Ion Trapping with Fast Response, Ion-Selective Microelectrodes Enhances Detection of Extracellular Ion Channel Gradients

Mark A. Messerli, Leon P. Collis and Peter J.S. Smith

BioCurrents Research Center
Cell Dynamics Program
Marine Biological Laboratory
Woods Hole, MA 02543

running title: ion channel mapping with ISMs
Abstract
Previously, functional mapping of channels has been achieved by measuring the passage of net charge and of specific ions with electrophysiological and intracellular fluorescence imaging techniques. However, functional mapping of ion channels using extracellular ion-selective microelectrodes has distinct advantages over the former methods. We have developed this method through measurement of extracellular K⁺ gradients due to efflux through Ca²⁺-activated K⁺ channels expressed in CHO cells. We report that electrodes constructed with short columns of a mechanically stable K⁺-selective liquid membrane, respond quickly and measure changes in local [K⁺] consistent with a diffusion model. When used in close proximity to the plasma membrane (< 4 μm) the ISMs pose a barrier to simple diffusion creating an ion trap. The ion trap amplifies the local change in [K⁺] without dramatically changing the rise or fall time of the [K⁺] profile. Measurement of extracellular K⁺ gradients from activated rSlo channels shows that rapid events, 10 –55 ms, can be characterized. This method provides a non-invasive means for functional mapping of channel location and density, as well as for characterizing the properties of ion channels in the plasma membrane.

Keywords: noninvasive ion-selective microelectrode, rSlo, single channel detection
Introduction

Plasma membrane ion channels are involved with maintaining ionic balance and controlling ionic changes for signaling pathways. Short-term blockade or inactivation of channels destroys signaling pathways while relatively long-term inactivation interferes with ionic homeostasis and cell viability (1). With the aid of voltage clamp methods, parameters such as conductance, permeability and mode of activation have been characterized for many different types of channels. However, current methods for the functional mapping of the location of ion channels on cells are tedious. There are at least three different types of methods that have been used to map the functional distribution of ion channels, 1) voltage clamp of channels using loose patch clamp (reviewed in 2); 2) scanning ion conductance (SICM) using whole cell voltage clamp with a scanning current passing electrode (3); 3) monitoring of intracellular fluorescent ion indicators (4, 5). The first two methods take advantage of the measurement of current due to the rapid movement of charged ions across the plasma membrane, although the measured current will be from any number of channels with different ionic permeabilities. Additional experimental manipulations are needed to obtain channel identity, including ion substitution and/or removal, addition of specific pharmacological blockers and identification of activation or reversal potentials. Loose patch-clamp methods are difficult to use on flattened morphologically distinct regions of cells compared to rounded-up regions, and generally are not practical on small cells (2). Ion substitution was used in conjunction with scanning ion conductance to functionally localize K\textsubscript{ATP} channels on cardiac myocytes. Specifically, K\textsuperscript{+} and other K\textsubscript{ATP} permeant ions were removed completely from both intracellular and extracellular solutions. K\textsuperscript{+} was supplied to the channel via a scanning extracellular pipette filled with 1M K\textsuperscript{+} (3). The success of this method was dependent on nonphysiological ionic conditions and whole cell voltage clamp. The third method, the use of fluorescent ion indicators to monitor ion concentration changes after passage through channels, allows long-term study of ion channels. However, it is at best restricted to Ca\textsuperscript{2+} and H\textsuperscript{+} influx through ion channels due to the low intracellular concentration of the ions as well as the high selectivity and sensitivity of their fluorescent indicators compared to other inorganic ion indicators (6).

Based on the insight given by these methods it became apparent that extracellular ion-selective microelectrodes (ISMs) could be used for functional mapping of ion channels and also for characterization of the channels. Extracellular ISMs have been used to measure steady and dynamic changes in extracellular ion gradients near single cells and tissues that arise as a result of ion passage across the plasma membrane via channels, transporters or exchangers (recently reviewed by 7, 8). Similar to fluorescent indicators, ISMs possess an inherent selectivity enabling measurement of channel activity for channels passing a specific ion in the presence of active channels passing other ions. However, they possess a greater signal to noise ratio (SNR) than their corresponding fluorescent ion indicators, are useful over a wider dynamic range and exist for a wide range of inorganic ions. Extracellular ISMs can be used in a scanning mode similar to the SICM and loose patch discussed above but do not contact the cell like loose patch and can be used under more physiologically relevant conditions than those used with SICM.

In this report we employed fast response, K\textsuperscript{+}-selective microelectrodes in order to capture rapid (10-50 ms duration) extracellular K\textsuperscript{+} transients that arise due to the activity of Ca\textsuperscript{2+}-activated K\textsuperscript{+} ion channels. We also report that ion trapping by the ISM increases the magnitude of the single channel gradients making it easier to detect and map the gradients.
**Materials and Methods**

**Electrode Construction**

Fast response ion-selective microelectrodes were fabricated by simple modifications to the previous design (9). The fast design K⁺-selective electrode incorporated a more mechanically stable liquid membrane that enabled construction with stable shorter columns, ≤ 10 μm. The K⁺-selective liquid membrane was made similar to the K⁺-selective mixture 60398, (Sigma-Aldrich (Fluka), St. Louis, MO), with the exception that 1,2-Dimethyl-3-nitrobenzene was replaced with 2-Nitrophenyl octyl ether (NPOE). The valinomycin-based liquid membranes have greater than 3 orders of magnitude selectivity for K⁺ over other common cations including Na⁺, H⁺, Ca²⁺ and Mg²⁺ (10). Microelectrode bodies for the liquid membrane were pulled from both borosilicate (WPI, Sarasota, FL) and aluminosilicate glass (Sutter Instruments, Novato, CA). Borosilicate (1.5/0.84 and 2.0/1.12 mm O.D./I.D.) and aluminosilicate (1.5/1.0 mm O.D./I.D.) glass capillaries were pulled to obtain tips of 2-3 μm inner diameter. This resulted in tip wall thickness of 200 and 300 nm for the 1.5 and 2.0 mm O.D. pipettes respectively. Stable ion-selective microelectrodes with short columns of ionophore were not always possible with borosilicate glass but were made more consistently with aluminosilicate glass. Pulled pipettes were silanized with the vapor method by first drying glass at 230-240°C for 20 min under vacuum 28 inHg. The evacuated oven was purged with argon and the electrodes were introduced to 20 μL of N,N-Dimethyltrimethylsilylamine (Sigma-Aldrich (Fluka), St. Louis, MO) in an enclosed metal box at 230-240°C for 20 min. After silanization, pipettes were stored in a bell jar with desiccant in the bottom. Electrical connection was made with a Ag/AgCl wire connected to a backfilling solution of 100 mM KCl, 10 mM HEPES pH 7.4.

Concentric ISMs were constructed according to previous designs (11, 12). Borosilicate glass with a filament (1.0/0.75 mm O.D./I.D.) was inserted and immobilized into a pulled and silanized outer borosilicate pipette (1.5/1.12 mm O.D./I.D.). Immobilization was achieved with a UV curing glue, Norland Optical Adhesive #74 (Norland Products Inc., Cranbury, N.J.). The tip of the inner pipette was within 10 μm of the tip of the outer pipette. The inner pipette was filled with backfilling solution before tip loading the liquid membrane into the outer pipette. The liquid membrane was equilibrated with 100 mM KCl before loading as previously described (13).

ISM measure ionic activity. However, we will instead refer to ISMs as measuring ionic concentration, as ionic activity is directly proportional to ion concentration, via the activity coefficient, and the changes that occur to the activity coefficient due to changes in ionic strength are negligible during our measurements. We reserve the term active or activity to discuss the state of the ion channels.

**Cell Maintenance and Handling**

Chinese Hamstery Ovary (CHO) cells stably expressing the Ca²⁺-activated K⁺ channel, rSlo, were maintained in Iscove’s Modified Dulbecco’s Medium with 10% FBS, HT supplement and 1% of 10K I.U. Penicillin Streptomycin (Invitrogen, Carlsbad, CA). During recordings, medium was replaced with a physiological saline containing (in mM) 140 NaCl, 5 CaCl₂, 29 glucose, 25 HEPES pH 7.2 with 0.3 KCl. Electrodes used for whole cell voltage clamp were filled with (in mM) 32.5 KCl, 97.5 K⁺ gluconate, 5 EGTA, 10 HEPES pH 7.2 with either 2.5 or 3.5 CaCl₂ resulting in a final free [Ca²⁺] of about 200 and 400 nM, respectively, calculated with WEBMAXC Standard (14) http://www.stanford.edu/~cpatton/maxc.html.
K⁺ gradients were measured near the plasma membrane of CHO cells held under whole cell voltage clamp with varying [Ca²⁺]. Iberiotoxin (300 nM) was added while clamping [Ca²⁺] to 400nM in order to capture single channel gradients at a range of applied voltages. Voltage clamp was performed with an Axoclamp 200B amplifier while K⁺ gradients were measured using an Axoclamp 2B amplifier with the 0.0001 H headstage. Additional amplification was provided by a Warner Instruments LPF-100B (Hamden, CT). An offset was applied to the ISM signal in order to apply gain and maintain the signal within the dynamic range of the LPF-100B amplifier.

**Ion Gradient Modeling and Analysis**

**Construction of a realistic ISM**

In order to assess the extent to which an ISM can affect the diffusion profile of K⁺ ions, a 3D approach to modeling the movements of ions was undertaken. First it was necessary to incorporate a 3D representation of a typical ISM into the model environment. It was only necessary to incorporate a 10 µm length of the ISM from the tip, as we were analyzing the diffusion of ions from a single ion channel over millisecond time periods. Typical dimensions of tip diameter and taper angle were extracted from scanning electron micrographs of ISMs and used to construct a 3D model using the open-source 3D generation suite, Blender (v. 2.45). This software was run on a Linux platform (Fedora release 8) on a HP xw4400 workstation with 2GB RAM and a 2.1 GHz processor. ISMs were translated and rotated within the 3D environment in order to assess the effect of orientation and separation distance on the diffusion profile of ions.

**Simulation of an ion channel**

We used a Monte-Carlo method for simulating the Brownian motion of ions from a point source in a 3D environment. For this purpose, the Model Description Language (MDL) used within MCell (version 3.0; (15, 16)) was used to simulate the diffusion and random walk of K⁺ ions from a point source (a 270 pS K⁺ channel), with a diffusion coefficient of 1.96 × 10⁻⁵ cm² s⁻¹. In order to decrease simulation time, the extracellular background concentration of K⁺ was left at zero. Our graphs report the change to the local [K⁺]. The code was written and executed using the same platform described above. Simulation time was optimized by partitioning the 3D environment into 0.2 µm cubes and by increasing the time-step to 10 µs. While reducing the simulation time significantly, no significant change in diffusion profile was observed by optimizing these parameters (data not shown). The simulation generated ions at a rate of 6.75 × 10⁷ s⁻¹ from a single point source. In order to further reduce simulation time, a 10 µm boundary box enclosed the simulation and any ion that contacted the box was deemed to have left the simulation space and removed from the simulation. This had no effect on diffusion profiles sampled at ≤3µm from the point source (data not shown). The boundary box was omitted for the diffusion profiles collected at 10 µm from the source. After a set channel open time (10-50 ms) the simulation was stopped and simulation data was checkpointed to a second simulation program where the ion channel was closed.

Ion concentration was assessed using MCell sample boxes to count ions. Sample boxes were generated using Blender and consisted of either a 100 nm thick disk covering the aperture of the ISM, or a 100 nm diameter sphere at the center of the ISM aperture. The disk sample box was a reflection of the actual ion concentration that the ISM would detect and thus convert to a corresponding change in the electrode's Nernstian voltage. However, ion concentration was
dependent on the orientation of the cylinder. To eliminate the angle of the sample box as a variable, a sphere was used at varying distances from the point source to calculate ion concentration. The separation distance between the sample boxes and the point source was adjusted at incrementing distances from 0.2 µm to 10 µm. To reduce simulation noise and obtain an average of simulation trials, the software was executed with different initial seed values that generated different sequences of "pseudorandom" numbers for the Monte-Carlo simulation.

Visualization of MCell simulation

To validate and verify the output of the MCell simulation it was necessary to generate 3D renderings of the meshes (ISM and sample boxes) and particles (ions). MCell generated visualization files for use in the open-source 3D render software, DReAMM (version 3.3.). All simulation images in this manuscript were generated with DReAMM.

Mean open times were measured from single channel gradients by hand using the cursors in Clampfit (Molecular Devices, Sunnyvale, CA). These values were used to calculate NPo, the product of the number of channels (N) and the open probability (Po), at different voltages by summing the time of the events and dividing by the time range used to collect the events. NPo was calculated from transmembrane currents according to eq.(8) in (17). Best-fit alignment of single channel gradients was performed as described earlier (18) with incorporation of the ion-trap correction factor determined in this work.
Results

Response times of K⁺-selective microelectrodes were determined by measuring a controlled transient change in [K⁺]. Current injection (20 pA) through one K⁺-selective microelectrode (source) was used to generate a step rise in [K⁺] that was measured with a second K⁺-selective microelectrode, illustrated in Figure 1A. This method enables rapid generation of a K⁺ gradient of similar magnitude to those generated by the ion channels in this study. The noise and drift associated with flow systems prevented their use for such small changes in [K⁺]. Figure 2A shows the voltage recorded by the second K⁺-selective microelectrode at increasing distance from the source electrode. The profiles contain the electrical transient of the applied current step summed with the measured [K⁺]-dependent voltage change. The magnitude of the electrical transient was independent of the distance between the source and measuring electrode and was therefore subtracted to reveal only the [K⁺]-dependent voltage measurements in Figure 2B. The measuring electrode came from a batch of electrodes with inner tip diameters between 2.5-2.8 μm and was filled with a 30 μm column of K⁺-selective mixture made with NPOE. [K⁺] profiles are evident within a few microns from the source electrode but are barely detected at 11.5 μm from the source. The slopes of the rise times and amplitudes of the [K⁺]-dependent voltage decrease with increasing distance from the source electrode, consistent with diffusion of K⁺ away from the source electrode. We calculate a 95% response time (t₉₅%) of 27 ± 1 ms for this ISM based on the measured [K⁺] profiles.

Detection of Single Channel Ion Gradients

Extracellular changes in [K⁺] were monitored next to CHO cells overexpressing rSlo channels as diagrammed in Figure 1B. Gating of this type of channel is dependent on membrane voltage, pHi and [Ca²⁺]i (19-22). Under the conditions we used, channels require ≥ +40 mV transmembrane potential to open with 200 nM [Ca²⁺]i, but will open at 0 mV in the presence of 400 nM [Ca²⁺]i. Once successful whole cell voltage clamp was achieved the K⁺-selective microelectrode was positioned next to the edge of the cell as close as possible without touching the liquid membrane of the ISM to the plasma membrane of the cell. Measurements were obtained from cells still stuck relatively flat to the dish and on cells that had rounded up slightly. With this setup, depolarization of the plasma membrane created measurable extracellular K⁺ gradients in 9 of 14 cells. Failed measurements were due to either noisy ISMs in which the liquid membrane was not stable within the micropipette or when the ISM became physically blocked by debris in the culture dish. Positioning of the ISM near the membrane without touching was also a problem. Tapping the patched cell with the liquid membrane caused an instant efflux of K⁺ and tapping the patched cell with the side of the ISM broke the high resistance seal of the patch electrode.

When 200 nM [Ca²⁺], was used, a basal increase in K⁺ was measured along with very few single channel gradients even at +60 mV. Figure 3 shows a recording from a cell clamped with 200 nM [Ca²⁺]i, first depolarized to +60 mV and then polarized to −60 mV, showing the difference in extracellular [K⁺] between a membrane with active and then inactive channels. The basal rise in [K⁺] is in part due to the efflux of K⁺ through activated channels but may also be due to K⁺ efflux through transporters. Superimposed on the steady increase in [K⁺]-dependent voltage are large amplitude spikes due to higher localized efflux of K⁺ through the rSlo channels, which are not seen at the −60 mV holding potential when channels are not open. The single channel gradients were used to calculate a mean open time of 16.9 ± 10.3 ms (n = 65 events) for the cell shown in Figure 3.
With 400 nM [Ca$^{2+}$], the ISM measured large basal increases in [K$^+$] next to the cell >8.5 mV at +60mV, due to the high over expression and combined activation of many of these channels. Single channel gradients were not easily resolved under these conditions. Iberiotoxin was added to a final concentration of 300 nM to knockout a large fraction of these channels in order to capture single channel gradients over a range of holding potentials. Figure 4A, B and C show representative [K$^+$]-dependent voltage changes measured with an ISM near the same CHO cell, clamped to 0, +20 and +60 mV respectively. The cell remained clamped at each potential for the duration of each recording. The ISM was moved in a stop and hold mode with a period of 20s, alternating between a position near the cell and then 20 μm further away. This gave rise to the step function in the recording, measuring a greater [K$^+$]-dependent voltage near the cell. Similar to Figure 3, large amplitude spikes are superimposed on the steady increase in [K$^+$]-dependent voltage only when the ISM is near the cell. The frequency and mean amplitude of the spikes increases with increasing applied voltage as expected from the nature of these channels. The single channel ion gradients are sufficiently small in the away position that they are buried within the noise. Figure 4D, E and F show expansion of the regions under the gray bars in Figure 4A, B and C, respectively. The recording in the near position is at the same region of the cell. With low channel activity, gradients from single channels can be discretized, Figure 4 D, E. However with higher activity, K$^+$ gradients from different channel events sum together giving rise to much larger [K$^+$]-dependent voltages as shown in Figure 4F. The inset to Figure 4F shows expansion of a broad spike under the gray bar in Figure 4F, indicating peaks from 3 K$^+$ gradients or channel events. We use the term channel events because currently we cannot resolve whether the 3 ion gradients originated from the same channel firing rapidly in succession or from multiple channels. The use of multiple ISMs in close proximity could give better positional information and help resolve this issue.

The single channel gradients possess the information to enable calculation of mean open times and open probabilities. The mean open times for the events from the cell shown in Figure 4 are 33.5 ± 7.9, 42.3 ± 18.4 and 32.6 ± 12.7 ms when the membrane potential was held at 0, 20 and 60 mV respectively. NPo values can be determined from the ion gradient measurements as well, but only reveal the value for a small patch of plasma membrane while NPo values from transmembrane currents represent channels over the entire membrane. A direct comparison of these values requires normalizing the ISM measured region to the entire surface area of the membrane. The calculated NPo values from transmembrane currents are 0.1, 1.3 and 4.3 for 0, 20 and 60 mV, respectively. The corrected NPo values determined from the ion gradient measurements are similar, 0.2, 1.3 and 4.2 at 0, 20 and 60 mV, respectively, assuming a 3 μm diameter measuring region for the ISM and a cell diameter of 10 μm, revealing the voltage-activated nature of these channels.

**Modeling of Ion Trapping by ISMs**

Best-fit alignment of single channel gradients was performed using a simple diffusion model and taking into account electrochemical driving force, channel open time, distance of the K$^+$-selective microelectrode from the channel and response time of the measuring system as presented earlier (18). However, these parameters were not sufficient to account for the [K$^+$] profiles that we measured with fast response microelectrodes. Specifically, the simple diffusion model was undershooting the magnitude of the measured profiles by about 2-fold. By taking into account that the microelectrode is acting as a significant barrier to diffusion along with its close proximity to the plasma membrane, we investigated the possibility that the microelectrode
is acting as an ion trap. Figure 5 displays an illustration of the physical model and representative changes to \([K^+]\) profiles when an ideal, standard sized ISM is positioned such that the center of the active surface is 1.5 \(\mu m\) away from the ion channel / point source. The \([K^+]\) has been determined within a small spherical region of interest shown in Figure 5B-E in order to closely compare the extent of ion trapping for an ISM at 3 different angles, when the sensing surface of the ISM is parallel, at a 45° angle and perpendicular to the plane of the plasma membrane, Figure 5C, D and E, respectively. Modeling of the small spherical region was used to remove the changes in \([K^+]\) that would occur due to the 3 different orientations of the ISM while measuring in an ion gradient. Ion trapping leads to modifications in the measured profile, as displayed in Figure 5F, including steeper rises and falls and higher peak concentration. The ion trap is greatest when the active surface of the ISM is both parallel and at a 45° angle to the plasma membrane. The ion trap enhances the magnitude to a lesser extent when the ISM surface is perpendicular to the plasma membrane when compared to diffusion in the absence of the ISM.

In order to determine the more practical influence of ion trapping on measurements we determined the change in concentration for a thin disk (100 nm) near the active surface of the ISM. The concentration profiles for the thin disk were calculated in the presence and absence of the ISM and a ratio of the two was calculated to assess the influence of ion-trapping. Figure 6A shows the concentration profiles when the plane of the sensing disk is parallel to the surface of the membrane for two separation distances, 0.3 and 1.0 \(\mu m\). The ratio of the profiles (presence of ISM / absence of ISM) nearly follows a step function except for a small, brief (< 2 ms) overshoot that occurs at the beginning of event (data not shown). The magnitude of this step function is plotted in Figure 6B for the 3 different configurations of the ISM surface with respect to the plasma membrane surface. The ratios are plotted against the separation distance between the plasma membrane surface to the center of the active surface of the ISM. The ion trap is greatest when the active surface of the ISM is parallel and at a 45° angle to the plasma membrane surface.

These ratio values were incorporated into the simple diffusion model used in (18) to perform the best-fit alignments of single channel \(K^+\) gradients from two different cells. The alignments were performed by taking into account the driving force and background extracellular \([K^+]\) while varying distance of the channel from the ISM, open time and time constant of the measuring system. Figure 7 shows representative alignments of two channel gradients acquired under very different circumstances. The channel gradients in Figure 7A and B were acquired from cells clamped at 0 mV with 400 nM \([Ca^{2+}]_i\), and clamped at +60 mV with 200 nM \([Ca^{2+}]_i\), respectively. Best-fit alignment of the ion gradient in Figure 7A indicates that the time constant of the ISM was relatively slow (65 ms) and that the channel was 0.72 \(\mu m\) from the center of the ISM and was open for 30 ms. Best-fit alignment of a measurement made with a concentric ISM shows a faster time constant (29 ms) and reports that the channel was 1.03 \(\mu m\) from the center of the ISM and was open for 15 ms.
Discussion

In order to map ion channels with brief open times it is critical to use ISMs with short response times. Long response times are primarily due to the RC filtering that occurs from the ISM and the electronics (23, 24). Large inner tip diameters and shorter columns of ion-selective mixture produce lower resistance electrodes (23, 25) thus reducing response times. We have employed these considerations by making microelectrodes with 2-3 μm inner diameters and with column lengths of 30 μm for Ca\(^{2+}\) and H\(^+\) electrodes, and 100 μm for K\(^+\) electrodes (8). While lowered resistance has shortened response times, parasitic capacitances remain. By increasing the thickness of the glass pipette and using a more mechanically stable K\(^+\)-selective mixture we were able to produce ISMs with response times less than 30 ms with a 30 μm column. Shortening the column further produces electrodes with even faster response times but leads to less stable electrodes. A unique, more complicated design to reduce column length also has the side effect of reducing capacitance. Instead of simply placing a backfilling solution behind the ion-selective mixture, a second inner pipette, filled with electrolyte, is run up near the tip. This design produced K\(^+\)-selective ISMs with time constants of 1.5 – 7 ms for 3 different ion-selective mixtures (11). This method has recently been used to produce ISMs with response times less than 10 – 20 ms for Ca\(^{2+}\) and H\(^+\) electrodes, respectively, in the absence of any significant interfering ions (12). Mathematical methods such as deconvolution can also be employed to correct the recording for the relatively slow response times of the ISMs (26).

Functional mapping of ion channels with ISMs requires that the channels are active and that the electrochemical driving force for the permeant ions is not at equilibrium. Larger extracellular K\(^+\) gradients will be generated when the electrochemical potential is further away from the K\(^+\) equilibrium potential. Here we used voltage clamp to depolarize the plasma membrane while controlling cytosolic [Ca\(^{2+}\)] to activate the rSlo channels at less positive depolarizing potentials. By controlling the electrochemical driving force and knowing the channel conductance we were able to measure single channel gradients in a controlled manner. The information contained within the mapped ion gradients enabled relative localization of active rSlo channels and equivalent determination of NPo compared to the Npo determined from the transmembrane currents. However, the slow response time of the ISM prevented detection of events shorter than about 10 ms leading to overestimation of mean open time by 3-5 fold, when compared to earlier reports (22). This selection for longer events was also evident in Figure 4 as the mean open time did not increase with increasing applied voltage.

The ISM was moved in a step and hold mode, translating between a position close to the cell and further away. This method, known as self-referencing, enables characterization of the noise of the system so that it can be removed from the noisy signal. In conjunction with spectral filtering, self-referencing can be used to increase the signal to noise ratio by subtracting the average power spectrum of the noise from the signal of interest. Spatial self-referencing would be useful for measurements on cells and tissues ex vivo where there is extra space to move the ISM and characterize the system noise. Temporal self-referencing would be more useful on electrically excitable cells where there are intermittent signals, allowing for the collection of noise in between signal.

The ion gradients measured here with fast response ISMs did not fit a simple diffusion model as reported earlier for slow response ISMs (18). The primary difference is that the rapid response ISMs, record a more accurate [K\(^+\)] profile leaving less room for flexibility in the modeling parameters. The ion trap introduces amplification of the small, brief events that other ion mapping techniques do not possess. Not only should this enhance characterization of
channels that generate extracellular gradients through efflux, but it should have a similar effect of enhancing characterization of channels that generate inverted gradients due to ionic influx, such as Na\(^+\) and Ca\(^{2+}\) channels. While ion trapping makes it easier to detect single channel gradients it poses a potential artifact for measurements where simple diffusion is assumed to predominate. Specifically, ion trapping could lead to an overestimate of ion flux that is calculated when performing self-referencing of ISMs. By using the electrode in a configuration where the active surface of the ISM is perpendicular to the membrane surface, this overestimate can be minimized.

In future works it may be important to determine the exact position of the channel with respect to the position of the ISM(s). This will provide sufficient information to enable the identification of the channel conductance and conductance changes without previous knowledge of channel characteristics. Currently the best placement of an ion channel that we can perform with a single electrode will put the channel somewhere along the surface of a hemisphere with the ISM at the center of its radius of curvature. If the plane of the plasma membrane is known, the hemisphere reduces to the circle or oval where the hemisphere and plane intersect. This can be achieved with a double-barreled microelectrode where one barrel serves as the ISM and the other (an open barrel) is used to map distance to the membrane with impedance feedback. A more complicated multi-barrel design employs two ISMs to measure the gradient, and a third open barrel for impedance feedback. Information collected from this triple barreled configuration would enable placement of the channel along the line of intersection of the two ISM measuring hemispheres, a semi-circle, intersected by the plasma membrane plane generated by the impedance feedback microelectrode. This reduces the location of the channel to 2 possible points. Careful modeling must be used to determine the level of distortion that a multibarreled electrode would impose to the diffusion gradient. This mode of detection could prove very useful when mapping channels of similar ionic permeability but different conductance.

The method outlined here will work best on electrically excitable cells and tissues where relatively large signals would occur for even brief channel open times as the membrane potential changes over a wide range. Channel mapping and characterization could occur through the many forms of extracellular or intracellular electrical stimulation as well as chemical and mechanical activation. Continuous mixing on the outside of the cells must be avoided, as it will disrupt the ion gradient. The method is also suited for use on cells with extracellular layers that do not significantly interfere with diffusion of the ions. We were able to functionally map mSlo channels on *Xenopus* oocytes that were still enclosed in the coelomic envelope (18).

**Acknowledgements**

We thank Dr. L. Kaczmarek (Yale University, New Haven, CT) for providing the CHO cells stably expressing the rSlo channel. We also thank Joel Stiles (Carnegie Mellon University, Pittsburgh, PA) for providing and supporting the use of the modeling software, supported by NIH P41 RR06009 and RO1 GM068630. This research was primarily funded by NIH:NCRR grant P41 RR001395 to PJSS.
References

**Figure Legends**

Figure 1. Diagrams showing the use of K⁺-selective microelectrodes. A) Current clamp of a K⁺-selective microelectrode was used to generate a point source for an ion gradient of similar magnitude to those expected from single Ca²⁺-activated K⁺ channels. [K⁺] profiles were collected by the measuring K⁺-selective microelectrode at increasing distance from the source. B) Whole cell voltage clamp was used to control the cytosolic [Ca²⁺] while depolarizing the membrane in order to activate the rSlo channels in a controlled manner. The K⁺-selective microelectrode was moved in a stop and hold mode, coming to rest for ≤ 10 s in a position near the cell and also in a position 20 μm away from the cell.

Figure 2. [K⁺] profiles measured at increasing distance from a current-clamped source K⁺-selective electrode. A) Raw voltage measurement from a K⁺-selective microelectrode that contains the [K⁺]-dependent voltage summed with a transient charge across the return electrode. The transient charge across the return electrode was independent of position in the bath and was subtracted away to generate the [K⁺]-dependent voltages alone as shown in B. B) The amplitude and rising and falling slopes of the [K⁺] profiles decrease with increasing distance from the source electrode.

Figure 3. Measurement of extracellular [K⁺] near the surface of a CHO cell overexpressing rSlo channels at both depolarizing and resting potentials. Depolarization was used to activate rSlo channels giving rise to a slightly higher background [K⁺] superimposed with single channel gradients that appear as spikes. Voltage clamping the membrane back to –60 mV lead to inactivation of rSlo channels, a reduction in background [K⁺] and removal of single channel [K⁺] gradients. Breaking the GΩ seal allowed K⁺ to flood out of the patch electrode leading to an immediate rise in [K⁺]-dependent voltage measured by the ISM.

Figure 4. Extracellular [K⁺] measurements near activated rSlo channels overexpressed in a CHO cell. A fast response K⁺-selective microelectrode was repeatedly positioned at a position near the cell for 10s and then at a position 20 μm away from the cell for 10 s. The cell was voltage clamped at 0, +20 and +60 mV, A, B and C respectively, in order to activate more channels. K⁺ gradients from single channels (spikes) are only captured when the electrode is in the near position. D, E and F show expansion of regions in A, B and C respectively, that have been highlighted with an overlying gray bar. Single channel gradients are easily detected at lower depolarizing potentials when fewer channels are active (D, E). Single channel gradients sum together producing larger [K⁺]-dependent voltages, when greater channel densities are activated (F). The inset to F shows expansion of large amplitude event that is the sum of at least 3 single channel gradients.

Figure 5. Diagram of the working ion trap model and resultant [K⁺] profiles for different configurations of the K⁺-selective microelectrode. A) Expanded view of the rendered microelectrode near the cell surface during the middle of K⁺ efflux through an open channel. B) Side view of the small sampling region (white sphere, expanded in inset) used to determine the [K⁺] profile in the absence of the microelectrode. C, D and E show side views of the sampling region when the microelectrode active surface is parallel, at a 45° angle and perpendicular to the plasma membrane surface. F) The calculated [K⁺]-profiles for the different configurations of the microelectrode indicate the highest amount of ion trapping when the active surface of the microelectrode is parallel and at a 45° angle to the cell surface.

Figure 6. Determination of the extent of ion trapping for different microelectrode configurations with distance from the plasma membrane. A) Theoretical [K⁺] profiles in the presence
and absence of the microelectrode when the microelectrode surface is parallel to the plasma membrane at 0.3 and 1 μm away from the plasma membrane. B) Summary plot of the theoretical correction factors for different configurations of microelectrode at increasing distance from the plasma membrane.

Figure 7. Best-fit alignment of theoretical [K⁺]-dependent voltage profiles with single channel gradient voltage measurements. The best-fit alignment is overlaid on the recording to enable direct comparison. The model incorporated both simple diffusion and the distance-dependent correction factor due to ion trapping.
Figure 1
Figure 2

(A) 

(B)
Figure 3

whole cell clamp changed from +60 mV to -60 mV
broke seal
Figure 4

A. 0 mV
B. +20 mV
C. +60 mV
D. 
E. 
F. 
Figure 5
Figure 6

A

B
Figure 7

A

[B] - Measured
---
B - Best-fit Alignment

0 mV holding potential

-0.2 0 0.2 0.4 0.6 0.8 1 1.2 1.4

Time (ms)

0 50 100 150 200 250 300 350 400 450 500

[K+] -dependent Voltage (mV)

+60 mV holding potential

B

-0.2 0 0.2 0.4 0.6 0.8 1 1.2 1.4

Time (ms)

0 50 100 150 200 250 300 350 400 450 500

[K+] -dependent Voltage (mV)