PAPER



Cite this: Analyst, 2022, 147, 5518

Received 27th July 2022, Accepted 24th October 2022 DOI: 10.1039/d2an01238c

rsc.li/analyst

Introduction

Gestational diabetes monitoring

Gestational diabetes is the third most prevalent type of diabetes mellitus after Type I and Type II diabetes.¹ As the name suggests, it affects women during pregnancy. GDM most commonly develops between the 24th to 28th week of pregnancy^{1,2} and has a range of short and long-term adverse effects on both the pregnant woman and foetus. The short-term effects to the patient include pre-eclampsia, shoulder dystocia and increased risk of C-section.³ Short-term effects on the infant include neonatal hypoglycemia and neonatal jaundice. Further, type II diabetes may develop in both mother and infant in the long term.⁴ To

Monitoring gestational diabetes at the point-of-care via dual glycated albumin lateral flow assays in conjunction with a handheld reader[†]

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Chronic conditions like diabetes require monitoring of vital biomarkers over extended periods of time. Monitoring gestational diabetes mellitus (GDM) is crucial to avoid short- and long-term adverse effects on both mother and infant. Providing monitoring systems to patients at the point-of-care (POC) has the potential to help mitigate these effects. In this manuscript, we propose the use of a sensing system combining lateral flow assays (LFAs) with a handheld colorimetric reader for use in tracking the glycemic status of a GDM patient at the POC. Current strategies of GDM monitoring include glucose and HbA1c measurements. These are often too frequent or not frequent enough for effective monitoring. Hence, we have developed a sensor for an intermediate interval biomarker - glycated albumin (GA). Based on the half-life of the protein, GA is measured once every 2-3 weeks. Here we first present two lateral flow assays, one for GA and another for total serum albumin used in conjunction with a handheld reader to read the colorimetric signals. Both assays have a sandwich aptamer format and measure the target proteins in their physiologically relevant ranges. The GA assay has a dynamic range of 3-20 mg ml⁻¹ and the serum albumin assay has a range of 20-50 mg ml⁻¹ without any sample dilution. Both LFAs were then incorporated into a single dual assay cartridge such that both assays could run simultaneously and provide the % glycated albumin value from a single test. Thus, the dual assay cartridge plus reader system has the potential to provide an effective platform for measuring GA for tracking GDM at the POC.

avoid these effects, it is crucial to monitor GDM carefully during pregnancy up to a couple of months after the baby is born.

Currently, the standard strategy to monitor GDM is direct glucose measurement and/or HbA1c measurement. Due to the rate of glucose metabolism, it is typically recommended that patients with diabetes measure it 4-5 times a day. Since GDM is primarily characterized by consistent hyperglycemia⁵ with rare hypoglycemic episodes, multiple glucose measurements may not always be necessary. Since the recommendation is multiple readings per day, traditional glucose meters would require multiple fingerpricks every day, which can result in non-compliance from the patients' side. The other standardized biomarker for diabetes is glycated hemoglobin (HbA1c).⁶ Due to its longer half-life, HbA1c is measured only once in 2–3 months.⁷ In this case, the longer timeline is not ideal since the crucial monitoring period for GDM is about 6-7 months. Hence, specifically for GDM, an intermediate biomarker (interval: 2 weeks) would be advantageous for monitoring glycemic status. It would also be useful for monitoring the effects of therapeutic strategies like alterations in dietary regimens and exercise routines.8

One such intermediate interval biomarker is glycated albumin (GA).^{9,10} Free glucose in the bloodstream binds to

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[†]Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d2an01238c

serum albumin *via* non-enzymatic glycation reaction to form glycated albumin.¹¹ It reflects glycemic excursions of postprandial plasma glucose along with average plasma glucose.¹² Since glycated albumin is reported as a ratio of glycated to total albumin, unglycated protein also has to be measured in addition to the glycated version. In normal conditions, about 10 to 16% of serum albumin is glycated.¹⁰ However, in cases of prolonged hyperglycemia, up to 40% of albumin can be glycated.¹⁰ Since glycation of albumin as a phenomenon is not restricted to GDM, as a biomarker, it can be used to monitor type I and type II diabetes as well.¹³

Point-of-care monitoring with lateral flow assays

In the past two years, COVID-19 has demonstrated the importance of point-of-care (POC) tests. Testing at home is not just convenient for patients but also plays a major role in relieving the load on healthcare providers.^{14–16} Most commercially available at-home tests like Panbio rapid test (Abbott) or SD Biosensor rapid test (Roche) use the standard lateral flow assay (LFA) design.¹⁷ Lateral flow assays are one of the most established forms of paper based POC test platforms.^{18,19} Their compact size, portability, low-cost and ease-of-use make them ideal candidates for POC applications. To date, several LFAs have been developed for a range of applications like COVID-19, influenza, bacterial pathogens, heart failure *etc*.^{18,20–22}

The principal components of any LFA typically include recognition elements like antibodies^{23,24} for the capture of the target biomolecule onto the test strip and transduction elements like enzymes^{25,26} or nanoparticles^{21,23} to transduce the recognition event into a readable signal. Antibodies have been established as recognition elements in lateral flow assays owing to their high sensitivity and specificity.^{27,28} However, they also have drawbacks like high cost of production and thermal instability. On the other hand, aptamers, which are single-stranded DNA or RNA sequences, also have the potential to be used as recognition elements for LFAs due to their thermal stability, stability over a range of pH, low-cost and longer shelf life which make them more suited for use in point-of-care tests.^{29–31}

The other main component of any biosensor is the transduction element. For qualitative POC tests, the signal generated by the transduction element can be observed visually to get a yes/no answer.³² However, for quantitative tests, the transduced signal must be measured and quantified.^{24,33} Several optical techniques like colorimetry,³⁴ fluorescence³⁵ and surface-enhanced Raman spectroscopy³⁶ have been explored for lateral flow applications. The use of bare gold nanoparticles for colorimetric signal generation is a sensitive and cost-effective optical transduction technique since it does not require any dye and uses less sophisticated equipment for signal generation and measurement.^{37–40}

Ikeda *et al.*,⁴¹ Testa *et al.*,⁴² and Ko *et al.*⁴³ have published assays for glycated albumin detection using various enzymatic methodologies. Inoue *et al.*,⁴⁴ and Bohli *et al.*⁴⁵ have published enzyme and antibody-based sensors using electrochemical detection methods. Ki *et al.*^{46,47} have published colorimetric

antibody-based assays to measure glycated and serum albumin. Apiwat *et al.*⁴⁸ and Gosh *et al.*⁴⁹ have published fluorescence-based detection systems with aptamers as the recognition elements. While successfully detecting and measuring glycated albumin, all these systems have certain limitations in terms of POC applications. Enzymes and antibodies have low thermal stability and hence require low-temperature storage for effective use. Transduction methods like electrochemistry are not ideal for POC application since the associated equipment is likely not compact or portable. Lastly, most of the above-mentioned assays require an additional dilution step for sensing due to high concentrations of the target proteins. Currently the only commercially available assay for GA is the Lucica GA-L assay.⁵⁰ While this assay is a commercial standard, it is a lab-based enzymatic assay which uses clinical analysers.

In this manuscript, a sensing system has been developed for monitoring gestational diabetes at the point-of-care using glycated albumin as the target biomarker. The sensing system is comprised of a dual assay cartridge consisting of two aptamer-based sandwich assays on lateral flow platforms, for glycated and serum albumin (Fig. 1(a)). Initially both assays were developed separately, and their results have been reported. That was followed by incorporation of both assays into a single dual assay cartridge to run both assays simultaneously. The signal generated by the assays was read using a custom handheld colorimetric reader (Fig. 1(b)-(d)) following our earlier designs.^{51,52} The combination of lateral flow tests with a compact reader satisfies the RE-ASSURED criteria set by World Health Organization (WHO) for point-of-care testing.53 The combination of dual lateral flow assay cartridge and the handheld reader was designed specifically for simultaneous measurement of both forms of albumin (glycated and total) such that the % glycated albumin value can be obtained by the patient in 30 min at-home without diluting the samples.

Materials and methods

Materials

Gold(III) chloride trihydrate (HAuCl₄) and tri-sodium citrate dihydrate (Na₃C₅H₅O₇) for gold nanoparticle synthesis were purchased from MilliporeSigma (St. Louis, MO). Syringe filters (0.2 µm) were purchased from VWR (Pennsylvania, USA). All five aptamers were purchased from Integrated DNA Technologies (Iowa, USA). The oligonucleotide sequences are listed in the ESI (Table S1[†]). Tris (2-carboxyehtyl) phosphine hydrochloride (TCEP), sodium chloride (NaCl) and phosphatebuffered saline (PBS; pH 7.4) tablets were purchased from Sigma. Nanosep centrifugal filters were purchased from Pall Corporation (New York, USA). Serum albumin, glycated albumin and glucose were purchased from Sigma. Simulated blood serum (without albumin) was purchased from Biochemazone (Edmonton, Canada). Milli-O deionized (DI) water (18.2 M Ω cm⁻¹) was used in all experiments. Whatmann G4 chromatography paper and nitrocellulose membrane (FF170HP) were purchased from Cytiva (Marlborough, MA).

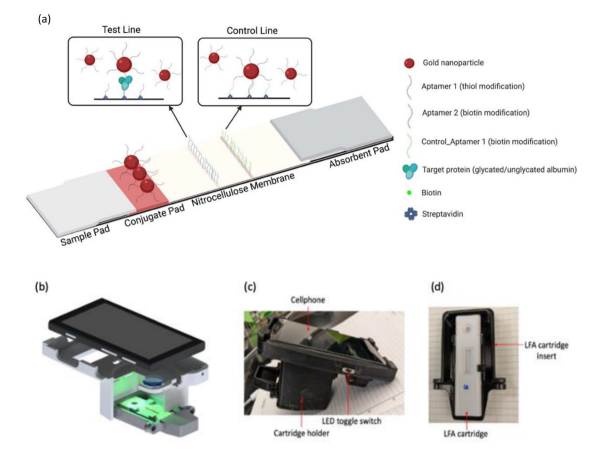


Fig. 1 (a) Lateral flow assay schematic (created with BioRender.com); (b) handheld colorimetric reader (CAD design); (c) handheld colorimetric reader and (d) lateral flow assay cartridge holder.

Glass fiber membrane was purchased from EMD Millipore (Burlington, MA). Thick absorbent paper (Trans-blot filter paper) was purchased from BioRad (Hercules, CA). Lateral flow cartridges and backing cards were purchased from DCN Dx (San Diego, CA). ESun pla+filament was purchased from Amazon. UV-Visible spectrometry was conducted using a Tecan Infinite 200 Pro (Männedorf, Switzerland) plate reader.

Zeta potential and dynamic light scattering measurements were conducted on a Zetasizer Nano ZS90 (Malvern, UK). Dual assay cartridges were printed on a Prusa i3 mk3s+printer. Colorimetric measurements were performed using a custom handheld reader developed by our team.

Methods

Synthesis and characterization of gold nanoparticles. Gold nanoparticles were synthesized using the protocol published by Bastus *et al.*⁵⁴ Briefly, 1 ml of 110 mM sodium citrate was added to 49 ml of boiling DI water. Two minutes after citrate addition, 335 μ l of 25 mM gold chloride was added. The solution was allowed to boil for 15–20 min for seed formation. Once the solution turned ruby red, the temperature was reduced and maintained at 90 °C for the remainder of the synthesis. After 1 hour, particle size was measured *via* absorbance measurements. If the size was insufficient, 335 μ l of 60 mM

sodium citrate was added, followed by 335 μ l of gold chloride. This step was repeated until the desired nanoparticle size was obtained. After the synthesis process, the nanoparticles were filtered using a 0.2 μ m syringe filter to remove aggregates. Filtered particles were stored at 4 °C until further use. The size and diameter of the particles were calculated from the absorbance spectrum. Hydrodynamic size was confirmed using dynamic light scattering. Zeta potential of the particles was also measured.

Conjugation of aptamers. Aptamers selective to glycated albumin and human serum albumin were conjugated to gold nanoparticles *via* thiol modifications on the aptamers (aptamer sequences listed in Table S1[†]). Selectivity of the aptamers was previously tested and confirmed by our group.⁵⁵ The aptamers purchased from Integrated DNA Technologies had a disulphide bond which was reduced before conjugation. This was achieved by adding 20 mM TCEP to the aptamers. After a one-hour incubation, the excess was removed by washing with a Nanosep centrifugal filter (10 kDa MWCO). The aptamers were finally resuspended to the original volume and folded by heating in a water bath at 85 °C. The concentrations of the aptamers were added to separate batches of gold nanoparticles in their respective ratios (3000:1 for the glycated albumin

assay and 2000:1 for the serum albumin assay). The aptamernanoparticle binding was carried out for one hour in a shaker and then left on the benchtop overnight. Salt aging was conducted after incubation using 2 M NaCl.⁵⁶ After salt addition was complete, particles were left on the benchtop overnight. The particles were then centrifuged at 10k rcf to remove the unbound aptamers and resuspended in 1× PBS. Particles were resuspended to a final OD 1 for the glycated albumin assay and OD 1.5 for the serum albumin assay. These particles were then stored at room temperature on the benchtop until further use.

Lateral flow assay (LFA) design. Lateral flow assay components used in this work followed the standard LFA design.^{27,30,57,58} Capture and control aptamers (sequences in Table S1[†]) were purchased with biotin modifications. The biotinylated aptamers were conjugated to streptavidin and immobilized onto the nitrocellulose membrane using a lateral flow dispenser. The striped aptamer was dried in a desiccator overnight. Glass fibre membrane was used for conjugate storage. It was pre-treated with 1× PBS + 0.7% casein + 7% sucrose + 0.05% Tween-20. Aptamer conjugated nanoparticles were loaded onto the glass fibre membrane and dried in a desiccator overnight. G4 chromatography paper, used as the sample pad, was pre-treated with $1 \times PBS + 5\%$ sucrose + 0.25% Tween-20. A thick trans blot absorbent pad was used to absorb excess sample, buffer, and nanoparticles. The chromatography paper, glass fibre membrane, nitrocellulose membrane and absorbent pad were attached to the backing card with a 2 mm overlap between consecutive layers (Fig. 1(a)). The backing card with the assembled lateral flow layers was placed into the cartridge, after which the test was ready to run. Cutting, pretreatment of membranes and assembly of the lateral flow strip were all a part of construction of the strips. All buffer formulations and membranes were identical for both glycated and serum albumin assays.

Handheld reader design. The reader was built based on the design of Mudanyali et al.51 A Nokia Lumia 1020 was used for the reader to capture images of the lateral flow test strips. A 20 mm diameter biconvex lens with a focal length of ~30 mm, was placed in front of the existing camera module of the smartphone, while the lateral flow test strips were placed ~35 mm away from the smartphone camera. Lateral flow test strips were illuminated using two LEDs as the light source (SunLED XSFWCB983W Broadcom and Limited ALMD-CM3F-Y1002), which were powered by two AAA batteries with a slide switch to turn it on/off. The mechanical body of the reader (Fig. 1(c)) was fabricated using the Stratasys Objet30 Pro 3D printer, which uses VeroBlackPlus photo resin as the building material to hold the smartphone, external lens, and its electrical components. A customized test-holder tray (Fig. 1(d)) was also designed specifically for the lateral flow test strip cartridge used in this work. This customized test tray was mounted with magnets to help secure the tray's position on the body of the reader.

Assay response and colorimetric measurement. Both lateral flow assays were run by adding $30 \ \mu$ l of the sample followed by

the addition of 50 μ l running buffer (1× PBS + 0.1% Tween-20) after 5 min. Thirty minutes after sample addition (optimization in next section), the signal was read using the handheld reader. The LFA cartridge was placed into its holder, which was then inserted into the reader. Once the holder was in place, the green LED (optimization in next section) was turned on and the image was captured. The images were then analysed using Image J software to get colorimetric intensities.

Determination of assay run time. The run time of the assay was the time interval measured starting from the addition of the sample to the development of a constant signal intensity at the test line. It determined by measuring the colorimetric signal intensity at different time points after sample addition: 5 min, 10 min, 20 min, 30 min, 40 min and 50 min.

Determination of colorimetric channel. The custom handheld reader was designed with two inbuilt LEDs: white and green. The ideal colorimetric channel was determined by capturing images with both white and green LED and analysing and comparing all colorimetric channels from both LEDs. The ideal channel was chosen to be the one that gave the highest signal sensitivity from lowest to highest target concentrations for both assays.

Assay response in serum. The response of both assays was tested in doped serum samples to evaluate their performance in a complex biological fluid. The simulated serum was doped with 8 and 20 mg ml⁻¹ glycated albumin and 35 and 50 mg ml⁻¹ serum albumin. 30 µl of the spiked samples was then added to the test strip followed by 50 µl of running buffer (1× PBS + 0.1% Tween-20). The handheld reader was then used to measure the signal at each test line. Each sample was tested in triplicate.

Dual assay cartridge design. The dual assay cartridge was designed in Solidworks and 3D printed in the lab. Two lateral flow assay strips: one for glycated albumin assay and the other one for serum albumin assay were placed on each side of the dual assay cartridge. Individual lateral flow assay strips were constructed with the same membranes and formulations as mentioned above. A G4 chromatography membrane pretreated with $1 \times PBS + 5\%$ sucrose + 0.25% Tween-20 was placed as the connector between both assays.

Dual assay response. The symmetry of flow in dual assay cartridge was first tested by placing glycated albumin assay strips in both channels of the dual cartridge. $50 \ \mu$ l of (20 mg ml⁻¹ glycated albumin) sample was added to the inlet followed by $50 \ \mu$ l of running buffer. The assay signals were read, and the intensities plotted and compared for both assays in the cartridge. Next, $50 \ \mu$ l of sample (S1/S2/S3) was added to the sample inlet to run both assays simultaneously. Sample 1 consisted of 3.5 mg ml⁻¹ GA and 25 mg ml⁻¹ HSA; sample 2 was comprised of 10 mg ml⁻¹ GA and 30 mg ml⁻¹ HSA; sample 3 was a mix of 20 mg ml⁻¹ GA and 30 mg ml⁻¹ HSA. All samples were made by doping simulated blood serum with respective combinations of glycated and serum albumin.

Five mins after sample addition, 50 μ l of running buffer (1× PBS + 0.1% Tween-20) was added to push all residual conjugates from the conjugate layers and flow through the nitrocel-

lulose membrane. Colorimetric intensities were measured 30 min after sample addition. Each sample was tested in triplicate.

Results and discussion

Nanoparticle characterization

Gold nanoparticles were synthesized by the seed growth method as described above. After synthesis was completed, the absorbance of the nanoparticles was measured using a plate reader. Using the absorbance spectrum, the size and concentration of the nanoparticles was calculated. The spectrum, as seen in Fig. 2(a), has an absorbance peak at 524 nm, which was used to calculate the size of the nanoparticles. These 40 nm nanoparticles were used in all experiments. After both aptamers were conjugated to separate sets of nanoparticle solutions, conjugation was confirmed using via absorbance measurements. An absorbance peak shift of 4 nm confirmed aptamer conjugation onto the nanoparticles for both assays. The peak shifts can be observed in Fig. 2(a). OD 1 particles were used for the glycated albumin assay, while OD 1.5 particles were used in the serum albumin assay owing to the higher concentration range. Nanoparticle size and conjugation

was further confirmed by measuring the zeta potential and hydrodynamic size. As seen in Fig. 2(b), a 10 nm increase in hydrodynamic size was observed after conjugation and a reduction of zeta potential from -31 mV to -42 mV was also seen (Fig. 2(c)).

Assay response

Determination of assay run time. The time interval for measurement of assay response was determined by measuring the signal intensities over a period of 50 min. As seen in Fig. 3(a), the intensity reached 90% of the maximum intensity in 20 min. Maximum signal intensity was reached at the 30 min time point, after which it stabilized. Hence, both assays were run for 30 min after which the final colorimetric intensities were measured. This was a marked improvement over the dipstick assays published by our group previously, in which signal development took between 60–75 min. The assay run time was measured for both glycated and serum albumin assays and 30 min was found to be the ideal run time for both assays.

Determination of colorimetric channel. The handheld reader used to measure colorimetric intensities was designed with two LEDs: white and green. Both assay responses were

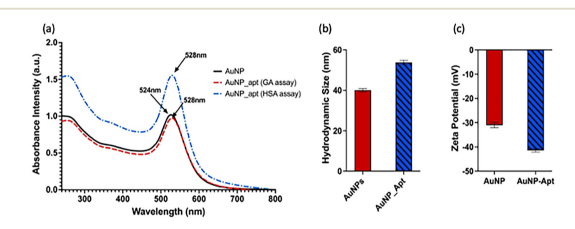


Fig. 2 (a) Absorbance spectrum of citrate capped AuNPs and aptamer conjugated AuNPs (GA assay and HSA assay); (b) hydrodynamic sizes of nanoparticles before and after conjugation; (c) zeta potential of particles before and after aptamer conjugation.

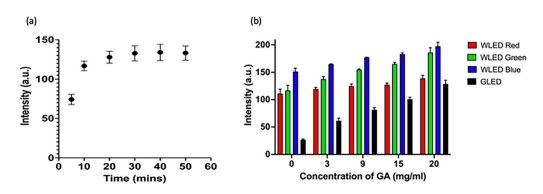


Fig. 3 (a) Development of colorimetric assay response signal intensity over time (n = 3); (b) colorimetric intensities of R–G–B channels (white LED) and green channel (green LED).

measured by capturing and analysing images using both LEDs to determine the optimum LED and channel to be used. As seen in Fig. 3(b), all three channels from the white LED as well as the green LED responded to the changing concentrations of the target protein. While the overall intensities of the colorimetric signals from the white LED images were higher, which was expected given the higher power of the white LED, it was observed that the green LED provided the highest sensitivity across the dynamic range, even higher than the green channel of the white LED. Hence, all intensity measurements for both assays were performed *via* green LED image captures. The colorimetric intensity measurements were performed for both assays and the green LED was observed to provide the best sensitivity for both assays.

Once the assay parameters were finalized, both assays were run to obtain responses in the relevant concentration ranges. Since the assays are colorimetric, qualitative responses of both assays were obtained and can be visually observed in Fig. 4(a) and (b). Intensities of colorimetric signals at the test lines (first red line) increased as the concentration of the target protein increased. However, for monitoring purposes, a qualitative result provides insufficient information since knowledge of disease severity is required. It has been established that up to 16% glycated albumin indicates a healthy non-diabetic person.¹⁰ However, if the % GA value is much higher, it indicates the need for intervention from healthcare professionals by prescribing alternative dietary or exercise regimens. Quantitative results were obtained by conducting analyses of images obtained using the handheld reader. The quantitative responses (calibration curves) in (Fig 5(a) and (b)) were obtained by analysing images from the green LED, 30 min after sample addition. Each concentration was tested in triplicate. A dynamic range of 3-20 mg ml⁻¹ was achieved for the glycated albumin assay and 20–50 mg ml⁻¹ for the serum albumin assay. It was observed from the plots that both assays have a linear correlation with the concentrations of the target proteins. Regression parameters for both assays have been listed in the ESI (Table S2†). Based on these parameters, the LoD for the glycated albumin assay was calculated to be 0.8 mg ml⁻¹ and for the serum albumin assay was 1.5 mg ml⁻¹.

Assay response in serum

Simulated blood serum was used to evaluate the response of both assays in a complex biological matrix. The serum was purchased without any albumin to control the amount of glycated and serum albumin present. This enabled the creation of samples containing different concentrations of glycated albumin and serum albumin to test both assays. The results are listed in Table 1 below. It can be observed that the recoveries are in the range of 93% to 104%, which implies that serum components do not have a major effect on the assays. Serum dilution was not required for either assay due to the running buffer volume of 50 μ l, which was instrumental in improving the flow and overcoming the higher complexity of the serum by preventing non-specific binding.

Table 1 Detection of target proteins in serum

Sample	Doped (mg ml ⁻¹)	Recovered $(mg ml^{-1})$	Recovery (%)	SD
GA 1	8	8.25	103.15	4.87
GA 2	20	19.48	97.4	2.7
HSA 1	35	33	94.28	3.61
HSA 2	50	46.89	93.78	4.04

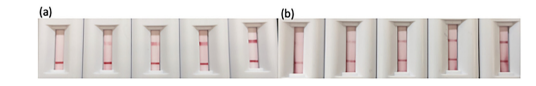


Fig. 4 Qualitative response for (a) glycated albumin and (b) serum albumin assays.

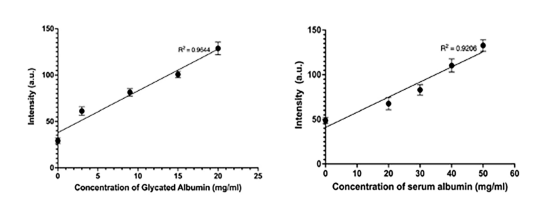


Fig. 5 Quantitative response (calibration curves) for (a) glycated albumin and (b) serum albumin assays.

Dual assay response

After both assays were optimized for the individual lateral flow configuration and tested in a complex biological medium, the final step was to incorporate them into a single cartridge. The dual assay cartridge was designed to enable the user to run both assays simultaneously and generate the % glycated albumin result in a single test instead of having to run two separate tests with two separate samples. The cartridge was designed in Solidworks and 3D printed in the lab. Fig. 6(a) illustrates the design of the cartridge. Fig. S1[†] includes top and bottom views of the cover and bottom of the cartridge. After designing and printing the cartridges, the symmetry of flow to both sides of the cartridge was confirmed by measuring and comparing signals for the same assay on both sides of the cartridge. It can be observed from Fig. 6(b) that signal intensities of assays on both sides of the cartridge were comparable implying that similar sample volumes flowed on both sides of the assays. Once flow symmetry was confirmed, dual assay response was tested via three samples with varying concentrations of glycated and serum albumin: S1 consisted of 3.5 mg ml⁻¹ GA and 25 mg ml⁻¹ HSA; S2 was comprised of 10 mg ml⁻¹ GA and 30 mg ml⁻¹ HSA; S3 was a mix of 20 mg ml⁻¹ GA and 30 mg ml⁻¹ HSA. The concentration levels were chosen such that the dynamic range of each assay was covered between the three samples. Fig. 7 demonstrates the dual assay response for all levels. Concentration levels of sample one (GA: 3.5 mg ml^{-1} and SA: 25 mg ml^{-1}) were chosen to demonstrate that lowest concentrations of both assay dynamic ranges could be successfully detected in the dual assay system. Sample 1 was equivalent to 12% glycation which corresponded to a non-diabetic healthy person. The second sample was at the midpoint of the glycated albumin assay range. For the serum albumin assay, the total serum albumin concentrations combined was equivalent to the mid-point of its dynamic range.

This sample corresponded to 25% glycation which was comparable to mild to moderately severe diabetes. Finally, in the last sample, the concentration of glycated albumin was chosen to be the upper limit of the dynamic range and serum albumin concentration was chosen such that the summation of both concentrations was equivalent to the highest relevant concentration for serum albumin. It was a 40% glycation sample which corresponded to a severely diabetic condition. In sample 3, unglycated serum albumin concentration was

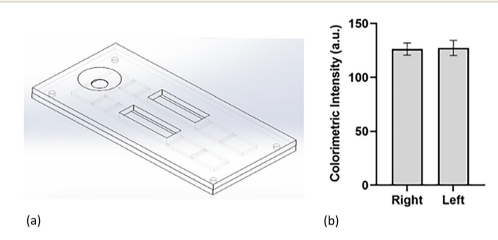


Fig. 6 (a) Dual assay cartridge design; (b) symmetry of flow in both channels of dual assay cartridge.

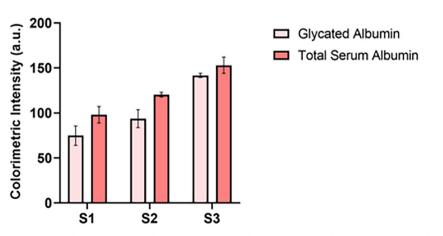


Fig. 7 Dual assay response (S1: 3.5 mg ml⁻¹ GA + 25 mg ml⁻¹ HSA; S2: 10 mg ml⁻¹ GA + 30 mg ml⁻¹ HSA; S3: 20 mg ml⁻¹ GA + 30 mg ml⁻¹ HSA).

maintained at the same concentration as sample 2 to demonstrate that only an increase in glycated albumin could also be detected by the serum albumin assay. It was observed that there was a 7-10% increase in signal intensities compared to individual assays run separately. However, this was expected since in case of the glycated albumin assay, non-specific binding from unglycated serum albumin to a small extent was anticipated. And in case of the serum albumin assay, aptamer binding to unglycated and glycated albumin was expected to differ to some extent resulting in minor differences in resultant signal intensities even at the same concentrations. In addition, all samples were doped in simulated blood serum which also contributed to the non-specific binding albeit to a small extent. However, it was noted that signal intensities for all concentration levels increased to a similar extent and hence the dual assay cartridge could in fact successfully be used to simultaneously measure glycated and total serum albumin to obtain the % glycated albumin value from a single sample and single test.

Conclusion

In this manuscript, two aptamer-based lateral flow assays were developed to measure glycated albumin and serum albumin in their relevant concentration ranges. The choice of aptamers was made over other recognition elements owing to their thermal stability, low cost, and relatively long shelf life. Both assays used gold nanoparticle-based colorimetry since it does not require any additional use of dyes which lowered the overall cost of the system. Both assays were immobilized onto lateral flow platforms, which afforded advantages like compact size, portability, and ease-of-use. A handheld colorimetric reader developed by the team was used in conjunction with lateral flow cartridges. When the sample was added to the cartridge, the target protein interacted and bound to the aptamer on the gold nanoparticles which were stored in the conjugate layer. The protein-particle complexes then flowed onto the nitrocellulose membrane, where they were captured at the test line. The number of particles at the test line was proportional to the concentration of the target protein in the sample, which was observed in the assay responses. The dynamic ranges for both assays were $3-20 \text{ mg ml}^{-1}$ and $20-50 \text{ mg ml}^{-1}$, which are both physiologically relevant, implying that sample dilution will not be required to run the assays. Finally, a dual assay cartridge was used to run both assays simultaneously to obtain the value of % glycated albumin from a single test. In conclusion, this research demonstrated that a dual lateral flow assay plus handheld colorimetric reader setup can effectively monitor glycated and serum albumin for ultimate use as a point-of-care sensor for monitoring gestational diabetes.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by the National Science Foundation Precise Advanced Technologies and Health Systems for Underserved Populations (PATHS-UP) Engineering Research Center through award no. EEC-1648451. The authors would also like to acknowledge the discussions with Dr Samuel Mabbott on lateral flow assay optimization.

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