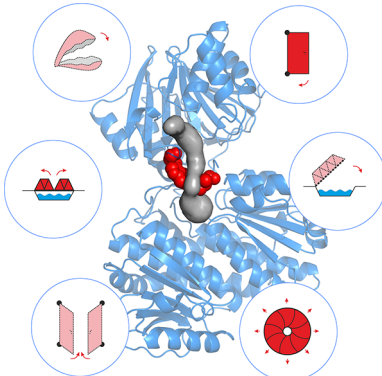


Gates of Enzymes

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1. INTRODUCTION

Enzymes are very efficient catalysts that are essential for the functioning of living organisms. The low efficiency of biocatalysts produced *de novo* relative to those that have

evolved naturally demonstrates that our understanding of enzymatic catalysis is still incomplete.^{1–4} The dynamic motion of enzymes during catalytic events is one of the many aspects of protein chemistry that are currently insufficiently well understood.^{5–9} On one hand, proteins need to have well-defined and organized structures in order to maintain stable functionality in the intracellular environment. On the other hand, some degree of flexibility is often required for catalytic activity. Molecular dynamics simulations have provided key insights into the importance of protein dynamics in catalysis, such as the observation of substrate access and product exit pathways that cannot be identified by inspecting crystal structures.¹⁰ Csermely et al. recently reported that mutations in regions that affect protein dynamics, such as hinge regions that are important in substrate binding, can have dramatic effects on catalytic activity.¹¹ In this review, we highlight the role of protein gates as another class of highly dynamic structures that play key roles in protein function.

Given the importance of gates for enzymatic catalysis, the number of studies that have examined them systematically is surprisingly small. Conformational gating in proteins was first described by McCammon and co-workers in 1981, but there have been relatively few systematic studies in this area since then.^{12–14} Moreover, much of the available data on gates in macromolecular systems is hidden or otherwise dispersed within the scientific literature, partly because there is currently no consensus regarding what defines a gate. Some authors describe all residues that affect the ligand's access to a target area as gating residues, whereas others apply the term exclusively to structural features that undergo large movements during the gating event. In this review, we define a gate as a dynamic system consisting of individual residues, loops, secondary structure elements, or domains that can reversibly switch between open and closed conformations and thereby control the passage of small molecules—substrates, products, ions, and solvent molecules—into and out of the protein structure. Under this definition, the anchoring residues that stabilize the open or closed conformations of a gate are not themselves gating residues. However, because of their various interactions with the gating residues, they can control the size and properties of the ligands that pass through the gate as well as the frequency of the exchange events.

Gates can be found in various systems, including enzymes, ion channels, protein–protein complexes, and protein–nucleic acid complexes.^{14,15} In this work, we focus specifically on gates in enzymes. We attempt to answer three basic questions—why, how, where—by describing the molecular function, structural

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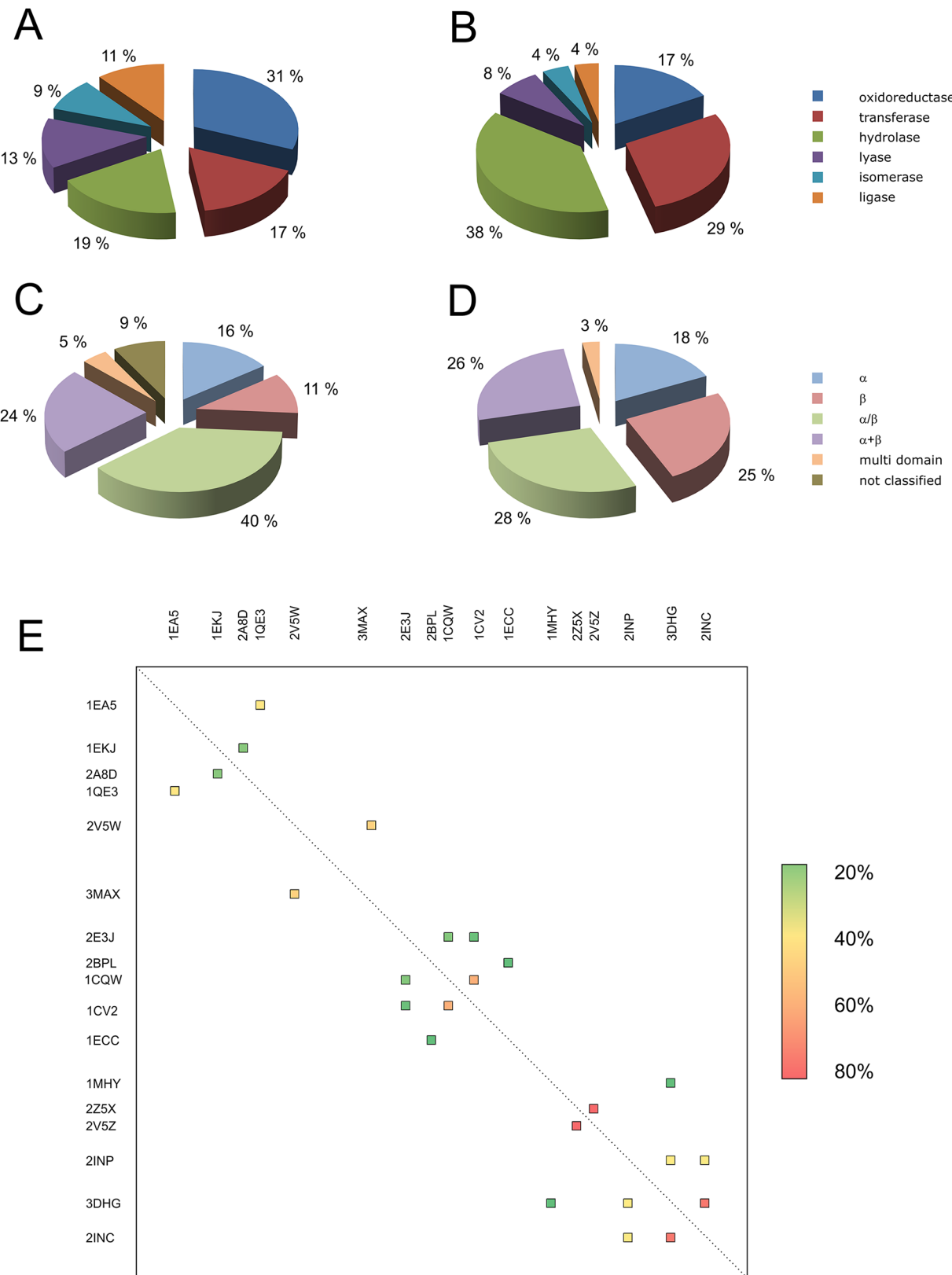


Figure 1. Distribution of (A) reviewed proteins and (B) proteins from the PDB database according to EC classes, (C) reviewed proteins and (D) proteins from the PDB database according to SCOP classification, and (E) identity matrix of reviewed enzymes (only the enzymes with sequence identities above 20% are shown for clarity).

basis, and location of gates within protein structures. We discuss 71 illustrative examples of enzymes that together contain 129 different molecular gates and propose a system for

their classification. Reviewed enzymes were chosen based on a literature search with a set of keywords corresponding to gates and conformational changes in enzymes. A preliminary set of

protein structures was filtered out, leaving only those entries for which both open and closed conformations were described. The final set represents different classes of enzymes (Figure 1A) and different protein folds (Figure 1C) and spans structurally and functionally the entire enzyme world. Among 71 chosen enzymes, only 17 have higher than 20% sequence identity with other set members (Figure 1E). The proposed classification system provides a useful framework for comparing gates of different enzymes and drawing general conclusions about gate function, structure, and position. Moreover, the classification scheme is easily extendable to describe the new gate types that will almost certainly be revealed by structural and functional analyses of newly isolated enzymes in the future.

2. MOLECULAR FUNCTION OF GATES

Analyses of protein dynamics have identified a number of enzymes with gates, suggesting that these structures are rather common. What is the molecular function of the gates? It seems that in enzymes they facilitate precise control over processes that are directly linked to catalysis. Enzyme gates can (i) contribute to enzyme selectivity by controlling substrate access to the active site, (ii) prevent solvent access to specific regions of the protein, and (iii) synchronize processes occurring in distant parts of the protein (Figure 2). The proper function of

site can also affect enzyme selectivity. Substrate access pathways, which often incorporate molecular gates, impose additional constraints on ligand binding to the active site.²⁰ The ability of ligands to traverse these access pathways can be controlled by (i) size discrimination at the narrowest point along the pathway forming a bottleneck, (ii) geometrical constraints, e.g., the curvature of the pathway, and (iii) specific molecular interactions such as hydrogen bonds, electrostatic interactions, and hydrophobic interactions with the residues comprising the access pathway. Protein gates can be regarded as molecular filters that discriminate between molecules as similar as molecular oxygen and carbon monoxide in NiFe hydrogenases^{21,22} or water and hydroperoxide in catalases.^{23,24} Gates act as filters in a wide range of enzymes, controlling the range of substrates that can be accepted by broad-specificity cytochromes P450,²⁵ the stereospecificity of epoxide hydrolases,²⁶ and product length in undecaprenyl-pyrophosphate synthases.²⁷

One of the first systematic descriptions of the influence of the gating process on substrate binding was reported by Szabo et al., who assumed that the switching between the open and the closed conformations of the gate was a stochastic process.^{28–30} This model was successfully used to demonstrate that despite conformation gating¹³ acetylcholinesterase can bind acetylcholine with a rate constant of $10^9 \text{ M}^{-1} \text{ s}^{-1}$ and predict the rate of formation of the enzyme–substrate complex in choline oxidase.³¹ Since gates create a barrier on the substrate access pathway, the kinetic rate constant for passage over the barrier can be obtained using Kramers' reaction rate theory or its later modifications.^{32–34} This methodology was used to compare the results of computational and experimental studies on the passage of the tetramethylammonium cation through acetylcholinesterase³⁵ and migration of ammonia through carbamoyl phosphate synthetase.³⁶

2.2. Control of Solvent Access

Spatial localization of the hydrophobic and hydrophilic regions within the structure of a protein is important in maintaining its proper fold and can also be crucial for catalytic function. The various steps of an enzymatic reaction may require different environments. These distinct environments can be generated by having the individual steps occur in spatially separate regions of the protein, but this does not eliminate the problem of transporting the substrate between these sites. There are important problems to be addressed, including transporting polar molecules from a polar environment to a nonpolar one and separating hydrophilic compartments from hydrophobic ones within the structure of a single protein.

In some proteins, these problems are addressed by the presence of selective barriers that permit passage of solutes but not water molecules. Crystallographic and NMR data can be used to identify cavities within a protein structure accessible to water molecules. Exclusion of water from some parts of the cavity, such as the active site or a specific tunnel, is essential for functioning of numerous enzymes. In simple cases, the gates may prevent the entrance of water molecules into the cavity when a substrate or a cofactor is not present, as occurs in rabbit 20a-hydroxysteroid dehydrogenase.³⁷ In more complex cases, the gates may permit access only to a specific part of the cavity, as occurs in carbamoyl phosphate synthetase³⁶ and imidazole glycerol phosphate synthase.³⁸ In the cytochromes P450, a “water channel” controls hydration of the substrate in the active site, which is extremely important for cytochrome activity.¹⁶

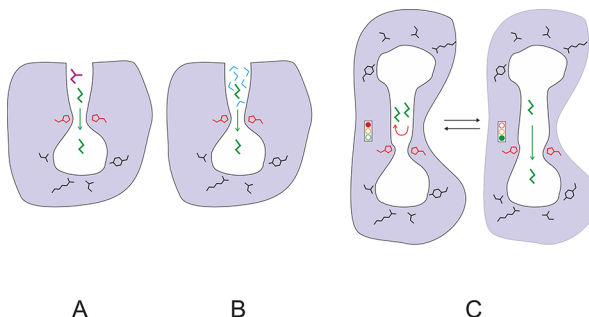


Figure 2. Schematic illustration of the molecular functions of protein gates: (A) control of substrate access, (B) control of solvent access, (C) control and synchronization of reactions. Protein is represented by the area colored in gray, active site cavity by the area in white, gating residues by red lines, substrate molecules by green or violet lines, and water molecules by blue lines.

even the simplest gates can potentially be essential for catalysis, and the gating event may even represent the rate-limiting step of the catalytic cycle. Interestingly, different gating residues within a single protein molecule may be responsible for restricting the access of specific substrates. High variability of the gating residues within an enzyme scaffold can lead to the evolution of enzyme families whose members are selective for specific substrate types. The best known example of such specialization within a single enzyme family is provided by the cytochromes P450.¹⁶

2.1. Control of Substrate Access

Enzyme selectivity has been traditionally explained by the “lock and key” model,¹⁷ which was subsequently complemented by the “induced fit” or “hand in glove”,¹⁸ “selected fit”,¹⁹ and “keyhole, lock, and key”²⁰ models. In many cases, these models provide an adequate description of enzyme selectivity based on adjustable complementarity between the active site and the cognate substrates. However, research conducted over the past decade has shown that regions located further from the active

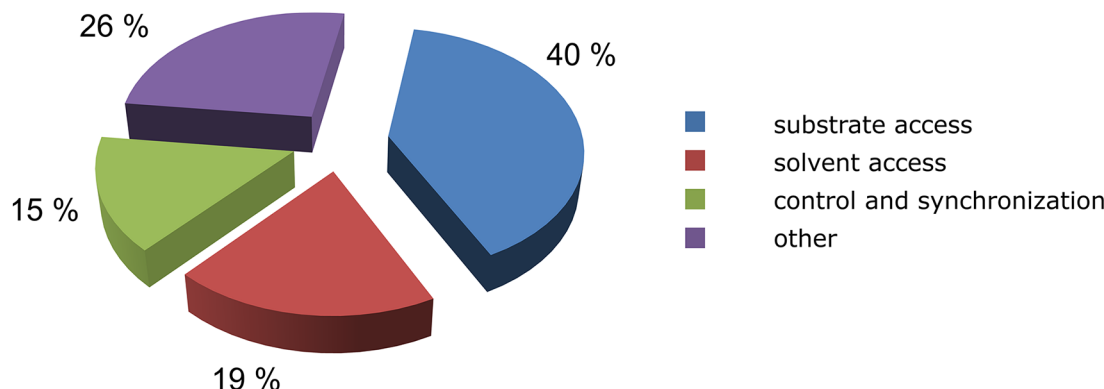


Figure 3. Distribution of gate functions; 71 proteins with 129 different gates were analyzed. Percentages shown in the figure are based on all of the identified functions of each gate; individual gate may perform multiple functions. Detailed description of the analyzed proteins is provided in Table 3.

The potential importance of gates that act as solvent barriers is further illustrated by the example of enzymes with ammonia tunnels. In these proteins, gates prevent water from entering the channel and protonating the ammonia, which is essential for maintaining its nucleophilic character.³⁹

Control of water access can be seen as a special case of the function described in section 2.1. However, when discussing water exclusion, the main emphasis is on the water permeability of the gates and their ability to distinguish water molecules from other ligands. The gate can simultaneously act as a barrier to passage of water molecules while acting as a selective filter for other molecules, allowing them to access the active site. Gates of this type resemble semiselective membranes that can distinguish between species such as water and ammonia, allowing only the second to pass. It is worth noting that passage of ‘permitted’ species can be facilitated by rearrangement of an individual gating residue, such as K99 in imidazole glycerol phosphate synthase. Conversely, passage of water molecules through gates of this type would often require significant conformational changes in all of the residues that comprise the gate.³⁸ This makes gates in enzymes far more sophisticated than semipermeable membranes.

2.3. Control and Synchronization of Reactions

Another function of gates becomes apparent when considering enzymes with two or more active sites. Many protein structures contain tunnels to facilitate efficient migration of intermediates and gates to synchronize chemical reactions. Such arrangements can be compared to a pair of workers on an assembly line. The second worker has to be ready before he can receive a product from the first one. Moreover, the products generated by the first worker must satisfy certain standards. The control gates located between the workers regulate the exchange of products over a well-defined period of time. Gates of this kind are common in ammonia-transferring enzymes, suggesting that they are old in evolutionary terms and functionally important.^{40–43} The need to efficiently transport ammonia within the interior of the protein may be related to its high cellular toxicity. We speculate that gates of this kind may be present in many enzymes that have multiple active sites connected by internal tunnels for the transport of intermediates. Many such enzymes have been studied in some detail, including carbamoyl phosphate synthetase, which has tunnels for ammonia and carbamate transportation;³⁶ asparagine synthetase,⁴⁴ glucosamine 6-phosphate synthase,⁴⁵ and glutamate synthase,⁴⁶ all of which have tunnels for ammonia transportation; tryptophan synthase for indole⁴⁷ and carbon

monoxide dehydrogenase/acetyl coenzyme A synthase for carbon monoxide transportation.⁴⁸

Systematic analysis of the functions of the known gates in the 71 proteins discussed in this article revealed the following distribution of gate types: 40% of the studied gates control substrate access, 19% control solvent access, 15% control and synchronize catalytic events, and 26% have other function (Figure 3).

3. STRUCTURAL BASIS OF GATES

Gates are dynamic systems that can make reversible transitions between open and closed states. They vary in size and complexity, from individual amino acid residues to loops, secondary structure elements, and even domains. The simplest gates consist of only one amino acid side chain that can close or open an access pathway by rotating. Opening and closing of more complicated systems can involve the synchronized movement of two or more residues, and the largest systems involve rearrangements of secondary elements or even entire domains (Table 1). For larger systems, movement of the gate may cause formation of a tunnel or enclosed cavity in addition to permitting or denying access to selected species.^{49,50} The following parameters can be useful for describing and discriminating between gates: (i) their constituent residues, (ii) their anchoring residues, (iii) the hinge region, i.e., the amino acids that make the structure flexible and allow it to move, (iv) the gate’s position, (v) the gate’s bottleneck diameter in the open and closed states, (vi) changes in the bottleneck’s size over time, (vii) the energy required to switch the gate from one state to the other, and (viii) the energy required for passage of specific molecules through the gate.

3.1. Residue Motion: Wings

The energetic barriers for residue rotation are quite small, 1–16 kcal/mol.⁵¹ While generally low, such barriers can nevertheless be large enough to significantly affect the probability that a given species will be able to pass through the gate or the rate at which they do so. Depending on the particular amino acid and its surroundings, one or both states of the gate may be stabilized by interactions with anchoring residues, e.g., hydrophobic interactions, H bonds, ionic interactions, salt bridges, and $\pi-\pi$ interactions. The strongest effect on the control of the passage is achieved when a large gating residue is located in the bottleneck of the pathway. The most common residues in this role are those whose side chains contain aromatic rings, i.e., W, F, and Y (Figure 4). Wing-type gates are common and can be found in enzymes such as imidazole

Table 1. Classifying Enzyme Gates According to Their Structural Basis

	1	2	3
Symbol	—●	●—●	●—●—●
Scheme of closed state			
Illustration of closed state			
Scheme of open state			
Illustration of open state			
Metaphor	wing	swinging door	aperture
Mechanism and moving part	side chain rotation	side chain rotation	backbone motion
Structural basis	1 residue	2 residues distant in sequence	2–4 residues distant in sequence
Amplitude of motion	< 2 Å	< 3 Å	< 3 Å
Time scale	ps – μs	ps – μs	ns – μs
Other features	anchoring residue	anchoring residues	hinge and anchoring free

Table 1. continued

	4	5	6
Symbol			
Scheme of closed state			
Illustration of closed state			
Scheme of open state			
Illustration of open state			
Metaphor	drawbridge	double drawbridge	shell
Mechanism and moving part	fragment movement	fragments movement	domain movement
Structural basis	~8-20 residues neighbour in sequence	2x ~8-20 residues neighbour in sequence	domain
Amplitude of motion	> 5 Å	> 5 Å	> 5 Å
Time scale	ns – μs	ns – μs	ms – s
Other features	hinge region, anchoring residues	hinge region, anchoring residues	hinge region, anchoring residues

glycerol phosphate synthase,³⁸ cytidine triphosphate synthetase,⁵² methane monooxygenase hydroxylase,⁵³ FabZ β-hydroxyacyl-acyl carrier protein dehydratase,⁵⁴ and cytochrome

P450.^{25,55} Even small gates of this type may require an activating agent to open. For example, the gate in the water channel of human monooxygenase CYP3A4 is created by the

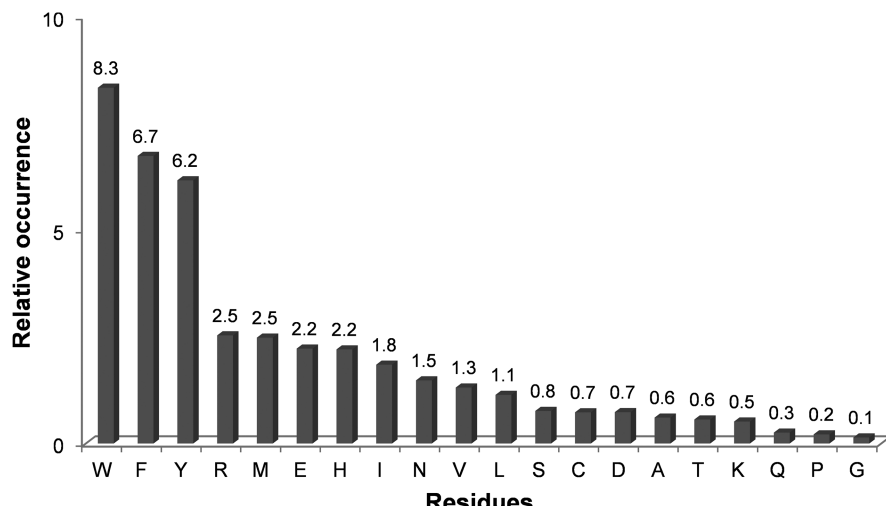


Figure 4. Relative occurrences of specific amino acid residues in wing and swinging door gates; 71 proteins with 129 gates were analyzed, and 154 residues that form wings or swinging doors were identified. Detailed description of the analyzed proteins is provided in Table 3. Values were normalized against the frequency with which each amino acid appears in all of the protein structures of the UniProtKB/Swiss-Prot database (2012_07).

interaction of the conserved residue R375 with the heme, which opens upon cytochrome P450 reductase binding to the enzyme.⁵⁶

3.2. Residue Motion: Swinging Doors

A more complex type of gate consists of two residues that can rotate but are stabilized in the closed conformation by a mutual interaction. Lario et al. introduced the phrase “swinging door” to describe gates of this type that were identified in cholesterol oxidase type I.⁵⁷ Some swinging door gates open by having both residues rotate in the same direction, while in others the two residues rotate in opposite directions. Common stabilizing interactions in swinging door gates include π stacking as occurs in the F–F pair of cytochrome P450_{3A4}^{58,59} and acetylcholinesterase,⁶⁰ ionic interactions as in toluene-4-monoxygenase⁶¹ and cytochrome P450_{cam}, P450_{BM3}, and P450_{eryF}^{25,55} aliphatic hydrophobic interactions such as those between the F–I, the F–V, and the F–L pairs of cytochrome P450_{3A4},^{58,59} aliphatic interactions such as those between the R–L and the L–I pairs of cytochrome P450_{3A4},^{58,59} and hydrogen bonds such as that between the R–S pair in cytochrome P450_{3A4}.⁵⁸ The open conformations of one or both of the gate residues may also be anchored, depending on the amino acids surrounding the gate. In comparison to wing gates, gates consisting of two residues can control wider tunnels and channels. It is worth mentioning that the individual residues that comprise a swinging door gate may simultaneously be components of another gate, as occurs in cytochrome P450_{3A4}.⁵⁸ Literature data indicate that most gates of this type consist of F–F pairs, and one way to screen for potential gates is to search for phenylalanine sandwiches.

3.3. Residue Motion: Apertures

Proteins undergo low-frequency breathing motions that may involve synchronized movements of bottleneck residues. In contrast to the previously described gates, the residues that form aperture type gates do not need to rotate and can maintain a rigid conformation. Their movements occur as a result of the synchronized relocalization of the enzyme backbone during its breathing motions. The ability of a given species to pass through gates of this type depends on the length of time the gate remains in the open state, which is determined

by the enzyme’s rigidity (especially in terms of the compartments housing the gating amino acids) and the strength of the interactions between the gating residues. Gates of this kind can therefore switch between states at different frequencies, which can be adjusted by mutating the gating residues. Aperture-type gates have been identified in several enzymes including carbamoyl phosphate synthetase,³⁶ choline oxidase,³¹ glutamate synthases,⁶² extradiol dioxygenases-homoprotocatechuate 2,3-dioxygenase,⁶³ cytochrome P450_{eryF},²⁵ and acetylcholinesterase.⁶⁴

3.4. Motions of Loops and Secondary Structure Elements: Drawbridges and Double Drawbridges

The movements of loops and secondary structure elements can provide an energetically favorable method of controlling access for larger ligands. The gates described above consist of individual residues and would not provide sufficient control for enzymes that have large substrates and correspondingly large active site cavities. In many cases, the loops involved in access control also contribute to formation of substrate/cofactor binding cavities. Alternatively, in enzymes with complex systems of internal tunnels such as the members of the cytochrome P450 family, the dynamic motion of the protein structure, especially the flexible B–C and F–G loops in the cytochromes P450, plays a vital role in the opening and closing of the tunnels.¹⁶ Protein motions of this type can also merge different tunnels, creating a wider opening. Here, gating elements control the access of large substrates by merging and dividing the space shared by the tunnels.¹⁶ However, in such cases the movements of the loops can cause formation of smaller and more selective gates such as the swinging doors described in the preceding sections.^{56,58}

Movements of loops and secondary structure elements can change the solvation of a cavity or the gate itself. The equilibrium between the open and the closed conformations depends on the anchoring residues and the flexibility of the hinge region. All of these elements play important roles in the movements of large gates. The conserved GxG motif found in most cytochrome P450 family members provides a good example.⁶⁵ Depending on cytochrome isoform, the motif flanks either one or both ends of the B–C loop. It increases the

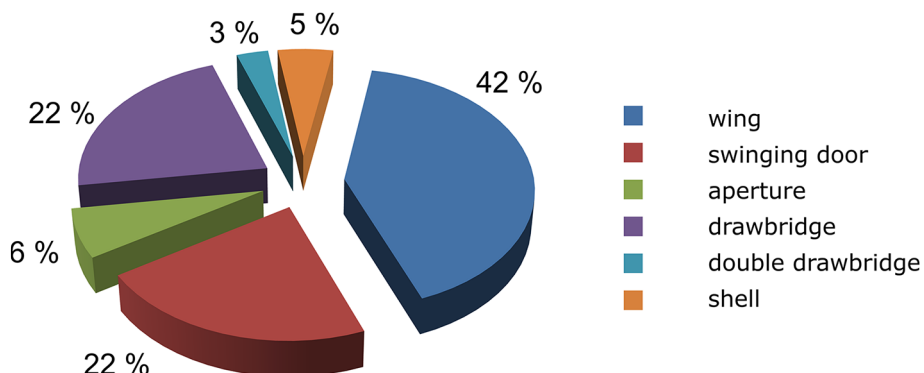


Figure 5. Frequencies of different gate types based on analysis of 71 proteins with 129 gates. Detailed description of the analyzed proteins is provided in Table 3.

flexibility of the loops, lowers the energy required for their motion, and facilitates tunnel opening and closing. The loops' variable lengths and levels of flexibility mean that each member of the P450 family has a gate with unique properties.¹⁶

3.5. Domain Motion: Shell

Large domain motions in enzymes are not generally regarded as gating systems. "Gates" of this scale are common in ion channels, which are beyond the scope of this review.¹⁴ However, one might expect that such large gates could be present in enzymes that catalyze reactions of very large substrates. Indeed, a gate of this kind has been observed in RNA polymerase, whose clamp domain opens to permit entry of promoter DNA during initiation, closes to establish a tight grip on the DNA during elongation, and then opens again to release the DNA during termination.⁶⁶ Interestingly, movement of large domains may protect enzymes from small molecule leakage and control their transport through long tunnel networks. This has been observed in carbon monoxide dehydrogenase/acetyl coenzyme A synthase, which operates as a tetrameric complex of distinct subunits.⁶⁷ The cap domain movement of epoxide hydrolase from *Mycobacterium tuberculosis* is another enzyme with a domain-scale gate that controls substrate access to the active site cavity.⁶⁸ Monomers of phospholipase A2 control access to their interface and the active site by adopting a different conformation during dimer aggregation.⁶⁹ Large domain movements often require an additional source of energy. For example, in the ATP-dependent protease HsIVU,⁷⁰ ATP hydrolysis is required to initiate conformational changes and propagate them to the residues that form the gate.

Structures of the gates found in 71 different proteins were analyzed systematically, yielding the results presented in Tables 2 and 3. The most common gate types are wings, swinging doors, and drawbridges, while apertures, double drawbridges, and shells are less common (Figure 5). However, these numbers may be distorted by the difficulty of identifying different gate classes by experimental methods or molecular dynamics simulations. Movements of only a few residues are more easily captured than those of secondary structure elements or domains due to the different time scales involved and the sizes of the moving structures. Moreover, gates may be controlled by overlapping processes that occur on different time scales and affect different structural regions.⁵⁰

4. LOCATIONS OF GATES

The roles of gates in the enzymatic catalysis discussed above suggest that these structures are natural hot spots for modifying enzyme properties. Identification of structural components of natural gates would therefore be very useful to protein designers. This raises a question: how and where should one look for the gates? Gates in proteins can be identified experimentally by protein crystallography and NMR spectroscopy and computationally by molecular dynamics simulations and normal-mode analysis.

71 芳基烷烃脱氯酶 LinB，⁷² L-氨基酸氧化酶，⁷³ 和对位甲苯酚单加氧酶。⁷⁴ 在某些情况下，仅一个构象将在解决的晶状结构中存在，这可能创建一个略显扭曲的图片，表明当开放构象被稳定时或当关闭构象被稳定时，不存在门控结构。^{31,61,75}

晶状分析的局限性可以通过先进的NMR光谱学克服，这使得同时研究多个蛋白构象成为可能，时间尺度从皮秒到毫秒不等。^{7,76} 这种分析提供了关于开放和关闭状态的信息，以及每个状态的分布和速率。NMR技术已被用于测量开放和关闭构象之间的交换率。^{77,78} HIV-1蛋白酶，⁷⁹ 和二氢叶酸还原酶。^{80,81} 总体而言，NMR对于研究远端效应突变对蛋白动力学的影响具有巨大潜力，从而有助于调查门控机制。

一些最有用的工具是为检测隧道、通道和蛋白结构中的腔室而开发的计算机程序。⁸² CAVER，⁸³ MOLE，⁸⁴ 和 MOLAXIS⁸⁵ 的输出可以分析以检测形成潜在门或识别最佳位置引入新门的瓶颈残基。这些“热点”突变可以赋予酶新的选择性。

Table 2. List of Enzymes Possessing Gates Described in the Scientific Literature with Indication of Their Function, Structural Basis, and Location

No.	Enzyme name	Substrate accessibility	Solvent accessibility	Function			Structural basis			Location			
				Synchronization of reactions	Other					Active site entrance and active site	Tunnel entrance and tunnel bottleneck	Cofactor cavity	Other
01	2-Amino-2-Desoxyisochorismate Synthase PhzE	+	+	+	+								
02	3-Hydroxybenzoate Hydroxylase MHBH		+	+	+	+!							+
03	4-Hydroxy-2-Ketovalerate Aldolase DmpG / Acylating Acetaldehyde Dehydrogenase DmpF	+	+	+	+	+							+
04	4-Hydroxybenzoate Hydroxylase PHBH		+			+!							+
05	Acetylcholinesterase AChE	+	+			+	+					+	
06	Acylaminoacyl Peptidase					+							+
07	α -Amylase TK1436	+				+	+						+
08	Asparagine Synthetase					+?							+
09	ATP-Dependent Proteases HslVU	+				+!!							+
10	Carbamoyl Phosphate Synthetase CPS – type II	+	+				+	+					+
11	Carbon Monoxide Dehydrogenase / Acetyl Coenzyme A Synthase	+		+	+								+
12	Carbonic Anhydrase β – type I						+						+
13	Carbonic Anhydrase β – type II	+	+	+		+		+				+	+
14	Carboxylesterase pnBCE					+	+						+
15	Catalase CAT-1 and CAT-3	+	+			+		+					+
16	Cellulobiohydrolase CEL 7A	+				+							+
17	Celllobiose Phosphorylase	+											+
18	Chalcone Synthase	+				+		+					+
19	Chloramphenicol Halogenase CmlS	+											+
20	Cholesterol Oxidase – type I SCHOX	+	+	+	+	+	+	+					+
21	Cholesterol Oxidase – type II BsChOx	+				+		+					+
22	Choline Oxidase	+							+	+?			+
23	Chondroitin AC Lyase	+				+					+		+
24	Copper-Containing Amine Oxidase	+				+							+
25	Cytidine Triphosphate Synthetase CTPS	+	+			+!		+			+	+	+
26	Cytochrome P450 CYP3A4	+	+			+!!	+				+	+	
27	Dihydrofolatereductase						+						+
28	Digeranylgeranylglycerophospholipid Reductase DGGR	+				+					+		+
29	Epoxide Hydrolase H37Rv	+											+
30	Epoxide Hydrolase M200	+				+?							+
31	FabZ β -Hydroxyacyl-Acyl Carrier Protein Dehydratase	+				+	+						+
32	Formiiminotransferase-Cyclodeaminase FTCD												+
33	Glucosamine 6-Phosphate Synthase GlmS	+	+	+	+	+	+	+			+		+
34	Glutamate Synthases GltS	+	+	+	+	+	+	+			+		+
35	Glutamine Phosphoribosylpyrophosphate Amidotransferase	+	+	+	+	+	+	+			+		+
36	Haloalkane Dehalogenase DhaA	+	+				+	+					+
37	Haloalkane Dehalogenase LimB	+											+
38	Histone Deacetylase HDAC1 and HDAC2	+					+	+					+
39	Histone Deacetylase HDAC8	+	+	+	+								+
40	HIV-1 Protease	+								+			+
41	Homoprotocatechuate 2,3-Dioxygenase	+					+	+					+
42	Hydrogenase FeFe	+											+
43	Hydrogenase NiFe								+!				
44	Imidazole Glycerol Phosphate Synthase IGPS	+	+			+		+					+
45	Inosine 5'-Monophosphate Dehydrogenase					+							
46	Ketoacyl Synthase KS	+	+				+						+
47	L-Amino Acid Oxidase	+				+	+						+
48	Lipase B	+				+	+						+
49	Lon Protease	+											+
50	Mannitol 2-Dehydrogenase					+	+						+
51	Methane Monooxygenase Hydroxylase MMOH	+	+	+	+	+	+			+	+		
52	Monoamine Oxidase A	+											+
53	Monoamine Oxidase B	+				+		+					+
54	Monooxygenase ActVAOrf6	+	+			+	+						+
55	NADH Oxidase					+!							
56	O-Acetylserine Sulphydrylase Cysteine Synthase	+				+							+
57	Oxidosqualene Cyclase SceOSC	+				+							+
58	Phenol Hydroxylase PHHY	+				+	+	+	+		+	+	+
59	Phospholipase A2	+											+
60	Phosphatidylinositol-Specific Phospholipase C	+											+
61	Quercetin 2,3-Dioxygenase	+				+	+	+					+

Table 2. continued

62	Rabbit 20a-Hydroxysteroid Dehydrogenase	+	+		+	+	+	+
63	Raucaffricine O- β -Dglucosidaseglucosidase	+		+			+	
64	RNA-Dependent RNA Polymerase		+		+			+
65	RNA Polymerase	+					+	+
66	Toluene-4-Monooxygenase T4MO	+	+	+	+		+	+
67	Toluene-Ortho-Xylene Monooxygenase ToMO	+			+		+	+
68	Triosephosphate Isomerase		+			+	+	
69	tRNA-Dependent Amidotransferase GatDE and GatCAB	+	+	+	+	+	+	+
70	Tryptophan Synthase	+	+	+	+	+	+	+
71	Undecaprenyl-Pyrophosphate Synthase		+			+	+	

+[!] Indicates gates interacting with cofactor. +^{!!} Cofactor assisted gating. +[?] Classification uncertain.



activities.⁸⁶ Zawaira et al.⁵⁹ used the CAVER software together with the Protein Interaction Calculator⁸⁷ for identifying gating residues within the cytochrome P450 family.

MD simulations are well suited for identification and analysis of gates and their behavior over time. Detailed descriptions of MD methods and their applications in simulating ligand migration can be found in recent reviews.^{9,88} Movements of large protein fragments on microsecond time scales can be investigated using Brownian dynamics,^{31,89} while Random Expulsion Molecular Dynamics and Steered Molecular Dynamics can be used to study pathways dedicated to transport of specific ligands.^{58,90} Some proteins have multiple pathways, each of which accommodates a different ligand or ligand class. This may in fact be a lot more common than is currently realized and can dramatically increase the complexity of gating systems arising from protein movement and the difficulty of identifying the true gating residues. For example, different residues control the ability of inhibitors E2020 and Huperzine A to access the active site of *Torpedo californica* acetylcholinesterase.⁶⁰ Similarly, in cytochrome P450_{3A4}, different residues in the same tunnel control access of temazepam and testosterone-6OH.⁵⁸ The importance of a gating residue identified by computational methods can be confirmed experimentally by site-directed mutagenesis and kinetic experiments.

Studies using the experimental and theoretical approaches for gate identification discussed above have demonstrated that their locations within the protein can vary widely. Gates have been observed (i) at the entry to the active site or even directly inside the active site, (ii) at the entry or in the bottleneck of the protein tunnel connecting the buried active site to the protein surface or connecting two active site cavities, and (iii) at the interface of the cofactor and active site cavities (Figure 6).

4.1. Active Site Entrance and Active Site

The entrance to the active site cavity is a suitable location for a gate, and gates situated here can have strong effects on enzyme activity. In some cases, the gating residues may even be a part of the active site.⁹¹ The simplest gates serve as filters that discriminate between potential substrates and thus play an important role in controlling enzyme selectivity. More advanced systems can prevent substrate entry when the active site residues are not properly oriented, e.g., in enzymes that require conformational changes before substrate binding. Many enzymes have gates at the entrance to their active sites, including acetylcholinesterase,⁶⁰ imidazole glycerol phosphate synthase,³⁸ glutamate synthase,⁴⁶ toluene-*o*-xylene monooxygenase,³⁸

nase,⁹¹ monooxygenase,⁹² choline oxidase,³¹ NiFe hydrogenases,²¹ carbonic anhydrases,⁹³ formiminotransferase-cyclo-deaminase,⁹⁴ type III polyketide synthases,⁹⁵ and FabZ β -hydroxyacyl-acyl carrier protein dehydratase.⁵⁴

4.2. Tunnel Entrance and Tunnel Bottleneck

The ability of ligands and solvent molecules to move from the media surrounding the protein to the active site can be controlled by gates located at any point along the tunnel. Gating residues may be situated at the tunnel entrance. However, it is more common to find them at the tunnel bottleneck. The tunnel entrance refers to the first shell of residues that define the tunnel and have contact with the bulk solvent. The tunnel bottleneck refers to the narrowest part that can be positioned anywhere along the tunnel (Figure 6). Even a single large residue whose side chain can project into the interior of the tunnel can exert efficient control over the access pathway. One might speculate that it might be favorable to have gates located inside tunnels because this allows their position to be more tightly controlled; their movements are restricted by the surrounding residues, and both the open and the closed conformations can be stabilized via interactions with neighboring amino acids. In contrast, residues located on the surface of the protein possess more degrees of freedom, and it is rare for both the open and the closed conformations to be stabilized. Examples of such gates inside the tunnels can be found in cholesterol oxidase type I,⁵⁷ toluene-4-monooxygenase,⁶¹ undecaprenyl-pyrophosphate synthase,²⁷ homoprotocatechuate 2,3-dioxygenase,⁶³ 4-hydroxy-2-ketovalerate aldolase/acylating acetaldehyde dehydrogenase,⁹⁶ epoxide hydrolase from *Aspergillus niger* M200,²⁶ and FabZ β -hydroxyacyl-acyl carrier protein dehydratase.⁵⁴ Similarly, gates can be situated in the bottlenecks of tunnels connecting two active sites. Gates in such positions are essential for enzymes that catalyze two reactions requiring different environments, such as glucosamine 6 phosphate synthase,⁴⁵ imidazole glycerol phosphate synthase,³⁸ cytidine triphosphate synthetase,⁵² carbamoyl phosphate synthetase,³⁶ and glutamate synthases.⁴⁶

4.3. Cofactor Cavity

Gates can be positioned at the interface of the active site and the cofactor cavity, allowing for more fine-grained control during the reaction. In NADH oxidase, the W47 residue acts as a gate that controls the accessibility of the FAD flavin ring and thus plays a crucial role during the catalytic cycle. The closed conformation is stabilized by hydrogen bonds between the cofactor and the peptide backbone, whereas stabilization of the open form may be advantageous during the initial steps of

Table 3. Detailed Description of Enzymes Possessing Gates Presented in the Scientific Literature

No.	Enzyme name	EC number
Gate function <i>Information about gate function</i>		
Gate location <i>Information about gate location</i>		
Gate structural basis <i>Schematic drawing of gate class</i>	GATE 1- font colour corresponds to the colour of residues on the picture Information about residues, open and closed conformation and mechanism of the changes of gate states ? GATE1 - ? indicates a hypothetical gate, some data missing to fulfil all requirements, e.g., lack of information about open and close conformation * GATE1 - * indicates important residues used for gates engineering	Enzyme function <i>Information about catalysed reaction</i> Small picture of whole enzyme with detected tunnels and gates protein – light blue surface tunnel – dark blue wire gates residues – red ball and stick Large picture – close-up on gate residues protein – light blue cartoon tunnel profile – grey spheres active sites residues (if shown) – green ball and stick gates residues (elements) – ball and stick representation colour correspond to colour of gate name cofactor – yellow ball and stick substrate – yellow ball and stick
Gate engineering <i>Information about mutants that change gate state or create new gates, including information about mutants closing and opening tunnels</i>		All pictures present results of CAVER 3.0 calculations of tunnels visualized by PyMOL. Tunnels corresponding to those reported in literature have been visualised. Reader should be aware that calculations were performed using available static structures from the PDB database, therefore calculated tunnels can differ from the tunnels described on the base of results from MD simulations.
PDB: Wild-type (WT) or mutants available in Protein Data Bank database	Tunnel	Name or part of the tunnel
References: All references used for table preparation	Length	Tunnel length in Å
	Bottleneck	Bottleneck diameter in Å; data taken from the literature
	Role	Information about transported molecules

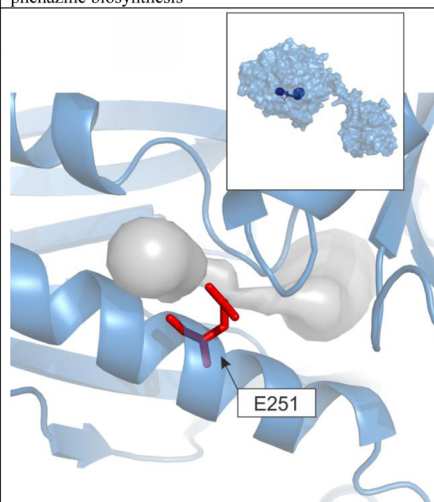
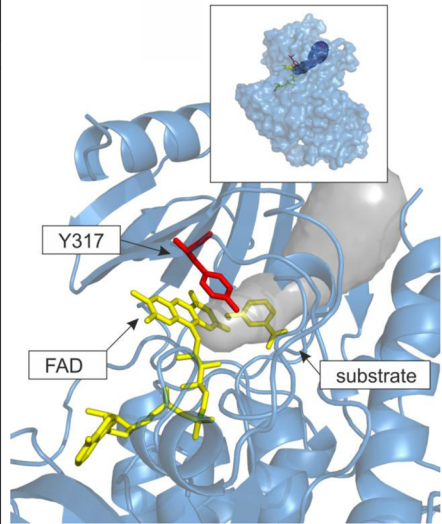
01	2-Amino-2-Desoxyisochorismate Synthase PhzE	4.1.3.27
Gate function GATE1 – Controls access of ammonia, synchronizes active sites		Enzyme function Utilizes chorismate and glutamine to synthesize 2-amino-2-desoxyisochorismate in the first step of phenazine biosynthesis
Gate location GATE1 – Between the MST (menaquinone, siderophore, tryptophan) domain and GATase1 active site		
Gate structural basis —● GATE1 E251 forms two hydrogen bonds with N149 in closed conformation interacts with K254 in open conformation		
Gate engineering		PDB ID: 3R74
	Tunnel U-shaped	
	Length	25
PDB: WT – 3R74 (open), 3R75 (closed)	Bottleneck	
References: ¹¹¹	Role	Ammonia transport

Table 3. continued

02	3-Hydroxybenzoate Hydroxylase MHBH		1.14.13.23
Gate function		Enzyme function	
GATE 1 – Controls the contact of NADPH with the isoalloxazine ring, protects FAD from the solvent		Conversion of 3-hydroxybenzoate to 3,4-dihydroxybenzoate	
Gate location			
GATE 1 – Between substrate tunnel and the FAD binding pocket			
Gate structural basis			
—●	GATE 1 Y317 The residue creates parallel π - π stacking interaction with FAD Opening – reorientation of both FAD and Y317		
Gate engineering			
			
		PDB ID: 2DKH	
		Tunnel	E1
		Length	20-22
		Bottleneck	4.5
		Role	O ₂ , substrate
		Product	
PDB: WT – 2DKH, 2DKI			
References: ⁹⁸			

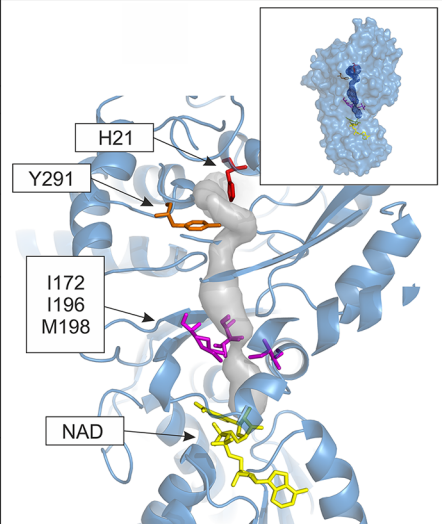
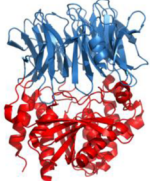
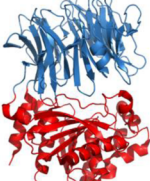
03	4-Hydroxy-2-Ketovalerate Aldolase DmpG / Acylating Acetaldehyde Dehydrogenase DmpF	4.1.3.39 / 1.2.1.10	
Gate function		Enzyme function	
GATE1 – Controls access of the substrate to the DmpG active site		Catalyses final two steps in degradation of toxic aromatic intermediates in the meta-cleavage pathway of catechol	
GATE2 – Proton transfer, synchronizing two active sites by controlling the passage of the acetaldehyde			
GATE3 – Controls intermediate entry to the DmpF active site; facilitates interaction between N171 and NAD			
Gate location			
GATE1 – Tunnel entrance in aldolase subunit			
GATE2 – Tunnel entrance in aldolase subunit			
GATE3 – Tunnel exit in dehydrogenase subunit			
Gate structural basis			
—●	GATE1 H21 Opening – reorientation of H21		
—●	GATE2 Y291 Opening – reorientation of Y291		
—●	GATE3 I172 + I196 and M198 I172, I196 and M198 block the tunnel exit in the dehydrogenase subunit Opening – in the structure with bound NAD ⁺ – an interaction of N171 with NAD induce unique orientation of I172, additionally I196 and M198 adopt open conformations		
Gate engineering			
H21A – acetaldehyde and propionaldehyde channelling reduced by more than 70%			
Y291F – reduced channelling efficiencies by >30%			
I196L, I196F – no significant changes in the channelling efficiency			
			
		PDB ID: 1NVM	
		Tunnel between active sites	
		Length	29
		Bottleneck	1.7
		Role	Acetaldehyde intermediate transport
PDB: WT – 1NVM			
References: ^{96,112}			

Table 3. continued

04	4-Hydroxybenzoate Hydroxylase PHBH		1.14.13.2
Gate function		Enzyme function	
GATE1 – Controls access of the solvent		Monoxygenation of <i>p</i> -hydroxybenzoate (<i>p</i> -OHB) to 3,4-dihydroxybenzoate	
GATE2 – Controls access of the solvent			
Gate location			
GATE1 – Close to the FAD cofactor			
GATE2 – One domain of the protein			
Gate structural basis			
—●	GATE1 R220 and FAD R220 modulates the dynamics of flavin movements Opening – reorientation of R220; out – solvent exposed; in – solvent excluded		
○	GATE2 Large domain movement As substrate (<i>p</i> -OHB) moves forward in the tunnel and reaches its high-affinity site, the $\beta\alpha\beta$ (1-180) and the sheet domains (180-270) are expected to rotate and close the active site onto the substrate		
Gate engineering			
R220Q – keeps enzyme in the open conformation – loss of selectivity and decrease of activity (100-fold)			
PDB: WT – 1IUW, Mutant R220Q – 1K0I, 1K0J, 1K0L		PDB ID: 1IUW	
References: ¹¹³		Tunnel	
		Length	
		Bottleneck	
		Role	Substrate access

05	Acetylcholinesterase AChE		3.1.1.7
Gate function		Enzyme function	
GATE1 – Controls access of the substrate to the active site		Hydrolysis of acetylcholine	
GATE2 – Controls escape of the acetic acid and/or the water molecule			
Gate location			
GATE1 – Main tunnel – 12 Å from the bottom of the gorge – entrance to the active site			
GATE2 – Back door tunnel – on the C67–C94 wall			
Gate structural basis			
—●	GATE1 F330 F330 controls the entrance of the natural substrate Second most important residue – Y121, followed by W84, F288, F290, F331, Y334 <i>Cationic substrate (Huperzine A):</i> – movement of F330, Y121 and D72 generate an electrostatic field affecting the substrates <i>Aromatic substrate (E2020):</i> residues grouped in 3 groups acting as “sender” and “receiver”, compose a “conveyer belt” via π – π stacking interactions with benzene ring of E2020: Group I contains W84, F330, and F331 Group II consists of F288, F290, and Y334 Group III includes Y70, Y121, and W279 <i>In mAChE</i> F338 -Y124		
—● —●	GATE2 W84, G441 and Y442 (TcAChE); W86, G448 and Y449 (mAChE) Opening – movement of the W84 indole ring, almost 90° rotation to a position where it interacts with Y442 <i>Alternative propositions</i> ?GATE2b – E82, P76, and G77 (with some small movement of D72, E73, and N85) ?GATE2c – Between V71, N85, P86, and M90 ?GATE2d – 90° rotation of the F78 and the displacement of V431 and W432 ?GATE2e – W84, V129, and G441 ?GATE2f – Q-loop (C67 – C94) with W84 could undergo a flap-like transition ?GATE2g – Facial rearrangement of the loop between W279 and S291		
Gate engineering			
V129W – 4-fold increase of K_m		Tunnel	main
V431C – 2-fold increase of K_m		Length	20
PDB: WT – 1W75, 1ACJ, 1ACL, 1EA5, 2ACE, 1MAH, 1QTI, 1DX6, 1EVE, 1OCE, 2XI4 (open back door)		Bottleneck	1.2-2.4
References: ^{13,35,60,64,114–123}		Role	Acetylcholine
			Small molecules

Table 3. continued

06	Acylaminoacyl Peptidase		3.4.19.1
Gate function GATE1 – Controls enzyme activity – only closed form is active		Enzyme function Removes acylated amino acid residues from the N terminus of oligopeptides	
Gate location GATE1 – Two domains of an enzyme monomer			
Gate structural basis			
	GATE1 Two domains may move away to form an opening of about 30°, with D376 being the hinge Open - accept substrate Closed - rearrange active site	 closed PDB ID: 3O4G	
Gate engineering		 open	
PDB: WT – 3O4G			Bottleneck
References: ^{124,125}			Role

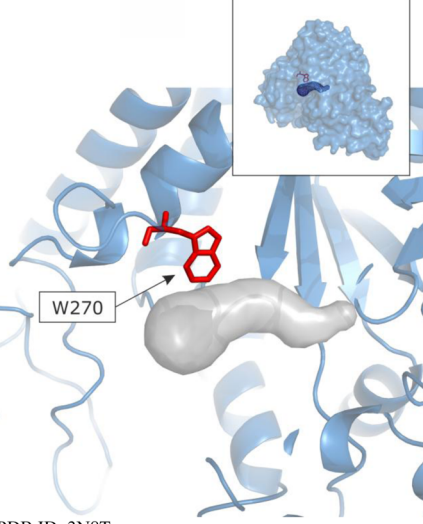
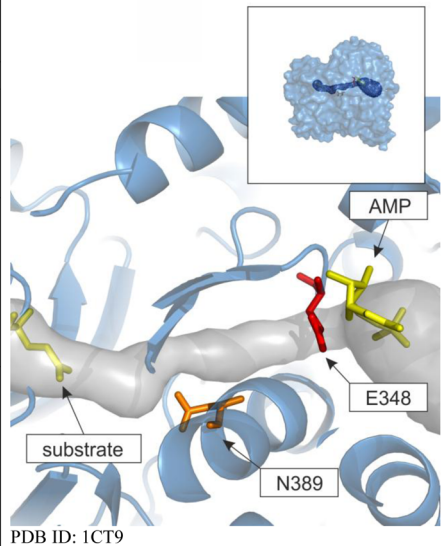
07	α -Amylase TK1436		2.4.1.18
Gate function GATE1 – Controls access of the substrate to the active site, regulates transglycosylation		Enzyme function Formation of branch points in glycogen and amylopectin by cleavage of α -1,4 glycosidic bonds and subsequent transfer to a new α -1,6 position	
Gate location GATE1 – At the entrance of the active site			
Gate structural basis			
	GATE1 W270 The residue displays different conformations depending on the presence or absence of ligands in the active-site pocket Other possible gate-keepers W28, W407, W416	 PDB ID: 3N8T	
Gate engineering			
PDB: WT – 3N8T, 3N92, 3N98			Bottleneck
References: ¹²⁶			Role Substrate recognition and binding

Table 3. continued

08	Asparagine Synthetase		6.3.5.4
Gate function ? GATE1 – Synchronizing active sites, establish the intramolecular tunnel for ammonia passage ? GATE2		Enzyme function ATP dependent synthesis of L-asparagine from L-aspartic acid	
Gate location ? GATE1 – C-terminal end of the ammonia tunnel linking the active site near to the ATP moiety ? GATE2 – C-terminal domain			
Gate structural basis			
—●	? GATE1 E348		
—●	? GATE2 N389		
Gate engineering E348D – impairs acyl-adenylate formation, tunnel is more solvent exposed leading to loss of the ammonia because of an impaired rate of β AspAMP formation			
PDB: WT – 1CT9		PDB ID: 1CT9	
References: ^{42,44,127,128}		Tunnel	
		Length	Bottleneck
			Role
			Ammonia transport

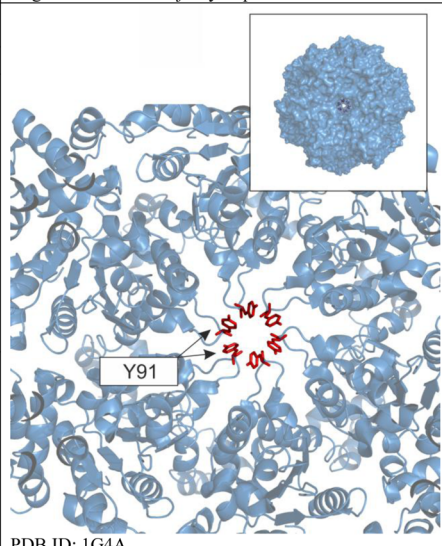
09	ATP-Dependent Protease HslVU		3.4.25.2
Gate function GATE1 – Controls access of the substrate to the active site located inside the chamber		Enzyme function Degradation of the majority of proteins in a cell	
Gate location GATE1 – Translocation tunnel			
Gate structural basis			
—●	GATE1 Y91 “Twist-and-open” mechanism - conformational changes induced by ATP hydrolysis are propagated to the gating sequence Y91 can move (180° rotation) from inside HslU toward HslV through the pore – closed pore has diameter 4.4 Å open pore has diameter of 19.3 Å HsIVU works as a hexamer therefore pore diameter depends on the number of Y91 pointed toward HslV		
Gate engineering Y91F, V92I, V92A, and V92S – decrease in protein degradation activity G90P, G93P, G90A, G93P, Y91I, Y91C, Y91S, Y91A, V92F, V92C – are not capable to support the proteolytic activity of HslV			
PDB: WT – 1G4A, 1G4B		PDB ID: 1G4A	
References: ^{13,129,130}		Tunnel	-
		Length	Bottleneck
			Closed - 4.4; open up to 19.3
			Role

Table 3. continued

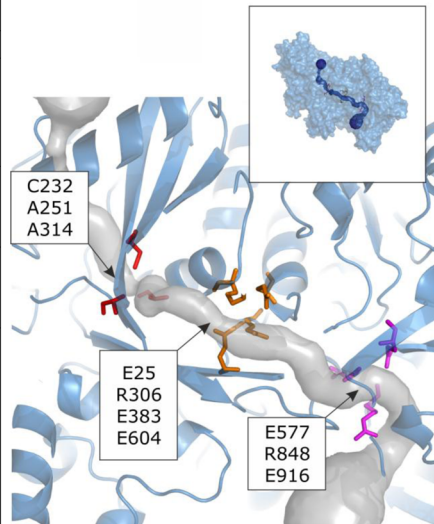
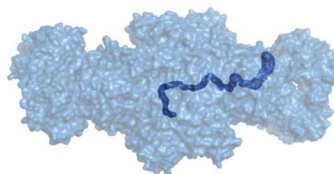
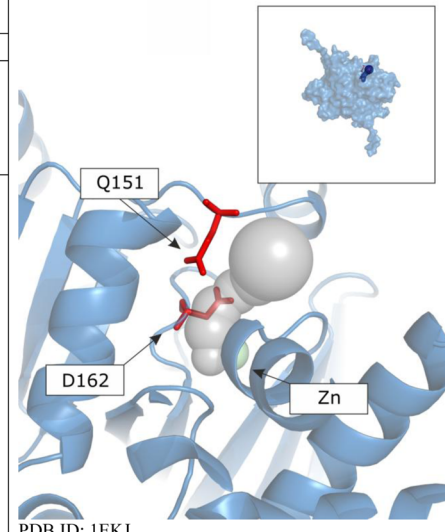
10	Carbamoyl Phosphate Synthetase CPS – type II			6.3.5.5
Gate function				Enzyme function Synthesis of carbamoyl phosphate
GATE1 – Desolvation of ammonia GATE2 – Controls entry of carbamate into the tunnel prior to phosphorylation to carbamoyl phosphate GATE3 – Controls entry of carbamate into the tunnel prior to phosphorylation to carbamoyl phosphate				
Gate location GATE1 – Large subunit/ammonia tunnel GATE2 – Carbamate tunnel near the carboxy phosphate binding site GATE3 – Carbamate tunnel near the site for the synthesis of carbamoyl phosphate				
Gate structural basis				
 GATE1 C232, A251, A314 (barrier 7.2 kcal/mol) Two more triad that may act as gate/switches for an ammonia passage are: I234-T249-L310 E217-T244-S307				PDB ID: 1JDB
 GATE2 R306, E25, ?E383, ?E604 R306 – ion pair with E25				
 GATE3 R848, E577 and ?E916 R848 – ion pair with E577				
Gate engineering C232V/A251V/A314V – closing ammonia tunnel – unable to synthesize carbamoyl phosphate using glutamine as a nitrogen source G359F, G359Y – decoupling separate chemical reactions via creation an escape route for the ammonia intermediate α P360A/ α H361A/ β R265A – unable to utilize glutamine for the synthesis of carbamoyl phosphate via creation of an escape route for the ammonia intermediate; full catalytic activity with external ammonia source E916Q – 10-fold decrease in the rate of carbamoyl phosphate synthesis E25Q/E383Q – carbamoyl phosphate synthetase activity was diminished 50-fold E25Q/E383Q/E604Q – glutaminase activity is decreased about 5-fold, and the bicarbonate-dependent ATPase activity is diminished at least 20-fold E577Q – decrease of carbamoyl phosphate synthesis (100-fold)				
PDB: WT – 1BXR, 1JDB	Tunnel	Whole	I part	II part
References: 36,41,103,131–134	Length	>100	45	
	Bottleneck		3.2	
	Role	Connects all 3 active sites	Ammonia transport	Carbamate transport
11	Carbon Monoxide Dehydrogenase / Acetyl Coenzyme A Synthase			1.2.7.4 / 1.2.99.2 / 2.3.1.169
Gate function				Enzyme function $\alpha\beta\beta$ tetramer; β subunit catalyses the reduction of CO ₂ to CO (C cluster), α subunit is responsible for the synthesis of acetyl-CoA from CO, coenzyme A (CoA) and a methyl group donated by a cobalamin-containing protein (A cluster)
GATE1 – Controls the reaction, protects from CO leakage and controls CO access to Ni _p -Ni _d -[Fe ₄ -S ₄] cluster (A cluster – acetyl-CoA synthase active site)				
Gate location GATE1 – 20 Å from the A cluster – on the A and C cluster border				
Gate structural basis				
 GATE1 Large conformation change of subunit α , gating residues – residues of the α subunit N-terminal domain 143-148 Open conformation – accessible active site – closed tunnel – F512 moves to a position within 4 Å of both Ni _p and Ni _d blocking putative ligand binding to the axial coordination site of Ni _p Close conformation – closed active site – open tunnel – F229 blocks axial ligand binding to Ni _d but not to the Zn ion				PDB ID: 1OAO Only one from two symmetrical tunnels is shown on a picture.
Gate engineering A110C, A222L, A265M – block the tunnel between the A and C-clusters; A222L complete blocking A578C, L215F, A219F – block the tunnel between the C clusters F70W, N101Q – block a region at the $\beta\beta$ subunit interface that might dynamically connect the tunnel with a newly discovered water tunnel	Tunnel			
	Length	130 between A clusters, additional two tunnels connecting C clusters 37 + additional water channel		
PDB: WT (<i>M. thermoacetica</i>) – 1OAO (open+closed form), WT (<i>C. hydrogenoformans</i>) – 1RU3 (closed form)	Bottleneck			
References: 48,67,135–137	Role	CO transport between two active sites		

Table 3. continued

12	Carbonic Anhydrase β – type I		4.2.1.1
Gate function ?GATE1 – Protects zinc cation site		Enzyme function Catalyses a reversible reaction to form bicarbonate from carbon dioxide and water	
Gate location ?GATE1 – Vicinity of the active site			
Gate structural basis			
● — ● ?	?GATE1 D162 donates a hydrogen bond to Q151 Acts as a gatekeeper residue by excluding anions from the zinc ligand environment that cannot donate a hydrogen bond at this position		
Gate engineering			
PDB: WT – 1EKJ		PDB ID: 1EKJ	
References: ⁹³		Tunnel	
		Length	
		Bottleneck	
		Role	Acetate ion, CO_2 transport

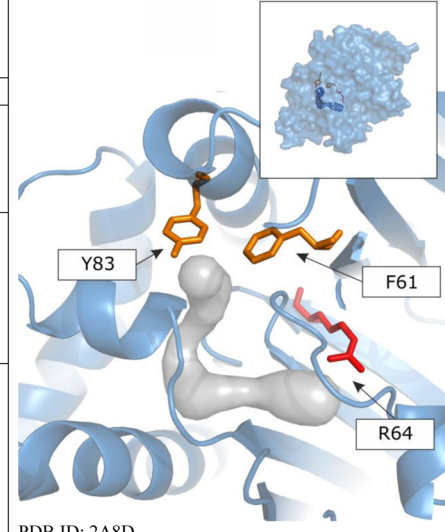
13	Carbonic Anhydrase β – type II		4.2.1.1
Gate function GATE1 – Controls access of the bicarbonate ion GATE2 – Controls access of the solvent, protects active site, controls transport of bicarbonate ion from the active site or from solution to the allosteric pocket		Enzyme function Catalyses a reversible reaction to form bicarbonate from carbon dioxide and water	
Gate location GATE1 – Middle of the tunnel GATE2 – Vicinity of the active site in a narrow hydrophobic active site cleft that lies along the dimer or pseudo-dimer interface			
Gate structural basis			
—●	GATE1 R64 – HICA Rotation of the guanidinium group allows migration of bicarbonate ion into the allosteric site		
● — ● ?	GATE2 F61, V62?, Y83, and V87 Other anhydrases PPCA – F422, I173, Y444, Y448 Rv3588 – F70, A75, Y89, V93 Nce103 – F97, L102, G111, D59		
Gate engineering			
PDB: WT (HICA) – 2A8D, WT (PPCA) – 1DDZ, WT (Rv3588) – 1YM3, WT (Nce103) – 3EYX		PDB ID: 2A8D	
References: ^{93,138}		Tunnel	
		Length	
		Bottleneck	
		Role	Acetate ion, CO_2 transport

Table 3. continued

14	Carboxylesterase pnbCE		3.1.1
Gate function		Enzyme function	
GATE1 – Controls exit of hydrolysis products by a side door GATE2 – Controls access to the active site gorge GATE3 – Controls access to the active site gorge		Detoxification of xenobiotics	
Gate location			
GATE1 – Between active site and side door GATE2 – Entrance to active site gorge GATE3 – Entrance to active site gorge			
Gate structural basis			
—●	GATE1 L362 Rotates 180° around its C-C bond and adopts two distinct conformations		
~~~~~	<b>GATE2</b> Loop coil_5 (residues 61-82) and coil_21 (residues 408-422) Move away from their equilibration conformation, the active site opens and can accommodate incoming substrate		
~~~~~	<b>GATE3</b> Loop coil_17(residues 308-323) and coil_21 Close for substrate hydrolysis, move away to release products		
Gate engineering			
Δ coil_5 – k_{cat} 4.5-fold smaller Δ coil_21 – k_{cat} 3-fold smaller			
PDB: WT – 1QE3		PDB ID: 1QE3	
References: ^{139,140}		Tunnel	
		Length	
		Bottleneck	
		Role	

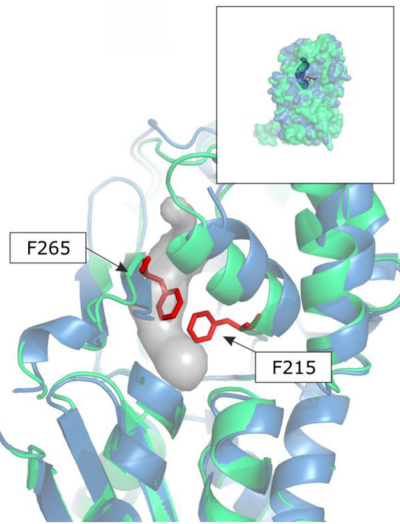
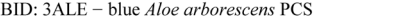
15	Catalase CAT-1 and CAT-3		1.11.1.6
Gate function		Enzyme function	
GATE1 – Blocks access to the heme, controls solvation ?GATE2 – ?		Decomposition of hydrogen peroxide to water and O ₂	
Gate location			
GATE1 – Entrance to the active site cavity from main tunnel ?GATE2 – Side tunnel			
Gate structural basis			
?	GATE1 H54, V95, and F132 Barrier to reach the active site: H ₂ O ₂ < 4.8 kcal/mol O ₂ ~2.2 kcal/mol H ₂ O ~ 3.6 kcal/mol Opening of the gate may be regulated according to the H ₂ O ₂ concentration in the small cavity before the gate; H ₂ O ₂ in the cavity would change the net of hydrogen bonds and trigger opening of the gate; water molecules interacting with amino acid residues in the cavity would determine closure of the gate		
—●	?GATE2 (not shown on picture) CAT-1 – hydroxyl group of S198 blocks tunnel CAT-3 – hydroxyl group of T208 blocks tunnel		
Gate engineering			
PDB: WT – 1NM0		Tunnel	Main
References: ^{23,24,88}		Length	31
		Bottleneck	
		Role	O ₂ and H ₂ O ₂ transport

Table 3. continued

16	Cellulobiohydrolase CEL7A		3.2.1.91
Gate function	GATE1 – Facilitates processing of crystalline cellulose degradation		
Gate location	GATE1 – Exo loop		
Gate structural basis	 GATE1 Exo loop + Y247 Y247 in closed conformations interacts with Y 371 from the short loop – acting like a button – it creates tunnel, loop covers the active site during reaction time		
Gate engineering	Y247F – removes hydrogen bond between Y247 and substrate - small effect on cellulose hydrolysis D241C/D249C – reduces mobility of the loop, disulphide bridge enhanced the activity on both amorphous and crystalline cellulose Deletion G245-Y252 – increases activity on amorphous cellulose, and half of the activity on crystalline cellulose		
PDB: <i>WT</i> – 1CEL			PDB ID: 1CEL
References: ^{49,141–143}			Tunnel
			Length 50
			Bottleneck 3.5
			Role Substrate cavity

17	Cellobiose Phosphorylase		2.4.1.20
Gate function	GATE1 – Controls access of the substrate		
Gate location	GATE1 – Flexible loop		
Gate structural basis	 GATE1 Flexible loop (495–513) Loop undergoes conformational changes during substrate binding and release		
Gate engineering	T508I – changes substrate specificity N156D and N163D – increase of the activity N156D/N163D/T508I/E649G/N667A – improves activity towards a whole range of β -glucosidic acceptors		
PDB: <i>WT</i> – 2CQT, 1V7X			PDB ID: 2CQT
References: ^{144,145}			Tunnel
			Length
			Bottleneck
			Role

Table 3. continued

18	Chalcone Synthase CHS		2.3.1.74
Gate function GATE1 – Controls orientation of the substrate		Enzyme function Decarboxylative condensations of malonyl-CoA with a CoA-linked starter	
Gate location GATE1 – Between the active-site cavity and the CoA binding tunnel			
Gate structural basis		 <p>GATE1 <i>Medicago sativa</i> CHS F215 and F265 Block the lower portion of the opening between cavities and help with folding and the internal orientation of the tetraketide intermediate during the cyclization reaction; F215 facilitate the decarboxylation of malonyl-CoA by maintaining the orientations of substrates and intermediates during the sequential condensation reactions</p>	
Gate engineering		<p><i>M. sativa</i> CHS F215S – changes the substrate specificity via opening a space at the cavity entrance</p> <p><i>A. arborescens</i> PCS *GATE2 M207G – opens a gate to two novel hidden pockets behind the active site of the enzyme - residue 207 controls the number of condensations of malonyl-CoA F80A/Y82A/M207G – provide further elongation of products</p>	
PDB: WT (<i>M. sativa</i> CHS) – 1CGK, WT (<i>A. arborescens</i> PCS) – 3ALE		PDBID: 3ALE – blue <i>Aloe arborescens</i> PCS PDBID: 1CGK – light green <i>Medicago sativa</i> CHS tunnel calculated for 3ALE structure	
References: ^{95,106,146}		Tunnel Length 16 Bottleneck Role Reaction cavity	

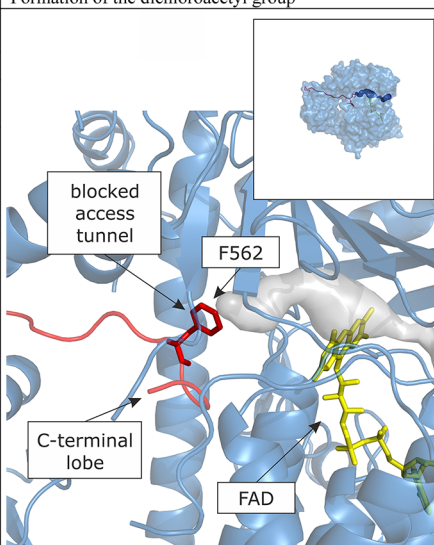
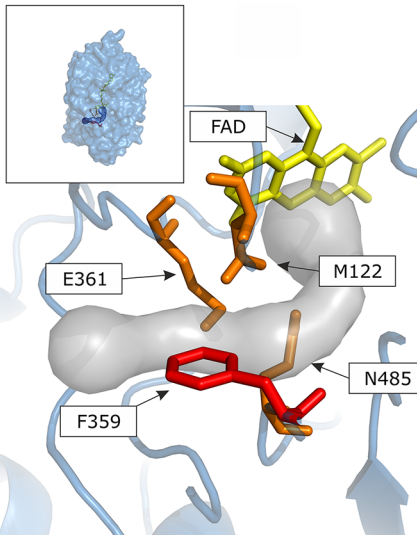
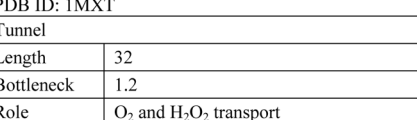
19	Chloramphenicol Halogenase CmlS		Not determined
Gate function GATE1 – Blocks access to the active site		Enzyme function Formation of the dichloroacetyl group	
Gate location GATE1 – Flexible loop (the C-terminal lobe of CmlS)			
Gate structural basis		 <p>Flexible loop (the C-terminal lobe of CmlS) + F562 acting as a plug</p>	
Gate engineering			
PDB: WT – 3I3L		PDB ID: 3I3L Tunnel Length 10 Bottleneck Role Connecting halogenation active site and FAD binding site	
References: ¹⁴⁷			

Table 3. continued

20	Cholesterol Oxidase – type 1 SCHOX		1.1.3.6
Gate function GATE1 – Controls access of the solvent GATE2 – Controls access of O ₂ to the flavin and assures that isomerization occurs before the oxidative half of the reaction, tuning the redox state of the cofactor		Enzyme function Catalyses the first step in the pathway of cholesterol degradation	
Gate location GATE1 – Frames the entrance to the tunnel GATE2 – Gate is in the tunnel leading to the isoalloxazine ring of flavin		 <p>PDB ID: 1MXT Tunnel Length 32 Bottleneck 1.2 Role O₂ and H₂O₂ transport</p>	
Gate structural basis			
<p>● GATE1 F359 F359 adopts two distinct alternate conformations separated by a 65° rotation of the benzene group Closed gate - maximizing hydrophobic packing interactions with V189, V124 and G347 Open conformation - the tunnel becomes solvent accessible Control of F359 gate appears to be dependent on the adopted conformation of N485</p> <p>● ● GATE2 N485, E361 and M122 Switch between their side-chain conformations, controls the access of O₂ to flavin 1. Binding of steroid → rotation of the methyl group of M122 → pushes N485 sealing the tunnel and creating an ideal environment for oxidation 2. After substrate oxidation, a strong hydrogen bond forms when N485 moves to conformation near the flavin, forcing M122 to a conformation that destabilizes the binding of the oxidized product. 3. When the substrate is oxidized and the FAD cofactor is reduced, the side chain of N485 rotates toward the cofactor → tunnel opening - regulates access of oxygen to the active site</p>		<p>Gate engineering F359W – rate of catalysed reaction decreases 13-fold G347N – could not be saturated with oxygen N485D – could not be saturated with oxygen</p> <p>PDB: WT – 1MXT, Mutant F359W – 3CNJ, Mutant N485D – 3GYI References: ^{57,99,148}</p>	

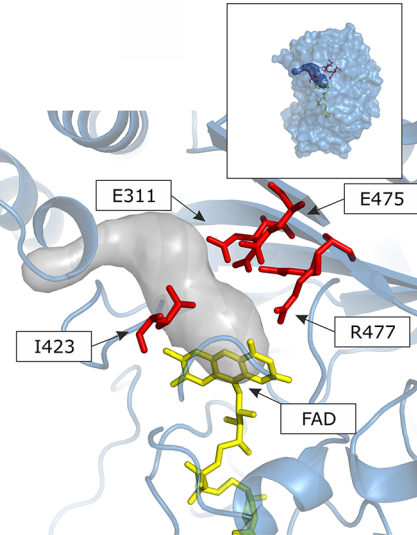
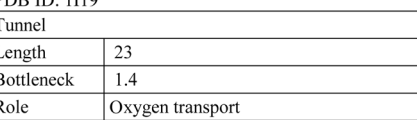
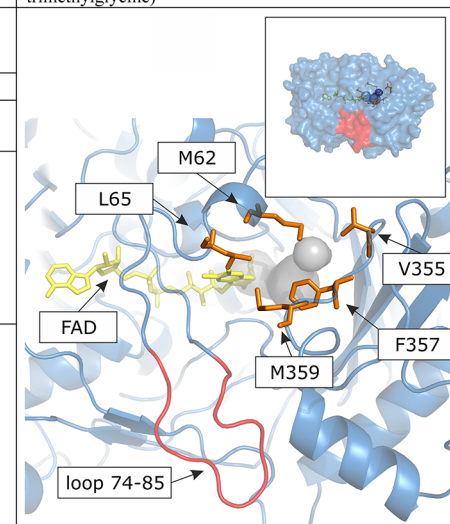
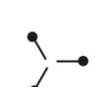
21	Cholesterol Oxidase – type II BsChOx		1.1.3.6
Gate function GATE1 – Modulates access/reactivity of dioxygen		Enzyme function Catalyses the first step in the pathway of cholesterol degradation	
Gate location GATE1 – Bottleneck of the tunnel		 <p>PDB ID: 1I19 Tunnel Length 23 Bottleneck 1.4 Role Oxygen transport</p>	
Gate structural basis			
<p>● ● GATE1 E311, I423, E475 and R477 Open conformation of R477 is stabilized by a salt bridge with E311 E311 tunes the E475—R477 pair</p> <p>● GATE2 E311D/Q/L – cause a switch in the basic kinetic mechanism of the reoxidation with dioxygen, while BsChOx wild type and most mutants show saturation behaviour with increasing O₂ For E311 – a linear dependence was found that would reflect a second-order process R477A – limits both oxidation and isomerization activities A204C, G309A, G309C, I423L, I423V, E475D, E475Q, R477K – no significant changes</p>		<p>Gate engineering E311D/Q/L – cause a switch in the basic kinetic mechanism of the reoxidation with dioxygen, while BsChOx wild type and most mutants show saturation behaviour with increasing O₂ For E311 – a linear dependence was found that would reflect a second-order process R477A – limits both oxidation and isomerization activities A204C, G309A, G309C, I423L, I423V, E475D, E475Q, R477K – no significant changes</p> <p>PDB: WT – 1I19 References: ^{99,149,150}</p>	

Table 3. continued

22	Choline Oxidase		1.1.3.17
Gate function ?GATE1 – Controls access of the positively charged substrate GATE2 – Controls access of the positively charged substrate		Enzyme function Oxidation of choline to glycine betaine (N,N,N-trimethylglycine)	
Gate location ?GATE1 – Loop adjacent to the active site of choline oxidase - residues 74-85 GATE2 – Located in the tunnel above active site			
Gate structural basis			
	? GATE1 Movement of the loop can open active site		
	GATE2 M62, L65, V355, F357, and M359 Breathing motion of M62, L65, V355, F357, and M359 The weakly hydrophobic interactions between the gating residues ensure that the positively charged substrate can easily slip to the highly electronegative active site The distribution of the residues just outside the five gating residues contains a considerable amount of negatively charged amino acids, which include E63, E66, E358, and E370, which may attract and guide the positively charged choline substrate to the active site		
Gate engineering			
PDB: WT – 2JBV		PDB ID: 2JBV	
References: ³¹		Tunnel	
		Length	
		Bottleneck	2.5
		Role	Substrate transport

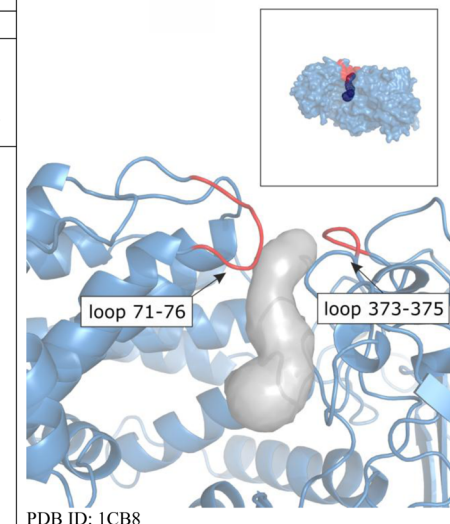
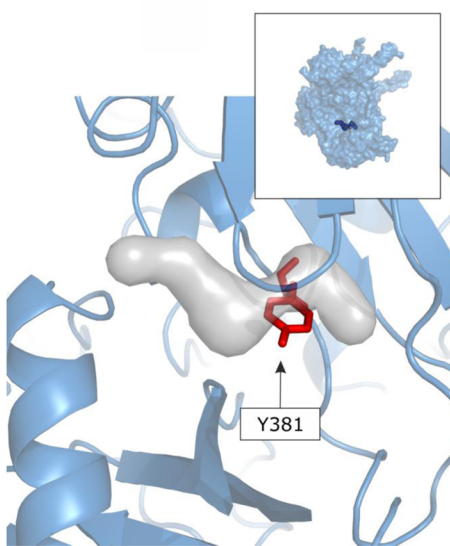
23	Chondroitin AC Lyase		4.2.2.5
Gate function GATE1 – Creates tunnel and the active site, controls access of the solvent		Enzyme function Degradation of glycosaminoglycans	
Gate location GATE1 – N-terminal domain and C-terminal domain			
Gate structural basis			
	GATE1 Two loops Loops are flexible and open periodically allowing the glycosaminoglycan chain to slide in; Gating involves the movement of the tips of one or two loops only, D71-W76 of the N-terminal domain and G373-K375 of the C-terminal domain		
Gate engineering			
PDB: WT – 1CB8, 1HM2, 1HMU, 1HM3, 1HMW		Tunnel	
References: ¹⁵¹		Length	
		Bottleneck	
		Role	Active site cavity

Table 3. continued

24	Copper – Containing Amine Oxidase		1.4.3.21
Gate function GATE1 – Blocks the back side of the active site from solvent access		Enzyme function Oxidation of primary amines to aldehydes reducing molecular oxygen to hydrogen peroxide	
Gate location GATE1 – Amine tunnel			
Gate structural basis			
—●	GATE1 Y381 in amine oxidase ECAO W156 in amine oxidase HCAO Y296 in amine oxidase AGAO F298 in amine oxidase PSAO Gating residues can form a π/π ring stacking interaction with the pyridine ring of the cofactor (TPQ)		
Gate engineering			
PDB: WT (ECAO) – 1OAC, WT (HCAO) – 2OQE, 2OOV, WT (AGAO) – 1RJO, 1RKY, WT (PSAO) – 1W2Z		Bottleneck	
References: ^{88,152–154}		Role	

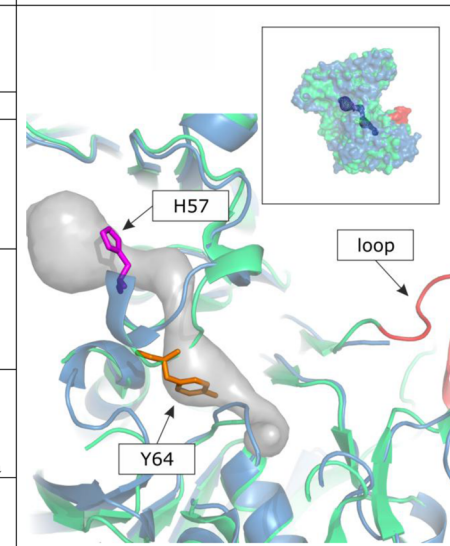
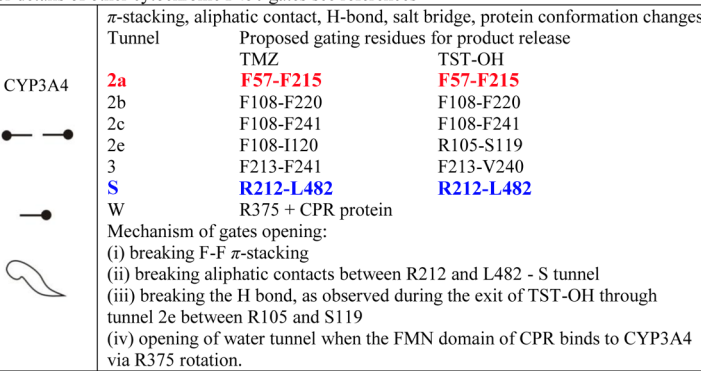
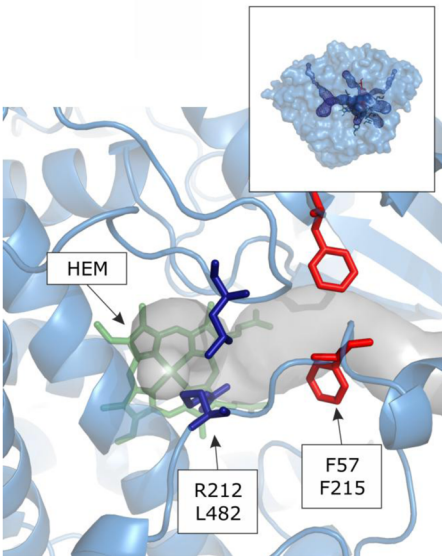
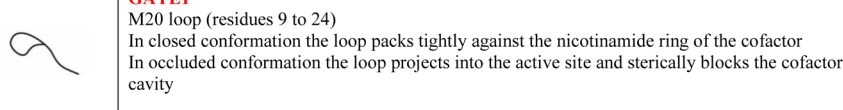
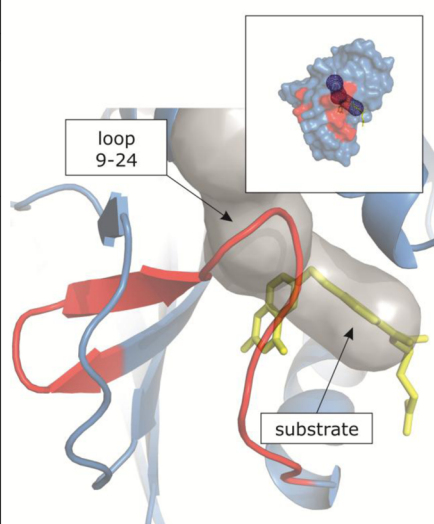
25	Cytidine Triphosphate Synthetase CTPS		6.3.4.2
Gate function GATE1 – Creates tunnel, protects tunnel from access of the solvent ?GATE2 – Controls passage of ammonia to the amidotransferase active site GATE3 – Controls passage of ammonia to the amidotransferase active site		Enzyme function Catalyses the formation of CTP from UTP, ATP and glutamine	
Gate location GATE1 – Loop of the glutaminase domain ?GATE2 – The domain interface, interacts with the amide group of the bound glutamine GATE3 – Between UTP site and the tunnel exit			
Gate structural basis			
—●	GATE1 (PDB ID: 1VCM) Loop The synthetase active site is exposed to the solvent and the binding pocket for the allosteric effector GTP is not properly formed After ATP and acceptor binding loop covers the entrance of the glutaminase site shielding glutamine and creates the tunnel		
—●	?GATE2 (PDB ID: 1VCM) Y64 Y64 might act as a door to the ammonia tunnel leading to the synthetase site		
—●	GATE3 (PDB ID: 2ADP) H57 1. Binding of the substrate UTP induces the rotation of H57 2. Rotation of H57 opens the tunnel for ammonia passage The ligand-induced change is postulated to regulate the timing for the translocation of ammonia		
Gate engineering			
PDB: WT – 1VCM, 1VCN, 1VCO, 2AD5, 1S1M		PDB ID: 1VCM – Light green PDB ID: 2AD5 – Blue Tunnel calculated for PDB ID: 2AD5 structure	
References: ^{52,155–157}		Tunnel between active sites	
		Length 25	
		Bottleneck 0.8–1.2	
		Role NH ₃ transport between active sites	

Table 3. continued

26	Cytochrome P450 CYP3A4		1.14.13.32																															
Gate function Controls access to the active site, controls selectivity and specificity of the enzyme	Enzyme function Catalyses mono-oxygenation reactions such as hydroxylation and epoxidation, major drug metabolizing enzyme in humans																																	
Gate location Depends on the P450 isoform, tunnel and the gate																																		
Gate structural basis Common gating mechanisms in cytochrome P450 family are: (i) associated with F-G-helix-loop-helix or B-C loop movement (tunnels 2a, 2ac, 2b, 2c, 2e); (ii) associated with swinging door type; (iii) associated with wing type; (iv) cofactor assisted (CPR) For details of other cytochrome P450 gates see references ^{16,59,90}																																		
 <p>CYP3A4 Tunnel Proposed gating residues for product release 2a F57-F215 F57-F215 2b F108-F220 F108-F220 2c F108-F241 F108-F241 2e F108-I120 R105-S119 3 F213-F241 F213-V240 S R212-L482 R212-L482 W R375 + CPR protein Mechanism of gates opening: (i) breaking F-F π-stacking (ii) breaking aliphatic contacts between R212 and L482 - S tunnel (iii) breaking the H bond, as observed during the exit of TST-OH through tunnel 2e between R105 and S119 (iv) opening of water tunnel when the FMN domain of CPR binds to CYP3A4 via R375 rotation.</p>	 <p>PDB ID: 1TQN - for clarity only gate of tunnel 2a and tunnel S are shown</p> <table border="1"> <thead> <tr> <th>Tunnel</th> <th>2a</th> <th>2b</th> <th>2c</th> <th>2e</th> <th>3</th> <th>S</th> <th>W</th> </tr> </thead> <tbody> <tr> <td>Length</td> <td>19</td> <td>18</td> <td>17</td> <td>12</td> <td>17</td> <td></td> <td></td> </tr> <tr> <td>Bottleneck</td> <td>4.1</td> <td>4.0</td> <td>3.9</td> <td>4.2</td> <td>4.1</td> <td></td> <td>4.5</td> </tr> <tr> <td>Role</td> <td colspan="3">Different products pathways</td> <td>Proton access/water/substrate</td> <td>Water</td> <td></td> <td></td> </tr> </tbody> </table>		Tunnel	2a	2b	2c	2e	3	S	W	Length	19	18	17	12	17			Bottleneck	4.1	4.0	3.9	4.2	4.1		4.5	Role	Different products pathways			Proton access/water/substrate	Water		
Tunnel	2a	2b	2c	2e	3	S	W																											
Length	19	18	17	12	17																													
Bottleneck	4.1	4.0	3.9	4.2	4.1		4.5																											
Role	Different products pathways			Proton access/water/substrate	Water																													
Gate engineering F108W, I120W – closure of tunnel 2e, switching of the midazolam entrance to another tunnel that may lead to a different active site orientation and product formation																																		
PDB: WT – 1TQN (CYP3A4), 1AKD (P450cam), 2D09 (CYP152A2), 1JPZ (P450-BM3), 1PQ2 (CYP2C8), 1IZO (CYP152A1), 1F4T (CYP119), 1NR6 (CYP2C5), 1OXA (P450eryF)																																		
References: ^{10,16,25,55,56,58,59,90,158,159}																																		

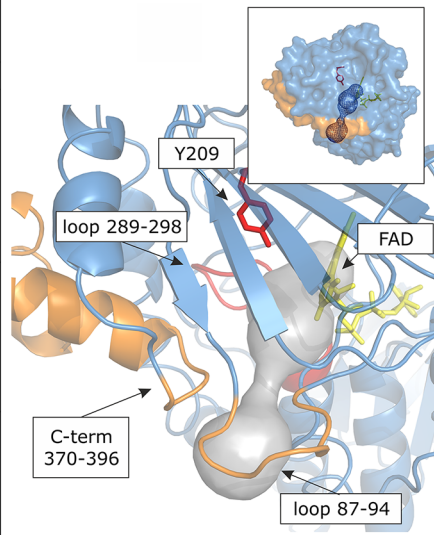
27	Dihydrofolatereductase		1.5.1.3
Gate function GATE1 – Controls the activity	Enzyme function Catalyses the stereospecific reduction of dihydrofolate to tetrahydrofolate		
Gate location GATE1 – Entrance to the cofactor cavity			
Gate structural basis  <p>GATE1 M20 loop (residues 9 to 24) In closed conformation the loop packs tightly against the nicotinamide ring of the cofactor In occluded conformation the loop projects into the active site and sterically blocks the cofactor cavity</p>			
Gate engineering N23PP – k_{cat} decreased 5-fold S148A – k_{cat} decreased 2-fold N23PP/S148A – k_{cat} decreased 6-fold			
PDB: WT – 1RX2, Mutants – 3QL0, 3QL3			
References: ^{80,81,160–162}			



PDB ID: 1RX2

Tunnel	
Length	
Bottleneck	
Role	

Table 3. continued

28	Digeranylgeranylglycerophospholipid Reductase DGGR		1.3.1
Gate function GATE1 – Opens tunnel, controls entry of the substrate, controls reduction of FAD GATE2 – Controls substrate binding/release		Enzyme function Converts 2,3-di-O-geranylgeranylglyceryl phosphate to 2,3-di-O-phytanylgluceryl phosphate (archaetidic acid)	
Gate location GATE1 – Vicinity of the cofactor binding site (FAD) GATE2 – C-terminal helical subdomain			
Gate structural basis			
	GATE1 Glycine-rich α 1– α 2 loop (residues 289–298) Cofactor FAD has two conformation IN (not accessible by solvent) and OUT (exposes the isoalloxazine ring allowing it to be reduced by NADH or NADPH). In IN conformation tunnel A is blocked, to open it has to turn into OUT conformation. The conformational changes in the glycine-rich α 1– α 2 loop disrupt interaction between Y209 and T292 allowing change of the FAD position. Conserved RxxFD and LxDG motifs may play a role in FAD's IN/OUT conformational switch.	 PDB ID: 3OZ2	
	GATE2 Two regions, the β 6– β 7 loop (residues 87–94) and the C-terminal helices (residues 370–396) The conformation changes of the C-terminal helical subdomain may be involved in substrate binding/release		
Gate engineering			
PDB: WT – 3OZ2			
References: ¹⁰⁰			

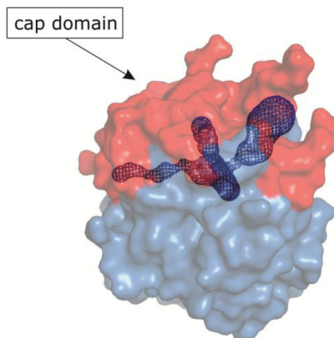
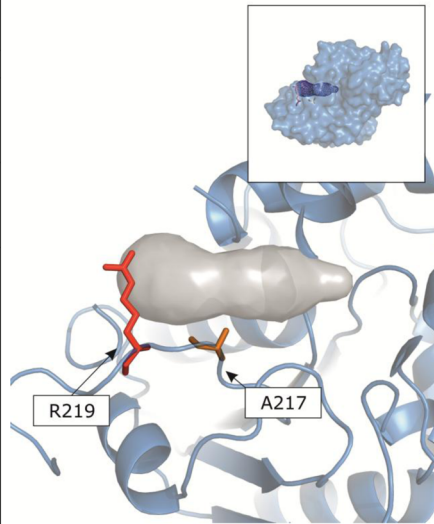
29	Epoxide Hydrolase H37Rv		3.3.2.3
Gate function GATE1 – Regulates access to the active site		Enzyme function Hydrolysis of epoxides	
Gate location GATE1 – Cap domain			
Gate structural basis			
	GATE1 (<i>Mycobacterium tuberculosis</i> H37Rv) Movement of the cap domain regulates access to the active site	 PDB ID: 2E3J	
Gate engineering			
PDB: WT (Aspergillus niger M200) – 1QO7			
References: ⁶⁸			

Table 3. continued

30	Epoxide Hydrolase M200		3.3.2.3
Gate function ?GATE1 – Controls enantioselectivity and activity ?GATE2 – Controls enantioselectivity and activity		Enzyme function Hydrolysis of epoxides	
Gate location ?GATE1 – Tunnel entrance ?GATE2 – Middle of the tunnel		 PDB ID: 1QO7	
Gate structural basis			
—●	?GATE1 (<i>Aspergillus niger</i> M200) R219		
?	—● ?GATE2 (<i>Aspergillus niger</i> M200) A217		
Gate engineering			
<i>Aspergillus niger</i> M200 A217C, A217E, A217G, A217L, A217P, A217Q, A217R, A217T, A217V – at the entrance to the tunnel result in different enantioselectivity and activity; e.g.: A217G mutation results in a 33-fold decrease of activity A217V 6,6-fold increase of activity with no changes in the products enantioselectivity for the reaction with allyl glycidyl			
PDB: WT (<i>Aspergillus niger</i> M200) – 1QO7			
References: ^{26,163}			

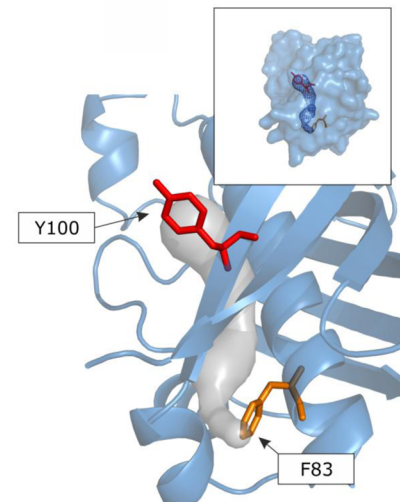
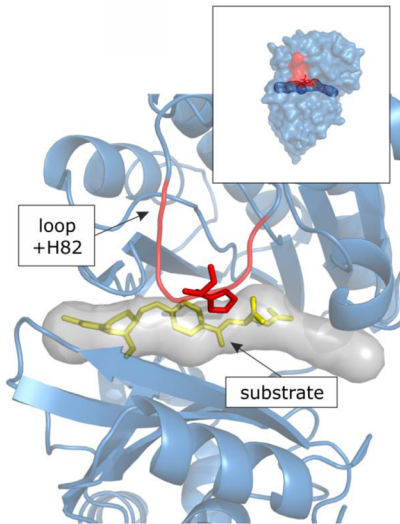
31	FabZ β-Hydroxyacyl-Acyl Carrier Protein Dehydratase		4.2.1.59
Gate function GATE1 – Controls access of the substrate, prohibits binding of small hydrophobic molecules to the unliganded enzyme GATE2 – Controls length of the substrate		Enzyme function Elongation cycles of both saturated and unsaturated fatty acids biosyntheses in the type II fatty acid biosynthesis system (FAS II) pathway	
Gate location GATE1 – Entrance to the tunnel GATE2 – Exit (back door) of the tunnel		 PDB ID: 2GLL	
Gate structural basis			
—●	GATE1 Y100 (Y88 in PaFabZ and L170 in PfFabZ) adopts either an open or closed conformation in the crystal structure In closed conformation - Y100 is stabilized by the Van der Waals interactions with M102, M154, and P112; in open one it flops ~120° and is stabilized by the Van der Waals interactions with M154, K62, and I64		
—●	GATE2 F83 Refined in two additional alternative conformations, leading the tunnel to form a L-shape or an U-shape, in closed conformation - F83 points toward 198, in open position it rotates ~120° points toward 193, exposing the exit to the bulk solvent T85 or F74 play a similar role in PaFabZ		
Gate engineering			
Y100A – drops the activity of the mutant to less than 50% of the enzymatic activity of the wild type, the new entrance has ~15 Å in width, completely exposes the active site to the bulk solvent – as a results ACP binds to the mutant more stronger than to the wild-type. In particular, the dissociation step of ACP from the HpFabZ mutant is extremely slow.			
PDB: WT (<i>HpFabZ</i>) – 2GLL, 2GLP, 2GLM Mutant Y100A – 2GLV, WT (<i>PfFabZ</i>) – 1ZHIG, WT (<i>PcFabZ</i>) – 1U1Z			
References: ^{54,164,165}			

Table 3. continued

32	Formiminotransferase-Cyclodeaminase FTCD		2.1.2.5
Gate function GATE1 – Controls position of H82 in proper orientation for reaction		Enzyme function Two independent, but sequential reactions in the histidine degradation pathway	
Gate location GATE1 – Blocks entrance from a short tunnel into active site		 PDB ID: 1QD1	
Gate structural basis  GATE1 Loop rearrangement including a change in the histidine H82 side-chain position			
Gate engineering			
PDB: <i>WT</i> – 1QD1		Tunnel	Main
References: ⁹⁴		Length	Second
		Bottleneck	8
		Role	4
		Intermediate transport	Formiminoglutamate substrate entry, glutamate product exit

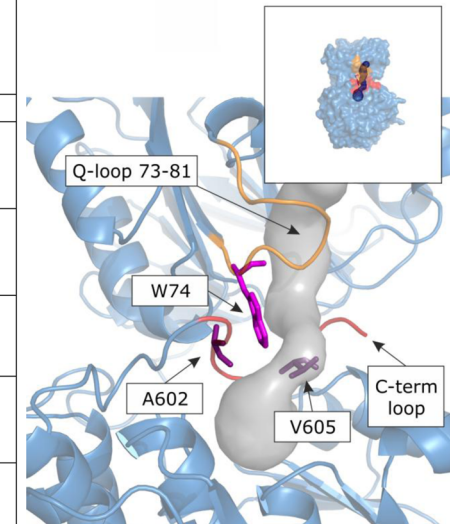
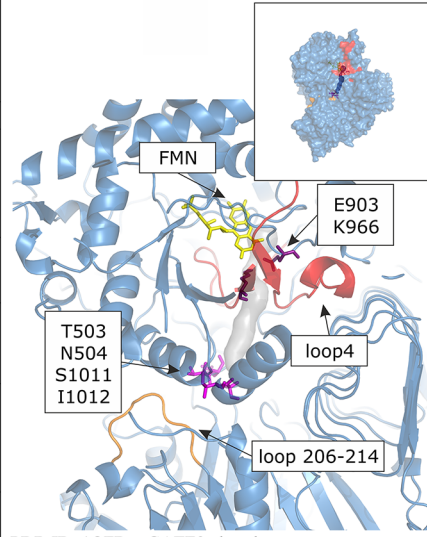
33	Glucosamine 6 Phosphate Synthase GlmS		2.6.1.16
Gate function GATE1 – Protects Fru 6P site from the solvent, creates the ammonia tunnel GATE2 – Protects glutaminase active site from the solvent GATE3 – Opens ammonia tunnel, ammonia enters the sugar site, acts as a solvent barrier GATE4 – Opens ammonia tunnel		Enzyme function Catalyses glucosamine-6P synthesis from fructose-6P and glutamine	
Gate location GATE1 – Part of the active site cavity GATE2 – Part of the active site cavity GATE3 – Tunnel between two active sites GATE4 – Bottleneck of the tunnel between active sites		 PDB ID: 2BPL	
Gate structural basis  GATE1 C-terminal loop (residues 600–608) Open state – relaxed loop Closed state – anchored by Y28 and W74 with Q loop  GATE2 Q-loop (residues 73–81) Open state - anchored by R539*, Close state - after Q enters the active site anchored by Y28 and W74 with C-terminal loop  GATE3 W74 x1 torsion angle of W74 changes by 75°  GATE4 A602 and V605 A shift of the side-chain of the A602 residue is concomitant to a re-orientation of its backbone carbonyl group, which can then form a strong H-bond with the hydroxyl group of the Y28			
Gate engineering W74A, W74L, W74F – inefficient ammonia transfer A602L, V605L – efficiency of ammonia transfer decreased 2-fold		Tunnel	
PDB: <i>WT</i> – 2J6H, 1JXA, 2VF4, 2BPL		Length	18
References: ^{14,45,50,155,166,167}		Bottleneck	
		Role	Ammonia transport

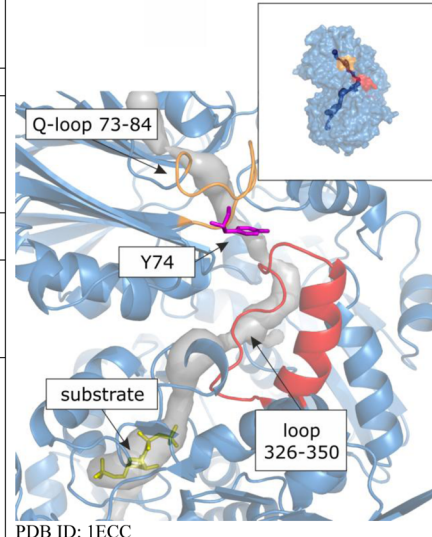
Table 3. continued

34	Glutamate Synthases GltS		1.4.7.1
Gate function		Enzyme function	
GATE1 – Controls correct conformation of active site for L-glutamine binding and hydrolysis, crucial for glutaminase activation and coupling of the glutaminase and synthase sites, creates ammonia tunnel		Formation of 2 L-glutamate from L-glutamine and 2-oxoglutarate	
GATE2 – Shields substrate from bulk solvent			
GATE3 & GATE4 – Controls transport of ammonia			
Gate location			
GATE1 – Loop protecting Fru 6P site from solvent			
GATE2 – Loop protecting glutaminase site from solvent			
GATE3 – Entrance to the interdomain tunnel			
GATE4 – Entrance to the synthase site (end of the tunnel)			
Gate structural basis			
	GATE1 Loop 4 (residues 933–978) Shifts to the active conformation – C-terminal residue E1013 forms a hydrogen bond with C1 and keeps correct conformation of C1 and loop 29–34 for L-glutamine binding and hydrolysis. E1013 side chain may also play a role in the precise geometry of the tunnel entry point		
	GATE2 Loop 206–214 Closing of loop 206–214 after L-glutamine binding		
	GATE3 T503, N504, S1011 and I1012 Small conformation changes of T503, N504 of the central domain, S1011 and I1012 of loop 4		
	GATE4 E903 and K966 Small conformation changes of E903 and K966 Additionally residues T507 and N508 and S976 and I977 may function as a gates for signalling between active sites		
Gate engineering			
E1013D – 100-fold decrease of activity, a sigmoid dependence of initial velocity on L-glutamine concentration			
E1013N – 1000-fold decrease of activity, exhibited hyperbolic kinetics			
E1013A – 1000-fold decrease of activity			
PDB: WT – 1OFD, 1OFE, 1EA0			
References: 46,62,155,168			



PDB ID: 1OFD – GATE3 closed

35	Glutamine Phosphoribosylpyrophosphate Amidotransferase		2.4.2.14
Gate function		Enzyme function	
GATE1 – Closes synthase site and creates a narrow, solvent-inaccessible tunnel between active sites		Catalyses the initial reaction in <i>de novo</i> purine nucleotide biosynthesis	
GATE2 – Covers glutaminase site			
GATE3 – Protects tunnel from the solvent access and avoids wasteful release of ammonia into solution			
Gate location			
GATE1 – Loop near the synthase active site			
GATE2 – Loop near the glutaminase active site			
GATE3 – On Q-loop			
Gate structural basis			
	GATE1 Loop (residues 326–350) The active site cavity is created by closure of the loop (residues 326–350) Closure of the active site creates a narrow, solvent-inaccessible tunnel between active sites I335 interacts with Y74		
	GATE2 Q-loop (residues 73–84)		
	GATE3 Y74 A key residue in coupling the glutamine and acceptor sites upon acceptor binding may play a similar gate role as W74 in GlmS		
Gate engineering			
PDB: WT – 1ECF, 1ECC			
References: 155,169,170			



PDB ID: 1ECC

Tunnel between two active sites

Length 20

Bottleneck

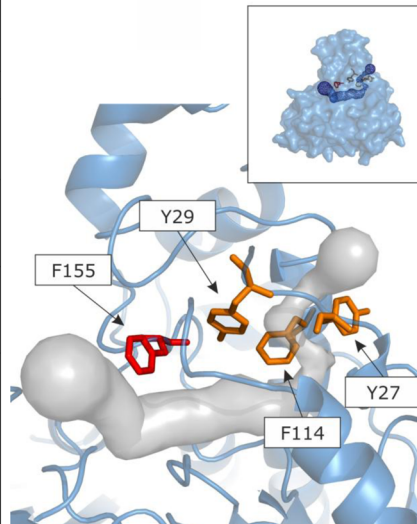
Role Ammonia transport

Table 3. continued

36	Haloalkane Dehalogenase DhaA			3.8.1.5
Gate function	GATE1, GATE2, GATE3, GATE4, GATE5 – Controls access to the tunnel			
Gate location	GATE1 – Tunnel p1, GATE2 – Tunnel p2a, GATE3 – Tunnel p2b, GATE4 – Tunnel p2c, GATE5 – Tunnel p3			
Gate structural basis				
● — ●	GATE1 F144 and F149			
— ●	GATE2 I135			
● — ●	GATE3 V245, W141			
● — ●	GATE4 β-bridge interaction between P210 and A212 of the CC loop and I135 of the NC loop			
— ●	GATE5 W138 blocks tunnel p3			
Gate engineering	C176Y, V245F, A172F, A145F – limiting access of water to the active site			
PDB: WT (<i>DhaA</i>) – 1CQW, 1BN6, 1BN7				
References: ^{109,110}				
				PDB ID: 1CQW
Tunnel	p1	p2a, p2b, p2c	p3	
Length	20	22	24	
Bottleneck	2.8	1.8	1.6	
Role	Product	Product	Product	

37	Haloalkane Dehalogenase LinB			3.8.1.5
Gate function	GATE1 – Controls access to the active site GATE2 – Controls selectivity			
Gate location	GATE1 – Tunnel bottleneck GATE2 – Tunnel bottleneck			
Gate structural basis				
— ●	GATE1 D147 Motion of this residues is necessary for entry of large substrates			
— ●	GATE2 L177 Partially blocks the entrance of the main tunnel			
Gate engineering	L177A, L177C, L177G, L177F, L177K, L177T, L177W, L177D, L177H, L177M, L177Q, L177R, L177S, L177V, L177Y influence the substrate specificity and activity			
PDB: WT (<i>LinB</i>) – 1CV2				
References: ^{72,108,171}				
				PDB ID: 1CV2
Tunnel	p1			
Length	12			
Bottleneck	2.6			
Role	Product			

Table 3. continued

38	Histone Deacetylase HDAC1 and HDAC2		3.5.1.98			
Gate function	Enzyme function Modifies chromatin structures by removing the acetyl group from the ε-amino lysine residues					
GATE1 – Controls access to the active site through tunnel A GATE2 – Controls access to the active site through tunnel B1 GATE3 – Controls access to the active site through tunnel B2 (only HDAC1)						
Gate location						
GATE1 – Bottleneck of tunnel A GATE2 – Bottleneck of tunnel B1 GATE3 – Bottleneck of tunnel B2 (only HDAC1)						
Gate structural basis						
—●	GATE1 F155 (F150) Side chain of F155 has to rotate ~ 180°					
●—●	GATE2 F114 + Y27 and Y29 (F109 + Y22 and Y24) F114 residue is located between two tyrosine rings in closed conformation Opening requires unzipping of the gate					
—●	GATE3 Y303 Observed only in HDAC1 – Y303 residue may rotate opening access through tunnel – anchoring residue in closed position M30					
Gate engineering						
PDB: WT (HDAC2) – 3MAX						
References: ¹⁷²						
						
PDB ID: 3MAX Tunnel A B1 B2 Length 11 14 16 Bottleneck						
Role Access to the active site Access to the active site Water exchange						

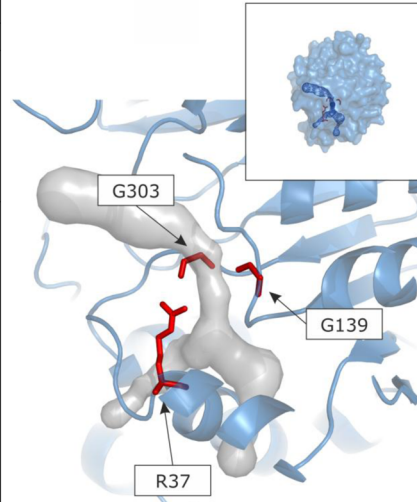
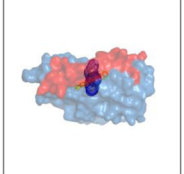
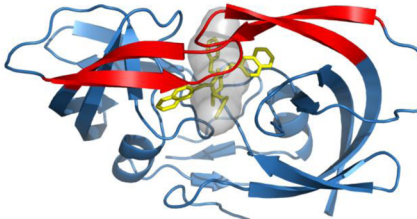
39	Histone Deacetylase HDAC8		3.5.1.98			
Gate function	Enzyme function Deacetylation of the ε-amino group of specific lysine residues within histones and other proteins					
GATE1 – Regulates water or product (acetate) transit from the active site through the internal tunnel						
Gate location						
GATE1 – Centre of the internal tunnel						
Gate structural basis						
—●	GATE1 R37 A structural reorientation of R37 and the loop is required for opening the access to the active site via the 14 Å ‘internal’ tunnel. R37 forms multiple hydrogen bond interactions with the backbone carbonyl oxygen atoms of conserved G303 and G305 positioned in a loop between the β8sand α10-helix Gating interaction between G139 and G303					
Gate engineering						
R37A – the values for k_{cat}/K_M decrease 530-fold						
PDB: WT – 2V5W						
References: ¹⁷³						
						
PDB ID: 2V5W Tunnel Length 11 14 Bottleneck						
Role Access to active site						

Table 3. continued

40	HIV-1 Protease	3.4.23.16
Gate function GATE1 – Controls access to the active site	Enzyme function Central role in processing HIV-1 viral polypeptide precursors	
Gate location GATE1 – β -turn flaps		
Gate structural basis		
	GATE1 β -turn flaps (residues 43–58) Movement of two β -turn flaps controls access to the active site	
Gate engineering		
WT – open 14% of time (Brownian dynamics simulations) G48V/V82A – open 2% of time (Brownian dynamics simulations) I84V/L90M – open 2% of time (Brownian dynamics simulations) L90M, G48V – open 14% of time (Brownian dynamics simulations) F53L – unstabilised semi open conformation due to lack of F53-I50 interaction		PDB ID: 1HVR
PDB: WT – 1HHP, 1HVR	Tunnel	
References: ^{79,89,174–177}	Length	
	Bottleneck	
	Role	Active site gorge

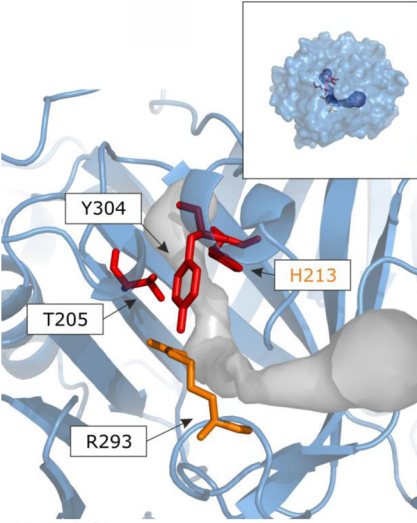
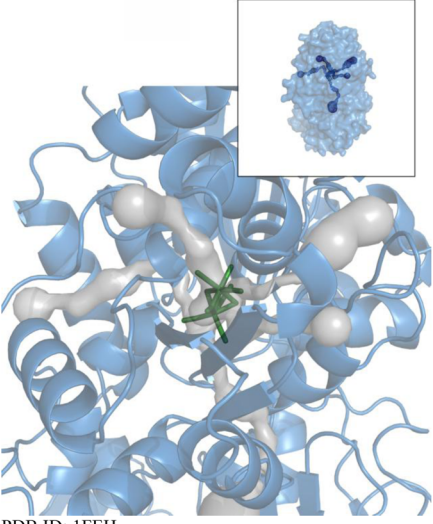
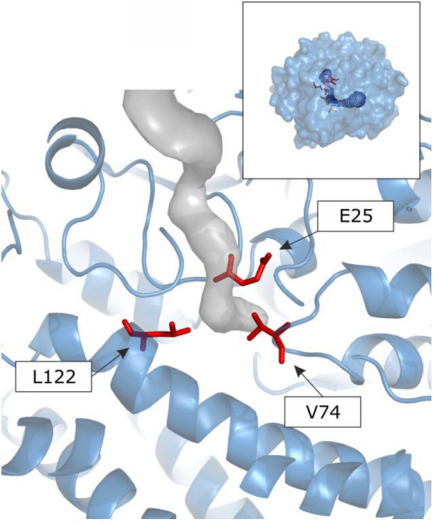
41	Homoprotocatechuate 2,3-Dioxygenase	1.13.11.15
Gate function GATE1 – Controls O ₂ diffusion pathway GATE2 – Controls alternative O ₂ diffusion pathway	Enzyme function Degradation of catechol and its derivatives by cleavage of aromatic rings	
Gate location GATE1 – Just below the protein surface GATE2 – Below the protein surface		
Gate structural basis		
	GATE1 T205, H213, and W304 Breathing motion of protein causes synchronizing movement of residues	
	GATE2 R293, H213 The fluctuation of R293 along with H213 could result in opening of the pathway	PDB ID: 2IG9
Gate engineering	Tunnel	
	Length	
	Bottleneck	
PDB: WT – 2IGA, 2IG9	Role	O ₂ transport
References: ⁶³		

Table 3. continued

42	Hydrogenase FeFe		1.12.7.2		
Gate function Controls access of oxygen and hydrogen to the active site			Enzyme function Catalyses the reversible oxidation of molecular hydrogen		
Gate location Not specified, between dynamic cavities					
Gate structural basis					
	No permanent tunnel – O ₂ moves from cavity to cavity as the cavities fluctuate inside the protein				
Gate engineering					
					
PDB: WT – 1HFE, 1FEH					
References: ^{88,178–180}					
43	Hydrogenase NiFe		1.12.2.1		
Gate function ?GATE1 – Protects the active site against O ₂			Enzyme function Catalyses the reversible oxidation of molecular hydrogen		
Gate location ?GATE1 – Vicinity of NiFe cluster, bottleneck close to the active site					
Gate structural basis					
	?GATE1 E25, V74 and L122 The size of the amino acids at positions 122 and/or 74 may determine the accessibility of the active site and therefore the resistance to O ₂				
Gate engineering V74Q, V74M, V74E, V74N, V74W, V74F, V74D, L122M/V74M, L122F/V74I, L122A/V74M – two contributions, size and polarity, are independent and have different effects to H ₂ , CO, O ₂ molecules, different influences on reaction rate and inhibition effect					
					
PDB: WT – 1YQW Mutants – 3CUR, 3CUS, 3H3X					
References: ^{21,22}					

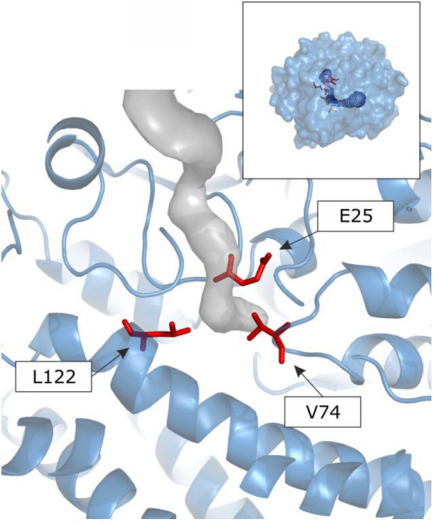
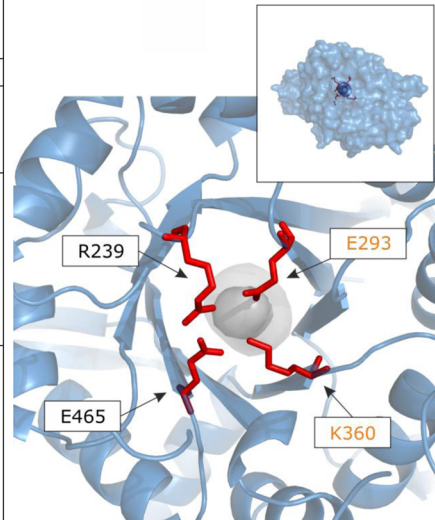
43	Hydrogenase NiFe		1.12.2.1		
Gate function ?GATE1 – Protects the active site against O ₂			Enzyme function Catalyses the reversible oxidation of molecular hydrogen		
Gate location ?GATE1 – Vicinity of NiFe cluster, bottleneck close to the active site					
Gate structural basis					
	?GATE1 E25, V74 and L122 The size of the amino acids at positions 122 and/or 74 may determine the accessibility of the active site and therefore the resistance to O ₂				
Gate engineering V74Q, V74M, V74E, V74N, V74W, V74F, V74D, L122M/V74M, L122F/V74I, L122A/V74M – two contributions, size and polarity, are independent and have different effects to H ₂ , CO, O ₂ molecules, different influences on reaction rate and inhibition effect					
					
PDB: WT – 1YQW Mutants – 3CUR, 3CUS, 3H3X					
References: ^{21,22}					

Table 3. continued

44	Imidazole Glycerol Phosphate Synthase IGPS		2.4.2
Gate function		Enzyme function	
GATE1 – Prevents penetration of bulk water molecules into chambers I and II		Catalyses the closure of the imidazole ring within histidine biosynthesis and provides 5-aminoimidazole-4-carboxamide ribotide (AICAR) for use in the <i>de novo</i> synthesis of purines	
GATE2 – Controls ammonia transfer, discriminates between ammonia and water			
Gate location			
GATE1 – Near the entrance of the $(\beta\alpha)_8$ barrel of hisF			
GATE2 – Between chamber I and chamber II			
Gate structural basis			
	GATE1 (for water) R5 (R239), E46 (E293), K99 (K360), and E167(E465) numbers from yeast or (<i>Thermotoga maritima</i>) Four strictly conserved gate residues act as the wall barrier for water molecules		
	GATE2 (for ammonia) K99 (K360) and E46 (E293) Conformational change of all residues – high barrier 25 kcal/mol for ammonia transfer K99 (K360) side chain rotation barrier 10 kcal/mol Both K99 and R5 – fully open gate barrier 2 kcal/mol Ammonia enters to chamber II through a side-opening between residues K99 and E46, the only requisite is the slight bending of the side chain of K99 (K360) - eliminates the need for an energetically costly gate-opening mechanism		
Gate engineering			
Y138F – experimentally no change of kinetics and stoichiometry, in simulations Y138 is not a gate but prevents bulk water from entering the interface during a reaction, keeping ammonia sequestered within the tunnel			
R5A (R239A) – loss of ammonia through new hole and results in a 10^3 decrease in k_{cat}/K_M values for the cyclase reaction, in simulations possibly because the mutation allowed water molecules to access chamber II; these additional water molecules increased the energetic barrier to ammonia entry and passage through the tunnel			
K99A (K360A) – experimentally 3-fold decrease in the overall reaction stoichiometry, in simulations larger opening between chamber I and chamber II, and simultaneously deprives E167 and E46 of a salt-bridge partner T78A – allowed a rapid and unhindered conduction of ammonia through the tunnel			
T78F, P76F – block the ammonia conduction			
Any mutation K181 (K196), D98 (D359), and Q123 (Q387) – decouples the two reactions			
PDB: WT – 1KA9, 1JVN			
References: ^{38,43,155,181–186}			



45	Inosine 5'- Monophosphate Dehydrogenase		1.1.1.205
Gate function		Enzyme function	
GATE1 – Conformational change converts the enzyme from a dehydrogenase into hydrolase		Transformation of inosine 5'-monophosphate into xanthosine 5'-monophosphate	
Gate location			
GATE1 – Loop covering active site cavity			
Gate structural basis			
	GATE1 C319 loop (313-328) adopts different conformations during the dehydrogenase and hydrolase reactions		
Gate engineering			
PDB: WT – 1ME9, 1LRT, 1ME8, 1PVN			
References: ¹⁸⁷			

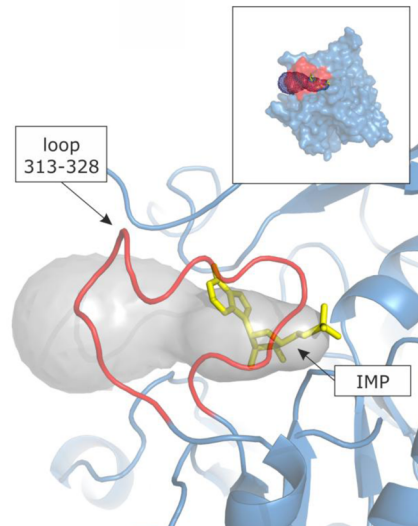


Table 3. continued

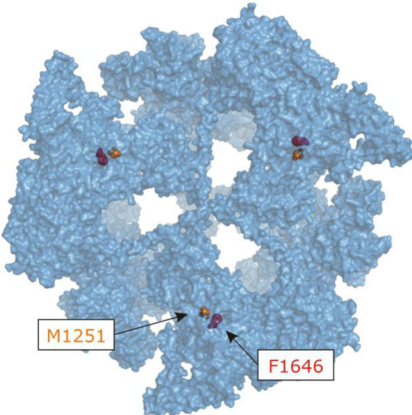
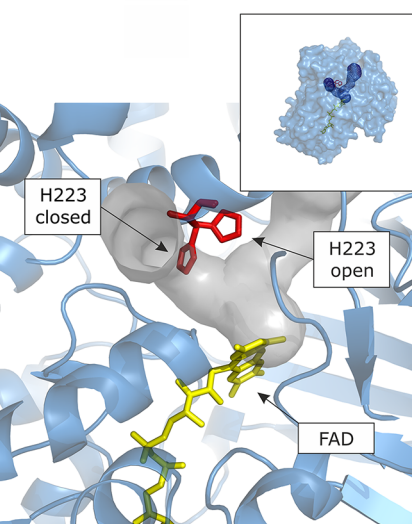
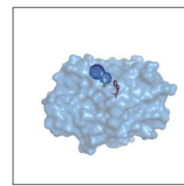
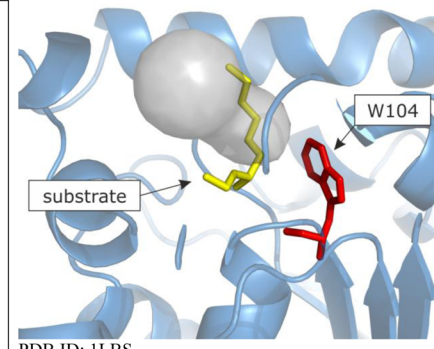
46	Ketoacyl Synthase KS		2.3.1.86
Gate function GATE1 – Shields the active site GATE2 – Controls access of the substrate		Enzyme function Elongates an ACP- or CoA-associated acyl chain by adding C2 units through a ping-pong decarboxylating condensation mechanism	
Gate location GATE1 – Entrance to the active site GATE2 – Centre of the acyl-binding tunnel			
Gate structural basis			
—●	GATE1 F1646 Shields the active site, flips and allows access to the nucleophilic cysteine		
—●	GATE2 M1251 rotates and unlocks the inner part of the fatty acid binding cavity		
Gate engineering			
			
		PDB ID: 2VKZ	
		Tunnel	
		Length	
		Bottleneck	
		Role	
47	L-Amino Acid Oxidase		1.4.3.2
Gate function GATE1 – Controls access to the oxygen tunnel, binds the substrate initially		Enzyme function Oxidative deamination of L-amino acid substrates	
Gate location GATE1 – 15 Å from surface in main funnel			
Gate structural basis			
—●	GATE1 H223 H223 has two conformations and can act as a gate and binds the substrate initially H223 conformation A closing oxygen tunnel → substrate entry and deprotonation of the zwitterion → substrate into Michaelis position → H223 turn into conformation B → opening of oxygen tunnel		
Gate engineering			
		PDB ID: 1F8S	
		Tunnel	Main funnel
		Length	25
		Bottleneck	4.2
		Role	Access to active site
		O_2 access and H_2O_2 release	
PDB: WT – 1F8R, 1F8S			
References: ^{73,189}			

Table 3. continued

48	Lipase B		3.1.1.3
Gate function (lipase from <i>Candida antarctica</i>) GATE1 – Controls regiospecificity, controls length of the substrate		Enzyme function Hydrolyse triacylglycerols and a broad range of other substrates, important for asymmetric synthesis	
Gate location GATE1 – Bottom of the substrate tunnel			
Gate structural basis			
<p>—●</p> <p>GATE1 W104 The stereospecificity pocket is defined by T42, S47 and W104 Pocket is buried under a surface helix and delimited by the side chain of W104 <i>R</i> enantiomer of butanoic ester fits well into the active site pocket after a small movement of the side chain of W104</p>		 <p>PDB ID: 1LBS</p>	
Gate engineering W104A, W104Q – change in substrate specificity		Tunnel	
Gate creation in other lipases Lipase from <i>Burkholderia cepacia</i> – mutation in bottleneck for increased enantioselectivity – the best double mutant L17S/L287I - 15-fold increased activity and a tenfold enhanced enantioselectivity		Length	
<i>Candida rugosa</i> lipase 1 – mutation of amino acids in different position inside the tunnel – P246F, L413F, L410W, L410F/S300E, L410F/S365L – different chain length selectivity		Bottleneck	
PDB: WT (<i>Candida antarctica</i>) – 1LBS, 3ICV, 3ICW, WT (<i>Burkholderia cepacia</i>) – 3LIP, WT (<i>Candida rugosa</i>) – 1LPO		Role	Access to active site
References: ^{86,190–192}			

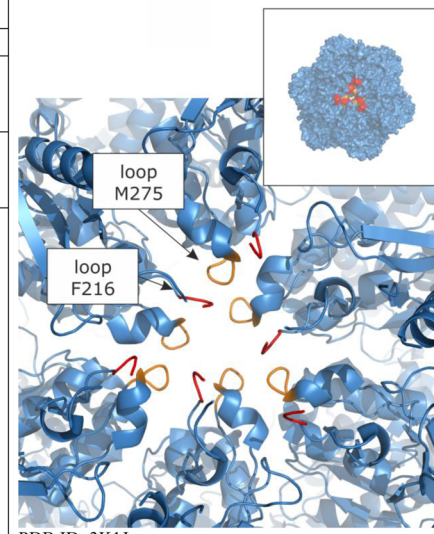
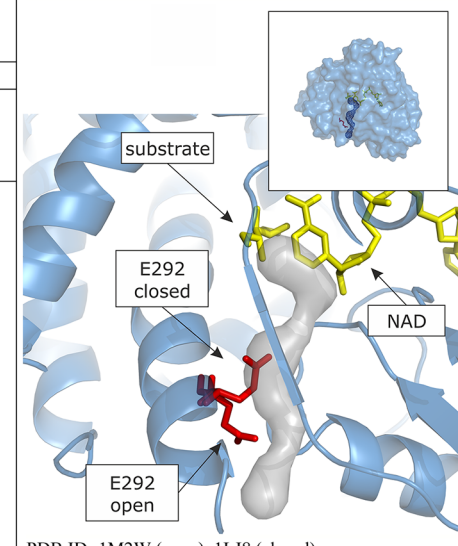
49	Lon Protease		3.4.21.53
Gate function GATE1 – Controls the access of the substrate		Enzyme function ATP-dependent proteolysis	
Gate location GATE1 – At the entrance to an internal unfolding and degradation chamber			
Gate structural basis		 <p>PDB ID: 3K1J</p>	
<p>—●</p> <p>GATE1 F216 loop</p> <p>GATE2 M275 loop</p>		Tunnel	
Gate engineering F216A – lost almost all of the ATP-dependent proteolytic activity against a casein and the aromatic peptide		Length	
		Bottleneck	
PDB: WT – 3K1J		Role	
References: ¹⁹³			

Table 3. continued

50	Mannitol 2-Dehydrogenase		1.1.1.67
Gate function	GATE1 – E292 functions as a gate in water chain mechanism of proton translocation		Enzyme function
Gate location	GATE1 – Bottleneck of the tunnel connecting active site with protein surface		Dehydrogenation of mannitol
Gate structural basis	 GATE1 E292 may adopt two conformation open and closed		
Gate engineering	E292A – 120-fold decrease in a rate of microscopic steps preceding catalytic oxidation of mannitol		
PDB: WT – 1LJ8 (closed), 1M2W (open)			
References: ¹⁹⁴			
 PDB ID: 1M2W (open), 1LJ8 (closed)			
Tunnel			
Length			
Bottleneck			
Role	Proton transfer		

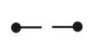
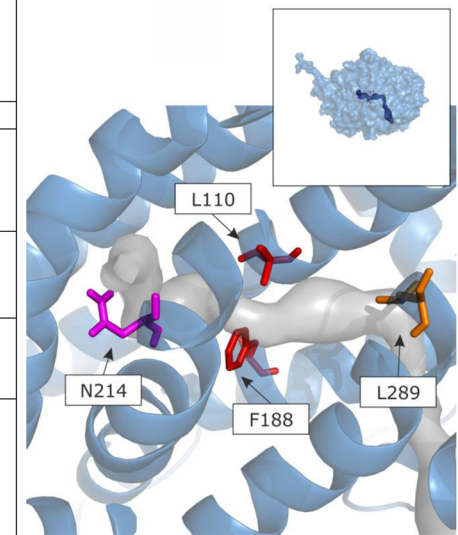
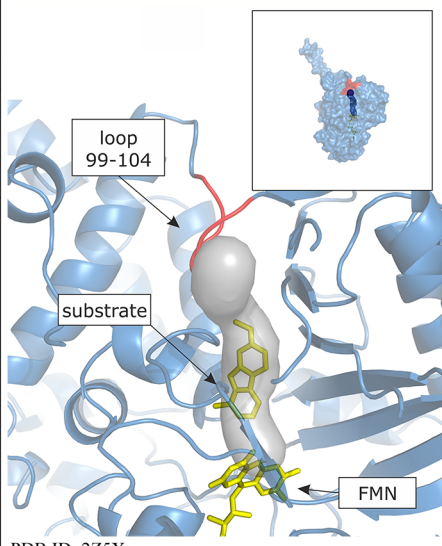
51	Methane Monooxygenase Hydroxylase MMOH		1.14.13.25
Gate function	GATE1 – Controls transport of the substrate between cavities 1 and 2 GATE2 – Allows substrate to traverse the protein cavities GATE3 – Controls entrance of small substrates such as O ₂ , H ₃ O ⁺ , CH ₄		Enzyme function
Gate location	GATE1 – Separates cavities 1 and 2 GATE2 – Separates cavities 2 and 3 GATE3 – One of the residue from active site is positioned at the interface between the cavity and the surface above the iron-coordinating residue		Converts hydrocarbon substrates either to alcohols or epoxides
Gate structural basis	 GATE1 L110, F188 Conformation changes of L110 and F188 opens the access between the cavities 1 and 2, other residue involved T21		
	GATE2 L289 Moves to allow the substrate analogues to traverse between the cavities 2 and 3, other residues creating the bottleneck F109, V285 and Y291		
	GATE3 N214 Conformation changes – movement of N214 forms a deep crevice in the four-helix bundle		
Gate engineering			
PDB: WT – 1MHY, 1XVG, 1XVF, 1XVE, 1XVB, 1XVC, 1XU5, 1XVD, 1XU3			
References: ^{53,74,195}			
 PDB ID: 1MHY			
Tunnel			
Length			
Bottleneck			
Role			

Table 3. continued

52	Monoamine Oxidase A		1.4.3.4
Gate function GATE1 – Controls the access of the substrate		Enzyme function Oxidative deamination of biogenicamines and amine neurotransmitters (serotonin, dopamine and epinephrine)	
Gate location GATE1 – At the entrance to the active site cavity			
Gate structural basis			
	GATE1 Loop 99-104 The loop movement is regulating access to the active site cavity and the loop flexibility is critical for opening the entry for substrates/inhibitors	 loop 99-104 substrate FMN PDB ID: 2Z5X	
Gate engineering G110A – increase of K_m of 5-fold G110P – increase of K_m of 19-fold			
PDB: WT – 2Z5X, Mutant G110A – 2Z5Y			
References: ^{196,197}			

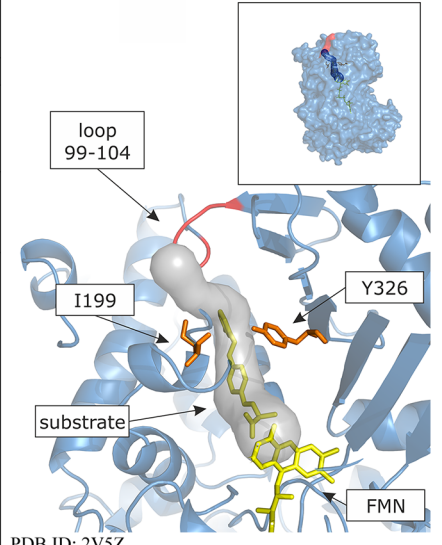
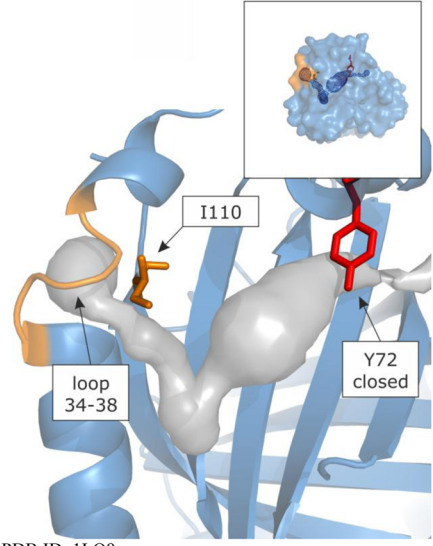
53	Monoamine Oxidase B		1.4.3.4
Gate function GATE1 – Controls the access of the substrate GATE2 – Controls the size of the active site cavity by separating or merging two smaller cavities		Enzyme function Oxidative deamination of biogenic amines and amine neurotransmitters (serotonin, dopamine and epinephrine)	
Gate location GATE1 – At the entrance to the active site cavity GATE2 – Separates two internal cavities			
Gate structural basis			
	GATE1 Loop 99-104 F103 side chain conformation movement is synchronized with the conformation changes of I199 residues (I199 open → F103 closed)	 loop 99-104 I199 Y326 substrate FMN PDB ID: 2V5Z	
Gate engineering I199A/Y326A – exhibits inhibitor binding properties more similar to those of monoamine oxidase A			
PDB: WT – 2V5Z, Mutant I199A/Y326A – 3ZYX			
References: ^{197,198}			

Table 3. continued

54	Monooxygenase ActVAOrf6		Not determined
Gate function		Enzyme function	
GATE1 – Controls access of the substrate, hydrogen bond donor and acceptor/proton gate		Oxidation of a phenolic compound 6-deoxydihydrokalafungin at the C-6 position into the corresponding quinone dihydrokalafungin	
GATE2 – Controls opening of the narrow tunnel, can also control exit for H ₂ O, O ₂ , H ₂ O			
Gate location			
GATE1 – Entrance to the active site			
GATE2 – Opposite side of the bound substrate			
Gate structural basis			
—●	GATE1 Y72 The residue possess two possible conformation that can act as a gate and act as hydrogen bond donor and acceptor		
—●	GATE2 I110 + loop (residues 34-38) Conformation changes close and open the narrow tunnel; gate can also control an exit path for H ₂ O		
Gate engineering			
PDB: WT – 1LQ9		PDB ID: 1LQ9	
References: ^{14,92}		Tunnel	
		Length	
		Bottleneck	
		Role	Proton transfer
			Oxygen/water transport

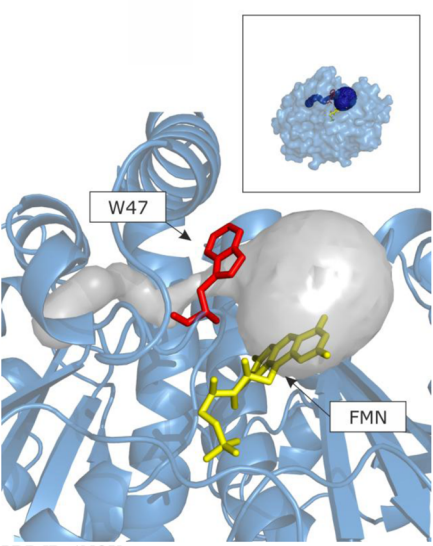
55	NADH Oxidase		1.6.99.3
Gate function		Enzyme function	
GATE1 – Controls accessibility of the flavin ring and plays a crucial role during the catalytic cycle		Hydride transfer from NADH to the intrinsic flavin cofactor	
Gate location			
GATE1 – Above the active site			
Gate structural basis			
—●	GATE1 W47 and cofactor FAD Cofactor-assisted gating mechanism, W47 moves from the original position toward the pyrimidine section Closed conformation is stabilized by the hydrogen bonds between cofactor and peptide backbone Stabilization of the open form may have advantages during the initial steps of the substrate binding, it may slow down the product dissociation		
Gate engineering		PDB ID: 1NOX	
PDB: WT – 1NOX		Tunnel	
References: ^{97,199}		Length	
		Bottleneck	
		Role	

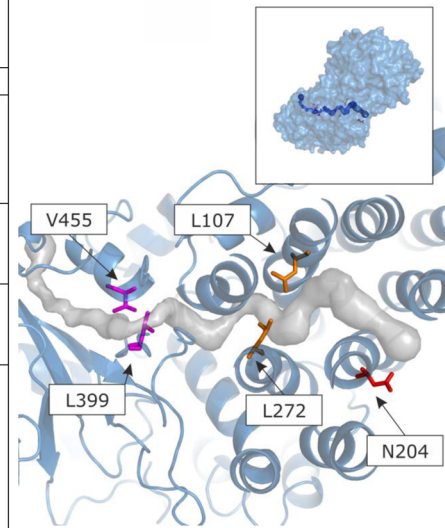
Table 3. continued

56	O-Acetylserine Sulphydrylase Cysteine Synthase		2.5.1.47
Gate function GATE1 – Controls access to the active site GATE2 – Controls access to the active site		Enzyme function Production of cysteine from the O-acetyl-serine intermediate	
Gate location GATE1 – Bottleneck of tunnel 12–20 Å from the active site cavity GATE2 – Bottleneck of tunnel 8–12 Å from the active site cavity			
Gate structural basis			
	GATE1 Motion of side chains in addition to backbone movements M101 or S100 on the loop of the upper domain	 PDB ID: 1Z7Y	
	GATE2 Motion of side chains in addition to backbone movements S75 on the Asn-loop for the inner gate		
Gate engineering			
PDB: WT – 1Z7Y, 1Z7W			
References: ²⁰⁰		Role Access to the active site	

57	Oxidosqualene Cyclase SeeOSC		5.4.99.7
Gate function GATE1 – Controls access of the substrate ?GATE2 – Controls access of the substrate		Enzyme function Cyclization of the 2,3-oxidosqualene into lanosterol	
Gate location GATE1 – Bottleneck between tunnel and active site cavity ?GATE2 – Bottleneck between tunnel and active site cavity			
Gate structural basis			
	GATE1 Y239 (Y237 in <i>HsaOSC</i>) A rotation of the side chain stabilized by a hydrogen bond bridge with P228 could open the channel and enable the substrate to enter the active site	 PDB ID: 1W6K	
	?GATE2 T235 (C233 in <i>HsaOSC</i>)		
?			
Gate engineering			
Y239F – 5-fold decrease in enzyme activity, mutation keeps the channel in a closed conformation Y239A – 2-fold decrease in enzyme activity, mutation keeps the channel in an open conformation			
PDB: WT (<i>HsaOSC</i>) – 1W6K			
References: ²⁰¹		Role Connects active site cavity with bulk solvent	

Table 3. continued

58	Phenol Hydroxylase PHH		1.14.13.7
Gate function GATE1 – Controls access to the active site GATE2 – Closes cavity 2 GATE3 – Closes cavity 2		Enzyme function Hydroxylates aromatic compounds	
Gate location GATE1 – Entrance to the tunnel GATE2 – Between active site pocket and cavity 2 GATE3 – Between cavity 3 and cavity 2			
Gate structural basis			
—●	GATE1 N204 The residue shift is redox-dependent Helix E orients N204 away from the active site and opens the tunnel N204 in closed conformation forms a hydrogen bond with S72		
● — ●	GATE2 L107 and L272		
● — ●	GATE3 L399 and V455		
Gate engineering			
PDB: WT – 2INP, 2INN			
References: ^{74,195}			



59	Phospholipase A2		3.1.1.4
Gate function GATE1 – Controls access to the interface and active sites		Enzyme function Catalyses the hydrolysis of acyl bonds in sn-3-phospholipids	
Gate location GATE1 – Position of monomers			
Gate structural basis			
—	GATE1 The dimer interface might act as a hinge The homodimer can be observed in open and closed conformations formed by different angle between monomers		
Gate engineering			
PDB: WT – 1CLP (open), 1PP2 (closed)			
References: ^{69,202,203}			

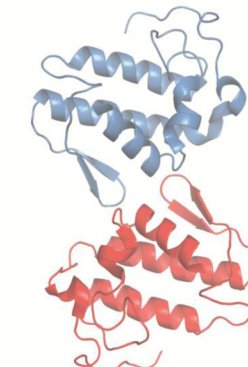
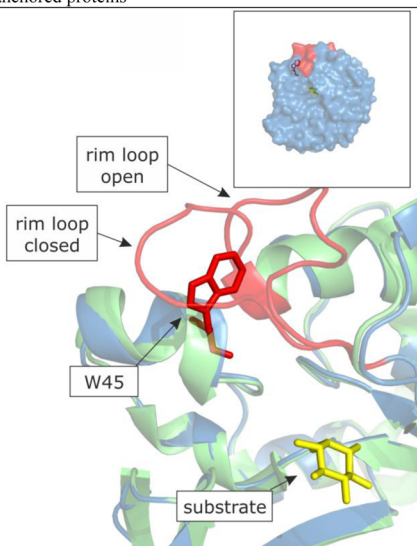
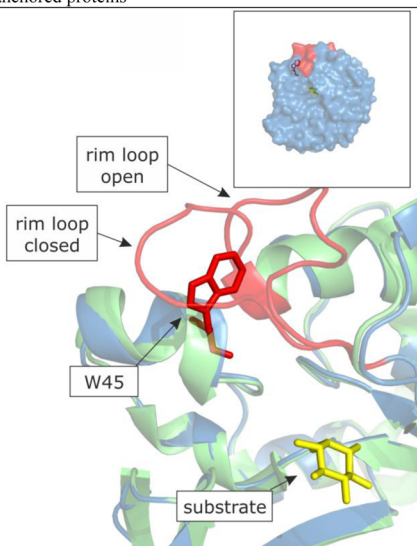
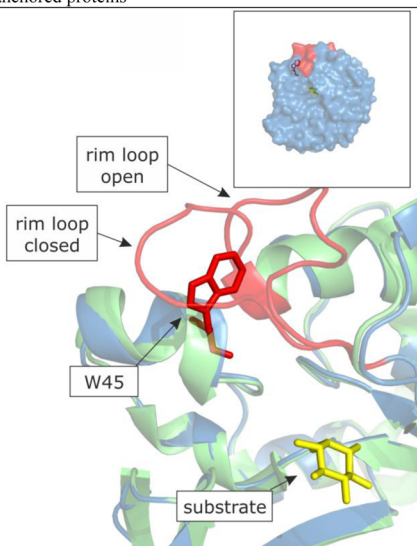


Table 3. continued

60	Phosphatidylinositol-Specific Phospholipase C		4.6.1.13
Gate function GATE1 – Controls access to the active site, controls product exit		Enzyme function Catalyses the cleavage of glycan-phosphatidylinositol anchored proteins	
Gate location GATE1 – Entrance to the active site			
Gate structural basis			
Gate engineering H258Y – keeps the rim mobile loop in extended conformation			
PDB: WT – 3V16, 3V18, <i>Mutants</i> H258Y – 3V1H		PDB ID: 3V16 – blue PDB ID: 3V18 – light green	
References: ²⁰⁴		Tunnel Length Bottleneck Role	

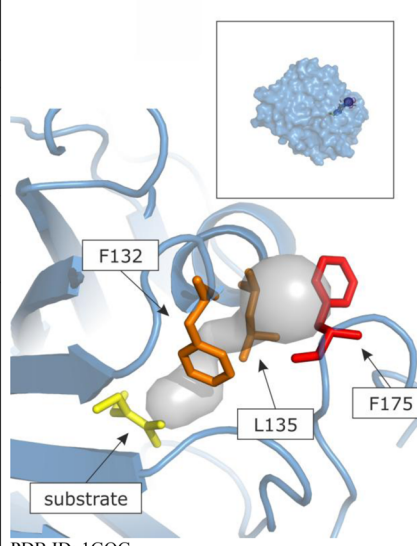
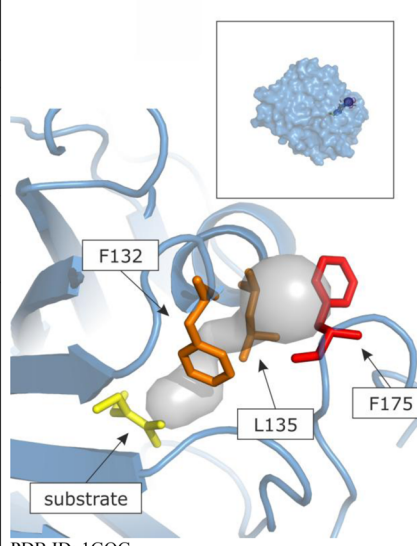
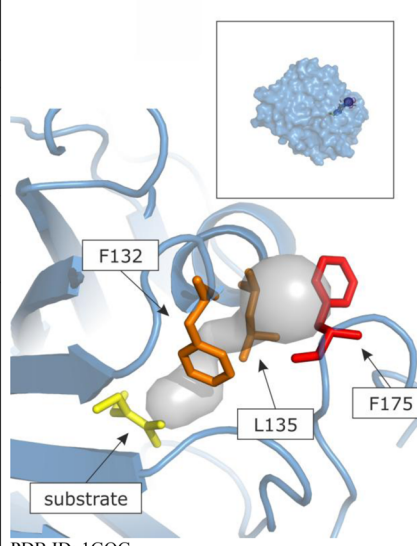
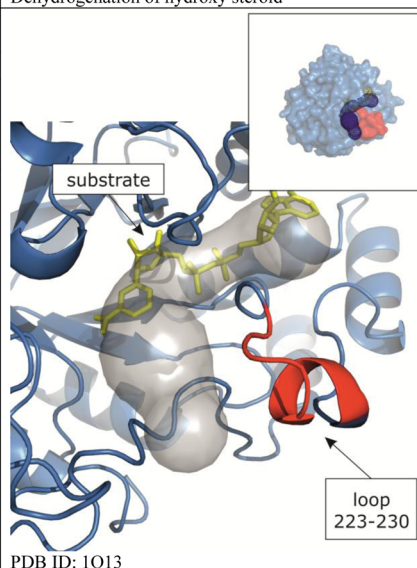
61	Quercetin 2,3-Dioxygenase		1.13.11.24
Gate function GATE1 – Controls access of small molecules into the active site GATE2 – Controls access of small molecules into the active site		Enzyme function Degradation pathway of flavonoids	
Gate location GATE1 – Tunnel connecting the bulk solvent and the active site cavity GATE2 – Tunnel connecting the bulk solvent and the active site cavity			
Gate structural basis			
Gate engineering			
PDB: WT – 1GQG, 1JUH, 1HII		PDB ID: 1GQG Tunnel Length 19 Bottleneck 1.6 Role O ₂ transport	
References: ⁷⁵			

Table 3. continued

62	Rabbit 20a-Hydroxysteroid Dehydrogenase	1.1.1.149
Gate function GATE1 – Protects cavity from the solvent in the absence of a steroid		Enzyme function Dehydrogenation of hydroxy steroid
Gate location GATE1 – Loops making-up the active site cavity		
Gate structural basis		PDB ID: 1Q13
 GATE1 Flexible loop B (residues 223–230) Movement of the loop partly controlled by the nature of Q230 Loops A and C mostly contribute in the pocket creation changing depth and size of the cavity		Tunnel
Gate engineering E230P – changes in the selectivity by reduced loop flexibility K274R – increases flexibility of loop by and eliminating K274–E227 contact, high k_{cat} for NADPH-dependent reduction of xylose		Length
PDB: WT – 1Q5M, 1Q13, WT (AKR5H1) – 2WZT 2WZM small loops		Bottleneck
References: ^{37,205–208}		Role
		Substrate steroid binding and orienting towards cofactor NADP(H)

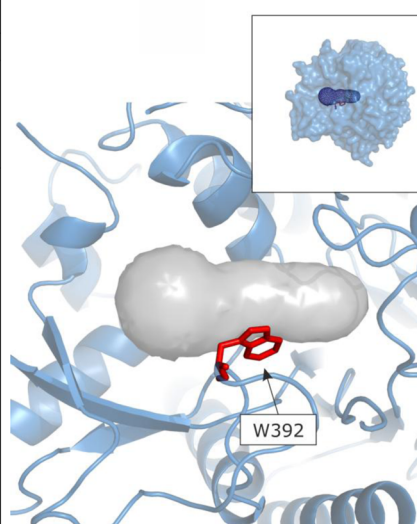
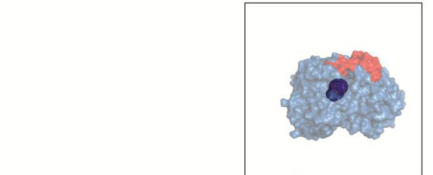
63	Raucaffricine O- β -Dglucosidaseglucosidase	3.2.1.125
Gate function GATE1 – Controls the access of the substrate		Enzyme function Hydrolyses alkaloid raucaffricine to aglycone vomilenine
Gate location GATE1 – At the entrance to the active site cavity		
Gate structural basis		PDB ID: 3U5U
 GATE1 W392 Conformation of W392 is controlled by S390		Tunnel
Gate engineering S390G – leads to more flexible conformation of W392 F485W – results in more fixed conformation of W392		Length
PDB: WT – 4A3Y, Mutants E186Q – 3U5U, 3U57, 3U5Y		Bottleneck
References: ²⁰⁹		Role

Table 3. continued

64	RNA-dependent RNA polymerase		2.7.7.48
Gate function GATE1 – Controls enzyme activity			
Gate location GATE1 – Surface loop			
Gate structural basis			
	GATE1 Δ1-loop + L30 Δ1-loop is involved in keeping the enzyme in a closed conformation, makes interaction with L30 (closed conformation is active, open is inactive)		
Gate engineering L30R and L30S – open conformation R222E – exhibited 2-fold reduction in activity, positively charged residue inside the tunnel K151E – activity was 7–10-fold higher at 21 °C, and 2–3-fold higher at 37 °C, positively charged residue near the tunnel entrance			 PDB ID: 2XWH
PDB: WT – 2XWH, 2XXD, 3MWV	Tunnel		
References: ²¹⁰	Length	19	
	Bottleneck		
	Role	NTP transport	

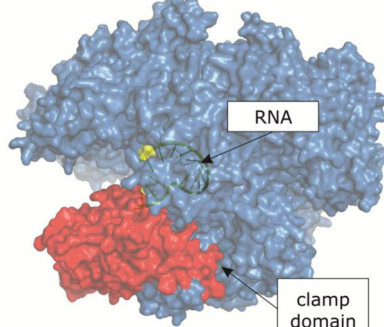
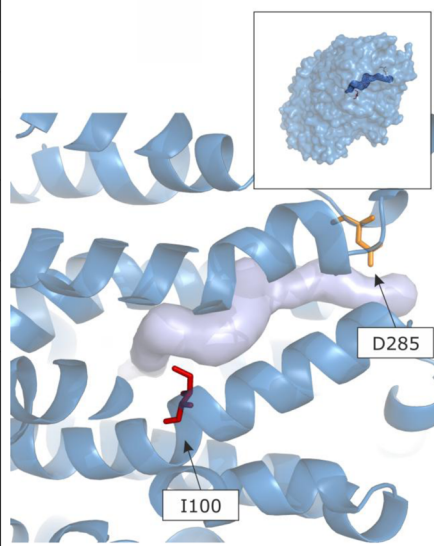
65	RNA Polymerase		2.7.7.6
Gate function GATE1 – Controls the access to the DNA-binding tunnel			
Gate location GATE1 – Entrance to the DNA binding tunnel			
Gate structural basis			
	GATE1 Clamp domain may open to permit entry of promoter DNA during initiation, close to establish the tight grip on DNA during elongation and then open again to allow release of DNA during termination		
Gate engineering			 PDB ID: 1I6H
PDB: WT – 2O5J, 1I6H, 1I6V	Tunnel		
References: ^{66,211}	Length		
	Bottleneck		
	Role		

Table 3. continued

66	Toluene-4-Monoxygenase T4MO	1.14.13
Gate function		
GATE1 – Controls access to the active site, controls selectivity		
GATE2 – Controls access to the tunnel		
GATE3 – Protects cavity from access of the solvent		
Gate location		
GATE1 – Boundary between the active site pocket and the tunnel		
GATE2 – Tunnel entrance		
GATE3 – Cavity entrance		
Gate structural basis		
—●	GATE1 I100 Side chain rotation	
—●	GATE2 D285 Steric blocking by negatively charged residue D285	
●—●	GATE3 R60, E64, E75, R69 Interactions between E64 and R60 and between E75 and R69, ionic gate for the cavity	
Gate engineering		
D285I, D285Q – improves oxidation of bulky substrates (11-fold)		
D285S – improves oxidation of styrene (1.7-fold)		
I100A – improves oxidation of bulky substrates (35-fold)		
I100G – improves oxidation of methyl-p-tolyl sulphide (11-fold)		
PDB: WT – 3DHG, WT (T4moC) – 1VM9, WT (T4moD) – 1G10		
References: ^{61,105}		
Enzyme function		
Hydroxylates toluene primarily at the para position to cresol		
		
PDB ID: 3DHG		
Tunnel		
Length		
Bottleneck		
Role	Substrate binding cavity	

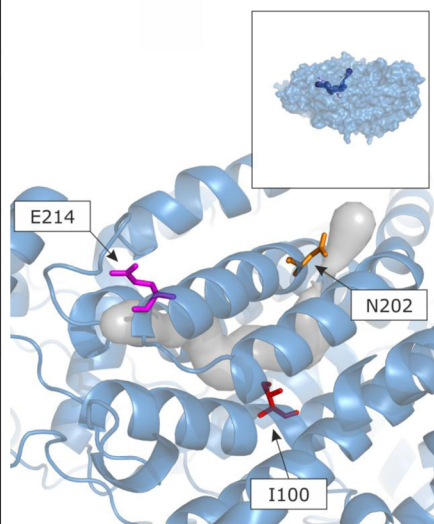
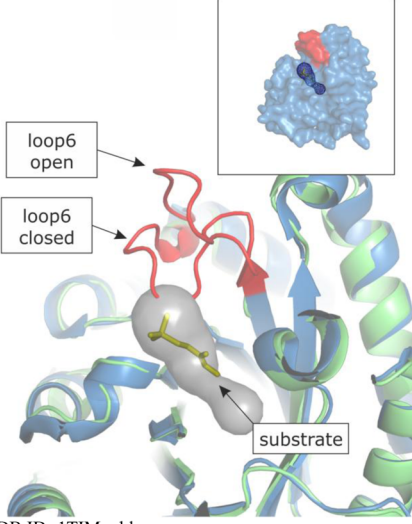
67	Toluene-Ortho-Xylene Monooxygenase ToMO	1.14
Gate function		
GATE1 – Controls access to the active site, controls selectivity		
GATE2 – Open pore allows access to the diiron centre for substrate (maybe also dioxygen or protons)		
Gate location		
GATE1 – Border of the pocket near Fe atoms – entrance to active site		
GATE2 – Entrance to the tunnel		
Gate structural basis		
—●	GATE1 I100 Residues H96, I100, T201, F205, and F196 border the pocket – I100 is an analogue of L110 (gate) in MMOH Ile100 contributes to defining the hypothetical para site, but it is also at the boundary between the active site pocket and the tunnel which connects the pocket to the surface of the protein	
—●	GATE2 N202 The conserved tunnel is gated by N202. The residue shifts in a redox-dependent manner. Its side chain is oriented away from the active site in the oxidized form and points inward in the reduced or Mn(II) reconstituted forms of the hydroxylase. This motion correlates with the carboxylate shift that occurs upon reduction of the dimetallic centre.	
Gate engineering		
I100A, I100W, F205L, A107V, A107I – decreased or removes activity		
Creation of new GATE3		
E214 – hot spot for new gate creation localized on helix E at the entrance to the tunnel 23 Å from the active site, mutation of those residue regulates access to the tunnel (E214A, E214G, E214V, E214W, E214F, E214Q, E214P) E214G – 15-fold improvement for p-nitrophenol oxidation		
PDB: WT – 2INC, 1T0Q		
References: ^{74,91,101,195,212,213}		
Enzyme function		
Catalyses hydroxylation of aromatics, oxidize benzene to phenol, catechol and trihydroxybenzene		
		
PDB ID: 2INC		
Tunnel		
Length	30-35	
Bottleneck		
Role	Substrate and product transport	

Table 3. continued

68	Triosephosphate Isomerase		5.3.1.1
Gate function GATE1 – Controls access of the water to the active site		Enzyme function Catalyses isomerisation D-glyceraldehyde 3-phosphate to dihydroxyacetone phosphate	
Gate location GATE1 – Entrance to active site			
Gate structural basis			
	GATE1 Loop 6 (residues 166 to 176). Residues 169–173 move as a rigid body, which position is controlled by flexible three residues N-terminal and C-terminal hinge regions In closed conformation loop sequesters the enzyme reaction from solvent. Closed conformation is stabilised by conserved anchoring residues from loop 7	 PDB ID: 1TIM – blue PDB ID: 1TPH – light green	
Gate engineering P(166)VW-AIGTG-KTA to P(166)GG-AIGTG-GGG mutant – k_{cat} decreases 2500-fold Y208F – k_{cat} decreases 2400-fold Y208T – k_{cat} decreases 200-fold			
PDB: WT – 1TIM (open), 1TPH (closed)			
References: ^{77,78,214–216}			

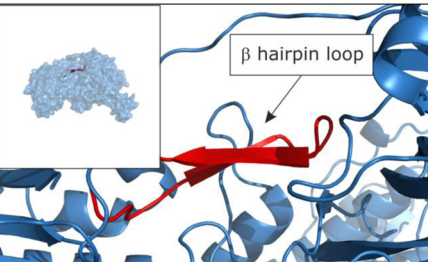
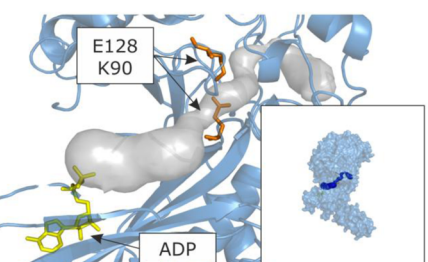
69	tRNA-Dependent Amidotransferase GatDE and GatCAB		6.3.5
Gate function GATE1 – Protects active site from access of the solvent, avoids wasteful release of ammonia into solution ?GATE2 – Controls ammonia transport		Enzyme function Conversion of Glu-tRNA ^{Gln} into Gln-tRNA ^{Gln} or Asp-tRNA ^{Asn} into Asn-tRNA ^{Asn}	
Gate location GATE1 – β hairpin loop surrounding asparaginase active site (in GatDE) ?GATE2 – Inside ammonia tunnel (in GatCAB)			
Gate structural basis			
	GATE1 GatDE β hairpin loop D β 7-D β 8, residues 100–118 in GatD, G100, the first glycine in the strictly conserved GGT motif, active conformation substrate in contact with T102	 PDB ID: 1ZQ1 GatDE PDB ID: 3H0L GatCAB	
	GATE2 GatCAB in <i>A. aeolicus</i> (<i>S. aureus</i>) E128 (E125), K90 (K88) E128 (E125) side chain can blocks the tunnel via a salt bridge with K90 (K88)	 PDB ID: 1ZQ1 GatDE PDB ID: 3H0L GatCAB	
Gate engineering			
PDB: WT (GatDE) – 1ZQ1, WT (GatCAB) – 3H0L, 3H0M, 3H0R			
References: ^{155,217–219}			
		Tunnel	GatDE
		Length	35
		Bottleneck	
		Role	Ammonia transport

Table 3. continued

70	Tryptophan Synthase		4.2.1.20
Gate function		Enzyme function	
GATE1 – Creates the tunnel and protects substrate binding cavity from the access of the solvent GATE2 – Controls transport of the indol GATE3 – Controls access to the subunit β and activates the β active site		Synthesis of L-tryptophan; subunit α cleavage of 3-indole-D-glycerol 3-phosphate; subunit β pyridoxal phosphate dependent condensation of indole with L-serine	
Gate location			
GATE1 – Surface loop from subunit α at the border between the subunits β and α GATE2 – Tunnel wall in subunit β GATE3 – Subunit β			
Gate structural basis			
	GATE1 Loop L177 - A190 A disordered surface loop in the subunit α after substrate binding becomes ordered and clamps down over the active site, isolating this region from solvent		
	GATE2 F280, Y279 The residue F280 can adopt alternative conformations in closed interact with C170 in open one it adopts Y279 position, Y279 moves toward the subunit α and interacts with part of the flexible loop-2 residues (54-61) of the subunit α		
	GATE3 D305 In the closed conformation D305 creates the H-bonded salt bridge with R141 with the associated H-bonding network involving S197 and S199		
Conformation changes are initiated by binding IGP - the allosteric communication between the two sites that results in full coupling of the reaction at the subunits α and β			
Gate engineering			
E49F, G51L, D60Y – decreases activity (2-fold) F280C, F280S – increases transport C170W, C170F – chemical modification of C170 (C170-NEM, C170-MMTS), obstructs the tunnel and accumulates indole intermediate			
PDB: WT – closed 3CEP, open conformations: 1BKS, 1KFK, 1KFJ, 1TTP, 2CLL, 2CLM, 2CLO		PDB ID: 1BKS	
References: ^{47,71,102,220,221}		Tunnel	
		Length 25 – 30	
		Bottleneck	
		Role Indol transport between two active sites	

71	Undecaprenyl-Pyrophosphate Synthase		2.5.1.31
Gate function		Enzyme function	
GATE1 – Controls length of the final product		Condensation reactions of isopentenyl pyrophosphate with allylic pyrophosphate to generate linear isoprenyl polymers	
Gate location			
GATE1 – Loop closing the entrance to the tunnel			
Gate structural basis			
	GATE1 71 – 83 loop The helix α 3 in open tunnel conformation is kinked by $\sim 30^\circ$ at E96, and the helix α 3 in closed tunnel conformation is kinked by $\sim 45^\circ$ at A92. Reaction starts with closed conformation \rightarrow when the chain length reaches C55, the dimethyl end is stopped at the end of the tunnel \rightarrow α 3 helix move away from the closed position into the open position (71 – 83 loop) \rightarrow fully synthesized C55 long product can exit easily through the open gate		
Gate engineering			
*GATE2 L137A – removes the floor of the tunnel and allows formation of a longer chain length products (normal length C55), bulky side chain of L137 serves to block further elongation of undecaprenyl-pyrophosphate			
A69L – results in long lived accumulation of a short chain intermediate C30 final product C55 A143V – similar to wild type, rate 3-fold lower			
S71A, N74A, or R77A – decreases in k_{cat} values (25–200-fold) W75A – increases in K_m for farnesyl pyrophosphate (8-fold) E81A, S71A – increases in K_m for isopentenyl pyrophosphate (22–33-fold)			
PDB: WT – 1JP3		PDB ID: 1JP3	
References: ²⁷		Tunnel	
		Length 30	
		Bottleneck	
		Role Active site cavity	

substrate binding since it is believed to slow down product dissociation.^{22,97} Other gates of this type have been reported in 3-hydroxybenzoate hydroxylase,⁹⁸ 4-hydroxy-2-ketovalerate aldolase/acylating acetaldehyde dehydrogenase,⁹⁶ and chole-

terol oxidase type I⁵⁷ and type II.⁹⁹ Moreover, cofactors themselves can also function as gates. The FAD cofactor of digeranylgeranylglycerophospholipid reductase has two different conformations, referred to as the “in” and “out”

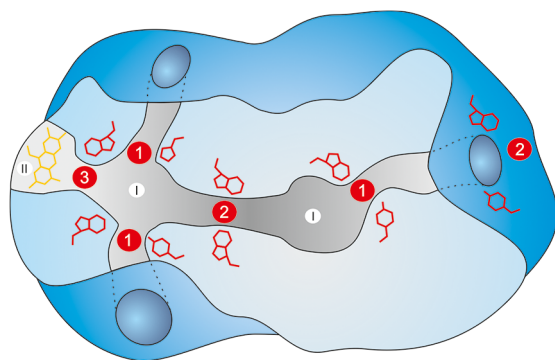


Figure 6. Locations of gates within a protein structure. Schematic representation of an enzyme with two active sites connected by a tunnel (I), a cofactor cavity (II), and multiple access tunnels. Gating residues in red may be located at the entrance to the active site (1), at the entrance or the bottleneck of the tunnel (2), and between the active site cavity and the cofactor cavity (3).

conformations. In the “in” conformation, the tunnel is blocked by FAD. To open the tunnel, FAD has to adopt the “out” conformation.¹⁰⁰

Gates are most commonly located at the tunnel entry and the tunnel bottleneck (51%). This is to be expected because the bottleneck represents the narrowest point of the tunnel, and its diameter often dictates the tunnel’s permeability. Another common location is the entrance to the active site cavity (28%). Gates at the entrance of the cofactor cavity are less common (5%), which is not surprising since not all enzymes have a cofactor cavity. In 16% of the cases studied, the gate was not located within any of these functional regions (Figure 7).

5. ENGINEERING OF GATES

The average rate of evolution of the gating residues in the cytochrome P450 family is significantly greater than that for the protein sequence as a whole.⁵⁹ The high rate of evolution at the gating residues suggests that gate engineering may be an attractive alternative to other rational enzyme design strategies. This idea is supported by a few observations: (i) the gates are often spatially separated from the active site, and so mutations at the gating residues should not be deleterious to protein function, (ii) the opening and closing of the access pathways can affect ligand exchange and thus enzyme activity and selectivity, and (iii) gate modification can modulate the solvent’s ability to access the active site, which in turn affects

solvation and stabilization of the transition state and also product release. In the following section, we describe selected cases in which an enzyme’s catalytic properties have been successfully altered by modifying its gates.

5.1. Gate Modification

Gates can be modified by substitution of the gating residues, hinge residues, or anchoring residues depending on the nature of the gate in question. Modification of the gating residues has been shown to change the selectivity and activity of toluene-*o*-xylene monooxygenase, with the E214G mutation improving oxidation of *p*-nitrophenol by a factor of 15.¹⁰¹ A similar improvement in overall activity was achieved in a lipase from *Burkholderia cepacia* by the mutations L17S + L28T. This double mutant also exhibited a 10-fold increase in enantioselectivity compared to the wild-type enzyme.⁸⁶ The T78F or P76F mutations in imidazole glycerol phosphate synthase override some of the control exerted by the wild-type gate and block the passage of ammonia through the tunnel.³⁸ In NiFe hydrogenases, mutations of the V74 and V74 + L122 residues changes the rates of transport for H₂, CO, and O₂, thereby modulating the overall rate of reaction.²¹

Gates that incorporate secondary structure elements are dependent on hinge and anchoring regions. It has been demonstrated that modification of the hinge region can change an enzyme’s activity and selectivity. Notably, the Q230P mutation in rabbit 20A-hydroxysteroid dehydrogenase decreases the flexibility of a key loop and thereby changes its selectivity.³⁷ Similarly, access to the active site in the HIV-1 protease is controlled by two β-turn flaps.⁸⁹ Results from Brownian dynamics simulations suggest that the G48V + V82A or I84V + L90M mutations in this enzyme reduce the likelihood that the active site will be exposed at any given point in time from 14% in the wild type to 2% in the mutants.⁸⁹ The importance of the anchoring residues in the HIV-1 protease was demonstrated by the F53L mutation, which generates a semiopen conformation due to removal of the stabilizing F53–I50 interaction.⁸⁹

5.2. Gate Removal

Removing gates typically increases substrate and product exchange rates but also allows more extensive access of water molecules to protein tunnels and cavities. The overall effect of gate removal is therefore equal to the combined effects of these two processes. Gate deletion has been shown to increase the rate of substrate binding to tryptophan synthase.¹⁰² The F280C and F280S mutations both increased the rate of indole binding

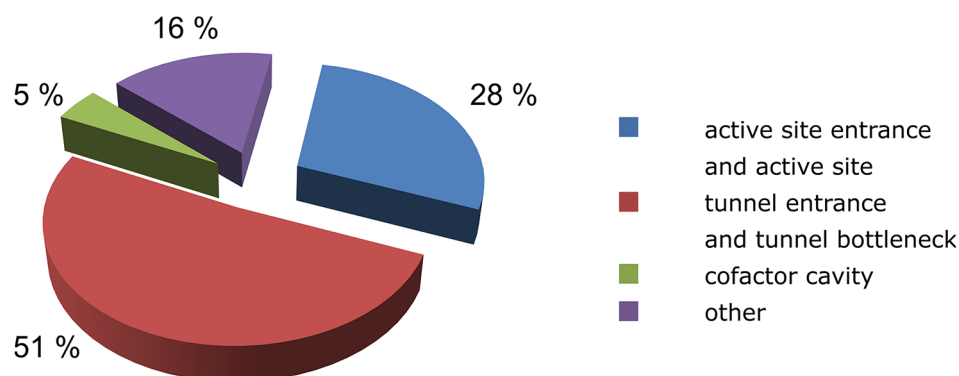


Figure 7. Distribution of gate locations within protein structures based on analysis of 71 proteins with 129 gates. Detailed description of the analyzed proteins is provided in Table 3.

by a factor of 2. Similarly, the T78A mutation allowed ammonia to rapidly pass through the tunnel in imidazole glycerol phosphate synthase.³⁸ The R239A mutation in the cyclase caused a 1000-fold decrease in the enzyme's k_{cat}/K_m value and decoupling of the reaction.³⁸ This dramatic change in enzyme catalytic efficiency was attributed to creation of a new route for ammonia release. A similar leakage of ammonia was caused by the G359F and G359Y mutations in carbamoyl phosphate synthetase.^{103,104} Negative consequences of gate removal were also observed for the FabZ- β -hydroxyacyl-acyl carrier protein dehydratase (HpFabZ),⁵⁴ in which the Y100A mutation leaves the active site completely exposed to the bulk solvent. As a result, the acyl carrier protein binds to the HpFabZ Y100A mutant much more strongly than to the wild-type HpFabZ, decreasing the mutant enzyme's activity by more than 50% due to the very slow dissociation of the acyl carrier protein.

In some cases, gate removal enables bulky substrates to access the active site cavity. Mutations D285I and D285Q in toluene-4-monooxygenase improved its ability to oxidize the large and bulky substrates 2-phenylethanol and methyl *p*-tolyl sulfide by factors of 8 and 11, respectively, while the D285S mutation improved the rate of styrene oxidation 1.7-fold.¹⁰⁵ The L137A mutation in undecaprenyl-pyrophosphate synthase removed the bottom of the tunnel in this enzyme, allowing formation of products with longer chain lengths.²⁷ A similar situation was encountered in type III polyketide synthases from *Aloe arborescens* PCS, in which the M207 residue controls the number of condensations of malonyl-CoA.¹⁰⁶ The M207G mutation opened a connection between the tunnel and two hidden pockets located behind the active site, resulting in formation of extended products. Further product elongation was achieved with the triple mutant F80A + Y82A + M207G.

5.3. Gate Insertion

To best of our knowledge, there have been no reports of an intentional introduction of a new gate into an enzyme structure. However, there have been studies in which an access tunnel was systematically modified with multiple substitutions, and it is reasonable to expect that some of these mutations might have created new gates. More research will clearly be needed to confirm this expectation. To verify successful intentional insertion of a new gate into a protein structure, it would be necessary to confirm the existence of both the open and the closed conformations at a position where previously only a single conformation could be adopted.

Site-directed mutagenesis targeting specific residues at various positions along the access tunnel of *Candida rugosa* lipase has been used to alter the acceptable substrate chain length for this enzyme. The mutants for which this was observed were P246F, L413F, L410W, L410F + S300E, and L410F + S36SL.¹⁰⁷ We note that the aromatic residues F and W, which are common in wing and swinging door gates, were introduced in each of these variants.

In another study, the residue L177 that is located near the entrance to the access tunnel of the haloalkane dehalogenase LinB from *Sphingobium japonicum* UT26 was substituted with all of the natural amino acids, yielding 19 mutants with significantly altered substrate specificity and activity.¹⁰⁸ Preliminary computational analyses of these variants using molecular dynamics revealed that the two residues possessing a single aromatic ring (F and Y) exhibited large fluctuations, as might be expected for gating.

Residue A217 is located at the entrance to the tunnel in the epoxide hydrolase EH from *Aspergillus niger* M200. This residue was substituted with C, E, G, L, P, Q, R, T, and V, and the effect of each mutation on the enzyme's activity and enantioselectivity was studied.²⁶ The mutants exhibited different enantioselectivity and activity relative to the wild type. For instance, the activity of the A217G mutation toward allyl glycidyl ether was lower than that of the wild type by a factor of 33, whereas the A217V mutation increased activity toward this substrate 6.6-fold.

Residues I135, W141, C176, V245, L246, and Y273 are positioned close to the entrance to the main and side tunnels of the haloalkane dehalogenase DhaA from *Rhodococcus rhodochrous* NCIMB 13064. These residues were simultaneously permuted in an attempt to improve this enzyme's activity against 1,2,3-trichloropropane. The most successful mutant, which featured the I135F, C176Y, V245F, L246I, and Y273F substitutions, showed 26-fold greater activity toward the target substrate than did the wild type.¹⁰⁹ In this mutant, three aromatic residues were introduced in place of aliphatic ones in the vicinity of the tunnels. Computational analysis of product release from the mutant suggests that substitutions introduced a transient rather than permanent structural feature and gating residues prevented access of water to the active site.¹¹⁰ Crystallographic analysis of the mutant revealed two distinct conformations for the Y176 side chain.¹¹⁰

6. CONCLUSIONS

This review highlights the importance of gates in enzymes. Gates play vital roles in controlling the catalytic activity and selectivity of enzymes and are more common in protein structures than is generally thought. In particular, gates control substrate access to the active site and product release, prevent or restrict solvent access to specific regions of the protein, and can synchronize processes occurring in distinct parts of the enzyme. Our literature survey of 129 gates in 71 enzymes revealed a large variety of systems with sophisticated structures. We presented a rigorous definition of gates and established a new scheme for their classification. The large number of inspected cases allowed us to build a catalogue of gates assigned to six distinct classes—wings, swinging doors, apertures, drawbridges, double drawbridges, and shells—with three different functions and three distinguishable locations. We also presented summary statistics that give a preliminary overview of the propensity of specific amino acid residues to occur in particular gate classes. The proposed classification scheme can be easily extended and updated but even in its present form can provide guidance for analysis and engineering of gates in biomolecular systems.

The biochemical relevance and specific location of gates within protein structures make them attractive targets for protein engineering. Attempts to rationally redesign gates typically involve computer-assisted gate identification followed by modification using focused directed evolution. This approach is compatible with a recent trend in protein engineering that stresses construction of small and smart libraries. Gate modification and deletion have been demonstrated in numerous cases, but the intentional insertion of new gates remains a challenge. Convenient methods for identifying gates in protein structures are essential prerequisites for their engineering. In silico, this can be achieved by coupling the software tools developed for describing pathways to tools developed for study of protein dynamics. Of the available

experimental techniques, NMR spectroscopy is particularly suitable for analysis of highly dynamic protein structures and can be expected to play an indispensable role in the study of gate dynamics at the atomic level. The field would also benefit from development of new experimental techniques for monitoring the passage of ligands through the protein pathways. One day it will be possible to control the catalytic properties of enzymes by rational engineering of their gates. To achieve this goal, we have to learn how gates evolved, how they interact with the other parts of the protein structure as well as with the ligand and solvent molecules, and how they fulfill their biological functions.

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