

# Automated microfluidic devices integrating solid-phase extraction, fluorescent labeling, and microchip electrophoresis for preterm birth biomarker analysis

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**Abstract** We have developed multichannel integrated microfluidic devices for automated preconcentration, labeling, purification, and separation of preterm birth (PTB) biomarkers. We fabricated multilayer poly(dimethylsiloxane)-cyclic olefin copolymer (PDMS-COC) devices that perform solid-phase extraction (SPE) and microchip electrophoresis ( $\mu$ CE) for automated PTB biomarker analysis. The PDMS control layer had a peristaltic pump and pneumatic valves for flow control, while the PDMS fluidic layer had five input reservoirs connected to microchannels and a  $\mu$ CE system. The COC layers had a reversed-phase octyl methacrylate porous polymer monolith for SPE and fluorescent labeling of PTB biomarkers. We determined  $\mu$ CE conditions for two PTB biomarkers, ferritin (Fer) and corticotropin-releasing factor (CRF). We used these integrated microfluidic devices to preconcentrate and purify off-chip-labeled Fer and CRF in an automated fashion. Finally, we performed a fully automated on-chip analysis of unlabeled PTB biomarkers, involving SPE, labeling, and  $\mu$ CE separation with 1 h total analysis time. These integrated systems have strong potential to be combined with upstream immunoaffinity extraction, offering a compact sample-to-answer biomarker analysis platform.

**Keywords** Microfluidics/microfabrication · Capillary electrophoresis/electrophoresis · Clinical/biomedical analysis · Separations/instrumentation · Extraction (SFE|SPE|SPME)

## Introduction

Microfluidics can miniaturize benchtop processes, promising fast, sensitive, inexpensive, and portable analysis systems suitable for point-of-care applications [1–4]. On-chip sample preparation is an important step in this direction, allowing multiple time-consuming off-chip steps such as preconcentration, purification, and labeling to be performed on the same platform [5]. Integration of on-chip sample preparation with miniaturized tools thus can simplify and automate the analysis process. Many recent studies focus on addressing this need, including preconcentration, purification, and separation of proteins in urine [6]; labeling and separation of preterm birth (PTB) biomarkers [7, 8]; affinity capture, tryptic digestion, and isotopic labeling of an apoptosis biomarker [9]; preconcentration and purification of core needle biopsy samples for steroid quantitation [10]; and on-chip immunocapture and sensing of proteins in whole blood and urine samples [11]. PTB biomarker analysis in miniaturized platforms is an area of interest because PTB is the most common complication of pregnancy, the primary cause of neonatal illnesses [12], and PTB risk can be correlated with specific molecular markers in bodily fluids [12]. Importantly, a panel of three serum peptides combined with six previously known biomarkers was shown to provide specific (~80%) and sensitive (~90%) prediction of the occurrence of a PTB several weeks in the future [13].

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Microfluidic sample preparation can utilize various methods including electrokinetic manipulation [14], solid-phase extraction (SPE) [15, 16], and size-selective membranes [6]. Reversed-phase SPE selectively retains analytes based on hydrophobic interactions [17]. Porous polymer monoliths for SPE [18, 19] are versatile and effective, particularly in microchips due to the convenience of on-chip monolith polymerization and the ability to control the porosity and chemical properties through the composition [15].

Microchip electrophoresis ( $\mu$ CE) is an effective analysis method for multiplexed biomarkers [20, 21]. Importantly,  $\mu$ CE increases in power and versatility when integrated with upstream sample processing like SPE [22]. Mohamadi et al. [23] developed an integrated microfluidic device combining immunoaffinity capture, preconcentration, and  $\mu$ CE of  $\beta$ -amyloid peptides in cerebrospinal fluid. Araya-Farias et al. [24] combined affinity SPE with  $\mu$ CE for the analysis of phosphopeptides. Nordman et al. [25] combined reversed-phase SPE,  $\mu$ CE, and electrospray ionization mass spectrometry for the analysis of verapamil. Sonker et al. [7] developed  $\mu$ CE and on-chip labeling modules for PTB biomarker analysis, but on-chip integration of SPE and  $\mu$ CE was not demonstrated. Notably, all these integrated microfluidic analyses used electrokinetic operation, which can simplify device fabrication, but applied voltages can be problematic for samples with high or low ionic strength. Alternatively, fluids can be manipulated on-chip with pneumatic pumps and valves [26, 27]. Sahore et al. [8] developed a pressure-injected  $\mu$ CE device for PTB biomarkers with improved signal and separation, but in that work, only  $\mu$ CE separation was demonstrated. Recently, Kumar et al. [28] reported a pressure-driven microfluidic device, integrating SPE and  $\mu$ CE for the preconcentration and separation of a PTB protein biomarker. This device used a single input reservoir and processed different solutions to perform the analysis of pre-labeled analytes, but it required the reservoir to be emptied and refilled in order to change solutions. Thus, two key improvements in automated analysis would be to interface multiple solution inputs with on-chip SPE and  $\mu$ CE, and to perform on-chip labeling in the analysis to eliminate off-chip processing steps.

In this work, we developed pressure-actuated, multi-channel, multilayer integrated microfluidic devices to analyze PTB biomarkers. We improved upon the design, operation, fabrication, SPE, and  $\mu$ CE conditions from a previous study [28] to make integrated poly(dimethylsiloxane)-cyclic olefin copolymer (PDMS-COC) microfluidic devices that preconcentrate, label, purify, and separate PTB biomarkers in an automated fashion. We optimized SPE retention and elution conditions to purify ferritin (Fer) and corticotropin-releasing factor (CRF) from unbound label and developed a  $\mu$ CE assay for these PTB biomarkers. Finally, we carried out a fully automated on-chip analysis of unlabeled PTB biomarkers, including the processes of

SPE, labeling, purification, and separation. This system is being developed as part of a larger integrated package that also entails upstream immunoaffinity extraction from a blood serum sample [22], with potential to provide a compact, integrated biomarker analysis platform.

## Experimental section

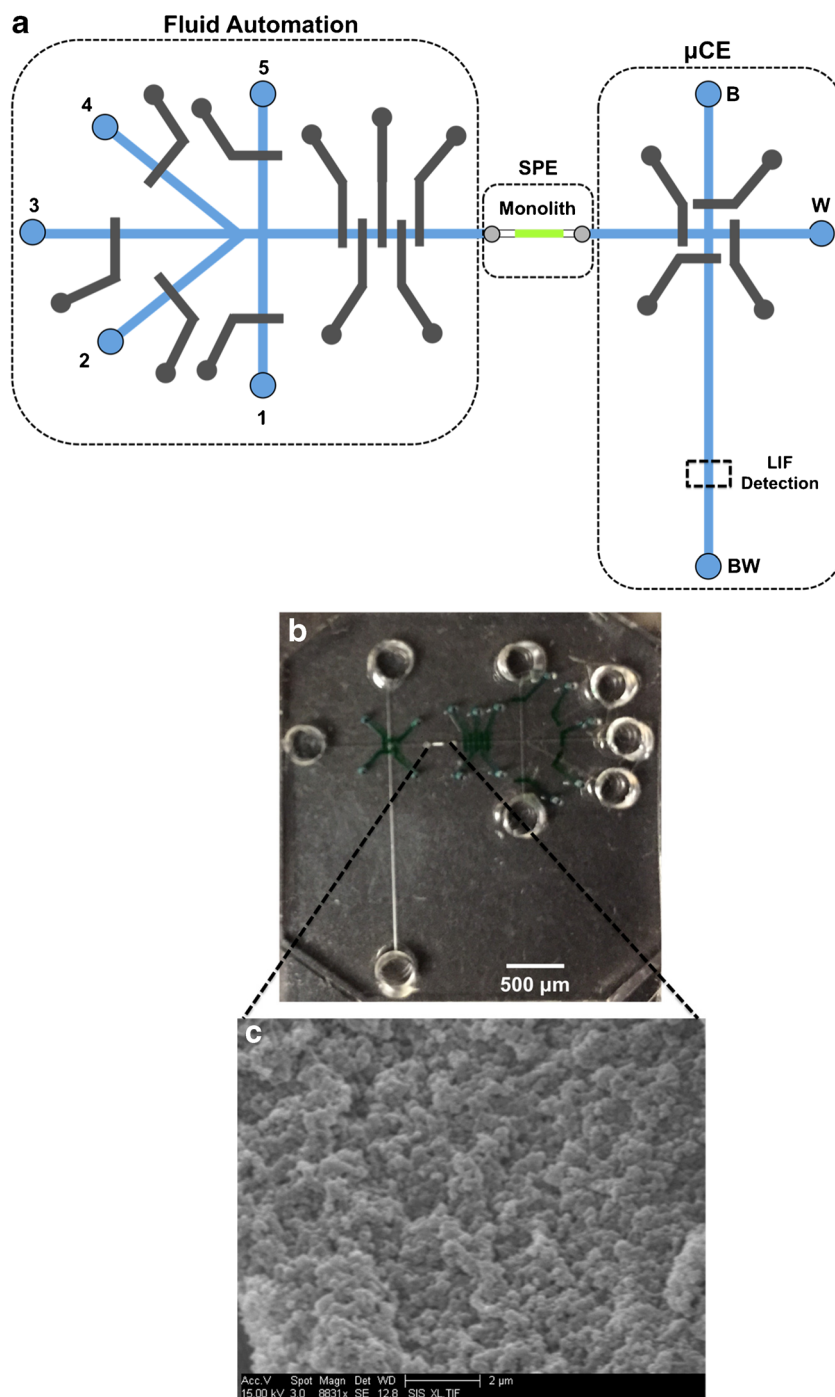
### Chemicals and materials

Methylcellulose (MC), ethylene glycol dimethacrylate (EGDMA), 1-dodecanol, and 2,2-dimethoxy-2-phenylacetophenone (DMPA) were purchased from Sigma-Aldrich (St. Louis, MO). Octyl methacrylate (C8) was obtained from Scientific Polymer Products (Ontario, NY). Acetonitrile (ACN) and cyclohexanol were purchased from Fisher Scientific (Pittsburgh, PA) and J.T. Baker (Phillipsburg, NJ), respectively. Sodium carbonate, sodium bicarbonate, and Fer were purchased from EMD Millipore (Billerica, MA). Human CRF was purchased from Biomatik (Wilmington, DE). PDMS base and curing agent were obtained from Dow (Midland, MI). Food coloring to visualize control layer channels was purchased from Walmart (Bentonville, AR). AZ P4620, SU-8 2025, and S1805 photoresists were purchased from MicroChem (Westborough, MA). COC (Zeonor 1020R) was obtained from Zeon Chemicals (Louisville, KY). Alexa Fluor 488-TFP Ester (AF 488) was purchased from Life Technologies (Carlsbad, CA). All solutions were made in deionized (DI) water either filtered by a Barnstead water purifier or purchased directly from Thermo Scientific (Waltham, MA).

### Integrated microfluidic device design

Microfluidic chip masks were designed using CleWin software (Informer Technologies, Shingle Springs, CA), with a device schematic shown in Fig. 1a. Each microfluidic chip had the top two layers made from PDMS and the bottom two layers constructed from COC. The control layer had a five-valve peristaltic pump for fluid injection, five pneumatic valves for fluid input control, and four pneumatic valves surrounding the  $\mu$ CE injection intersection for plug capture. The width of all pneumatic valves was 250  $\mu$ m, and the spacing between the peristaltic pump valves was 120  $\mu$ m. A four-valve design with 200- $\mu$ m spacing from the injection intersection was used to capture the sample plug [8]. The fluidic PDMS layer had 100- $\mu$ m-wide and  $18 \pm 1$ - $\mu$ m-deep ( $n = 23$ ) channels, with five input reservoirs connected to fluid channels for automation: (1) sample, (2) label, (3) rinse, (4) label eluent, and (5) labeled sample eluent; and three reservoirs connected to channels for  $\mu$ CE: waste (W), buffer (B),

**Fig. 1** Integrated microfluidic device. **(a)** Schematic view of the multichannel integrated microchip design, showing hydrodynamic controls, SPE column, and  $\mu$ CE section. Eight reservoirs (1 sample, 2 fluorescent label, 3 rinse, 4 label eluent, 5 labeled sample eluent, B buffer, BW buffer waste, W waste) were used. Color scheme: *blue* fluidic channels, *gray* control channels, and *green* monolith. Operation involves sample loading on the monolith through pump operation with valve 1 opened and valves 2–5 closed. Subsequently label is loaded with valve 2 opened and valves 1 and 3–5 closed. After labeling, the monolith is rinsed with solution from reservoir 3 with valve 3 opened and valves 1–2 and 4–5 closed. With valve 4 opened and valves 1–3 and 5 closed, the unattached label is eluted using solution pumped from reservoir 4. Upon pumping the labeled sample eluent with valve 5 opened and valves 1–4 closed, the eluted PTB biomarker fluid plug is transferred into the injection intersection and then separated by  $\mu$ CE with LIF detection. Except during  $\mu$ CE runs, the separation channel valves to the  $\mu$ CE intersection were kept closed while leaving the injection channel valves open. During  $\mu$ CE, the injection valves were closed while opening the separation valves. **(b)** Photograph of a microfabricated device, showing a PDMS pneumatic pump and valves filled with colored dye solution, fluid channels in PDMS-COC, and the porous polymer monolith in COC. **(c)** SEM image showing monolith morphology in a device



and buffer waste (BW). The distance between the B and W reservoirs to the injection intersection was 0.5 cm, and the distance from the BW reservoir to the intersection was 1.5 cm. The top and bottom COC layers were  $\sim 100$   $\mu$ m and 1 mm thick, respectively, with the top layer having  $\sim 200$ - $\mu$ m diameter through-holes that connected the COC to the PDMS and with the bottom layer having an 80- $\mu$ m-wide, 1.5-mm-long, and  $29 \pm 3$ - $\mu$ m-deep ( $n = 22$ ) channel in which the reversed-phase porous polymer monolith was formed.

## Device fabrication

Device fabrication followed a method previously developed in our laboratory [28]; see also Fig. S1 in the Electronic Supplementary Material (ESM). Figure 1b is a photograph of a fully assembled integrated microdevice. A C8 reversed-phase porous polymer monolith was fabricated inside the COC microfluidic channel using 24% C8 and 11% EGDMA, with 10% cyclohexanol and 55% 1-dodecanol as

porogens, and 1% DMPA as the photoinitiator. The monolith solution was exposed to UV light through the bottom COC layer through a Cr mask for 11 min, and a 0.6-mm-long monolith was obtained. The polymerized monolith was rinsed using 2-propanol for at least 30 min. Other details of the monolith fabrication procedure are reported in an earlier study [28]. Morphology of the C8 monolith was observed using scanning electron microscopy (SEM; Philips XL30 FEG, FEI, Hillsboro, OR) as shown in Fig. 1c; the details of the SEM sample preparation and imaging procedure have been published previously [29].

### Experimental setup

Details of the experimental setup can be found in our previous publications [7, 8]. Briefly, a Nikon Eclipse TE 300 microscope (Nikon, Melville, NY) with 488 nm laser excitation (JDSU, Shenzhen, China) and photomultiplier tube fluorescence detection was used. Separation voltages were applied using platinum electrodes connected to a variable power supply (Stanford Research Systems, Sunnyvale, CA). The LabVIEW (National Instruments, Austin, TX) control software and electrically controlled pneumatic system (Clippard Instrument, Cincinnati, OH) were expanded from our earlier work to accommodate 14 valves. The peristaltic pump could sustain a maximum of 18 psi before the PDMS-COC interface began to delaminate, so the air supply for solenoid actuation was split into two parts, one addressing the peristaltic pump (18 psi) and the other addressing the individual valves (25 psi).

### Device operation

Samples, AF 488, and different ACN eluents were prepared in 10 mM sodium bicarbonate buffer (BCB, pH 9.9). Labeling conditions such as buffer type, ionic strength, pH, and analyte/dye ratio were first optimized off-chip with analysis by pressure-injected  $\mu$ CE [8]. BCB (10 mM, pH 9.9) with 1:40 and 1:10 analyte-to-AF 488 ratios for Fer and CRF, respectively, provided the best off-chip labeling, and these conditions were adopted for on-chip labeling. Off-chip-labeled Fer (450 kDa) and CRF (4.7 kDa) were purified four times at 14,000 rpm for 15 min using 50- and 3-kDa centrifugal filters (EMD Millipore), respectively. The concentration of labeled Fer was measured using a NanoDrop Spectrometer (Thermo Scientific).

Some experiments were performed with 200- $\mu$ m valve spacing, three-layer PDMS-COC devices for pressure-injected  $\mu$ CE [8] of Fer (200 nM), CRF (750 nM), and a mixture of Fer and CRF at these same concentrations. Before experiments, pressure-injected  $\mu$ CE devices were washed with 2-propanol, DI water, and 10 mM BCB, with vacuum drying between each step. Subsequently, the devices

were filled with running buffer (25 mM BCB, pH 9.9, with 0.05% MC), and samples dissolved in 10 mM BCB were injected with 20 Hz peristaltic pump actuation for 30 s. We previously reported detailed operation procedures for pressure-injected  $\mu$ CE devices [8].

Integrated microfluidic devices were prepared for use in the following operational sequence. First, with empty channels, all valves were actuated and their control lines were filled using colored dye solutions; if the channels were filled with liquid before filling the control lines, it increased bubble formation due to the air permeability of PDMS. Next, the fluid automation reservoirs and corresponding channels were filled with 90% ACN in 10 mM BCB. Filling the COC channel and the monolith required the peristaltic pump to prime the monolith with 90% ACN solution for 2 min. Then, the solution in the reservoirs was replaced with 50% and then 30% ACN, and the peristaltic pump was operated for 1 min each time. Next, all reservoirs were filled with 10 mM BCB, pH 9.9, and the peristaltic pump was operated with solution pumped from each reservoir (keeping the other valves closed) for at least 30 s to fill all the  $\mu$ CE channels. Finally, with the  $\mu$ CE injection valves closed, the separation channel was filled with 25 mM BCB (pH 9.9) containing 0.05% MC.

With the device filled, preconcentration of Fer (50 nM), purification of CRF (750 nM), and separation of off-chip-labeled Fer (50 nM) and CRF (750 nM) were performed as follows. Samples were loaded for 15 min with the peristaltic pump (20 Hz) with valve 1 and the injection channel valves opened but all others closed, flowing solution to reservoir W. Solutions in reservoirs 1–5 were replaced with fresh 10 mM BCB (pH 9.9), and all the automation channels were rinsed individually, followed by filling reservoir 4 with the label eluent (50% ACN in BCB) and reservoir 5 with the labeled sample eluent (90% ACN in BCB). The label eluent was flowed (20 Hz) to elute the unattached dye. Finally, the labeled sample eluent flow (20 Hz) eluted the PTB biomarkers using discrete injections of 5 s each, followed by  $\mu$ CE separation with laser-induced fluorescence (LIF) detection 10 mm from the injection intersection as described previously [8].

To determine its elution and retention properties on the monolith, a 40- $\mu$ M AF 488 sample was loaded for 5 min. After 5 min, a 10 mM BCB (pH 9.9) rinse and 15–90% ACN solutions were flowed, while LIF was measured after the monolith.

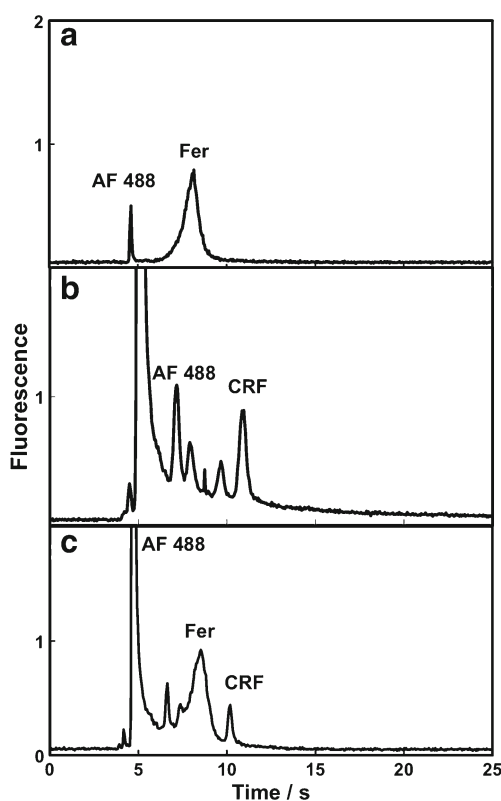
For on-chip labeling of PTB biomarkers, the integrated device was prepared and operated similarly. Either individual component [Fer (100 nM) or CRF (750 nM)] or a mixture of the two (150 nM Fer and 1  $\mu$ M CRF) was loaded for 15 min with 20 Hz peristaltic pump actuation. Next, 10 or 40  $\mu$ M AF 488 was loaded for 5 min at 13.3 Hz actuation, and the reaction was allowed to proceed for 30 min. After 10 mM BCB rinsing and 50% ACN label elution (20 Hz actuation), the labeled samples were eluted and transferred to the injection

intersection using 90% ACN in 5-s increments, with each injection probed by  $\mu$ CE separation and LIF detection as before. Three to eight 5-s injections were needed to clear the dead volume between the monolith and  $\mu$ CE intersection. Between 5 and 15 additional injections were probed by  $\mu$ CE. The cumulative injection time (including clearing the dead volume) is reported for each experiment in its figure caption. Additional device operation details can be found in the Fig. 1 caption.

## Results and discussion

### Microchip electrophoresis of PTB biomarkers

Initially, experiments were performed to analyze the separation of AF 488-labeled Fer and CRF, using a pressure-operated PDMS-COC  $\mu$ CE device [8]. Microchip electropherograms are shown in Fig. 2; most of the unreacted label is

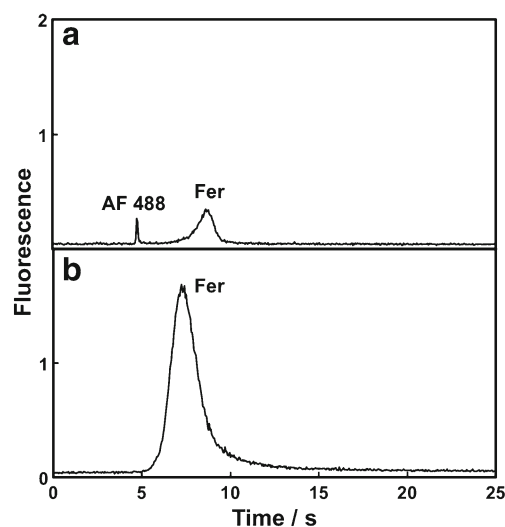


**Fig. 2** Pressure-driven injection  $\mu$ CE for off-chip AF 488-labeled (a) Fer (200 nM,  $N = 21,000$  plates/m), (b) CRF (750 nM,  $N = 77,000$  plates/m), and (c) Fer (200 nM,  $N = 11,000$  plates/m)/CRF (750 nM,  $N = 270,000$  plates/m) mixture using a three-layer PDMS-COC device with  $18 \pm 1 \mu\text{m}$  ( $n = 16$ ) channel depth, 24 mm separation channel length, and 200  $\mu\text{m}$  valve spacing. Additional experimental conditions: 20 Hz pump actuation rate, 30 s injection time, 10 mm detection distance, 1500 V  $\mu$ CE voltage, and 25 mM BCB (pH 9.9) with 0.05% MC separation buffer

removed after 30 kDa filtering of the labeled Fer (Fig. 2a), but even 3 kDa filtration did not completely remove AF 488 and possible decomposition byproducts from labeled CRF (Fig. 2b). Consequently, the CRF sample had additional peaks at faster migration times than CRF; nonetheless, the individual PTB biomarkers were well resolved from these peaks. In the separation of the PTB biomarker mixture, Fer co-migrated with some of the AF 488 peaks, but it was still well resolved from the CRF peak (Fig. 2c). The number of theoretical plates per meter (N/m) for PTB biomarker peaks is given in each figure caption; these N/m values are comparable to published work for PDMS  $\mu$ CE devices [8]. On the basis of this separation data, we decided to use similar buffer conditions for  $\mu$ CE in our integrated microfluidic devices.

### Sample Preconcentration

Preconcentration is useful for improving limits of detection or aiding in further sample preparation. We used the C8 reversed-phase monolith in the COC channel of our integrated microfluidic device to preconcentrate AF 488-labeled Fer. After loading, rinsing and label elution, the preconcentrated Fer was incrementally eluted, injected, and analyzed by  $\mu$ CE. The pressure-injected microchip electropherogram in Fig. 3a shows a labeled Fer peak that is well resolved from AF 488 still present in the off-chip filtered sample. Figure 3b shows Fer preconcentrated on-chip by SPE and analyzed by  $\mu$ CE using an integrated microfluidic device. Because of the monolith

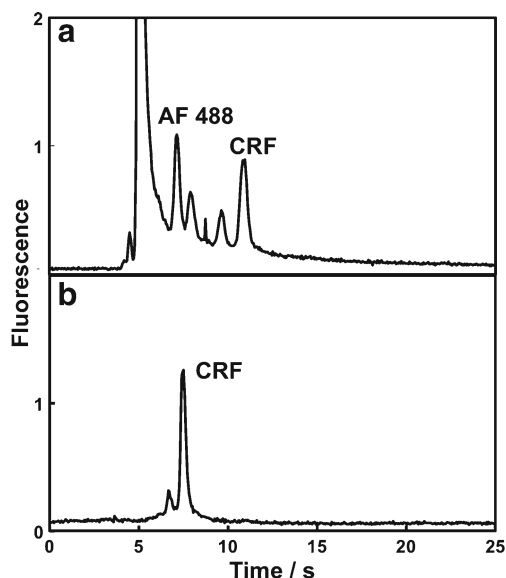


**Fig. 3** PTB biomarker preconcentration using an integrated microfluidic device. Pressure injection  $\mu$ CE for (a) off-chip AF 488-labeled Fer (100 nM,  $N = 19,000$  plates/m) using a three-layer PDMS-COC device without preconcentration and (b) off-chip AF 488-labeled Fer (50 nM,  $N = 5600$  plates/m) preconcentrated on the monolith, eluted, injected, and separated by  $\mu$ CE using a four-layer PDMS-COC-integrated microfluidic chip. In (a), all parameters were the same as in Fig. 2. In (b), all parameters except  $\mu$ CE voltage (1200 V) and separation channel length ( $19 \pm 1$  mm) were the same as in Fig. 2

enrichment and label elution steps, the extra dye peak was removed and  $\mu$ CE of Fer yielded a peak with a 13-fold greater area than in Fig. 3a. Comparing Fig. 3a to the  $\mu$ CE data for all eight runs, we calculated a  $\sim$ 50-fold preconcentration factor in the integrated microfluidic device. To account for minor variations in operational conditions, a spiked reference analyte could be used to determine a normalization factor for more accurate quantitation with our preconcentration approach. Importantly, the use of pneumatic valves allows multiple injections of the small-volume enriched sample that would be difficult to achieve using electrokinetic methods [29]. In addition to accomplishing preconcentration, the experiments also demonstrate the ability of our integrated microfluidic devices to do automated analysis.

### Sample purification

In integrated microfluidics, sample purification can reduce the interferences present. We used our integrated microfluidic devices to purify AF 488-labeled CRF using differences in retention on the monolith. Unconjugated AF 488 was eluted first using 50% ACN, while the purified and labeled CRF was eluted from the monolith with 90% ACN. CRF that was labeled with AF 488 and 3 kDa filtered off-chip was analyzed by pressure-injected  $\mu$ CE. Dye peaks in addition to labeled CRF were visible, even after filtration (Fig. 4a). The reversed-

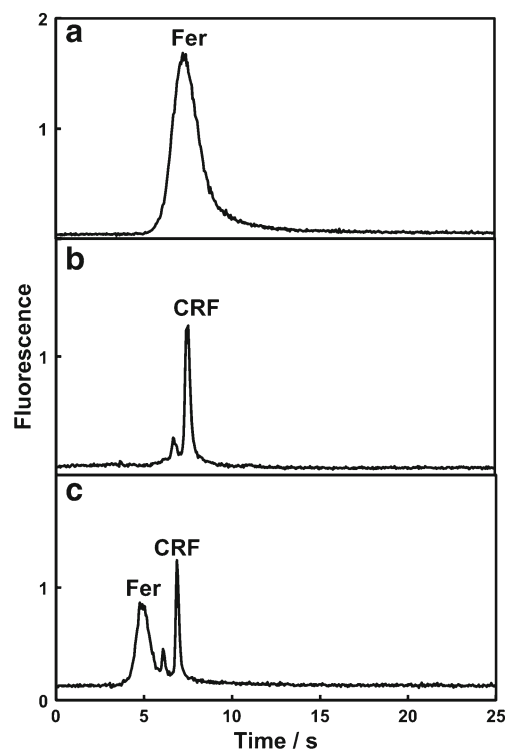


**Fig. 4** PTB biomarker purification in an integrated microfluidic device. Pressure-injected  $\mu$ CE of off-chip AF 488-labeled and 3 kDa filtered CRF (750 nM) (a) without on-chip purification in a three-layer PDMS-COC device ( $N = 77,000$  plates/m) and (b) with on-chip SPE purification after a 40-s cumulative injection time ( $N = 180,000$  plates/m). In (a), all parameters were the same as in Fig. 2. In (b), all parameters except  $\mu$ CE voltage (1500 V) were the same as in Fig. 3b. The difference in migration times for CRF can be explained by different separation voltages, the use of different devices [ $\mu$ CE only in (a) vs. integrated in (b)], and different captured sample solutions [10 mM BCB in (a) and 90% ACN in (b)]

phase monolith in our device purifies more efficiently than the filter (compare 15 and 90% ACN elution traces in ESM Fig. S2), such that when we preconcentrated the same sample in our integrated microfluidic device followed by selective label and sample elution, sharp peaks for CRF and one minor impurity were obtained (Fig. 4b). This clearly demonstrates the benefits of our devices for automated on-chip SPE- $\mu$ CE sample purification and analysis.

### Integrated microfluidic device analysis of pre-labeled PTB biomarkers

Disease diagnosis often requires determination of a panel of biomarkers; as a first step in this direction, we have analyzed a mixture of two off-chip-labeled PTB biomarkers in our integrated system. Fer and CRF were loaded individually, extracted, and eluted, and the corresponding  $\mu$ CE results are shown in Fig. 5a, b. Selective elution in our integrated microfluidic device yielded electropherograms free of label with just a minor impurity in Fig. 5b along with the biomarker peak. Different  $\mu$ CE voltages (1200 V in Fig. 5a and 1500 V in Fig. 5b) cause the Fer and CRF peaks to have similar migration times. A mixture containing Fer and CRF was similarly



**Fig. 5** Off-chip AF 488-labeled PTB biomarker analysis in an integrated microchip for (a) Fer (50 nM, 30-s cumulative injection time,  $N = 5600$  plates/m), (b) CRF (750 nM, 40-s cumulative injection time,  $N = 180,000$  plates/m), and (c) Fer (100 nM,  $N = 8800$  plates/m)/CRF (750 nM,  $N = 220,000$  plates/m) mixture (30-s cumulative injection time). All parameters except  $\mu$ CE voltage [1500 V for (b)] were the same as in Fig. 3b

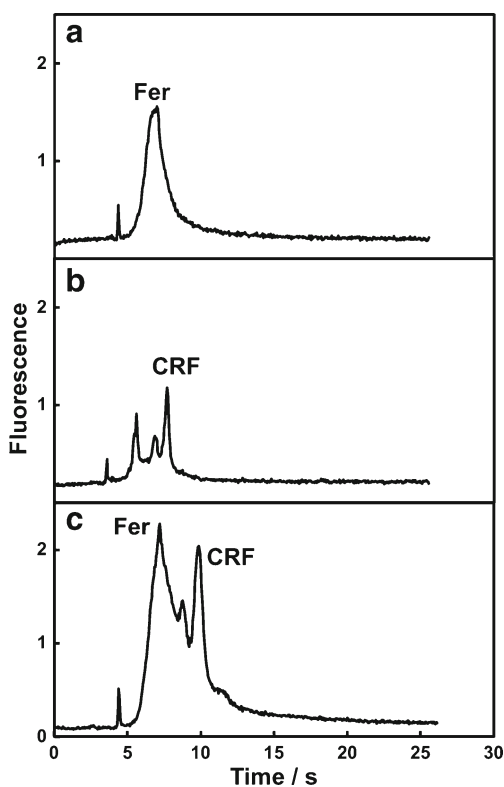
processed in an integrated microfluidic device, and the electropherogram in Fig. 5c shows the purified Fer and CRF peaks clearly resolved. The combination of  $\mu$ CE with upstream SPE improves the biomarker separation, which would be difficult to achieve using only SPE [7]. These results demonstrate the ability of our devices to process and separate multiple PTB biomarkers.

### On-chip labeling of PTB biomarkers

A key advantage of our integrated system is the ability to do automated labeling followed by separation of PTB biomarkers. Off-chip results were optimal with a molar label-to-analyte ratio as high as 40, so we tested the retention and elution characteristics of the C8 monolith with 40  $\mu$ M AF 488. Elution of AF 488 was studied using 15–90% ACN, and we found that 50% ACN was the best eluent for AF 488 since it yielded the largest peak (see ESM Fig. S2).

To demonstrate integrated PTB biomarker analysis, unlabeled Fer and CRF were loaded and labeled on-chip as described in the “Experimental section.” Figure 6a shows  $\mu$ CE

of on-chip-labeled Fer with a peak shape similar to that observed for off-chip-labeled Fer in Fig. 3b. Figure 6b shows  $\mu$ CE of on-chip-labeled CRF with a peak shape similar to the data obtained using off-chip-labeled CRF in Fig. 4b. A tenfold higher CRF concentration compared to Fer was used because CRF (3 kDa) has fewer amine groups for labeling than Fer (450 kDa), and the retention of CRF on the monolith is lower than that for Fer. Although we used 40  $\mu$ M AF 488 for on-chip labeling, the very small unattached label peak at 4 s clearly demonstrates the ability of our devices to label as well as purify the labeled PTB biomarkers. Finally, we loaded a mixture containing unlabeled Fer and CRF, and labeled, purified, and separated the PTB biomarkers by  $\mu$ CE, as seen in Fig. 6c. Figure 6c shows the resolution of the on-chip-labeled Fer and CRF, with a small CRF impurity peak between them and a smaller AF 488 peak at  $\sim$ 4 s. Data in Fig. 6c match well with the off-chip-labeled sample in Fig. 5c; the minor offset in migration times is due to different separation conditions. These results demonstrate fully automated preconcentration, labeling, purification, and separation of multiple PTB biomarkers in our integrated microfluidic devices.



**Fig. 6**  $\mu$ CE after on-chip SPE, labeling [with 40  $\mu$ M AF 488 for (a) and (c) and 10  $\mu$ M AF 488 for (b)], and elution of PTB biomarkers in an integrated microfluidic device. (a) Fer (100 nM, 40-s cumulative injection time,  $N = 6000$  plates/m). (b) CRF (750 nM, 35-s cumulative injection time,  $N = 110,000$  plates/m). (c) Fer (150 nM)/CRF (1  $\mu$ M), with 80-s cumulative injection time. Additional separation buffer was loaded in B and BW reservoirs before this injection. All parameters except the applied voltage [800 V for (c)] were the same as in Fig. 3b

### Conclusions

We have developed pneumatically controlled, integrated microfluidic devices for automated on-chip labeling and  $\mu$ CE separation of PTB biomarkers. The devices had a four-layer PDMS-COC structure and were fabricated using soft lithography and hot embossing techniques. Fluidic controls in the top PDMS layer included a five-valve peristaltic pump, five pneumatic valves for flow control, and four pneumatic valves for  $\mu$ CE injection control. The second PDMS fluidic layer had five input reservoirs connected to fluid channels and a  $\mu$ CE setup. The bottom two layers were made from COC and had a reversed-phase C8 porous polymer monolith fabricated inside the microchannel for SPE and fluorescent labeling. We used our integrated microfluidic devices to preconcentrate, purify, and separate off-chip-labeled PTB biomarkers in an automated fashion. Furthermore, we demonstrated the ability of our integrated microfluidic devices to do an analysis of a mixture of PTB biomarkers. Finally, we carried out a fully automated analysis including preconcentration, labeling, purification, and  $\mu$ CE of a mixture of PTB biomarkers on our microdevices in 1-h total analysis time.

Key advantages of our approach are the integration of sample preparation and  $\mu$ CE in a single platform to on-chip label and separate biomarkers in an automated fashion. Additionally, the use of pneumatic fluid controls allows the use of stronger eluents and eliminates complications like mobility bias that are associated with electrokinetic methods. The use of an on-chip peristaltic pump further reduces dead volumes associated with external syringe pumps, for example. The use of pneumatic

valves offers additional control in sample manipulation, allowing plugs to be injected multiple times in a discrete and precise manner. We are now working to develop a fully integrated sample-to-answer microfluidic device for PTB biomarker analysis that further combines upstream immunoaffinity extraction with these processes. Multilayer device fabrication using conventional clean room techniques, while feasible, can be a limiting factor in these analyses due to long fabrication times. Three-dimensional printing may provide an attractive alternative to reduce fabrication times, provided truly microfluidic (<100  $\mu\text{m}$ ) features can be created [30, 31]. Finally, although this work is directed towards the analysis of preterm birth biomarkers, the general nature of our approach makes it broadly applicable to other disease-related biomarkers.

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#### Compliance with ethical standards

**Conflict of interest** The authors declare that they have no conflict of interest.

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