Two-dimensional infrared (2D IR) spectroscopy for elucidating ion occupancies in the selectivity filter of ion channels

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Abstract. This article reviews work presented at the ESCBM 2017 on using two-dimensional infrared (2D IR) spectroscopy to probe ion binding configurations in the potassium ion channel KcsA. We discuss two studies in which we use 2D IR spectroscopy in conjunction with protein semisynthesis and molecular dynamics (MD) simulations to test against two competing models of ion permeation and to investigate how changes in the conformation of the intracellular gate affect the structure and ion affinities of the selectivity filter. In our initial study, the 2D IR spectrum of an isotopically labeled KcsA filter reveals two spectral features that correspond to different structures and ion binding configurations. MD simulations help us link the experiments to atomistic structures, allowing us to determine the prevailing mechanism of ion conduction. In a follow-up study, we probe how ion occupancies in the filter change with the opening and closing of the intracellular gate. We experimentally modulate the conformation of KcsA using different sample conditions, and are able to show that the ion occupancies in the filter change with the state of the gate, revealing complexities in the conformational landscape of the potassium ion channel. Our work shows the potential of these techniques in addressing the fundamental biophysical questions in membrane protein structure and function.

Keywords: 2D IR spectroscopy, potassium ion channels, ion permeation, protein semisynthesis

1. Introduction and background

Potassium ion channels are a family of ion channels that conduct potassium ions across cell membranes with high selectivity and fidelity. These ion channels play a pivotal role in many physiologically relevant processes, including signal transduction and sensation, and are essential for maintaining the resting membrane potentials of excitable cells [11,13,17,24,28,40]. While these channels are diverse in their channel activity and gating mechanisms, the architecture of these potassium channels are strikingly similar [13,40]. Proteins in this superfamily are canonically composed of multiple subunits with several transmembrane-spanning α-helices and a conserved pore-loop domain that assemble to form the ion permeation pathway through which potassium ions cross the cell membrane. In the ion permeation pathway, the ions typically traverse a narrow central pore, known as the selectivity filter, which is lined with backbone carbonyls that coordinate ions during conduction. The electronegative carbonyls from the signature sequence motif ... TVGYG... point towards the central axis and form the binding sites that

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interact with passing cations [18]. This selectivity filter is highly conserved among K⁺ channels and is the structural element that enables these channels to rapidly conduct K⁺ ions.

K⁺ channels conduct ions at a rate near the diffusion limit (∼10⁸ s⁻¹) with high selectivity for K⁺ over Na⁺, so understanding the underlying molecular basis of ion conduction is critical to figuring out how these channels conduct at such high throughput rates without compromising fidelity. The underpinnings for the accepted model for conduction, the knock-on model, were initially laid out by Hodgkin and Keynes in their landmark studies of squid giant axon [26]. In order to recapture the near-diffusion limited conduction rates that were measured from their electrophysiology studies, they postulated that the K⁺ ions must propagate through a chain of potassium selective sites in a single file [26]. For every K⁺ ion that enters through the intracellular side, another leaves the pore on the extracellular side, resulting in a net translocation of one K⁺ ion [26]. MacKinnon’s crystal structure of the bacterial potassium ion channel, KcsA, confirmed what Hodgkin and Keynes hypothesized about the ion permeation mechanism through potassium ion channels [18]. The 3.2 Å X-ray crystal structure revealed a four-fold symmetric channel with a central pore lined with backbone carbonyls that form four distinct binding sites (S1-S4) that coordinate potassium ions during conduction [18]. Indeed, the structure showed that the pore was wide enough for a single file of ions, but interestingly, the measured distance between ions suggested that they did not occupy adjacent binding sites [18]. Anomalous scattering from structural studies carried out using electron dense Tl⁺ as a surrogate for K⁺ revealed the absolute occupancies of the binding sites in the filter [57]. Results from this study showed that each site has an ion occupancy of ~0.5, which was also interpreted to reflect an equal population of [S2, S4] and [S1, S3] ion configurations [57,58]. The knock-on model was further refined by streaming potential measurements that revealed the net translocation of one water molecule with the passage of a K⁺ ion [1,27]. Molecular dynamics (MD) simulations and multi-ion potential of mean force (PMF) calculations further corroborated the experiments and revealed that small fluctuations in the filter facilitate barrierless ion permeation [8,9,48]. Figure 1(E) shows the two ion binding configurations that comprise the knock-on model of ion conduction. In this model, water and ions alternate through the filter and the two configurations (i.e. [W, S2, W, S4] and [S1, W, S3, W]) are essentially energetically equivalent [9]. The filter alternates between these two configurations during conduction: As a K⁺ ion enters from the intracellular cavity, the single file of water and ion advances towards the extracellular side, resulting in the efflux of a single K⁺ ion and water molecule on the other side of the filter. Although there is strong support for the knock-on mechanism of permeation, these experiments and simulations cannot completely rule out other competing models as the experiments are ambiguous enough to allow for alternative interpretation [2,19,21]. Results from a recent MD simulation study suggest that the near diffusion-limited conduction rates observed experimentally cannot be recapitulated by a permeation mechanism in which water and ions alternate through the filter [32]. Removing water from the filter, they were able to recover measurable conduction rates, and, interestingly, they were able to reproduce the high throughput rates of typical K⁺ channels when the ions were in direct contact [32]. In this hard-knock model, shown in Fig. 1(E), ions traverse the filter in pairs and water does not cotranslate with the ions [32]. Refinement of the absolute occupancies from existing X-ray crystal structures demonstrated that the model is supported by the body of experimental evidence [32]. The knock-on and the hard-knock models are competing mechanisms of permeation. In the former case, ions propagate through the filter with intervening water molecules, contrary to the latter which suggests that ions go through the pore in pairs without water. Without new experimental evidence, it is difficult to tell which of these models truly captures the molecular mechanisms of conduction through potassium ion channels.
Gating, or how these channels allow ions to access the pore, is characterized by large conformational changes in the protein as well as changes in the structure and ion occupancy in the selectivity filter. For the potassium ion channel KcsA, specifically, these structural transitions between the conducting and non-conducting states of the channel are controlled at both the selectivity filter and the intracellular helix bundle. Previous structural studies have pointed to two unique structural conformations of the filter referred to as the conductive and constricted states. In the conductive conformation (Fig. 1(B)), the backbone carboxyls that form the binding sites point directly into the cavity of the filter to coordinate passing ions. The constricted conformation (Fig. 1(D)) is characterized by the pinching in of the pore into an hourglass shape at Gly77, which splays the backbone carboxyls of nearby residues away from the central axis and destabilizes the S2 and S3 binding sites. Likewise, the intracellular helix bundle gate goes between two different conformational states, as illustrated in Fig. 1(A) and 1(C). In the closed conformation, the inner helices of the channel form an inverted pyramid, occluding ions from the intracellular side of the membrane from the pore. The transition to the open state causes these helices to rotate and splay open (Fig. 1(C)), separating the helices and permitting permeation. The structural conformations of both the intracellular helix bundle gate and the selectivity filter are modulated by pH and K⁺ concentrations, re-
spectively [16,18,57,58]. Not surprisingly, these two separate domains are allosterically coupled. When the helix gate opens as the channel transitions from the closed to the open conformation to enable ion flux, it promotes the conformational change of the selectivity filter [14–16]. Therefore, to elucidate the molecular determinants of gating, it is imperative to determine the degree of coupling between the inner helix gate and the selectivity filter and how this interplay ultimately affects ion conduction. Previous experiments and simulations have suggested a mechanical mechanism by which the collective motions of the inner helices affect the conformation of the selectivity filter [14–16,45]. These studies point to the Phe103 residue, which resides near the hinge of the inner helix, sterically clashing with the bottom of the selectivity filter when the channel transitions between the closed to the open position [15,45]. This then causes a conformational change in the selectivity filter, subsequently leading to inactivation of the channel [14,16,35,39]. To this end, the intricacies of the communication pathway between the selectivity filter and the helix bundle gate are complex, and remain to be delineated.

In this work, we describe the experimental implementation of two-dimensional infrared (2D IR) spectroscopy to address fundamental biophysical questions in ion channel conduction and gating. The theoretical details of third-order nonlinear spectroscopy have been described elsewhere [23,42], so, instead, we will focus on the aspects of this technique that make it an invaluable tool for studying these ion channels and for answering questions about conduction and gating. 2D IR spectroscopy is a relatively new technology that utilizes molecular vibrations to gain insight into the structure and dynamics of biomacromolecular systems [3,12,22,47,51,56]. The inherent time resolution of 2D IR spectroscopy is on the order of picoseconds, which is much faster than the timescales of many physiologically relevant biological processes. This means that when we take a 2D IR spectrum of a protein, we are fundamentally taking a snapshot of the structural distributions of the protein and its local environment [22,25]. In terms of the ion channels, this means that we can probe the instantaneous ion binding configurations in the selectivity filter. Furthermore, vibrations are intrinsically sensitive to their local electrostatic environment, and, as such, their frequencies and lineshapes are influenced by nearby ions, water, and even the structure and dynamics of the protein itself [20]. In other words, the magnitude and direction of spectral shift for the amide I mode, the vibrational mode typically measured in protein IR spectroscopy, is highly dependent on the chemical surroundings. The spectral signature for a carbonyl in close proximity to an ion is different than that of a carbonyl hydrogen-bonded to water, and so, we can use this to our advantage to experimentally probe ion occupancies in the selectivity filter. It is this combination of the inherent time resolution and the sensitivity of vibrations to the local electric fields that makes 2D IR spectroscopy a particularly powerful tool in probing ion binding configurations in the pore of potassium ion channels.

2. Objective

The data we present at ECSBM 2017 represent an overview of two studies in which we used 2D IR spectroscopy to probe ion occupancies in the selectivity filter of the bacterial potassium ion channel, KcsA. In our initial study, we use a combination of protein semisynthesis, 2D IR spectroscopy and MD simulations to probe the instantaneous ion binding configurations in the pore of KcsA in the closed-conductive configuration to test against the two existing models of ion conduction [33]. We semisynthesized a KcsA channel with a total of twelve $^{13}$C$^{18}$O heavy isotope labels in the filter, allowing us to spectroscopically isolate and probe those residues and determine the ion occupancies in the S1, S2, and S3 binding sites. From the 2D IR experiment, we were able to resolve two elongated features which we
attribute to different ion binding configurations and structural distributions. Analysis using MD simulations and 2D IR calculations reveal that the spectral features could only arise from ion binding configurations from the knock-on mechanism of conduction, effectively ruling out the hard-knock model as the exclusive mechanism for ion conduction. Our work also reveals that water cotranslates through the filter with the ions. We not only provide new experimental evidence that support the knock-on model of ion conduction but also present new spectroscopic signatures for a Val76-flipped state which has only been observed in simulations. Our second study focuses on the coupling between the selectivity filter and the helix bundle gate, inferred through changes in the ion occupancy of the filter under different experimental conditions [34]. Using the same techniques, we probed the ion occupancies in the selectivity filter of KcsA in the closed-constricted and the open-constricted states. Our experiments identify two spectral features which correspond to states with an ion in S1 and states without an ion in S1. The relative intensities of these two features are dependent on whether the intracellular gate is closed or open, suggesting that the state of the gate influences the relative populations of the two ion binding configurations. MD simulations and 2D IR calculations reveal that transitions of the intracellular gate leads to structural changes in the selectivity filter as observed through changes in the ion occupancy at the S1 binding site. This work highlights the molecular complexities in the allosteric coupling between the intracellular bundle gate and the selectivity filter.

3. Methods

3.1. Preparation of KcsA samples

The potassium ion channel in our study, KcsA, is semisynthesized through methods that have been previously utilized to incorporate non-natural amino acids into the selectivity filter [30,31]. This approach allows for the introduction of three $^{13}$C$^{18}$O isotope labels at the Val76, Gly77, and Gly79 positions, which enables us to specifically probe the S1, S2, and S3 binding sites in the selectivity filter by shifting these amide I vibrational modes away into a relatively clear region of the spectrum. The semisynthesis of the channel involves the chemical synthesis of the KcsA pore peptide (residues 70–81) with an N-terminal thiazolidine and a C-terminal thioester that incorporates the $^{13}$C$^{18}$O labels at the appropriate positions. Both the N-terminal and C-terminal flanking polypeptides (the thioester N-peptide and N-terminal Cys C-peptide, respectively) are recombinantly expressed in E. coli and purified using established methods [7,30,31,53]. Assembly of the KcsA polypeptide occurs over the course of two sequential native chemical ligation (NCL) reactions. The first NCL reaction yields an intermediate peptide of the pore peptide and the C-peptide. Subsequent deprotection of the intermediate and reaction with the N-peptide generates the KcsA polypeptide, which has been shown to refold in vitro into lipids or micelles to form functional KcsA channels. In these 2D IR studies, both the labeled and unlabeled channels are prepared at different pH and KCl salt concentrations in DM micelles, allowing us to experimentally modulate the conformation of the helix bundle gate and the selectivity filter. The sample conditions that we use are critical to capturing KcsA in its different conformational states. For the initial study where we probe the channel in the closed-conductive conformation, following purification and folding, the channel is resolubilized in 0.25% DM micelles in 50 mM Tris-HCl, pH 7.5, 150 mM KCl buffer. In our subsequent study, the low K$^+$ sample (1 mM KCl) was obtained by dialyzing the purified KcsA channel against 50 mM Tris-HCl and 1 mM KCl, giving us the closed-constricted conformation of the channel. Likewise, the open-constricted conformation was isolated by dialysis against 50 mM 1,4-diethylpiperazine (pH 4.0) and 150 mM KCl.
3.2. Spectroscopic measurements

2D IR spectroscopic measurements are carried out using rapid-scan 2D IR technology, which has been developed around the mid-IR pulse shaper [50–52]. The output of a Ti:Sapphire oscillator is amplified in a Nd:YLF pumped regenerative amplifier before it is sent into a β-barium borate optical parametric amplifier and AgGaS₂ difference frequency generation setup to produce the femtosecond mid-IR pulses that are essential for the 2D IR experiments. Using a ZnSe beamsplitter, a large fraction of the mid-IR beam is focused into a germanium acousto-optic modulator which facilitates the shaping of the two pump pulses by modulating both the phase and amplitude of incoming beam through computer-generated acoustic waves that propagate through the crystal. The two pump pulses are then spatially and temporally overlapped with the probe pulse at the sample to produce the 2D IR signal that propagates in the direction of the probe beam, which is directed into a spectrograph and measured by a single array mercury cadmium telluride detector. The mid-IR pulse shaper gives us the ability to generate these pulse pairs with each laser shot and to control relative delay between the two pump pulses \( t_1 \) electronically, which significantly reduces the data collection time. Furthermore, with the pulse shaper, we can implement phase cycling of the pump pulses to remove unwanted scatter, critical when working with biological samples in lipids and detergents.

3.3. MD simulations and 2D IR calculations

While the details of the MD simulations are provided elsewhere [33,34]. Briefly, for our initial studies, we performed MD simulations using Drude oscillator derived polarizable force fields starting from previously described equilibrated structures [38]. For each of knock-on and hard-knock simulation systems, 100 configurations were generated for each ion binding configuration, which were energy minimized in a GROMOS 53a6 force field that was conducive for calculating 2D IR spectra [43]. 20 ps MD trajectories were run for each of these configurations and sampled with a 2 fs integration time step. Ion configurations of interest were constrained to the filter using flat bottom potentials. The amide I 2D IR spectra are calculated using established methods [36,37,55]. Briefly, the frequencies are calculated from the MD simulations as projections of the electric fields on point charges and shifted by 66 cm\(^{-1}\) to match the experimental shifts observed for the \(^{13}\)C\(^{18}\)O isotope labels. Nearest-neighbor frequency shifts parameterized by Ramachandran angles and transition dipole coupling schemes were used to calculate the couplings between the amide I modes [36,54]. Following the parameterization of the frequencies and couplings, the third-order response functions are calculated which are used to generate the 2D IR spectra.

In our follow-up investigation, we use a slightly different method for the MD simulations of the closed-constricted and open-constricted states, which are discussed elsewhere [44]. Each of these 532 ns long MD simulations is run using a CHARMM36 force field. To maintain the open state, we took care to protonate the appropriate residues along the activation gate. All of the structures from the MD simulations show that the channel structure is conserved between lipids and micelles when compared to the X-ray crystal structures. From these 532 ns trajectories, the initial configurations for the open- and closed-constricted states were extracted every 250 ps, effectively giving us 2128 snapshots which are then used to generate the response functions for the 2D IR calculations, as described in the previous study. Hamiltonian clustering analysis is performed in the analysis of the 2D IR spectra. In this methodology, a parameter \( \Delta_{ij} \) is calculated to describe the couplings between Val76 and Gly77 for each snapshot. The \( \Delta_{ij} \) for each starting configuration is then sorted using a \( k \)-means clustering algorithm, allowing
Fig. 2. Comparison of experimental spectrum to calculated spectra of different ion permeation models. (A) Experimental spectrum shows two inhomogeneously broadened peaks. (B) Linear combination of the knock-on states (including a Val-76 flipped state) exactly reproduces the frequencies and lineshapes observed in the 2D IR experiment. (C) No linear combination of the hard-knock spectra could reproduce the experimental spectrum. The spectrum reveals a single pair of peaks that are homogeneous in lineshape. (D) No linear combination of the knock-on spectra without water could recapture the frequencies and lineshapes observed in the experiment, indicating that water must also be in the pore of the filter.

us to advantageously group the structures based on their observables rather than on defined structural parameters [4,5].

4. Results and discussion

Our initial study focused on the question of ion conduction through the selectivity filter [33]. In Fig. 2(A), we show the experimental 2D IR difference spectrum of KcsA (obtained by subtracting the unlabeled spectrum from the isotope labeled spectrum) compared to simulated spectra of the linear combination of knock-on states (Fig. 2(B)), the linear combination of hard-knock states (Fig. 2(C)), and the linear combination of knock-on states without water (Fig. 2(D)). In the experimental difference spectrum, we see two elongated peaks at $\omega_{\text{pump}} = 1603$ and 1580 cm$^{-1}$, which we attribute to a weighted average of different ion binding configurations and structural states of the protein. To link the experimental observables to the atomistic structures and ion binding configurations, we turn to MD simulations and 2D IR calculations. Simulations of the [W, S2, W, S4] and [S1, W, S3, W] states of the knock-on model reveal spectral features at 1608 and 1600 cm$^{-1}$, respectively. Taking a linear combination of the two states, we can exactly reproduce the frequency and the lineshape of the $\omega_{\text{pump}} = 1603$ cm$^{-1}$ peak in the experimental spectrum. Interestingly, from our analysis of the knock-on model, the $\omega_{\text{pump}} = 1580$ cm$^{-1}$ peak we observe in the experiment arises from what is called the Val76-flipped state of the [S1, W, S3, W] ion binding configuration. The Val76-flipped state, which has only been previously observed in MD simulations, occurs when a single Val76 residue in the tetramer flips 180° outside of the filter, away
from the conduction pathway [10,19]. The calculated 2D IR spectrum of the Val76-flipped state recaptures the experimentally observed lower frequency peak. By taking a linear combination of all of the knock-on states and the Val76-flipped state, we are able to reproduce the experimental 2D IR spectrum. When we simulate and calculate the 2D IR spectra for the states in the competing model, the spectra for each of the hard-knock states is surprisingly similar to each other. Figure 2(C) shows that the linear combination of the hard-knock states, which is in poor agreement to our experimental spectrum. The calculated spectrum of the hard-knock states reveals a single peak centered at a frequency that falls between the two peaks in our experimental spectrum. Furthermore, the peak is significantly more homogeneous (i.e. rounded) in lineshape relative to the experimental spectrum. Because no linear combination of the hard-knock states can recapitulate the experimentally observed frequencies and lineshapes, we conclude that our data is consistent with the knock-on model and rule out the hard-knock model as the exclusive mechanism of ion conduction. To test whether water cotranslates with the ions, we simulate the states in the knock-on model without the intervening waters. Notable changes to the spectra when we eliminate water from the filter are a blueshift of about 4 cm$^{-1}$ and a change in the lineshape of the spectrum for the [0, S2, 0, S4] state. Because of this, no linear combination of the knock-on states without water can reproduce the experimental spectra, which indicates that water must be present in the filter during conduction as implicated by the streaming potential measurements [1,27]. Our results not only provide new experimental evidence for the knock-on model of conduction, but also show that water and ions are conducted by the KcsA channel.

In our subsequent study, we probed the effects of gating on the ion occupancy in the selectivity filter [34]. Figures 3(A) and 3(B) shows the experimental difference spectra for the open-constricted and

![Fig. 3. Experimental and simulated 2D IR spectra of the open-constricted and closed-constricted states. (A) Experimental spectrum of the open-constricted state shows a dominant pair of high frequency peaks. (B) Experimental spectrum of the closed-constricted state shows a pair of low frequency peaks. (C)–(D) Simulated 2D IR spectrum of the open-constricted and closed constricted states, respectively. (E), (G) Hamiltonian clustering analysis of the open-constricted simulations reveal two main groups in which (E) an ion is absent from the S1 binding site and (G) an ion is present. The majority of the states lack an ion in the S1 binding site. (F), (H) Hamiltonian clustering analysis of the closed-constricted states reveal the same two groups: (F) an S1-unoccupied state and (H) an S1-occupied configuration. In the closed-constricted conformation, the majority of states favor the presence of the ion in the S1 binding site.](image-url)
the closed-constricted states of the channel. The spectrum for the open-constricted state reveals an intense pair of high frequency peaks at $\omega_{\text{pump}} = 1618$ cm$^{-1}$ while the spectrum of the closed-constricted state shows a single pair of low frequency peaks at $\omega_{\text{pump}} = 1599$ cm$^{-1}$. The results are unexpected because in both of these conformational states, the conformation of the selectivity filter does not change: The only difference between these two KcsA samples is the state of the intracellular gate. Again, we turn to MD simulations to determine the molecular origins of the two spectral features. We simulate and calculate the 2D IR spectra for the open-constricted and the close-constricted conformations from sampled 532 ns trajectories, and our calculated spectra are remarkably similar to the experimental spectra (Fig. 3(C) and 3(D)). The calculated spectrum for the open-constricted state reveals two pairs of peaks, with one set of intense peaks at $\omega_{\text{pump}} = 1617$ cm$^{-1}$. Likewise, the calculated spectrum for the closed-constricted state reproduces the experimental spectrum. To define the ion binding configurations that give rise to these spectral signatures, we perform Hamiltonian clustering analysis [34], in which we binned the sampled MD snapshots used to generate the spectra shown in Fig. 3(E) and 3(G) according to their vibrational Hamiltonian. By defining a quantity $\Delta_{ij}$ that describes the coupling between Val76 and Gly77 labels and calculating a $\Delta_{ij}$ score for each snapshot, we are able to group these structures according to their observables using a $k$-means clustering algorithm [41]. Thus, using this method, we show that two distinct structural groups emerge for the open-constricted state of KcsA. Figure 3(E) and 3(G) shows the two main groups that comprise the instantaneous snapshots in the MD trajectory: The first group, which represents 73% of the structures, has a 2D IR spectrum that is dominated by a high frequency peak at $\omega_{\text{pump}} = 1617$ cm$^{-1}$. 14% of the other structures fell into the second group which is characterized by a 2D IR spectrum of a low frequency peak at $\omega_{\text{pump}} = 1602$ cm$^{-1}$. Investigation of the structures that give rise to these spectra reveal that the high frequency peak comes from states in which the S1 binding site is empty and the low frequency peak comes from states in which the S1 binding site is occupied by a potassium ion. Based on the examination, in the open-constricted conformation, a majority of the states lack an ion in the S1 binding site. Likewise, for the closed-constricted state, we see a similar breakdown of into two subensembles with one spectrum exhibiting a pair of peaks at $\omega_{\text{pump}} = 1617$ cm$^{-1}$ and the other at 1602 cm$^{-1}$, but, interestingly, the populations of these states are reversed. In this case (Fig. 3(F) and 3(H)), we see that only 21% of the states do not have a K$^+$ ion in the S1 binding site, compared to the 73% we observed for the open-constricted state. In other words, in the closed-constricted state, there is a higher population of structures with an ion in the S1 binding site. Our analysis uncovers two spectral parameters that define the ion occupancy of the S1 site: peak frequency and intensity. The peak frequency tells us whether or not an ion is occupying the S1 binding site and the peak intensity tells us about the relative populations of the two states. Because of the remarkable agreement between simulations and experiments, we are able to extract additional information from the MD simulations. We show from our analysis that there are structural differences between the two selectivity filters when the ion is present or absent at the S1 site. When the ion is present, the filter is very rigid, which is contrary to what we observe when the ion is absent. In this case, the filter is more dynamic and flexible as measured by the deviations of the $\phi$ and $\psi$ angles about the Gly77 carbonyl. The disparity also implies that the state of the gate is somehow tied to these conformational differences. These structural differences led us to examine the structures of the two subensembles, which reveal two residues in the channel that may play a role in the coupling of the motions of the helix to the structural changes in the selectivity filter: Phe103 and Arg64. Although additional residues may be involved in the communication pathway between these two domains, only Phe103 and Arg64 showed substantial changes in distance to the filter between the closed and open states. Along with data from previous mutational studies [15,45], this finding leads us to postulate that these two residues play an important role in the
coupling between the helix and the filter. Our work effectively shows that the large structural motions of the intracellular gate also affect the conformation of the selectivity filter and alters its binding affinity for ions at the S1 binding site.

5. Conclusions

As illustrated in our two studies, using a combination of protein semisynthesis, 2D IR spectroscopy, and MD simulations, we were able to elucidate the ion occupancies in the selectivity filter of KcsA. Because of the ultrafast time resolution of the technique, we are able to get an instantaneous snapshot of the ensemble of ions, water, and protein structures. Furthermore, the sensitivity of vibrations to their local environment makes it an appropriate technique if we want to discern between two competing ion permeation models. In our initial studies, we provide new experimental evidence for the knock-on model of ion permeation. Our experimental 2D IR spectrum reveals two spectral features that can only be reproduced by structures and ion binding configurations of knock-on states where water and ions alternate through the filter. Additionally, we observe the Val76-flipped state, which was predicted to exist for the [S1, W, S3, W] from MD simulations. Furthermore, we show that none of the hard-knock ion binding configurations could recapture the two elongated peaks we observe experimentally, indicating that the hard-knock model is not the main mechanism for ion conduction. In our follow-up study, we probed the ion occupancies of the selectivity filter of KcsA under different experimental conditions. Using the different sample conditions, we were able to modulate both the conformational state of the inner gating helix and the selectivity filter, thus allowing us to examine the ion binding configurations of the closed-constricted and the open-constricted states. Our experimental spectra reveal two different spectra for these two states, which is intriguing given the distinction between these two states is whether the gate is open or closed. Further analysis using MD simulations and Hamiltonian clustering analysis indicates that the frequencies and intensities of the peaks describe the ion occupancy at the S1 binding site. We show that there is a greater population of states with an S1 ion in the closed-constricted state, which is contrary to what we observe in the open-constricted state where the majority of states lack an ion in the S1 binding site. Additional analysis suggests structural differences between the filters with and without K⁺ in the S1 binding site, implying that the state of the gate affects the binding affinity of the S1 binding site. Ultimately, we postulate that several residues like Phe103 and Arg64 outside of the selectivity filter couple the motions of the inner helix gate to the conformational changes of the filter. As we have shown in this work, the methodology we describe has been successfully implemented to address fundamental questions of ion permeation and gating in potassium ion channels. We anticipate that this combination of techniques will be a powerful set of tools in the investigations of functional mechanisms of other complex biomolecular systems.

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