Particle Manipulation Using Electric Field Gradients in Microdevices

Andrea Diane Rojas

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Rafael V. Davalos
Alex O. Aning
Abby R. Whittington
Eva M. Schmelz

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ABSTRACT

Electrokinetics is a family of effects that induces motion of a liquid or a particle within a liquid in response to an external electric field. Using the intrinsic electrical properties of bacteria and of breast cancer cells, electrokinetics can be used to manipulate these particles for two different types of applications: tissue engineering and breast cancer detection. The first application studied the effects of electric fields on bacteria cells as well as calcium ions to potentially create a meniscus scaffold with hydroxyapatite ends for anchoring. In response to the electric field, calcium ions were able to deposit locally and simultaneously with cellulose growth. Bacteria cells were also studied to determine their response under an AC field. At low frequencies, bacteria demonstrated controlled movement caused by electroosmosis and dielectrophoresis with a net motion caused by a dielectrophoretic force.

In the second application, the separation capabilities of different stages of breast cancer cells from the same cell line were tested using contactless dielectrophoretic (cDEP) devices. The electric field gradients in cDEP devices were altered to optimize selectivity and to determine an estimated membrane capacitance for each. From the results, the membrane capacitance of the early to intermediate stages proved to be very similar; however, late stage breast cancer cells have potential in being separated from early and intermediate stages.
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Chapter 1. Introduction

All particles will experience forces in response to applied electric fields as a consequence of their intrinsic electrical properties [1]. This was first discovered by Reuss in 1809 when charged clay particles suspended in a fluid migrated under an influence of an applied electric field which has been known as electrophoresis [2]. Another facet of electrophoresis, simply by adding a non-uniform electric field to a system, allowed uncharged particles to migrate in a fluid. The particle response to this application of non-uniform fields was identified by Pohl and defined as dielectrophoresis [3] and further presented in his 1978 book. Since then many applications have used this electrokinetic technology, including industrial applications that deposit a patterned coating of nanoparticles on a substrate using both DC and AC fields [4], environmental applications that sort bacteria from water [5], and biomedical applications that aggregate cells for co-culture systems [6].

This document describes two potential applications for using electrokinetics: tissue engineering and cell separation of breast cancer cells. For tissue engineering, bacterial cellulose is explored as a material for menisci purposes because of its hydroexpansive properties that resemble natural menisci. The cellulose is produced by bacteria that secrete fibers as part of their metabolic processes. Dielectrophoresis could be used as a method to control the motion of bacteria to create a meniscus scaffold. Furthermore, a meniscus scaffold needs to be anchored into the environment, and the addition of hydroxyapatite on the ends of the scaffold would promote integration. Therefore, electrokinetics will be used to attempt deposition of calcium ions onto a bacterial cellulose meniscus scaffold.

The second study for electrokinetics will be on understanding breast cancer cell response to electric field gradients. The cell line progression model from the same parent cell, MCF10A, has not been studied using electrokinetics and will provide insight on the capabilities for separating breast cancer cells by stage. Previous work has proven to separate aggressive cells from early and intermediate cells using dielectrophoresis [7]. However, the cell lines used were all from different people. To study cancer detection techniques, it is ideal to use cells from the same person. Therefore, the following work strives to separate stages of breast cancer cells by designing and testing dielectrophoretic microdevices with different electric field gradients.
Chapter 2. Background & Significance

2.1 Directed Motion of Bacteria and Calcium Ions to Create a Meniscus

2.1.1 What is Bacterial Cellulose (BC)?

In the Philippines, a gel-like dessert exists, Nata de Coco, which is produced in coconut water from bacteria cultures. The gel-like substance is referred to as bacterial cellulose and can be generated by the species *Acetobacter xylinum*. The production of cellulose begins with the movement of millions of bacteria cells in an acidic liquid at the air interface. The oxygen present at the air-liquid interface allows for continuous cellulose production. Cellulose nanofibrils form from cell metabolic secretions of glucose chains that entangle with other cellulose nanofibrils, resulting in a complex network of fibers that forms a blanket of hydrophilic material [8, 9]. The hydrophilic pellicle has been found to have great mechanical properties compared to other polymers [10] and proven to be biocompatible [11]. BC can easily be molded into many shapes if unagitated, therefore, has potential in medical applications including skin substitutes [12], artificial blood vessels [13], bone[12, 14], and other tissue engineering projects. Some include manipulating the bacterial cellulose porosity with wax particles to induce cell proliferation [15].

2.1.2 Alignment of Fibers Using Electric Fields

In the tissue engineering aspect, top-down fabrication of materials are limited on their feature size and therefore clinical translations of the products are limited due to their inability to reproduce on the nano or micro level. Complications with feature size arise when cells cannot infiltrate scaffolds leading to cell proliferation failure. Bacterial cellulose is a potential scaffolding material but is limited in its application due to their porous structure. The cellulose naturally forms a tight, random network of fibers creating small pores that are not large enough for cells to infiltrate. Furthermore, top-down fabrication techniques on bacterial cellulose such as cutting it into a desired shape compromises the mechanical integrity. Therefore, any bottom-up approach to fabricating larger pores and/or controlling fiber orientation would increase the applications for bacterial cellulose in the medical field.
Sano et al. have been able to approach manufacturing issues by creating a bottom-up fabrication method that controls the motion of the cellulose-producing bacteria. The *Acetobacter xylinum* are one or two microns in size and therefore can feel the effects of electrokinetics [16]. Their size assists with electrokinetics having a stronger effect than Brownian motion and negligible effects from gravity [17]. Furthermore, since *Acetobacter xylinum* thrive in acidic solutions, their net charge is negative [18]. The phenomena of electrophoresis, or motion of a charged particle relative to a fluid under a uniform field, can play a dominant role having precise control over bacteria cells and has been utilized to align bacterial cellulose fiber orientation. Using low DC fields in the range of 0.25 V/cm to 1 V/cm, bacterial cells continued to secrete cellulose nanofibrils and result in fiber alignment with their electrophoretic mobility estimated to be 4.68x10^{-9} \text{ m}^2\text{V}^{-1}\text{s}^{-1}. The ability to direct movement of bacteria potentially offers controlled fiber alignment and pore size while maintaining mechanical integrity.

### 2.1.3 Deposition of Ions onto Bacterial Cellulose Fibers

The bacterial cellulose fiber network presents a good template to deposit or mineralize nanoparticles for several biomedical applications. These include creating antimicrobial wound dressings which are impregnated with silver nanoparticles [19] and bone regeneration by creating a composite of cellulose and hydroxyapatite [12, 14]. Methods described for creating these composites involve coating after the bacterial cellulose pellicle is formed using immersion techniques. For bone mimics, the cellulose is first immersed in calcium solution then immersed in a phosphorus solution to create a hydroxyapatite structure. Concerns arise when the porosity of the cellulose limits the infiltration of the fluid into the pellicle resulting in solely surface deposition.

Previous experiments in the Bioelectromechanical systems laboratory has shown to simplify deposition processes. Using similar experimental protocols for aligning fibers, the deposition of materials onto cellulose fibers using applied DC fields was also discovered. Altering electrode materials and media constituents allowed deposition of free ions on to the cellulose fibers [20]. Deposition on individual cellulose fibers was seen using FESEM analysis and EDS verified deposition of the following electrode materials: copper, graphite, aluminum, and platinum.
Phosphorus, a component not needed in the liquid medium, was added to the system before the application of an electric field and FESEM verified the deposition of phosphorus. The electric fields used during these experiments ranged from 0.15 V/cm to 1 V/cm. With this technique, one can grow a BC pellicle while depositing desired materials onto individual fibers, thereby, simplifying the fabrication process.

Figure 2. Bacterial cellulose coated with copper [20]

2.1.4 Significance of Bacteria Studies

Applying an electric field to bacterial cellulose resulted in two discoveries: alignment of fibers and the deposition of ions. Both of these discoveries direct the field to fabricate a meniscus scaffold. Studies have shown that bacterial cellulose is comparable to the collagen fibrils of menisci; however, at high compression strains the ordered and arranged structure of the collagen fibrils clearly surpasses the unorganized network of cellulose fibers [21]. Using an electric field on the bacterial cellulose may orientate the fibers in the directions desired to mimic the collagen structure and mechanical properties. The meniscus has an ordered layer structure mainly containing circumferential fibers with some integrated radial fibers as seen in Figure 3.

Figure 3. A meniscus made of circumferential fibers with radial fibers integrated within the network
To secure the manufactured bacterial cellulose meniscus, it should attach to the intercondyloid fossa which is located between the condyles of the tibia. Therefore, the meniscus should be fabricated with hydroxyapatite at the horns to initiate osteogenesis on the tibia. Hydroxyapatite, a natural mineral component of bone, is composed of calcium and phosphate and is formulated Ca$_5$(PO$_4$)$_3$(OH). In this work, electric field gradients will be used to optimize design considerations that will simplify the fabrication process. Meanwhile, deposition of calcium ions on cellulose fibers will be tested and optimized. The ideal solution would be to create a meniscus scaffold in only a few steps by applying an electric field for both alignment of fibers and deposition of calcium ions, which would be followed by a phosphorus bath to create the hydroxyapatite structure.

2.2 Separation of Breast Cancer Cells for Personalized Medicine

The motivation to create a technology that can personalize medicine for breast cancer management and treatment is preceded by the negative results of blindly recommended therapies. Oftentimes, these blindly recommended therapies lead to the immune system becoming more resistive to further alternative treatments that if used originally could have killed the cancer and/or prevented reoccurrence. Furthermore, patients with the same diagnosis treated with the same therapeutic course may result with varying responses. The differences in their responses depend on unidentified biomarkers, tumor microenvironment, tumor stage, and other unknown factors which make up a cancerous cell’s genetic fingerprint [22]. This genetic fingerprint creates a hurdle for personalizing medicine.

Personalizing medicine to manage and treat cancer will be optimized from experimental therapeutic treatments in vitro using cancer cells from the patient. Studies have shown that cancer from a primary tumor sheds cells into the blood stream even at early stages [23, 24]. The cells known as circulating tumor cells (CTCs) circulate in the blood stream and may either be attacked and killed by the immune system or initiate a secondary tumor site to begin metastasis. More studies have shown that there is a direct correlation to the density of CTCs in the peripheral blood and the progression of cancer [24-26]. Thus, the level of CTCs in blood can be used as a predictor for patient prognosis [27]. Detection techniques that focus on finding CTCs using blood samples provides a minimally invasive method to constantly screen for reoccurrences as well as test for early stages of cancer. The following sections will discuss current methods and approaches for detecting CTCs in vitro.

2.2.1 Techniques for Detecting Circulating Tumor Cells

The main challenge for detecting circulating tumor cells (CTCs) is the extremely low concentration in blood (WBC 5-10×10$^6$ mL$^{-1}$, RBC 5-9×10$^6$ mL$^{-1}$, and platelets 2.5-4×10$^6$ mL$^{-1}$), as well as the identification of whether they are tumor initiating cells or not [28]. Several technologies have been implemented to target these challenges, including the state of the art magnetic-activated cell sorting (MACS) [29] and fluorescence-activated cell sorting (FACS) [30], which are high throughput screening methods that label target cells using antibody conjugated magnetic beads or fluorphore conjugated antibodies. The most reliable method for CTC detection in blood is the CellSearch System, an automated enrichment and immunocytochemical detection system first approved for use on monitoring patients with
metastatic breast cancer. The system uses magnetic beads coated with an antibody that targets the epithelial cell adhesion molecule (EpCAM). Targeting CTCs with antibodies and labels has its disadvantages of compromising cell structure and function as well as having limited sensitivity. Prior knowledge of biomarkers are needed to target cells and even with negative labeling, where undesired cells are targeted for removal, the sample may result in low yield and purity [31]. A recent study questioned the sensitivity of CellSearch because the normal genotype of invasive cancer is typically negative for the EpCAM expression, therefore the system will give false negatives [32-34].

A method for CTC detection that excludes labeling is magnetophoresis. Separations based on magnetophoresis use intrinsic magnetic properties of cells. When magnetophoretic separators are scaled down to the microscale, red blood cells and white blood cells are sorted from whole blood [35] and breast cancer cells are sorted from red blood cells [36]. The size was scaled-down because macro-scale separators generated low magnetic flux gradients on the biological cells. Another label-less detection method includes hydrodynamic sorting which is based on inertial microfluidics. Park et al used a multi-orifice structure to move a particle laterally through inertial lift force and vorticity and based the lateral movement on particle size [37]. However, the multi-orifice structure alone does not sufficiently separate CTCs. Hydrodynamic sorting uses cell size to separate particles from one another, but oftentimes cells are heterogeneous in their size even though they are of the same cell type [38]. Therefore, Moon et al combined it with dielectrophoresis to purify the output [39].

2.2.2 Dielectrophoresis

Dielectrophoresis (DEP), the motion of a particle due to its polarization in the presence of a non-uniform electric field, can be used to differentiate between cells based on their intrinsic properties [3, 40]. A particle’s response to a dielectrophoretic force may either be positive or negative depending on the permittivity of the cell compared to the suspending fluid. A non-uniform electric field polarizes the cell and the response time depends on the cell’s complex permittivity, which is frequency dependent. Therefore, at some frequencies the cell’s may respond to an applied force negatively or positively, i.e. repelled or attracted. DEP uses a cell’s positive or negative response to selectively trap, separate, and sort polymer microspheres, bacteria, and several types of eukaryotic cells in microchip applications [41-44]. The disadvantage of using microdevices with interdigitated electrodes was the susceptibility of electrode delamination and fouling during experimentation. Also, the electrode array patterned on the bottom of the channel results in a DEP force that is weaker towards the top of the channel where cells may not feel the force [45]. Turning up the electric field may increase the DEP force at the top of the channel, but may cause cell damage to those closer to the electrode. To reduce the disadvantages of traditional DEP microdevices and maintain the selectivity, insulator-based dielectrophoretic (iDEP) devices were invented by Cummings et al [46]. The iDEP devices simplified the fabrication process of DEP microdevices because the interdigitated electrodes were replaced by insulating geometries, i.e. posts. A DC or AC electric field is applied through electrodes that straddle the insulating structure array, consequently, creating non-uniform electric field gradients around the posts. iDEP proved to be a successful adaptation and a robust microdevice. Modifying microchannel geometry enhanced the performance of iDEP microdevices [47-50] and lead to single cell trapping of MCF7 breast cancer cells [51],
separation of live and dead HeLa cells [52], separation and concentration of two species of live bacteria [5], and separation of polystyrene beads in a multipart microdevice [53]. The disadvantages for iDEP devices are the relatively high current flow within the channel causing joule heating [51] and contamination from electrode/sample contact. When using dielectrophoresis for CTC detection and enrichment, it is ideal to limit contamination from electrode contact.

2.2.3 Contactless Dielectrophoresis

A fairly new rendition of dielectrophoresis is contactless dielectrophoresis (cDEP). The microdevice design eliminates sample/electrode contact and proves to be ideal for handling CTCs or any other precious cells. The metal electrodes are replaced by fluid electrode channels filled with phosphate buffer saline PBS that are separated from the main channel by a polydimethylsiloxane (PDMS) barrier. The insulating barriers capacitively couple the fluid electrodes to the sample channel creating frequency dependent electric field gradients that varies in response to geometrical and material properties of the device [54]. cDEP has been used to mix 0.5 µm fluorescent beads [55] and, more relevantly, has been used to separate live from dead THP-1 human leukemia cells [56]. The following sections describe two types of devices that operate at different frequency ranges with the original cDEP device working in high frequency ranges.

2.2.3a High Frequency

The first few cDEP microdevices presented a main sample channel with electrode side channels positioned directly above and below the area of cell capture. Numerical studies were experimentally validated when below 100 kHz, cells did not respond to the applied electric field gradients [56]. Furthermore, Shafiee et al found that frequencies above 700 kHz incurred leakages of the side channels to the main sample channels. The minimum electric field gradient squared on a particle necessary to induce movement was found to be on the order of 10^{12} [57]. Increasing the frequency and voltages improved cell capture, but cell lysis occurred when the voltage exceeded a threshold [56]. A more selective cDEP microdevice resembled traditional iDEP devices where in the main sample channel was an array of insulating posts. The electrode side channels were positioned 20 microns above and below the section of the main sample channel containing the array. This particular device was used in multiple experiments, first, to trap live human leukemia cells around insulating posts while dead cells flowed through [56] and, second, for the separation of breast cancer cells by stage [7].
The results from this device demonstrated that separation of cells from different stages of breast cancer using cDEP is possible, although, overlap in trapping efficiencies occurred at some frequencies. Keeping in mind that design factors affect the selectivity, it is possible for a new cDEP design working in the same frequency range may be more selective. Studies on separation of cancer cells by stage from different cell lines has also shown some success in other DEP studies, Mulhall et al determined that different stages of oral squamous cells have distinct electrical properties [59] and Gascoyne et al sorted different MDA breast tumor cell lines using a dielectric affinity column [44]. Furthermore, studies on the impedance signatures of the same cells used in Henslee et al were found to be unique [60]. These studies suggest that factors other than cell size, a dominating effect for cell DEP response, can be exploited using DEP.

2.2.3b Low Frequency

A recent development of cDEP technology has permitted lower frequencies to reach the threshold of $10^{12}$, the minimum electric field gradient squared needed for cell response. A new design idea increased lengths of the electrode barrier and distances between the source and sink, which is about 1 cm from each other instead of 500 microns as seen in high frequency devices. The increased barrier length increases capacitance which decreases the total impedance of the barriers at lower frequencies. Furthermore, the increased distance between the electrodes increases resistance in the sample channel and results in a higher proportion of voltage drop across the sample [61]. Now that cDEP devices are capable of working at low frequencies, the 1st crossover frequencies of mammalian cells may be analyzed. Sano et al sorted red blood cells experiencing negative dielectrophoresis from THP-1 human leukemia cells experiencing positive dielectrophoresis [54] and have used this cDEP device to determine the membrane capacitance of mammalian cells [61].
The purpose of this research is to optimize cDEP microdevices to separate different stages of breast cancer from the same parent cell line, MCF10A, so that they may be cultivated for personalizing medicine purposes. Each stage of cancer will then be tested against different therapeutic treatment options to determine the best one. It is possible that a mixture of different therapies may be best to fight the cancer; therefore, separating them by stage is a step forward towards personalizing medicine. Previous work using high frequency cDEP studied the separation efficiency for stages of breast cancer cells obtained from different people. The cDEP device isolated aggressive MDA-MB-231 cells from intermediate and early stage cells, MCF7 and MCF10A, respectively [7]. The selectivity of the device can be improved by altering the electric field gradients, but to mimic an ideal situation, the sample must come from the same person. The progression of the MCF10A cell line continues with MCF10AT1 cells followed by MCF10CA1 cells, which are early/intermediate and aggressive. Another cell line derived from the MCF10AT1 line is the ductal carcinoma in situ (DCIS) cells which are intermediate cells. The cell lines can be described more in detail from Miller et al and Santner et al [62, 63].
Chapter 3. Electrokinetic Theory

Electrokinetics is a family of effects that involves the motion of a liquid or particle within a liquid in response to an external electric field. Most dielectric substances possess a surface electric charge distribution that will attract ions of opposite sign (counter-ions) and repel ions of like charge (co-ions) when brought into contact with an aqueous (polar) medium. This attractive and repulsive process, when balanced by thermal diffusion, leads to the formation of the electric double layer (EDL). The EDL is composed of two layers of ions. The first layer is known as the Stern layer and consists of bound surface charge. The second layer consists of unbound charge, formed by a balance between Coulombic attraction and diffusion, and is known as the diffuse layer. The EDL forms the basis for which electrokinetic phenomena occur. If an electric field, for example, is applied tangentially along a charged surface, the field will exert a force on the charge and produce particle or liquid motion in the form of electrophoresis or electroosmosis, respectively. As such, the electrokinetic phenomena presented here depend on two factors: charge and an external electric field. The discussion below highlights the basic theory of these two phenomena.

3.1 Electrokinetics in a Uniform Electric Field

The application of a voltage difference across some distance in a liquid will generate an electric field, \( E \)

\[
E = -\nabla \phi \quad (1)
\]

where, \( \phi \) is the applied voltage. The response of a particle within the fluid is dependent on the distribution of the electric field. For example, a uniform direct current (DC) electric field can induce particle movement if the particle surface contains bound charge, but will induce no movement of a neutral particle. An influence by a uniform electric field on particle movement is called electrophoresis. The charged particle will move towards the electrode that is of opposite sign of the surface charge due to the Coulombic force attraction. Thus, a positively charged particle will navigate towards a negatively charged electrode. The force on the particle is directly proportional to the net charge of the cell, \( q \).

\[
F = qE \quad (2)
\]

Under a uniform DC field, the net forces on a particle are dependent on the motions induced by electrophoresis and electroosmosis. For a fluid in a DC field, the electric field interacts with free charges in the EDL, which forces this charged region into motion to drive fluid flow. Figure 5 shows the velocity profile of electroosmotic flow compared to laminar flow. Electroosmotic flow moves at the same rate along the width of the channel minimizing sample dispersion while pressure-driven flow causes the liquid in the center of the channel to move faster. Under a uniform AC field, however, no net fluid motion is observed as the net effect of fluid movement over one field cycle is zero [18].
Electroosmotic Flow  Pressure-Driven Flow

Figure 5. Electroosmotic flow is more uniform across the width of the channel while pressure driven flow is parabolic

Using Hückel’s equation [64], the sum of the electrophoretic mobility and electroosmotic mobility is used to determine the electrokinetic velocity, $V_{EK}$, of a spherical cell, where electrophoresis and electroosmosis are the only effects due to the applied electric field.

\[ V_{EK} = (\mu_{EP} + \mu_{EO})E \]  

\[ V_{EK} = \mu_{EK}E \]  

The electroosmotic mobility of a fluid under a uniform field may be calculated using

\[ \mu_{EO} = -\frac{\varepsilon \zeta}{\eta} \]  

Where $\zeta$ is the electrostatic potential between the ions in the EDL, $\eta$ is the viscosity of the fluid, and $\varepsilon$ is the permittivity of the fluid [64].

3.2 Electrokinetics in a Non-uniform Electric Field

Depending on the electric field distribution, more complex electrokinetic phenomena can occur. One such type is known as dielectrophoresis (DEP). Unlike electrophoresis, DEP uses non-uniform electric fields to induce particle motion. The key difference between DEP and electrophoresis is that while electrophoresis relies on bound surface charge to produce a net body force, the surface charge used to create DEP is induced by the electric field. As such, DEP can be used to drive movement of a neutral particle, whereas electrophoresis requires a charged particle. Hence, to adequately describe DEP one must consider how a particle polarizes under the influence of a electric field. The polarization of the particle is taken into account by calculating the net dipole force on a particle.
Figure 6. The net force on a dipole (a) and a polarized particle (b)

The figure shows two opposing charges separated by a distance, \( d \). The arrows projecting off the charges represent the electric field at those charges. At a random spatial distance, \( r \), is the location of the negative charge, \(-q\), and at \( r+d \) is the location of the positive charge, \(+q\). The following equation describes the net force on the dipole:

\[
F = qE(r + d) - qE(r) \tag{6}
\]

The electric field can be expanded about the position of \( r \) using the vector Taylor series expansion:

\[
E(r + d) = \sum_{k=0}^{\infty} \frac{1}{k!} (d \cdot \nabla)^k E \tag{7}
\]

\[
F = qE(r) + q \left( d_x \frac{\partial}{\partial x} + d_y \frac{\partial}{\partial y} + d_z \frac{\partial}{\partial z} \right) E + \ldots -qE(r) \tag{8}
\]

Since \( d \) is much smaller, only the zero and first term of the expansion is considered and results in the following equation:

\[
F = (qd \cdot \nabla)E = (p \cdot \nabla)E \quad \text{since } p = qd \tag{9}
\]

where \( p \) is the dipole moment. Using Coulomb’s law and the two-point dipole, the electric potential, \( \Phi_{\text{dipole}} \), can be calculated with a dipole moment \( p \) in a dielectric medium of permittivity \( \varepsilon_L \) [1, 65]:

\[
\Phi_{\text{dipole}} = \frac{p \cos \theta}{4\pi\varepsilon_L r^2} \tag{10}
\]

Using this equation, the effective field-induced dipole moment of a particle, \( p \), can be determined by comparing the potential of a dipole with the potential of an induced particle which is
calculated using a spherical single-shelled particle model [66]. This comparison can be made because the ion distribution for a single-shelled model under an electric field resembles that of a dipole. Other models including a homogeneous spherical particle, ellipsoidal single-shelled particle, and multi-shell particle have all been used to determine the induced dipole term of biological particles [18, 65, 67].

![Diagram of a single shell model of a cell](image)

Figure 7. Single shell model of a cell comprised of a cytoplasm core and membrane as an outer shell with radii $r_1$ and $r_2$

The single-shell model represents the cytoplasm core and the cell membrane of a mammalian cell. Using this model, each layer of the cell is modeled as an individual electrical shell as seen in Figure 7. The induced electrical potential, $\Phi$, of the cytoplasm, cell membrane, and suspending medium is calculated by solving the Laplace equation using two sets of boundary conditions:

\[
\begin{align*}
\Phi_L - \Phi_{\text{mem}} &= 0, \text{ at } r = r_2 \\
\Phi_{\text{mem}} - \Phi_{\text{cyt}} &= 0, \text{ at } r_2 = r_1
\end{align*}
\]

And

\[
\begin{align*}
\varepsilon_{\text{mem}}^* \frac{d\Phi_{\text{mem}}}{dn} - \varepsilon_L^* \frac{d\Phi_L}{dn} &= 0, \text{ at } r = r_2 \\
\varepsilon_{\text{cyt}}^* \frac{d\Phi_{\text{cyt}}}{dn} - \varepsilon_{\text{mem}}^* \frac{d\Phi_{\text{mem}}}{dn} &= 0, \text{ at } r_2 = r_1
\end{align*}
\]

where the first set of conditions state that the potential is continuous across the interfaces and the second set describes the total current at the interfaces [65]. Once the induced electrical potential for a single shell particle is determined it can be compared to Equation (10) to determine the induced dipole moment of the particle

\[
p = 4\pi\varepsilon_L r^3 \left( \frac{\varepsilon_{\text{eff}} - \varepsilon_L^*}{\varepsilon_{\text{eff}} + 2\varepsilon_L^*} \right) E
\]
where $\varepsilon_{\text{eff}}^*$ is the effective complex permittivity that captures the charging characteristics of both interfaces. Taking into consideration exponential time dependence $e^{i\omega t}$ for the electric field, the instantaneous applied field at time $t$ and location $r$ is given by

$$E(r, t) = \text{Re}[E(r)e^{i\omega t}]$$

where $i^2 = -1$ and $\text{Re}[...]$ is the real part of the complex phasor $E = E \cos \omega t + iE \sin \omega t$. This notation accounts for the magnitude and phase of the electric field that induces a dipole moment as the field is moving sinusoidally in time. The net dielectrophoretic force on the dipole can now be determined by combining Equations (9), (15), and (16) as well as taking the time-average [1]

$$F_{\text{DEP}} = 4\pi \varepsilon_L r^3 \text{Re}[K(\omega)]E \cdot \nabla E$$

(17)

Using the vector relation, $\nabla (A \cdot B) = (A \cdot \nabla)B + (B \cdot \nabla)A + B \times (\nabla \times A) + A \times (\nabla \times B)$, and knowing the electric field does not rotate ($\nabla \times E = 0$) gives us $(E \cdot \nabla)E = \frac{1}{2} \nabla(E \cdot E)$; therefore, the dielectrophoretic equation becomes

$$F_{\text{DEP}} = 2\pi \varepsilon_L r^3 \text{Re}[K(\omega)]\nabla(E \cdot E)$$

(18)

The cell in a fluid will also experience a drag force:

$$F_{\text{Drag}} = 6\pi \eta r v$$

(19)

where $\eta$ is the viscosity of the particle and $v$ is the velocity. For cell motion to occur, the magnitude of the $F_{\text{DEP}}$ must be greater than $F_{\text{Drag}}$.

### Table 1. Components of the DEP Force: Units and Magnitudes

<table>
<thead>
<tr>
<th>$F_{\text{DEP}}$</th>
<th>$\varepsilon_L$</th>
<th>radius</th>
<th>$\nabla(E \cdot E)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N = \left(\frac{A^2 \cdot s^4}{kg \cdot m^3}\right) \left(\frac{kg \cdot m}{A^2 \cdot s^6}\right)$</td>
<td>$\frac{F}{m}$</td>
<td>$m$</td>
<td>$\frac{V^2}{m^3}$</td>
</tr>
</tbody>
</table>

| COMSOL Units | $\frac{kg \cdot m}{s^2}$ | $\frac{A \cdot s}{V \cdot m} = \frac{A^2 \cdot s^4}{kg \cdot m^3}$ | $m$ | $\frac{kg^2 \cdot m}{A^2 \cdot s^6}$ |

| Magnitude | $10^{-15} - 10^{-20}$ | $10^{-12}$ | $10^{-6}$ | $10^{10} - 10^{15}$ |

### 3.3 Clausius-Mossotti (CM) Factor

The CM factor is a key component in determining the dielectrophoretic force on a particle and is defined by the single-shell model as follows
The CM factor measures the effective polarization strength of a particle in response to an angular frequency $\omega$. The effective polarization strength assumes the target to be a lossy, spherical particle. The breast cancer cells used are considered lossy due to their two interfaces, cytoplasm-membrane and membrane-suspending liquid, which create a phase lag when a cell polarizes and depolarizes. The loss of a particle is frequency dependent, therefore, their dielectric constants, $\varepsilon$, become complex quantities defined by $\varepsilon^* = \varepsilon + \frac{\sigma}{i\omega}$ where $\sigma$ is the conductivity. The single-shell model considers this with the effective complex permittivity term $\varepsilon_{eff}^*$:

$$\varepsilon_{eff} = \varepsilon_{eff}^* = \frac{\varepsilon_{mem}^*}{\frac{(r_2^*)^3}{(r_1^*)^3} + \frac{2(\varepsilon_{cyt}^* - \varepsilon_{mem}^*)}{\varepsilon_{cyt}^* - 2\varepsilon_{mem}^*}}$$

The CM factor varies from -0.5 to 1 across the frequency spectrum. When $[K(\omega)] < 0$, a cell will experience a negative DEP force and will be repulsed from areas of high fields. This occurs when the particle’s permittivity is less than the permittivity of the suspending media ($\varepsilon_p < \varepsilon_L$). When $[K(\omega)] > 0$, a cell will experience a positive DEP force and will attract to areas of high fields because the cell is more polarizable than the fluid ($\varepsilon_p > \varepsilon_L$). The point where $[K(\omega)] = 0$ is known as the crossover frequency, $f_{x01}$. At this crossover frequency, a particle will experience neither a negative nor positive force.

The crossover frequency provides a useful way to characterize cells using dielectrophoresis. When making the assumption that the cell membrane thickness is extremely thin ($r_2 - r_1 \ll r_2$), the membrane capacitance $C_m$ can be calculated with a measured crossover frequency [68]

$$C_m = \frac{\sqrt{2}}{2\pi rf_{x01}} \sigma_L$$

Figure 8 shows a plot of the CM factor for an early stage breast cancer cell, MCF10A, modeled using the single shell model. Because the cell contains two interfaces, there will be an inverse RC charge time of the cell membrane, which will then provide two crossover frequencies. According to the plot, higher frequencies on the order of $10^8$ will allow fields to penetrate into the cytoplasm but before then the low frequencies will fail to have polarization effects on the cytoplasm.
Figure 8. The Clausius-Mossotti curve of an early stage breast cancer cell modeled using the single shell model.
Chapter 4. Methods and Materials

4.1 Microdevice Fabrication

4.1.1 Polycarbonate Master Stamp Fabrication

Master stamps for cellulose fiber alignment were fabricated from polycarbonate boards that were milled using a computer numerical controlling (CNC) machine. Fabrication was outsourced for gratis by the mechanical engineering department at Virginia Tech. Rectangular channels were milled into a variety of different lengths and widths, but similar depths.

![Sample polycarbonate stamp to create channels](image)

4.1.2 Silicon Wafer Master Stamp Fabrication

In the Micron cleanroom, a <100> grade silicon wafer (University Wafer, MA) was cleansed of surface contaminates in BOE for 5 seconds then rinsed with deionized water and blown dry using pressurized nitrogen. To prepare the surface for photoresist, hexamethyldisilizane (HDMS) is used to promote adhesion and works best when HDMS is applied as a gas phase on a heated substrate. Using a spinner, 5 drops of HDMS solution was spun on the wafer at 4000 rpm for 90 seconds then it was transferred to a hot plate set at 115°C for 90 seconds. The wafer was then centered on the spinner again and a quarter-sized amount of AZ 9260 (AZ Electronics, NJ) positive photoresist was poured onto the wafer and spun at 4000 rpm for 40 seconds. The wafer was transferred to the hot plate to bake the photoresist at 115 °C for 30 seconds.

Upon curing of the photoresist, the MA-6 Mask Aligner (SUSS MicroTec, Germany) was used to imprint the device design onto the photoresist covered wafer. The 2D device was designed using AutoCAD Mechanical (AutoDesk, CA) and the high-quality photomask was printed by CAD/Art Services, Inc. The photomask was printed with the emulsion side up; therefore, features would be imprinted onto the device with the correct dimensions. The photomask is taped onto a 4” x 4” glass substrate fitted for the mask aligner and placed in the aligner so that the photomask comes in contact with the photoresist-covered wafer. The aligner was set to soft contact at a distance of 40 µm with a UV exposure for 60 seconds. Following exposure to the
UV lamp, the silicon wafer was placed in a 2:1 AZ 400K (AZ Electronics) developer to water solution for 30 seconds then rinsed with deionized water to remove the photoresist from the microdevice features.

Once all the photoresist was removed from the features, the Deep Reactive Ion Etching (DRIE) machine was used to etch the silicon wafer. The duration was calculated after measuring the rate at which it etched between the time points of 3 minutes and 5 minutes using the Dektak (Bruker AXS, WI). The Dektak measures the depth and is accurate for reading until a depth of 50 to 60 µm. Following DRIE etching, the wafer was placed in acetone for at least 5 minutes to remove the remaining photoresist. Due to a scaffolding effect from the etch method, further processing of the silicon wafer was needed to smooth the walls of the channels. Tetramethylammonium hydroxide (TMAH), an anisotropic wet etchant of silicon, was heated to 70°C and used for the removal of extra photoresist and the smoothing of the walls of the channels. The wafer was placed in the heated TMAH for 5 minutes and then rinsed with deionized water followed by pressurized nitrogen drying. A final coating on the silicon wafer was needed so that molds of PDMS could be easily peeled off the silicon wafer. The coating used is PTFE, a fluoropolymer with the chemical formula of C$_2$F$_4$. The DRIE applied the PTFE in gas form for 5 minutes.

4.1.3 Mold Formation of Devices

Molds of the devices were created using a SYLGARD 184 polydimethylsiloxane (PDMS) kit from Dow Corning. A 10:1 ratio of the elastomeric base and hardener was measured out by weight and mixed together by stirring vigorously. Depending on the size of the device, 25 grams to 40 grams of polymer was used. The polymer was placed in vacuum for 30 minutes to remove the bubbles formed from mixing. Meanwhile, the master stamp was wrapped in an aluminum foil boat and placed on a hot plate at 115 °C. After 30 minutes in vacuum, the majority, if not all, of the bubbles were removed, the polymer was slowly poured over the heated master stamp and left to cure for about 15-20 minutes. After the polymer cured, it was removed from the heat source and cooled for 5 minutes before peeling from the stamp. Holes for electrodes and for the inlets and outlets were punctured in the PDMS using 1.5 mm punchers (Ted Pella, CA).

4.1.4 Bonding of PDMS Device to Glass Slide or PDMS

The PDMS mold was cleansed with a mixture of soap and water, deionized water, and ethanol followed by pressurized air to dry off the PDMS. Further removal of any contamination was accomplished using scotch tape. The same cleansing process was used for glass slides. A plasma cleaner (Harrick Plasma, NY) was used to bond the PDMS to either glass or another PDMS layer. The samples were placed surface-side up under vacuum for two minutes to create a low-pressure gas chamber. After two minutes, the plasma was turned on for two minutes to ionize the gas atoms using high frequencies, creating a magenta-colored glow, which in turn bombarded the
surfaces of the PDMS and glass slides to break apart bonds of organic surface contaminants. To effectively bond PDMS to PDMS, they were exposed to plasma for ten minutes. Immediately after exposure to plasma, the PDMS was removed with the plasma-exposed surface facing up then the glass slide, or the other PDMS layer, was flipped over and placed firmly against the PDMS layer. Uniform pressure was applied by gently rolling a cylindrical object over the surface of the PDMS. The bonded device was placed in vacuum until experimentation.

Figure 10. Microfluidic cDEP devices bonded to glass

4.2 Cell Culture Protocol

4.2.1 Bacteria Cells

*Gluconacetobacter xylinus*, also known as *Acetobacter xylinum*, were purchased in a freeze-dried form from ATCC (Manassas, VA) and stored at -5°C until ready for revival. The bacteria pellet was revived by rehydrating with a few drops of the broth medium, 50 g glucose, 5 g yeast extract, 12.5 g CaCO₃, and 1 L deionized water, and then aseptically transferred into broth medium. Agar plates were made by cooling half-filled petri dishes containing fresh broth medium with the addition of 1.5% agar. Using microbiology techniques, a sterile loop was dipped into the solution of revived bacteria. The loop was swabbed back and forth in a zigzag manner from the outer rim to the center of the petri dish. Once that center was reached, the agar was punctured by the loop. The petri dish was rotated 90 degrees and the lifter started by crossing over the previous section and finally forming zigzags without crossing back. The lifter is then punctured again into the agar plate and the process is repeated one more time. Several agar plates were created and placed in the incubator at 27°C to allow cell colonies to form. Once cell colonies formed, 6 colonies were collected at a time and frozen with 20% glycerol at -80°C until ready for use.
To grow bacterial cellulose from the frozen state, a vial containing 6 colonies was retrieved from the freezer and brought to room temperature. Once colonies were at room temperature, they were transferred into a T-75 flask with 100 mL of media and placed in an incubator set at 27°C for about 3 days until a cellulose pellicle formed at the air/liquid interface. If no apparent formation of cellulose occurred then a new frozen vial was used and the process repeated. When a pellicle formed at the liquid/air interface, the flask was shaken up to release bacteria cells from the pellicle. Then 2 mL of media was removed and placed in a new container with 100 mL of fresh media.

4.2.2 Bacteria Cell Media

The media used for all experiments was a modified Hestrin-Schramm medium. The constituents of the media were a mix of the following in 1 liter of deionized water: 40 g fructose, 1 g potassium phosphate (KH$_2$PO$_4$), 0.25 g magnesium sulfate heptahydrate (MgSO$_4$•7H$_2$O), 3.3 g ammonium sulfate (NH$_4$)$_2$, 5 g yeast extract, 5 mL vitamin solution, 10 mL salt or trace metal solution, and 20 mL corn steep liquor. The solution was sterile using filtration and the pH was balanced to 5.5 with sodium hydroxide (NaOH).

4.2.3 Mammalian Cells

From a frozen state, a cell vial was removed from liquid nitrogen and sprayed with ethanol followed by loosening the lid. The vial was immediately submerged (with the cap above waterline) in a 37°C water bath until thawed (< 5 minutes). Under a cell hood, fresh 37°C media was quickly introduced into the vial with a pipette and mixed in with the old media containing DMSO. All contents of the vial were transferred into a 15 mL conical tube, which was then
topped with 10 mL of fresh media. The cells were centrifuged down at 120 rpm for 5 minutes and old media was aspirated using a Pasteur pipette. Using a pipette, 5 mL of fresh warm media was pipetted up and down to break up the cell clump that had formed while centrifuging and then all of the media was transferred into a T-25 Flask for incubation at 37°C.

After cells adhered to the bottom of the flask, they were ready for passaging. Trypsin, PBS, and media were placed in the 37°C water bath. After about 20 minutes, media was aspirated from the cell flask by holding the flask upside down making sure the Pasteur pipette did not come in contact with the adhered cells. The cells were washed with about 5-10 mL of PBS and then PBS was aspirated. This process was repeated once more. Then 1-1.5 mL (for T-25 flask) or 3 mL (for T-75 flask) of trypsin was added to the flask and placed in the incubator for 3-7 minutes, but no more than 10 minutes. If some cells were still adhered to the bottom of the flask, then the bottom or sides of the flask were lightly tapped until they become loose. About 5 mL of fresh media (for T-25 flask) or 8 mL of fresh media (for T-75 flask) was added to neutralize the media then transferred to a conical tube for centrifuging. The cells were spun down for 5 minutes and mixture of media and trypsin was aspirated with a Pasteur pipette with care taken to not aspirate the cell clump. Media was added to the clump of cells and pipetted up and down until the clump was not apparent. A small amount from the inoculated media was taken and plated into a new T-75 flask with 15mL of media (a T-25 flask will only have 5 mL of media). The amount taken was dependent on current number of cells and how fast or slow the cells grew. The flask was placed in the incubator until feeding, passaging, or experimentation.

Feeding occurred 2-3 days after the last passage or feeding time or whenever the media drastically changed color. To feed the cells, media was heated up in the water bath and, optionally, PBS was also heated. Once the fresh media was heated, the old media in the cell flask was aspirated using the Pasteur pipette (making sure the cells were adhered to the bottom prior to). If using the PBS, the cells were washed and then 15 mL of media was added.

For experiments, the cells were passaged as mentioned previously. However, after cells were re-suspended in fresh media, one mL of inoculated media was removed and analyzed using the Vi-Cell (Beckman Coulter, IN) to count the number of cells and average diameter. Once the cell count was determined, some cells would be used to plate a new flask and the remaining cells would be re-suspended in DEP buffer (8.5% sucrose wt/vol, 0.3% glucose wt/vol, 0.725% DMEM/F12 wt/vol) [57] to create the appropriate concentration needed for the experiment. Staining of cells was done either with Calcein AM or Calcein AM red/orange. About 4-6 µL of stain was added into the conical tube with inoculated media and placed in a warm water bath for 10 minutes. The cells were spun down for 5 minutes and re-suspended in fresh DEP buffer. The conductivity was measured using a SevenGo Pro conductivity meter (Mettler-Toledo,Inc., OH) and if the conductivity was not near 100 µS/cm then they would be spun down for 5 minutes and re-suspended until the desired conductivity was reached.
Note: Cells were not used for experiments after 10-15 passages. New cells were thawed and cultured.

For freezing cells, it was best to use cells that have had only a low passage number. After passaging, several cryovials were filled with 1 mL of the inoculated media (with no less than $1 \times 10^6$ cells/mL) and then finally no more than 10% sterile DMSO was slowly added to each of the cryovials. The DMSO helps prevent cell death. The cryovials were placed in a CoolCell® (BIOCISION LLC, CA) then directly in a -80°C freezer to freeze at a rate of -1 degree per minute.

### 4.2.4 Mammalian Cell Culture Media

The MCF10A and MCF10AT1 cells were cultured with 500 mL of DMEM/F12 (Invitrogen, NY), 25 mL of horse serum (Invitrogen), 5 mL of penicillin/streptomycin (Invitrogen), 500 µL of insulin (Sigma-Aldrich), 50 µL of cholera toxin(Sigma-Aldrich), 100 µL of epidermal growth factor (Sigma-Aldrich) and 250 µL of hydrocortisone (Sigma-Aldrich). MCF10CA1 and DCIS were cultured with DMEM/F12 and 5% horse serum. Additionally, 1% of syringe-filtered L-glutamine (Invitrogen) was added every two weeks in each culture medium.
Chapter 5. Bacterial Cellulose Studies

5.1 Experimental Procedures

5.1.1 Applied DC fields for Controlling Deposition of Ions onto Cellulose Fibers

5.1.1a Device Setup for Deposition

Inoculated media was inserted into PDMS channels using 1000 µL pipette tips which were used as reservoirs. Extra inoculated media was injected into each tip to equalize the flow in the channel. The channels were 3 cm long and 0.5 cm wide. Platinum electrodes were placed in the pipette tip reservoirs and connected to a DC power supply (Tenma Electronics, DE). The experiments were left to run for 72 hours at several voltages: 3.5 V, 4 V, 5.5 V, 6 V, and 7 V.

5.1.1b Media Alterations for Ion Deposition

The culture media for the bacteria cells was altered using 25% 1M CaCl₂ to 75% modified Hestrin-Schramm (HS) Medium. For each experiment, 20 mL of the media mixture was inoculated with 1 mL of bacteria.

5.1.1c Sample Preparation for ESEM

PDMS bonded to PDMS or glass was peeled apart to remove the pellicles formed during experiments. The pellicles were placed in test tubes of deionized water, and then placed in the sonicator for one hour at 60°C. Afterwards, the deionized water was removed and replaced with fresh deionized water. The sample was then placed in liquid nitrogen and quickly removed and placed in a freeze dryer (Labconco Corp., MO) for 48 hours. Once the samples were lyophilized, they were cut and mounted onto an SEM specimen holder via copper tape. The samples were sputter-coated with a mixture of platinum and palladium for 45 seconds to make the sample conductive.

5.1.1d Alizarin Red Staining for Presence of Calcium

After the application of a DC field, samples that were not prepared for ESEM analysis were submerged in deionized and shaken for 5 minutes in a conical tube. The deionized water was replaced with fresh deionized water. Two drops of Alizarin Red Staining (Electron Microscopy Sciences, PA), a tool to identify if calcium is present in a sample, was added to the submerged pellicle for 10 minutes. The pellicle was then removed from the staining process and placed in another conical tube with fresh deionized water and shaken for 5 minutes. Finally, the stained pellicle was placed on a petri dish and analyzed. If the sample contained calcium, the areas would turn a pinkish-red.
5.1.2 Applied AC Field Experiments to Create a Meniscus Scaffold

5.1.2a Preparation for Fluorescent Optical Microscopy

The devices used were PDMS molds bonded to glass with holes punctured on each end with 1.5 mm hole punchers (Ted Pella, Inc., CA). Bacteria cells were suspended in modified HS media and stained using Bactlight™ (Invitrogen, CA), so that they would be easily seen under the fluorescent microscope. A pipette was used to inject inoculated media into the meniscus-shaped mold until it was completely filled. Aluminum electrodes were placed directly into the punctured holes and pressure flow was allowed to settle for several minutes before applying an AC electric field.

5.1.2b Applied AC Fields on Fluorescent Bacteria

A function generator (GFG-3015, GW Instek, Taiwan) was used to apply an AC field to the electrodes and was monitored using an oscilloscope (TDS-1002B, Tektronix, USA). Frequencies used in these experiments were below 1 Hz and the magnitudes used were 10 V and below. An inverted fluorescent microscope (Leica DMI 6000B, Leica Micro-systems, IL) equipped with a camera (Leica DFC420, Leica Micro-systems, IL) was used to record video footage of bacteria cell response to the applied AC fields.

5.1.2c Numerical Modeling of a 2D Meniscus under AC field

Using the AC/DC module in COMSOL Multiphysics 4.1., a geometry designed in AutoCAD Mechanical was modeled to solve for the electric field distribution. This was accomplished by solving for the potential distribution, \( \Phi \), using the Laplace equation, \( \nabla^2 \Phi = 0 \). The boundary conditions used are prescribed uniform potentials at the inlet or outlet of the holes. The electrical properties used are presented in Table 2.

Table 2. Electrical Properties Used for Numerical Modeling of Meniscus Design

<table>
<thead>
<tr>
<th>Electrical Properties of Materials</th>
<th>Electrical Conductivity (S/m)</th>
<th>Relative Electrical Permittivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDMS</td>
<td>( 0.83 \times 10^{12} )</td>
<td>2.65</td>
</tr>
<tr>
<td>Water</td>
<td>( 5.5 \times 10^{-6} )</td>
<td>80</td>
</tr>
</tbody>
</table>

5.2 Results and Discussion

The use of electrophoresis on Acetobacter xylinum cells is an innovative approach for microweaving a potential tissue scaffold [16]. It is an alternative to top-down fabrication which increases in cost when decreasing in feature size. Initial studies with electrophoresis allowed potential for inexpensive bottom-up fabrication methods that would create feature sizes on the nanoscale using the cellulose-producing bacteria. The following studies of controlling bacterial cellulose with DC and AC fields lead to further understandings of cell and environment
response. The results presented here demonstrate the potential for creating a meniscus structure using electric fields.

5.2.1 Deposition of Ions Using Applied DC Electric Fields.

The main goal for these experiments was to determine whether deposition of calcium could occur simultaneously with fiber alignment to create a bottom-up approach for tissue engineering. This would not only eliminate a step in the processing, but also improve the technique for thoroughly depositing a material such as hydroxyapatite within the scaffold. As mentioned previously in Chapter 2, current work to create a bacterial cellulose scaffold with hydroxyapatite is limited to the cellulose surface.

The voltages applied during these experiments were to represent the electric fields within the ranges given from a previous study [16]. However in these experiments, an electric field of 1.83 V/cm (5.5 V) provided the best results by producing cellulose for ESEM analysis. Voltages above 5.5 V in the same chambers failed to produce a cellulose pellicle in the 6 attempts. A higher electric field than 1.83 V/cm is a factor that may cause bacteria to stop producing cellulose possibly because at these fields the velocity of the cell may be too high for secretion of glucose chains [69] or other metabolic processes. At lower voltages the experiments failed to show deposition.
Table 3. EDS Analysis of Bacterial Cellulose Control

<table>
<thead>
<tr>
<th>Materials</th>
<th>Atomic Percent</th>
<th>Mass Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>42.74</td>
<td>53.26</td>
</tr>
<tr>
<td>Palladium</td>
<td>2.349</td>
<td>0.34</td>
</tr>
<tr>
<td>Calcium</td>
<td>0.15</td>
<td>0.06</td>
</tr>
<tr>
<td>Oxygen</td>
<td>49.08</td>
<td>45.92</td>
</tr>
</tbody>
</table>

Figure 12 shows a control of a BC pellicle in 25% 1M CaCl₂ and 75% HS media. An EDS analysis was taken and demonstrated that the amount of calcium in atomic percent was 0.15. The palladium amount presented in the EDS results accounts for the palladium coating used for sample preparation. Figure 13 shows an ESEM image of a BC pellicle that underwent 5.5V in the same solution. The EDS analysis results show that there is an increase in calcium deposited compared to the control.
Figure 13. Bacterial cellulose under 5.5 V in CaCl\(_2\) solution with depositions

Table 4. EDS Analysis of Bacterial Cellulose under 5.5 V

<table>
<thead>
<tr>
<th>Materials</th>
<th>Atomic Percent</th>
<th>Mass Percent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>43.67</td>
<td>33.16</td>
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<tr>
<td>Palladium</td>
<td>0.83</td>
<td>5.61</td>
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<tr>
<td>Platinum</td>
<td>0.36</td>
<td>4.39</td>
</tr>
<tr>
<td>Calcium</td>
<td>0.70</td>
<td>1.77</td>
</tr>
<tr>
<td>Oxygen</td>
<td>54.44</td>
<td>55.07</td>
</tr>
</tbody>
</table>

It was noted that Figure 13 was taken near the electrode hole where deposition was seen. Moving further away from the electrode hole, Figure 14 shows that deposition failed to occur. Even though deposition failed to occur, cellulose was aligned. Thus, at 1.83 V/cm (5.5V) deposition occurred solely at the cathode hole.
Figure 14. Bacterial cellulose image taken far from electrode hole showing alignment but no deposition under the applied field

Figure 15 further confirms these results when Alizarin red was used to detect calcium in the pellicle with an applied field of 1.66 V/cm (5 V).

Figure 15. Bacterial cellulose pellicle showing the presence of calcium at one electrode hole
There was localized deposition seen on the BC pellicle. It is possible that once the cellulose was growing and calcium ions in the local region of the cathode were deposited, the calcium ions in other regions had difficulty migrating towards the cathode. Therefore, calcium was only deposited locally instead of throughout the entire pellicle. If the experiment was set to run for longer than 72 hours, then the amount deposited may have increased.

At lower electric fields, many of the experiments did not produce cellulose and when they did, no deposition of calcium was seen. The experiments that failed to produce cellulose were a result of the location: the fume hood. Since the experiments did not use much media, the exhaust of the fume hood dried them up. With deposition failing to occur on the samples that did produce cellulose, further studies were not completed at these electric fields.

Although the deposition was not seen to be uniform throughout the pellicle, deposition did occur simultaneously with alignment of fibers. With that in mind, this technology has potential to be used in creating a meniscus with aligned fibers and with hydroxyapatite ends to anchor in position.

5.2.2 Bacteria Manipulation for Creating a Meniscus Using AC Fields

Since the bacterium, *Acetobacter xylinum*, creates a fiber network that is biocompatible, hydroexpansive, and mechanically suitable, then the ideal tissue to mimic is the meniscus. Controlling the orientation of the fiber network has potential with electric fields; however, the movement has not been thoroughly studied using AC fields. The following study used a meniscus-shaped mold to carry out experiments that would mimic future experiments. The design was used to guide the bacteria in the circumferential direction that mimics the orientation of native menisci collagen fibers. In this mold, channels were designed to be 50 microns wide and spaced apart by 150 microns of PDMS. The bacteria movement was monitored while adjusting frequencies and voltage.

![Figure 16. Schematic of meniscus pattern used in the experiments. Outer diameter is 46 mm with a width of 10mm. Each channel was 50 microns wide and 50 microns deep](image-url)
The results of the experiments proved that bacteria cells did respond to an AC field at low frequencies such as 1 Hz. Higher frequencies were tested but no movement was observed. The response of the bacteria at low frequencies showed that they moved backwards and forwards due to electroosmotic flow. The movement did have a net motion, which is a result of a dielectrophoretic force. It is safe to assume that the time-average velocity for the electroosmotic flow is equal to zero.

A study on the electric field gradients created within the device was accomplished using numerical modeling. The model uses a slightly different meniscus shape, but experimental conditions were applied and the overall result is similar. Looking at the surface plot of the electric field gradients, one can see that they are highest at the locations closest to the electrodes. Also, the gradients are highest where the PDMS barriers end, which is in the shape of the upside down ‘v’. The field minimum is presented in the center of the device. Therefore, bacteria cells on the left side of the meniscus will have a net motion towards the left, whereas, the bacteria cells on the right side will have a net motion to the right.

![Surface plot showing the electric field gradients to be highest towards the ends. The plot was normalized using 1 Hz and 20 V.](image)

**5.3 Future Work**

As this was only a study on the response of bacteria cells to the fields, further work must be completed to understand whether the mechanism of using these electric fields will be beneficial in this field. First, the mechanism for depositing calcium ions needs to be studied further by increasing deposition time from 48-72 hours to 7 days or more. From preliminary experiments, it was noted that electrode material would degrade within several hours or days and therefore a field would be terminated. Experiments on electrode coatings such as using Nafion could prevent electrode ions from being removed and thus allow long term studies to occur. After deposition of calcium ions, the cellulose will need to be immersed in a phosphate solution, such as simulated body fluid [14], to initiate hydroxyapatite formation. Characterization will determine if the hydroxyapatite formation embodies the bone structure and if it can promote osteogenesis.
Secondly, it was proven that bacteria cells respond to a DEP force under an AC electric field at low frequencies. Since the experiments were observed under an optical microscope, cellulose fibers were undetectable with the naked eye. Studies should be done over several days to see if cellulose can grow under these conditions and whether fibers are growing within the center of the meniscus. It may be possible that the DEP force will drive all the bacteria cells towards the horns of the meniscus, which can create a dense network in that area compared to the center of the meniscus. Altering the device geometry and electrode setup may solve the potential issue of non-uniform density in the structure. However, device ideas that allow an open surface would be ideal in creating the meniscus, since the cells need oxygen to produce the cellulose and the cellulose needs to maintain its mechanical integrity. Even if the device contained a closed surface, the geometry will only be replicable within the first few layers of the pellicle. Lastly, mechanical testing on the bacterial cellulose will need to be performed to determine the effects of electric fields on cellulose growth. The strength of aligned cellulose has not been tested thoroughly enough to confirm any improvements to mechanical properties. Positive results of this would be a major milestone in engineering meniscus scaffolds.
Chapter 6. Separation of Breast Cancer by Stage

6.1 Experimental Procedures

6.1.1 High Frequency Experimental Setup

Microdevices were placed in vacuum for 30 minutes prior to experimentation. Once the cells were almost ready, the microdevice was removed from vacuum and prepped for experiments. Pipette tips were placed in the inlets and outlets of the electrode side channels and phosphate buffered saline (PBS) was injected into these side channels using a pipette. The microdevice was viewed under the microscope to verify no leakage occurred. Thin-walled PTFE tubing (Cole-Parmer Instrument Co., IL) was used and the ends were cut at an angle into one 3-inch piece and one 6-inch piece. The 3-inch piece of tubing was inserted into the outlet of the device. Once the cells were ready and suspended at a concentration of $10^6$ cells/mL, the 6-inch tubing was attached to a 1 mL plastic syringe and used to draw up half a milliliter of cell suspension. The open end of the tubing was inserted into the microdevice and the syringe was placed in a micro-syringe pump (Harvard Apparatus, MA). The syringe pump applied flow rates as low as 0.02 mL/hr.

An AC electric field was created by a combination of waveform generation and amplification equipment. The output signal of a function generator was amplified with a wideband power amplifier (Amp-Line Corp., NY). The output voltage was enough to power a high-voltage transformer (Amp-Line Corp., NY) which was connected to the electrode side channels and an oscilloscope to monitor the frequency and amplitude. Note: The transformer was placed on a grounded surface and attached to devices using high-voltage wiring.

An inverted fluorescent microscope equipped with a camera was used to record 50 second videos of the breast cancer cells flowing through the channel. The first two microdevices experimented with were analyzed by quick observations. The final high frequency cDEP device was analyzed using a MATLAB code. The code analyzed the intensity of the fluorescent cells flowing into the device and out of the device. This measured the percent trapping of the cells.

6.1.2 Low Frequency Experimental Setup

The same process was used to prepare the microdevice for experimentation, but instead of one outlet the microdevice contained two outlet channels. Each cell line was cultured in three different flasks for each set of sample testing (n=3) to maintain viability throughout the experiments. The concentration for each sample size was set around $3\times10^6$ cells/mL and the
conductivity measured before and after each set of experiments. The electronics used to operate at lower frequencies were developed by another graduate student, Michael Sano. These electronics were designed to operate from 0 to 70 kHz and no more than 300 V\textsubscript{rms}.

For the low frequency experiments, a flow rate of 0.005 mL/hr was used for sorting the cells. For all experiments, an applied voltage of 200 V was maintained while the frequency was adjusted. For each frequency tested, there was a five minute waiting period before video recording began. This was the amount of time the cells took to flow through the entire channel and to experience all the effects of the electric field gradients. Once the five minutes were up, video recording began and lasted for two minutes. The next frequency applied was chosen at random by drawing a number out of hat. The videos were analyzed using a similar MATLAB code as the high frequency, but instead the code was used to measure the fluorescence intensity of each cell as it crossed a specific point on one line.

![Figure 18. MATLAB analysis measuring the fluorescence intensity of breast cancer cells as they cross the red line. The spread of the red line is from 0 to 300 and the peaks shown on the graph measure the fluorescence intensity where cells cross the line. For this example, many cells have crossed the red line in the 0 to 25 range which corresponds to the top of the channel.](image)

### 6.1.3 Numerical Modeling of Devices

Microfluidic devices were modeled using the AC/DC module in COMSOL Multiphysics 4.1. This was accomplished by solving for the potential distribution, $\Phi$, using the Laplace equation, $\nabla^2 \Phi = 0$. The boundary conditions were prescribed uniform potentials at the inlet or outlet of the electrode side channels. Table 5 lists the electrical properties used for modeling.
Table 5. Electrical Properties Used for Numerical Model of cDEP Devices [57]

<table>
<thead>
<tr>
<th>Electrical Properties of Materials</th>
<th>Electrical Conductivity (S/m)</th>
<th>Relative Electrical Permittivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEP Buffer</td>
<td>0.01</td>
<td>80</td>
</tr>
<tr>
<td>PBS</td>
<td>1.4</td>
<td>80</td>
</tr>
<tr>
<td>PDMS</td>
<td>$0.83 \times 10^{12}$</td>
<td>2.65</td>
</tr>
</tbody>
</table>

### 6.2 Results and Discussion

Contactless dielectrophoretic devices were used for the following experiments to remove the electrode/sample contact. Especially with handling breast cancer cells, the need to reduce contamination is important when working with concentrations on the order of 10 circulating tumor cells per mL of whole blood [70]. Any sort of contamination will affect culturing of these cells. The following results walk through the design processes taken to change the selectivity of the cDEP microdevices for breast cancer cells of different stage. It begins with alterations in the high frequency devices, frequencies ranging from 100 kHz to 300 kHz, and ends with the newly developed low frequency cDEP microdevice which can be used at frequencies of 70 kHz and below.

#### 6.2.1 High Frequency cDEP Experiments

There were three sets of devices that were used to study the breast cancer cell response to electric field gradients. The designs were chosen to use the frequencies and voltages similar to previous work [7]. In designing the models, the gradient of the electric field squared on the order of $10^{12}$ and above needed to be present to affect cells flowing through the channel. The first set of experiments were initially conducted in a device used to successfully separate RBC and human leukemia cells in the Bioelectromechanical systems laboratory at Virginia Tech [71].
However, due to the inherited nature of breast cancer cells being a form of epithelial cells that attach to surfaces and to each other easily, the cells fouled the device quickly. The fouling disrupted fluid flow and the observed trapping of cells. To relieve clogging of the device, the design was altered by increasing the space between each row of insulating posts.
For preliminary results of device 1, the early and early/intermediate cells, MCF10A and MCF10AT1, were tested and values were recorded for onset and 90% of trapping under a flow rate of 0.02 mL/hr. The onset of trapping was observed when cells began to show an obvious attraction to the insulating posts by either slowing their velocity in the vicinity or by ‘sticking’ briefly to the post for a couple of seconds. The 90% trapping was observed when most of the cells were attracted to the posts and formed pearl chains around them. Figure 19 shows the pearl chain formation of THP-1 human leukemia cells around the insulating posts. The results showed for both cell stages that the onset and 90% trapping were alike making the device not selective enough. The frequencies and voltages recorded for the onset of the MCF10A cell was 320 kHz and 95 V, while for the MCF10AT1 cells they were 320 kHz and 100 V. The frequency and voltage noted for 90% trapping was exactly the same for both cell types. It is safe to state that this device for breast cancer cells is not selective enough even though it may be for a mixture of THP-1 cells and red blood cells. One of the main factors for selectiveness is the difference in cell radius as pointed out in Equation 18 for dielectrophoretic force.

It was noted that the parameters used for these experiments were not exactly the same parameters used in the work of Henslee et al with breast cancer cells [7]. It is possible that the barrier thickness and depth has changed slightly due to processing. A quick study on the effect of barrier thickness on the electric field gradient squared was modeled using COMSOL Multiphysics 4.1. Figure 21 demonstrates the effect of a 10 micron barrier to a 20 micron barrier. The frequency needed to reach the electric field gradient threshold decreases with decreasing barrier thickness, so any slight differences in the thickness of the barrier may affect the frequency by 100 kHz or more. Thus, it explains the differences in the frequencies used from previous work to current work.

![Graph](image)

**Figure 21.** The maximum electric field gradient squared created in identical cDEP devices with different PDMS barrier thicknesses.
A new device, **device 2**, was designed using two different insulating post sizes: 100 µm and 75 µm in diameter. The design was tested to determine whether the selectivity was an issue due to clogging of the device around the corners of the electric field gradients. This design contains large posts surrounded by small posts, which creates a higher gradient within the center of the device. This would create a stronger force for attracting cells in the center of the device rather than on the edges.

![Diagram of device 2 showing the schematic and electric field gradients normalized with 100 V and 200 kHz. The 100 µm diameter posts are bordered by 75 µm diameter posts.](image)

**Figure 22.** Device 2 showing the schematic and electric field gradients normalized with 100 V and 200 kHz. The 100 µm diameter posts are bordered by 75 µm diameter posts

The early/intermediate cell line and the aggressive cell line, MCF10AT1 and MCF10CA1, were studied in this particular microdevice design. The fluid flow rate was set to 0.05 mL/hr. During the experiments, the area of analysis was focused solely within the center of the device. However, the response of the cells to this device did not change and the onset of trapping for each cell line was exactly the same.

The final high frequency cDEP design, **device 3**, was designed with a smaller channel width. The width of the sample channel and the number of post rows were reduced to improve fluid flow and eliminate fouling. The early and late stage breast cancer cells, MCF10A and MCF10CA1, were tested in the device at a flow rate of 0.02 mL/hr. The data was analyzed using a MATLAB code described in the beginning of this chapter. Figure 23 shows at frequencies below 300 kHz, MCF10A cells have an increasing % trapping trend at 160 V and 200 V. The aggressive cell line,
MCF10CA1, at 160 V seems to maintain a lower percent trapping range (10-30%) at higher frequencies than the MCF10A cells. The percent trapping increases to 50-70% when increasing voltage at those same frequencies. Overall at 160 V and 200 V, the MCF10A cells begin trapping at much lower frequencies than the MCF10CA1 cells. Lastly, 90% of MCF10A cells can be trapped at a lower frequency compared to MCF10CA1 cells.

![Graphs showing trapping percentage at different voltages and frequencies.](image)

**Figure 23.** Percentage trapping is compared between the early and late stages of breast cancer. At 160 V, the early stage has a steady incline in response to increased frequencies (a). At 200 V, the voltage is strong enough to induce trapping of late stages at higher frequencies (b). MCF10A cells need a lower frequency for 90% trapping to occur compared to MCF10CA1 (c).

Figure 24 demonstrates that at the same applied frequency and slightly different voltage, the MCF10CA1 cells do not pearl chain as well as the MCF10A cell line. When pearl chaining occurs, the dipole-dipole interaction between cells offers another binding force to a stationary object like the interaction between the cell and insulating post caused by the electric field gradients. The longer the chains, the more pronounced their interactions [1]. Therefore, fluid
flow cannot as easily break the interactions caused by the electric field gradients and carry the cell passed this region. However, increasing the frequency or voltage for the MCF10CA1 cells will eventually create the pearl chain effect. Thus, the cells inherently have different electrical properties because for the same device it takes a different frequency and voltage to reach the onset and 90% trapping efficiency or even to create the pearl chain effect.

![MCF10A 340 kHz 240 V](image1.png) ![MCF10CA1 340 kHz 200 V](image2.png)

Figure 24. Device 3 demonstrates that the early stage breast cancer cell line, MCF10A, traps very well at a frequency of 340 kHz compared to the late stage breast cancer cell line, MCF10CA1

The issue continues to be selectivity of the device, and with higher frequencies this selectivity seems limited if the particles have similar electrical properties. Using devices with frequencies near the cell’s crossover frequency will help ascertain whether cDEP alone can be used for separating out the different stages of breast cancer.

6.2.2 Low Frequency cDEP Experiments

A new cDEP device was recently designed to work at lower frequencies. The source electrodes were positioned in the beginning of the device and spaced 1 cm away from the sink. The barriers stretched alongside the main sample channel, located near the source electrodes, extend half a centimeter long while the barriers near the sink electrodes stretch one and a half cm long. As mentioned before in Chapter 2, the increased barrier length creates an increase in capacitance which in turn decreases the total impedance of the barriers at low frequencies, thus, allowing this device to operate at low frequencies. Figure 25a shows the electrodes, shaded in grey, are labeled source and sink. The main sample channel does not necessarily have insulating posts to create electric field gradients as was done for high frequency devices. Instead the microdevice contains ‘sawtooth’ structures that distort the electric field. Figure 25c shows the electric field gradients with a maximum around these ‘sawtooth’ structures.
The device design altered the electric field gradients to increase at lower frequencies, especially frequencies in which mammalian cells experience a crossover frequency, or a frequency in which the cell response switches from positive DEP to negative DEP. Figure 26 compares the electric field gradients for the low frequency device to a high frequency device. As mentioned before, the electric field gradients must reach a minimum of $10^{12}$ for the cells to respond to the DEP force. The graph proves that the low frequency cDEP device is capable of achieving this minimum starting at frequencies of 3 Hz. In previous work, human leukemia cells, THP-1 cells, were separated from red blood cells because the THP-1 cells experienced a positive DEP force between 20 and 70 kHz [54]. The DEP force directed cells into streamlines closer to the maxima of the electric field gradients and, therefore, caused them to flow across the top of the device. At the same frequencies, the red blood cells experienced a negative DEP force and were repelled from the maxima of the electric field gradients and therefore flowed towards the bottom of the channel.
Figure 26. The maximum electric field gradient squared across a frequency spectrum was determined for a high frequency and low frequency device using COMSOL Multiphysics 4.1. The red line pinpoints the lowest value that a cell needs to experience a DEP force. The low frequency device operates above this threshold across a large span of frequencies.

The breast cancer cells experienced similar behavior to the THP-1 cells. At low frequencies including 5 kHz, all stages of breast cancer were repelled away from the ‘sawtooth’ structure. Figure 27 demonstrates cell response under no electric field, 5 kHz, and 50 kHz.

Since the breast cancer cells were derived from the same cell line, MCF10A, it was expected that their electrical properties were to be very similar. However, it was hypothesized that they could be distinguishable at the first crossover frequency, $f_{x01}$, using cDEP. At 30 kHz and above, all cell lines were responding to some level of a positive DEP force by increasing cell-to-cell
interactions that became apparent with pearl chains forming as frequencies increased. The cell-to-cell interactions can play a major role especially when determining the crossover frequency in this device. At low flow rates and frequencies within the crossover range, the interactions may attract more cells from one streamline into another, therefore, influencing where the cells cross the line of analysis.

For these results two crossover frequencies were estimated using interpolation, \( f_{x01} \) was calculated based on the width of the channel and \( c - f_{x01} \) was calculated based on the control response. The cell’s response at the crossover frequency is neither negative nor positive, and the observed response imitated the control. Each of these frequencies were then normalized to the suspension conductivity which are comparable to results from Gascoyne et al where he determined this ratio for several cell types including other breast cancer cell lines [44].

<table>
<thead>
<tr>
<th>Stages of Breast Cancer</th>
<th>( f_{x01} ) (Mhz-m/S)</th>
<th>( c - f_{x01} ) (Mhz-m/S)</th>
<th>( \frac{f_{x01}}{\sigma} )</th>
<th>( \frac{c - f_{x01}}{\sigma} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCF10A</td>
<td>18.4</td>
<td>20.4</td>
<td>1.79</td>
<td>1.98</td>
</tr>
<tr>
<td>MCF10AT1</td>
<td>26.4</td>
<td>20.5</td>
<td>2.54</td>
<td>1.96</td>
</tr>
<tr>
<td>DCIS</td>
<td>21.4</td>
<td>21.8</td>
<td>1.99</td>
<td>2.03</td>
</tr>
<tr>
<td>MCF10CA1</td>
<td>18.84</td>
<td>13.75</td>
<td>1.75</td>
<td>1.28</td>
</tr>
</tbody>
</table>

For the early to intermediate stages, the estimated crossover frequencies were very similar. With the added control measure, the crossover frequencies adjusted slightly almost to the point of overlap. The control measured the movement of the cells when the electric field was turned off, this was to determine whether fluctuations in fluid flow or cell concentration was skewing the results. Therefore, these two crossover frequencies suggest that the measured values can vary significantly with fluctuations in flow rate and cell concentration, limiting the selectivity. For this specific device, separation cannot occur for the different stages of cancer. However, there is potential for the MCF10CA1 cells to separate from the others since the crossover frequency is much lower. This particular device was tested to separate the cells but small fluctuations had an unfavorable effect, thereby, limiting its overall selectivity.

Using the results of the crossover frequency normalized by the control, the cell membrane capacitance may be estimated for MCF10CA1 cells using Equation 22 and can be compared to MCF10A estimated and reported membrane capacitance values.
Table 7. Calculated Membrane Capacitance of Breast Cancer Cells

<table>
<thead>
<tr>
<th></th>
<th>$c - f_{x01}$</th>
<th>$\sigma$ (µS/cm)</th>
<th>Radius (µm)</th>
<th>Membrane Capacitance (F/m²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCF10A [7]</td>
<td>12.54</td>
<td>100</td>
<td>9.25</td>
<td>0.0194</td>
</tr>
<tr>
<td>MCF10A</td>
<td>20.4</td>
<td>103</td>
<td>7.96 ±0.99</td>
<td>0.0143 ± 0.0036</td>
</tr>
<tr>
<td>MCF10AT1</td>
<td>20.5</td>
<td>104.5</td>
<td>6.81 ±0.90</td>
<td>0.0168 ± 0.0045</td>
</tr>
<tr>
<td>DCIS</td>
<td>21.8</td>
<td>107</td>
<td>7.79 ±0.62</td>
<td>0.0141 ± 0.0023</td>
</tr>
<tr>
<td>MCF10CA1</td>
<td>13.75</td>
<td>107</td>
<td>6.88 ±0.58</td>
<td>0.0255 ± 0.0042</td>
</tr>
</tbody>
</table>

The membrane capacitance was calculated using each cell’s radius with their standard deviations. The CM factor is plotted with these standard deviations to determine if the membrane of each cell plays a significant role in the electrical properties of the breast cancer cells. To plot the single shell model of each cell, the cytoplasm conductivity was assumed to be 1 S/m and the cytoplasm permittivity was assumed to be $50\varepsilon_0$. With these assumptions, the second crossover frequency will not change and therefore is not shown.

According to the figure, the membrane capacitance for MCF10A, MCF10AT1, and DCIS is not significant. With such a large variation in cell size, the first crossover frequency is not suitable for separating these stages of breast cancer cells, especially within this current cDEP device. The
1st crossover frequencies are the same between early to intermediate stages of breast cancer, suggesting that the membrane electrical properties are not significantly different enough to produce a difference in DEP mobility and to drive effective separation. However, the potential to separate MCF10CA1 cells from early and intermediate stages at these low frequencies remains possible.

6.3 Future Work

Although the devices presented fell short of isolating different stages of breast cancer, the limits have not been reached. Future preliminary studies should be done on these four different breast cancer stages to determine their electric properties prior to moving on with testing the selectivity of a device. The cells properties can be determined using electrorotation spectroscopy and electroporation treatments which can measure, through direct analysis of the cell, the membrane capacitance and conductance, as well as the polarization charging time.

Knowing at the first crossover frequency that the selectivity of these breast cancer cells is unattainable using current cDEP designs only motivates for progress to be made studying either the second crossover frequency or material aspects of the microfluidic device itself. According to several authors, the membrane capacitance, cell radius, and media conductivity generally affect the first crossover frequency, but the conductivity of a cell’s cytoplasm plays a role in the second crossover frequency\[38, 61\]. Currently, experiments using cDEP at high frequencies in the MHz range has not been studied, and perhaps for these breast cancer cells the cytoplasm conductivities have more variation near the 2nd crossover frequency.

Peering at the design aspect of the microdevice, the electric field gradients play a major role in the selectivity of the cells. The devices used for ‘high frequency’ experiments were limited in that range because of the material properties of PDMS. If another material with a higher dielectric constant could be used to replace it, then the ‘high frequency’ devices may operate at low frequencies especially near the crossover frequency. More experiments may also be conducted with varying fluid conductivities and determining breast cancer cell stage response.

Another approach for using cDEP to separate breast cancer cells is to include a magnetic component. Han et al previously explored magnetic fields to attract RBC to a center channel while cancer cells were repelled to outer channels [36]. A preliminary experiment with cDEP would be to incorporate a magnetic center channel filled with a solution of ferromagnetic materials such as Fe, Ni, or Cobalt. Combining this with cDEP will help reduce the prep time needed to remove white blood and cancer cells from blood.
Chapter 7. Conclusions

Electrokinetics has demonstrated its capability in two separate biomedical applications: tissue engineering and cancer detection. The first application studied the effects of electric fields on bacteria cells as well as calcium ions. In response to the electric field, calcium ions were able to deposit simultaneously with cellulose growth. The benefit of using this technique is that alignment of cellulose fibers can occur as nucleation sites for mineralization are forming. Furthermore, there was localized deposition of ions at the electrode holes. The localization allows for a meniscus design to form horns that will initiate osteogenesis to the tibia that will anchor the structure. The other study in this application was determining the effects of an AC field on bacteria cells. At low frequencies, bacteria were showing a response to electroosmosis and DEP with a net motion caused by a DEP force created by the electric field gradients formed from the meniscus design.

In the second application, dielectrophoresis in a contactless DEP device was studied as a method for separating different stages of breast cancer all from the same person. Several geometries that altered the electric field gradients were studied to determine their selectivity. With both ‘high’ and ‘low’ frequencies, the cells were incapable of being distinguished. The cell’s membrane capacitance resulted in being very similar for each cell line and insignificant. Calculated membrane capacitances overlapped simply by the variation of cell size within each cell line. However, the most aggressive cell line showed potential for isolation from the other cells. Therefore, the current cDEP devices were not selective enough to show this isolation. Future work will be needed to study each cell’s cytoplasm response to electric fields, or in other words studies at high frequencies that near each cell’s 2nd crossover frequency.
References


Appendix A: MATLAB Code to Analyze Results from High Frequency cDEP Experiments

function image_analysis(file_name, varargin)
% this function analyzes cDEP devices to measure the number of cells entering and leaving a trapping region
% there are two lines

clc     % clear matlab command window (clean up)

    % get input variables if any
p = inputParser;
p.addRequired('file_name');
p.addParamValue('entrance_x_offset', 0);
p.addParamValue('entrance_y_offset', 0);
p.addParamValue('exit_x_offset', 0);
p.addParamValue('exit_y_offset', 0);
p.addParamValue('entrance_scale', 1);
p.addParamValue('exit_scale', 1);
p.addParamValue('threshold', 95);
p.addParamValue('line_length', 440);
p.addParamValue('video_length', 53);
p.addParamValue('start_analysis', 10);
p.addParamValue('end_analysis', 40);
p.addParamValue('dir', 'G:\cDEP Experiments Breast Cancer\06-13-11\')
p.parse(file_name, varargin{:});
p.Results;

% simplify variable names
entrance_x_offset = p.Results.entrance_x_offset
entrance_y_offset = p.Results.entrance_y_offset
exit_x_offset = p.Results.exit_x_offset;
exit_y_offset = p.Results.exit_y_offset;
entrance_scale = p.Results.entrance_scale;
exit_scale = p.Results.exit_scale;
cell_threshold = p.Results.threshold;
line_length = p.Results.line_length;
video_length = p.Results.video_length;
start_analysis = p.Results.start_analysis;
end_analysis = p.Results.end_analysis;
file_name = p.Results.file_name;
dir = p.Results.dir;

line_length

% Parameters

% cut line for cell evaluation
x(1,:) = [25 + entrance_x_offset, 25 + entrance_x_offset];
y(1,:) = [100+entrance_y_offset, (100+ line_length +entrance_y_offset)];
%exit
x(2,:) = [850 + exit_x_offset, 850 + exit_x_offset];
y(2,:) = [100+ exit_y_offset, (100+ line_length +exit_y_offset)];

% Read movie file in
read_file = strcat(dir, file_name, '.avi')
% addpath read_file
path(path,dir)
movie = mmreader(read_file);
% file_info = aviinfo('control.avi')
num_frames= movie.NumberOfFrames

%display the first frame and the cut line
mov(1).cdata = read(movie, 2);
frame = mov(1).cdata;
first_frame = rgb2gray(frame);
%plot frame
figure(1), imshow(first_frame);
hold on
%plot data line
plot (x(1,:),y(1,:),'-rx')
plot (x(2,:),y(2,:),'-bx')
% hold off
tic
distribution = [];
old_dist = [];
%use this to control the plotting of some
data later on
total_in = 0;
total_out = 0;
trapping_efficiency = 0;

fps = num_frames/ video_length %Frames Per Second
start_frame = ceil(fps* start_analysis)
end_frame = ceil(fps* end_analysis)

%split movie into individual frames
for cur_frame = start_frame:5:end_frame

%movie and frame data
clear mov;
mov(cur_frame).cdata = read(movie, cur_frame);
frame = mov(cur_frame).cdata;
A = rgb2gray(frame);

% A = A - first_frame; %uncomment this line to remove the image background

%plot frame
if(mod(cur_frame, 2)==1)
   figure(3), imshow(A);
   hold on
   %plot data line
   plot (x(1,:),y(1,:),'-rx')
   plot (x(2,:),y(2,:),'-bx')
   hold off
end

%% Analyze data here
%original pixel data
input_profile = improfile(A,x(1,:),y(1,:));
output_profile = improfile(A,x(2,:),y(2,:));
%plot pixel data
figure(4)
subplot(6,1,1),title(file_name),plot(input_profile, '-r')
hold on, plot(output_profile, '-b'),plot([0 length(input_profile)], [cell_threshold cell_threshold])
hold off

%smoothed pixel data
input = smooth(input_profile, 10);
output = smooth(output_profile, 10);

%plot smoothed pixel data
subplot(6,1,2),
if(mod(cur_frame, 2)==1)
   plot(input, '-r')
   hold on
   plot(output, '-b'), plot([0 length(output)], [cell_threshold cell_threshold]), hold off
   ylim([0,cell_threshold])
end

%thresholded pixel data
input_threshold = input>cell_threshold;
output_threshold = output>cell_threshold;

%time analysis
input_time(cur_frame) = sum(input_threshold);
output_time(cur_frame) = sum(output_threshold);

%plot time analysis
subplot(6,1,3)
   plot(input_time, '-r');
   hold on
   plot(output_time, '-b')
title(file_name);
   hold off

%trapping efficiency analysis
total_in = total_in + sum(input_threshold)
total_out = total_out + sum(output_threshold)
trapping_efficiency = ((total_in - total_out) / total_in) * 100;

subplot(6,1,5)
    bar([total_in, total_out])
    set(gca,'XTickLabel', ['cells in', 'cells out'])

subplot(6,1,6)
    bar(trapping_efficiency)
    ylim([0,100])
    set(gca,'XTickLabel', ['trapping efficiency'])
    title(int2str(trapping_efficiency))

%input distribution
if ~exist('input_distribution')
    input_distribution = zeros(size(input_threshold));
    input_distribution = input_distribution + input_threshold;
    old_input_dist = input_distribution;
else
    input_distribution = input_distribution + input_threshold;
end

%output distribution
if ~exist('output_distribution')
    output_distribution = zeros(size(output_threshold));
    output_distribution = output_distribution + output_threshold;
    old_output_dist = output_distribution;
else
    output_distribution = output_distribution + output_threshold;
end

if ( isequal(input_distribution,old_input_dist) && isequal(output_distribution, old_output_dist))
    %dont replot if distributions haven't changed
else
    %if distribution has changed, plot
    subplot(6,1,4)
    plot(input_distribution, '-r');
    hold on
    plot(output_distribution, '-b')
title(file_name);
    hold off
    pause(.1)
end

old_input_dist = input_distribution;
old_output_dist = output_distribution;

end
hold off

%
directory = 'output files/';
file = strcat(dir,directory,file_name, '.txt');
xLtitle = {'input_distribution ', 'output_distribution ', 'input_time ', 'output_time ', 'total_in ', 'total_out ', 'trapping_efficiency '}
xLdata = padcat(input_distribution, output_distribution, input_time, output_time, total_in, total_out, trapping_efficiency);
size(xLtitle)
size(xLdata)

% data = [xLtitle; xLdata]
% xlswrite(file, xLtitle, 'Sheet1', 'A1');
% save(file, 'xLdata', '-ascii')

% file = strcat(dir, directory,file_name, '.mat');
% save(file, 'input_distribution', 'output_distribution', 'input_time', 'output_time', 'total_in', 'total_out', 'trapping_efficiency', '-mat')

beep; pause(1);beep;

end

function analyze_all  % Calls the imagine_analysis function for each video
clear all;
close all;
clc

image_analysis('340 160V - 1','line_length', 280,'entrance_y_offset',75, 'exit_x_offset',8,'exit_y_offset',80,'threshold', 70,'dir', 'G:\cDEP Experiments Breast Cancer\06-13-11\')
Appendix B: MATLAB Code to Analyze Results from Low Frequency cDEP Experiments

function image_analysis(file_name, x_offset, y_offset, threshold)
%         close all
clc
%         clear all

cell_threshold = 80;
switch nargin
  case 1
    x_offset=0;
y_offset=0;
  case 2
    y_offset=0;
  case 4
    cell_threshold = threshold
end

% Parameters
display_every = 100;
%cut line for cell evaluation
x2 = [450+x_offset, 450+x_offset];
y2 = [155+y_offset, 460+y_offset];

% Read movie file in
%dir = 'H:\cDEP Experiments Breast Cancer\08-11-11\';
% dir = 'G:\cDEP Experiments Breast Cancer\08-13-11\';
%dir = 'F:\cDEP Experiments Breast Cancer\08-15-11\';
dir = 'H:\cDEP Experiments Breast Cancer\08-16-11\';

read_file = strcat(dir, file_name, '.avi')
%         addpath read_file
path(path,dir)
movie = mmreader(read_file);
% file_info = aviinfo('control.avi')
num_frames= movie.NumberOfFrames

frame = read(movie, 1);
A = rgb2gray(frame);
first_frame = A;
% plot frame
figure(1), imshow(A);
hold on
%plot data line
plot (x2,y2,'-rx')
% hold off

start_frame = 1;
end_frame = num_frames;
tic

distribution = [];
old_dist = [];
%use this to control the plotting of some data later on

%split movie into individual frames
for cur_frame = start_frame:10:end_frame-100

%movie and frame data
% clear mov;
% mov(cur_frame).cdata = read(movie, cur_frame);
% frame = mov(cur_frame).cdata;
% frame = read(movie,cur_frame);
A = rgb2gray(frame);
% A = A - first_frame; %This removes the background
% plot frame
if(mod(cur_frame, display_every)==1)
    figure(3), imshow(A);
    hold on
    %plot data line
    plot (x2,y2,'-rx')
    hold off
end

%xlabel(strcat(int2str(cur_frame), '/', int2str(end_frame)));

%%% Analyze data here
%original pixel data
c = improfile(A,x2,y2);

%plot pixel data
if(mod(cur_frame, display_every)==1)
    figure(4)
    subplot(4,1,1),title(file_name),plot(c)
    hold on, plot([0 length(c)], [cell_threshold cell_threshold])
    hold off
end

%smoothed pixel data
output = smooth(c, 10);

%plot smoothed pixel data
subplot(4,1,2),
if(mod(cur_frame, display_every)==1)
    plot(output, '-r'),
    hold on
plot([0 length(output)], [cell_threshold cell_threshold]), hold off
end

% threshold pixel data
threshold = output>cell_threshold;

if(size(distribution) == 0)
    distribution = zeros(size(threshold));
    distribution = distribution + threshold;
    old_dist = distribution;
else
    distribution = distribution + threshold;
end

if (distribution == old_dist)
else
    subplot(2,1,2)
    plot(distribution, '-b');
    title(file_name);
    hold off
end
    old_dist = distribution;
end
hold off
toc

directory = 'output files/';
file = strcat(dir,directory,file_name, '.txt');
save(file, 'distribution', '-ascii')

file = strcat(dir,directory,file_name, '.mat');
save(file, 'distribution', '-mat')

beep; pause(1);beep;
end

function analyze_all %Calls the image_analysis function to analyze the each
    clear all;
close all;

    % AT1 08-16-11
    %image_analysis('10 kHz 200V AT1_3',-70,40,20)
    %image_analysis('20 kHz 200V AT1_3',-70,40,20)
    %image_analysis('20 kHz 200V AT1_3 short',-70,40,20)
    %image_analysis('30 kHz 200V AT1_3',-70,40,20)
    %image_analysis('40 kHz 200V AT1_3',-70,40,20)
    %image_analysis('50 kHz 200V AT1_3',-70,40,20)
    %image_analysis('60 kHz 200V AT1_3',-70,40,20)
    %image_analysis('70 kHz 200V AT1_3',-70,40,20)
    %image_analysis('Control AT1_3',-70,40,20)
Appendix C: MATLAB Code for Plotting the Clausius-Mossotti Curve

```
function model_used_for_AndreaThesis_plus_minus
clc
clear all
close all
w=2*pi*logspace(2,9,100);
w2 = w.*w;

rnorm= 9.25e-6;  %radius of MCF10A from Henslee et al
rn10Ap=7.96e-6;  %radius of MCF10A found by Andrea
rn10Am=7.96e-6;
rn10A=7.96e-6;

rAT1=6.81e-6;    %radius of MCF10AT1 found by me(Andrea)
rAT1p=6.81e-6;   %radius of MCF10AT1 found by me(Andrea)
rAT1m=6.81e-6;   %radius of MCF10AT1 found by me(Andrea)
rDCIS=7.79e-6;   %radius of DCIS found by me(Andrea)
rDCISp=7.79e-6;  %radius of DCIS found by me(Andrea)
rDCISm=7.79e-6;  %radius of DCIS found by me(Andrea)
rCA1=6.88e-6;    %radius of MCF10CA1 found by me(Andrea)
rCA1p=6.88e-6;   %radius of MCF10CA1 found by me(Andrea)
rCA1m=6.88e-6;   %radius of MCF10CA1 found by me(Andrea)

conductivity = 0.01;% [S/m]

% Calculate C-M for each cell
for i=1:length(conductivity)
    knnorm(i,:) = norm(rnorm, w,conductivity(i));
    kn10A(i,:) = n10A(rn10A, w,conductivity(i));
    kn10Ap(i,:) = n10Ap(rn10Ap, w,conductivity(i));
    kn10Am(i,:) = n10Am(rn10Am, w,conductivity(i));
    kAT1(i,:) = AT1(rAT1, w,conductivity(i));
    kAT1p(i,:) = AT1p(rAT1p, w,conductivity(i));
    kAT1m(i,:) = AT1m(rAT1m, w,conductivity(i));
    kDCIS(i,:) = DCIS(rDCIS, w,conductivity(i));
    kDCISp(i,:) = DCISp(rDCISp, w,conductivity(i));
    kDCISm(i,:) = DCISm(rDCISm, w,conductivity(i));
    kCA1(i,:) = CA1(rCA1, w,conductivity(i));
    kCA1p(i,:) = CA1p(rCA1p, w,conductivity(i));
    kCA1m(i,:) = CA1m(rCA1m, w,conductivity(i));
end

% Plot the CM factor based on membrane capacitance std deviation pos and neg
figure('name', 'CM of Breast Cancer Cells by Stage');
semilogx(w/(2*pi),knorm(1,:), 'k', w/(2*pi), kn10Ap(1,:), 'r',w/(2*pi), kn10Am(1,:), 'r', w/(2*pi), kAT1p(1,:), 'b',w/(2*pi), kAT1m(1,:), 'b',w/(2*pi), 0, 'k', w/(2*pi), kDCISp(1,:), 'g',w/(2*pi), kDCISm(1,:), 'g',w/(2*pi), kCA1p(1,:), 'c',w/(2*pi), kCA1m(1,:), 'c')
xlabel('Frequency (Hz)')
ylabel('Re[K(\omega)]')
```

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hold off
end

function Knorm= norm(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10Aref Cells based on the conductivity of the media
% r = cell radius
% w = angular frequency
% sig_media = conductivity of the media
rnorm=9.25e-6;
w2 = 2*pi*logspace(2,9,100);
w = w.*w;
% radial frequency squared
e0=8.85e-12;
e1=80*e0;
sig1=100e-4; %sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.0194; %value from Henslee et al
tm=Cm*rnorm/sigc;
tc=ec/sigc;
t1=e1/sig1;
tmstar=Cm*rnorm/sig1;
j=sqrt(-1);
t1tm=t1*tm;
tctms=tc*tmstar;
top=w2.*(t1tm-tctms)+j.*w..*(tmstar-t1-tm)-1;
bottom=w2.*(2*t1tm+tctms)-j.*w..*(tmstar+2*t1+2*tm)-2;
Knorm=(-1).*top./bottom;
size(Knorm);
end

function Kn10Ap = n10Ap(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10Ap std dev (positive)Cells based on the conductivity of the media
w2 = w.*w; %radial frequency squared
e0=8.85e-12;
e1=80*e0;
sig1=103e-4; %sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.01269; %value calculated
tm=Cm*r/sigc;
tc=ec/sigc;
t1=e1/sig1;
tmstar=Cm*r/sig1;
j=sqrt(-1);
t1tm=t1*tm;
tctms=tc*tmstar;
top=w2.*(t1tm-tctms)+j.*w..*(tmstar-t1-tm)-1;
bottom=w2.*(2*t1tm+tctms)-j.*w..*(tmstar+2*t1+2*tm)-2;
Kn10Ap=(-1).*top./bottom;
size(Kn10Ap);
end

function Kn10Am = n10Am(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10Ap std dev (negative) Cells based on the conductivity of the media
w2 = w.*w; %radial frequency squared
e0=8.85e-12;
e1=80*e0;
sigl=103e-4; %sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.0163; %value calculated
tm=Cm*r/sigc;
tc=ec/sigc;
tl=e1/sig1;
tmstar=Cm*r/sig1;
j=sqrt(-1);
t1tm=tl*tm;
tctms=tc*tmstar;
top=w2.*(t1tm-tctms)+j.*w.*(tmstar-1-tm)-1;
bottom=w2.*(2*t1tm+tctms)-j.*w.*(tmstar+2*tl+2*tm)-2;
Kn10Am=((-1)*top./bottom;
size(Kn10Am);
end

function KAT1p= AT1p(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10AT1 (pos std dev) cells based on the conductivity of the media
w2 = w.*w; %radial frequency squared
e0=8.85e-12;
e1=80*e0;
sigl=104.5e-4; %sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.01488; %value calculated
tm=Cm*r/sigc;
tc=ec/sigc;
tl=e1/sig1;
tmstar=Cm*r/sig1;
j=sqrt(-1);
t1tm=tl*tm;
tctms=tc*tmstar;
top=w2.*(t1tm-tctms)+j.*w.*(tmstar-1-tm)-1;
bottom=w2.*(2*t1tm+tctms)-j.*w.*(tmstar+2*tl+2*tm)-2;
KAT1p=((-1)*top./bottom;
size(KAT1p);
end

function KAT1m= AT1m(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10AT1 (negative std dev) Cells based on the conductivity of the media
w2 = w.*w; %radial frequency squared
e0=8.85e-12;
e1=80*e0;
sigl=104.5e-4; %sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.01941; %value calculated
function KDCISp = DCISp(r, w, sig_media)
% this function returns an array of values for the Clausius-Mossotti factor
% for DCIS (positive std dev) cells based on the conductivity of the media
w2 = w.*w;  % radial frequency squared
e0=8.85e-12;
e1=80*e0;
sig1=107e-4;  % sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.01314;  % value calculated
tm=Cm*r/sigc;
tc=ec/sigc;
t1=e1/sig1;
tmstar=Cm*r/sig1;
j=sqrt(-1);
t1tm=t1*tm;
tctms=tc*tmstar;
top=w2.*(t1tm-tctms)+j.*w.*(tmstar-t1-tm)-1;
bottom=w2.*(2*t1tm+tctms)-j.*w.*(tmstar+2*t1+2*tm)-2;
KAT1m=-(1)*top./bottom;
size(KAT1m);
end

function KDCISm = DCISm(r, w, sig_media)
% this function returns an array of values for the Clausius-Mossotti factor
% for DCIS (negative std dev) cells based on the conductivity of the media
% r = cell radius
% w = angular frequency
% sig_media = conductivity of the media

w2 = w.*w;  % radial frequency squared
e0=8.85e-12;
e1=80*e0;
sig1=107e-4;  % sigma of DEP buffer ~100uS/cm
ec=50*e0;
sigc=1;
Cm = 0.01541;  % value calculated
tm=Cm*r/sigc;
tc=ec/sigc;
t1=e1/sig1;
tmstar=Cm*r/sig1;
j=sqrt(-1);
tltm = t1*tm;
tctms = tc*tmstar;
top = w2.*(tltm-tctms)+j.*w.*(tmstar-t1-tm)-1;
bottom = w2.*(2*tltm+tctms)-j.*w.*(tmstar+2*t1+2*tm)-2;
KDCISm = (-1)*top./bottom;
size(KDCISm);
end

function KCA1p = CA1p(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10CA1 (positive std dev) Cells based on the conductivity of the media
w2 = w.^2;
%radial frequency squared
e0 = 8.85e-12;
e1 = 80*e0;
sig1 = 107e-4;  %sigma of DEP buffer ~100uS/cm
ec = 50*e0;
sigc = 1;
Cm = 0.02349;  %value calculated
tm = Cm*r/sigc;
tc = ec/sigc;
t1 = e1/sig1;
tmstar = Cm*r/sig1;
j = sqrt(-1);
tltm = t1*tm;
tctms = tc*tmstar;
top = w2.*(tltm-tctms)+j.*w.*(tmstar-t1-tm)-1;
bottom = w2.*(2*tltm+tctms)-j.*w.*(tmstar+2*t1+2*tm)-2;
KCA1p = (-1)*top./bottom;
size(KCA1p);
end

function KCA1m = CA1m(r, w, sig_media)
%this function returns an array of values for the Clausius-Mossotti factor
%for 10CA1 (negative std dev) cells based on the conductivity of the media
w2 = w.^2;
%radial frequency squared
e0 = 8.85e-12;
e1 = 80*e0;
sig1 = 107e-4;  %sigma of DEP buffer ~100uS/cm
ec = 50*e0;
sigc = 1;
Cm = 0.02780;  %value calculated
tm = Cm*r/sigc;
tc = ec/sigc;
t1 = e1/sig1;
tmstar = Cm*r/sig1;
j = sqrt(-1);
tltm = t1*tm;
tctms = tc*tmstar;
top = w2.*(tltm-tctms)+j.*w.*(tmstar-t1-tm)-1;
bottom = w2.*(2*tltm+tctms)-j.*w.*(tmstar+2*t1+2*tm)-2;
KCA1m = (-1)*top./bottom;
size(KCA1m);
end