

CryoEM structures reveal allosteric regulation of the catalytic activity of the multi-protein human MAT enzyme complexes

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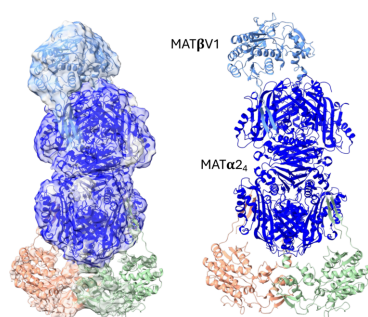
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S-Adenosyl methionine (S-AdoMet), the biological methyl donor essential for sustaining the life of most complex organisms, is the second most widely used cofactor, after ATP, in biochemical reactions and is synthesized by the enzyme methionine adenosyl transferase (MAT) from ATP and methionine. MAT, also known as S-adenosylmethionine synthetase, is found in almost every organism. S-AdoMet is employed universally by different methyltransferases that catalyze the methylation of biomolecules such as nucleic acids, proteins and lipids. In plant cells S-AdoMet produced by MAT enzymes controls the level of critical metabolites such as ethylene, polyamines and biotin, and regulates essential cellular processes such as cell division and synthesis of cell wall, chlorophyll and membrane. MAT enzyme complex MAT α 2 β , comprising the catalytic unit MAT α 2 and the regulatory protein MAT β , is found in nearly all human tissues and is essential for providing the necessary S-AdoMet flux for methylation of DNA and various proteins including histones. The enzymatic activity of MAT α 2 is enhanced by several fold upon complexation with both variants of MAT β (β V1 and β V2). Using cryogenic electron microscopy, we determined the high-resolution resting-state structures of the MAT α 2 β V1₂ and MAT α 2 β V2₂ complexes, providing insights into the allosteric regulation of MAT catalytic activity, revealing how MAT β V association could facilitate substrate binding, stabilize the transition state and promote product release to drive the catalytic cycle, and opening new possibilities for inhibitor binding.

1. Introduction

S-Adenosylmethionine (S-AdoMet), a universal methyl donor essential for cellular metabolism, is synthesized by methionine adenosyltransferase (MAT), also known as S-adenosylmethionine synthase (Markham & Pajares, 2009; Kotb & Geller, 1993). This enzyme catalyses the conversion of methionine and ATP into S-AdoMet, which serves as the principal methyl donor for a wide range of methyltransferases (MT) (Markham & Pajares, 2009). As such, MAT enzymes are critical for maintaining cellular homeostasis by supporting DNA and protein methylation, regulating mRNA m⁶A modification, and preserving molecular stability (Schlesier *et al.*, 2013; Strahl & Allis, 2000; Dominissini *et al.*, 2012; Grillo & Colombatto, 2008; Markham & Pajares, 2009). Beyond methylation, S-AdoMet functions as a key intermediate in the transsulfuration pathway, contributing to glutathione biosynthesis and redox balance, and plays an essential role in polyamine biosynthesis,



thereby supporting cell growth, differentiation and apoptosis (Fernández-Ramos *et al.*, 2025) (Fig. S1).

MAT enzymes exist in multiple isoforms with distinct roles, and their expression and complex formation are tightly regulated by cellular conditions, including metabolic state, methylation demand, oxidative stress and cell proliferation states (Ramani & Lu, 2017; Murray *et al.*, 2019). The tissue-specific distribution of MAT isozymes, together with the ability of cells to switch between isoforms, underscores the importance of regulatory mechanisms in controlling MAT function and cellular differentiation.

MAT1A encodes MAT α 1, which is predominantly expressed in the adult liver, where it maintains hepatic SAME levels and supports liver function and regeneration (Mato *et al.*, 2008). Its downregulation is associated with liver diseases such as cirrhosis and hepatocellular carcinoma (HCC) (Barbier-Torres *et al.*, 2022; Llovet *et al.*, 2021; Murray *et al.*, 2019). In contrast, *MAT2A* encodes MAT α 2, which is expressed in extrahepatic tissues and proliferating cells, and is frequently upregulated in cancers to meet increased SAME demand (Murray *et al.*, 2019). Two variants of MAT β V, the regulatory subunit, are encoded by *MAT2B*, which on forming a complex with MAT α 2 enhance SAME biosynthesis (Martínez-Chantar *et al.*, 2003; Murray *et al.*, 2019). *MAT2B* is overexpressed in aggressive cancers, such as triple-negative breast cancer (TNBC), where it contributes to tumour progression through increased proliferation and epigenetic alterations (Murray *et al.*, 2019; Peng *et al.*, 2015; Yang *et al.*, 2013; Xu *et al.*, 2019).

The MAT β V isoforms share 94% sequence identity but differ in their N-terminal regions by 20 amino-acid residues. MAT β V1 contains an extended N-terminus predicted to form a β -hairpin motif, whereas MAT β V2 possesses a shorter, unstructured loop (Murray *et al.*, 2014). These structural differences are accompanied by distinct tissue-specific expression patterns: MAT β V1 is predominantly expressed in the prostate, foetal liver, thyroid, lung, brain and adrenal gland, while MAT β V2 is mostly found in skeletal muscle and the heart (Yang *et al.* 2008; Murray *et al.*, 2019). Both isoforms are present in the thymus and kidney (Peng *et al.*, 2015; Yang *et al.*, 2013). Dysregulation of MAT β V isoforms is implicated in liver disease, cancer and neurological disorders, making them important therapeutic targets. The increasing levels of obesity have led to a high prevalence of non-alcoholic fatty liver disease (NAFLD), where higher levels of SAME are present in serum consistent with an increased synthesis of SAME in liver (Sáenz de Urturi *et al.*, 2022). Thus, MAT isozyme gene expression and subcellular localization have often been used as prognostic markers for liver, renal, breast and pancreatic carcinomas (Maldonado *et al.*, 2018).

MAT α 1 and MAT α 2 share 84% sequence identity with high structural similarity, and form a tetrameric assembly with four active sites, each composed of residues contributed by neighbouring subunits, reflecting the cooperative nature of catalysis (Kotb & Geller, 1993; Murray *et al.*, 2014, 2016; Panmanee *et al.*, 2020; González *et al.*, 2003). The enzymatic reaction proceeds via an S_N2 mechanism in which L-methio-

nine acts as a nucleophile to attack ATP to generate SAME and intermediate tripolyphosphate (PPPi). This intermediate is subsequently hydrolysed at the β - γ bond, yielding inorganic phosphate (Pi) and pyrophosphate (PPi) (Fig. S1) (O'Hagan & Schmidberger, 2010; Bailey *et al.*, 2021; Firestone & Schramm, 2017). The reaction products, SAME, PPi and Pi, are then released from the enzyme. While MAT α tetramers are catalytically active on their own, SAME production is markedly enhanced upon complex formation with MAT β V (Wan *et al.*, 2024; Panmanee *et al.*, 2019). Notably MAT α 2 activity increases by 450% and 250% in the presence of MAT β V1 and MAT β V2, respectively (Murray *et al.*, 2014). Mutation of Gln113 and Ser114 in the MAT α 2 gating loop severely impairs the enzymatic activity, which can only not be restored but enhanced compared with the wild type MAT α 2 alone upon interaction with MAT β V isoforms, underscoring the importance of protein-protein interactions in regulating catalysis (Panmanee *et al.*, 2019, 2020; Murray *et al.*, 2016).

How such regulation and control occur through protein-protein interaction and complex formation remains largely unknown due to the scarcity of structural data on protein complexes. To date, structural information on MAT complexes is limited to a crystal structure of the MAT α 2-MAT β V2 heterohexamer that we elucidated more than ten years ago (Murray *et al.*, 2014). The active site in this complex was occupied by a variety of ligands. The structure revealed an unexpected 4:2 stoichiometry and suggested that the binding of MAT β V2 to MAT α 2 is mediated by its C-terminal motif. Crystallographic efforts have not succeeded either for a resting state (ligand-free) structure of the MAT α 2-MAT β V2 complex or complexes with MAT β V1.

Here, we employed cryogenic-sample electron microscopy (cryoEM; Henderson & Hasnain, 2023) to overcome limitations associated with crystallization and determined the apo structures of the heterohexameric MAT α 2₄-MAT β V1₂ and MAT α 2₄-MAT β V2₂ complexes. These structures provide the first resting-state snapshots of human MAT enzyme complexes, revealing key features of their architecture and offering new insights into isoform-specific allosteric regulation. Our findings advance the understanding of the first step of the methionine cycle and provide a structural framework for targeting MAT β V-mediated metabolic dysregulation in disease.

2. Results

2.1. CryoEM structures of the MAT enzyme complexes (MAT α 2₄ β V1₂ and MAT α 2₄ β V2₂) in their resting states

To shed light on how the activity of tetrameric MAT α 2 increases in the presence of MAT β V1 and MAT β V2, cryoEM structures of the MAT enzyme complexes MAT α 2₄ β V1₂ and MAT α 2₄ β V2₂ were determined in their resting states. The heterohexameric cryoEM structure of the MAT α 2₄ β V1₂ complex is the first structure of this complex, consisting of a MAT α 2 tetramer flanked by a MAT β V1 dimer, and was resolved at a resolution of 2.6 Å in the core region to

approximately 6.0 Å in the peripheral regions (Fig. 1). The core, which represented a MAT α 2₄ tetramer as well as one of the MAT β V1 (referred to as the northern end) had well resolved EM density; however, MAT β V1 bound at the other terminus (referred to as the southern end) displayed a high degree of conformational flexibility evident in the EM maps.

The explanation for this rotational flexibility or wobbly conformation near the southern end is indeed offered by the cryoEM structure of the MAT α 2₄ β V1₂ complex, where the N-terminal of MAT β V1 adopts a β -hairpin motif that folds back and lodges into a hydrophobic pocket on the surface of one MAT α 2 monomer (Chain B) (Fig. 2). Several key intermolecular interactions, including ionic bonds, hydrogen bonding and cation– π interactions, are observed between the

N-terminal of MAT β V1 and the surface of MAT α 2 Chain A, increasing its buried surface area by approximately 50%. We found that a key hydrophobic interaction between MAT β V1-Gly15 and MAT α 2-His89 in one monomer (Chain B) holds MAT β V1 at one end, thereby preventing its association with the adjacent monomer (Chain A). Structural superposition of MAT β V1 onto the remaining two MAT α 2 monomers (Chains C and D) further reveals pronounced steric clashes between MAT β V1-Gly15 and MAT α 2-His89, which preclude N-terminal engagement at the opposite monomeric interfaces (Fig. 2). Consequently, at the opposite southern end, MAT β V1 is anchored solely through its C-terminal residues Val321-Phe322-His323, while the symmetric architecture of the MAT α 2 dimeric interface allows this subunit to wobble on

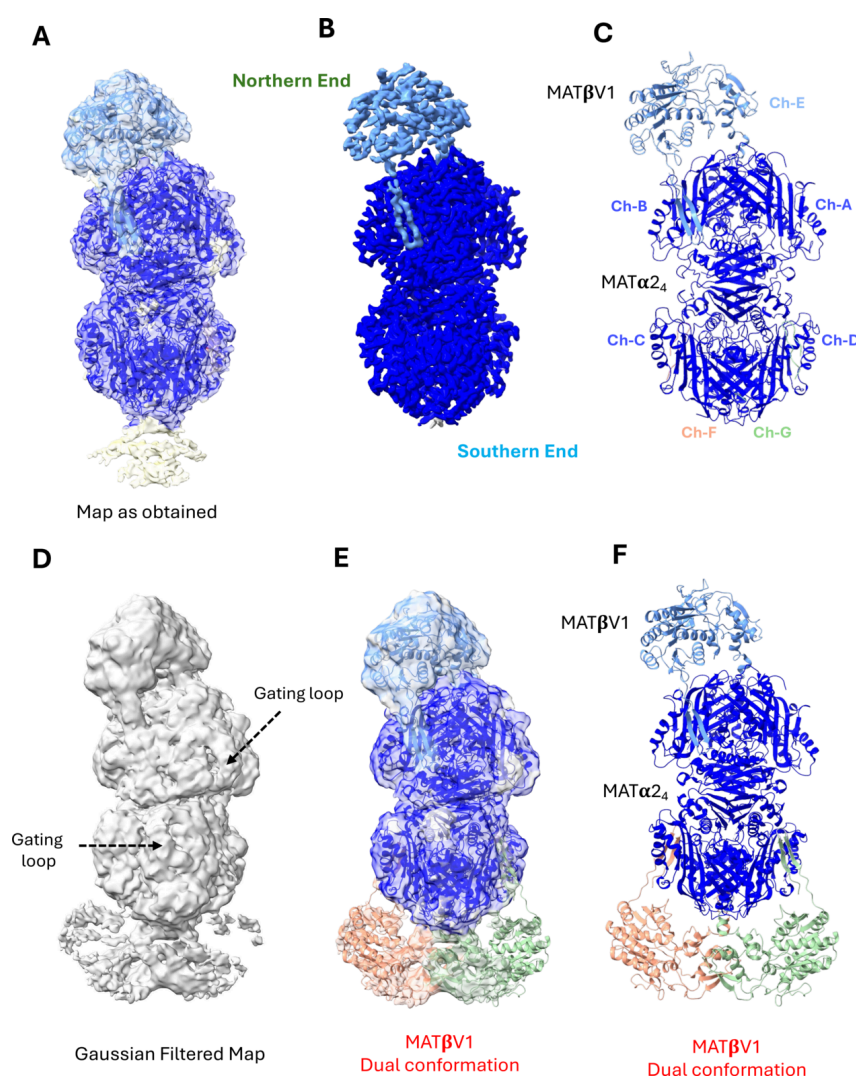


Figure 1

High-resolution cryoEM structure of the MAT α 2₄ β V1₂ complex. (A) CryoEM Coulomb potential map of the MAT α 2₄ β V1₂ complex determined by single-particle cryoEM at 2.6 Å resolution. (B) Composite cryoEM density map showing high-resolution density for the MAT α 2₄ core (blue) and high-to-medium resolution density for MAT β V1 at the northern end (cornflower blue). (C) Cartoon representation of the complex, highlighting the well defined MAT α 2₄ core (blue) and a single, well resolved MAT β V1 at the northern end (cornflower blue), alongside partially built MAT β V1 subunits at the southern end adopting dual conformations shown in salmon and green. (D) Gaussian-filtered cryoEM map revealing the two conformations of MAT β V1 at the southern end. (E) Cartoon model of MAT α 2₄ with the northern MAT β V1 (cornflower blue) and two MAT β V1 subunits (salmon and light green) at the southern end rigid-body fitted into the Gaussian-filtered cryoEM map. (F) Overall view of the MAT α 2 tetramer (blue) interacting with MAT β V1. MAT β V1 at the northern end is shown in cornflower blue, while the southern end exhibiting a dual or wobbly conformation is represented in salmon and light green.

either side (see *Methods: Model building*). However, if transiently stabilized by weaker interactions, the complex may adopt either a *syn* or *anti* conformation, depending on which side of the MAT α 2 dimer interface it engages. In both orientations, the MAT β V1 tail is stabilized by the interactions involving identical residues of the MAT α 2 dimeric interface. This rotational flexibility was evident during data processing, where in addition to the predominant high-resolution ‘wobbly’ conformation a small fraction of MAT α 2 β V1 $_2$ complexes were also observed in both *syn* and *anti* conformational states, with the peripheral MAT β V1 subunits in the southern end either aligned on the same side or on opposite sides of the MAT α 2 $_4$ core (Fig. S2). Given the dynamic nature of the complex and the conformational flexibility suggested by these cryoEM maps, 3D variability analysis (3DVA) in *CryoSPARC* was applied to explore both continuous and discrete motions

across the particle populations. This approach enabled the identification of correlated domain movements and alternative structural states within each class, providing insight into the range of conformations adopted by the MAT α 2 β V1 $_2$ complex in solution (Supplementary movies 1–5). To further examine whether the observed heterogeneity reflects coordinated motions within the complex, we performed chain-wise structural superposition using MAT α Chain B as a reference to compare the relative positioning of protomers across these classes (Fig. 3). This analysis revealed coordinated movement across MAT α 2 protomers, indicating that motions at the peripheral regulatory subunits are coupled to rearrangements within the catalytic core.

Notably, the amplitude of motion is asymmetric, and the largest conformational difference is observed between the *syn* and *anti* states, where the MAT β V1 subunit at the southern

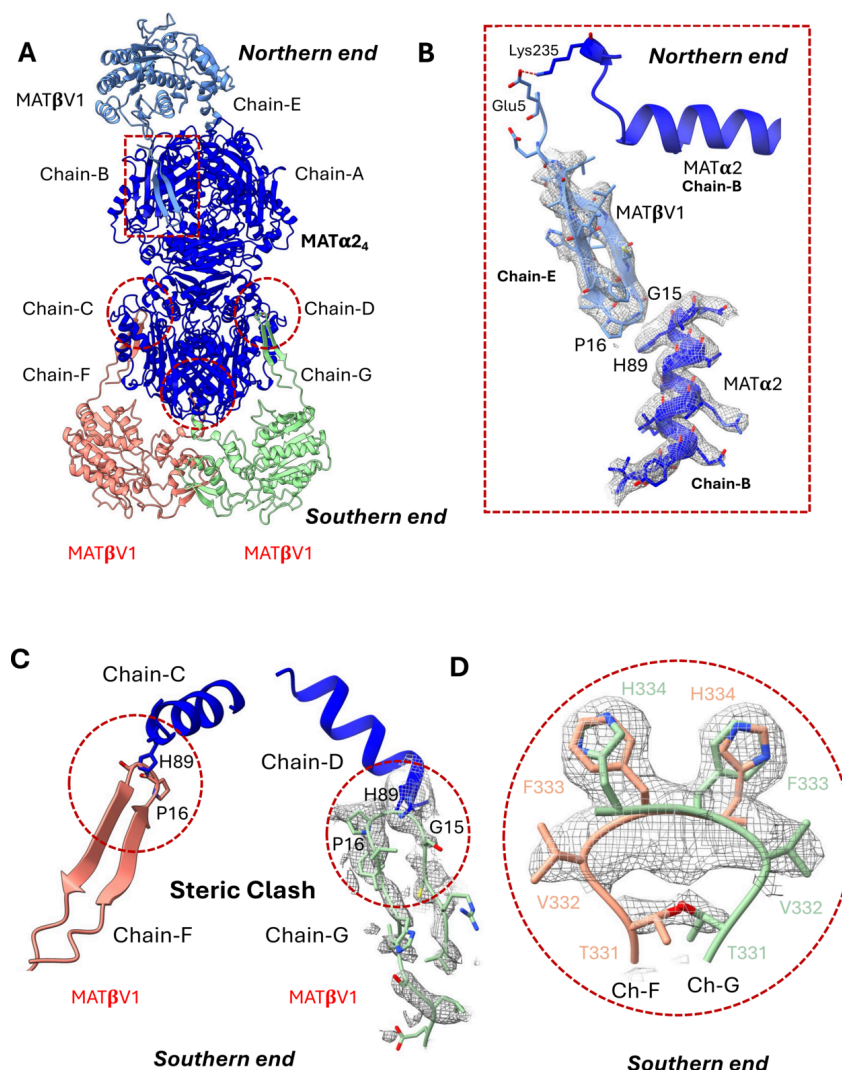


Figure 2

MAT α 2 $_2$ interaction with the MAT β V1 N-terminus. (A) Cartoon representation of MAT α 2 tetramer (marine blue) interactions with MAT β V1 (cornflower blue, salmon and light green). (B) Zoomed-in view of the MAT α 2 chain A interaction interface with the N-terminal hairpin of MAT β V1, located within the MAT α 2 groove at the northern side; the ionic bond between Lys235 MAT α 2 and Glu5 MAT β V1 is indicated by the dotted red line. His89 MAT α 2 in Chain A is in the conformation accommodating MAT β V1 for the complex formation. (C) The steric clash of His89 MAT α 2 in Chain C and Chain D preventing full binding of the MAT β V1 N-terminus (Coulomb density showing β -hairpin of one MAT β V1 at the southern end modelled with 50% occupancy). (D) Coulomb density at the southern end interface showing two possible conformations of MAT β V1 with equal probabilities of 50%.

interface undergoes a pronounced side-to-side reorientation across the MAT α 2 dimer interface. In contrast, the wobbly class represents a continuum of intermediate conformations that bridge these two extremes, resulting in an apparent reduction in displacement due to ensemble averaging. This indicates that the *syn* and *anti* conformations define the endpoints of a continuous rotational motion. *AlphaFold3* modelling is also consistent with these dual orientations of MAT α 2 β V1₂ with nearly equal probabilities (Fig. S3).

An important ionic interaction may exist between MAT β V1-Glu5 and MAT α 2-Lys235. Lys235 is part of a critical secondary structure that includes Phe250, a residue identified as important for substrate binding and product release during catalysis (Fig. 2B). Interestingly, the position of this secondary structure, and consequently that of Phe250, shifts during different stages of catalysis, highlighting the significance of this ionic interaction, which could play a role in allosteric regulation. Also, a common structural feature of the MAT enzyme complex is the ‘gating loop’ (amino acids 113–131) that flanks the active site. When the active site is occupied, this loop adopts a helical closed conformation, forming a gate over the catalytic pocket; in contrast, when the site is unoccupied, the loop becomes disordered or remains in an open conformation. The position of the gating loops in the MAT α 2 β V1₂ complex is well defined at low threshold, with all four active sites of the MAT α 2 dimeric interface existing in an open conformation state consistent with the absence of any ligand/substrate bound to the active site (Fig. S4).

To shed light on why the increase in activity of MAT α 2₄ is lower when it associates with β V2 compared with β V1 (250 vs 450%) despite 94% sequence identity, we also determined the

heterohexameric structure of the MAT α 2 β V2₂ complex in the ligand-free resting state, at 2.6 Å, comprising a MAT α 2 tetramer flanked by a MAT β V2 subunit on each end (Fig. 4). Like MAT β V1, MAT β V2 interacts with MAT α 2 by inserting its conserved C-terminal tail into the cleft at the MAT α 2 dimer interface, conferring rotational flexibility that enables engagement on both sides of the MAT α 2 active site (Fig. S5). Its N-terminal forms a disordered loop that could not be fully traced, but the MAT β V2 at the northern end is partly stabilized in the 3D map, suggesting a weaker interaction with MAT α 2 compared with MAT β V1. This explains the higher quality of the EM density near the northern end and why the dynamic conformation was predominantly observed at the southern end during data processing (Fig. S11). The greater stability of MAT α 2 β V1₂ is consistent with its higher enzymatic activity compared with MAT α 2 β V2₂, suggesting that these interactions play an important role in allosteric regulation and in enhancing the overall activity of the catalytic unit. In the MAT α 2 β V2₂ complex, all gating loops were observed in the open conformation at low threshold (see *Methods: Model building*) (Fig. S4).

The *AlphaFold3* model of MAT α 2 β V2₂ predicted MAT β V₂ in both *syn* and *anti* conformations, while the previously determined ligand-bound crystal structure captured it in the *anti* conformation only (PDB ID: 4ndn) (Fig. S3). This discrepancy underscores the importance of cryoEM data, which not only confirm the dual conformation of MAT β V but also reveal conformational heterogeneity of the MAT enzyme complexes that might be missed in crystal structures due to the selective process imposed by crystallization conditions.

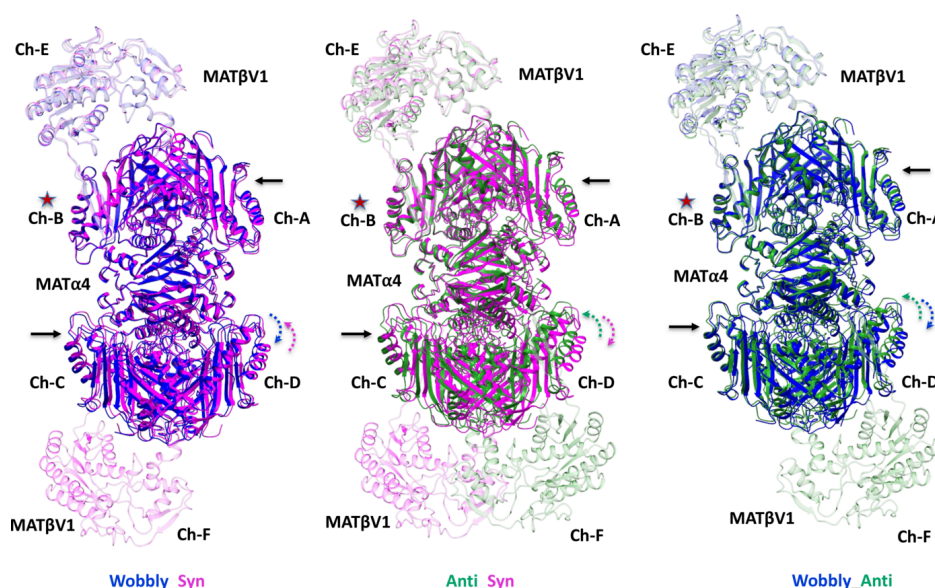


Figure 3

Chain-wise structural superposition reveals discrete conformational states and coordinated motion in the MAT α 2 β V1₂ complex. Structural superposition of the three cryoEM classes of MAT α 2 β V1₂, wobbly, *syn* and *anti*, was performed using MAT α 2 Chain B as a reference (red star) to assess relative conformational changes across the complex. The three pairwise comparisons are shown: wobbly versus *syn* (left), *anti* versus *syn* (middle) and wobbly versus *anti* (right). MAT α 2 protomers (Chains A–D) and MAT β V1 subunits (Chains E and F) are labelled. In all comparisons, the MAT β V1 subunit at the northern end (Chain E) and the adjacent MAT α 2 protomers (Chains A and B) superimpose closely, indicating a rigid, well anchored interface. In contrast, the MAT β V1 subunit at the southern end (Chain F) exhibits pronounced positional variability, accompanied by subtle but consistent shifts in the neighbouring MAT α 2 protomers (Chains C and D), highlighting propagation of motion into the catalytic core.

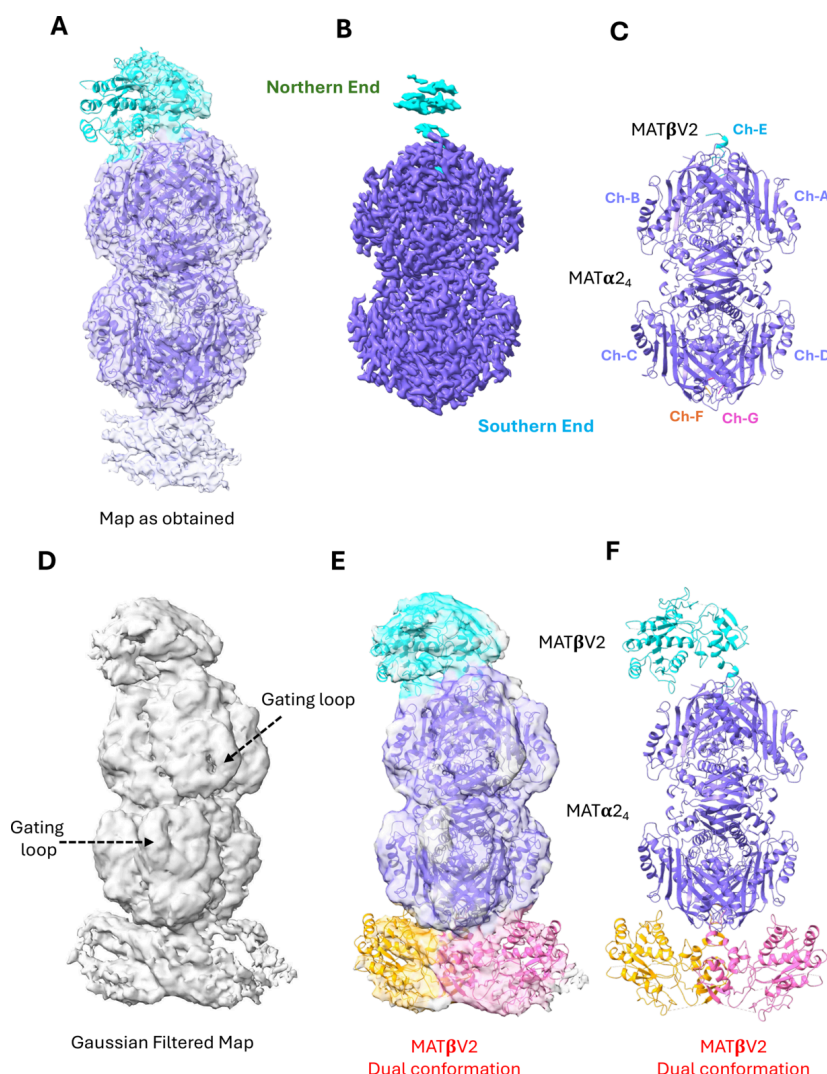


Figure 4

High-resolution cryoEM structure of the MAT α ₂₄βV₂₂ complex. (A) CryoEM Coulomb potential map of the MAT α ₂₄βV₂₂ complex determined by single-particle cryoEM at 2.6 Å resolution. (B) Composite cryoEM density, displaying high resolution for the MAT α ₂₄ core (medium slate blue) and lower resolution density for MATβV₂ at the northern end (cyan). (C) Cartoon of the enzyme complex showing better-defined MAT α ₂₄ (medium slate blue) and single MATβV₂ (cyan). (D) Gaussian-filtered cryoEM map showing a single conformation of MATβV₂ at the northern end and dual or wobbly conformations at the southern end. (E) Cartoon representation of MAT α ₂₄, with a single MATβV₂ at the northern end and two conformations of MATβV₂ rigid-body fitted into a Gaussian-filtered cryoEM map. (F) Cartoon representation of the MAT α ₂ tetramer (medium slate blue) interacting with MATβV₂. MATβV₂ at the northern end is shown in cyan, while the southern end exhibiting a dual conformation is represented in light orange and magenta.

The EM density in the tetrameric core, corresponding to MAT α ₂₄, was well resolved, permitting a high confidence in model building. However, the peripheral region exhibited varying degrees of structural flexibility, with MATβV₂ bound to one MAT α ₂ dimer adopting a partially stable conformation at the northern end while the other, bound to the second MAT α ₂ dimer at the southern end, displayed dynamic sampling even greater than that in MAT α ₂₄βV₁₂ with limited visibility in the final 3D reconstruction. Gaussian-filtered density maps clearly revealed a single conformation of MATβV₂ at one end while a dual conformation at the other (Fig. 4). Similar to MAT α ₂₄βV₁₂, 3D variability analysis (3DVA) of the MAT α ₂₄βV₂₂ complex revealed coordinated domain rearrangements indicative of a cooperative breathing

motion. These dynamics are clearly visualized in the gating loop, which alternates between ordered and disordered states and transitions between open and closed conformations in response to rotational movements of the MATβV₂ subunits surrounding the active site (Supplementary movies 6–9).

2.2. Structural comparison of cryoEM apo MAT enzyme complexes with the ligand-bound MAT α ₂₄βV₂₂ crystal structure

The crystal structure of the MAT α ₂₄βV₂₂ holoenzyme complex was obtained in its post catalytic state where the products, such as SAMe, PPNP (an imidotriphosphate, a non-hydrolyzable triphosphate analogue) and metal ions, were

trapped in the active site (Murray *et al.*, 2014). Superposition of the MAT α 2 β V1₂ apo cryoEM structure onto the SAME + PPNP-bound crystal structure of the MAT α 2 β V2₂ complex revealed structural differences with an overall r.m.s.d. of ~ 1.1 Å. Noticeable conformational differences were observed in the MAT β V subunits, where MAT β V1 appeared more stable than MAT β V2 and is shifted downward towards the MAT α 2₄ core by establishing additional contacts through its N-terminal β -hairpin motif. For example, subtle shifts were observed in MAT β V2 positioning, and the MAT α 2 dimer interface cavity appeared more dilated in the ligand-bound complex compared with the apo structure. As a result, the three C-terminal residues of MAT β V2 (Val321, Phe322, His323) in the ligand-complexed structure are repositioned deeper into the MAT α 2 dimeric interface, strengthening both ionic and hydrophobic interactions (Fig. S6). MAT β V2-His323 interacts with MAT α 2-Gly273 in Chain A and MAT α 2-Gly275 in Chain B, both of which are located within a critical loop containing key active site residues, including Lys285 and Lys265, that recognize the tripolyphosphate of ATP at the active site and play a direct role in catalysis. In the product-bound crystallographic structure, the distance between MAT β V2-His323NE2 and MAT α 2-Gly273 (Chain A) is 2.8 Å, compared with 4.1 Å in the apo MAT α 2 β V1₂ cryoEM structure, resulting in the loss of a hydrogen bond. A pronounced structural rearrangement was observed in the gating loop (residues 113–135), which was poorly resolved in the crystal structure for the open conformation but clearly visible in the cryoEM maps of MAT α 2 β V1₂ and MAT α 2 β V2₂ at low threshold. In the apo state, the distance between Phe250 (Chain B) and Ile117 (Chain A) is ~ 24 Å, whereas in the SAME-bound closed conformation, this distance decreases to ~ 7.5 Å, reflecting a shift of ~ 18 Å (Fig. S6). This movement is particularly evident in the position of Phe250, which participates in π – π stacking with the SAME adenine ring in the product-bound structure.

Despite large changes in the SAME binding region, particularly the position of Phe250, the positions of Glu23, His29, Lys181, Lys265, Lys285 and Asp291 which are involved in PPP binding (PPNP in the structure) are preserved in the product-bound and apo structures (Fig. S6). This suggests that the overall structural rearrangement occurs primarily in the SAME binding/producing region, while the PPNP-binding region remains structurally conserved, explaining previous findings showing that SAME production and ATP hydrolysis are independent of each other (Panmanee *et al.*, 2019). The dynamic motion in this region is further elucidated by comparing frames 1 and 20 of the volume series derived from 3D variability analysis (3DVA) of the final consensus map of the MAT α 2 β V2₂ complex. The analysis reveals reversible remodelling of gating loops across all protomers, inter-converting between open and closed states. These concerted transitions are coupled to rotational rearrangements of the MAT β V2 subunits flanking the active site (Fig. S6; Supplementary movie 9).

These observations are also consistent with the comparison between the apo cryoEM and SAME + PPNP-bound crystal

structures of MAT α 2 β V2₂, suggesting that the mode of substrate binding and product release is conserved between MAT α 2 β V1₂ and MAT α 2 β V2₂ and any differences in the rate of SAME production are due to differences in protein–protein interaction with different regulatory MAT β V subunits (β V1 and β V2) differentially facilitating substrate entry/binding and product release from MAT α 2 active sites. These structural changes upon complex formation in MAT α 2 β V2₂ and MAT α 2 β V1₂ are thus responsible for enhanced SAME production compared with MAT α 2₄ alone. Furthermore, greater stability of MAT α 2 β V1₂ may help to provide a better configuration for catalysis and thus higher activity than MAT α 2 β V2₂.

2.3. Allosteric regulation by MAT β V

The binding of MAT β V2 to MAT α 2 induces a series of conformational changes that propagate from the C-terminal cavity to the active site, influencing substrate binding, catalysis and product release. Previous studies have proposed that MAT β V2 regulates the catalytic activity of MAT α 2 by altering its kinetic properties, specifically enhancing its affinity for L-methionine while reducing its sensitivity to SAME inhibition (Wan *et al.*, 2024; Panmanee *et al.*, 2019; LeGros *et al.*, 2001). However, conflicting findings from a separate study suggest that MAT β V2 binding primarily stabilizes MAT α 2 without significantly influencing its catalytic activity or SAME inhibition (Bailey *et al.*, 2021).

In this study, we explored the structural basis of MAT β V-mediated regulation of MAT α 2 by comparing the apo MAT α 2 (PDB ID: 6faj) and the product-bound (SAME + PPNP) MAT α 2 β V2₂ (PDB ID: 4ndn) crystal structures with the cryoEM MAT α 2 β V2₂ and MAT α 2 β V1₂ structures in their ligand-free (resting) states. Pairwise structural comparisons revealed that MAT β V binding induces significant rearrangements in key catalytic residues near the active sites (Fig. S7). A central conformational shift involves Phe250, which, together with Ser247, Arg249 and Ile322, forms a critical pocket that stabilizes the adenine ring of SAME during product formation. In the cryoEM structures, these residues adopt intermediate conformations, bridging the apo MAT α 2 (PDB ID: 6faj) and product-bound (MAT α 2 β V2₂; PDB ID: 4ndn) structures. These intermediate rearrangements likely prime the active sites that can support more effective substrate binding, product formation and its release, highlighting a dynamic cycle of allosteric modulation contributing to efficient catalysis *i.e.* a higher level of SAME formation compared with MAT α 2₄ alone. Our findings demonstrate that the N-terminal loop and C-terminal tail of MAT β V work in concert for repositioning Phe250, Arg249 and surrounding catalytic residues, effectively fine-tuning the active site geometry.

A markedly impaired SAME synthesis has been demonstrated through mutational analysis of residues within the MAT α 1 active site, which is highly conserved and structurally similar to MAT α 2, (Fernández-Irigoyen *et al.*, 2010). Structural mapping of these mutations at Arg249, Ile252, Gly257, Asp258 and Ala259 resulted in defective methionine binding

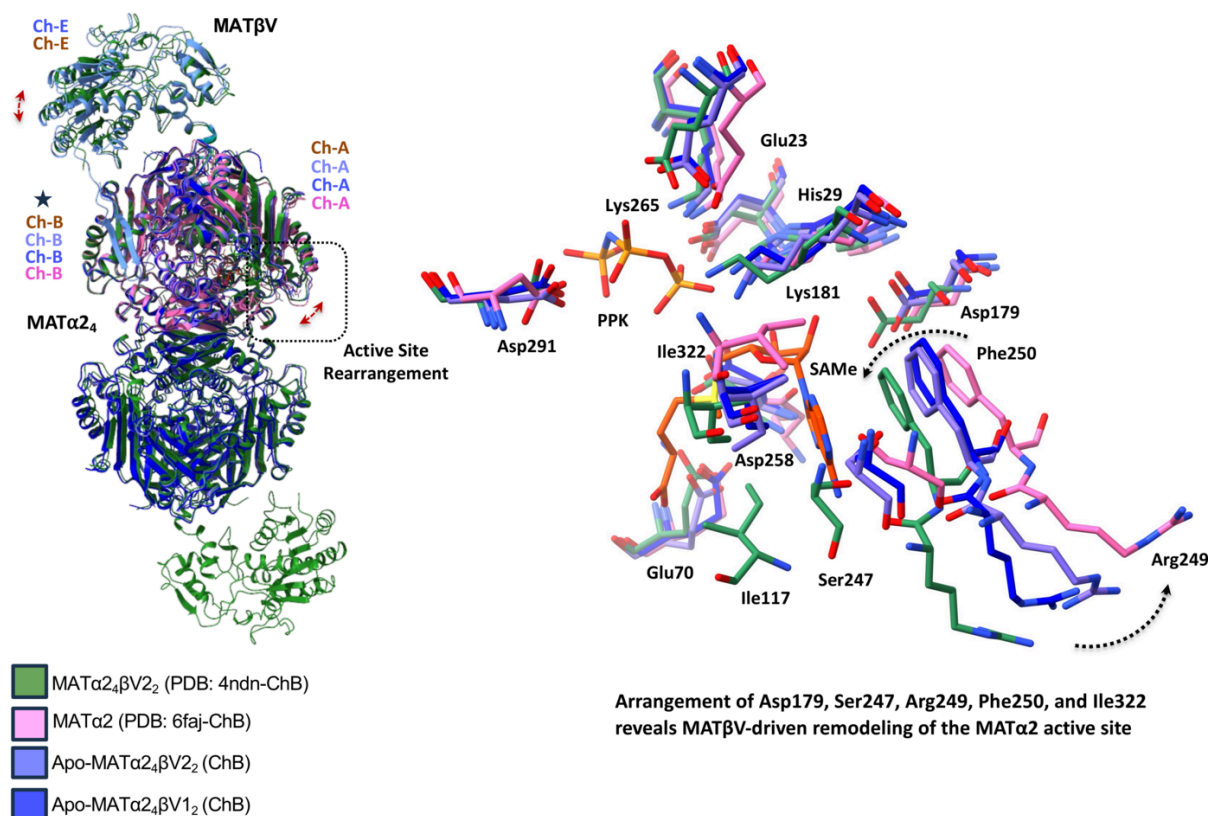


Figure 5

Key structural insights into control, regulation and allosteric mechanism of MAT enzyme complexes. Chain-specific structural superposition of the MATα2 protomer in apo MATα2 (PDB ID: 6faj, Chain B), the SAMe + PPNP-bound MATα2βV2 complex (PDB ID: 4ndn, Chain B), and the resting-state cryoEM structures of MATα2βV2 (Chain B) and MATα2βV1 (Chain B) illustrates allosteric conformational changes near the active-site pocket upon MATβV binding. The right inset presents an expanded stick view of the active-site pocket, highlighting the spatial orientation of key residues and showing how MATβV modulates the architecture of the MATα2 active site. The MATα2 protomers used as references for chain-specific structural superposition are indicated with asterisks, and all structures are colour-coded as indicated.

in MATα1. Arg249 is positioned between Phe250 and Ser248, within a flexible loop implicated in methionine entry and positioning within the active site.

Based on cryoEM maps of the MATα–MATβV complexes captured in multiple conformational states and 3D variability analysis, we propose that MATβV functions as an allosteric regulator, whose structural flexibility facilitates catalysis by modulating substrate entry, binding and product release, with each orientation (*syn* or *anti*) potentially favouring a distinct step of the reaction in response to different binding partners and varying metabolic or cellular conditions (Fig. S8).

Notably, the N-terminal β-hairpin of MATβV1, present in the MATα2βV1 complex which forms additional stabilizing interactions at the MATα2–MATβV1 interface, likely contributes to higher enzymatic activity compared to the complex with MATβV2.

3. Conclusion

Although it has been recognized that protein–protein interactions are fundamental to many control mechanisms *in vivo*, our understanding of how protein–protein interactions regulate enzyme activity, for example, has remained limited owing to a scarcity of high-resolution structures of complexes

comprising the catalytic enzyme and regulatory protein (Keskin *et al.*, 2008; Jones & Thornton, 1996; Garner & Janda, 2011; Luck *et al.*, 2020). This is also the case for methionine adenosyl transferase (MAT) enzyme complexes that catalyse the formation of SAMe by utilizing methionine and ATP. Higher levels of SAMe formation are achieved when the catalytic subunit MATα complexes with different isoforms of the regulatory subunit MATβ. The ligand-bound (with SAMe, PPNP and metal ions bound) crystallographic structure of the MATα2βV2 complexes (Murray *et al.*, 2014) together with the cryoEM structures of MATα2βV1 and MATα2βV2 complexes, resolved here in their apo forms, provide a structural roadmap of the enzyme's catalytic cycle, allowing us to shed light on the structural elements in MATβV that are likely to be important for complex formation and allosteric regulation.

Structural comparison between ligand-bound crystal structures and the cryoEM apo structure of the MATα2βV2 complex reveals a pronounced repositioning of MATβV2, accompanied by dilation of the cavity at the MATα2 dimer interface, consistent with observations from 3D variability analysis for both heterocomplexes. These interactions promote structural rearrangements that affect the gating-loop conformation that could reposition Arg249 and Phe250, key

residues for controlling substrate access and product release (Fig. 5). These comparisons also allow us to conclude that the greater stability of MAT α 2–MAT β V1 compared with MAT α 2–MAT β V2 arises due to the presence of the N-terminal β -hairpin, which interacts with a hydrophobic patch on the MAT α 2 surface. MAT β V2 lacks the N-terminal β -hairpin and relies solely on the C-terminal tail for binding to MAT α 2, resulting in a less stable complex.

The stability of the complex correlates well with the enhancement of enzymatic activity of MAT α 2, which is increased substantially but to varying levels when complexed with MAT β V1 and MAT β V2, respectively (Murray *et al.*, 2014). The stable nature of these complexes, admittedly of varying degree of stability, and their direct impact on the catalytic activity suggest that these complexes represent obligatory associations of MAT α 2 and β V1/ β V2 with profound functional impact and as such they are also likely to be more relevant for drug development and *in vivo* testing. The obligatory associated complexes differ from the transient encounter complexes, such as the electron transfer complexes, where the heterocomplex interfaces are generally less extensive and the partner proteins have weaker associations (Keskin *et al.*, 2008). Unlike the obligatory enzymatic complexes, the interaction between partner proteins in an electron transfer complex, such as cytochrome *c* and cytochrome *c* oxidase, is weak and dynamic, enabling frequent on-off exchange of partner proteins (Sakamoto *et al.*, 2011). The presence of multiple conformational states (wobbly, *syn* and *anti*) suggests that the β -hairpin insertion in MAT β V1 not only enhances overall complex stability but also restricts the extent of conformational sampling, thereby modulating long-range allosteric communication between distal regulatory sites and the active site.

The differential activity of MAT α 2–MAT β V1 and MAT α 2–MAT β V2 suggests distinct regulatory roles of MAT β V in SAME production, potentially adapting to specific metabolic demands and cellular conditions. The stability of the MAT α 2–MAT β V1 complex may provide a robust platform for sustained enzymatic activity, ensuring a steady supply of SAME for methylation reactions. This is particularly important in rapidly dividing cells with high metabolic demands, such as cancer cells, embryonic stem cells and the immune cells during activation. In contrast, the transient nature of the MAT α 2–MAT β V2 complex may limit its ability to sustain high levels of SAME production over extended periods, making it less suitable for tissues with constant high metabolic demands. This tiered regulation – with MAT β V1/V2 serving as metabolic rheostats – ensures precise SAME allocation, while its dysregulation creates therapeutic vulnerabilities. For example, the liver's MAT1A-to-MAT2A switch in hepatocellular carcinoma (HCC) and the overexpression of MAT β V1 in aggressive cancers such as triple-negative breast cancer (TNBC) exemplify how shifting complex composition mirrors pathological SAME requirement (Murray *et al.*, 2019; Xu *et al.*, 2019).

The cryoEM structures in this study also help to explain why MAT α 1 selectively forms a stable complex *in vitro* with MAT β V1 but not with MAT β V2, despite sharing high

sequence and structural identity with MAT α 2 (Panmanee *et al.*, 2020). Structural studies of hetero complexes of MAT α 2 show that stable complex formation requires two key interaction modules in MAT β V: the C-terminal tail, which inserts into the MAT α dimeric interface, and the N-terminal β -hairpin, which engages a hydrophobic surface patch. Because MAT β V1 contains both elements, whereas MAT β V2 lacks the β -hairpin, MAT β V1 consistently binds more strongly to MAT α subunits and enhances catalytic output to a greater extent. *In silico* modelling of MAT α 1–MAT β V complexes further revealed that although MAT β V2 can fit into the MAT α 1 interface without steric clashes, variant residues in MAT α 1 – such as the substitutions of Arg192→Asn and Ile196→Leu – disrupt critical ionic and hydrophobic interactions that normally stabilize the MAT α 2–MAT β V2 complex (Fig. S10). Together, these findings suggest that subtle sequence and structural divergence among MAT β isoforms modulates their interaction landscape, thereby fine-tuning MAT α activity under specific cellular conditions. Despite substantial conformational heterogeneity, these complexes remain stable, indicating that MAT α 2 forms obligatory functional associations with β V1 and β V2. These complexes, therefore, represent the biologically relevant state of the enzyme and are appropriate targets for drug development and *in vivo* studies, particularly in cancer, where MAT α 2 and the two major MAT2B variants (MAT β V1 and MAT β V2) are overexpressed, conferring growth advantage.

4. Materials and methods

4.1. Protein expression and purification of MAT α 1, MAT α 2, MAT β V1 and MAT β V2

All the MAT enzyme constructs (MAT α 1, MAT α 2, MAT β V1 and MAT β V2) were produced in *E. coli* BL21 (DE3 strain) and grown in lysogeny broth (LB) with 50 mg ml^{−1} of kanamycin for antibiotics selection, as described in detail before (Panmanee *et al.*, 2019). Cells were lysed using a hydraulic press cell disruptor at 28 000 p.s.i and cell debris was then removed by centrifuging at 20 000g for 45 minutes at 4°C. The proteins were eluted with elution buffer [25mM HEPES pH 7.5, 500mM NaCl, 250mM imidazole, 5%(v/v) glycerol and 1mM DTT]. The fractions containing the MAT α 1 or MAT α 2 were pooled and incubated overnight with tobacco etch virus (TEV) protease in a ratio of 100:1 (protein:protease) at 4°C in dialysis buffer [25mM HEPES pH 7.5, 500mM NaCl, 5%(v/v) glycerol and 1mM DTT]. The fractions containing MAT β V1 or MAT β V2 were incubated with Sentrin Specific Protease 2 (SEN2) and dialyzed overnight. Final proteins were eluted in SEC buffer [25mM HEPES pH 7.5, 500mM NaCl, 5%(v/v) glycerol and 0.5mM TCEP], concentrated, and stored at −80°C.

4.2. Formation of the MAT α 2 β V2₂ and the MAT α 2 β V1₂ complexes

The complex formation protocol was performed as previously reported (Murray *et al.*, 2014). Briefly, MAT α 2 was

incubated with MAT β V2 or MAT β V1 in the buffer (50mM HEPES pH 7.5, 150mM NaCl and 1mM DTT) for 2 h at 4°C. Samples were loaded onto a Superdex 200 10/300 gel filtration column and the complexes were eluted. Fractions containing each complex were pooled and concentrated to 5 mg ml⁻¹ and quantified using the Bradford assay.

4.3. CryoEM grid preparation and data collection

MAT α 2 β V1 and MAT α 2 β V2 were diluted from 5 mg ml⁻¹ to optimal concentrations of 0.125 mg ml⁻¹ and 0.25 mg ml⁻¹, respectively. Quantifoil R 1.2/1.3 300 mesh grids (Quantifoil™) were glow discharged for 2 minutes at 12 mA, within a PELCO easiGLOW glow discharging device. Vitrification and grid formation was undertaken using the Vitrobot Mark IV vitrification device. 3.5 μ l of sample was applied to the grids, with a wait time of 10 s before blotting, and with a blot time of 6 s and blot force of 1. Imaging was performed with a Titan Krios G2 microscope, operated at 300 kV with an X-FEG. Movies were collected in *EPU* software (version 3.8), using a Falcon 4i detector with a Selectris energy filter, at a pixel size of 0.74 Å (165 000 magnification) (Grollios *et al.*, 2024). For MAT α 2 β V1 and MAT α 2 β V2, 10 375 and 10 707 movies were collected in counting mode, using a total dose of 40 e⁻ Å⁻² and dose rate of 7.25 e⁻ pixel⁻¹ s⁻¹ and 7.41 e⁻ pixel⁻¹ s⁻¹, respectively (Supplementary Table 1).

4.4. CryoEM data processing

The cryoEM data-processing workflows for the MAT α 2 β V2 and MAT α 2 β V1₂ datasets are illustrated in Supplementary Figs. 11–19. The datasets were processed using an integrated approach combining *CryoSPARC* (version 4.6.0) and *RELION* (version 4.0.2), with *crYOLO* for particle picking (Punjani *et al.*, 2017; Scheres, 2012; Wagner *et al.*, 2019). For the MAT α 2 β V2 complex, 10 707 movies were motion corrected, followed by contrast transfer function (CTF) estimation using *CTFFIND-4.1* and 2D classification to select ~4.5 million particles (Rohou & Grigorieff, 2015). These were used for *ab initio* model building and heterogeneous refinement. A major class displaying the MAT α 2 tetramer and partial MAT β V2 density was subjected to 3D classification, yielding a well defined MAT α 2 β V2 reconstruction comprising ~135 219 particles (Class II) that was refined to 2.61 Å resolution, with focused local refinement improving the MAT β V2 density near the northern end to ~4.2 Å (see the supporting information).

Processing for the MAT α 2 β V1 dataset followed a similar procedure, where 10 375 movie stacks were motion-corrected, followed by CTF estimation, 2D classification, *ab initio* model building and heterogeneous refinement to select ~1 million particles. These were used for *ab initio* model building and heterogeneous refinement. A major class displaying the MAT α 2 tetramer and clear MAT β V1 density near the northern end, along with the dual conformation near the southern end, was subjected to 3D classification. This yielded a well resolved MAT α 2 β V1₂ reconstruction comprising ~122 584 particles (Class II) that was refined to 2.61 Å reso-

lution. Subsequent focused local refinement improved the MAT β V1 density at the northern end to ~3.3 Å. A subset of ~51 300 particles from Class IV, featuring MAT β V1 subunits in *anti* conformation relative to the MAT α 2₄ core, was refined to a resolution of 3.2 Å. Using an alternative focused 3D classification strategy, ~95 000 particles (Class VI) corresponding to the MAT α 2 β V1₂ complex in a *syn* conformation – where both MAT β V1 subunits are aligned in the same direction – were identified and refined to a final resolution of 2.85 Å. 3D variability analysis in *CryoSPARC* was conducted on the final particle sets to characterize both discrete and continuous structural heterogeneity of the reconstructed volumes (see the supporting information) (Punjani *et al.*, 2017).

4.5. Model building

Model building for the resting state of MAT α 2 β V2₂ was initiated by segmenting the high-resolution ligand-bound MAT α 2 β V2₂ crystal structure (PDB ID: 4ndn) into MAT α 2 and MAT β V2 subunits. The MAT α 2 subunits were rigid-body fitted into well resolved regions of the cryoEM density corresponding to the tetrameric core using *ChimeraX* (Meng *et al.*, 2023). All gating loops were observed in an open conformation at low threshold (Fig. S4), and model building was performed manually in *COOT* only for the gating loop in Chain D, while those in the remaining chains were left unmodelled (Emsley & Cowtan, 2004). The EM density at the northern end enabled modelling of only the C-terminal residues of MAT β V2 (residues 311–323), whereas the N-terminal region and the remainder of the protein (residues 1–310) were fitted as a rigid body into the EM map and were not included in the deposited structure (PDB ID: 9qpo). Notably, the last four C-terminal residues of MAT β V2 at the southern end (Thr320, Val321, Phe322 and His323) were modelled in two distinct conformations, each with 50% occupancy. Subsequent refinement was carried out in *Phenix* (Liebschner *et al.*, 2019). The remaining portions of the MAT β V2 subunit near the southern end were rigid-body fitted into the low-threshold map in dual conformation at equal occupancies (50%) for structural interpretations (not included in the deposited structure), reflecting the conformational heterogeneity and wobbly nature of this region.

Model building for the MAT α 2 β V1₂ resting state EM map followed a similar strategy, using segmentation of the ligand-bound MAT α 2 β V2₂ crystal structure (PDB ID: 4ndn) into MAT α 2 and MAT β V1 subunits, followed by rigid-body fitting into the cryoEM density using *ChimeraX* (Meng *et al.*, 2023). The gating loop was manually built in *COOT* for Chain B using low-threshold EM maps, while those in the remaining chains were left unmodelled. The MAT β V subunit at the northern end was better resolved and fitted the EM density more clearly, allowing modelling of missing N-terminal residues in *COOT* (Emsley & Cowtan, 2004). Notably, at the southern end, the N-terminus of MAT β V1 (Chain F and Chain G; residues 7–19) was built into weak EM densities on both sides of the MAT α 2 dimeric interface (Fig. 2C), showing steric

clash and unstable interaction, while the C-terminal end residues (332–334) were modelled in two distinct conformations. Subsequent refinement was performed using the Servalcat *REFMAC5* controller implemented in the *CCP-EM* package (Burnley *et al.*, 2017). The remaining portions of each MAT β V1 subunit near the southern end were rigid-body fitted into the low-threshold map at equal occupancies (50%) only for structural interpretations and were not included in the deposited structure (Figs. 1E and 4E), reflecting substantial flexibility and conformational heterogeneity in this region.

Model building of the MAT α ₂ β V1₂ complexes in the *syn* and *anti* conformations included the MAT α ₂ core and two MAT β V1 subunits rigid-body fitted into their respective positions using *ChimeraX* and *iSOLDE*, followed by real-space refinement in *Phenix* (Meng *et al.*, 2023; Croll, 2018; Liebschner *et al.*, 2019). The final structures (PDB IDs: 30gd and 30gh) excluded gating loops, which could not be modelled owing to insufficient density. The stereochemical quality of all structures was assessed using *COOT* and *MolProbity*, and comprehensive model validation was performed using *Phenix* (Supplementary Tables 1 and 2) (Emsley & Cowtan, 2004; Davis *et al.*, 2007). Figures were generated using *PyMOL* (<https://www.pymol.org>) and *ChimeraX* (Meng *et al.*, 2023).

The MAT α ₂ protomer used as a reference for chain-specific structural superposition is indicated with an asterisk in Fig. 5. Notably, the active site of MAT α ₂ (Chain B) in the product-bound MAT α ₂ β V2₂ complex (PDB ID: 4ndn) is unoccupied; therefore, this protomer was selected as the reference for structural comparison with apo MAT α ₂ (PDB: 6faj, Chain B) and the resting-state cryoEM structures of MAT α ₂ β V2₂ (Chain B) and MAT α ₂ β V1₂ (Chain B). MAT β V adopts a similar conformation across these structures, enabling identification of structural changes and comparison of active-site residues in the ligand-bound MAT α ₂ β V2₂ complex (PDB ID: 4ndn, Chain A), apo MAT α ₂ (PDB ID: 6faj, Chain A), and the cryoEM structures of MAT α ₂ β V2₂ and MAT α ₂ β V1₂ (Chain A).

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