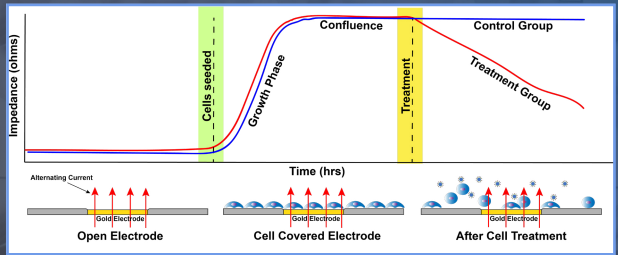


Barrier Function • Proliferation • Cytotoxicity • Migration Invasion • Signal Transduction • Differentiation

How ECIS Works



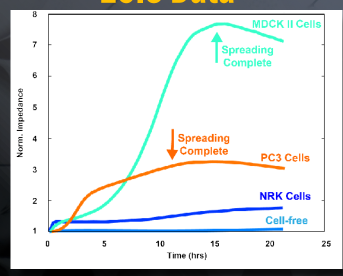
Schematic of an ECIS experiemnt starting with a cell-free electrode. Once cells are added, their attachment and proliferation cause a rise in impedance which plateaus once cells have reached confluency. At this point the cells can be perturbed and the resulting changes in impedance due to changes in cell behavior are montiored

Cell function modulates cell morphology. ECIS® is capable of detecting and quantifying cell behavior changes in the sub-nanometer to micrometer range. In ECIS® a small alternating current (I) is applied across the electrode pattern at the bottom of the ECIS® arrays (direct current cannot be used). This results in a potential (V) across the electrodes which is measured by the ECIS® instrument.

The impedance (Z) is determined by Ohm's law Z = V/I. When cells are added to the ECIS® Arrays and attach to the electrodes, they act as insulators increasing the impedance. As cells grow and cover the electrodes, the current is impeded in a manner related to the number of cells covering the electrode, the morphology of the cells and the nature of the cell attachment.

When cells are stimulated to change their function, the accompanying changes in cell morphology alter the impedance. The data generated is impedance versus time.

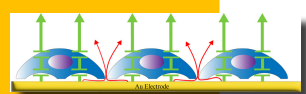
ECIS Data



The instrument can also use a range of AC frequencies from 100-100kHz and complex impedance measurements to determine different cell morphology parameters including barrier function, close contacts, and membrane capacitance.

Au Electrode

Current flow at low AC frequencies is via paracellular pathways



Current flow at high AC frequencies is via transcellular pathways

How Frequencies Reveal Cell Behavior

To understand why AC frequency is important in ECIS® we have to consider how frequency affects the current paths of cell-covered electrodes. (Note: the total current is maintained constant and voltage changes are measured.)

At relatively low frequencies (< 4,000Hz) most of the current flows in the solution channels under and between adjacent cells (red lines).

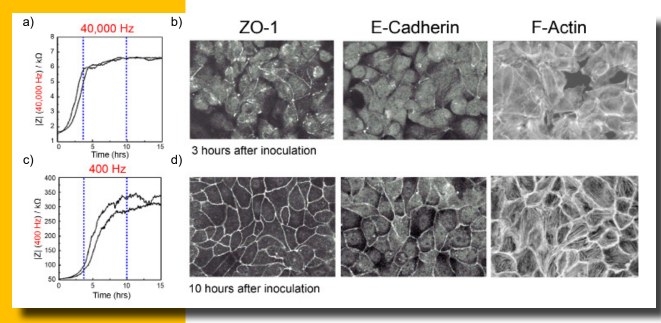
At higher frequencies (> 32,000 Hz) more current now capacitively couples directly through the insulating cell membranes (green lines).

The high frequency impedance is more affected by cell-coverage, whereas the low frequency responds more strongly to changes in the spaces under and between the cells.

With the ECIS® Z-Theta instrument, the impedance is broken down into its components (resistance and capacitance), and quantitative information about the cells can be obtained by modeling (Giaever and Keese PNAS 1991).

Using impedance data at multiple AC frequencies the ECIS® model calculates time course changes in:

- · The barrier function (permeability) of the cell layer
- · The degree of constricted flow of current under the cells
- · The cell membrane capacitance



Comparison of confluence and barrier formation of MDCK II cells. (a) ECIS® readings at High AC frequency (40,000 Hz) reveals plateauing impedance at ~3 hours suggesting confluence of the monolayer versus (b) low AC frequency showing plateauing at ~10 hours suggesting maturation of cell-cell juncitonal formation. Fluorescence images taken of the same ECIS® wells for zona occludens, E-cadherin, and F-actin (c) 3 hours after incoulation and (d) 10 hours after inoculation. Data courtesy of Professor Joachim Wegener, Univ. of Regensburg

APPLICATIONS

Publications available at www.biophysics.com

ATTACHMENT AND SPREADING

Traditional "counting attached assays" can only quantify the number of cells attached to any ECM coating. ECIS® assays give feedback on the strength of the attachment of the cells to the ECM.

CELL PROLIFERATION

As cells proliferate two factors act to change the impedance: cell number and cell morphology. In most instances the cells grow asynchronously and the impedance gradually increases until a maximum when cells become confluent. The impedance change is approximately linear with cell number while the cells are sub-confluent.

DIFFERENTIATION AND STEM CELL BIOLOGY

When cells differentiate they change their behavior allowing ECIS® to follow the events of cell differentiation. While most tools available to characterize stem cells preclude their further use, the label-free non-invasive nature of ECIS® allows for subsequent use of characterized stem cells.

BARRIER FUNCTION

Epithelial cells and endothelial cells regulate the passage of molecules across cell layers. Diseases, especially vascular disease, occur when this function is impaired.

SIGNAL TRANSDUCTION

ECIS® is especially useful to monitor the signal transduction pathways activated by G protein coupled receptors (GPCR). GPCR activation, regardless of the second messenger, results in alterations of the cell's cytoskeletal elements, causing morphological changes.

CELL INVASION

ECIS® distinguish between transmigration mechanisms that leave the monolayer intact from those that disrupt the cell layer. Published examples include metastatic cell and leukocyte trans-endothelial migration, as well as the migration of pathogens such as yeast, streptococcus, plasmodium, trypanosomes, and spirochetes.

CELL MIGRATION

ECIS instruments include an elevated field mode allowing for electroporation and wounding. The ECIS® wound is precisely defined, as it includes only those cells on the electrode. Additionally, with ECIS® the ECM protein coating is not scraped off and is unaffected by the current.

INFLAMMATION

ECIS® recovery-after-wounding allow for the discovery of molecules which aid in the process of tissue repair. ECIS® barrier function assays specifically measure the response of epithelial and endothelial cells to secreted cytokines and can give indirect information about the binding of immune cells to the epithelium or endothelium.

TEER

Continuous long-term measurement of TEER from under 10 to 10,000 ohmcm² in up to 16 wells using commercially available 6mm membrane inserts. Fast barrier function dynamics can be accurately monitored.

140

CELL TOXICITY

5.0 Wounded The ECIS® system has been used 4.5 Migration Rate specifically to assess the cytotoxicity of a 4.0 variety of toxicants. ECIS-based toxicity 3.5 70 BSC-1 Cells - DMSO tests are far superior to simple cell death Cell-free 3.0 assays, because cell function is also 60 2.5 monitored. 느 50 6.0 MDCK II Cells 2.0 Fibronectin Laminin 1.5 5.0 1.0 in D (ug/ml) 0.5 tance 3.0 2.0 80 100 120 (hrs) 1.0 10 15 20 Time (hrs) Cell-free



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Time (hrs)