Crystal structure of the dopamine N-acetyltransferase–acetyl-CoA complex provides insights into the catalytic mechanism

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The daily cycle of melatonin biosynthesis in mammals is regulated by AANAT (arylalkylamine N-acetyltransferase; EC 2.3.1.87), making it an attractive target for therapeutic control of abnormal melatonin production in mood and sleep disorders. Drosophila melanogaster Dat (dopamine N-acetyltransferase) is an AANAT. Until the present study, no insect Dat structure had been solved, and, consequently, the structural basis for its acetyl-transfer activity was not well understood. We report in the present paper the high-resolution crystal structure for a D. melanogaster Dat–AcCoA (acetyl-CoA) complex obtained using one-edge (selenium) single-wavelength anomalous diffraction. A binding study using isothermal titration calorimetry suggested that the cofactor bound to Dat first before substrate. Examination of the complex structure and a substrate-docked model indicated that Dat contains a novel AANAT catalytic triad. Site-directed mutagenesis, kinetic studies and pH-rate profiles confirmed that Glu47, Ser182 and Ser186 were critical for catalysis. Collectively, the results of the present study suggest that Dat possesses a specialized active site structure dedicated to a catalytic mechanism.

Key words: arylalkylamine N-acetyltransferase (AANAT), catalytic triad, dopamine N-acetyltransferase (Dat), melatonin, single-wavelength anomalous diffraction.

INTRODUCTION

Melatonin is a major hormonal mediator of light-induced photoperiodic changes in circadian biological events [1–3] and is found in bacteria, protozoa, macroalgae, plants, fungi, invertebrates and vertebrates. In vertebrates, melatonin, a pineal hormone, regulates seasonal and circadian cycles, including the sleep/wake cycle. In invertebrates, melatonin appears to initiate such physiological processes as reproduction and diapause [4,5]. In insects, amine acetylation by AANATs (arylalkylamine N-acetyltransferases) occurs during melatonin formation [6], sclerotization [7] and neurotransmitter inactivation [8–11]. These many different activities suggest that several insect AANAT isoforms exist. Drosophila melanogaster has two AANAT isoforms encoded by distinct genes [12,13], with Dat (dopamine N-acetyltransferase; EC 2.3.1.87) encoded by aaNAT1a.

Dat, as an AANAT, is a member of the GNAT (GCN5-related N-acetyltransferase) superfamily. GNATs are universally distributed in Nature and use an acyl-CoA to acylate their cognate substrates. Dat catalyses the transfer of the acetyl group in AcCoA (acetyl-CoA) to various arylalkylamines, including dopamine, serotonin, phenylethylamine and tryptamine. GNATs, including Dat, appear to have evolved from a common ancestor, as their sequences all contain four moderately conserved motifs denoted C, D, A and B (listed from the N- to the C-terminus) [14]. Up to the present study, a three-dimensional structure for an insect AANAT had not been solved. Notably, only one other AANAT structure has been solved, that of the Ovis aries (sheep) AANAT, SNAT (serotonin N-acetyltransferase) in its apo form and complexed with a bisubstrate analogue [15,16]. In SNAT, His118, His122 and Tyr168 were suggested to participate in the catalytic reaction [17]. The Dat sequence is not similar to that of SNAT, suggesting that the Dat catalytic residues and those of SNAT might differ, as might their catalytic mechanisms. In the present study, we describe the 1.46 Å (1 Å = 0.1 nm) resolution crystal structure of a truncated D. melanogaster Dat (Dat21–230)–AcCoA complex obtained from one-edge (selenium) single-wavelength anomalous diffraction data [18]. By combining examination of the crystal structure and a ternary (Dat21–230)–AcCoA–docked-substrate (tryptamine) model, and site-directed mutagenesis in conjunction with a kinetic study, we identified a probable catalytic triad and could propose a catalytic mechanism. This is the first structural study of an insect AANAT and has therefore provided insights into the catalytic mechanism and the structure–function relationships of GNATs.

EXPERIMENTAL

Protein expression and purification

The Dat gene was obtained from the DGRC (Drosophila Genomics Resource Center). PCR-based mutagenesis techniques were employed to make the N-terminal and C-terminal truncation mutant [19]. DNA encoding Dat21–230 was ligated into a pGEX-6P-3 vector and transformed into Escherichia coli BL21 (DE3) cells for subsequent expression of GST (glutathione transferase)-tagged Dat21–230. The Dat mutants were achieved using the QuikChange* site-directed mutagenesis kit (Stratagene). The cells were grown in 2-litre flasks that contained 400 ml of Luria–Bertani broth and ampicillin (100 μg/ml), which were then incubated at 37 °C with shaking until the D600 of the culture reached 0.5–0.6. The flasks were cooled to room temperature (25 °C) and then treated with isopropyl β-D-thiogalactopyranoside (final concentration, 1 mM). The cultures were then incubated at 24°C for 7 h.

Abbreviations used: AANAT, arylalkylamine N-acetyltransferase; AcCoA, acetyl-CoA; Dat, dopamine N-acetyltransferase; DTT, dithiothreitol; GNAT, GCN5-related N-acetyltransferase; GST, glutathione transferase; ITC, isothermal titration calorimetry; RMSD, root mean square deviation; SNAT, serotonin N-acetyltransferase.

The atomic co-ordinates and structural factors have been deposited in the PDB under accession code 3TE4 for the Dat21–230–AcCoA complex structure.

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The cell pastes were suspended in 30 ml of lysis buffer [PBS, pH 6.9, 10 mM DTT (dithiothreitol), 10 % (v/v) glycerol and 10 mM EDTA], and the cells were lysed by passage through a High-Pressure Homogenizer EmulsiFlex-C3 at 15 000 psi (1 psi = 6.9 kPa). Insoluble protein and cell debris were removed by centrifugation (4 °C, 48,000 g, 30 min), and the supernatant was applied to a 5 ml column of glutathione–Sepharose 4B (GE Healthcare). After elution from the glutathione–Sepharose 4B column, fractions that contained the fusion protein were combined and dialysed overnight against 20 mM Tris/HCl, pH 7.0, 150 mM NaCl, 1 mM DTT, 0.5 mM EDTA and 0.6 % Triton X-100. The GST tag was removed by PreScission protease (2 units/mg of fusion protein, Sigma), and the reaction was monitored by SDS/PAGE. Then the GST tag and the protease were removed by passage through 5 ml of glutathione–Sepharose 4B equilibrated with lysis buffer. The recovered protein was dialysed against 50 mM Tris/HCl, pH 7.0, 10 mM DTT, 1 mM EDTA and 10 % (w/v) glycerol, and then concentrated using a 5000 Da molecular mass cut-off (PES) Vivaflow 50 ultrafiltration apparatus (Sartorius Stedim Biotech). The concentrated protein was then equilibrated with lysis buffer. The recovered protein was dialysed against 50 mM Tris/HCl, pH 7.0, 10 mM DTT, 1 mM EDTA and 10 % (w/v) glycerol, and then concentrated using a 5000 Da molecular mass cut-off (PES) Vivaflow 50 ultrafiltration apparatus (Sartorius Stedim Biotech). The concentrated protein was then subjected to anion-exchange chromatography through 5 ml of HiTrap Q HP column (GE Healthcare); fractions containing the protein in 50 mM sodium phosphate, pH 6.8, 5 mM DTT, 1 mM EDTA and 0.198 mg/ml BSA and 2 mM AcCoA that was generated by the enzymatic acetyl-transfer reaction. Crystallization trials were performed at 15 °C after 7 days at 4 °C using a mixture of 1 μl of concentrated protein in 50 mM sodium phosphate, pH 6.8, 5 mM DTT, 1 mM EDTA and 1 μl of reservoir solution (0.1 M imidazole, pH 8.5, 1.0 M NaH₂PO₄, 1.6 M K₂HPO₄ and 0.2 M NaCl), which was subjected to SDS/PAGE and quadrupole time-of-flight MS (Sephacryl S-100 HR, GE Healthcare). The purified protein was subjected to SDS/PAGE and quadrupole time-of-flight MS (Quattro Ultima, Micromass). The same protocol was used to purify wild-type Dat, Dat mutant and selenomethionine-labelled Dat₂₁–₂₃₀ from an M9-medium expression culture [20].

### Crystallization and data collection

The purified selenomethionine-labelled Dat₂₁–₂₃₀ was mixed with a 4-fold molar excess of AcCoA. Crystallization trials were performed at 15 °C using the hanging-drop vapour-diffusion method, and crystals were obtained using the Wizard I random sparse matrix crystallization screen (Emerald Biosystems). The crystals most suitable for X-ray data were harvested after 7 days at 4 °C using a mixture of 1 μl of concentrated protein in 50 mM sodium phosphate, pH 6.8, 5 mM DTT, 1 mM EDTA and 1 μl of reservoir solution (0.1 M imidazole, pH 8.5, 1.0 M NaH₂PO₄, 1.6 M K₂HPO₄ and 0.2 M NaCl), which was equilibrated against 200 μl of reservoir solution. Diffraction data (1.46 Å resolution) were collected at 100 K at the SPXF beamline BL13B1, National Synchrotron Radiation Research Center (Taiwan) using an ADSC Quantum-315r CCD Area Detector. A complete single-wavelength anomalous dispersion dataset was collected at the selenium-absorption high-remote wavelength (0.96369 Å, determined by fluorescence scan). The datasets were processed and scaled using HKL2000 [21]. Crystallographic and refinement statistics are provided in Table 1.

### Structure resolution and refinement

The seven selenium sites in the asymmetric unit of the complex were located using SHELX C/D/E [22]. The overall figure of merit was 0.75. The electron density map was of good quality, such that 99 % of the structure could be built automatically by wARP [23]. The remaining 1 % was manually built with the aid of Coot [24]. Alternating model-building and refinement cycles using REFMAC5 [25] were performed with appropriate entries made in the dictionary for AcCoA. The final R-factor was 0.140 (R̄₂ = 0.198). The overall geometry of the final structure was assessed by PROCHECK [26]. Structural representations and models were generated using PyMol (Schrödinger). CD

CD spectra were acquired using an Aviv 202 spectropolarimeter (AVIV Biomedical) and a 1 mm pathlength cuvette. The far-UV CD spectra of the proteins, each corrected for the contribution of the buffer (7.5 μM protein, 5 mM sodium phosphate, 0.5 mM DTT and 0.1 mM EDTA, pH 6.7) are reported as mean residue ellipticity ([θ] deg-cm²-dmol⁻¹). Ellipticities were measured between 190 nm and 260 nm at 1 nm intervals at 25 °C. The CD data were displayed using GraphPad Prism (GraphPad Software).

### Acetyltransferase kinetic assay

The assay detected the reaction of Eillman’s reagent (5,5’-dithiobis-(2-nitrobenzoic acid)) [27] with the thiolate anion of CoA that was generated by the enzymatic acetyl-transfer reaction. Reactions were carried out in 0.3 ml of 50 mM sodium phosphate, pH 7.2, 500 mM NaCl, 2 mM EDTA, 0.05 mg/ml BSA and 2 mM AcCoA, with a variable tryptamine concentration (0–2 mM) at 30 °C in 1.5 ml microfuge tubes. Reactions were initiated by addition of enzyme (30 nM final concentration) that had been diluted 1000-fold with reaction buffer immediately before use and had been kept on ice prior to use in the assay. Reactions were

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**Table 1 Data collection and refinement statistics**

Values in parentheses are for the highest-resolution shell. $R_{merge} = \frac{\sum_I(\sum(|I(hkl)| - <I(hkl)>)^2)}{\sum_I(\sum|I(hkl)|^2)}$, where $R_{merge}$ was calculated over 5 % of amplitudes that were chosen at random and not used in refinement. Ramachandran analysis was from PROCHECK [26].

| Parameter | Selenomethionine derivative |
|-----------|-----------------------------|
| PDBe code | 3TE4                        |
| Data collection | |
| Protein | Dat₂₁–₂₃₀–AcCoA |
| Space group | P2₁/2/2₁ |
| Cell dimensions | |
| $a$, $b$, $c$ (Å) | 44.03, 56.62, 83.94 |
| $\alpha$, $\beta$, $\gamma$ (°) | 90, 90, 90 |
| Wavelength (Å) | 0.96369 |
| Resolution (Å) | 30–1.46 (1.51–1.46) |
| $R_{merge}$ | 0.058 (0.209) |
| $I/\sigma(I)$ | 23.4 (9.25) |
| Completeness (%) | 98.7 (98.7) |
| Redundancy (%) | 5.5 (5.3) |
| Refinement | |
| Resolution (Å) | 23.6–1.46 (1.50–1.46) |
| Number of reflections | 36,626 |
| $R_{merge}$ | 0.140 (0.198) |
| Number of atoms | |
| Protein | 1687 |
| AcCoA | 51 |
| Water | 945 |
| B-factors | |
| Protein | 13.7 |
| AcCoA | 8.9 |
| Water | 30.5 |
| RMSD | |
| Bond lengths (Å) | 0.019 |
| Bond angles (°) | 1.494 |
| Ramachandran plot statistics (%) | |
| Residues in most favoured regions (%) | 97.6 |
| Residues in additional allowed regions (%) | 2.4 |
| Residues in generously allowed regions (%) | 0 |
Table 2 Kinetic parameters for Dat in vitro

The activities of the mutants were normalized to that of the wild-type enzyme and all activities were expressed as percentages. The assays were performed at 30 °C in 50 mM sodium phosphate buffer, pH 7.2, with 500 mM NaCl, 2 mM EDTA, 0.05 mg/ml BSA and 2 mM AcCoA using tryptamine as the substrate with different concentrations (0, 0.05, 0.1, 0.2, 0.4, 0.8, 1.5 or 2 mM).

| Dat form | k_{cat} (s\(^{-1}\)) | Tryptamine | AcCoA | Tryptamine | k_{cat}/K_m (mM\(^{-1}\).s\(^{-1}\)) | Enzyme activity (%) |
|----------|---------------------|------------|-------|------------|---------------------------------|-------------------|
| Wild-type Dat | 21.9 ± 1.4 | 0.47 ± 0.09 | 0.43 ± 0.09 | 47 ± 9 | 100.0 |
| S182A | 1.8 ± 0.1 | 0.65 ± 0.09 | 0.36 ± 0.05 | 2.7 ± 0.4 | 5.7 |
| S186A | 4.8 ± 0.4 | 0.55 ± 0.07 | 0.59 ± 0.05 | 8.7 ± 1.2 | 18 |
| E47A | 0.13 ± 0.01 | 1.2 ± 0.1 | 0.04 ± 0.01 | 0.11 ± 0.01 | 0.23 |
| Y185F | 22.0 ± 1.3 | 0.47 ± 0.08 | 0.43 ± 0.02 | 47 ± 8 | 100.0 |
| Y185A | 22.0 ± 1.2 | 0.49 ± 0.07 | 0.40 ± 0.03 | 45 ± 6 | 96 |

RESULTS AND DISCUSSION

Characterization of a truncated Dat

We truncated the D. melanogaster Dat gene because the PredictProtein server [31] had predicted that residues 1–20 and 231–240 would be disordered in the native structure, which might have interfered with crystallization. The truncated Dat (called Dat\(_{21–230}\)) contains the four highly conserved motifs and has a molecular mass of 24411 Da, which is consistent with its theoretical molecular mass. Moreover, Dat\(_{21–230}\) and full-length Dat have very similar CD spectra and identical enzymatic activities. The ITC experiments revealed the similar affinity of substrate and cofactor for Dat\(_{21–230}\) and full-length Dat (Supplementary Figure S1 at http://www.BiochemJ.org/bj/446/bj4460395add.htm). In summary, these results showed that the truncation form of Dat is similar to native Dat both functionally and structurally, without detectably having an impact on the structural integrity of the enzyme.

Crystal structure of the Dat\(_{21–230}–\text{AcCoA}\) complex

The crystal structure of the Dat\(_{21–230}–\text{AcCoA}\) complex was solved at 1.46 Å resolution; the diffraction data having been acquired by one-edge (selenium) single-wavelength anomalous diffraction. Each asymmetry unit contains one Dat\(_{21–230}–\text{AcCoA}\) complex. Dat\(_{21–230}\) is a globular protein containing a seven-strand mixed β-sheet surrounded by nine α-helices (Figure 1A). The β-sheet is highly twisted and can be considered as two β-sheets (β1–β4 and β5–β7), which are partially separated at β4 and β5. The strands in each neighbouring pair are orientated in opposite directions (antiparallel), except for β4 and β5, which are parallel with each other. The β-sheet separates the α-helices into two groups that reside on opposite sides of the β-sheet. Motif C (Tryr\(^{157}\)–Phe\(^{165}\)) is located in β1 and α1. Motif D is located in β2 and β3 (Lys\(^{176}\)–Asn\(^{186}\)). Motif A (Leu\(^{179}\)–Gly\(^{177}\)) is located in β4, α5 and α6. Motif B (Val\(^{179}\)–Phe\(^{186}\)) is located in β5, α7 and β6.

ITC (isothermal titration calorimetry)

Binding of dopamine and AcCoA to Dat was measured by ITC using an Micro VP-ITC microcalorimeter (MicroCal). Aliquots of 6 μl of 5 mM AcCoA or dopamine were titrated at 25 °C by injection into protein (40 μM in 1.4 ml) in 50 mM sodium phosphate (pH 6.8), 5 mM DTT and 1 mM EDTA. The titration with 5 mM dopamine was performed in the presence of 1 mM CoA to avoid enzyme turnover. Background heats from ligand to buffer titrations were subtracted, and the corrected heats were measured under initial conditions (<10% turnover of the limiting substrate). The concentration of CoA produced was calculated on the basis of a standard curve of CoA. All assays were performed in triplicate, and rate values generally agreed within 10%. Kinetic constants were calculated by fitting the data to the Michaelis–Menten equation using a non-linear curve fit (GraphPad Prism), and the values are shown as means ± S.D. in Table 2. The error bars in the substrate–velocity plot represent the S.D. of each data point. The kinetic constants in the pH range 7–11 were measured as described above, with the exception that the reactions were carried out in sodium phosphate buffer at the indicated pH values (5.0, 6.0, 7.0, 8.0, 9.0 and 10.0). The results plotted were the average of three separate experiments. Curve fits for k_{cat} and k_{cat}/K_m data were calculated as described previously [28].

Molecular docking

The Dat\(_{21–230}–\text{AcCoA}\) structure was docked with tryptamine using GOLD [29]. Hydrogen atoms were added to the tryptamine using Discovery Studio 2.0 (Accelrys). The program automatically docked tryptamine into the active site of the Dat\(_{21–230}–\text{AcCoA}\) complex in all possible orientations. GoldScore was used as the fitness function with standard default settings employed for all calculations. The configuration file was defined as follows. The docking site was defined as all atoms within 10 Å of a specified centroid (x, y, z co-ordinates: 80.43, 84.41, 90.48). The best-ranked 25-docked poses were computationally evaluated and visually inspected. The pose with the most favourable energy was selected and further refined by steepest descendent minimization. Substrate–protein interactions were analysed by Discovery Studio 2.0 and Ligplot v.4.0 [30].
The crystal structure of the Dat21–230–AcCoA complex and interactions between Dat21–230 and AcCoA

(A) A stereo ribbon diagram of the Dat21–230–AcCoA complex. Dat contains nine α-helices and seven β-strands. The conserved GNAT motifs C, D, A and B are coloured cyan, green, yellow and red respectively. AcCoA is shown as a stick model and is situated in its binding cavity. (B) A stereo 2Fo − Fc electron density map for AcCoA contoured at 1σ within the stick model of the AcCoA-binding site. Hydrogen bonds are shown as broken lines. A certain number of the Dat21–230–AcCoA interactions are mediated by hydrogen bonding with water molecules (W1–W5, small red balls).

The AcCoA-binding site

The position of AcCoA was clearly defined in the electron density map and is located in a cleft that appears to be part of the funnel-shaped active site. The AcCoA conformation is approximately S-shaped. The cleft is almost completely covered by motifs A and B, which include the aforementioned secondary structural elements and the loop between α5 and α6 (Figure 1A). The AcCoA pantetheine is bound in this cleft at a position where β4 and β5 diverge. AcCoA forms hydrogen bonds with the backbone atoms of Leu146, Val148, Arg153, Gly154, Gly156 and Ala158 and one salt bridge with Lys192. Additionally, it also forms water-mediated hydrogen bonds with Asp46, Ser147, Tyr152, Leu155, Ile157, Gly159, Ser182 and Val186, and other van der Waals interactions with Val148, Arg153, Gly154, Ala158, Val179 and Val189 (Figure 1B and Supplementary Table S1 at http://www.BiochemJ.org/bj/446/bj4460395add.htm). Residues in the turn that joins α5 and α6 of motif A form a positively charged pocket that contains the pyrophosphate (Supplementary Figure S2A at http://www.BiochemJ.org/bj/446/bj4460395add.htm). Notably, the residues in the turn are the most highly conserved within motif A of insect AANATs (Figure 2). The AcCoA acetyl is located between β5 and a β-bulge in β4 and appears to be stabilized by contacts with the protein. Interestingly, in this β-bulge, the Lys144 and Ile145 side chains are on the same face of the β-sheet (Supplementary Figure S3 at http://www.BiochemJ.org/bj/446/bj4460395add.htm). The carbonyl oxygen of the AcCoA acetyl is within hydrogen-bonding distance of the Leu146 backbone nitrogen, which is positioned distincitively as a result of the β-bulge. Additionally, the main-chain carbonyl oxygen of Leu146 is within hydrogen-bonding distance of the nearest pantetheine nitrogen (N4P). By constraining AcCoA at two points via these two possible Leu146 backbone hydrogen bonds, the AcCoA sulfur atom is within hydrogen-bonding distance of the Ser186 hydroxy group (discussed below). Of note, Lys192 can form a distinct salt bridge with 3′-phosphate or α-phosphate of the ADP moity. This feature has not been observed in other AANAT complex structures to date.

Substrate and cofactor binding study by ITC

To investigate the binding property of Dat for its substrate and cofactor, ITC measurements were used to accurately determine the binding parameter and thermodynamic characterizations of Dat with dopamine and AcCoA respectively (Figure 3). The ITC data revealed that the interaction of Dat with dopamine appeared to be an exothermic reaction with enthalpy \( \Delta H = -7.85 \text{ kcal/mol} \), and entropy \( \Delta S = -5.32 \text{ cal/mol per K} \). Fitting to one set of the site-binding model, the \( K_s \) of dopamine binding to Dat was...
Dat structure reveals a novel catalytic triad

Figure 2 Sequence alignment of insect AANATs

Alignment of Dat and eight other insect AANAT sequences. Identical and conserved residues found in ≥ 80% of the protein sequences at a given position are highlighted in black and grey respectively. Dat21–230 secondary structural elements are shown above the sequence alignment. *, Dat catalytic-triad residues; †, residues implicated in the binding of Dat 21–230 and AcCoA; and $\Delta$, non-polar residues predicted to be involved in tryptamine binding according to the ternary-docking model. The conserved motifs C, D, A and B are shown as boxes. Motif C (Tyr22–Phe43) is located in $\beta_1$ and $\alpha_1$. Motif D is located in $\beta_2$ and $\beta_3$ (Lys76–Asn90). Motif A (Leu139–Gly173) is located in $\beta_4$, $\alpha_5$ and $\alpha_6$. Motif B (Val179–Phe199) is located in $\beta_5$, $\alpha_7$ and $\beta_6$. Species: Acyrthosiphon pisum, Antheraea pernyi, Apis mellifera, Bombyx mori, Culex quinquefasciatus, Drosophila melanogaster, Glossina morsitans, Periplaneta americana and Tribolium castaneum.

calculated to be $3.88 \times 10^4$ M$^{-1}$, and the binding constant $K_d$ was 25.9 μM. The interaction of Dat with AcCoA had a higher affinity, with a $K_a$ of $1.88 \times 10^5$ M$^{-1}$. As shown in Table 3, a significant amount of heat was released when Dat associated with AcCoA, indicating that the binding interactions had significant enthalpic contributions ($\Delta H = -11.5$ kcal/mol), which were dictated by hydrogen bonding and van der Waals interactions (Figure 1B and Supplementary Table S1). Analysis of the ITC data revealed a slightly unfavourable entropic contribution ($\Delta S = -14.4$ cal/mol per K), possibly indicating that the Dat structure was slightly stabilized upon binding to AcCoA and that very few solvent molecules were freed from the binding pocket (solvent entropy). These results were in agreement with crystallographic data, as indicated by: (i) the significant reduction of the $B$ values for three regions upon formation of the Dat–AcCoA complex (conformational entropy) (Supplementary Figure S4 at http://www.BiochemJ.org/bj/446/bj4460395add.htm); and (ii) there were not only a few water molecules in contact with AcCoA through hydrogen bonding, but also there were some water molecules in the binding pocket. To determine the binding order for the substrate and cofactor towards Dat, the dopamine titration experiment indicated that there was no significant binding to Dat in the absence of CoA (results not shown). This ITC result suggested that the cofactor binds first to Dat followed by substrate binding.

Comparison of Dat21–230 with other N-acetyltransferase domains revealed a structurally conserved core domain

As shown by iSARST [32], the Dat21–230 tertiary structure is similar to those of N-acetyltransferase domains found in other GNATs (Supplementary Figure S5 at http://www.BiochemJ.org/bj/446/bj4460395add.htm), including SNAT [16], tabtoxin-

resistance protein [33], aminoglycoside 6-N-acetyltransferase [34], histone acetyltransferase [35], and glucosamine-6-phosphate N-acetyltransferase 1 [36], which had structure diversity scores in comparison with the Dat21–230 structure of 3.47, 4.16, 4.58, 5.75 and 6.63 respectively (Table 4). The pairwise alignments of the C, D, A and B motifs in Dat21–230 with those
in the other five N-acetyltransferase domains [37] are generally consistent, although some loop structures are not well aligned. Comparison of the values for the Ca RMSDs (root mean square deviations) of structurally equivalent residues in Dat_{21–230} and in each of the other proteins indicates that the core fold structures are very similar despite a low level of sequence identity and similarity (Table 4). The conformation of the bound AcCoA in Dat_{21–230} and its orientation compared with those of the other five GNAT proteins is virtually indistinguishable. Additionally, each structure possesses a β-bulge at a position equivalent to Lys^{144} and Ile^{145} in Dat_{21–230}. The presence of the β-bulge is remarkably well conserved in active sites among GNATs, suggesting a critical role in the formation of the AcCoA-binding site [38,39]. However, one striking difference between Dat_{21–230} and the other five structures exists: the tyrosine residue found in each of those five structures, whose phenolic hydroxyl group is within hydrogen-bonding distance of the AcCoA sulfur atom and may therefore stabilize the thiolate anion of the departing CoA molecule, is replaced by Ser^{186} (see below and Supplementary Figures S5 and S6 at http://www.BiochemJ.org/bj/446/bj4460395add.htm). The critical role of this tyrosine residue in catalysis was also reported in these acetyltransferases of GNATs [17,40]. A structure-based alignment of the Dat and SNAT sequences shows that the sequences of the conserved GNAT motifs were unambiguously aligned (Supplementary Figure S7 at http://www.BiochemJ.org/bj/446/bj4460395add.htm). Inspection of their structures revealed that Dat has a deeper funnel leading into its active site (Supplementary Figure S2). A remarkable feature that distinguishes the Dat structure from that of SNAT is that the Dat substrate-binding site is more deeply buried. The geometry and non-polar nature of the funnel interior provide a basis for the enzymatic properties of Dat. The ITC results revealed Dat had a much higher affinity for substrate than that of SNAT (Dat, $K_d = 222.5 \mu M$ for serotonin, results not shown; SNAT, $K_d = 1299 \mu M$ for serotonin) [41]. The deeply embedded cavity provides a non-polar environment that can bind the substrate more avidly. The shape and size of the cavity must therefore dictate substrate selectivity and specificity. Some common protein–acetyl and protein–pantheine interactions were found in both structures. Nevertheless, Dat_{21–230} has more residues involved in the interactions with AcCoA than those of SNAT (17 residues in Dat; nine residues in SNAT) [16]. The ITC data also provide evidence of an increased binding affinity of approximately 45-fold between Dat and AcCoA compared with that of SNAT (Dat, $K_d = 5.43 \mu M$; SNAT, $K_d = 242 \mu M$) [41].

### A ternary docking model

To begin to understand how a substrate binds in Dat, tryptamine was docked into the Dat_{21–230}–AcCoA crystal structure (Figure 4A). Our docking strategy was to bring the tryptamine amino group in close proximity to the acetyl carbonyl group of AcCoA as would be expected during the reaction. A non-polar pocket, denoted the dopamine-binding pocket, that is mainly formed by Phe^{41}, Leu^{61}, Leu^{65}, Tyr^{64}, Ile^{116} and Ile^{145}, was found inside the funnel-shaped active site and contained the indole ring of the docked tryptamine, which made extensive non-polar contacts with the side chains of these residues. Given the geometry and non-polar nature of the pocket, steric factors and non-polar interactions should dominate substrate–Dat binding. The six residues are highly conserved in insect AANATs (Figure 2). Van der Waals interactions and packing of non-polar residues therefore are probably responsible for the binding of arylalkylamine substrates in insect AANATs. The dopamine-binding site is solvent-accessible via the funnel, so that a substrate can enter and an N-acetylated product can leave, and it is adjacent to the acetyl in AcCoA.

### Potential catalytic residues

Alignment of nine insect AANAT sequences revealed that the sequence identity among them is substantial, between 30 and 63% (Figure 2). There are many regions in which >80% of the residues are conserved. Examination of the Dat_{21–230}–AcCoA structure and the substrate-docked model showed that certain residues are important for substrate and AcCoA binding, and others may be involved in catalysis (Figures 2B and 4A). The conserved insect AANAT Glu^{67}, Ser^{182} and Ser^{186} may be the catalytic residues, because they are situated near the leaving and acceptor groups.

### Table 3 Thermodynamic analysis of binding of the substrate and cofactor to Dat

| Molecule | $K_d$ (10^{4} \text{M}^{-1}) | $K_a$ (\mu M) | $\Delta H$ (kcal/mol) | $\Delta S$ (cal/mol per K) | $\Delta G$ (kcal/mol) | $\sigma$
|----------|-----------------|-------------|-----------------|-----------------|-----------------|------|
| Dopamine | 3.88 ± 0.26     | 25.9 ± 2.5  | −7.85 ± 0.17    | −5.32           | −6.26           | 1.03 ± 0.03 |
| AcCoA    | 18.8 ± 2.6      | 5.43 ± 1.09 | −11.5 ± 0.3     | −14.4           | −7.20           | 0.88 ± 0.01 |

### Table 4 Pairwise comparisons of the Dat_{21–230}–AcCoA structure with five N-acetyltransferases by the iSARST server [32]

| N-acetyltransferase | Aligned residues | Identity* | Similarity† | RMSD for the Ca carbons (Å) | Structure diversity‡ | iSARST score | PDB code |
|---------------------|-----------------|-----------|-------------|-----------------------------|---------------------|-------------|----------|
| SNAT                | 125             | 12 % (20/166) | 23 % (38/166) | 1.89                        | 3.47                | 1CJW        |
| TTR                 | 128             | 6.4 % (14/166) | 26 % (43/166) | 2.33                        | 4.16                | 1GHE        |
| AAC (6’)-li         | 120             | 12 % (22/181) | 25 % (45/181) | 2.20                        | 4.58                | 1B6T        |
| HPA2                | 127             | 15 % (22/150) | 29 % (44/150) | 3.41                        | 5.75                | 100M        |
| GNA1                | 126             | 15 % (23/154) | 31 % (47/154) | 3.82                        | 6.63                | 1I12        |

*Values in parentheses: (number of identical residues in the aligned regions/inquiry protein sequence length) × 100.
†Values in parentheses: (number of conservatively exchanged residues in the aligned regions/inquiry protein sequence length) × 100.
‡Structure diversity: RMSD/(number of residues in the aligned sequence/total number of residues in the shorter-length protein) × 100.
in AcCoA and tryptamine respectively (Figure 4B). Besides, the Dat sequence contains a Tyr185 just before Ser186. As tyrosine is essential for the catalysis in some acetyltransferases of the GNAT superfamily, the potential involvement of Tyr185 in catalysis also requires elucidation. Therefore we individually mutated each of these residues to either alanine or phenylalanine (only for Tyr185) and examined their effects on Dat catalysis.

Comparison of the kinetic parameters of E47A, S182A, Y185F/Y185A, S186A and wild-type Dat

The secondary structures of all mutants and wild-type Dat were identical according to far-UV CD spectroscopy (Figure 5A). Given the highly variable effects on both the turnover and the Michaelis–Menten constants for the enzymatic activity of the mutants except for Y185F/Y185A (Figure 5B and Table 2), Glu47, Ser182 and Ser186 probably function as a catalytic triad. Interestingly, the relative impact of each mutation on the activity of Dat was quite different. Compared with the value for Dat, mutation of Ser186 and Ser182 caused an apparent decrease in the $k_{cat}/K_m$ value for tryptamine, by 82% and 94% respectively. E47A had a $k_{cat}/K_m$ value for tryptamine that is only 0.23% of that found for Dat, and is the only mutation of a Dat active-site residue identified to date that has such a dramatic effect on enzymatic activity. Furthermore, E47A reduced the $k_{cat}$ value 168-fold and increased the tryptamine $K_m$ value 2.5-fold. A smaller decrease in the $k_{cat}$ value was observed for S186A (4.5-fold reduction), whereas S182A exhibited a greater decrease in the $k_{cat}$ value (12-fold reduction). It is clear that these three mutants showed increases in the $K_m$ for tryptamine and decreases in the $K_m$ for AcCoA in the reaction. Notably, the drastically reduced activity for E47A also suggests that Glu47 initiates catalysis (see below). Of note, the Y185F and Y185A variants show no significant effect on Dat enzyme activity. Structural analysis (Supplementary Figure S6) combined with kinetic studies exclude the possibility of Tyr185 involved in catalysis.

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The residues in Dat, tryptamine and AcCoA that are directly involved in catalysis are included in the Figure and are labelled. (a) The proposed mechanism involves Glu47, acting as a base catalyst, to initiate the reaction by abstracting a proton from the hydroxy group of Ser182. (b and c) The hydroxylate of Ser186 can then act as a general base catalyst for deprotonation of the substrate amino, thereby forming a tetrahedral intermediate. (d) The hydroxy group of Ser186, which is separated by 3.8 Å from the AcCoA sulfur, could then serve as a general acid catalyst to protonate the thiolate anion of the leaving CoA. (e) The proton is transferred from the carboxylate group of Glu47 to the alkoxide group on Ser186 through a water molecule (see Supplementary Figure S10 at http://www.BiochemJ.org/bj/446/bj4460395add.htm). The backbone amide of Leu146 can form a hydrogen bond with the carbonyl oxygen of the acetyl group to impose the appropriate stereochemistry on the tetrahedral intermediate.

The pH-dependence of steady-state kinetic measurement for wt Dat, S182A, S186A and E47A

Investigations of the pH rate profile can sometimes provide information about the properties of ionizable groups that participate in enzymatic catalysis. Analyses of log(kcat/Km) compared with pH should reflect pKa values of ionizable groups present in the free enzyme and/or substrate [42]. The pH-dependence of log(kcat/Km) of the wild-type Dat with tryptamine and AcCoA, an indicator of the state of the free enzyme or the substrate, revealed two ionizable groups with pKa values of 6.9 ± 0.1 (pKa1) and 9.0 ± 0.2 (pKa2) (Supplementary Figure 8A at http://www.BiochemJ.org/bj/446/bj4460395add.htm). The log(kcat/Km) compared with pH profiles of S182A and S186A revealed a distinct change in the higher pH values (Supplementary Figures 8B and 8C). Although the acidic limb with a pKa of ~6.9 was maintained, the basic limb of the wild-type profile was absent, suggesting that the higher pKa value (pKa2) reflects the ionization of the hydroxy group of Ser182/Ser186. Therefore we could reasonably refer the ionizable groups to Ser182 and Ser186, which could act as an acid/base and nucleophile for catalysis in the deprotonated form. Such a pKa would be close to that of a previously proposed serine residue in catalysis of fatty acid amide hydrolase [43,44]. Even though Ser182/Ser186 would have a pKa value of approximately 9.0, which is 4.0 units lower than a free serine residue in solution (pKa = 13), it is well known that protein active sites can change the pKa values of ionizable groups on catalytic residues compared with their values in free solution [45]. The pKa can be altered immensely by the microenvironment, including electrostatic effects, hydrogen bonding or a hydrophobic surrounding provided by an enzyme, to the extent that residues which are basic in solution may act as proton donors, and vice versa. The high resolution of the Dat complex structure has allowed us to define a hydrogen-bonded network that includes water-mediated hydrogen bonds between Ser182 and Ser186, which would probably be one of the factors required to depress the pKa value significantly (W5 in Figure 1B). There are also many examples of catalytic residues that have significantly lower pKa values as compared with the native pKa of the amino acid [45–47]. Moreover, the pH-dependence of log(kcat/Km) for the E47A mutant showed a linear dependence on [OH−] with a slope of 0.3 (Supplementary Figure 8D). The E47A mutation resulted in a pH-dependence profile of log(kcat/Km), which abolished the acid limb seen with the wild-type enzyme. This suggests that the pKa value reflects the ionization of the carboxylate group of Glu47. The greatly reduced enzyme activity of the E47A mutant coupled with its altered pH-rate profile was consistent with a role for Glu47 to act as a base in the catalysis reaction. There are also several examples of glutamate residues that play a catalytic role as a general base [48,49].

Proposed catalytic mechanism for Dat

The structural and kinetic activity data that are available for GNATs support a mechanism that proceeds by direct nucleophilic attack on the AcCoA acetyl by an acceptor substrate [28,38]. For Dat, the crystallographic evidence in conjunction with the altered pH-rate profile support the simplest acetyl-transfer mechanism involving one of the two serine residues acting as a general
Dat structure reveals a novel catalytic triad

Conclusions

The present study represents the first structure of an insect AANAT. We have also confirmed that Dat is a member of the GNAT superfamily and suggested that Dat from D. melanogaster utilizes a Glu-Ser-Ser catalytic triad mechanism. For members of a protein superfamily to be able to catalyse the same chemical reaction but target different substrates requires a common catalytic domain, but also structural diversity within the domain. It is interesting to research how specificity for many different substrate types can be generated given this common central domain. Such studies should increase our knowledge of Dat in relation to other GNAT superfamily members and should allow for the development of specific insecticides and potential lead compounds that might treat mood and sleep disorders induced by abnormal melatonin production.

AUTHOR CONTRIBUTION

Kuo-Chang Cheng and Ping-Chiang Lyu designed the research, analysed the data and wrote the paper. Kuo-Chang Cheng performed all of the experiments. Jhen-Ni Liao purified the recombinant proteins and carried out enzyme kinetics and ITC experiments. Kuo-Chang Cheng determined the structure. Ping-Chiang Lyu commented on the paper.

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SUPPLEMENTARY ONLINE DATA

Crystal structure of the dopamine N-acetyltransferase–acetyl CoA complex provides insights into the catalytic mechanism

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Figure S1 Comparisons of Dat and Dat<sub>21–230</sub>

(A) CD spectra and (B) enzymatic kinetic profiles of Dat and Dat<sub>21–230</sub>. Release of the CoA was used to measure the enzymatic activity. Titration of dopamine (C) and AcCoA (D) against Dat<sub>21–230</sub> were performed. Results are shown for heat change (upper panels) and peak integration (lower panels). The continuous line represents the best fits to a single-site binding model. The binding constants (K<sub>d</sub>) for dopamine and AcCoA were 12.6 μM and 2.82 μM respectively.

The atomic co-ordinates and structural factors have been deposited in the PDB under accession code 3TE4 for the Dat<sub>21–230</sub>–AcCoA complex structure.

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Figure S2  Substrate- and AcCoA-binding pockets in Dat21–230 and in SNAT

(A and B) Surface electrostatic potential representations of the active sites in Dat21–230 and SNAT respectively. Tryptamine and AcCoA in their respective binding pockets in Dat21–230 are shown as stick and ball-and-stick models respectively. In SNAT, the bisubstrate analogue, CoA-S-acetyltryptamine, is shown as ball-and-stick model. The atoms of AcCoA are coloured green (carbon), blue (nitrogen), yellow (sulfur), red (oxygen) and orange (phosphorous). The colour code for the electrostatic potential is blue for positive charges and red for negative charges. The bottom panels are close-ups of the active sites.

Figure S3  The Dat21–230 β-bulge

The residues of β3 and β4 are shown as stick models with the β-bulge formed by Lys144 and Ile145, which positions their side chains on the same face of the β-sheet. AcCoA is shown as a ball-and-stick model. The atoms of AcCoA are coloured green (carbon), blue (nitrogen), yellow (sulfur), red (oxygen) and orange (phosphorous). The broken lines indicate hydrogen bonds.

Figure S4  Temperature factor plots of Dat crystal structures in apo form and complex form

Plot of main-chain temperature factors for the apo form (continuous line) and complex form (broken line) of Dat crystal structures. In the complex structure, three regions, residues 35–54, 95–117 and 215–220, show significantly lower temperature factors compared with that of the apo form.
Dat structure reveals a novel catalytic triad

Figure S5 Structures of Dat21–230 and other GCN5 N-acetyltransferase domains

The conserved GNAT motifs C, D, A and B are coloured light blue, green, yellow and red respectively. AcCoA is shown as a ball-and-stick model. The conserved catalytic tyrosine residues (serine in Dat) are shown as red stick models (highlighted in the broken circle). AAC (6′)-II, aminoglycoside 6′-N-acetyltransferase type II, Enterococcus faecium; GNA1, glucosamine-6-phosphate N-acetyltransferase 1, Saccharomyces cerevisiae; HPA2, histone acetyltransferase, Saccharomyces cerevisiae; SNAT, serotonin N-acetyltransferase, Ovis aries; TTR, tabtoxin-resistance protein, Pseudomonas syringae.
Figure S6  Structure of Dat21–230–AcCoA complex with a docked tryptamine: spatial relationships between potential residues and AcCoA at the active site

The Dat sequence contains a Tyr^{185} residue just before Ser^{186}. The residues Tyr^{185} (red) and Ser^{186} (black) are represented as stick models. The broken lines identify distance between the AcCoA sulfur atom and target residues. The numbers represent distance separations between various atoms in angstroms. Tryptamine is shown as a stick model and AcCoA is shown as a ball-and-stick model.

Figure S7  Structure-based sequence alignment of Dat and O. aries SNAT

To structurally align the sequences, first their three-dimensional structures were manually aligned with gaps introduced as needed. Then, the sequences of the common secondary structural elements were manually aligned with the positions of those in SNAT adjusted so that they were aligned with the corresponding secondary structures in Dat21–230. The secondary structure elements in Dat and SNAT are indicated above and below the sequences respectively. The conserved motifs C, D, A and B are shown as boxes coloured cyan, green, yellow and red respectively.
Figure S8  The pH profiles of log(\(k_{\text{cat}}/K_m\)) for wild-type Dat, S182A, S186A and E47A

The pH-dependence of log(\(k_{\text{cat}}/K_m\)) observed for Dat (A) are shown with curve fits obtained by non-linear least-squares regression. The pKₐ values are given on the graphs. pK₁ (6.9 ± 0.1) refers to the acidic limb of the profile, and pK₂ (9.0 ± 0.2) represents the basic limb. pH values compared with log(\(k_{\text{cat}}/K_m\)) profiles of S182A (B) and S186A(C) indicate a ionizable residue which is active only in the deprotonated form involved in catalysis with a pKₐ of approximately 6.9. The pH profile for the E47A mutant (D) was fitted to a line with a slope of 0.3.

Figure S9  Comparison of the catalytic triads of AANATs from D. melanogaster and O. aries

The structural superimposition of Dat (blue) and SNAT (grey). The ligands and catalytic residues for Dat and SNAT are labelled and shown in stick representation in blue and grey respectively. The two structures share a very similar structural fold despite low sequence identity (12 %), with an RMSD value of 1.89 Å.
Figure S10  Comparison of the catalytic triad residues of Dat in apo form and complex form

The catalytic triad residues Glu47 and Ser182 are represented as stick models in the apo form (A) and complex form (B) of the Dat structure. The broken lines identify the distance between the catalytic residues. The numbers represent distance separations between various atoms in angstroms. (C) The catalytic triad residues Glu47 and Ser186 form a hydrogen bond network with each other by a proton-conducting water molecule (shown as a red ball). The density, shown as a blue mesh, is a 2Fo – Fc map, contoured at 1.1σ. Note that, by comparing the Dat structure between apo form and complex form, two crystallographic facts could implicate the catalytic role of Glu47: (i) the distance between the Glu47 carboxylate anion and Ser182 hydroxy group decreased after formation of the Dat–AcCoA complex structure, which enabled Glu47 to abstract a proton from the hydroxy group of Ser182. Moreover, the carboxylate of Glu47 forms a hydrogen bond to a proton-conducting water molecule. This water molecule is co-ordinated by hydrogen bonding and is 2.8 Å away from the hydroxy oxygen of Ser186 in an appropriate position to transfer a proton (crystallographic and refinement statistics of the apo form structure are not shown. K.-C. Cheng and P.-C. Lyu, unpublished work).
Table S1  Detailed protein–cofactor contacts

The numbers represent distance separations between cofactor atoms and protein residue atoms in angstroms. See Figures S11 and S12 for more details.

(i) AcCoA salt bridge

| Atom | Salt bridge |
|------|-------------|
| O8A  | Lys153 NZ 2.7 |
| O2A  | Lys153 NZ 2.7 |

(ii) AcCoA hydrogen bonds

| Atom | Hydrogen bond | Water-mediated hydrogen bond |
|------|---------------|-------------------------------|
| O7A  | W1 2.7        | Gly154 O 2.9                  |
| O1A  | Gly156 N 3.3  |                               |
|      | Gly156 N 2.8  |                               |
|      | W2 3.4        | Gly156 N 3.1                  |
|      | W2 3.4        | Leu155 N 2.8                  |
|      | W2 3.4        | Ile155 N 2.9                  |
|      | W2 3.4        | Tyr152 O 2.7                  |
| O2A  | W3 2.7        | Gly159 N 2.8                  |
| O4A  | Ala154 N 2.8  |                               |
|      | W2 2.7        | Gly159 N 3.1                  |
|      | W2 2.7        | Leu155 N 2.8                  |
|      | W2 2.7        | Ile155 N 2.9                  |
|      | W2 2.7        | Tyr152 O 2.7                  |
| O5A  | Gly154 N 2.9  |                               |
| O9P  | Arg155 NH2 3.3|                               |
|      | Val154 N 3.1  |                               |
|      | W4 2.9        | Arg155 NH2 3.2                |
|      | W4 2.9        | Ser147 OG 2.7                 |
|      | W4 2.9        | Asp106 OD1 2.8                |
| O5P  | W5 2.9        | Ser147 OG 2.8                 |
|      | W5 2.9        | Ser106 OG 2.7                 |
| N4P  | Leu146 O 2.9  |                               |
| 0    | Leu146 N 3.0  |                               |

(iii) AcCoA van der Waals interactions

| Atom | Van der Waals interaction |
|------|---------------------------|
| C2A  | Arg155 CB 3.5             |
| C5A  | Gly156 CA 3.8             |
| C6A  | Arg155 CB 3.8             |
| C8A  | Gly156 CA 3.7             |
| C5B  | Val156 CG2 3.7            |
| CCP  | Val156 CG2 4.2            |
| CDP  | Leu146 CG 3.6             |
| CEP  | Val156 CG 3.8             |
| C8P  | Arg155 CZ 3.6             |
| CH3  | Val157 CG1 3.8            |