementary Fig. 8b). The side chain of Asp 46 forms a hydrogen bond with the hydroxyl of Tyr 124, which is equivalent to the Tyr 108 residue in LeuT that has a role in substrate recognition (Fig. 2c). Mutations at this aspartate result in substantial losses in transport activity and reduced binding affinities for cocaine. Ser 421 (TM8), which coordinates a sodium ion at site 2 (Na2), is within 3.5 Å of the propylamine group of the TCA and also forms a hydrogen bond with the carbonyl of Phe 43. Ser 421 therefore participates in a network of hydrogen bonds that interconnects norryptiline with the Na2 site and was also found to be crucial for high-affinity recognition of antidepressants by human SERT.

The N methyl group of norryptiline is 3.1 Å away from the main-chain carbonyl of Phe 319 and sterically prevents Phe 319 and TM6a from closing the extracellular gate above the drug, thereby stabilizing the outward-open state of the transporter. Phe 319 is the equivalent of Phe 253 in LeuT, which gates the substrate-binding pocket (Fig. 2c). The relative position of Phe 319 is markedly different from Phe 253 in the substrate-bound, occluded structure of LeuT, and instead resembles the positions of Phe 253 in the substrate-free and inhibitor-bound structures. To address the question of whether norryptiline could bind to DATcrys in a LeuT-like, occluded conformation, we superimposed DATcrys onto the occluded state of LeuT and found that Phe 319 and Phe 325 would clash with the dibenzocycloheptene ring of the TCA (Fig. 2c and Supplementary Fig. 4c). Identification of norryptiline bound in the substrate-binding pocket of DATcrys provides the first structural evidence that TCA antidepressants inhibit neurotransmitter transporters by preventing substrate binding and stabilizing the outward-open conformation. The DATcrys-norryptiline complex, together with the LeuBAT antidepressant complexes (Wang, H., et al., unpublished observations), demonstrate conclusively that antidepressants inhibit NSSs by acting at the primary or S1 site, in stark contrast to how TCAs inhibit LeuT via a non-competitive mechanism by binding within the extracellular vestibule.

### Ion-binding sites

Locations of ions essential for transport could be identified in DATcrys with electron densities (>4.0σ) at three locations near the non-helical hinge-like regions of TMs 1 and 6, and close to the TCA. Densities at the two sites coincided exactly with Na1 and Na2 sites identified in LeuT (Fig. 3a, b). A chloride ion was positioned at the third position of high omit density nestled in between TM2, 6 and 7 and close to Na1 (Fig. 3a). Placing ions in the omit densities during model building led to a concomitant loss of Fe2-Fc density during refinement. The atomic

---

**Figure 2** | Antidepressant-binding site.

- **a.** Overall view of the norryptiline bound DATcrys Fe2–Fc densities (blue mesh) for drug and ions are contoured at σ levels of 2.5 and 3.0, respectively.
- **b.** Close up view of the drug-binding pocket. Na+ and Cl− ions are shown as spheres. Norryptiline is represented as sticks (magenta). The amino group of norryptiline is 2.7 Å from the carbonyl oxygen of Phe 43 (TM1a) and the N-methyl group of norryptiline is 3.1 Å from the carbonyl oxygen of Phe 319. Residues lining the drug binding pocket have interfacial areas greater than 10 Å2 represented as sticks.
- **c.** Comparison of the drug or substrate binding pocket of DATcrys with that of LeuT (PDB code 2A65). The distance between the carbonyl group of leucine and Tyr 108 (spheres) is 2.7 Å in the occluded state (2A65) and 5.1 Å in the inhibitor bound state (3F3A) of LeuT, whereas the equivalent interaction in DATcrys between Asp 46 and Tyr 124 is 5.1 Å.

---

**Figure 3** | Ion-binding sites.

**a.** Na1 and chloride ion binding sites. Na1 is purple and Cl− is green and both are modelled as spheres. **b.** Coordination at the Na2 site is trigonal bipyramidal with the water molecule (w, red sphere) 3.3 Å from the sodium ion. Distances are in angstroms for residues that are in the coordination sphere and interactions are shown by dashed lines. Residues are coloured according to their respective TMs.
displacement factors of the ions matched the B-values of surrounding atoms. The sodium at site 1 is located ~5.2 Å away from the amino group of nortriptyline and is coordinated with an octahedral geometry by side-chain oxygens of Asn 49, Ser 520 and Asn 352, and main chain carbonyls of Ala 44 and Ser 520 (Fig. 3a and Supplementary Fig. 8b). Interestingly, the sodium at Na1 is also coordinated by one water molecule which in turn is within hydrogen bonding distance to Asp 46, thus showing that the Asp in TM1 indirectly participates in the sodium ion coordination. The mean ion coordinating distances (2.6 Å) at this site are longer than the distances (2.42 Å) reported for Na⁺ ions in solution but shorter than the distances reported for K⁺ ions (2.84 Å; Supplementary Table 3) ⁴⁰).

The chloride ion is located 5.0 Å away from the Na1 site at a position previously identified by computational and mutational studies based on LeuT ⁴¹ and the GABA transporters ⁴¹. A recent structural study of a chloride-dependent E290S mutant of LeuT also identified a chloride ion at this location ⁴¹. Chloride is coordinated in a tetrahedral fashion through residues in TM6 (Ser 320, Gln 316), TM7 (Ser 356) and TM2 (Tyr 69) (Fig. 3a). Interestingly, the hydroxyl group of Ser 320 bridges the Na1 and Cl⁻ sites and is positioned to interact with both ions. The mean ion-ligand distances at the Cl⁻ site are 3.0 Å (Supplementary Table 3) and the B-factors of surrounding atoms are similar to that of chloride, supporting the placement of chloride at this site.

The sodium at the Na2 site is located below the plane of the drug towards the cytoplasmic face, in between TM1 and 8, and is coordinated in a trigonal bipyramidal fashion by main chain carbonyls from Gly 42 (TM1), Val 45 (TM1-hinge) and Leu 417 (TM8) and the side chain oxygens from Ser 421 and Asp 420 (TM8) (fig. 3b). The mean ion-oxygen distances are 2.4 Å, in line with reported values for sodium coordination in solution (Supplementary Table 3). Although the interconnected network of interactions between TMs 1, 6, nortriptyline, sodium and chloride provides a structure-based mechanism for the coupling of ion and inhibitor binding ⁴⁲, we do not have a comprehensive understanding of the ion dependence of inhibitor binding in NSs.

Figure 4 Cholesterol site. a. Cholesterol (yellow sticks) shown with F O – F c density (light magenta) contoured at 2.0σ. Residues that interface with the cholesterol group are represented as sticks. b. Potential role of cholesterol in maintaining an outward-open state of transporter. Cholesterol (sticks with transparent surface) sterically clashes with the position of TM1a in the inward-open conformation of LeuT (PDB code 3TT5) ⁴³.

Figure 5 Extracellular and cytoplasmic gates and the C-terminal latch. a. Relative locations of the open extracellular gate (red box), closed cytoplasmic gate (blue box) and C-terminal latch (green box) in DAT ⁴⁴. b. The width of the extracellular gate is depicted by the distances between Tyr 124 and Phe 319 (10 Å), and Arg 52 and Asp 475 (10 Å). Nortriptyline, ions and helices are coloured as in Fig. 1. c. The cytoplasmic gate is closed by polar and electrostatic interactions between TM1a, TM2, IL1, TM6b and TM8. d. The C-terminal helix following TM12 is bound to the cytoplasmic face of the transporter via polar interactions with IL1. Polar and electrostatic bonds are represented as grey dashed lines.
**Cholesterol-binding site**

A cholesterol molecule is lodged in a trough-shaped cavity bordered by TM5, TM7 and TM1a at a depth equivalent to the inner leaflet of the membrane (Fig. 4a). Branched aliphatic residues are primarily involved in forming the protein–cholesterol interface (359 Å²), thus allowing cholesterol to bury ~57% of its solvent-accessible surface area. E_b – E_l density for this site clearly demarcated the orientation of the isocholesterol groups of cholesterol anchored at the junctions of TM5 and 7 by residues Leu 236, Leu 277 and Ile 358. The β-face of the sterol ring primarily faces residues Tyr 273, Leu 270 and Trp 266 in TM5 and also interacts with residues Val 34, Leu 37, Leu 38 and Ile 41 on TM1a. The α-face of cholesterol interfaces with residues Leu 347 and Ile 351 in TM7 (Fig. 4a).

Cholesterol has an important role in modulating the function of NSS members, stabilizing an outward-open state of DAT in concomitant increase in maximum binding or R_max for cocaine. In LeuT, TM1a undergoes a large conformational change upon transition from the outward-facing open and occluded states to the inward-open state. If a similar conformational change were to occur in DAT, it would disrupt the cholesterol site (Fig. 4b). We propose that one mechanism for the action of cholesterol on DAT is that by occupying its binding site in the outward-open, inhibitor-bound state, cholesterol stabilizes the outward-open conformation of the transporter.

**Extracellular and cytoplasmic gates**

The ion and ligand binding sites in DAT_cryo are accessible to solvent from the extracellular face owing to the open gate above the primary binding pocket. The distance between Tyr 124 of TM3 and Phe 319 in TM6a is 10 Å, whereas in the substrate-bound, occluded state of LeuT the corresponding distance is half as long (Fig. 5a, b). Similarly, the 10 Å separation between Arg 52 on TM1b and Asp 475 on TM10 renders the primary binding site accessible to extracellular solvent. The ionic bulk of the triecyl moiety combined with the extended N-methylpropylamine chain of nortriptyline prevents both TM1b and TM6a from approaching TM3 and TM8 to cap the putative substrate pocket and close the gate.

In contrast to the extracellular gate, extensive polar interactions at the intracellular face of the transporter form a thick barrier of ~24 Å between the ligand and ion pockets and solvent to keep the cytoplasmic gate shut. At the cytoplasmic face of the transporter, the indole nitrogen of Trp 30 caps the carbonyl oxygen of Tyr 331 in TM6b, and Arg 27 forms a salt bridge to Asp 435 of TM8 (Fig. 5c). Arg 27, Trp 30 and Asp 435 are strictly conserved in NSS orthologues and LeuT, suggesting that these intracellular gate interactions are general and important facets of the transport mechanism for this family of sodium symporters. Tyr 334, the residue corresponding to Tyr 335 in human DAT, was previously shown to be responsible for shifting the conformational equilibrium of the DAT towards an inward-open state.

**C-terminal latch**

Two novel attributes at the C terminus of DAT_cryo were immediately evident from the structure. Helix 12 is shifted by 22° in comparison to its position in LeuT, resulting in the exposure of TM3 to solvent and lipid (Supplementary Fig. 5). Pro 572, conserved in most eukaryotic NSS members, is probably at the root of the kink between TMs 12a and 12b, and thus has an important role orienting the second half of the helix away from the rest of the transporter. The hairpin between TM12b and the intracellular C terminus of DAT_cryo is stabilized by hydrogen bonding between the ε-nitrogen of Arg 589 and carbonyl oxygen of Thr 582 (Fig. 5d). The second feature is the C-terminal helix, which contains 2.5 turns from residues 586 to 595, where several hydrogen bonds and a carbon–carbon interaction between Trp 597 and Arg 101 restrain this C-terminal helix near intracellular loop 1 (IL1) at the cytoplasmic face of DAT_cryo. Although sequence conservation within TM12 and the C terminus is rather low across NSS orthologues, Gly 584 is located at the hairpin hinge and strictly conserved, and only lys or arg is present at the position equivalent to Arg 589 of DAT_cryo, suggesting that the conformation of the C terminus in the structure is a conserved feature among NSS orthologues. Studies of human DAT have identified the region following TM12 to contain sites for protein kinase C-mediated endocytotic trafficking and it is plausible that phosphorylation may alter the conformation or accessibility of the C terminus, allowing it to interact with cellular machinery for internalization. We also note that the latch participates in interactions with IL1, which in turn interacts with TM1a, thus suggesting that the C-terminal latch may modulate transporter activity.

**Conclusion**

The structure of DAT_cryo captures the transporter in an inhibitor-bound, outward-open conformation. The TCA nortriptyline targets the primary substrate site and stabilizes the open conformation by sterically preventing closure of the extracellular gate (Fig. 6a). One chloride and two sodium ions are located adjacent to the ligand, suggesting that the binding of ions and inhibitor are directly coupled. A cholesterol molecule bound to a crevice flanking TM1a probably stabilizes the outward-open, inhibitor-bound conformation (Fig. 6b). The structure reveals a C-terminal latch that makes extensive interactions with the cytoplasmic face of the transporter, proximal to the cytoplasmic gate, and thus in a position to modulate transport activity. Taken together, the structure of a eukaryotic DAT reveals novel insights into antidepressant recognition and structural elements implicated in the regulation of neurotransmitter transport, providing a foundation for drug design strategies.

**METHODS SUMMARY**

The Drosophila DAT_cryo construct (Supplementary Fig. 1) was expressed in virus-infected mammalian cells and purified by affinity and size-exclusion chromatography. Fab 9D5 was added before crystallization along with nortriptyline (1 mM) at a DAT:F(ab′)2 molar ratio of 1:1.1 and concentrated down to 3 mg ml⁻¹. Crystals of the complex were obtained in the presence of 100 mM glycine, pH 9, and 38% polyethylene glycol (PEG) 350 monomethyl ether (MME). The structure was solved by molecular replacement using a polyalanine model of LeuT (PDB code 3F3A) and an ensemble of Fab variable and constant domains. Data

---

**Figure 6 | Mechanisms of antidepressants and cholesterol.**

(a) The TCA nortriptyline (magenta) wedges between scaffold helices 3 and 8 and the core helices 1 and 6, preventing the movement of TMs 1b and 6a from closing the extracellular vestibule. Cholesterol (yellow) is bound in an intracellular pocket and hinders the movement of TM1a, thereby stabilizing the outward-open conformation of DAT. The C-terminal latch interacts with IL1 as part of the cytoplasmic gate.
processing, model building and refinement were performed using standard crystallographic software (Supplementary Table 1).

Full Methods and any associated references are available in the online version of the paper.

Received 2 June; accepted 7 August 2013.

Published online 15 September 2013.


Acknowledgements We thank J. Cavley for generating monoclonal antibodies and S. Amaral for providing the wild-type Drosophila DAT construct. We would like to thank H. Dong and O. Deaton for comments and suggestions along with other Gouaux laboratory members for discussions during manuscript preparation. We thank L. Vasksis for assistance with figures and J. Owen for help with manuscript preparation. We thank the staff of the Northeastern Collaborative Access Team (NECAT) at the Advanced Photon Source (APS) for assistance with data collection. This work was supported by a postdoctoral fellowship from the American Heart Association (A.P.), a National Institute of Mental Health research award (K.H.W.) and by the National Institutes of Health (E.G.). E.G. is an investigator with the Howard Hughes Medical Institute.

Author Contributions A.P., K.H.W. and L.G. designed the project. A.P. and K.H.W. performed protein purification, crystallography and biochemical assays. A.P., K.H.W. and E.G. wrote the manuscript.

Author Information The coordinates for the structures have been deposited in the Protein Data Bank under the accession code 4UAA. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to E.G. (gouaux@ohsu.edu).
METHODS

Screening, construct optimization and protein expression. The Drosophila DAT was selected as a promising candidate for structural studies after screening multiple orthologs of DATs and NETs by fluorescence detection site-exclusion chromatography (FSEC). In addition, FSEC was used to screen other parameters such as detergent efficacy, thermostability, lipid effects, tertiary epitope-specific monoclonal antibodies and sample homogeneity following purification. The DNA encoding the D. melanogaster DAT was provided by S. Amara and was modified by removal of the first 20 amino acids (M1-20), by a deletion in E1L (A164-206) and by point mutations to enhance thermostability (V74A, V275A, A311A, L415A, G338L) by PCR-based methods. This modified DAT sequence, denoted DATmouse was fused to a C-terminal GFP–His6 tag with a thrombin cleavage site (LEVPRGS) in place of residues 602-607. DATmouse GFP–His6 was produced by virus-mediated expression in mammalian cells.12

Antibody production. Monoclonal antibodies against DATmouse were raised by D. Cawley using standard methods. Antibodies were screened by FSEC and western blot to select clones that recognized native folded DATmouse protein. Sequencing of Fab regions was performed on mouse hybridoma cells (Fusion Antibodies) and on the intact antibody protein by Edman degradation (by M. A. Gavrilovic). Antibody was purified from hybridoma supernatant using 4 mercapto ethylpyridine resin. Fab protein was generated by papain cleavage of full-length antibody, followed by Fc capture on Protein A resin and cation exchange. Fab was stored in 20 mM sodium acetate, pH 5.5, 250 mM NaCl and 10% glycerol.

Purification of DATmouse Membranes. Membranes were solubilized in TBS (20 mM Tris, pH 8.0, 150 mM NaCl) containing n-dodecyl-β-D-maltoside (DDM) at a w/v ratio of 0.1 g detergent per 1 g membrane. The detergent-soluble fraction was incubated with cobalt charged metal ion affinity resin, and DATmouse GFP–His6 was eluted with 100 mM imidazole in 20 mM Tris, pH 8.0, 300 mM NaCl, 5% glycerol, 14 mM lipids (1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine (POPC), 1-palmitoyl-2-oleoyl-sn-glycero-3-phosphoethanolamine (POPE) and 1-palmitoyl-2-oleoyl-sn-glycero-3-phospho(1'-methylglycerol) (PAPG) at a weight ratio of 3:1:1, 1 mM DDM and 0.1 mM cholesteryl hemisuccinate. After thrombin digestion to remove the GFP–His6 tag, DATmouse was isolated by size-exclusion chromatography in 20 mM Tris, pH 8.0, 100 mM NaCl, 5% glycerol, 14 mM POPE, 4 mM dodecyl-maltoside and 0.1 mM cholesteryl hemisuccinate. The purified DATmouse protein was mixed with Fab SDB at a molar ratio of 1:3:1 and used for crystallization trials at 3 mg/ml.11, 14

Crystallization. Crystals grew in 100 mM glycine, pH 9.0 and 38% PEG 350 Mw using a drop ratio of 1 μl protein and 0.5 μl reservoir solution by hanging drop vapour diffusion. Initial crystals appeared at 4 °C after 2 days, reaching full size after 7 days. Crystals were flash frozen in liquid N2 directly and used for X-ray diffraction data collection.

Structure determination. X-ray data were collected at the Advanced Photon Source (Argonne National Laboratory, beamline 24-ID). Data were indexed, integrated, and scaled using HKL2000 (ref. 54) (Supplementary Table 1). The structure was solved using molecular replacement, with ensembles of constant (constant domains of heavy and light chains as one set) and variable domains (variable domains of heavy and light chains as a second set) of Fab coordinates in the PDB along with a polyalanine model of Lc (PDB code 3FJA). A multi-model search was done using Phaser. Initial phases were Improved by iterative steps of manual model building, refinement and maximum likelihood density modification using COOT, Phenix Refine and Phenix Phase and Build, respectively. Multiple rounds of refinement led to the placement of a majority of main-chain and side-chain atoms for both the Fab and DATmouse. The structure was refined to acceptable R-factors (Supplementary Table 1) with residues 25-24 and 600-665 in DATmouse and 135-138 in the heavy chain unmodelled owing to poor density. Nortripyrine, ions and cholesterol molecules were placed into Fc–Fc density contoured at 2σ or greater in the putative substrate pocket, ion sites and at the periphery of the transporter. Stereochemistry was evaluated using MolProbity.15

1. Ligation binding and uptake measurements. Scintillation proximity assays using transporter solubilized in detergent16 were carried out using copper yttrium silicate (Cu-Y2SiO5) beads (PerkinElmer) at 0.5 mg ml−1, 30 mM H2O2, 10 mM DATmouse–GFP–His6 protein in the same buffer as that used for size-exclusion chromatography. Unlabelled nortripyrine was used as the competitor ligand. Assay plates were read using a MicroBeta Trilux 1450 LSC & Luminescence counter. Data were fitted being a standard single site competition and, KI values were calculated from the IC50 values using the Cheng-Prusoff equation.

Uptake assays were performed using HEK293 cells expressing respective mutant constructs. Cells were re-suspended in 10 μM [3H]-dopamine (140-191 H) containing uptake buffer made with 25 mM Hepes-Tris, pH 7.1, 130 mM NaCl, 1 mM MgSO4, 5 mM KCl, 1 mM CaCl2, 5 mM d-glucose and 1 mM L-asparagine. Control samples were pre-incubated with 10 μM cold desipramine before addition of label. Assays were quenched with cold uptake buffer containing 1 μM desipramine after 10 min, cells were washed twice with cold uptake buffer and activity was measured from solubilized cells by scintillation counting. Data were plotted using Origin 7.0.

Thermotolerance screening of DAT. Sites for mutagenesis were selected on the basis of a model of Drosophila DAT built on the template of LeuT and residues were altered to Ala, Leu or Phe.12 Individual mutants along with the wild-type construct were transfected into HEK293 cells and kept in culture for 48 h, then tested for binding activity after detergent solubilization. Samples were split and one part was kept at 4 °C, and the other portion of lysate heated at 40 °C for 10 min. [3H]-nortripyrine was added before heating to select for mutants that stabilize an inhibitor-bound state of the transporter. Scintillation proximity assay was used to monitor activity in a high-throughput format. Mutants that consistently had an increased melting temperature (Ta) compared to wild-type (Ta ≈ 35 °C) were chosen and pooled into one construct, which yielded a four mutant construct with a Ta of ~60 °C.