Potentiometric-scanning ion conductance microscopy for measurement at tight junctions

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Scanning Ion Conductance Microscopy (SICM) has been developed originally for high-resolution imaging of topographic features. Recently, we have described a hybrid voltage scanning mode of SICM, termed Potentiometric-SICM (P-SICM) for recording transmembrane ionic conductance at specific nanostructures of synthetic and biological interfaces. With this technique, paracellular conductance through the tight junction – a subcellular structure that has been difficult to interrogate previously – has been realized. P-SICM utilizes a dual-barrel pipet to differentiate paracellular from transcellular transport processes with nanoscale spatial resolution. The unique combination of voltage scanning and topographic imaging enables P-SICM to capture paracellular conductance within a nominal radius of several hundred nanometers. This review summarizes recent advances in paracellular conductance recording with an emphasis on the P-SICM based approach, which is applied to detect claudin-2 mediated permeability changes at the tight junction.

Introduction

Movement of ions, molecules and solutes across the epithelium is an intricate and well-controlled process. Two primary avenues of transport exist, energy-dependent transcellular pathways or passive paracellular pathways.1-3 The cellular organelle known as the tight junction is responsible for formation of a paracellular barrier between the apical and basal compartments.4 Because the paracellular pathway is located at the cell-cell boundary, common techniques (e.g., patch-clamp) that rely on formation of a gigaohm seal between the pipet and cell membrane are not feasible. Alternate approaches to the study of paracellular transport are thus required. Most common approaches interrogate an area-averaged response of thousands, or even millions of channel conductances, with current density reaching μA/cm². Potentiometric-Scanning Ion Conductance Microscopy (P-SICM) (Fig. 1), promises to provide a method to record paracellular conductance at subcellular resolution, which will advance understanding of micro- and nanoscale heterogeneity in transport at paracellular pathways.5

Biophysical Techniques for Macroscopic Epithelial Studies

Traditional transepithelial electrical resistance (TER) measurements in conjunction with molecular biology techniques are used widely to inform the present understanding of transport at tissue interfaces. These techniques, which include the well-established Ussing chamber setup, provide an excellent opportunity to study transepithelial ion transport and ion permeability.6-12 However, these measurements represent the aggregate response of thousands of transport events across the epithelium, which can obfuscate studies of unique transport processes. A number of approaches have been developed to separate components of conductance across cellular barriers.13 For instance, to allow epithelial resistance measurements independent of capacitance, one-path impedance spectroscopy was applied to differentiate capacitive components, which exist only at the epithelial cell membrane and are dependent on the frequency of externally applied electric stimuli, such as transepithelial potentials.14-16 To further differentiate between trans- and paracellular resistances, a two-path impedance method has also been described, which combines impedance measurements of Ca²⁺ dependent tight junction openings and flux measurements with paracellular specific permeable markers.17-19 However, these impedance recordings still reflect the average response from trans- or paracellular pathways over an area that is at cm² scale.

Conductance Scanning

Conductance scanning (also voltage scanning) employs a scanning electrode to measure local variations in current density close to a cell surface and has been applied to determine paracellular conductance. In 1972, Frömter first introduced the principle for conductance scanning, where heterogeneity in the electric field above different conductive pathways of an epithelium was observed.20,21 Continuing efforts by Fromm22-28 and Cereijido29,30 dramatically improved measurements of localized trans- and paracellular conductance over epithelia. In these recordings, one pair of electrodes was used to apply a transepithelial potential across a cell monolayer, while a second pair of electrodes was...
used to measure the resultant changes in electric field at discrete locations above the sample. The electric field above the cell layer was measured by the scanning electrode (microcapillary) at two depths (close to and far away from the surface).\textsuperscript{22,23,32} or by a pair of pipets mounted in fixed positions separated by 30–50 μm.\textsuperscript{13}

Local conductance was then be calculated from Equation 1,\textsuperscript{23,32}

\[
G = \frac{E}{\rho \cdot V^e}
\]  

Equation 1

where \(E\) is the electrical field (the potential difference measured at two discrete points relative to the vertical movement of pipet, or pipet separation distance), \(\rho\) is the bath solution resistivity and \(V^e\) is the transepithelial potential. These techniques are very useful and provide unique vistas of epithelial conductance changes, but a major technical challenge remains – more precise spatial positioning of pipets is required to improve the accuracy of recorded changes in the electric field \(E\). In these recordings, an optical microscope was utilized to position pipets over cell junctions or cell bodies with the help of a micromanipulator. Vertical position was determined when the pipet first touched the cell surface as a virtual zero and then the pipet was manually withdrawn from the surface. For these types of measurements, precision in pipet positioning was influenced highly by operator’s judgment and samples could be damaged during pipet manipulation (samples with visible damage were subsequently discarded).\textsuperscript{23}

**Scanning Ion Conductance Microscopy**

Scanning ion conductance microscopy (SICM) is a novel technique based on scanning probe microscopy (SPM), which raster-scans non-conductive samples to record ion currents and generate topographic images. First introduced by Hansma and colleagues in 1989,\textsuperscript{34} SICM utilizes a hollow pipet with nanoscale tip dimensions as a probe to image samples. Pipets are pulled from capillaries with methods commonly used in patch-clamp studies. Through the manipulation of pulling parameters such as temperature, pull velocity, and delay, pipets with a variety of tip diameters and geometries can be fabricated. After fabrication, the pipet is filled with electrolyte and a silver/silver chloride electrode is back inserted. A potential difference is applied between the pipet electrode and a reference electrode placed in the bath (PE) in one barrel detects current passing between PE and a reference and pipet electrodes through the pipet tip opening. When the pipet is moved close to a surface, ion current changes according to the pipet-surface distance. Thus the pipet position relative to a surface can be controlled precisely with a piezoelectric positioner. This phenomenon forms the basis of current feedback regulation in SICM and allows SICM to dynamically scan surfaces in a non-invasive fashion. Ion current images can then be generated from the current measured, while the recording of vertical probe displacement generates topographic images.\textsuperscript{35}

SICM is an attractive tool for biological imaging because of the high spatial resolution afforded. SICM resolution can be as low as ~10 nm vertically and ~50 nm laterally,\textsuperscript{36} but this value is highly dependent upon pipet tip geometry as well as probe-surface distance. A limiting factor in SICM resolution is pipet tip diameter, as finite element simulations have shown these two parameters are directly proportional.\textsuperscript{37} Additionally, probe-sample distance is an important consideration to achieve optimal SICM resolution. Simulations reveal the observed ion current drop over a sampling distance is a function of tip geometry, cone angle, and inner-to-outer diameter ratio.\textsuperscript{38} The best recorded lateral resolution is 3–6 nm, which was achieved on S-layer proteins from *Bacillus sphaericus* with a 13 nm inner diameter pipet.\textsuperscript{39} An example of high resolution SICM topography image is shown in Figure 2. Here, a Madin-Darby canine kidney (MDCK) epithelial monolayer is shown. Cell bodies, cell junctions, and microvilli are clearly observed and differentiated.

In comparison to other SPM techniques, SICM has several distinct advantages for biological applications. As previously described, SICM is a non-invasive method, which generates topographic images of live cells with high fidelity and under physiological conditions. Compared with atomic force microscopy (AFM), chemical fixation of biological samples is not required in SICM, and thus deformation of cellular structures and loss of biological activity can be avoided. Unlike scanning electrochemical microscopy (SECM), SICM can image biological samples without the use of a redox mediator, which is advantageous for live cell measurement under normal physiological conditions. SICM also has a more robust feedback system than most other SPM techniques, which provides greater control over the pipet position, an important factor for conductance measurements (vide infra).\textsuperscript{35,40–43}

SICM has undergone several significant modifications since 1989. A number of hybrid-SICM platforms have been described for enhanced measurements. Our group has developed 3 and 4-electrode systems specifically for measurements of biological samples.\textsuperscript{44,45} The 3-electrode system places an electrode beneath a porous substrate (e.g., membrane or tissue). This modification allows for more precise control of transmembrane potentials. The 4-electrode system adds a platinum counter electrode above the membrane. The additional electrode is beneficial for studies of biological samples with multiple conductances present on the apical surface. These modifications have paved the way for the development of P-SICM for conductance recording of tight junctions.

**Potentiometric-Scanning Ion Conductance Microscopy (P-SICM)**

To incorporate voltage scanning into SICM, a dual-barrel probe design was utilized (Fig. 1), where the first SICM barrel was for imaging and positioning of the pipet; the second barrel was for recording the potential variations over cell bodies or cell junctions (Fig. 1).\textsuperscript{5} In a perfusion cell culture system (e.g., Transwell) where epithelial cells grow to form a monolayer, the dual-barrel pipet is placed into the upper chamber. The current electrode (PE) in one barrel detects current passing between PE and a reference electrode (RE) for pipet positioning, serving as the common reference and pipet electrodes through the pipet tip opening.
The potential probe (UE) in the second barrel serves to measure localized potential differences at cell surfaces relative to the RE. Potential differences at UE are generated by a transepithelial potential \((V_T)\) between RE and a working electrode (WE, bottom chamber) over heterogeneous conductive pathways within the recorded sample.

To evaluate local conductance change with P-SICM, potential deflections at the pipet tip were measured at two fixed probe-surface distances \((D_p)\), which were precisely controlled by a piezoelectric positioner. One position is close to the surface (0.2 \(\mu\)m), which is on the order of the dimension of the pipet tip (50 \(\mu\)m inner diameter for each barrel) and is controlled through the robust feedback signal to maintain a constant \(D_p\) (detailed SICM feedback mechanism can be found in reviews: refs. 35, 43). Here, \(D_p\) was kept constant during imaging or fixed position measurements, and could be determined experimentally through approach curves. A second position used in P-SICM is 12.5 \(\mu\)m above the sample, which in this case is half of the range of z-piezo movement. A longer vertical movement of the pipet probe can be achieved with the combination of piezo and Z stepper motor, but local potential variations are limited within approximately 10 \(\mu\)m above the cell surface. Therefore, these two positions are ideal for accurate evaluation of cell monolayer conductance and work well with the hardware capabilities of the P-SICM platform implemented here. Importantly, for measurements at these two positions, pipet feedback was turned off to avoid any convection due to pipet movement or possible crosstalk between electrodes.

As in the case of scanning conductance measurements described above, in the vicinity of the cell layer, the local transepithelial conductance can be determined based on Equation 1 and be further described with the following equation:

\[
G = \frac{E}{\rho \cdot V_e} = \left( \frac{\Delta V_{0.2 \mu m} - \Delta V_{12.5 \mu m}}{\rho \cdot V_e} \right)
\]

in which, \(E\), the electric field is obtained by measuring potential deflection \((\Delta V)\) induced by transepithelial potential \((V_T)\) at two probe-surface distances \((D_p)\); \(V\) is the range of potential swept (\(V = 100 \text{ mV, } -50 \text{ mV to } +50 \text{ mV}\)); \(\rho\) is the cell medium resistivity; and \(\Delta Z\) is the vertical displacement of the pipet. To avoid polarization in the cell plasma membrane, an alternating transepithelial potential \((V_T)\) was applied, the frequency of which was determined from impedance measurement to minimize the capacitive contributions. The magnitude of the transepithelial potential was less than 50 mV to avoid cell damage and disruption of the cell membrane.\(^{33,46,47}\)

SICM was originally designed to study biological samples, so pipet can be filled with a wide range of electrolytes, such as phosphate buffered saline (PBS) for cell physiological studies, or normal cell culture medium helping preserve the viability of cells.\(^{40,48}\) A constant distance between pipet and sample surface allows non-invasive topographic imaging of the cell apical surface. On these high-resolution topographic images, the positions of cell body and cell junction can be clearly identified. In the case where each image contains 512 pixels \(\times\) 512 pixels for a scan area of 40 \(\mu\)m \(\times\) 40 \(\mu\)m, the dimension of each pixel (lateral movement of pipet) is about 78 nm. At smaller scan sizes, pixel dimensions can be reduced to as small as tens of nm. When a cell junction is located from the topographic image, the pixel coordinate for that position can be obtained, which allows the pipet to be positioned at a desired location with high spatial precision. The measured conductance over each identified spot therefore correlates with the underlying membrane permeability. SICM provides a more efficient and accurate way to position the pipet, as compared with conventional diffraction-limited optical microscopy.

**P-SICM Studies of MDCKII Cells**

Aided by high resolution control of vertical and lateral pipet position, P-SICM allows differentiation of conductive pathways in the apical cell membrane. We have applied this novel technique to recordings of the wild type Madin-Darby canine kidney strain II (MDCKII-WT) cells. First, a topographic image was obtained under conventional SICM mode in absence of transepithelial potentials. The locations of cell bodies (CB, transcellular pathway) and cell junctions (CJ, paracellular pathway) were then pinpointed from the image to extract their spatial coordinates. The recording pipet was positioned over CB or CJ based on these coordinates to record the potential deflection \((\Delta V)\) induced by the applied transepithelial potential \((V_T\) ~ 50 mV to +50 mV) at two discrete probe-surface distances (0.2 \(\mu\)m and 12.5 \(\mu\)m) respectively. Multiple potential measurements were performed on three MDCKII-WT monolayers with a transepithelial electrical potential difference of 400 mV.
transcellular conductance (G_t) displays Gaussian distributions with averages of 2.53 ± 1.49 mS/cm² (n = 49) and 6.20 ± 2.54 mS/cm² (n = 62), respectively. The transcellular conductance (G_t) significantly differs from the paracellular conductance (G_p) (p < 0.001). These conductance values are very close to results from Gitter et al., who measured transcellular conductance of 2.5 ± 0.5 mS/cm² (n = 10) and paracellular conductance of 9.7 ± 1.2 mS/cm² (n = 10) from MDCK C11 cells (a resemblance of MDCKII cells) with a TER of 83 ± 6 Ω·cm² (n = 10). The G_t and G_p measured by P-SICM agree well with previously reported values without further mathematical deductions, a caveat required in other current/voltage scanning studies. These results demonstrate the ability of P-SICM to measure the heterogeneous potential distribution induced by trans- and paracellular pathways in epithelial cells, and prove the utility of P-SICM in differentiation of conductive pathways in cell membranes.

To demonstrate the ability of P-SICM in capturing paracellular conductance changes, a MDCKII cell line transfected with claudin-2 siRNA molecule (hereafter referred to as MDCKII-C2 cell) was carefully examined. In the MDCKII-C2 cells, the expression of claudin-2 (a known cation selective paracellular channel protein of tight junctions) was depressed by RNA interference. In absence of claudin-2, the TER of MDCKII-C2 increased significantly to 247.95 ± 30.35 Ω·cm², as compared with MDCKII-WT (113.68 ± 24.98 Ω·cm²). The G_p in MDCKII-C2 cells significantly decreased to 2.63 ± 1.26 mS/cm² (n = 78), compared with 6.20 ± 2.54 mS/cm² (n = 62) in WT cells. The 58% decrease in paracellular conductance was attributed to the loss of claudin-2 channels in the tight junction. Figure 3 plots apparent conductance measurements of cell junction (G_p) with P-SICM for MDCKII-WT and -C2. Notably a broader distribution and larger mean value was found in MDCKII-WT, which indicates a more heterogeneous paracellular pathway conferred by claudin-2 channels. These results not only confirmed claudin-2 as the channel protein underlying paracellular conductance, but also demonstrated the ability of P-SICM in capturing the paracellular conductance changes within the tight junction. The recorded G_p for MDCKII-C2 also decreased compared with MDCKII-WT. This small, but non-negligible decrease may be due to interference from surrounding paracellular pathways.

Finally, we aimed to use P-SICM to determine the ion selectivity of paracellular pathways. As reported by Yu et al., claudin-2 channel are aqueous, narrow and cation selective channels. To study the ion selectivity of claudin-2 channel with P-SICM, MDCKII-WT and -C2 cells were recorded in three types of solutions: (1) a control - Ringer’s solution in both chambers; (2) a cation replacement solution - Na⁺ in basolateral solution substituted by isomolar N-methyl-D-glucamine (NMDG⁺); and (3) an anion replacement solution - Cl⁻ in basolateral solution substituted by gluconate (Glu⁻). Both NMDG⁺ and Glu⁻ are known to be impermeable to cell membrane and tight junctions. From the recordings of P-SICM, the only significant change after ion replacement (p < 0.001) is a 33% decrease in the paracellular conductance of MDCKII-WT cells (with claudin-2 channels) under cation replacing conditions. In contrast, C2 cells (without claudin-2 channels) show no difference in conductance after ion replacement. The details of recorded values under ion replacing conditions were presented in Table S3. These results not only confirm the cation selectivity of claudin-2 channel but also extend the application of P-SICM to a broader spectrum of electrophysiological studies for tight junctions.

**Limitations and Future Directions**

Despite the ostensible advantages in biological recordings, there are several limitations associated with P-SICM. Although the use of a dual-barrel pipet allows pipet positioning and potential measurement to be controlled simultaneously, this design inevitably introduces a registry error in the spatial localization of recorded conductance. Although predicted to be small due to a ~50 nm separation between the two barrel tip centers, this factor must be taken into account in future recordings when spatial resolution is further improved to under 100 nm. As with most scanning probe techniques, the ultimate resolution attainable by P-SICM is largely a function of the probe employed, the feedback mechanism utilized and reductions in thermal drift. For P-SICM the geometry of the pipet tip and the probe-surface distance (D_p) represent two primary resolution-limiting factors. For experiments described above, signals recorded were localized to nominal radii on the order of 265 nm. Further, the multitude of competing conductance pathways and the use of a nonzero probe-surface distance (D_p) may result in systematic errors that arise from transcellular artifacts. Reduction of D_p and the use of smaller scanning probes are two practical approaches to improve spatial resolution. Smaller pipets are also more effective for potential measurements because a smaller probe-surface distance.

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**Figure 2.** High resolution topographic image of Madin-Darby canine kidney epithelial cell monolayer. Cell body and cell junction are clearly identified. Figure 3 shows the apparent conductance measurements of cell junction (G_p) with P-SICM for MDCKII-WT and -C2.
and variations in local potential are steeper at these closer working distances.

As for present study, P-SICM only makes point measurements that are then summarized to describe an average behavior of distinct pathways on cell membranes, which is time consuming and laborious. In the future, P-SICM can be modified to record potential images of the entire cell membrane under various transepithelial potentials. This improvement will allow differentiation of individual paracellular channels along the length of each tight junction structure. Tight junction dynamics is another focal point of P-SICM based research in the future. It pertains to two layers of tight junction regulation. (1) Tight junction stability. Signaling pathways such as MLCK (myosin light-chain kinase) regulate ZO-1 protein turnover within the tight junction and the transepithelial resistance in time-dependent manners. The recording frequency used in current study – 1 Hz, although not sufficient to capture fast opening and closing events within the tight junction, has already revealed heterogeneous conductance distribution at each recording location (Fig. 3). Increasing the recording frequency, coupled with drugs such as TNF (tumor necrosis factor) to manipulate tight junction stability, will reveal the time constant in paracellular channel kinetics, a key measure for understanding the dynamic behavior of tight junction. (2) Tight junction interaction. Both the cis and trans interactions between claudin proteins are important for tight junction permeability. Previous recording approaches lack the resolution to zoom into the discrete pairs of claudin interaction. Aided by fluorophore tagged protein tracking, P-SICM can reveal permeability differences induced by local claudin interactions within a few hundred nanometers under co-expression or co-culture conditions. Because P-SICM can position the recording pipet precisely over the tight junction, it enables direct stimulation or inhibition of discrete paracellular channels by highly localized application of ions, agonists or antagonists to the tight junction via the same pipet. This approach essentially avoids unwanted secondary effects caused by systemic application of these reagents. Finally, if a certain level of automation can be applied to current P-SICM configuration, large-scale screening for paracellular channel inhibitors will be made possible. Such drugs will not only provide new treatments to many barrier diseases but also facilitate structural analyses of the paracellular channel.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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Figure 3. Histogram of the conductance recorded over cell junctions (CJs) for MDCKII-WT (red) and MDCKII-C2 (blue). Note that the conductance for MDCKII-WT displays a broader distribution and a larger mean value than MDCKII-C2 (Adapted from reference 5, with permission).
