X-ray and neutron crystallography are powerful techniques utilized to study the structures of biomolecules. Visualization of enzymes in complex with substrate/product and the capture of intermediate states can be related to activity to facilitate understanding of the catalytic mechanism. Subsequent analysis of small molecule binding within the enzyme active site provides insight into mechanisms of inhibition, supporting the design of novel inhibitors using a structure-guided approach. The first X-ray crystal structures were determined for small, ubiquitous enzymes such as carbonic anhydrase (CA). CAs are a family of zinc metalloenzymes that catalyze the hydration of CO$_2$, producing HCO$_3^-$ and a proton. The CA structure and ping-pong mechanism have been extensively studied and are well understood. Though the function of CA plays an important role in a variety of physiological functions, CA has also been associated with diseases such as glaucoma, edema, epilepsy, obesity, and cancer and is therefore recognized as a drug target. In this review, a brief history of crystallography and its impact on CA research is discussed.

1. Introduction

1.1. A Brief History of Crystallography. Röntgen discovered a form of radiation in 1895 while analyzing the range of cathode rays in vacuum tubes. He termed this radiation X-rays and determined that the permeability of an object to such radiation directly correlates to its density [1]. Laue hypothesized that radiation of short wavelengths, such as X-rays, would diffract when passed through a crystal if the wavelength is of similar magnitude to the distance between planes of the crystal lattice. Friedrich and Knipping confirmed Laue’s hypothesis and successfully demonstrated diffraction from crystals of copper sulfate in 1912 [2]. Based on their findings, Bragg recognized that the treatment of a diffraction pattern as reflections from parallel planes within a crystalline lattice could be used to relate the angle of the incident beam to the wavelength and distance between the planes, now known as Bragg’s Law [3]. Subsequent experiments led to Bragg’s development of the first X-ray spectrometer and determination of the crystal structure of sodium chloride [4, 5].

The growth of protein crystals dates back as early as 1840 with the observation of hemoglobin crystals in blood samples [6]. However, the first crystal of an enzyme, urease, was not achieved until 1926 [7]. X-ray diffraction of a pepsin protein crystal was collected in 1934 following the optimization of conditions to ensure hydration of the crystal during data collection. However, the pepsin structure was not determined for several more decades [8]. The first protein crystal structures determined include myoglobin in 1957 and hemoglobin in 1960 [9, 10].

Nearly a decade after the first observation of X-ray diffraction, interest in obtaining neutron diffraction from single crystals increased. X-ray crystallography requires electrons in the sample to interact with the incoming X-ray beam to generate a diffraction pattern. The scattering factor of an atom is the likelihood of a diffraction event occurring and is dependent on how many electrons are in the atom. Electron rich atoms have a high scattering factor, meaning they are easily distinguishable from the diffraction pattern [11]. Therefore, a resulting limitation of X-ray crystallography is the inability to see hydrogens due to an inherently low scattering factor. However, neutron scattering lengths of Hydrogen and Deuterium are comparable to other atoms. Neutron crystallography can therefore be used to identify the position and accessibility of H atoms, providing insight into side chain protonation states and hydrogen bonding networks that may improve the understanding of catalytic
mechanisms [12–14]. Early experiments were performed at the Argonne and Clinton Laboratories, sources used to collect data for the Manhattan Project. Successful neutron diffraction was collected from calcite crystals in 1944 and NaCl crystals in 1945 [15, 16]. The first single crystal neutron structures were determined in 1951 [17].

As interest in protein structure and function continued to grow, the need for a database of crystal structures led to the development of the Protein Data Bank (PDB, rcsb.org) in 1971. The PDB was started with 7 depositions, including the structures of myoglobin and hemoglobin, and has grown to over 125,000 structures to date [18]. This statistic highlights the importance of X-ray and neutron crystallography as techniques for understanding enzyme structure in relation to catalytic mechanism and the use of such knowledge to guide drug design.

1.2. Discovery of Carbonic Anhydrases. In the 1920s, two theories were proposed concerning the transport of CO₂ in the blood. The most common hypothesis was termed the HCO₃⁻ theory which stated that CO₂ is transported to the lungs in the form of HCO₃⁻. The HCO₃⁻ is then converted to carbonic acid by proteins in the blood, which dehydrates to release CO₂. However, the rate of spontaneous carbonic acid dehydration was calculated to be significantly lower than the observed physiological rate of CO₂ respiration [19]. The rate of CO₂ production was therefore proposed to be increased by a catalyst present in red blood cells [20]. This catalyst was isolated from ox blood in 1932, determined to be distinct from hemoglobin, and given the name carbonic anhydrase (CA) [21]. Thus, it was concluded that CA catalyzes the reversible hydration of CO₂ to produce HCO₃⁻ and a proton.

\[
\text{CO}_2 + \text{H}_2\text{O} \rightleftharpoons \text{HCO}_3^- + \text{H}^+ \quad (1)
\]

Initial experiments identified two distinct forms of CA (CA B and CA C) [22]. CA B was discovered to be ~7-fold more abundant whereas CA C exhibited higher catalytic activity [23]. Fractionation experiments identified a third form, CA A, that was originally thought to be an artifact of purification due to the similarities of amino acid composition between CA A and CA B [24]. The nomenclature of CA A, CA B, and CA C were reassigned as CA III, CA I, and CA II, respectively.

1.3. CAs in the Human Genome. CA I, CA II, and CA III were confirmed as unique CA isoforms through sequence comparison and gene mapping [23, 25–28]. The three genes were mapped onto a cluster of genes on chromosome 8 in the order CA II, CA III, and CA I [29]. With the completion of the human genome, a total of 15 CA isoforms have now been identified [30]. These CA isozymes differ by both cellular localization and catalytic efficiency. CA I, CA II, CA III, CA VII, CA VIII, CA X, CA XI, and CA XIII are cytosolic; CA IV, CA IX, CA XII, and CA XIV are membrane-bound; CA Va and CA Vb are mitochondrial; and CA VI is secreted [31].

1.4. CA Families. Since the discovery of mammalian CAs in 1932, 7 genetically unique families have been identified: α, β, γ, δ, ζ, η, and θ [21, 32–37]. The CA families all catalyze the reversible hydration of CO₂ and are therefore classified by differences in structural fold. Furthermore, the tetrahedral coordination of a metal ion to three active site residues and a water molecule is conserved between the classes, but the identity of the metal ion and amino acids differs. The α CAs are the most well-studied class as these are expressed in mammals, though examples have been identified in bacteria and protozoa as well. The α CAs have a catalytic zinc coordinated by three His residues (x, x+2, x+25, where x=94 in human CA II). The activity and/or overexpression of several isoforms in the α class have been associated with human diseases and are therefore recognized as therapeutic targets.

β CAs were discovered soon after the α class in 1939. The β CAs were isolated from plant chloroplasts and have since been identified in eubacteria, algae, and archaea. β CAs are dimeric in nature and are the only CA class to exhibit allosteric regulation. β CAs are also zinc metalloenzymes; however the zinc is coordinated by two Cys and one His in the active site (x, x+56, x+59, where x=68 in Cam2) [38]. Because the catalytic activity of β CAs has been shown to be essential to the survival of bacteria and this class is structurally unique from the CAs expressed in humans, β CAs are targeted in the design of new antibiotics [39].

γ CAs have been found in archaea and bacteria. They are catalytically active as trimers with the metal ion coordinated between the monomers by a His residue from each monomer (x, x+36, x+41, where x=81 in Cam) [40]. Previous studies have hypothesized iron as the physiologically relevant metal ion [41]. However, the enzyme is most active when bound to cobalt and maintains activity in coordination with zinc [42].

The δ and ζ classes of CA are both expressed in marine diatoms. δ CAs were first identified in *Thalassiosira weissflogii* in 1997 [43]. The active site structure of δ CAs is hypothesized to be similar to that of the α class with a catalytic zinc ion coordinated by three His residues (x, x+3, x+12, where x=114 in TweCA) [32, 44]. ζ maintains activity with cadmium or zinc bound, though is expected to preferentially bind cadmium in seawater with low concentrations of zinc. The ζ CA active site resembles that of β CAs coordinating the metal to a His and two Cys residues (x, x+52, x+62, where x=263 in CDCα1) [45].

The η and θ CAs represent the most recently identified families and therefore little information is known about these classes. The η CAs are closely related to and previously thought to belong to the α class. It was not until 2014 that sequence and phylogenetic analyses identified the η CAs as a unique genetic class [33]. η CAs are also unique in that the catalytic zinc is hypothesized to be coordinated by two His and one Gln [46]. To date, η CA has only been identified in the Plasmodia species of protozoa, the pathogen that causes malaria, and is therefore recognized as a potential target of antiparasitic agents. θ CA was isolated from *Phaeodactylum tricornutum* in 2016 and identified as a CA zinc metalloenzyme due to its CO₂ hydration and esterase activity [34].

1.5. CA Crystal Structures. To date, crystal structures have only been determined for the α, β, γ, and ζ CA families. There
are over 900 CA structures deposited in the PDB that have
been solved using X-ray or neutron crystallography (Figures
1 and 2). The majority of these structures represent the α
class, including structures for all the catalytically active human
isoforms with the exception of CA V (Figure 2, Table 1) [47–
56]. A large number of structures also include α CAs in
complex with various classes of inhibitors, highlighting the
emphasis placed on the α CAs as drug targets.

The first α CA crystal structure of CA II was determined
to 2.0 Å resolution by Liljas et al. in 1972 and further
refined by Eriksson et al. in 1988 to the space group P2₁
with cell dimensions \( a = 42.7, b = 41.7, c = 73.0 \) Å, and \( \beta = 104.6^\circ \). The overall shape of the
molecule is an ellipsoid with dimensions \( \sim 40 \times 40 \times 50 \) Å. The conically shaped active
site is \( \sim 15 \) Å deep with the catalytic zinc situated at the
base. Seven right-handed α-helices are present on the surface
of the enzyme, surrounding a 10 stranded \( \beta \)-sheet core.
The zinc is tetrahedrally coordinated to three His residues
and a water molecule (Figure 3(a)). Electron density was
observed for six ordered water molecules in the active site
that are stabilized via hydrogen bonding with hydrophilic
residues [57, 58]. Many CA II structures have since been
solved and recently sub-angstrom resolution was achieved,
providing further insight into the CA catalytic mechanism
[48].

The first crystal structure of a β CA isolated from
\( P.\) purpureum red algae was determined in 2000 [35]. Now,
nearly 80 structures have been deposited to the PDB repre-
senting plant, yeast, bacterial, and archaeal β CAs. The β CAs
form oligomers from dimeric fundamental units and most
commonly exist as tetramers with the tetrameric interface
perpendicular to that of the individual dimers. All β CA
monomers exhibit a four or five stranded parallel \( \beta \)-sheet
core. Upon dimerization, the \( \beta \)-sheets of each monomer align
to form an extended \( \beta \)-sheet. Several α-helices surround
the core, two of which interact with the other monomer to
support dimerization. A hydrophobic cleft along the dimer
interface leads to the catalytic zinc and is predicted to bind
substrate \( \text{CO}_2 \) (Figure 3(b)) [38]. There are two subclasses of
β CA characterized by the coordination of the zinc ion. In
Type I, zinc is coordinated by two Cys residues, one His, and
a water molecule/ligand. In Type II the coordination sphere
consists of two Cys, one His, and one Asp, which replaces
the water. While Type I are active under most pHS, Type II
are catalytic below pH 8.3. This is due to the Asp residue
preventing the catalytic hydroxyl from coordinating to the
zinc. At basic pHS above 8.3, the Asp residue is removed from
the zinc, allowing a water molecule to bind and activate the
enzyme [59]. This also results in the formation of an Asp-
Arg dyad, similar to the Type I structures, that serves as a
hydrogen bond acceptor of the ligand. As the pH reaches 8.3,
Type II enzymes undergo conformational changes to mimic
the active sites of Type I. A noncatalytic \( \text{HCO}_3^- \) binding
site containing a Trp-Arg-Tyr triad has also been observed
in several β CAs located approximately 8 Å from the zinc.
Figure 2: Human CA isoform structures: (a) CA I, (b) CA II, (c) CA III, (d) CA IV, (e) CAVI, (f) CA VII, (g) CA VIII, (h) CA IX, (i) CA XII, (j) CA XIII, and (k) CA XIV.
| # Structures in PDB | PDB ID | Resolution (Å) | Space group | Cell dimensions (Å, °) | Oligomeric state | % identity to CA II | Rmsd to CA II (Å) |
|---------------------|--------|----------------|-------------|------------------------|-----------------|-------------------|------------------|
| 24                  | 2CAB   | 2.0            | P2₁,2₁,2₁   | a = 81.5 b = 73.6 c = 37.1 β = 104.4 | Monomer         | 61                | 0.9              |
| 703                 | 3KS3   | 0.9            | P2₁         | a = 42.3 b = 41.4 c = 72.3 β = 104.4 | Monomer         | -                 | -                |
| 5                   | 1Z93   | 2.1            | P6₃         | a = 44.7 b = 44.7 c = 72.3 β = 104.4 | Monomer         | 58                | 0.8              |
| 10                  | 1ZNC   | 2.1            | C2          | a = 47.7 b = 44.7 c = 72.3 β = 104.4 | Monomer         | 34                | 1.3              |
| 1                   | 3FE4   | 2.8            | P₂₁,₂₁,₂₁   | a = 57.7 b = 80.0 c = 141.0 β = 90.0 | Monomer         | 35                | 1.4              |
| 2                   | 3MDZ   | 1.9            | C₂2₂₁      | a = 69.7 b = 97.7 c = 133.4 β = 109.0 | Monomer         | 56                | 0.6              |
| 6                   | 5DVX   | 2.3            | P₂₁,₂₁,₂₁   | a = 57.9 b = 102.7 c = 82.4 β = 109.0 | Monomer         | 34                | 1.4              |
| 16                  | 1JCZ   | 1.6            | C₂          | a = 146.7 b = 44.6 c = 85.2 β = 94.1 | Dimer           | 36                | 1.2              |
| 13                  | 3D0N   | 1.6            | P₂₁         | a = 57.8 b = 58.2 c = 72.1 β = 92.4 | Monomer         | 61                | 0.8              |
| 2                   | 4LU3   | 2.0            | P4₁,2₁,2₁  | a = 88.6 b = 88.6 c = 108.9 β = 92.4 | Monomer         | 37                | 1.2              |

To date, no crystal structures of CA VA, CAVB, CA X, or CA XI have been deposited in the PDB.
Occupancy of this site is hypothesized to restrict the Type II class from undergoing conformational change, inactivating the enzyme [60].

Fifteen γ CA crystal structures have been deposited in the PDB to date. These structures have all been determined from enzyme purified from *M. thermophila* and *P. horikoshii* of the archaea domain. γ CAs are active as homotrimers and each monomer is characterized by a left-handed parallel β-helix. The catalytic metal ion binds in the trimer interface and is coordinated to three His residues; H81 and H122 contributed by one monomer and H117 by the other (Figure 3(c)) [41]. This coordination geometry is distorted if zinc or cobalt is bound in place of the physiologically relevant iron. Dual conformations of E84 were identified in the electron density, suggesting a role in proton transfer [36].

Crystals structures of ζ CAs are limited to CDCA1. CDCA1 has three tandem repeats (R1-R3), each of which is enzymatically active and mimics the structure of a β CA dimer. Each repeat is ellipsoidal in shape with a total of 7 α-helices and 9 β-strands. The overall structure can be divided into two lobes with a funnel shaped cleft that leads to the active site. The metal ion is tetrahedrally coordinated to three conserved residues (C263, H315, and C325) and a water molecule (Figure 3(d)). A conserved Asp-Arg dyad (D265–R26) contributes to the hydrogen bonding network in the active site and is predicted to contribute to the catalytic mechanism. Interestingly, the apo-CDCA1 structure is stable and has a more open conformation than holo-CDCA1. Rotation of the Cys residues that are no longer coordinating a metal ion contributes to this structural change [37].

2. Mechanism

2.1. Ping-Pong Mechanism. CAs are some of the most catalytically efficient enzymes known with CA II exhibiting a turnover rate of 1.1 µs⁻¹ and the fastest, SazCA, a rate of 4.4 µs⁻¹ [61, 62]. CA II and all subsequent α CAs exhibit a classic two-step ping-pong mechanism where the first step is a nucleophilic attack of CO₂, generating bicarbonate. The second step is the regeneration of the catalytic zinc-bound hydroxyl [63].

\[
\text{ZnOH}^- + \text{CO}_2 \rightleftharpoons \text{Zn(OH)}^- \text{CO}_2 \rightleftharpoons \text{ZnHCO}_3^- = \\
\text{ZnH}_2\text{O} + \text{HCO}_3^- \\
\text{ZnH}_2\text{O} = \text{Zn(OH)}^- \text{H}^+ + \text{H}_6\text{O} = \\
\text{ZnOH}^- + \text{H}_6\text{O}^+ 
\]

The full mechanism was first reported in 1988 by Silverman and Lindskog where they described the interworking
of the α CAs through structure activity relationships and kinetic data (Figure 4) [63]. In the hydration direction, the reaction starts with an open active site, a primed zinc-bound hydroxyl, and an empty hydrophobic binding pocket (Figure 4(a)). CO$_2$ diffuses into the active site, binding in a hydrophobic pocket approximately 2.8 Å away from the catalytic zinc-bound hydroxyl [64]. As CO$_2$ binds, H64 initiates a conformational change from the “out” to the “in” position in preparation for proton transfer [65]. CO$_2$ binding triggers a nucleophilic attack from the free pair of electrons on the hydroxyl group onto the partially positive carbon of the CO$_2$ (Figure 4(b)) [66]. This forms HCO$_3^−$ anchored to zinc through the same nucleophilic oxygen that performed the attack (Figure 4(c)). HCO$_3^−$ is displaced/released via the binding of a water molecule in its place (Figure 4(d)). This first step is nearly instantaneous due to the efficient nucleophilic attack of the hydroxyl and quick on/off rate of the HCO$_3^−$. It therefore does not significantly contribute to the overall rate of the reaction [63].

The second, rate limiting step is the regeneration of the zinc-bound hydroxyl via proton transfer from the zinc-bound water to bulk solvent [67]. While HCO$_3^−$ is released, H64 completes its conformational change to fully occupy the “in” conformation [67]. An ordered water network in the active site acts as a proton wire, allowing intramolecular proton transfer [68]. This proton wire is made up of six water molecules (ZnH$_2$O/OH$^−$, DW, W1, W2, W3a, and W3b) that are stabilized through a hydrogen bonding network [68]. Multiple residues located in the hydrophilic region of the active site stabilize these waters, including Y7, N62, N67, T199, and T200 [69]. The zinc-bound water is deprotonated and the proton is moved 8 to 10 Å out of the active site through these clustered waters [69]. These hydrogen bond interactions result in rapid proton shuttling out of the active site and regeneration of the catalytic hydroxyl [68]. Proton transfer is a two-part process. The first being the shuttling of the proton through the six waters to H64. The second is the transfer of the proton between H64 to bulk solvent [70].

2.2. Substrate Binding. Due to the highly efficient turnover rate of CA II and the limited solubility of CO$_2$ in solution, the binding sites of substrates for CAs remained unknown until hypothesized via molecular dynamics (MD). Using free energy perturbation simulations, Liang and Lipscomb identified three potential CO$_2$ binding sites in the hydrophobic region of the active site. One site was identified as the potential catalytic site and two other binding locations likely sites for CO$_2$ replenishment [71]. The primary binding pocket was shown to have the highest predicted affinity ~4 Å from the catalytic zinc and interact with residues H119, V121, V143, L198, T199, V207, and W209. Additionally, the determined orientation of CO$_2$ in this site supported the mechanism of nucleophilic attack by the zinc-bound hydroxide. To test this proposed binding site, site directed mutagenesis was performed by Fierke et al. [72]. Residues within this hydrophobic pocket were mutated and subsequent kinetic tests were performed. A total of twelve V143 variants of varying size and charge were tested, all of which showed a decrease in catalytic rate. Large aromatic groups such as Phe and Tyr showed the largest impact on activity, resulting in 10$^5$ fold decreased rates [72]. Of note, the V143Y mutant decreased catalytic activity below 0.02% of the wild type, confirming that a large, sterically hindering residue in the hydrophobic pocket prevents CO$_2$ binding [72]. While convincing, the CO$_2$ binding site had yet to be confirmed via X-ray crystallography.

The first CA structures confirming the MD predictions were obtained via cryocooling holo- and apo-CA II under 15 atm of pressurized CO$_2$ [73, 74]. Technological advances at synchrotrons allowing data collections under high pressures in conjugation with acidifying of the crystal permitted these structures to be obtained. Under 15 atm of CO$_2$, the buffer surrounding the CA crystals becomes acidic, rendering the reaction inactive due to the zinc-bound water preventing nucleophilic attack. As predicted by the MD simulations, the hydrophobic pocket was determined to be the site of CO$_2$ binding, approximately 4 Å away from the zinc. This was also true of the apo-CA II. From these structures, it was apparent that T199 forms a weak hydrogen bond with CO$_2$ orienting it for optimal nucleophilic attack [73].

Similarly, the binding site of HCO$_3^−$ in CA II was unknown due to its weak binding affinity. Previously reported work did see success in cobalt substituted structures of CA II, however attempting crystal soaks with wild type CA II at high concentrations of HCO$_3^−$ failed due to the millimolar binding affinity [75]. In 1993, work done by Xue et al. showed CA I exhibited a greater binding affinity for HCO$_3^−$ in relation to CA II. They hypothesized that this difference in affinity was due to the H200 residue, which is a Thr in CA II [76]. Site directed mutagenesis was performed on CA II to make a T200H variant. This variant demonstrated an increase in anion affinities in relation to wild type and also greatly increased HCO$_3^−$ binding affinity [77]. Therefore, the T200H CA II variant was used to determine the first crystal structure of CA II in complex with HCO$_3^−$. The HCO$_3^−$ was observed to bind directly to the zinc, with one of the oxygen atoms occupying the site of the hydroxyl. The other two oxygen atoms bind in the hydrophobic pocket oriented similarly to the previously mentioned CO$_2$, one of which displaces an ordered water and forms a hydrogen bond with the NH of T199 [76]. The T199 gatekeeper residue was shown to be important for orienting the substrate as seen previously in the CO$_2$ binding studies. T199 is responsible for orienting HCO$_3^−$ for the dehydration reaction [78]. Therefore, it was hypothesized that mutating this residue would have detrimental impacts on CA activity.

To further investigate HCO$_3^−$ binding, West et al. revisited the hydrophobic pocket V143 mutation studies. Combining kinetic assays with $^{18}$O exchange mass spectrometry, they concluded that a V143A mutation only resulted in a 2-fold decrease in activity, while V143I and V143L resulted in a larger 20-fold decrease [67]. Traditional X-ray crystallography was conducted on all three of these variants; however no changes in the active site were seen. To determine the mutations’ influence on the mechanism, the V143I mutant was cryocooled and put under 15 atm of CO$_2$ in accordance with previous CO$_2$ binding studies [74]. Diffraction data was collected and
showed similar CO₂ binding with the previously reported wild type. However, HCO₃⁻ was also captured in the active site with an occupancy of 0.35 and CO₂ with an occupancy of 0.65. This data suggested that the V143I mutation caused a slower conversion of CO₂ into HCO₃⁻ with an increased affinity for HCO₃⁻, allowing it to be captured in the active site. Compared to the previously reported T200H variant and HCO₃⁻ complex, the V143I showed a 0.4 Å movement in the unbound oxygen atoms and an overall 18° rotation [67]. While similar, this slight movement and rotation were due to increased steric hindrance of the added Ile. This study concluded that the V143I increased an apparent energy barrier of catalysis by ~1.7 kcal/mol owing to the capture of HCO₃⁻. This is possibly the result of tighter binding of HCO₃⁻ in the variant as previous CO₂ capture experiments reported no bound HCO₃⁻ [67, 74].

2.3. Proton Transfer. The second step of the α CA mechanism, proton transfer, was first hypothesized in 1975 by Steiner et al. Here, they reported that for a realistic mechanism for α CAs, the rate limiting step was the regeneration of the zinc-bound hydroxyl [61]. They predicted that there must be an intramolecular transfer of the proton from the zinc-bound water onto a nearby residue, most likely an ionizable His. This concept was tested in 1988 by Silverman and Lindskog during their structural and kinetic testing, hypothesizing that the ordered water network was responsible for the transfer of the proton onto H64 [63]. However, the concept of the proton wire and transfer was just a hypothesis based on the ordered water network found in all CA II crystal structures. There was little to no experimental proof confirming the function of the proton until a series of MD experiments and mutation-based structure activity relationship studies.

The initial MD predictions for the proton wire were performed in 2006 by Fisher et al. Here, they performed a simplified MD simulation, modeling the CA II structure coordinates with 220 locations of water molecules inside a solvated cubic simulation box [70]. The simulation proposed the most likely proton transfer occurring through the 6-membered ordered water network and transferred the proton to H64. The zinc-bound water is stabilized through both T199 and a water termed “deep water” (DW) adjacent to the hydrophobic pocket. This stabilization along with zinc coordination allows a proton to be removed from the water molecule with negligible energy input [70]. The proton is then transferred to W1 and stabilized by the oxygen atom of T200. It is then taken by W2, where it is further stabilized by W3a and W3b. These subsidiary waters do not actually take the proton but instead were predicted to help stabilize the protonated W2. Finally, the simulation estimated that H64 in the “in” confirmation accepts the proton and undergoes a conformational change into the “out” position where the proton is donated to the bulk solvent and out of the active site [70]. While accurate and groundbreaking, this simulation was only modeled using the 6 water cluster, H64, and the zinc-bound solvent, ignoring the rest of the enzyme. This work proved the proton wire was the likely mechanism of proton transfer, but more work was needed to confirm (Figure 4).

Subsequent MD simulations at a higher degree of complexity were performed in an attempt to model the experimentally observed rates of CAs. However, in the early 2000s, many of these experiments produced varying results due to differences in MD methodology [79–81]. In 2009 Maupin et al. used a new technique known as multistate empirical valence bond MD (MS-EVB) to model proton transfer throughout the CA active site that could accurately predict the kinetic rates. Compared to other methods, MS-EVB fully describes the dynamical charge defect delocalization and Grothuss shuttling of the proton and multiple water molecules [81, 82]. This work also incorporated ionizable moieties that accurately portray zinc-bound water/hydroxide of CA II. Analysis of their MD simulation indicates that the proton travels along a similar path as previously predicted.

\[ \text{Zinc-bound } H_2O \rightarrow W1 \rightarrow W2 \rightarrow H64 \text{ “in” } \rightarrow H64 \text{ “out” } \rightarrow \text{Bulk Solvent} \] (3)

First, the zinc-bound H₂O interacts with W1 to form a Zundel cation (H₂O₃⁺). The proton is then transferred onto W1, forming a higher energy Eigen cation (H₂O₄⁺). The proton is further transferred to W2 as it reverts back to the low energy Zundel cation [81]. However, the highest energy barrier was observed before the proton was transferred to H64, indicating another complex formation with a final Eigen cation generated with W3a. This indicates that, for proton transfer, the limiting rate is Eigen cation formation upon proton transfer to W3a [81].

This MS-EVB MD simulation also accurately predicts H64 proton acceptance and donation. When H64 is in the “out” conformation the free energy barrier for proton transfer is 14.6 ± 0.4 kcal/mol [81]. However, once in the “in” conformation this barrier is lowered to 10.0 ± 0.4 kcal/mol. This decrease in the free energy barrier is due to H64 stabilizing interactions with W2 and the rest of the water network, priming the waters for proton transfer. This energy barrier translates to an estimated catalytic rate of 1.0 µs⁻¹, matching known catalytic rates of 1.1 µs⁻¹ [81]. Building on the accuracy of this simulation, the MS-EVB model also accurately predicts the effect of exogenous buffer on the rate of proton transfer. At physiologically relevant conditions, there is essentially an unlimited supply of exogenous buffer for CA to utilize for its proton transfer. At these levels of buffer, the rate limiting step of proton transfer is the transfer of protons through the water cluster and not the H64 release. However, once exogenous buffer is limited, the rate becomes more dependent on H64 proton release, as the limited buffer is less accepting of protons [81, 83]. The MS-EVB simulation accurately predicts this, as the predicted rate of buffer limited conditions drastically decreases to 0.03 µs⁻¹ compared to the rate of physiological buffer conditions of 1.0 µs⁻¹.

To confirm the MS-EVB MD simulations, further experiments were conducted to mutate H64 and confirm the importance of proton transfer. H64 was mutated in CA II to Ala and a combination of kinetic, structure, and MD experiments were conducted to determine the effect [84]. Initially,
Figure 4: Active site mechanism. The surface rendition of CA II with zinc bound to active site residues H94, H96, and H119. The hydrophobic side is depicted in orange while the hydrophilic side is depicted in purple. Blue arrows depict substrate flow, CO$_2$ and HCO$_3^-$ through the hydrophobic side and proton transfer through the hydrophilic side. A cartoon depiction of α CA ping-pong mechanism.

a kinetic test was done on the H64A variant to determine the effect on turnover rate. Using O$_{18}$ mass spectrometry, H64A variant exhibited a 20-50-fold decrease in rate. The effect of having a hydrophobic residue at the point of proton donation resulted in a destabilized water network with W3b being completely absent in the structure. This caused the rate to be dependent on H$_2$O deprotonation occurring spontaneously. MS-EVB simulations predicted that other “self-rescuing” pathways were used to shuttle protons, utilizing acidic residues near the active site for proton transfer, forgoing the typical 6 water cluster [84]. 4-Methylimidazole (4MI), a known activator of CAs and potential proton acceptor, was used as a chemical rescue test in kinetic and MD experiments. Again, O$_{18}$ mass spectrometry was used to determine the effects of rescuing H64A variant with 4MI. The 4MI was able to restore functional levels of enzyme activity without the normal H64. The variant was then crystallized in complex with 4MI to determine how its activating effects could rescue proton transfer. Using a combination of NMR and X-ray determined binding sites, the 4MI efficiently restored proton transfer in the MD simulation, further validating the importance of a proton acceptor residue [84].

2.4. Neutron Crystallography. While X-ray crystallography is utilized to determine the solvent molecule positions throughout the structure, the methodology is unable to determine their orientation. The clustered water orientations are needed to validate the proton wire as this mechanism largely depends on distinct orientations of the waters, allowing them to span a network of hydrogen bonds. Recent work done by Fisher et al. performed neutron crystallography on CA II at varying pHs to discover the orientations of water in the proton wire [85, 86]. Data collected from crystals at pH 10.0 showed a broken proton wire that lacked a hydrogen bond between W1 and W2. The W2, W3a, and W3b were oriented toward the imidazole ring of H64 occupying the in conformation but did not form hydrogen bonds as previously predicted [85]. However, at pH 7.8, the orientation of W1 changed. At pH 10, W1 served only as a hydrogen bond donor to both DW and T200 whereas at pH 7.8, W1 acts as a hydrogen bond acceptor of T200, forming a subsequent hydrogen bond with W2 [85, 86]. This additional hydrogen bond forces W2 and W3a to reorient in a concerted fashion to complete the proton wire. W2 now behaves as a hydrogen bond donor to H64, indicating that the most likely path for proton transfer is ZnH$_2$O/OH$^-$/W1-W2-H64 and confirming previous MD simulations and predictions [85, 86].

3. Drug Design

3.1. α CAs as a Drug Target. Multiple α CA isoforms are recognized as therapeutic targets in diseases such as glaucoma, edema, epilepsy, obesity, and cancer. Therefore, a significant portion of CA research has focused on the design of CA inhibitors (CAIs). CAI research began in 1940 with the observation that sulfanilamide (p-aminobenzene-sulfonamide) inhibited CA activity. Furthermore, this inhibitory effect showed a trend in activity with the observation that sulfanilamide showed an increase in affinity compared to benzene derivatives with a general increase in activity as the acidity of the compound increased [89]. These experiments led to the use of a drug design strategy termed the “ring approach” in which a ring system was added to a sulfonamide ZBG in order to increase affinity for the target CA [90]. Early
CAI research led to the development of first generation CAIs that are still available in the clinic, including acetazolamide (Diamox), methazolamide (Neptazane), ethoxzolamide, and dichlorphenamide (Keveyis). However, many of these compounds exhibited solubility issues. Subsequent CAI design therefore employed the “tail method” in which functional groups were added to an aromatic/heterocyclic sulfonamide scaffold in order to modulate the physicochemical properties and solubility of the compound [90]. This technique resulted in the development of second generation CAIs such as dorzolamide (Trusopt) and brinzolamide (Azopt). There are several additional sulfonamide-based CAIs clinically available for the treatment of the aforementioned diseases, including topiramate, celecoxib (Celebrex), sulpiride (Dogmatil), sulthiame, valdecoxib, zonisamide, irosustat (COUMATE), and esterone sulfamate (EMATE) (Figure 5). The role of CA in the onset or progression of each disease and the use of CAIs as a treatment will be discussed.

3.1.1. Glaucoma. Glaucoma is a disease characterized by high intraocular pressure (IOP) and loss of vision. An increase in IOP is most often associated with the retention of aqueous humor, a liquid between the cornea and lens, which is caused by an absence or decrease in humor drainage. A main component of this aqueous humor is sodium bicarbonate, the secretion of which is controlled by CA activity in the uvea of the eye. The inhibition of CA II, CA IV, and CA XII has been shown to decrease the secretion of humor, resulting in decreased IOP. Therefore, CA inhibitors are used in conjunction with adrenergic agonists or antagonists for the treatment of glaucoma [91].

The first CAIs used to treat glaucoma include acetazolamide, methazolamide, ethoxzolamide, and dichlorphenamide. However, these compounds act systemically, inhibiting multiple CA isoforms, and have been shown to cause unwanted side effects such as fatigue, abnormal tingling, and kidney stones [66, 92]. The second generation of compounds, brinzolamide and dorzolamide, exhibit nanomolar inhibition and show increased water solubility to allow for use as a topical agent, resulting in fewer unwanted side effects. However, the drug solution is acidic and can cause redness and irritation. Dorzolamide has also been recorded to exhibit some serious psychological side effects such as depression and dementia [93].

Recent developments in antiglaucoma inhibitors use the tail method with an aromatic sulfonamide scaffold to which amino, hydroxyl, or nitrate ester groups are added to increase solubility. These novel compounds have been tested as topical agents in animals and show good solubility and inhibitory effects, resulting in prolonged decreased IOP. In addition, compounds that contain a NO donating moiety can also aid in vasodilation and humor secretion, supplying more blood to the optic nerve and further decreasing IOP. These so-called “hybrid drugs” were also tested in animals and exhibited a greater reduction in IOP than either brinzolamide or dorzolamide, representing a promising class of lead compounds for antiglaucoma drug development [94].

3.1.2. Edema. Edema is a condition characterized by abnormal brain activity and seizures. It is hypothesized that CA is involved in the secretion of bicarbonate-rich cerebrospinal fluid, similar to CA function in the secretion of aqueous humor in the eye [98]. Although CA II and CA VII are both expressed in neurons, CAI design has mainly focused on the inhibition of CA VII [99]. CA VII is expressed more specifically in the cortex, hippocampus, and thalamus [100]. CA VII activity regulates neuronal pH and maintains bicarbonate gradients in the brain for the management of neuronal channels by GABA receptors (GABARs). GABARs are membrane proteins involved in an array of neuronal pathways and are sensitive to changes in pH, so these receptors are influenced by CA catalytic activity [99, 101]. CA VII is able to quickly replenish bicarbonate, resulting in a net uptake of chloride ions and depolarization, leading to excitation of the GABAR. GABAR excitation and signaling have been shown to promote seizures and CA VII activity has been associated with epilepsy [99]. While the function of CAIs in the brain is not well characterized, the inhibition of CA using acetazolamide has been shown to increase cerebral blood flow and CO\textsubscript{2} retention [102].

Classic CAIs such as acetazolamide and methazolamide have been clinically used as antiepileptics, but have more recently been replaced by topiramate [98]. Topiramate has a sulfamate moiety that interacts directly with the active site zinc and functions through the enhancement of GABA-mediated inhibition transmissions [103]. However, topiramate has been shown to cause metabolic acidosis with long-term treatments and possibly interfere with bone formation in children, when anticonvulsant therapies are most commonly started [104]. The ring method has been employed to improve antiepileptics by adding valproic acid, a well-known antiepileptic, to aromatic sulfonamide scaffolds such as acetazolamide and topiramate. Compounds with more lipophilic derivatives exhibited better anticonvulsant properties. The valproyl moiety is predicted to interact favorably with the
Figure 5: CA inhibitors structural formulae of clinically relevant CAIs.

hydrophobic pocket of the active site, making a more efficacious inhibitor. However, the efficacy of CA VII inhibitors greatly decreases if the compound is unable to penetrate the blood-brain barrier [98].

### 3.1.4. Obesity

Significant weight loss has been observed as a side effect of the antiepileptic CAIs topiramate and zonisamide. CA V research is therefore focused on the possibility of a link between CA V inhibition and weight loss. CA V is expressed in the mitochondria as two forms: Vb exhibits wide tissue distribution whereas Va is limited to liver tissue. CA V provides bicarbonate to serve as a substrate for pyruvate carboxylate in gluconeogenesis and oxaloacetate production in lipogenesis. The inhibition of CA V is therefore expected to induce weight loss by depleting a source of bicarbonate and decreasing fatty acid synthesis [105]. Furthermore, the use of a nonspecific CAI, acetazolamide, has been previously shown to decrease the production of fatty acids in human adipose tissue [106]. The majority of the CAIs studied thus far have been substituted benzene sulfonamides with substituents such as amino, hydroxyl, nitro, halogen, and carboxy moieties [107]. The tail method has also been employed to add lipophilic moieties to increase membrane permeability, ensuring the compounds are able to reach mitochondria within the cytoplasm [108]. Kinetic studies have shown that the CA V active site may accommodate bulkier inhibitors and that the location of the additional substituent has more of an effect on the inhibition constant than the chemical identity of the moiety [107, 108].

### 3.1.5. Cancer

CA IX overexpression has been observed in several cancer types including lung, renal, brain, colon, pancreatic, liver, breast, esophageal, ovarian, and skin cancer. In contrast, CA IX expression is limited to the GI tract in healthy tissue [109]. CA IX expression is regulated by hypoxia inducible factor (HIF-1). Rapidly proliferating tumor cells often develop regions of hypoxia characterized by low oxygen concentrations. In such conditions, the HIF-1α subunit is not hydroxylated, preventing recognition by the von Hippel Lindau (VHL) protein and subsequent ubiquitin degradation. HIF-1α accumulates and transports into the nucleus where it dimerizes with HIF-1β to form an active transcription factor. HIF1 regulates the expression of stress response genes, including CA IX, by binding to the hormone response element [110].

CA IX activity is hypothesized to be critical for the regulation of pH in cancer cells that must thrive in an acidic tumor microenvironment [111–113]. The rapid growth and proliferation of aggressive, metastatic cancers often induce a metabolic switch from oxidative phosphorylation to anaerobic glycolysis, termed the Warburg effect. This metabolic switch increases the amount of lactic acid that is produced and exported by the cell, acidifying the extracellular milieu to pH ~6.5 [114–116]. A decrease in pH impacts cell viability by disrupting biological activities such as ATP production, cell migration, proliferation, and protein synthesis [117]. The overexpression of CA IX enables neoplastic cells to survive such harsh conditions by catalyzing the production of bicarbonate, which can act as a buffer in the surrounding environment or be transported into the cell to maintain near physiological pH [66]. As such, CA IX has been identified as a biomarker and therapeutic target for various cancer types, with a primary focus on breast cancer. Furthermore, the inhibition of CA IX activity via knock down or small molecule inhibition has been shown to decrease tumor
volume and improve overall survival in breast cancer mouse models [118].

As CA IX is a membrane-bound isoform with an extracellular catalytic domain, CAI selectivity can be enhanced by designing membrane impermeable compounds to prevent the off-target binding of cytosolic CAs. For example, positively or negatively charged hydrophilic moieties can be added to promote impermeability. However, these properties make it unlikely that such compounds would enter the bloodstream or be developed into a drug to be taken orally. Therefore, inhibitors are designed as prodrugs with hydrophobic moieties that mask the desired, inhibitory substituents until the compound is present in the reductive conditions of a hypoxic environment where it will be hydrolyzed and become an active inhibitor [119].

 Artificial sweeteners have recently been identified as potential CAs for the development of cancer therapeutics. Saccharin and acesulfame potassium were shown to bind within the CA active site and exhibit binding affinities in the micromolar range [120, 121]. Moreover, two sulfonamide-based CAs, SLC-0111 (4-(4-fluorophenylureido)-benzenesulfonamide) and indisulam (N-(3-chloro-7-indolyl)-1,4-benzenedisulfonamide), have successfully completed phase I clinical trials and are currently entering phase II [122, 123].

3.2. Isoform Selective Drug Design. Although several CAs are currently in use for the treatment of diseases, these clinically available compounds do not exhibit sufficient isoform selectivity and therefore bind to multiple CA isoforms. Off-target binding can lead to sequestration of the drug, requiring higher doses for treatment and subsequently decreasing efficacy. However, the design of isoform specific CAs is complicated by the structural homology shared by the catalytically active CAs. X-ray and neutron crystallography are therefore utilized to analyze the binding of inhibitors in the CA active site and guide drug design of isoform specific inhibitors.

3.2.1. Structure-Based Drug Design. Structure-guided drug design is a technique that uses high resolution crystal structures of a molecular target to rationalize the design of high affinity, small molecule inhibitors. This process often begins with high throughput screening of different classes of inhibitors to identify lead compounds that inhibit CA activity. The most promising compounds, which exhibit binding affinities in the nano- to micromolar range, are then studied using X-ray crystallography. The crystal structure complex is analyzed to identify interactions between the compound and target molecule that promote selective binding. New derivatives can then be designed to promote such interactions via the addition of functional groups to the lead compound [124].

The tail approach, previously described as a method to improve inhibitor solubility, can also be applied in structure-guided drug design. The mapping and comparison of active site residues between isoforms have identified residues unique to the target isozymes. Elongation or derivatization of a compound tail can promote interactions with these isoform unique residues, improving selectivity [66]. New inhibitors can also be designed using a fragment approach in which small molecules that bind different areas of the active site are chemically linked to synthesize a single compound. This technique has the potential to combine a molecule with high affinity for the active site zinc to a fragment that binds to isoform unique residues at the entrance of the active site [125].

It is important to recognize that the affinity of a CAI is dependent on the free energy of binding (ΔG = ΔH - T ΔS) with both enthalpic and entropic contributions. A ligand loses rotational freedom upon binding, decreasing ΔS. This entropic penalty can be counteracted by enthalpic gains upon the formation of interactions between the ligand and target molecule in addition to increases in entropy as water molecules are displaced from the active site [126]. Both terms of the free energy calculation must therefore be considered during the selection of chemical moieties in the drug design process. For example, the addition of hydrophobic functional groups has been shown to induce entropic penalties that are not balanced by the increase in enthalpy, decreasing the binding affinity for the target [127]. One must also consider how compound derivatization will impact drug delivery and pharmacological properties such as absorption, distribution, metabolism, and excretion. Compounds that are to be delivered orally should follow Lipinski’s rule of five: fewer than 5 Hydrogen bond donors and 10 acceptors, molecular weight < 500 g/mol, and the log P < 5 [128]. However, these are not steadfast rules and many FDA approved drugs do not fulfill each of the five rules [129].

3.2.2. Selective Pocket. The combination of high resolution X-ray and medium resolution neutron crystal structures has led to a thorough characterization of the α CA active site as described above. Nearly 500 structures of CA II in complex with CAs have been deposited into the PDB, allowing the differentiation of compound binding that can be attributed to functionalization or derivatization of lead compounds. For example, shorter inhibitors that contain aromatic functional groups orient into the hydrophobic region of the active site (I91, V121, F131, V135, L141, V143, L198, P202, L204, V207, and W209) and are often involved in parallel stacking interactions with F131. Alternatively, compounds with longer tails or charge were shown to bind preferentially in the hydrophilic region (N62, H64, N67, Q92, T199, and T200) (Figure 6(a)) [130].

The hydrophobic half of the active site was then further divided into two pockets in which the tails of inhibitors orient. Pocket 1 consists of residues L198, F131, V135, and L204 whereas Pocket 2 contains I91, V121, and F131 (Figures 6(b) and 6(c)). The bulky, aromatic side chain of F131 separates the two pockets and often dictates inhibitor binding. The superposition of approximately 30 CAs identified that the majority of compound tails that bind in the hydrophobic region orient into Pocket 2. However, compound binding affinities were not observed to correlate to pocket preference [131]. Crystallographic studies showed that more flexible linkers, such as ureido fragment (NHCONH), promote tail binding in Pocket 1 and increase affinity [132].
Pockets in α-CAs. (a) Hydrophilic and hydrophobic regions of the active site have been colored purple and orange, respectively. The hydrophobic pockets (b) 1 and (c) 2 have been shaded yellow and green, respectively. (d) Isoform unique residues that constitute the selective pocket are shown in blue (Table 2).

In 2013, a structural comparison of the binding of all nonredundant inhibitors in complex with CA II was performed. Of the 145 compounds, only 14 were observed to orient toward a region between the hydrophobic and hydrophilic areas of the active site. This region was termed the “selective pocket” due to the variability of residues 67, 69, 91, and 131 between isoforms (Figure 6(d)) [133]. Therefore, this pocket can be targeted during the design of CAIs to improve isoform selectivity.

3.2.3. Neutron Crystallography. The use of neutron crystallography in conjunction with X-ray crystallography allows the visualization of both “heavy” (non-H) and “light” (H) atoms. It is therefore possible to determine the protonation state of amino acid side chains and inhibitors using joint refinement. For example, the standard CAI acetazolamide exhibits three possible protonation states in solution. Initial solution state NMR studies hypothesized that sulfonamide-based compounds bind in the deprotonated state and interact with catalytic zinc through the sulfonamide nitrogen [134]. Joint neutron/X-ray studies confirmed that acetazolamide binds in the anionic form with the negatively charged sulfonamido coordinating to the zinc. Neutron crystallography can also provide insight into hydrogen bonding and the displacement of water molecules, which can be used to guide the design of new CAIs [135]. For example, the neutron structure of acetazolamide was compared to methazolamide. The addition of the hydrophobic methyl group prevented the formation of a hydrogen bond and resulted in the displacement of four additional water molecules in comparison to acetazolamide binding. However, the gain in entropy is hypothesized to counteract the loss of hydrogen bonding, justifying the similar binding affinities of the two CAIs [136].

3.2.4. Nonclassical CAIs. The search for isoform selective CAIs has also led to the discovery of nonsulfonamide-based inhibitors that exhibit unique mechanisms of inhibition, including compounds that anchor to the zinc-bound water and CAIs that bind outside the active site, occluding entrance of substrate. These compounds have the additional benefit of preventing adverse effects in individuals with sulfa allergies [88, 137]. Compounds that anchor through the zinc-bound water deselect from zinc binding that is well established for sulfonamide-based inhibitors. Therefore, the binding affinity relies primarily on interactions with residues in the active
|   | CA I   | CA II  | CA III  | CA IV  | CA VA  | CA VB  | CA VI  | CA VII | CA VIII | CA IX  | CA X   | CA XI  | CA XII | CA XIII | CA XIV |
|---|--------|--------|---------|--------|--------|--------|--------|--------|---------|--------|--------|--------|--------|---------|--------|
| 5 | W      | W      | W       | W      | C      | C      | W      | W      | W       | W      | W      | W      | W      | W       | W      |
| 7 | Y      | Y      | Y       | Y      | W      | Y      | W      | Y      | Y       | Y      | Y      | Y      | Y      | Y       | Y      |
| 62| V      | N      | N       | N      | T      | N      | N      | N      | N       | D      | N      | T      | T      | N       | S      |
| 64| H      | H      | K      | H      | Y      | Y      | H      | H      | H       | R      | R      | H      | H      | H       | H      |
| 65| S      | A      | T      | S      | S      | L      | T      | S      | T       | S      | H      | S      | S      | T       | T      |
| 67| H      | N      | R      | M      | M      | Q      | Q      | Q      | Q       | Q      | Q      | S      | S      | K       | N      |
| 92| Q      | Q      | Q      | Q      | Q      | Q      | Q      | Q      | Q       | Q      | Q      | Q      | Q      | Q       | Q      |
| 121| A      | I      | V      | V      | V      | V      | V      | V      | V       | V      | V      | V      | V      | V       | V      |
| 141| L      | L      | I      | I      | L      | L      | L      | L      | L       | I      | L      | L      | L      | L       | L      |
| 143| V      | V      | V      | V      | V      | V      | V      | V      | V       | I      | V      | I      | I       | V       | V      |
| 198| L      | L      | F      | L      | L      | L      | L      | L      | L       | L      | M      | L      | L       | L      | L      |
| 200| H      | T      | T      | T      | T      | T      | T      | T      | I       | I      | T      | I      | T       | V       | T      |
| 207| V      | V      | I      | V      | V      | V      | V      | V      | V       | A      | V      | V      | V       | V      | V      |
| 209| W      | W      | W      | W      | W      | W      | W      | W      | W       | W      | W      | W      | W      | W       | W      |
| 244| N      | N      | N      | N      | N      | N      | N      | N      | N       | N      | N      | N      | N      | N       | N      |
| 10-15 Å | | | | | | | | | | | | | | | |
| 60| I      | L      | L      | Q      | W      | W      | V      | T      | T       | R      | Y      | Y      | T       | S       | H      |
| 69| N      | E      | V      | L      | E      | E      | S      | D      | I       | T      | R      | L      | N       | D       | S      |
| 91| F      | I      | R      | K      | K      | Q      | K      | Y      | L       | E      | S      | T       | R       | A       | A      |
| 131| L      | F      | F      | V      | Y      | F      | Y      | I      | V       | V      | L      | A       | F       | L       | L      |
| 135| A      | V      | L      | Q      | V      | A      | Q      | A      | V       | L      | A      | T       | S       | A       | A      |
| 201| P      | P      | P      | P      | P      | P      | P      | P      | P       | P      | P      | P      | P       | P       | P      |
| 202| P      | P      | P      | T      | P      | P      | P      | P      | P       | P      | P      | P      | P       | P       | P      |
| 204| Y      | L      | E      | D      | T      | S      | T      | S      | S       | A      | Y      | S       | N       | L       | Y      |
| 15-20 Å | | | | | | | | | | | | | | | |
| 4 | D      | H      | E      | H      | S      | T      | D      | G      | E       | H      | W      | W      | K      | S       | H      |
| 19| L      | D      | L      | P      | V      | H      | L      | V      | V       | V      | V      | V      | K      | F       | S      |
| 20| Y      | F      | F      | V      | V      | D      | Y      | Y      | F       | S      | N      | N      | Y       | F       | Y      |
| 21| P      | P      | P      | S      | L      | P      | P      | P      | P       | S      | A      | P      | P       | P       | P      |
| 22| I      | I      | N      | V      | V      | V      | A      | I      | D       | A      | A      | A      | S       | I       | E      |
| 57| K      | L      | K      | W      | L      | L      | F      | L      | C       | L      | G      | G      | F       | K       | L      |
| 58| E      | R      | T      | T      | Y      | H      | P      | S      | E       | R      | T      | T      | L       | I       | D      |
| 72| D      | D      | D      | N      | D      | D      | S      | D      | S       | P      | K      | P      | S       | D       | S      |
| 123| W      | W      | W      | K      | W      | Y      | W      | W      | W       | L      | Y      | F      | Y       | W       | Y      |
| 130| S      | D      | T      | N      | N      | N      | S      | T      | S       | R      | N      | N      | D       | S       | S      |
| 132| A      | G      | K      | K      | K      | E      | D      | G      | D       | D      | T      | S      | S       | V       | S      |
| 136| S      | Q      | K      | D      | V      | L      | D      | S      | G       | G      | K      | R      | N       | H       | E      |
| 170| K      | K      | K      | K      | P      | K      | P      | K      | E       | K      | K      | K      | K       | K       | K      |
| 171| G      | G      | G      | E      | D      | D      | G      | G      | G       | G      | N      | N      | G       | G       | D      |
| 173| R      | S      | E      | S      | R      | L      | R      | K      | S       | E      | A      | A      | E       | Q       | K      |
site. In comparison to compounds that bind directly to the catalytic zinc, inhibitors that anchor through the water extend an additional 2.5–3 Å from the zinc, increasing the probability of forming interactions with isoform unique residues in the selective pocket. Examples of such compounds include phenol- and carboxylic-based inhibitors. Both classes anchor to the zinc-bound solvent through the hydroxyl moiety. Phenol-based compounds exhibit binding affinities within the micromolar range and are hypothesized to inhibit activity via obstruction of the CO$_2$ binding site caused by interactions of the phenyl group with hydrophobic residues in the active site [138]. Increasing the length or complexity of phenols, as seen in phenol-based natural products, has been shown to improve inhibitory properties against CAs [139]. Similarly, the alteration of scaffold size and chemical properties in carboxylic acid CAIs has been shown to dictate the orientation of the tail in the active site, promoting isoform selectivity based upon steric hindrance of active site residues. Again, compounds of increasing length that extend toward the selective pocket have been shown to improve the binding affinity over 100-fold [140].

The majority of isoform unique residues encircle the entrance of the active site. Therefore, compounds that inhibit activity by occlusion of the active site are more likely to interact with these unique residues, improving isoform selectivity of the CAIs. Examples of this class of inhibitors include disaccharides and artificial sweeteners, such as sucrose and sucralose. Interestingly, sucralose inhibits CA activity in the micromolar range whereas sucrose binds but does not inhibit CA activity. Based on the crystal structures, sucralose binds along the hydrophobic region and is predicted to prevent entry of CO$_2$. In contrast, a small opening into the active site exists when sucrose is bound, allowing movement of CO$_2$ [141, 142].

3.3. CAIs as Antibiotics. Bacterial resistance is a growing issue worldwide and pathogenic strains such as S. aureus, M. tuberculosis, H. pylori, B. suis, and S. pneumoniae no longer respond to classic antibiotics [143]. The need to identify new molecular targets essential to the life cycle and virulence of the pathogen has led to a focus on inhibition of bacterial CAs. Because the majority of bacterial CAs belong to the β CA class that is not expressed in humans, these enzymes represent promising drug targets. Additionally, CAIs would represent a different mechanism of action compared to current antibiotics and would therefore be less likely to result in resistance [39].

Sulfonamide-based compounds have been shown to inhibit the activity of β CAs both in vitro and in vivo in the H. pylori, S. pneumoniae, and B. suis bacterial [144–147]. However, many CAIs that exhibit favorable inhibition profiles in vitro have not been observed in vivo, which is hypothesized to be due to the fact that the CAIs are impermeable to the cell wall [148]. Because sulfonamides also inhibit the α CAs expressed in humans, other classes of small molecules have been explored such as phenols, boronic acids, and anions [39]. A promising class includes dithiocarbamate inhibitors that coordinate to the metal ion through the CS$_2$$^-$ moiety. Aryl, aryalkyl-, heterocyclic, and aliphatic derivatization were shown to increase affinity for β CAs isolated from M. tuberculosis with some compounds binding in the subnanomolar range [149]. Although these compounds exhibit micromolar binding affinities, carboxylic acids have also been under investigation due to the ease of derivatization. Increasing the hydrophobicity of this group of CAIs could potentially improve cell wall permeability [150].

4. Conclusion

CAs are an essential class of enzymes in every class of life, from marine diatoms and bacteria to humans. The crystal structure of CA II was amongst the first structures to be determined and was one of the seven structures that contributed to the development of the PDB. Structural characterization has not only led to the classification of the seven CA families but has also improved the understanding of the catalytic mechanism. X-ray and neutron crystallography studies have identified the binding sites of both substrate/product and elucidated proton transfer through observation of an ordered water network and dual conformations of the proton shuttle residue. These techniques have also driven the design of CAIs as antiglaucoma, antiepileptic, antinfective, and anticancer therapies in addition to antibiotics.

Although structure-guided drug design has led to several classes of high affinity CAIs, the majority of these compounds still do not exhibit sufficient isoform selectivity to prevent off-target binding. Therefore, interactions between an inhibitor tail and the four residues of the selective pocket do not adequately differentiate between the fifteen human α CAs. Structural biologists are now mapping residues beyond the active site. CA can be divided into three zones, 5–10 Å, 10–15 Å, and 15–20 Å extending radially from the catalytic zinc (Figure 7, Table 2). With the exception of residue 67, the majority of residues in the 5 – 10 Å zone remain conserved between the isozymes. This observation is reinforced by the lack of selectivity observed for compact CAIs that bind deep within the active site. The 10 – 15 Å zone contains the selective pocket and has therefore been the targeted region for drug design over the past several years. However, the number of isoform unique residues increases further in the 15 – 20 Å zone. In addition to the functionalization of a compound tail, the extension of the linker and/or tail components of CAIs is predicted to further increase selectivity by promoting interactions with the diverse residues outside the active site.

Our current knowledge of the catalytic mechanism is limited to static crystal structures and a complete understanding of the dynamics of the reaction has yet to be achieved. Insight into the dynamics of the proton wire has been recently expanded by the determination of X-ray crystal structures from crystals under high CO$_2$ pressure. Varying the time of incubation at room temperature prior to data collection provides a method to visualize CO$_2$ release. This study identified an extension of the active site water network termed the entrance conduit waters (EC1, EC2, EC3, EC4, and EC5). Analysis of the alternate conformations of these
dynamic water molecules led to the proposal of a restoration of the zinc-bound water following HCO$_3^-$ diffusion [151].

The development of X-ray free electron laser (XFEL) sources now allows the design of serial femtosecond crystallography (SFX) experiments on a time scale capable of capturing intermediate states of a catalytic mechanism [152]. With one of the fastest known biological reactions, CA is an ideal candidate for such SFX experiments to map the movements of substrate/product through the active site. Microcrystals of CA II suitable for XFEL data collection have recently been produced and confirmed to diffract at the Pohang Accelerator Laboratory XFEL [153]. A further benefit is the collection of diffraction data at room temperature before the onset of radiation damage. Data collection at cryogenic temperatures may limit occupancy of alternate conformations relevant to the catalytic mechanism [154]. Alternatively, the collection of multiple data sets at increasing temperatures, known as multitemperature or temperature jump crystallography, allows the sampling of conformational heterogeneity [155].

Disclosure

The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health.

Conflicts of Interest

The authors declare that there are no conflicts of interest regarding the publication of this paper.

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