Tunable resistive pulse sensing as a tool to monitor analyte induced particle aggregation

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Abstract
Analytical technologies based upon superparamagnetic beads, SPBs, offer a rapid, simple and inexpensive way of separating and purifying the target analyte prior to detection. The SPBs can perform the capture, purification and the signal transduction stages, producing a simple, fast, and sensitive label-free format. Particle aggregation in the presence of the analyte is a common example of such a detection strategy. Herein we demonstrate the key parameters which lead to aggregation. We utilize a tunable resistive pulse sensing, TRPS, technology to follow the aggregation using three different methods of analysis. TRPS allows a comparison in the data treatment using average population values, particle concentration, and a more detailed analysis monitoring the change in aggregate size and frequency. To validate the approach, we use the well-known biotin-avidin binding assay to demonstrate the advantages and limitations of each type of analysis. Also presented are the key parameters that contribute to assay sensitivity such as bead concentration, size, binding capacities and data analysis.

Keywords: Superparamagnetic beads, resistive pulse sensing, aggregation, binding capacity

Introduction
The emergence of nano-materials and technology has catalyzed the search for new, low cost and robust detection and quantification methods. This research has been complemented and driven with the development, miniaturization and micro-fabrication of technologies available for point-of-care, POC, diagnostics and prognosis [1-5]. Detection strategies using nanomaterials are numerous and listed elsewhere in reviews [6] but can be generalized into spectroscopy [7-9], electrochemical [10,11] or magnetic [12].

Prior to analysis samples may require a pre-treatment stage involving purification or filtration to help ensure their compatibility with the technology platform. Superparamagnetic beads, SPBs, provide an efficient and cost effective way to separate and pre-concentrate analytes from solution [13-16] helping to simplify and facilitate the front-end of an assay. They are readily available through numerous commercial sources and synthesis strategies [17], offering a range of surface chemistries that can be quickly conjugated to any capture probe of interest [13,18,19]. SPBs have demonstrated their versatility for use in conjunction with techniques such as the polymerase chain reaction [20] and mass spectroscopy [21], and have been utilized in high-throughput linear [13,22-25] and nonlinear magnetophoresis assays [14].

If the target analyte contains multiple epitopes, then it is possible for the target to bind to multiple beads simultaneously, leading to aggregation [13,26-29]. In such agglutination assays, the concentration of the analyte is inferred from the changes in aggregate size and/or frequency [30]. Using aggregation as an analytical signal allows for a simple “label-free” detection strategy and has been employed in conjunction with Au particles [31,32], fluorescence, or fluorescence resonance energy transfer [33] often in conjunction with optical, cytometry [34], magneto resistive sensing [35], and dynamic light scattering detection platforms [27,36]. Aggregation of SPBs does not typically result in a change in optical signal due to their optically dense iron oxide cores. Alternative detection strategies for monitoring aggregation using SPBs have relied upon light scattering techniques to monitor the size of the aggregates using chain length [36] or turbidity [27,37] and changes in magnetic properties as particles are forced to be in close proximity to each other [28]. These technologies have been demonstrated with detection levels down to picomolar levels, with assay times as low as 5 minutes, however light scattering and optical techniques can have a limited dynamic range and could prove difficult to multiplex [38].

The characterization of colloid and nanoparticle based systems has been aided in recent years by the resurgence and development of Coulter counting techniques such as micro- to nanopore based technologies [39-47]. Known collectively as resistive pulse sensing, RPS, they offer an attractive technology format because the measurements provide information on individual particles within their natural environment. RPS has been used to study numerous types of particles in different contexts with a very high resolution, including single molecules biophysics, protein sensing, biological detection [41,48,49] and synthetic nanoparticle characterization [47,50].

An elegant and novel adaptation to RPS using a tunable elastomeric pore technology, TRPS [46,49], allows for further versatility as the pore can be stretched in real time to suit the sample [50]. Techniques have been developed using TRPS to accurately determine the concentration [47], size and surface
charge [51] of dispersed inorganic particles, TRPS is capable of detecting biological particles [44,49,52,53] as well as discriminating between particles of different surface chemistry and derivatives [29,54-56], and is typically used to measure particles with one dimension larger than 100 nm, making them suitable for a wide range of applications [46]. Recently we utilized a TRPS technology to monitor the aggregation of a special class of nanomaterial termed nanorods in the presence of analytes [29], other groups have monitored the aggregation of gold nanoparticles in the presence or absence of DNA to detect single nucleotide polymorphisms [57].

Whilst techniques have utilized the TRPS technology to monitor aggregation, the signal can often be complicated by particles traversing the pore in close proximity [56,58], and require beads with a uniform dimater. Here we present a comparison of the different strategies offered to interpret the aggregation signal using TRPS and have identified key parameters that control the aggregation of SPBs for the streptavidin-biotin assay, i.e., bead concentration, bead size, binding capacity, reaction time, and application of magnetic force. The avidin-biotin interaction was used as a model as it is among the strongest known non-covalent specific molecular interactions, i.e., $K_d \approx 10^{-15}$ M. Under the optimal assay conditions the assay could be used to detect aggregation in the presence of a biotinylated protein across five orders of magnitude. We also present results that correlate the concentration of analyte and the most stable and reproducible sizes of aggregates. We note that increases in the number of dimers for larger 3 micrometer beads best correlate with the increase in analyte concentration, whereas the smaller 300 nm particles form larger aggregates 8, 9 and 10mers in the presence of the analyte. This insight will help users optimize the TRPS setup for monitoring the aggregation signal, and offers an insight into the aggregation state of particles in the presence of the analyte.

**Experimental Materials**

The particles used in this study were, Dynal particles carboxyl (1 micrometer - 650.12 and 3 micrometer -143.05D, Invitrogen UK), and Bio-Adembeads, 300 nm (2003233, Ademtech, Pessac, France). For the 3 micrometer beads 1 mg = 5 x $10^7$ beads, for the 1 micrometer beads 1 mg = 6 x $10^8$ beads, and the 300 nm beads 1 mg = 1 x $10^{10}$ beads. Bovine serum albumin, BSA, 2-(N-Morpholino) ethanesulfonic acid (MES), Tween-20, and Sodium Dodecyl Sulphate (SDS) were all purchased from Sigma Aldrich (Gillingham, UK). 1 step ABTS solution, biotin-HRP, Biotin-BSA (BBSA), 1-Ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC), Sulfo-n-hydroxysuccinimide, N-hydroxysulfosuccinimide (sulfo-NHS) and avidin, were purchased from Thermo Scientific (Loughborough, UK). All chemicals and proteins were used as purchased without further purification. All particles were first collected with a magnet before being washed three times in the reaction buffer prior to use.

**Buffers used**

PBS (0.01 M phosphate buffer, 0.0027 M Potassium Chloride and 0.137 M Sodium Chloride, pH 7.4), PBST (PBS with 0.05 (v/v)% Tween). MES buffer (50 mM MES, pH 6.0). All separations were performed using a magnetic separator, “magrack 6” purchased from GE healthcare (Hatfield, UK).

**Modification of the beads with avidin**

Both the 1 and 3 micrometer diameter beads were modified with the avidin protein prior to the assay being performed. The beads (1 mg/ml) were washed three times with MES buffer, after the third wash step EDC (1 mg) and NHS (1 mg) were added to the solution in 1 ml of MES buffer, the particles were sonicated gently for 30 s and left on a rotating wheel allowing end-over-end mixing for 30 mins. This resulted in NHS activated beads. After 30 mins the particles were washed with MES buffer twice before a 1 ml PBS solution containing the required amount of avidin protein was added to the particles. The particles were sonicated gently for 30 s and left on a rotating wheel allowing end-over-end mixing for 3 hours. After this time the beads were washed with PBST three times before being resuspended in PBST containing 0.05% (wt/wt) BSA protein to block any nonspecific adsorption.

**Protein coverage of beads**

The binding capacities of all beads were measured using the standard biotin-HRP and ABTS quantification, the detailed procedure for which can be found elsewhere [59]. Briefly, 50 μl of beads (at 1 mg/ml) where washed with PBST once and incubated with a fresh PBST (0.05% wt/wt BSA) solution for 30 mins to block the surface and eliminate any non-specific adsorption. After 30 mins the beads were washed twice with PBST and incubated with 0.05 μg of Biotin-HRP protein in PBST for 30 mins. After 30 mins the beads were washed three times with PBST and resuspended in 100 μl of PBST. To this solution 200 μl of ABTS solution was added and the vial was placed onto a rotating wheel for 1 min. After 1 min 200 μl of 1% (wt/wt%) SDS “STOP” solution was added to the beads and the absorbance at 410 nm was recorded. This final absorbance value was converted into a binding capacity using a calibration curve (standards) of known biotin-HRP protein.

**Assay format**

The SPBs (1 mg/ml) were routinely stored in PBST buffer containing 0.05 (wt/wt%) BSA. This storage served to act as a blocking solution reducing non-specific adsorption. When required the SPBs were diluted to the required concentration, and sonicated briefly (<10 s). Typical assay volumes were 500 – 1000 μl. The final dilution of the beads and its resultant concentration is reported in all graphs and calculations. Samples were prepared by diluting the target BBSA into PBST. The solution was incubated at room temperature on
a rotating wheel allowing end-over-end mixing. After the required hybridization time, the sample was either analyzed immediately, or subject to the magnetic assisted aggregation. Each experiment was run multiple times and the standard deviation of each point is displayed in the errors bars on the graphs.

**Magnetic assisted aggregation, MAS**

For the assisted aggregation - the sample was placed next to a permanent magnet, until all the beads were judged to have come out of solution after ∼1 min. The magnet had a field strength of 1 mT present up to 6 mm from the neodymium magnet bar on its surface. The initial solution was removed leaving all the beads on the side of the container and 500 μl PBS was added to the beads. The solution was stirred and the magnet was replaced separating the beads for a second time. When required the magnet was replaced for a third and fourth time. During the last separation, the magnet was removed allowing the pellet to settle to the bottom of the vial for 30 s before finally being vortexed ensuring the beads were evenly dispersed in the solution for analysis. This process aims at increasing the number of particle-particle interactions and thus increase the number of particle aggregates.

**Effects of adding additional particles to the reaction after the MAS**

During the magnetic assisted aggregation process described above, after the first MAS the analyte solution was removed and an additional 500 μl solution of beads in PBST was added to the sample vial. The aliquot contained the same concentration of beads as the original solution, thus doubling the concentration of beads. They were then treated with the same method described above.

**TRPS setup**

Measurements were made using the qNano system obtained from Izon Science (Christchurch), which incorporates the fluid cell, stretching apparatus, data recording and analysis software (v.2.2) and the pore specimens themselves [46,47]. The pores used were designated 'NP4000' for use with the 3 micrometer beads and most suitable, as determined by the manufacturer, for detecting particles in the range 2000 – 8000 nm, 'NP1000' for use with the 1 micrometer beads and most suitable for detecting particles in the range 500 – 2000 nm and 'NP400' for use with the 300 nm beads and most suitable for detecting particles in the range 200 – 800 nm. The tunable pores can be reused and in each set of experiments only one pore for each size of particle was needed. A typical setup for each membrane was: NP4000, Stretch 3.5 mm, Voltage 0.04V; NP1000, Stretch 5 mm, Voltage 0.04V; NP400, Stretch 5 mm, Voltage 0.32V [29]. Once a baseline current of approx. 100 nA ± 10 was established for each membrane the potential and stretch were kept constant during the experiment i.e., it never changed during the data capture or changed between each sample. Once the instrument was setup under the above conditions, it was stable over a period of hours. After each sample was analysed only the upper fluid cell needed to be washed with PBST to ensure no residual particles remained and the membrane stability meant that it did not need to be relaxed after each measurement. To further verify the upper cell was clean of residual aggregates, a PBST solution was placed into the cell and the signal recorded for 2 mins to ensure a stable baseline with no pulses observed. Izons calibration beads (obtained with the instrument) were used to calculate the initial concentration and size of all the particles, particles 955 nm in diameter, and 350 nm in diameter were used for this purpose.

**Data analysis**

Each particle that traverses the pore is recorded as a pulse event, termed ∆I_p, the magnitude of the pulse or peak magnitude, can be related to the particle size, and their frequency used to determine the concentrations. Data capture was performed using Izon's control suite 2.2 software. The number of data points captured for each experiment exceeded 500. In all measurements a noise threshold of 0.1 nA was used and all recorded events were clearly distinguishable from this noise level in the baseline current.

During the first step of the assay, avidin functionalized SPBs were added to a solution containing biotinylated-BSA, each BSA protein contains an average of 9 biotin molecules per BSA protein (data from supplier). The solution was incubated on a rotating wheel for 25 minutes and in the second step, an external magnetic gradient was applied to the bead suspensions to separate the beads and accelerate the rate of bead interaction with each other, in all experiments unless stated the number of MAS stages was always two. In the final step a 40 microlitres aliquot of the sample was placed into the TRPS technology for analysis. During the incubation period the instrument was setup as described above, a stable baseline current was established prior to running the samples. When required calibration beads were passed through the pore at the start and at the end of each experiment to allow the concentration and size of the beads to be determined.

**Method 1: Average size**

Upon processing the raw data with the software a "txt" file was created which contained information on each pulse recorded during the experiment. The peak height and time was extracted from the data and copied into a spreadsheet. Further analysis of the data was carried out using Microsoft Excel 2010. The average peak height was calculated from all the data points captured and plotted against concentration.

**Method 2: Determining the change in bead concentration**

The initial concentration of beads was calculated using the calibration beads. During each experiment a "blank" sample, a solution containing no analyte, was then used as a calibration
file for the remaining experiments. To monitor the changes in bead concentration as a function of the analyte concentration, each subsequent sample was then compared against the concentration of beads in the “blank” file.

**Method 3: Monitoring the change in aggregate size**

The data was extracted as a .txt file as described above. The blank sample described above was used to determine the location and size of the monomer and larger aggregate peaks. As the blank sample contained no analyte and therefore the observed signal represented monomers, a bin size which accounted for these data points was determined. The bin size ranged from the lowest peak current, lpc, upto 94% of the max peak current, mpc, observed for the monomers. A value of 94% was chosen to omit the slight tail in the data set seen in Figure 1B. As the peak height is proportional to the volume displaced by the traversing particle \([44,47]\) a dimer would produce twice the signal and therefore any signal that was observed within \(0.94\text{mpc} + (0.94\text{mpc} - \text{lpc})\) was counted as a dimer, thus every subsequent window was simply an increase of \((0.94\text{mpc} - \text{lpc})\). A histogram could then be constructed using multiples of the monomer peak to determine the number of larger aggregates. Once the number of data points for each bin had been calculated from the raw data, the number was

![Figure 1](http://www.hoajonline.com/journals/pdf/2053-0927-1-1.pdf)

**Figure 1.** Assay using 3 μm diameter beads, binding capacity of 14 μg/mg, assay time of 25 mins, 5 x 10^8 beads/ml.

(A) Change in average pulse height, \(\Delta ip = \text{blue}\), and bead concentration = green, against concentration of biotinylated analyte, N.B. error bars are plotted but smaller than the symbol on the graph.

(B) Example of raw data from an individual experiment, red histogram displays the pulse height (\(\Delta ip\)) for a blank sample, the black distribution shows the change in \(\Delta ip\) for an analyte concentration of 1 x 10^{-9} \text{ mol dm}^{-3}. Overlaid is a schematic (N.B not to exact scale or position) gate size for the monomer, dimer.

(C) Plot of percentage of signal of monomer = red, dimer = green, trimer = blue and tetramer = purple.
converted into a percentage by dividing it by the total number of data points for the experiment. The ratio was then plotted as a function of analyte concentration. In each experiment the fraction of aggregates up to and including the 12mers was calculated, even if they are not plotted in the figures. All experiments were repeated at least once and in some cases triplicate, where error bars are shown they represent the standard deviation around the mean value taken from at least triplicate repeat experiments.

Results and discussion
Comparing methods for analyzing the aggregation of beads
Method 1: The simplest way to analyze the data, is to plot the mean peak size, $\Delta i_p$, signal versus analyte concentration, as shown in Figure 1A (blue diamonds). As the concentration of the analyte is increased the average $\Delta i_p$ remains steady until a sharp increase around $1 \times 10^{-9}$ mol dm$^{-3}$ of analyte. The mean pulse height continues to increase up to an analyte concentration of $1 \times 10^{-8}$ mol dm$^{-3}$ before decreasing back to its original value. The increase in $\Delta i_p$ can be attributed to the particles forming larger aggregates in the presence of the analyte, at lower concentrations of analyte the number of aggregates is too low to be detected as the majority of the signal comes from the individual monomers. The decrease in signal from aggregation at high concentrations of analytes has been well documented for agglutination assays. It results from the saturation of the binding sites on the particle surface with the analyte and is known as the ‘hook’ effect. Plotting the mean $\Delta i_p$ as a function of concentration seems insensitive to analyte concentrations below $1 \times 10^{-9}$ mol dm$^{-3}$. Method 2: An alternative method is to follow the change in particle concentration. The TRPS method can be calibrated to calculate the concentration of the particles, as the flux, $J$, of particles through the pores is directly proportional to their concentration [44]. As the particles aggregate the total number of beads in solution decreases, as monomers are consumed into growing particle clusters. The TRPS technology was calibrated to measure the change in particle concentration. The same data used for the above $\Delta i_p$ measurements was re-analyzed, the results are plotted in Figure 1A (green triangles). Measuring the change in bead concentration seems to be more sensitive, and a decrease in bead count can be observed at concentrations as low as $1 \times 10^{-12}$ mol dm$^{-3}$. However with this method the measured bead concentrations varied in repeat runs by up to 45%, shown in Supplement figure S1 As a comparison the triplicate measurements for methods 1 are shown in Supplement figure S2. If insufficient time is given for the beads to come out of solution in the presence of the magnet, or beads are washed out of the vial during the MAS stage the variation from run to run may be effected. Therefore, if the technique is to be used across multiple labs the variation from the technician’s patience in waiting for all the beads to be extracted from the solution could lead to variations in the results. Given that we hope to develop this technique for the detection of analytes, the user reproducibility must be high, and a more robust method of data analysis is needed.

In this study commercial Dynal and Ademtech beads with relatively uniform diameters, <2% and <10% CV, respectively, were used. When the SPBs are highly uniform in diameter, it is possible to isolate the signal from the larger aggregates from that of the individual beads. Method 3: Figure 1B shows the raw data from the Izon software for the “Blank/Background” (red) and a “Sample” exposed to the analyte (black). In the blank sample the beads produce an individual peak which we term “monomers”. The gate was titled “monomer”, as the $\Delta i_p$ is proportional to volume of the particle traversing the pore, a dimer should have twice the signal of a monomer. Therefore, by doubling the gate size the number of dimers could be counted. A similar process can be done to account for the trimers, tetramers etc. Figure 1C summarizes the quantitative analysis of the percentage of monomers, dimers, trimers and tetramers formed in the assay using the 3 micrometer beads, as a function of analyte concentration. At concentrations between $10^{-12}$ and $10^{-9}$ M there was a direct correlation between the number of dimer aggregates and the analyte concentration, $R^2 = 0.936$. Again at concentrations greater than 1 nM there was a decrease in number of all sizes of aggregates and increase in monomers was observed due to the onset of the hook effect. The variation in results, from assay to assay was lower using this analysis, the average coefficient of variation for data points across the entire concentration was no more than 25%, whereas for methods one and two the CV range which from 35-65%. The reasons being 1 – that the average population value is no longer being used, and therefore the small number of larger aggregates is not lost on the analysis, and 2 – we use the fraction of the beads that are remaining, if any particles are lost during the wash and MAS stages it does not impact upon the ratio between monomers and larger aggregates. The assumption would be that we lose an equal proportion of monomers and dimers during any washing steps which may not hold true as larger aggregates tend to separate out from solution at faster rates. However the reduction in errors bars and the mirroring of the dimer and monomer peak suggest that we do not have a strong sample bias. We therefore continued to use this method to monitor particle aggregation.

Influence of magnetic assisted aggregation, MAS, on the hook effect
Figure 2A presents the number of monomers as a function of analyte concentration and number of MAS performed. The influence of magnetic extraction on the assay itself was determined by analyzing the number of monomers with and without the MAS. Several observations can be made from these results, firstly in the absence of the MAS (green triangles) fewer aggregates appear to form, and the number of monomers is high across the range of analyte concentrations. In the absence
of the MAS bringing the beads together, aggregates are only formed if two beads collide in solution with the correct orientation and alignment to form a bond. Upon the addition of the magnetic field the number of interactions increases, leading to an increase in aggregation and the decrease in the number of monomers. Increasing the number of MAS stages from 2 to 4 does not seem to drastically decrease the number of monomers across the concentration range and allows the user to pick the most convenient number for their assay conditions and time allowances.

The hook effect could be problematic during an analytical measurement as two concentrations of analyte in solution could in theory produce the same aggregation ratio. To explore the possibility of removing the hook effect, we added a second aliquot of beads to the solution after the first MAS stage. The MAS stage was repeated two more times and the sample analyzed. Figure 2B displays the results and the fraction of the sample that was measured to be mono, di, tri and tetramers. It is evident from Figure 2B that the characteristic decrease in aggregate signal and increase in monomer signal typically observed at high concentrations of analyte in previous experiments has been shifted beyond the concentration range being measured here, and a gradual decrease in monomer signal is observed across the entire range of the assay. The explanation for this is that at high concentrations of analyte the majority of the binding sites on the beads are occupied. Therefore when brought into contact with each other no free binding sites are available to form aggregates. The addition of fresh beads after the first MAS stage introduces vacant binding sites leading to aggregation.

**Influence of binding capacity, concentration, and diameter**

To further investigate the onset of aggregation and dynamic range of the assay, a set of experiments was devised in which the binding capacity was systematically varied. Figure 3A and 3B present the percentage of monomers and dimers, respectively, produced by the reaction of 3 micrometer SPBs across a range of analyte concentrations. In each experiment the binding capacity of the beads was gradually decreased from 14 μg/mg to 3 μg/mg. As the binding capacity was lowered, the magnitude of the change in the numbers of monomers and dimers, decreases. Hence as the binding capacity is reduced the beads are less efficient at forming aggregates and therefore the signal is diminished.

Figure 4A illustrates the effect of lowering the number of beads in the assay, and thus reducing the total number of binding sites. This has the effect of shifting the onset of the hook effect to a lower concentration. Whilst the hook effect appears at a lower analyte concentration it is also worth noting that a greater decrease in monomer signal is also observed changing from 70% for the 2.5 × 10^8 beads/ml down to 40% of the signal for the 1 × 10^7 beads/ml, effectively making the onset of aggregation easier to observe. The onset of the hook effect was found to be closely linked to both the concentration of avidin in the reaction and the density of avidin on the beads. To test the effect of particle size a similar set of experiments were then performed with biotinylated-BSA
for avidin functionalized 1 micrometer beads and the results are shown in Figure 4B. Qualitatively similar trends were observed, i.e., both the intensity and position of the maximum number of aggregates can be tailored by changing the bead number and binding capacity. Quantitatively, however, the results for the 1 micrometer beads differed from the 3 micrometer beads. It is clear from Figure 4B that when a similar concentration of beads are used (green and blue)
the peak in aggregation appeared at a lower concentration of analyte for the 1 micrometer beads, and a larger fraction of beads formed aggregates. The interpretation of this result is that when the same number of 1 and 3 micrometer beads are used the 1 micrometer beads aggregate more efficiently. This is most likely due to the fact that the 1 micrometer beads have a higher rotational diffusion coefficient. When the total binding capacity of the 1 micrometer beads was increased to 20 μg/mg, more 1 micrometer beads were added to solution (blue and red) the onset of the hook effect again is shifted to a higher concentration of analyte, reflecting the change in total binding capacity of the beads surface.

The change in behavior from lowering the binding capacity of the beads – Figure 3, and changing the total number of

![Figure 5](http://www.hoajonline.com/journals/pdf/2053-0927-1-1.pdf)

**Figure 5.** (A) Distribution of monomer = red, dimer = green, trimer = blue and tetramer = purple. 1 μm avidin modified beads 25 min assay time 6 x 10^9 beads /ml, binding capacity 20 μg/mg. (B) Distribution of monomer = red, dimer = green, and tetramer = purple using 0.3 μm Streptavidin modified beads 25 min assay time 1 x 10^9 beads /ml, binding capacity 4 μg/ mg. (C) Distribution of 5mer = blue, 6mer = purple, and 7mer = green and 8mer = red using 0.3 μm Streptavidin modified beads 25 min assay time 1 x 10^10 beads /ml binding capacity 4 μg/mg. (D) Correlation, R^2 value, plotted along the axis between the increase in aggregate size versus the decrease in monomer signal for the three different bead sizes.
As we change the number of beads in solution, we can shift the onset of the hook effect by simply changing the total concentration of the target analyte. A comparable set, we cannot say with absolute certainty that the different sizes of aggregates for the 1 micrometer beads when compared to the 3 micrometer beads where typically larger, i.e., larger numbers of gates were required to count all the data points. The smaller particles seem to form larger and more readily measurable aggregates, and one possible explanation is that the smaller beads have an ability to nucleate and grow in aggregate size, mimicking crystallization of nanoparticles. Micron sized particles have previously been used as model systems to aid with nucleation and growth studies. Similar observations of cluster size might have proven difficult using flow cytometry or light scattering techniques and the TRPS may offer the possibility of studying the aggregation and growth of colloidal particles in solution.

### Conclusions

In the current study, we have used TRPS technology to characterize the aggregation state of avidin functionalized particles in the presence of a biotinylated target. We have attempted to present several methods of data interpretation, aimed at increasing the reproducibility of the experiment and allowing for a detailed and informative study of the aggregation state. We have then used the TRPS technology to investigate the key experimental variables in designing a magnetic aggregation assay. A summary of all the factors that were found to influence the onset of aggregation can be found in Figure 6. The resulting assay platform was capable of operating over several orders of magnitude of concentration; the ability to measure aggregation can be improved by increasing the density of receptors on the beads, decreasing the size of the beads used, and/or decreasing the total number of beads used in the assay. However, lowering binding capacity and bead number resulted in a more rapid saturation of the beads with analyte and thus the hook effect appeared at lower analyte concentrations. By controlling the particle size, binding capacity, and particle concentration, the aggregation of SPBs in the presence of the target analyte can be tailored and predicted to produce a simple and sensitive analytical method.

### Additional files

- Supplement figure S1
- Supplement figure S2

### Competing interests

The authors declare that they have no competing interests.
Authors’ contributions

Emily Billinge performed experiments, prepared the manuscript and provided scientific input. Julien Muzard provided scientific advice, supporting in designing the experiments and writing the manuscript. Mark Platt performed the experiments, designed the experiments wrote the document, performed the preliminary data analysis and managed the project.

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