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# Electrical Detection of Cancer Biomarker using Aptamers with Nanogap Break-Junctions

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# Abstract

Epidermal Growth Factor Receptor (EGFR) is a cell surface protein overexpressed in cancerous cells. It is known to be the most common oncongene. EGFR concentration also increases in the serum of cancer patients. The detection of small changes in the concentration of EGFR can be critical for early diagnosis, resulting in better treatment and improved survival rate of cancer patients. This article reports an RNA aptamer based approach to selectively capture EGFR protein and an electrical scheme for its detection. Pairs of gold electrodes with nanometer separation were made through confluence of focused ion beam scratching and electromigration. The aptamer was hybridized to a single stranded DNA molecule, which in turn was immobilized on SiO<sub>2</sub> surface between the gold nanoelectrodes. The selectivity of the aptamer was demonstrated by using control chips with mutated non–selective aptamer and with no aptamer. Surface functionalization was characterized by optical detection and two orders of magnitude increase in direct current (DC) was measured when selective capture of EGFR occurred. This represents an electronic biosensor for the detection of proteins of interest for medical applications.

# Keywords

Epidermal Growth Factor Receptor (EGFR); Focused Ion Beam (FIB) milling; electromigration; DNA; Aptamer; Break-junction

# 1. Introduction

Evolution of proteomics has enabled better understanding of disease progression, and helped in establishing new biomarkers for early diagnosis, appropriate treatment and faster drug discovery. Overexpression of Epidermal Growth Factor Receptor (EGFR) is considered to

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be involved in several types of cancers. Its overexpression has been reported as an indication for many cancers like breast, lung, cervical, bladder, esophageal and ovarian cancer [1–6]. The presence of elevated or reduced levels of a biomarker can point out the presence or onset of a disease, specify the phase of that disease and enable monitoring of changes in homeostasis [7]. Elevated levels of EGFR in the serum of lung cancer patients has been reported before. The normal EGFR concentration can be in the range of 45–78 ng/ml, whereas it may be as much as an order higher in the lymph node metastasis of lung cancer patients (850 ng/ml) [8, 9]. The early detection of a few tens to hundreds of ng/ml EGFR can be important to improve the prognosis of the patients.

A number of approaches have been reported for the detection of EGFR and its interactions. These techniques incorporate immunohistochemical, flow cytometric, amperometric, mechanical and optical detection modalities [10–15]. Optical waveguides have also been used for label-free sensing to study cell biology and for biomolecular detection [16–19]. These methods of detecting proteins are generally categorized as labeled detection and labelfree detection methods. In the case of labeled detection, presence of proteins is confirmed by some secondary molecule (organic, inorganic or radioactive) that binds to the proteins. Optical detection utilizes a fluorescent tag attached to the proteins and the change in fluorescent intensity after binding signals the presence of proteins. Whereas electrical detection directly interrogates the presence of protein molecules. It entails efficient device fabrication in order to define contact structures with a molecular dimension separation. Charge transport properties of a single molecule have also been studied with such structures. Such works have been primarily to identify molecules that may be used as functional electronics components [20–22]. It was recently reported that a single molecule was captured by a probing aptamer between nanogapped single-walled carbon nanotubes (SWCNTs) and charge-transport properties were characterized [23].

There have been reports of many types of metal-molecule-metal structures to measure electrical behavior of molecules [22]. Such structures can be broadly classified as vertical device structures (VDS) and lateral device structures (LDS). VDS utilize a self-assembled monolayer of molecules grown on a metal surface which acts as one contact. The second contact is made through conductive probes of atomic force microscopes, scanning tunneling microscopes or crosswire arrangements [24-26]. LDS involves a pair of metal contacts with nanometer sized separation and the probes selective to protein molecules are immobilized between the contact structures. Such devices that contain a thin metal strip, in that a gap is created by some technique, are known as break-junctions. Such gaps can be formed through e-beam lithography, electrodeposition of metals or electromigration induced "breaking" of metal lines [27-31]. These techniques are either slow and costly, mainly because of e-beam writing of the whole structures/lines or have very low yield of functional devices. This article reports electrical detection of EGFR molecules by making use of a new class of break-junctions where a focused ion beam (FIB) is used to scratch a thin metal strip (made with optical lithography and lift-off), followed by conventional microscale probing and electromigration induced breaking to produce break-junctions with high throughput, greater yield, low cost and at exact spot, as recently reported [32].

Antibodies are mostly used to functionalize surfaces for protein binding and detection. The antibody-based devices are not well-suited for field-deployable or point-of-care modalities because these require specific ranges of temperature, humidity and ionic strengths of the buffer solution in order to retain their function [33]. Low ionic strength is necessitated to prevail over surface Debye screening but it causes poor interaction between target and surface probes [33]. Recent works have shown significant advancements introducing aptamers for several types of target molecules [34–41]. Aptamers are reported to be as

selective and sensitive as antibodies but exceedingly stable over changing temperatures, humidity and ionic concentrations [42].

This article reports cheap and rapid nano-electrode manufacturing and direct current detection of low concentration of EGFR (50  $\mu$ g/ml) binding to RNA aptamers. The effective attachment of RNA aptamers on a chemically modified silicon dioxide surface and EGFR binding to aptamer was detected from direct current electronic measurements. Fig. 1 shows a sketch of the sensor that describes the mechanism of the transduction. The inset shows a scanning electron microscope (SEM) micrograph of the actual device layout. Attachment of RNA aptamer to SiO<sub>2</sub> surface and RNA-protein complex interactions were verified with optical measurements as well.

# 2. Materials and methods

#### 2.1. Materials

The chemicals used for this work were 3'-Aminopropyltriethoxysilane (APTES); 1,4-Phenylene diisothiocyanate (PDITC); Ethanol; Isopropyl alcohol (IPA); Acetone; Dimethyl sulfoxide (DMSO); Pyridine; 6-amino-1-hexanol; N,N-dimethylformamide (DMF); N,Ndiisopropylethylamine (DIPEA); Phosphate buffered Saline (PBS); Magnesium chloride; Hybridization buffer solution and Diethylpyrocarbonate (DEPC) treated de-ionized (DI) water. The chemicals were obtained from Sigma-Aldrich (St. Louis, MO, USA). The 5'amine modified single-strand DNA (ssDNA) were purchased from Alpha DNA (Montreal, Quebec, Canada).

The target molecule was recombinant human EGFR/ErbB1 Fc Chimera (R&D Systems, Minneapolis, MN). The receptor, as per the supplier, was able to bind recombinant human EGF in a functional ELISA at  $K_d$ <8nM.

#### 2.2. Methods

The reported work included three distinct aims that were accomplished towards the final goal of a device that can detect EGFR in real-time. The first section involves the fabrication of nanogap break-junctions. The second one elucidates the immobilization of selective probes (aptamers) on the  $SiO_2$  substrates. The third and most important part demonstrates the integration of protein binding and its optical and electrical detection.

2.2.1. Controlled fabrication of nanogap break-junctions—A two-step photolithography process was carried out to define metallic structures on  $SiO_2$  chips. The first layer of pattern was used to create 3 µm wide lines of titanium/gold (thickness 50 Å/ 150 Å) using metal lift-off. The second step of photolithography defined  $200 \times 200 \ \mu m^2$ probing pads that were made of Ti/Au (thickness 100 Å/500 Å) and aligned to the far edges of lines made in first lithography/metallization/lift-off. Inset to Fig. 1 shows the device at this step. The metal lines were partially scratched by FIB milling process (at spot highlighted with arrow in inset to Fig. 1). These scratched parts were where the nanogap break-junctions were ultimately made. FIB scratching with gallium ions was done at 30 kV acceleration voltage, 1 pA milling current and 2 sec per micron scan rate of the beam to achieve optimum scratch depth. The current-voltage (I-V) characteristics across the metal lines were determined using Agilent 4155C semiconductor parameter analyzer on a probe station. The *I*–V measurements before and after the nanomanufacturing of metallic breakjunctions were recorded to demonstrate the electromigration effect. The essential advantage of FIB scratching followed by electromigration is to increase the yield of useful devices with desired gap between electrodes, make nanogap breaks at precise location on the line, and

narrow the distribution of the nanogap sizes. These are not possible with approaches involving only electromigration [21, 43–45].

**2.2.2. Aptamer preparation**—Aptamer was isolated with a well-known process called "Systematic Evolution of Ligands by Exponential Enrichment" (SELEX) as reported before [46, 47]. An iterative selection of binding species against purified human EGFR (R&D Systems, Minneapolis, MN) was done to isolate anti-EGFR RNA aptamer [48, 49]. The high affinity ( $K_d$  = 2.4 nM) anti-EGFR RNA aptamer and a non-functional mutated aptamer were extended with a capture sequence. The extended anti-EGFR aptamer (5'-GGC GCU CCG ACC UUA GUC UCU GUG CCG CUA UAA UGC ACG GAU UUA AUC GCC GUA GAA AAG CAU GUC AAA GCC GGA ACC GUG UAG CAC AGC AGA GAA UUA AAU GCC CGC CAU GAC CAG-3'); extended mutant aptamer (5'-GGC GCU CCG ACC UUA GUC UCU GUU CCC ACA UCA UGC ACA AGG ACA AUU CUG UGC AUC CAA GGA GGA GUU CUC GGA ACC GUG UAG CAC AGC AGA GAA UUA AAU GCC CGC CAU GAC CAG-3') and amino modified substrate-anchored probe ssDNA (5'-amine-CTG GTC ATG GCG GGC ATT TAA TTC-3') were used. The capture sequence has been highlighted in bold font. The capture sequence was the modification of isolated anti-EGFR aptamer and the mutant aptamer. This was made by extending the DNA template at its 3' end and had 24 nucleotides. The extended capture sequence didn't disrupt the aptamer structure but was used to immobilize aptamers on the surface through duplexformation with the surface-bound ssDNA probe molecules.

#### 2.2.3. Immobilization of ssDNA and probing aptamers on the SiO<sub>2</sub> chip

surface—A thermally oxidized silicon wafer was diced into 4×4 mm<sup>2</sup> small chips. The chips were initially cleaned in oxygen plasma for 15 minutes using  $Ar+O_2$  at 200 W. This also resulted in a hydrophilic SiO<sub>2</sub> surface. The chips were then immediately immersed in a solution of 2% APTES in ethanol for 1 hour at room temperature to get the surfaces silanized. The chips were then sequentially rinsed with IPA and DI water and dried in nitrogen flow, followed by curing at 115 °C for 30 minutes. The chips were then submerged in a solution of 1 mM PDITC in DMSO containing 10% pyridine for 5 to 7 hours at 45 °C in order to get PDITC homobifunctional linker molecules attached onto the silanized surface. The chips were then washed sequentially with ethanol and DI water and dried in nitrogen flow. A volume of 8  $\mu$ l of amine modified ssDNA solution (5  $\mu$ M concentration of ssDNA in DI water with 50% DMSO, 1% Pyridine) was placed on each substrate and allowed to incubate in a humidity chamber for 16 to 18 hours at 45 °C to attach amino modified ssDNA to PDITC. After ssDNA attachment, each chip was rinsed sequentially with ethanol and DEPC treated DI water and dried with nitrogen. The functionalized surface was then deactivated and unbound reactive groups of PDITC capped by submerging the chips in a blocking buffer (BB) solution that contained 50 mM 6-amino-1-hexanol with 150 mM DIPEA in DMF for 1 hour. The chips were then rinsed with DMF, ethanol, DEPC treated DI water and dried in nitrogen stream. After that, a volume of 8 µl of anti-EGFR RNA aptamer solution was placed on every substrate for hybridization by incubating the chips in the buffer solution (100 nM of anti-EGFR aptamer with 5:1 of DI water and hybridization buffer) at 42 °C for 1 hour. The chips were then washed thoroughly with DEPC treated DI water and dried with nitrogen gas. The capture extension of the anti-EGFR RNA aptamer hybridized with the surface bound complementary ssDNA. In one set of controls, the mutant aptamer was hybridized to surface-bound ssDNA on separate chips using identical protocol. The presence of ssDNA and RNA aptamers immobilized on the SiO<sub>2</sub> surface was determined by fluorescence measurements of Acridine Orange (AO) stain at an excitation wavelength of 460 nm and the emission wavelength of 650 nm using a confocal microscope. Another set of controlled chips had no aptamer but underwent all other steps like the aptamer chips.

**2.2.4. Protein binding to the probing aptamer**—Chips with the attached RNA aptamer were then incubated in 50 ng/ $\mu$ l of EGFR protein prepared in PBS with 5 mM Mg<sup>2+</sup> at 37 °C for 45 minutes by putting 8  $\mu$ l of solution on each substrate. The chips were washed thoroughly with PBS and DEPC treated DI water and dried in the nitrogen flow. The chips were analyzed for the captured EGFR protein by optical detection of fluorescent stain *Sypro Ruby Protein Blot* at 488 nm wavelength. The fluorescence intensity analysis was done with *ImageJ* software [50]. Essentially, the surface bound ssDNA was used to immobilize an anti-EGFR aptamer and subsequently the aptamer was used to capture the EGFR protein.

After confirming the attachment chemistry for RNA immobilization and selectivity of anti-EGFR aptamer on plain chips, electrical detection of EGFR protein was performed using nanomanufactured metallic break-junction devices on oxidized silicon chips. An exactly identical attachment chemistry as for plain chips was employed for these devices. Although the concentration of EGFR used was high but the theoretical limit of detection (LOD) of the device is much smaller. As a first order estimate, the protein footstep is about 11 nm and in a nanogap 50 nm long and 3  $\mu$ m wide, we would need about 400 EGFR copies to fill the whole area of nanogap [51]. However, surface charges at nanoscale, time of detection i.e. time it would take for EGFR to reach the capture region just by diffusion [52], steric hindrance, conformation of molecules will all contribute to LOD of this framework.

Semiconductor parameter analyzer recorded the direct current tunneling through the nanometer scale electrode separation before and after the protein binding to the probing aptamers in between these nano-electrodes.

# 3. Results and discussion

#### 3.1. Fabrication and characterization of nanometer-sized break-junctions

Optimum FIB scratches were obtained by utilizing our previously reported protocol [32]. The I–V data recorded after FIB scratching of the metal lines showed a linear ohmic behavior that confirmed electrical and physical continuity of the metal lines (Fig. 2(a)). A voltage sweep from 0 to 4 V broke the metal lines due to electromigration. Fig. 2(b) shows a sudden drop in current that depicts an absolute break in the metal line. The SEM micrographs also showed a complete break in the metal lines with nano-scale separation (shown clearly in the inset of Fig. 2(c)). Figure 2(d) demonstrates the comparison of I-V data recorded for the same voltage range (-1 to +1 V) before and after the application of electromigration-inducing bias. All data and micrographs of Fig. 2 are representative for the rest of the devices on the chips.

A very small and random current flow was observed after the electrode separation that showed tunneling current characteristics. The tunneling current in an arrangement of the nanogap electrodes with vacuum as the insulator between them can be approximated as a function of the average barrier height relative to the Fermi level of the negative electrode, the barrier width and the applied voltage across the nanoelectrodes [53].

#### 3.2. Optical detection of immobilized aptamer and captured protein

The surface binding and the presence of ssDNA and RNA aptamer immobilized on the plain  $SiO_2$  chips was confirmed by AO stain and the fluorescence measurements. Data of Fig. 3(a) confirms the binding of ssDNA to the functionalized plain  $SiO_2$  surface while Figure 3(b) depicts the hybridization of RNA aptamer to the surface-bound ssDNA. AO stain gives green fluorescence when it interacts with ssDNA whereas a flame-red fluorescence is observed when it interacts with double stranded nucleic acids [54]. The AO stain binds electrostatically to the nucleic acids as it carries positive charge. Electrostatic

communication with non-specific polyanions was avoided by using a very low concentration of AO stain (0.2% v/v) and providing a buffer solution with  $Mg^{+2}$  and  $Na^+$  cations that competed binding to the nucleic acids [55].

EGFR binding to the probing aptamer onto the plain chip surface was confirmed with the Sypro Ruby protein gel stain. Sypro Ruby is a ruthenium based stain used to detect the amino acids lysine, arginine & histidine [56]. Figure 3(c) shows a clear enhancement of fluorescent intensity measured from the chips with anti-EGFR RNA aptamer probes and EGFR protein as compared to that for negative control chips that included probe-functionalized chips not exposed to the target proteins and the chips exposed to the target protein but without probe aptamer immobilized on the surface. The fluorescent intensity was at a maximum when use of a blocking buffer (BB) was omitted and therefore, unreacted PDITC moieties also contributed towards protein capturing. On the other hand, probe-functionalized chips with BB treatment selectively captured the EGFR protein as revealed by a marked increase in fluorescence intensity. The BB was thus used in subsequent experiments to avoid non-specific adsorption/binding on gold or oxide surfaces,

#### 3.3. Electrical detection of EGFR binding

Fig. 4(a) shows representative data from one device that shows the electronic detection of selective EGFR binding. The control chips with exactly same devices but with mutant aptamer or no aptamer at all showed no change in conductivity (Figure 4(b)). The scrambled sequence of a mutant aptamer prevented it from recognizing the EGFR whereas anti-EGFR RNA aptamer selectively captured EGFR. The I-V measurements recorded from -1 to +1 V across the metal break-junctions, showed a significant increase in current after the capture of EGFR by surface immobilized aptamers. There were two orders of reduction in the resistance through the the nanoelectrodes indicating the conducting behavior of proteins that bridged the nanogap between the electrodes. The yield of devices was 60% that can be possibly increased by tagging the protein molecules with conducting nanoparticles, e.g. of gold. Serum EGFR levels elevate in many cancers and aptamers isolated through SELEX show high-affinity binding to these. The non-specific binding of the aptamer with non-target serum proteins has already been studied and found to be very atypical [8, 57–59]. Therefore the presented data shows the power of this electronic biosensor for direct detection of EGFR present in human serum or plasma.

#### 4. Conclusion

A metallic break-junction framework with precise nanometer-sized separation is presented that utilizes aptamers for selective EGFR capture. The aptamer specifically binds, recognizes and captures EGFR in break-junctions. The capture is electrically detected. The integration of FIB scratching with electromigration provides rapid, cheap and controlled manufacturing of nanoscale break-junctions with high yield. The presence of protein between the electrodes shows a robust increase in conductivity between otherwise insulated nanoelectrodes. The detection of a disease linked protein can help in early diagnosis and improved therapy. Such proteonic chips can find applications in many electrical sensors for detection of biologically relevant molecules and species.

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# Figure 1.

A 3D sketch of the device that demonstrates the mechanism of the electronic detection (not to scale). Inset shows an SEM micrograph of the thin Au line on  $SiO_2$  chip.



Figure 2(a)



Figure 2(b)



Figure 2(c)



Figure 2.

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Characterization of the device (a) Representative I-V data after FIB scratching of metal line shows linear ohmic behavior. (b) A sudden drop in current confirms the complete breaking of metal line due to electromigration and results in nanogap break-junctions. (c) SEM micrograph showing a clean break between nanoelectrodes. Inset shows magnified view of a break-junction with a gap smaller than 30 nm. (d) Comparison of I-V data for a representative break-junction before and after the electromigration.







#### Figure 3.

(a) Acridine Orange (AO) stain intensity measurements (background subtracted) depicts the presence of surface-bound DNA by comparing it with control chips. (n=10). (b) Fluorescence intensity measurements (background subtracted) using AO confirms the RNA hybridization to surface bound ssDNA when compared to control chips (n=10). (c) Sypro staining shows the selective capture of EGFR on functionalized SiO<sub>2</sub>. From L to R: Chip # 1: Control chip treated with blocking buffer (BB) but no aptamer; Chip # 2: Control chip

with no BB treatment and no aptamer; Chip # 3: Control chip functionalized with mutant aptamer and treated with BB; Chip # 4: Functionalized chip with EGFR aptamer and captured protein after BB treatment; Chip # 5: Functionalized chip with EGFR aptamer and captured protein without using BB. (Error bars represent the standard deviation for n = 10)



#### Figure 4.

Comparison of I-V data for a representative break-junction before and after the surface modification and exposure to EGFR: (a) Anti-EGFR aptamer chip shows current increase due to the capture of EGFR bridging the nanogap; (b) Mutant aptamer functionalized control chip shows no change in conductivity, inset shows data for control chip with no aptamer and thus no change in conductivity. Both controls show no capture of EGFR.