Multiplexed, label-free detection of biomarkers using aptamers and Tunable Resistive Pulse Sensing (AptaTRPS).

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Multiplexed, label-free detection of biomarkers using aptamers and tunable resistive pulse sensing (AptaTRPS)
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Abstract:
Diagnostics that are capable of detecting multiple biomarkers are improving the accuracy and efficiency of bioassays. In previous work we have demonstrated the potential of an aptamer-based sensor (aptasensor) utilising Tunable Resistive Pulse Sensing (TRPS). Here, we have advanced the technique identifying key experimental designs for potential POC assays. The assay utilised superparamagnetic beads, and using TRPS monitored their translocations through a pore. If the surfaces of the beads are modified with an aptamer, the frequency of beads (translocations/minute) through the pore can be related to the concentration of specific proteins in the solution. Herein, we have demonstrated the successful use of TRPS to observe the binding of two proteins, to their specific aptamers simultaneously. We describe a series of experiments illustrating key factors which we believe are integral to bead-based assays and demonstrate a general method for a multiplexed assay. In summary, we have explored the effects of beads size, concentration, potential bias, pH and aptamer affinity to enhance the sensitivity and practically of a TRPS aptasensor. The method utilises the fact the binding of the aptamer to the protein results in a change in charge density on the bead surface, the isoelectric point of the protein then dominates the mobility of the beads, creating a multiplexed assay termed AptaTRPS. By alteration of the applied potential to the instrument it is possible to produce a positive signal in a simple multiplexed assay.

Keywords: Aptasensor, aptamers, biomarkers, Tunable Resistive Pulse Sensing, Point of Care
1. Introduction:

In recent decades there has been an increase in interest in rapid, affordable and user-friendly diagnostic techniques; driven by the rise in interest in personalised medicine, reduced assay times and the desire to provide home diagnostics (Chan et al., 2013). To make personalised medicine available to everyone, it is necessary to minimise the cost and labour intensity of assays so that they are affordable and available regardless of geographical location (Yager et al., 2008). Additionally, it is also desirable to minimise the time taken to yield results of an assay and to move towards point-of-care (POC) diagnostics and monitoring to improve decision-making. Key considerations in the choice of an appropriate technology are the ability to multiplex, assay time, cost, reliability and simplicity (Song et al., 2014).

Gold standard methods of analysis still heavily rely on PCR or antibody based technologies. A number of concerns remain with antibodies, as they: add cost and complexity to any assay; require the use of animals or cell culture for their generation; are susceptible to batch-to-batch variations; cause difficulties in handling and storage; and can have less than ideal sensitivity (Ledur et al., 1995; Tombelli et al., 2005). Their integration in POC technologies often gives the device a limited shelf life, and test substrates require refrigeration. A viable and accessible alternative to antibodies, addressing these concerns, would be a transformative technology. Aptamers are rapidly gaining interest as one alternative to antibodies (Fang and Tan, 2010; Levy-Nissenbaum et al., 2008; Mayer, 2009; Rowe et al., 2009). They offer an increased stability and ease of synthesis, they are far more cost-effective and reliable than their antibody counterparts and can easily be incorporated into diagnostic platforms. Full reviews on aptasensors can be found (Chiu and Huang, 2009; Cho et al., 2009; de-los-Santos-Álvarez et al., 2008; Famulok and Mayer, 2011) and whilst they have been criticised for repetition of choice in targets (Cho et al., 2009), this is likely due to authors keeping with tried and tested high affinity ligands to enable greater comparative power between new platforms.

One of the more recent technologies to implement aptamers is Resistive Pulse Sensing (RPS) (Rotem et al., 2012). A more recent form of RPS uses a polyurethane elastomeric membrane and is known as Tunable Resistive Pulse Sensing (TRPS) (Platt et al., 2012) in which the pore is able to be mechanically manipulated in
real time to alter pore geometry (Roberts et al., 2010; Vogel et al., 2011). In brief, the set up and theory for RPS technologies is as follows: a stable ionic current is established by two electrodes, separated by a pore; as beads/analytes translocate the pore they temporarily occlude ions, leading to a transient decrease in potential known as a “blockade event”, examples of which can be seen in Figure 1C. In the current arrangement, the pore is mounted laterally so that particles typically move from the upper fluid cell into the lower fluid cell, aided by an inherent pressure head due to 40 µl of liquid in the upper fluid cell of approximately 50Pa (Willmott et al., 2012), and a positive or negative bias is applied via an electrode under the pore (see figure 1B). By monitoring changes in full width half maximum (FWHM), peak magnitude ($\Delta i_p$) and peak frequency (events/min) it is possible to elucidate the zeta potential (Vogel et al., 2012), size (Vogel et al., 2011), and concentration (Willmott et al., 2010) of colloidal dispersions in situ, as described in more detail elsewhere (Vogel et al., 2011). TRPS is well suited to detect DNA and DNA-protein interactions; previous work has provided proof-of-concept studies on the technology (Billinge et al., 2014) in which the velocity and frequency of aptamer-tagged beads changes upon the addition of the target analyte as the aptamer folds to a specific conformation whereby the target binds, shielding the surface charge and altering double layer structure, as has been demonstrated with the target thrombin (Billinge et al., 2014). A significant development to this method would be the ability to detect multiple proteins at once.

In the present study we develop an aptasensor to haemostatic proteins Vascular Endothelial Growth Factor (VEGF) and Platelet Derived Growth Factor (PDGF) which are both involved in various disease processes such as cancer (Wei et al., 2003), atherosclerosis (Matsumoto and Mugishima, 2006), and Alzheimer’s (Tarkowski et al., 2002). Although circulating levels of VEGF have been found to be variable between individuals, between 9 – 150 ng/L (Jelkmann, 2001), VEGF levels have been demonstrated to be elevated in several aggressive types of cancer, such as ovarian (Li et al., 2004), haematological (Belgore et al., 2001) and gastric carcinoma (Karayiannakis et al., 2002) and VEGF measurement has been highlighted as both a potential prognostic or diagnostic marker (Jelkmann, 2001) and for comparison between pre- and post-operative cancer states (Karayiannakis et al., 2002; Li et al., 2004). PDGF levels have also been found to be elevated in several
different forms of cancer and has recently been added to the Glasgow Prognostic Index, which measures inflammatory mediators, as an important indication of tumour and significant predictor of survival (Hamilton et al., 2014). Previous studies have measured both VEGF and PDGF levels in cancer patients undergoing anti-VEGF treatment (Madsen et al., 2012), however this was performed over two different systems, increasing the cost and time considerations of the assay.

To monitor two analytes simultaneously in one-step analysis we use the ability of TRPS to provide accurate particle-by-particle analysis. Its ability to accurately calculate the beads size as it translocates the pore, allows the data from each particle set to be separated according to their size. By using two differently sized beads and separately functionalising them with anti-VEGF or anti-PDGF aptamers, as displayed in figure 1A, it is possible to separate the individual signals based on $\Delta i_p$, figure 1C, and monitor changes to their frequency and speed, relating them back to the target analyte concentration.
Figure 1: A Schematic of the multiplexed experimental process; two beads of different sizes are conjugated to either VEGF or PDGF. B Beads are incubated with a combination of concentrations of VEGF and PDGF for 30 minutes; as they bind to the target their velocity and frequency through the pore is monitored. C Recorded “resistive peaks” for two different sized beads; the blockade magnitude ($\Delta i_p$) of each peak is proportional to the volume of each bead and can therefore be calibrated to provide the diameter of each individual bead.
2. Materials and Methods:

**Chemicals and reagents**

The following chemicals were sourced from Sigma Aldrich, UK, without any further purification unless otherwise stated: Phosphate Buffered Saline (PBS – P4417), Tween 20 (P1379), Bovine Serum Albumin (BSA – A2153), and thrombin (T7513). DNA sequences were obtained from Sigma Aldrich’s custom oligonucleotides service as lyophilised powders and made up to a stock concentration of 100pmol/µL: 5’ CAG GCT ACG GCA CGT AGA GCA TCA CCA TGA TCC TG-Biotin-3’ (anti-PDGF-BB aptamer)(Deng et al., 2013), 5’ ATA CCA GTC TAT TCA ATT GGG CCC GTC CGT ATG GTG GGT GTG CTG GCC AG-Biotin-3’ (VEa5 anti-VEGF aptamer)(Hasegawa et al., 2008). 5’ TGT GGG GGT GGA CGG GCC GGG TAG ATT TTT-Biotin-3’ (V7t1 anti-VEGF aptamer)(Nonaka et al., 2010). Human recombinant PDGF-BB and human VEGF165 were purchased from Life Technologies, UK (PHG0044 and PHC9394, respectively). Water purified to a resistivity of 18.2 MΩ cm (Maxima) was used to make all solutions unless otherwise specified and 1xPBST (0.05% Tween) was used as the buffer.

**Beads**

Streptavidin modified superparamagnetic beads of 120 nm (bio-adembeads streptavidin plus 0321) and 300 nm (bio-adembeads streptavidin 0313) in diameter were purchased from Ademtech (France). Binding capacities of 120 nm and 300 nm beads according to the manufacturer were 3512 pmol/mg and 477 pmol/mg respectively. Carboxyl beads of known concentration and diameter (SKP200; mode diameter 200nm) were sourced from Izon Sciences (Oxford, UK) to be used as a calibrant.

**Aptamer functionalization**

Aptamers were received as a lyophilised powder and made up to a stock concentration of 100µM with deionised water, these were then separated into aliquots for storage in the freezer so that sequences were only frozen and thawed once. The 120 and 300 nm beads were vortexed for 1 minute, sonicated for 1 minute and diluted to a concentration of 3 x 10^9 beads/mL in PBST. The streptavidin coated beads were incubated with a 10-fold excess of the desired biotinylated aptamer at room temperature on a rotary wheel for 30 minutes to allow the streptavidin to react with the biotinylated ends of the DNA. The functionalised beads were then removed from the rotary wheel and centrifuged for 2 minutes at 10 000 rpm before being
placed onto a MagRack (Life Sciences). After 2 minutes the beads had formed a visible cluster in the sample vial adjacent to the magnet and the solution was carefully removed and replaced with an equal volume of PBST – this wash stage was performed twice to remove any excess aptamer which would lower the sensitivity of the binding assay. These could then be stored in the fridge.

**Individual protein measurements**

Beads functionalised with the aptamers were divided into several vials, to which varying volumes of the protein stock solution was added to each vial to create samples of aptamer modified beads mixed with a range of protein concentrations. In all cases the volume of the solution was adjusted and kept constant by adding PBST. Each sample was left on a rotary wheel for 30 minutes to allow the protein to associate before being introduced into the TRPS instrument.

**Multiplexed analysis of PDGF-BB and VEGF**

120 nm beads with V7t1 aptamer and 300 nm beads with PDGF-BB aptamer were prepared as described above at a bead concentration of $6 \times 10^9$ beads/mL. 50 µL of each of the functionalised beads were placed into vials and combinations of concentrations of PDGF-BB and VEGF$_{165}$ were added. Each sample was left on a rotary wheel for 30 minutes to allow the protein to associate before being introduced into the TRPS instrument. Each sample was analysed via TRPS for the maximum run time of 10 minutes. The peak traces were calibrated and separated according to their size for independent analyses. Each individual blockade trace was examined for shape and directionality before being included in the sample set.

**Tunable Resistive Pulse Sensing (TRPS)**

Measurements were made with the Izon qNano system, purchased from Izon Sciences (Oxford, UK) which incorporates the fluid cell, stretching apparatus, data recording and analysis software (Control Suite V2.2.2.117). The pores used were designated “np200” by the manufacturer and are described as most suitable for detecting beads in the range 100 – 300 nm. A stretch of 46 mm and voltage of +/-0.4 V were chosen so that blockade events were clearly visible above the level of noise (<10pA); these conditions were maintained throughout the experiments so that data was comparable. 80µL of electrolyte buffer (PBS) was carefully pipetted into the lower fluid cell, taking care not to introduce bubbles. The upper fluid cell was then attached to the instrument and 40 µL of PBST was added to enable confirmation of a stable baseline and clean pore. When a positive current is applied (+0.4 V) peaks
are visible as a downward deflection from the baseline, when a negative current is applied (-0.4 V) peak were visible as upward deflections of the baseline.

**Data Analysis**

**Removal of reverse peaks**

Raw data was analysed by examining each individual blockade trace visually using the Izon ControlSuite V2.2.2.117 Software. When resistance was highest at the beginning of this peak and tailed toward the baseline with time this was considered a “normal” translocation. When the current increased gradually and then dropped sharply this was considered a “reverse” translocation and removed from subsequent data processing. Examples of these are shown in Supplementary Figure 1.

**Extraction of the particle rate for each bead population**

Each bead records a blockade magnitude that reflects its volume, for example 120 nm Ademtech beads typically produce a blockade magnitude of 0.17 nA with the pore setup used here. Data which had been visually inspected as described above was then calibrated with beads of known diameter and narrow distribution. Prior to any analysis the Ademtech beads were calibrated separately to accurately measure the range of particle sizes in the provided batches and these ranges were used to provide filter information for data processing of experimental samples. Peaks recorded as being between 100 and 200 nm were recorded as “120 nm” and peaks between 210 and 400 nm were recorded as “300 nm”. Data was then extracted as a .csv file containing data for each particle and the particle rate was then calculated by counting the number of beads observed divided by the recording time.
3. Results and Discussion:

**A**

![Figure 2: A scatter-plot displaying average rate of four experiments using 120nm, $3 \times 10^9$ beads/mL, beads functionalised with VEa5 (anti-VEGF aptamer) against concentration of VEGF$_{165}$, circles represent raw data, diamonds are values where the double peaks have been removed as described within the methods section. The error bars show 1 Standard deviation from the mean.](image)

**B**

![Example peak trace for (i) a bead travelling in top-to-bottom (see supplementary figure 1 for more examples) direction and (ii) a bead travelling bottom-to-top, or one not completely traversing the pore i.e. Bottom-pore-bottom.](image)

**Figure 2:** A scatter-plot displaying average rate of four experiments using 120nm, $3 \times 10^9$ beads/mL, beads functionalised with VEa5 (anti-VEGF aptamer) against concentration of VEGF$_{165}$. Circles represent raw data, diamonds are values where the double peaks have been removed as described within the methods section. The error bars show 1 Standard deviation from the mean. **B** Example peak trace for (i) a bead travelling in top-to-bottom (see supplementary figure 1 for more examples) direction and (ii) a bead travelling bottom-to-top, or one not completely traversing the pore i.e. Bottom-pore-bottom.

**VEGF and PDGF detection**

In the final assay the aim is to quantify both VEGF and PDGF simultaneously, we therefore chose to anchor each aptamer onto a different sized bead to allow the signals to be separated. Typically the VEGF aptamers were immobilised onto 120nm beads and the PDGF aptamers onto 300nm beads, as shown in figure 1.
Initially the TRPS instrument was set up with a positive bias applied to the electrode under the nanopore (lower fluid cell) similarly to that described in our previous work (Billinge et al., 2014). Samples containing 120 nm beads functionalised with VEa5 aptamer which had been incubated with several concentrations of VEGF_{165} (as described in the experimental section) were introduced to the TRPS instrument; the resultant data is displayed in Figure 2A. Initially the rate was predicted to diminish with increasing protein concentration, as observed for thrombin protein during a similar experimental design (Billinge et al., 2014); this was thought to due to the protein binding to the DNA and shielding the charge on the phosphate backbone. As we increase the concentration of VEGF from 0 to 10nM we see the predicted decrease in particle rate, however at higher concentrations, > 10nM, the particle rate increases before reaching a steady value of ~220 beads per minute. We examined the current traces in more detail and observed numerous double peaks, shown in figure 2Bii. The conical pore used in TRPS gives rise to a characteristic peak shape, displayed in figure 1 and explained in detail elsewhere (Willmott et al., 2012). The shape of the peak gives information on the direction which the particle is travelling. For beads travelling from the top of the fluid cell entering the narrow sensing zone first before then traveling through the pore into the expanding conical region they produce a peak shape as shown in figures 1 and 2Bi (more examples are displayed in supplementary figure 1). If a particle was to travel in the reverse direction the peak shape would be inverted (figure 2Bii and supplementary figure 1). Below 10 nM no reverse peaks were observed throughout the experiment, however when the concentration of VEGF increased, > 10nm, there was an increase in the number of peak traces, termed “reverse peaks” which represent beads coming back up from the base of the pore (figure 2Bii). The presence of these reverse peaks led to an increase in the rate recorded by the software. When these erroneous peaks were then removed from the data set, the rate is seen to diminish, as expected, with increasing VEGF_{165} concentration, see figure 2A black diamonds. We repeated the experiment with a second aptamer, termed V7t1, and the same effect was observed (supplementary figure 2). The two selected anti-VEGF aptamers bind to different regions of the VEGF protein – VEa5 binds the heparin binding domain (Hasegawa et al., 2008) whereas V7t1 forms a g-quadruplex and interacts with the receptor binding domain of VEGF (Nonaka et al., 2010), both of these aptamers produced reversed peaks at higher analyte concentrations and provide evidence that this observation
was not due to a specific aptamer-epitope interaction. We performed and rechecked an assay using the previously published thrombin aptamer (data not shown) and no reverse peaks were observed at higher thrombin concentrations.

In a typical TRPS experiment it has been shown that the number of bead translocations is a function of the bead concentration, diffusion coefficient of the beads, applied pressure, electroosmotic and electrophoretic mobility. Thus within the current experimental setup, ensuring concentration, pressure and osmotic contributions remain constant, changes in particle translocations can be inferred to a change in the electrophoretic mobility of the beads (Billinge et al., 2014). We attributed our observation of the beads re-entering the pore and moving from the lower fluid cell back up to the upper to the addition of the protein and its charge. VEGF\textsubscript{165} has a reported isoelectric point at pH 8.5 (Ferrara et al., 1992), suggesting that at the pH of 7.4 the protein should have a net positive surface charge, whereas thrombin (pI 7.1 (Righettl and Tudor, 1981)) would be largely neutral. Upon binding to the aptamer and shielding the phosphate backbone the charge upon the protein must therefore dominate the electrophoretic mobility. As this occurred with both VEGF aptamers, which as stated above bind to different regions of the protein, it does not seem to affect the results which epitope is occupied.

To test the hypothesis we switched the applied voltage bias to a negative value (negative electrode, in the lower fluid cell). In this configuration DNA modified beads usually do not traverse the pore opening, or have a much reduced frequency of translocation, here for the 120nm particles no translocation events were recorded. Upon incubation with the VEGF protein the frequency of beads was directly proportional to the concentration of the protein, shown in Figure 3. This is a major improvement and observation from previous thrombin work (Billinge et al., 2014). In thrombin studies the number of translocation events per minute decreases as target analyte concentration increased, in effect switching off the signal. Here we have a more typical and convenient setup, wherein an increase in analyte concentration produced an increase in signal.

A series of experiments were then performed to confirm that PDGF could also be detected using this technique. First a positive bias was applied to the lower fluid cell of + 0.4 V, and a diminution of particle rate was observed as PDGF concentration increases. Unlike within the VEGF experiment where beads were observed translocating the pore in reverse i.e. from the lower fluid cell to the upper, as
displayed within supplementary figure 3 using 300 nm beads. To ascertain if this effect was dependent upon the bead size i.e. due to additional mass of the 300nm beads creating bias given the larger gravity forces, the same experiment was performed using 120 nm beads. Again we failed to see any reverse translocations suggesting gravitational effects did not impact upon the measurement. The same trend was observed and is displayed in supplementary figure 4, confirming that the lack of reversed peaks is not an artefact of using a different bead size. As PDGF-BB has an isoelectric point of 9.8 it is assumed that the protein would have a positive surface charge in the chosen electrolyte. In order to confirm that the two bead populations could be analysed together under the same conditions, experiments were then performed under a fixed negative bias of -0.4 V. Supplementary figure 5 displays the results for these experiments in which a clear and proportional increase in rate is visible with increasing PDGF concentrations. As the behaviour of PDGF and VEGF bound beads was found to be similar, providing an increase in rate under a negative bias, it can be concluded that both proteins can be observed binding their aptamers under these conditions.

As each PDGF-BB molecule can bind two aptamers, aggregation is sometimes observed in their assays(Huang et al., 2005). In contrast, addition of VEGF does not cause aggregation as the protein only has one epitope for the aptamer(Hasegawa et al., 2008). In our set of experiments we also observe an increase in the number of aggregates as PDGF-BB concentration increases up to 10 nM, as displayed in supplementary figures 6 and 7. However, an increase in rate in relation to PDGF-BB concentration is still observable, thus the formation of aggregates did not affect the AptaTRPS technique.
Factors which influence the sensitivity of the assay

To determine if the bead size and concentration would have any impact on the sensitivity and ease of the assay we investigated their effects using the VEGF aptamers VEa5 and V7t1. Data summarising the effects of bead diameter and concentration are displayed in figure 3A and 3B. To investigate the effect of bead concentration VEa5 modified beads at 3, 2.5 and 2 × 10⁹ beads/mL were added to solution containing the VEGF protein (figure 3A/B curves 2, 3 and 4). As the bead concentration decreases, so does the concentration of VEGF at which the bead rate remains constant 100 nM (line 2) to 25 nM (line 4), this has an impact upon the dynamic range of the assay and is likely due to a decrease in the number of available binding sites in solution, leading to the saturation of each individual bead.
with VEGF at much lower concentrations of protein. As indicated above the number of beads moving through the pore is proportional to their concentration, hence lower concentrations of beads will result in a lower overall rate, at bead concentrations below $2.5 \times 10^9$ beads/mL it could be suggested that the bead count is so low (<10 beads a minute) it is no longer practical for accurate measurement and lengthens the required assay run-time unnecessarily.

To investigate the impact of bead size, the bead concentration was kept the same ($3 \times 10^9$ beads/mL) and the diameter was increased to 300 nm (figure 3A and B line 1 and 2). The rate of 300 nm beads at 0 nM VEGF was initially higher than the 120 nm beads with average rates of 35 and 0 beads/min respectively. It is likely that this is due to both the increased mass of the beads, leading to a higher gravitational effect, and also due to a lower relative loading of DNA per area of beads suspended. Both beads have comparable binding capacities (see method section) therefore we conclude the 300 nm beads have a lower packing density of streptavidin, and therefore DNA, on their surface.

The differences in packing density might also be responsible for the differences in rate signal between 120 and 300 nm beads. It is observed that there is a sharp initial increase in rate upon addition of VEGF and the rate signal appears to saturate sooner for the 300 nm beads; this could indicate that the sensitivity could be tailored by altering the packing density of capture probe on the surface of beads. For the purposes of this assay, these results demonstrate that the 300 nm beads are able to provide a rate-related signal change in response to protein binding which can be analysed via TRPS, further supporting the notion of the multiplexed AptaTRPS assay. Figure 3C represents the difference in the rate signal for two anti-VEGF aptamers: VEa5 (line 2) and V7t1 (line 5). There are two observations from this data; firstly, the rate signal for the V7t1 aptamer increases at low concentrations of VEGF and appears to have levelled out by 10 nM, whereas the rate of VEa5 beads continues to increase up to 100 nM of added protein. Initially we explored the same VEGF concentrations for both aptamers, however due to the rate remaining constant for the V7t1 aptamer at concentrations of VEGF > 1nM we chose to significantly reduce the amount of VEGF added. To illustrate this effect, Figure 3D displays an expanded view of the data for the V7t1 aptamer which exemplifies the ability of this aptasensor to detect VEGF at concentrations down to 0.05 nM. This is due to differences in aptamer binding mechanism and affinity. VEa5 has a reported $K_D$ of 130
nM (Hasegawa et al., 2008) and is postulated to bind due to the presence of stem-loop structures (Kanakaraj et al., 2013). V7t1 has a reported $K_D$ of 1.4 nM for VEGF$_{165}$ and due to its lack of selectivity over VEGF$_{121}$ is postulated to bind the receptor binding domain present on both VEGF analogues by forming a G-quadruplex (Nonaka et al., 2010); the superior affinity of V7t1 allows for the measurement of smaller concentrations of VEGF$_{165}$ on this platform.

The second observation from figure 3C is the marked difference in maximum rate between VEA5 and V7t1 which reach circa 60 beads/minute and 120 beads/minute, respectively at higher VEGF concentrations. As the overall number of aptamer molecules and beads should be the same, the difference in rate must be due to the difference in length of DNA. As demonstrated in our previous work, additional to the concentration of DNA added to the beads surface, the length of DNA sequence used is also a key factor in determining the maximum bead rate through the pore (Billinge et al., 2014). The V7t1 aptamer is 20 nucleotides shorter than VEA5 and with a negative bias under the pore, and the concentration of the beads being identical in all experiments the rates vary ~40 beads/minute (V7t1) and ~1 bead/minute (VEA5) respectively, when VEGF concentration is 0 nM. This is attributed to the effects of the length of the DNA, with the short DNA sequence able to pass through the pore more easily. Upon the protein binding to the aptamer, we believe there are two possible reasons for the higher total rate for the shorter aptamer sequence. The first could be due to the protein shielding the phosphate backbone to a greater extent for the shorter DNA, and secondly the binding mechanism. The conformation of the DNA aptamer, the location it holds the protein with respect to the bead, could impact on the bead zeta potential and drag, however further work is needed to infer implications of any mechanistic differences.

To further support the theory that bead rate is dominated by the charge of VEGF, we then altered the pH of our buffer to 9.3. It was expected that at this pH the VEGF would have an overall negative surface charge, and with a negative voltage bias applied under the pore the previously observed increase in particle rate with increasing protein concentration would be eliminated. As shown in table 1 this was indeed found to be the case, with bead rate remaining at 0.5 beads/min. When the applied potential was revered to an equal positive potential, the rate decreases with increasing protein concentration, reverting back to the trend
observed previously with thrombin (Billinge et al., 2014), i.e. a diminution, suggesting a charge-shielding mechanism is taking place.

<table>
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<th>pH 9.3</th>
<th>pH 7.4</th>
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<td>0.5</td>
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<td>0.5</td>
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Table 1: comparative summary of the effect of buffer pH on recorded particle rate using the same pore and instrument conditions.
**Filtering the data sets from two different particles**

In order to successfully multiplex the rate signals for two different bead sizes, it first must be possible to differentiate the beads from one another. Figure 4A displays the calibrated diameter distributions for the two different bead sets measured separately and together (inset) which confirms that the resolution of TRPS is able to distinguish these two bead populations without overlap. The distribution of 120 nm bioadembeads streptavidin plus (Figure 4Ai) is much narrower than for the 300 nm bioadembeads streptavidin; this is believed to be due to a difference in the stringency of bead manufacture – it is possible to see the distribution of bead diameters with great accuracy due to the particle-by-particle analysis of TRPS which has been shown to have far greater resolving power over conventional sizing techniques, such as Dynamic Light Scattering, when analysing non-uniform dispersions (Henriquez et al., 2004; Roberts et al., 2012; Willmott et al., 2010).
Figure 4: A Distribution of particle diameters for (i) 120 nm bio-adembeads streptavidin plus (Ademtech, France) and (ii) 300 nm bio-adembeads streptavidin (Ademtech, France). B 1 Combined rate of equal concentrations of 120 and 300 nm beads passing through the pore, 2 rate of 120 nm beads taken from 1, 3 rate of 300 nm beads, taken from 1.

**Multiplexing PDGF-BB and VEGF measurement**

Figure 4B line 1 is an example of the raw data plot showing particle count versus time for a sample containing both 120 and 300 nm beads. Following the experiment we can reanalyse the data and divide the total bead count into the two different beads sizes, lines 2 and 3. The 120 nm beads (line 2) are seen to run through the pore at a greater frequency than the 300 nm beads, despite being at the same concentration; this has been attributed to the effect of the ratio of the diameter of the pore constriction vs the diameter of the beads. As the pore size is reduced the particle rate decreases(Kozak et al., 2012) and this will be seen to a greater extent for the larger 300nm beads. We were unable to increase the size of the pore any
further as the signal from the smaller beads would have been lost in the baseline noise.

Figure 5: A bar graph of triplicate experiments comparing the average particle rates of 120 nm beads functionalised with V7t1 as VEGF \textsubscript{165} concentration increases with and without the addition of 10 nM PDGF-BB in the presence of 300 nm beads functionalised with anti-PDGF aptamer. B bar graph of triplicate experiments comparing the average particle rates of 300 nm beads functionalised with anti-PDGF aptamer as PDGF-BB concentration increases with and without the addition of 10 nM VEGF\textsubscript{165} in the presence of 120 nm beads functionalised with V7t1. C 3D surface plot displaying the effect of PDGF and VEGF on particle rate of 120 nm beads. D 3D surface plot displaying the effect of PDGF and VEGF on particle rate of 300 nm beads. Error bars represent 1 standard deviation from the mean.

In order to demonstrate its ability to multiplex, a control experiment was devised to ensure that the presence of additional proteins and beads do not interfere with the signal. Figure 5A shows the rate of 120 nm beads in the presence of different concentrations of VEGF with and without the secondary analyte - 10 nM PDGF-BB. Figure 5B displays the rate for 300 nm beads in the presence of different
concentrations of PEGD with and without the secondary analyte - 10 nM VEGF\textsubscript{165}. These experiments illustrate that the signal from one particle does not change or deviate in the presence of additional beads and target proteins. There does seem to be a small effect of the VEGF protein on the PDGF assay, figure 5b, where at 0nM PDGF and 10nM VEGF there is an increase in particle rate, and a decrease in sensitivity to PDGF at lower concentrations recorded as a small change in particle rate for PDGF concentrations 2-7 nM. This might be attributed to nonspecific interactions between the PDGF aptamer and the VEGF target, and that these nonspecific or weaker interactions are overcome at higher concentrations of PDGF.

This is further illustrated in supplementary figure 8 which displays the rate of 120 nm beads functionalised with Vt71 aptamer against concentration of VEGF\textsubscript{165} in the presence of an equal concentration of 300 nm anti-PDGF aptamer loaded beads with several concentrations of PDGF-BB ranging from 0 – 10 nM. An increase in rate is still visible for the 120 nm beads regardless of the addition of 300 nm beads and PDGF-BB concentrations. As illustrated in figure 3, figure 5A verifies that no significant difference between the values for 0.5 and 10 nM of VEGF, suggesting that at this point all of the available binding sites on the 120 nm bead are occupied. The frequency of both bead sets was measured in the presence of several combinations of concentrations of PDGF-BB and VEGF\textsubscript{165}. The data from all of these experiments are represented as surface plots in Figure 5C and D. For both bead sets (C and D) rate is seen to increase when their specific protein target concentration is increased. This trends remains largely unaffected by the presence of the other bead population and protein and the two can be measured simultaneously.

These results suggest that the simultaneous measurement of two different growth factors by TRPS is possible by using differently sized beads to provide a differentiable signal which is able to be selectively filtered by blockade magnitude to pull out the desired individual rate data. The observed increases in particle rate are believed to be due to an increase in electrophoretic mobility toward the negative electrode at the base of the pore when the positive proteins bind selectively to their respective DNA aptamers via a conformational change. Multiplexing the TRPS method illustrates the potential for the platform to be used as a biosensor. As demonstrated in our previous work (Billinge et al., 2014) the
technique offers comparable sensitivity and assay times when compared to Surface Plasmon Resonance techniques or flow cytometry platforms. The TRPS technique does not have the throughput of a modern flow cytometer but does offer an improved signal and resolution sub 500nm particles, without the need of fluorescent labels.

4. Conclusions
VEGF$_{165}$ and PDGF-BB were successfully monitored binding to aptamer-laden beads of 120 and 300 nm in diameter. The signals generated were able to be monitored in situ and also in tandem with both proteins measured independently from one another using the same sample using the beads themselves as a label, without the need for any additional identifiers such as fluorophores. Parameters regarding bead concentration and size were considered and a multiplexed assay undertaken.

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