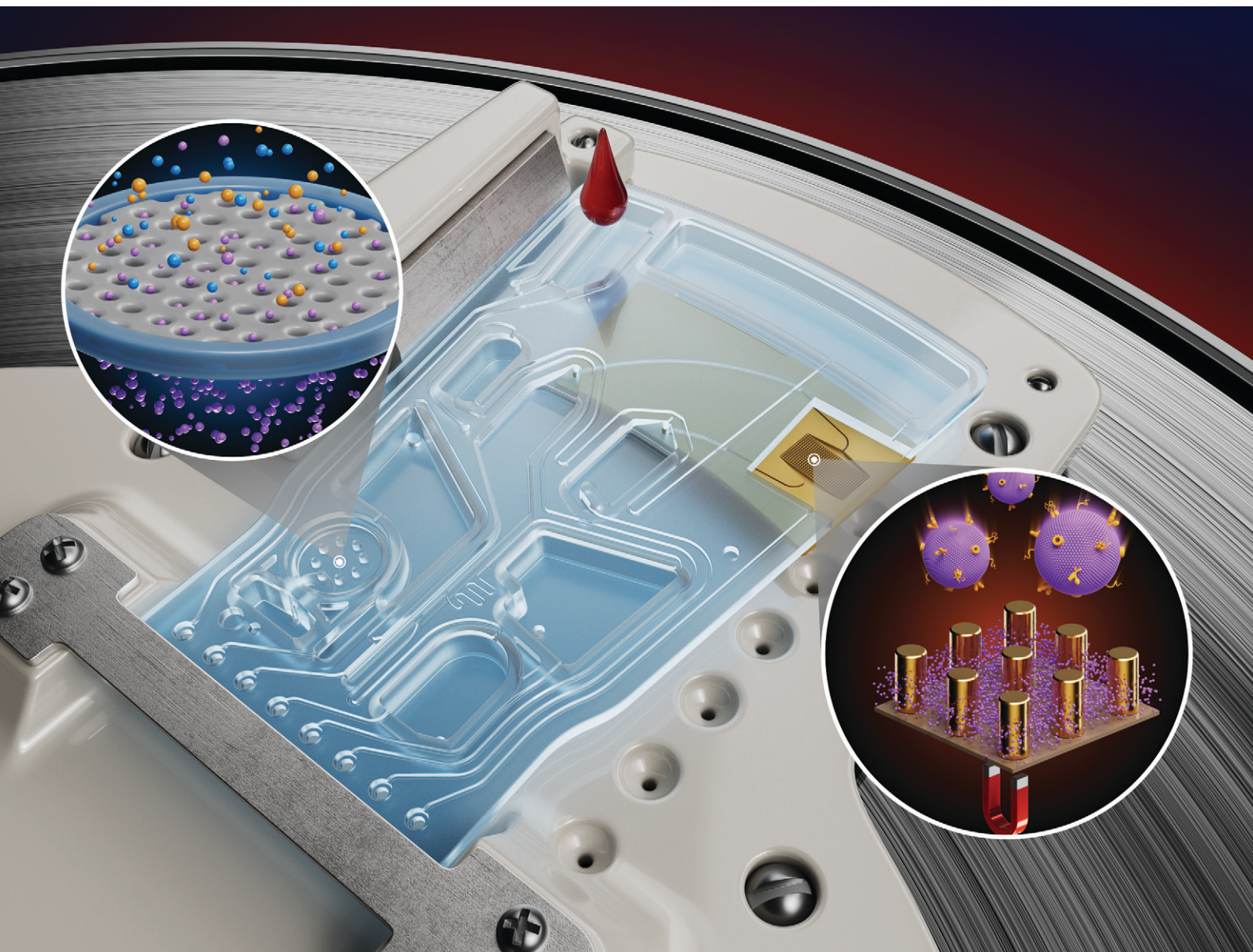


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***EV-Blade*: an automated centrifugal-pneumatic cartridge for size- and affinity-based exosome isolation from whole blood**

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Extracellular vesicles (EVs), especially the exosome-sized subset are increasingly exploited as minimally invasive cancer biomarkers. These small vesicles are abundant in biofluids and play crucial roles in intercellular communication and disease progression by transporting bioactive molecules. Exosomes offer distinct diagnostic and prognostic advantages over traditional cancer biomarkers, but purifying and extracting exosomes from blood remains challenging. There is a need to simply and cost-effectively isolate exosomes from milliliter quantities of whole blood for transcriptional and other omics-based research. Addressing this gap, we propose a microfluidic cartridge, the *EV-Blade*, for size and affinity-based purification of exosomes on a centrifugal microfluidic platform. We demonstrate a method to automate exosome purification from whole blood samples on a single microfluidic cartridge. The *EV-Blade* system combines blood centrifugation, plasma filtration for EV size selection and immunomagnetic capture using functionalized superparamagnetic nanoparticles targeting CD9, CD63, and CD81 exosomal surface proteins. We report on the device performance, purity of exosome recovery and the quality of RNA collected following on-chip EV lysis. We use this automated method to detect relevant long coding and non-coding RNA transcripts in circulating blood exosomes, showcasing the *EV-Blade* for use in cancer patient risk stratification. The system presented herein represents a significant advancement in exosome purification, offering a robust and automated platform for liquid biopsy-based cancer research and clinical applications. This innovation holds promise for cancer diagnosis, prognosis, and monitoring through non-invasive biomarkers.

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Introduction

Extracellular vesicles (EVs), including exosomes, have emerged as promising vehicles for biomarker discovery in cancer research.¹ Here we define exosomes as a subset of small EVs (sEVs)² with an outer diameter under 200 nm and bearing various combinations of the CD9, CD63 and CD81 transmembrane proteins.³ They are secreted by various cells in the body and can be found abundantly in biofluids such as blood, urine, and saliva.⁴ They play critical roles in intercellular communication, disease progression, and therapeutic resistance by shuttling various bioactive molecules, including proteins, metabolites, and RNA.^{5–9} Consequently, exosomes have attracted significant attention as potential sources for non-invasive diagnostic and prognostic information in cancer patients.^{10–12}

In the context of cancer, exosomes are implicated in multiple processes, including modulation of the tumor microenvironment, drug resistance, and the establishment of pre-metastatic niches.^{5,7–9,13,14} Furthermore, exosomes offer several advantages over traditional cancer biomarkers, such as tumor-associated antigens and circulating tumor cells. Exosomes are more abundant in peripheral blood and can be detected at earlier disease states,¹⁵ making them highly promising for liquid biopsy-based strategies in cancer diagnosis, prognosis, and monitoring. However, the purification of exosomes from peripheral blood remains a challenging and time-consuming process due to their small size and complex physical properties. While differential ultracentrifugation has long been considered the gold standard,¹⁶ it is often accompanied by further purification steps based on density gradient or affinity capture techniques. Other purification methods, such as size exclusion chromatography,^{17,18} precipitation with volume-excluding polymers¹⁹ and filtration,²⁰ have also been explored. Notably, size-based pre-selection combined with immuno-affinity capture have shown promising results for biomarker discovery using total RNA sequencing (RNA-seq).^{21,22} Multiple

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microfluidic devices have been developed to purify EVs from various types of biofluids. Whole blood is a complex matrix, so most devices require initial sample pre-treatment by centrifugation and then use plasma or serum as inputs in EV filtration,^{23–25} deterministic lateral displacement²⁶ or immuno-affinity^{27–31} protocols. Other devices have implemented sized-based only^{32–34} or affinity-based only³⁵ EV purification starting from whole blood. Only two devices currently combine both selection methods^{36,37} from whole undiluted blood, with starting volumes in the 150 to 300 μL range. EVs derived from blood contain a low amount of RNA, usually within the nanograms per milliliter (ng mL^{-1}) range.³⁸ Blood volumes between 1 and 3 mL may be required to obtain quality RNA-seq data for biomarker discovery and disease monitoring.^{18,39} Despite the progress made in exosome purification techniques, there is still a need for a simple, cost-effective workflow capable of efficiently isolating specific subsets of sEVs from milliliter volumes of whole blood. The complexity of current, labour-intensive EV separation and purification methods hinders the widespread adoption of exosome-based liquid biopsies in a clinical setting.

Building on our previous work in active pneumatic, centrifugal microfluidics^{40–42} and microfluidic-based immuno-magnetics,^{43,44} we present here the *EV-Blade*, implementing an automated method to purify exosomes from milliliter quantities of whole blood performed wholly on a low-cost microfluidic cartridge compatible with large-scale manufacturing. During continuous centrifugal rotation, the *EV-Blade* cartridge combines size-based EV pre-selection, using both whole blood centrifugation and plasma cross-flow filtration, together with EV immunomagnetic capture with functionalized superparamagnetic nanoparticles (MNPs) targeting well-established exosomal surface proteins CD9, CD63, and CD81.^{45,46} The automated device enabled repeated recovery of pure exosomes lysates, with yields equivalent to manual reference protocols for combined size and affinity-based selection. Following on-chip lysis of exosomes purified from whole blood samples, we recovered high quality RNA. Using RT-qPCR we detected relevant long coding and non-coding exosomal RNA transcripts representative of B-cell acute lymphoblastic leukemia (B-ALL),²² highlighting the potential for translation of our method in the field of exosome-based liquid biopsies for non-invasive cancer biomarkers monitoring.

Experimental

Active pneumatic control in centrifugal microfluidics

In standard centrifugal microfluidics, the centrifugal force generated by rotation drives on-chip fluid displacement. Valves are used for temporization of protocol steps and spatial control of both samples and reagents. Commonly, capillary valves,⁴⁷ hydrophobic valves⁴⁸ or passive siphons⁴⁹ enable or restrict fluid transfers at specific rotation speeds. Other dynamic approaches to centrifugal valving include magnetic⁵⁰ or mechanical actuation⁵¹ and dissolvable

materials⁵² which significantly increase cartridge complexity. Moreover, these valves typically operate irreversibly and once open the entire fluid content dispenses from one reservoir to the next. Mixing actions can be achieved using complex microstructures,^{53,54} active agitation with magnetic particles or shaking cycles.⁵⁵ Given the inherent limitations of passive and mechanically complex approaches to valving, mixing, and fluid handling, alternative solutions such as active pneumatic pumping have been proposed. Previous work by Clime *et al.*⁴⁰ describe the implementation of the PowerBlade system, enabling active pneumatic pumping within a centrifugal microfluidics platform while in rotation. Briefly, a computer-controlled pressure manifold is integrated on the platform's rotor, capable of delivering positive and negative pressure to the device cartridge through individually addressable ports.

On the PowerBlade system fluid displacement can now be controlled by balancing centrifugal forces and pneumatic pressure gradients. This capability enables movement of liquids against the centrifugal force (reverse pumping), from the device periphery back to the center of rotation, and by extension, back and forth movements through fluidic structures which are particularly useful for operations involving incubation or washing. Pneumatic actuated siphon valves can be reused repeatedly, enabling precise dispensing of fluid aliquots from the same reservoir. Injecting air into liquid filled device reservoirs through the pneumatic manifold generates bubbles that rise in the liquid against the centrifugal force field enabling fast and efficient mixing. Finally, routing flow from a single starting reservoir towards two or more different paths (flow switching) is easily achieved. Overall, active pneumatic pumping on the PowerBlade platform offers great flexibility for fluid handling inside centrifugal microfluidic devices.

Microfluidic cartridge fabrication

The main body of the *EV-Blade* fluidic cartridge was designed with Solidworks (Dassault Systèmes, FR) and fabricated with a CNC milling machine (Q350, Menig Automation, USA) from custom bio-compatible thermoplastic blanks in-house injected, cyclo olefin polymer (COP) Zeonor 1060R (Zeon Chemicals LP, USA) for low volume prototyping, followed by a small volume injection molding run (1000 units) of the microfluidic cartridges in Zeonor 1060R (Protolabs, USA).

The magnetic capture sub-unit uses a previously developed magnetic gradient concentrator called the *M-Chip*.⁴³ It consists of an array of micron-sized, high aspect ratio cylindrical pillars (20 μm diameter, 50 μm pitch) hot embossed in a polymer substrate. A master mold with multiple insert dies was defined on a silicon wafer using photolithography and deep reactive ion-etching (80 μm depth). The micropattern was first replicated by hot embossing in a 300 μm thick fluorinated ethylene polymer (FEP) film to create a re-usable working stamp for serial replication of the microstructured pillar array in 1 mm thick,



150 mm diameter Zeonor 1060R blanks. After embossing, individual arrays were die-cut and coated with $5.5 \pm 0.5 \mu\text{m}$ of nickel followed by 200 nm of gold *via* electroless deposition to protect and passivate the nickel surface. The fluidic layer was fabricated by hot embossing in-house, extruded, thermoplastic elastomer (TPE, Mediprene OF 400M) sheets using an epoxy mold replicated from a PDMS mold, itself replicated from a 50 μm thick negative photoresist (SU8 1075, Gersteltec, CH) patterned on a silicon wafer. After punching inlet and outlet through hole *vias* in the TPE fluidic layer, it was bonded to the microstructured pillar array by contact at room temperature (RT). Finally, the through hole *vias* were aligned with the top of the *EV-Blade* cartridge and bonded by contact. The complete assembly was then annealed at 60 °C for 12 h in an oven to increase adhesion and release residual stress in the injection molded parts.

The cross-flow filtration sub-unit uses a PVDF 0.22 μm filtration membrane (Sterlitech, USA) bonded to a seat structure inside the filtration chamber and fixed using a laser-cut ring of pressure sensitive adhesive (PSA, Adhesive Research, USA).

The *EV-Blade* cartridge microfluidic channels and reservoirs were sealed by laminating a PSA (Adhesive Research, USA) together with a thin COP (Zeon Chemicals LP, USA) backing layer.

Cell culture

REH (ETV6-RUNX1⁺ human B cell precursor leukemia; ATCC CRL-1567) and 697 (TCF3-PBX1⁺ human B cell precursor leukemia; DSMZ ACC-42) cells were first cultured in RPMI 1640 medium (Wisent, CA) supplemented with 10% heat-inactivated fetal bovine serum (FBS) (Wisent, CA) and 1% penicillin/streptomycin (Wisent, CA). They were then transferred gradually to 100% X-VIVO 15 serum-free hematopoietic cell medium (Lonza, CH) over the course of 4 passages. All cell lines were cultured in a 37 °C incubator with 5% CO₂. Conditioned cell culture medium (CCM) was collected 48 h after passage and apoptosis was kept under 5%.

Blood collection

Blood samples were collected in K2 EDTA coated tubes (BD, USA) by venipuncture from healthy donors. The institutional review board of the CHU Sainte-Justine Research Center approved the research protocol and written informed consent was obtained from all participants. The study was performed in accordance with the Declaration of Helsinki.

Manual EVs isolation from CCM for blood spiking

For all the cellular models, 45 mL of CCM was collected at 70% cell confluency and centrifuged at 300×g for 10 min at RT to remove suspended cells followed by a second round of centrifugation at 2500×g for 10 min at RT to remove large cell debris. The supernatant was collected and concentrated

down to 900 μL by ultrafiltration using Amicon Ultra-15 centrifugal filter units with 100 kDa molecular weight cut-off (Millipore Sigma, USA). Concentrated CCM was used as input for EVs isolation by size-exclusion chromatography SEC using qEV original 70 nm columns (Izon Science, NZ). For each sample, 4 × 500 μL fractions were collected, filtered through a 0.22 μm centrifugal filter unit at 12 000g for 4 min (Ultrafree-MC, Millipore Sigma, USA) to remove large EVs (>200 nm in diameter) and concentrated down to 0.5 mL using Amicon Ultra-0.5 centrifugal filter units with 30 kDa molecular weight cut-off spun at 14 000×g. The volume of concentrated sEVs added to the blood samples was adjusted to align with C_q values of B-ALL biomarker candidates we previously measured in patient samples *via* RT-qPCR.²²

Manual exosomes and sEVs purification from blood

Freshly collected blood was pooled and divided into 1 mL aliquots. Aliquots used for modeling exosome based liquid biopsies were spiked with concentrated sEVs to align with C_q values of B-ALL biomarker candidates we previously measured in patient samples *via* RT-qPCR.²²

For all manual methods, blood samples underwent centrifugation at 1100×g for 10 min for plasma separation. For size-exclusion chromatography (SEC), the collected plasma was filtered using a syringe and a 0.8 μm cellulose acetate filter membrane (GVS, Italy) following recommendations of Gaspar *et al.*¹⁸ Then, 500 μL of filtered plasma was used as input for EVs isolation *via* SEC using qEV original 70 nm columns (Izon Science, New-Zealand). Four 500 μL fractions per sample were collected and centrifuged at 20 000×g to remove larger microvesicles.

For manual immunomagnetic exosomes purification, the plasma was filtered through 0.22 μm centrifugal filter units at 12 000×g for 4 min (Ultrafree-MC, Millipore Sigma, USA) to remove large EVs (>200 nm in diameter). Each filtered plasma sample was then processed with the EasySep Human Pan-Extracellular Vesicle Positive Selection Kit (Stemcell, CA), purifying EVs bearing CD9, CD63 and CD81 surface proteins following the manufacturer's protocol. After the final wash step, bead bound EVs were either lysed in RL buffer (Norgen, CA) for RNA extraction or in RIPA buffer (Sigma) for protein analysis.

Tunable resistive pulse sensing (TRPS)

TRPS measurements were conducted using a qNano instrument (Izon Science, NZ). Various polyurethane nanopores appropriately stretched (NP100, NP150, NP200, NP400, NP800, Izon Science, NZ) and the matching diluted calibration polystyrene beads (CPC100, CPC200, CPC400, CPC800, Izon Science, NZ) were used to obtain the particle size distribution (PSD) of the EV samples. All samples were diluted in phosphate buffered saline (PBS) with 25 mM trehalose and 0.05% Tween20 to minimize EV aggregation. Each measurement was performed at two different pressures



(5 and 10 mbar). Data were processed and analyzed using the Izon Control Suite software v3.3.2.2001.

Optimization of magnetic capture element

The capture efficiency of magnetic nanoparticles (MNPs) using the *M-Chip* device was assessed by optical absorbance measurements as described previously.^{43,56} Briefly, stand-alone *M-Chip* devices with various gap sizes between the nickel/gold coated pillars were assembled onto in-house, injection molded luer-lock cover plates and connected to a syringe pump. See Poncelet *et al.*⁴⁴ for more details on the test device assembly, noting that a single, larger N52 magnet was used for the current study to model the configuration of the *EV-Blade* cartridge. A custom setup, consisting of a pair of tube holders each containing a photodiode and an 850 nm wavelength led coupled to a signal amplifier and associated computer software was used to record the measurements in real-time. Optical absorbance in the outlet tubing was compared to a reference tube filled with the MNPs dilution buffer. For each measurement, 100 μL unlabelled EasySep MNPs (Stemcell, CA) were diluted in 1 mL PBS. The *M-Chip* array was first filled with the dilution buffer to record a baseline then switched to diluted MNPs. Once maximal absorbance at the outlet was reached MNPs capture was initiated in the *M-Chip* device by placing it either between a pair of permanent NdFeB magnets (K&J Magnetics, USA) or along the side of a single magnet. After reaching an absorbance plateau corresponding to the maximal capture capacity for a set flowrate and magnet configuration, dilution buffer was flowed through the *M-Chip* to simulate washing steps and the recording was ended.

MNPs preparation for on-chip processing

The manufacturer's original protocol starts with incubation of the pre-processed plasma sample with 50 $\mu\text{L mL}^{-1}$ of antibodies cocktail for 10 min at RT, followed by incubation with 100 $\mu\text{L mL}^{-1}$ of MNPs for 10 min at RT before the washing and EVs lysis steps. Due to limited availability of pressure ports on the PowerBlade instrument, constraining the total number of steps possible to automate on a single device, we introduced the following variation. The MNPs and antibodies cocktail were incubated together off-chip for 10 min at RT before loading in the *EV-Blade* device.

Assessment of CD9⁺ CD63⁺ CD81⁺EV recovery and purity with the *EV-Blade*

Concentration of known EV protein markers CD9, CD63, CD81 and syntenin-1 (Synt-1) as well as cytochrome-C (Cyt-C) as a marker of cellular organelles contamination and potential contamination by various apolipoproteins (Apo-A1, Apo-A2, Apo-B, Apo-C2, Apo-E) were measured using bead-based immunoassay panels (exosome characterization 6-plex human and human apolipoprotein ProcartaPlex panels, Invitrogen, USA) on a CS 1000 Autoplex Analyzer (PerkinElmer, USA) after lysis of purified EVs in radioimmunoprecipitation assay

(RIPA) buffer. Starting from aliquots of the same pooled blood sample, we compared protein content in sEVs purified by SEC, manually purified with the EasySep Human Pan-Extracellular Vesicle Positive Selection Kit (Stemcell, CA) and EVs purified from whole blood with the *EV-Blade* cartridge on the PowerBlade. All samples were lysed using 100 μL of RIPA buffer and topped-up to 200 μL with PBS.

RNA extraction and quantification

For the samples purified with the *EV-Blade* microfluidic cartridges, bead-bound exosomes trapped in the magnetic pillar array were lysed on-chip with RL buffer (Norgen, CA). The lysate was collected from the cartridge at the outlet of the processed sample chamber for off-chip RNA purification. Total RNA was extracted using the Total RNA Purification Micro Kit (Norgen, CA) following the manufacturer's instructions with on-column DNase digestion (Norgen, CA) following the RNA binding step. For conventionally purified samples, the magnetic beads were removed from the lysate by centrifugation before further processing through the purification columns. For all samples, RNA yield and quality was assessed by electrophoresis on a 2100 bioanalyzer with the RNA 6000 nano or pico kits (Agilent Technologies, USA).

Results and discussion

EV-Blade microfluidic cartridge

The *EV-Blade* cartridge (Fig. 1) consists of multiple fluidic sub-units designed to automate sequential steps of the full EV purification protocol. These sub-units are connected by linking microfluidic channels. Handling and flow of sample and reagent fluids is controlled by active pneumatic pumping using eight pressure ports on the microfluidic cartridge. These cartridge ports interface directly with eight individually controlled pressure manifold ports on the PowerBlade instrument, as described previously.⁴⁰ The blood fractionation chamber is designed to accommodate up to 1 mL of whole blood. After white and red blood cells are sedimented by on-chip centrifugation, the plasma is transferred to the subsequent cross-flow filtration step through an outlet channel to collect just the upper plasma fraction. The filtration unit uses a two-chamber design. One chamber houses a 13 mm diameter membrane filter with 0.22 μm pore size, supported on a seat structure consisting of an array of 1 mm diameter posts and fixed in place using a laser-cut ring of pressure sensitive adhesive. The second chamber serves as a plasma overflow reservoir during the transfer process. The wash (700 μL) and lysis (100 μL) buffer reservoirs store reagents on-board until needed at specific time points in the protocol. During the protocol, waste products are directed to a collection reservoir (up to 3 mL). At the end of the protocol, lysate from purified EVs (100 μL) is sent to a dedicated processed sample reservoir. All the reservoirs have a dedicated port to allow manual reagent loading by pipetting before sealing with 4 mm diameter discs of PCR compatible adhesive foil prior to loading the



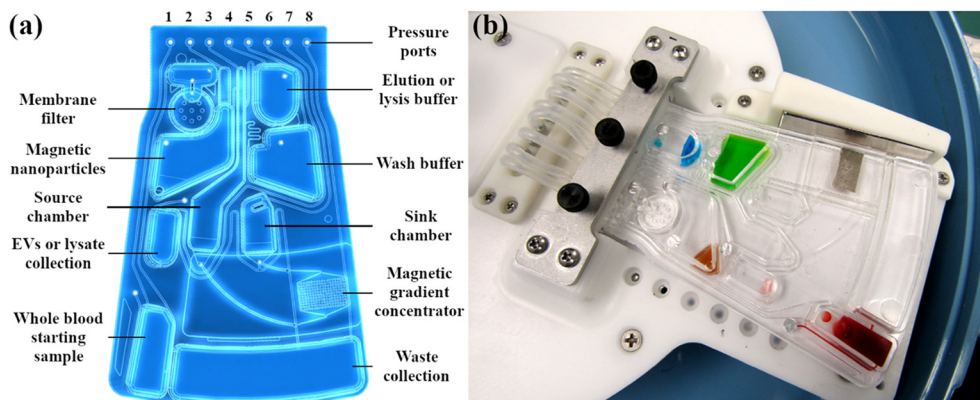


Fig. 1 Design and configuration of the *EV-Blade* cartridge. (a) 3D model detailing the disposition of the channels, reservoirs and pressure ports (labelled 1 to 8). (b) Photograph of an assembled cartridge filled with whole blood, MNPs, wash and lysis buffers and fitted on the PowerBlade instrument with a permanent magnet on the side. The main body of the cartridge is injection molded and the filter membrane and magnetic gradient concentrator are mounted during assembly. Blue and green food dyes were used as contrasting agents.

microfluidic *EV-Blade* cartridge on the PowerBlade instrument. Finally, the magnetic capture unit has three design elements: (i) a source chamber, (ii) a magnetic capture insert and (iii) a sink chamber. During sample processing, each reagent or mixture is first transferred from its holding reservoir of origin into the source chamber. Next the sample is transferred through the magnetic capture insert channels and magnetic pillar array and into the sink chamber. After

filling of the sink chamber, the liquid can be transferred either to the waste collection chamber or to the EV lysate collection chamber. By appropriate actuation of the pressure ports, liquids can be transferred from the sink chamber back through the capture insert to the source chamber, allowing multiple passes through the magnetic capture array. This unique back and forth fluid handling capability of the *EV-Blade* platform not only maximizes magnetic capture

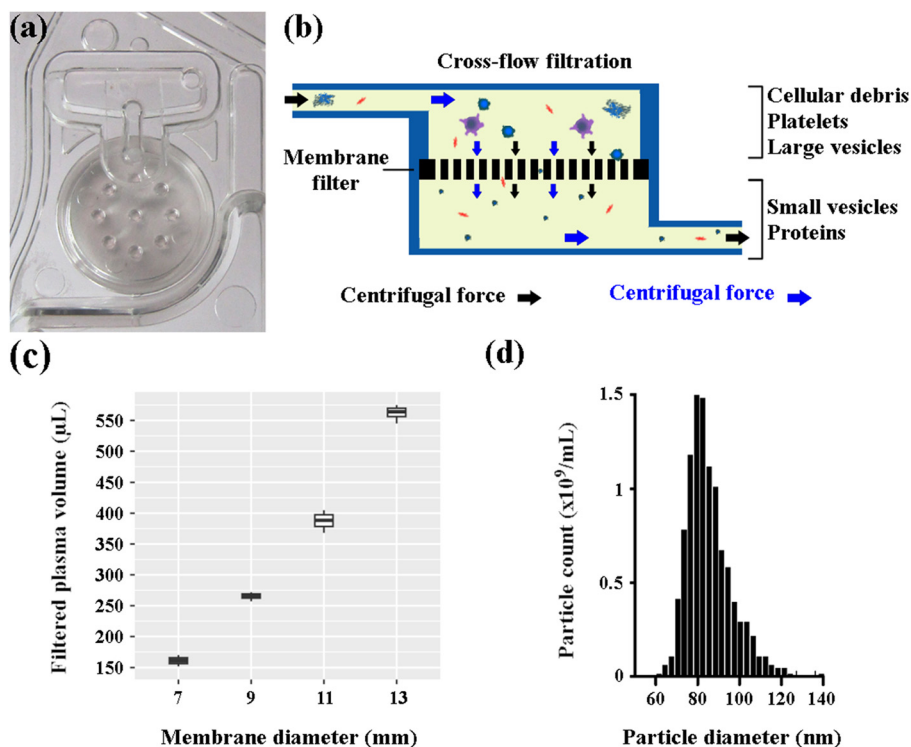


Fig. 2 Cross-flow filtration of plasma with a filter membrane insert. (a) Close-up photograph of the filtering chamber. (b) Schematics of the on-chip cross-flow filtration process. Large EVs, remaining platelets and cell debris are retained on top of the membrane allowing only smaller EVs and proteins to pass through. (c) Plasma volumes recovered after 0.22 μm filtration for various membrane diameters between 7 and 13 mm (d) particle size distribution and concentration of the small particles in plasma recovered after filtration with a 13 mm diameter 0.22 μm pore-size membrane.



efficiency, but maximizes washing and lysis efficacy as well. By harnessing the dual fluidic handling modes of centrifugal microfluidics and active pneumatic pumping together with the additional filtration and immuno-magnetic capture functional units, the *EV-Blade* enables the implementation of complex and clinically relevant EV preparation protocols on low-cost, industry scalable microfluidic devices.

Plasma filtering sub-unit

Fractionation of whole blood by centrifugation removes most of the cells, platelets and larger cell-debris from the sample. The filtering sub-unit shown in Fig. 2a is designed to remove the remaining platelets, cell debris and larger EVs from the plasma sample *via* cross-flow filtration (Fig. 2b) prior to immuno-capture. A membrane cut-off size of $0.22\ \mu\text{m}$ typically permits most sEVs to pass through. We assessed plasma recovery post-filtration for four different membrane diameters, ranging from 7 mm to 13 mm (Fig. 2c), determining that 13 mm membranes are suitable for processing up to 1 mL of whole blood on the *EV-Blade* cartridge. Additionally, we observed that filtration capacity is

directly proportional to the membrane area, providing design criteria for cross-flow filtration of plasma on centrifugal microfluidics platforms. Fig. 2d shows a total particle count in the solution after filtration on the cartridge. The resulting PSD is consistent with typical profiles observed for sEV preparations.^{17,18,57} At this stage, the sample is also expected to contain lipoproteins of comparable size.

Magnetic capture sub-unit

Fig. 3a shows a close-up photograph of the magnetic capture sub-unit with the source and sink chamber that regulate flow through the *M-Chip*. We used 400 nm diameter MNPs for exosome immunocapture within the *M-Chip*, a Ni/Au-coated micropillar magnetic array. Compared to the same mass of $2.8\ \mu\text{m}$ Dynabeads, these MNPs provide approximately nine times more capture surface area and 350 times more particles in suspension, significantly accelerating binding kinetics due to increased target-particle interactions. While the reduced magnetic content of the MNPs poses a challenge for separation, the *EV-Blade* design overcomes this by positioning a magnetic gradient concentrator (the *M-Chip*) adjacent to a

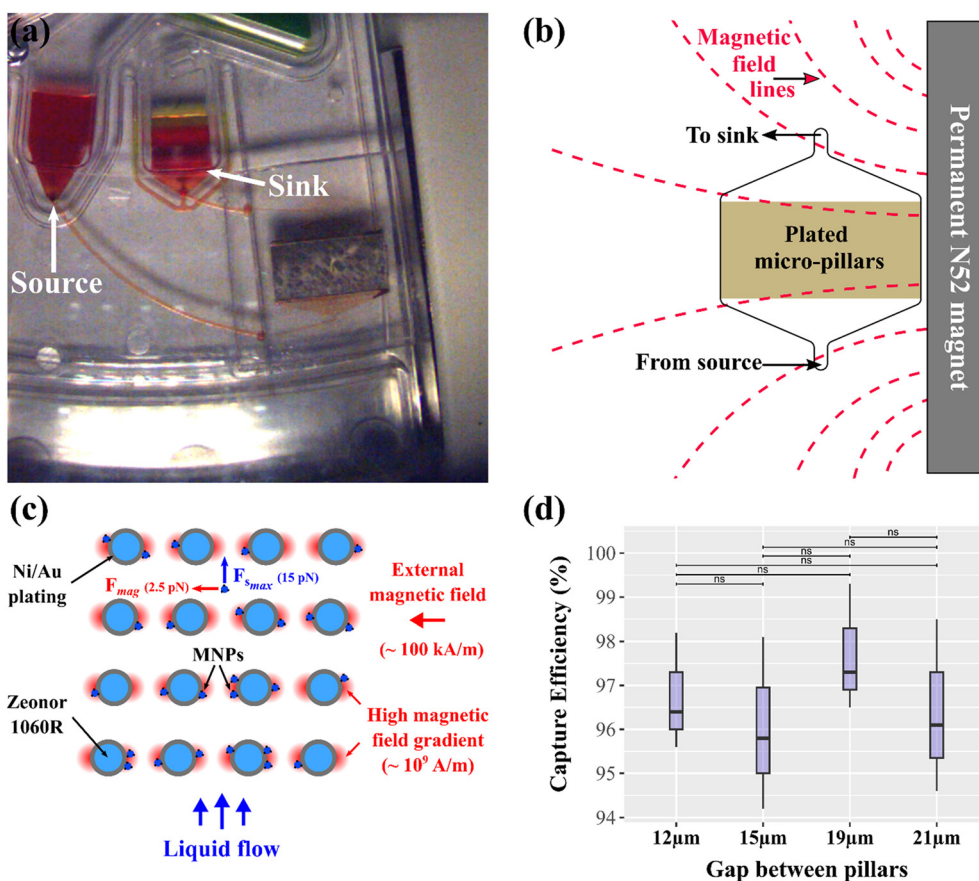


Fig. 3 Configuration and working principle of the magnetic capture sub-unit. (a) Close-up photograph showing the *M-Chip* insert, source and sink chambers and the permanent magnet mounted on the side of the microfluidic cartridge. (b) Schematic of the position of the *M-Chip* insert relative to the magnetic field lines generated by the permanent N52 magnet. (c) Schematic of a 4×4 array of Zeonor 1060R pillars plated with Ni/Au acting as strong magnetic field gradient concentrator (*M-Chip*) for the capture of MNPs. (d) Capture efficiency for 400 nm MNPs in solution while varying gap between pillars.



compact N52 permanent magnet (Fig. 3b). The pillar array generates local magnetic field gradients several orders of magnitude higher than the external magnet alone (Fig. 3c), enabling rapid MNP capture using magnets that can reasonably be mounted on a centrifuge rotor. All calculations regarding the MNP capture dynamics are detailed in the SI. The *EV-Blade* also incorporates a novel centrifugal-pneumatic filling strategy (Fig. 3c) that drives liquid into the *M-Chip* against the centrifugal force, efficiently displacing air and preventing bubble entrapment, a common limitation in previous syringe-pump implementations,^{43,44} thus enabling stable operation above 200 $\mu\text{L min}^{-1}$ without flow disruption. The absence of trapped bubbles ensures consistent and uniform liquid exchange, enabling faster washing steps and supporting a higher overall sample processing capacity within shorter timeframes. Further exploration of this deterministic filling phenomenon is presented in Fig. S1. Importantly, capture efficiency remains consistent despite minor manufacturing variations in pillar geometry (Fig. 3d and SI), underlining the robustness of the system for high-throughput, scalable immunocapture from complex biofluids at high flow rates.

Automated exosomes purification protocol

The sequential exosomes purification protocol including pressure values, rotation speeds, durations and open ports on the pressure manifold is summarized in Table 1. Fig. 4 shows selected stroboscopic images taken during an

experiment to illustrate the key steps of the automated protocol. A video of the entire experiment is available in the SI (Movie Fig. S4). The protocol starts with fractionation of the whole blood sample using just the centrifugal force function (step 1, Fig. 4a). The plasma fraction is then transferred to the filtration chamber by applying positive pressure through port #1. Plasma flow through the filtration membrane is driven both by the centrifugal force and a gentle negative pressure applied to ports #2 to #8 (step 2, Fig. 4b). The filtered plasma then flows directly into the incubation chamber containing a pre-loaded solution of MNPs functionalized with a cocktail of EV specific antibodies. By applying negative pressure to port #2, air is entrained through the plasma in a pressure differential with port #3 (at atmosphere), inducing bubble formation at the base of the incubation chamber. Buoyed to the chamber top by the centrifugal force, the air bubbles create a convection current within the chamber to efficiently mix the functionalized MNPs with the filtered plasma (step 3, Fig. 4c). This bubble mixing step is repeated periodically during the incubation step (10 min duration) to maintain MNP dispersion in the solution. Separately, at 5 min before the end of the incubation step, a small aliquot of pre-loaded wash buffer is transferred into the source chamber by applying positive pressure to port #5. Complete, bubble-free filling of the magnetic capture element is controlled by balancing the centrifugal force with active pneumatic pumping; this controlled wetting and filling primes the *M-Chip* micro-pillar array insert without trapping air, which

Table 1 Exosomes size- and affinity-based purification protocol on the *EV-Blade*

Step	Operation	Active ports	Applied pressure (psi)	Rotation speed (rpm)	Duration
Blood fractionation					
1	Centrifugation	—	—	800	30 min
Plasma filtration					
2a	Transfer plasma to filtration chamber	1	4.5	400	30 s
2b	0.22 μm plasma filtration	1, 2	1	500	60 s
Affinity-capture					
3a	Re-suspend MNPs ^a by bubble mixing	3	2	600	5 \times 1 s
3b	Plasma incubation with MNPs	—	—	100	10 min
	Bubble mixing during incubation	3	2	600	10 \times 2 s
Mchip priming					
4a	Send wash buffer ^b aliquot to sink	5	1	500	1 s
4b	Flow wash buffer in <i>M-Chip</i>	6, 7, 8	-0.5	200	60 s
4c	Empty drain to waste	8	-2	500	3 s
Magnetic capture					
5a	Send plasma + MNPs to sink	4	1	500	5 s
5b	Flow beads in <i>M-Chip</i>	6, 7, 8	-0.3	200	120 s
5c	Empty drain to waste	8	-2	500	3 s
Wash (repeat 3 times)					
6a	Send wash aliquot to sink	5	1	600	2 s
6b	Flow wash buffer in <i>M-Chip</i> (back and forth)	6,7,8	\pm 0.5	200	3 \times 45 s
6c	Empty drain to waste	8	-2	500	3 s
EVs lysis					
7a	Send lysis buffer ^c to sink	4	1.5	500	3 s
7b	Flow lysis buffer in <i>M-Chip</i> (back and forth)	6, 7, 8	\pm 0.5	200	6 \times 30 s
Lysate collection					
8	Send EVs lysate to processed sample chamber	6	-2	500	10 s

^a Suspension containing 50 μL EasySep MNPs incubated with 25 μL pan-EVs positive selection cocktail (Stemcell). ^b PBS. ^c Buffer RL (Total RNA Purification Micro Kit, Norgen).



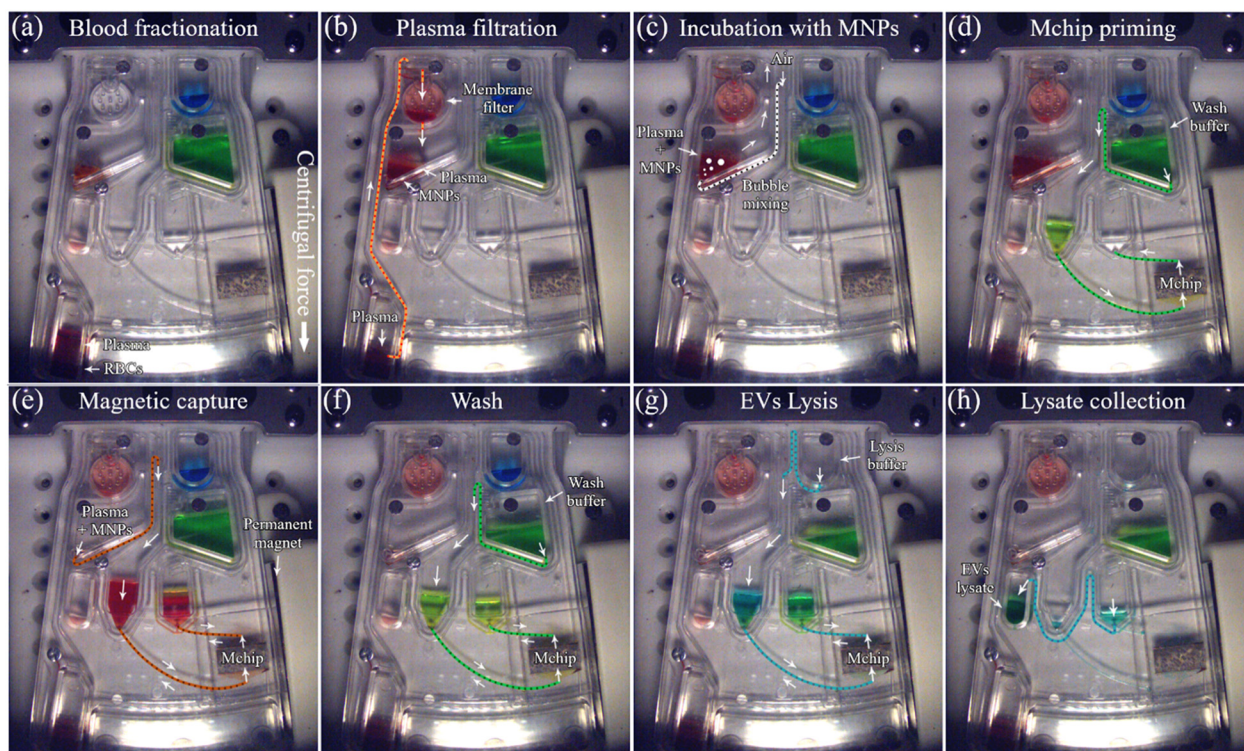


Fig. 4 Selected stroboscopic images of the *EV-Blade* cartridge at different stages of the EVs purification protocol. Operational parameters for each step are detailed in Table 1. (a) Fractionation of the blood sample by centrifugation. (b) The plasma fraction is transferred to the cross-flow filtration chamber. The liquid flow through the membrane is driven by negative pressure and the centrifugal force, both generated by the PowerBlade instrument. (c) Air bubbles are injected *via* a channel at the bottom of the incubation chamber to induce chaotic mixing of the filtered plasma with the MNPs. (d) An aliquot of wash buffer is used to fully wet the surface of the micropillar array. (e) The plasma/MNPs mixture is flowed through the pillars for instantaneous flow-through magnetic capture. The MNP-free plasma is then transferred to the waste chamber (f) multiple aliquots of wash buffer are flowed back and forth through the *M-Chip* and discarded into the waste chamber. (g) The lysis buffer is transferred to the source chamber and flowed back and forth through the *M-Chip* for incubation with the EVs bound to the captured MNPs. (h) The EVs lysate is transferred to the processed sample chamber for manual collection and off-chip downstream analyses.

would otherwise disrupt the flow through pillars and impact capture efficiency (step 4, Fig. 4d). The filtered plasma/MNPs mixture is transferred to the source chamber by applying positive pressure to port #2. As the suspended MNPs transit the primed *M-Chip*, the entire population is trapped on the Ni/Au coated pillars, leaving the plasma in the sink chamber depleted of target EVs. The depleted plasma is then itself discarded into the waste collection chamber by applying negative pressure to port #8 (step 5, Fig. 4e). After capture, multiple, sequential aliquots of wash buffer are transferred to the source chamber by applying positive pressure to port #5 and then subsequently transferred through the *M-Chip* to remove most non-vesicular contaminants (step 6, Fig. 4f). By oscillating ports #6 to #8 between positive and negative pressure, buffer is shuttled back and forth through the *M-Chip* to increase overall washing efficiency. Each wash buffer aliquot entering the sink chamber is then also discarded into the waste chamber. Next, the lysis buffer is transferred to the source chamber by applying positive pressure to port #4. Over the course of 5 min, the lysis buffer is cycled back and forth through the *M-Chip* to lyse the captured EVs (step 7, Fig. 4g) without needing to release the MNP from the microstructure pillars. Finally, the lysate collected in the sink chamber is

transferred to the lysate collection chamber by applying negative pressure to port #6 (step 8, Fig. 4h). The EVs lysate can then be manually extracted from the device for downstream analyses. The *EV-Blade* cartridge enables automated purification of EVs sub-populations and lysate recovery in 60 min.

Exosome purification efficiency

We compared the exosome purification performance of the *EV-Blade* against SEC and a manual size- and affinity-based exosomes purification protocol that combines centrifugation-based blood fractionation, 0.22 μm filtration using a centrifugal filtering column and immunomagnetic capture in a tube. Freshly collected whole blood samples were pooled and divided into 1 mL aliquots. We processed 3 blood samples with each method (SEC, manual size- and affinity-based purification and *EV-Blade*), and measured the protein concentration of known exosomes-enriched transmembrane proteins (CD63, CD81, CD9 and Synt-1) in the recovered lysates. Both methods yielded almost identically pure exosomes populations, with negligible cellular contamination, as indicated by the absence of Cyt-C (Fig. 5a).



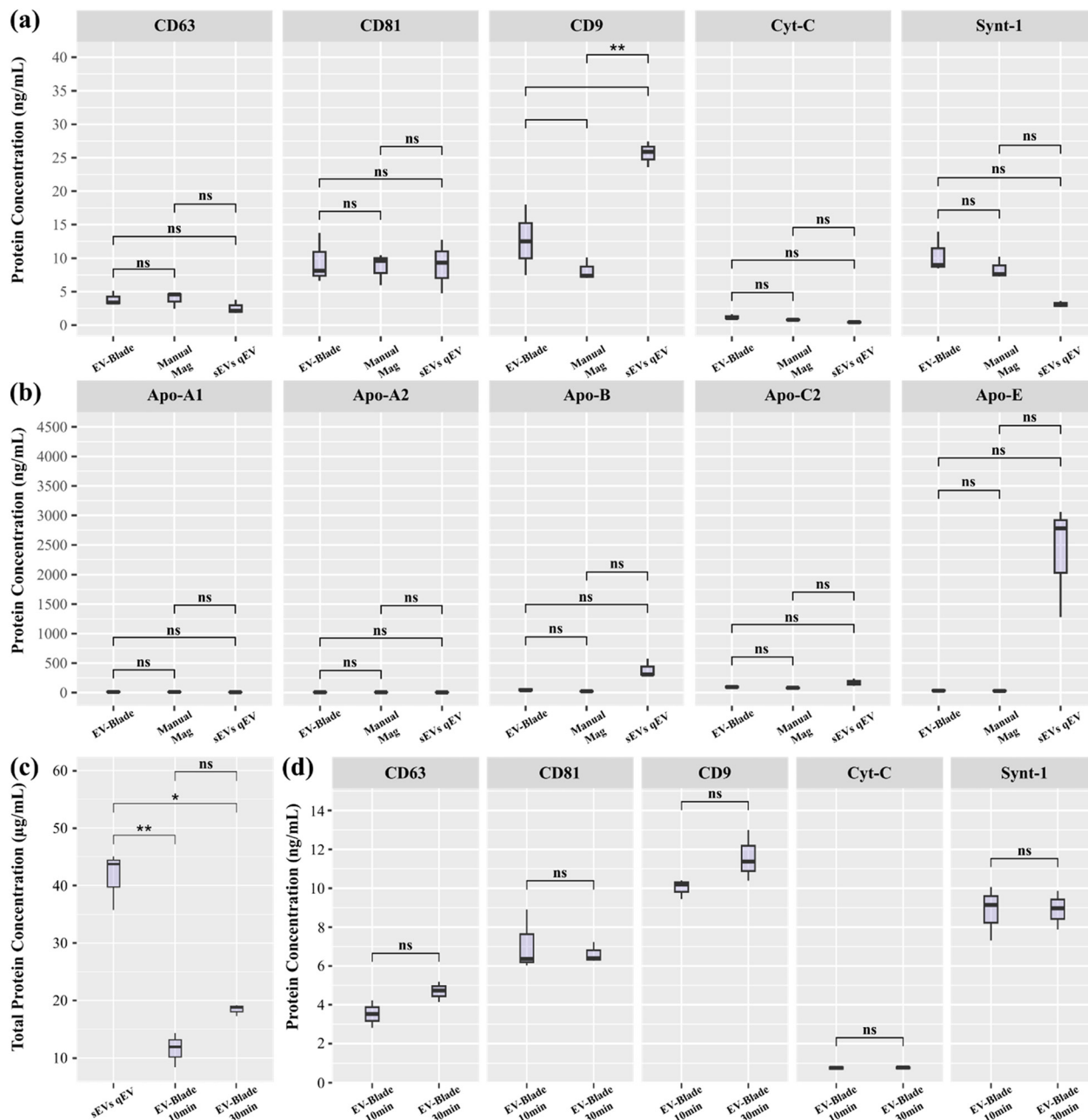


Fig. 5 Characterization of exosomes purified using the *EV-Blade* microfluidic device. Boxplots of (a) the protein concentrations for known exosomal transmembrane proteins (CD63, CD81, CD9 and syntenin-1) and the intra-cellular organelle marker cytochrome-C measured with the ProcartaPlex exosomes characterisation immunoassay comparing *EV-Blade* purified exosomes, manual immunomagnetic-based purified exosomes and SEC purified sEVs; $**p$ -value < 0.01, (b) lipoproteins concentrations (Apo-A1, Apo-A2, Apo-B, Apo-C2 and Apo-E) measured with the ProcartaPlex human apolipoprotein panel immunoassay comparing *EV-Blade* purified exosomes, manual immunomagnetic-based purified exosomes and SEC purified sEVs, (c) total protein concentrations measured with the microBCA protein assay for SEC purified sEVs, *EV-Blade* purified exosomes with a 10 min incubation step and *EV-Blade* purified exosomes with a 30 min incubation step; $*p$ -value < 0.05; $**p$ -value < 0.01, (d) protein concentrations for known exosomal transmembrane proteins (CD63, CD81, CD9 and Syntenin-1) and the intra-cellular organelle marker cytochrome-C measured with the ProcartaPlex exosomes characterisation immunoassay for *EV-Blade* purified exosomes with a 10 min incubation step and *EV-Blade* purified exosomes with a 30 min incubation step.

Additionally, preparations from both size- and affinity-based methods (Manual Mag and *EV-Blade*) were free from 5 lipoproteins (Apo-A1, Apo-A2, Apo-B, Apo-C2 and Apo-E) whereas SEC preparations (sEVs qEV) contained traces of Apo-B and a significant amount of Apo-E, indicating

contamination by high-density lipoprotein (HDL) particles in SEC purified EVs (Fig. 5b). The total protein yield was higher for SEC purified EVs compared to *EV-Blade* purified exosomes (Fig. 5c), which can be attributed to HDL particle contamination in SEC samples and the higher specificity of



the *EV-Blade* capture protocol targeting a subset of blood derived sEVs enriched in CD63, CD81 and CD9.

Increasing the immuno-capture incubation time in the *EV-Blade* method resulted in more total protein (Fig. 5c) but did not significantly alter exosomes-specific protein levels (Fig. 5d), suggesting 10 min incubation is sufficient. Extending incubation does not further increase exosome yield and may reduce purity through non specific binding. Reference values measured for exosomal transmembrane proteins and lipoproteins in filtered plasma samples are shown in Fig. S2. Combined with the PSD shown in Fig. 2c, these results confirm that the *EV-Blade* cartridge effectively purifies exosomes from whole blood. Processed samples are free from size-similar intra-cellular organelle contaminants and yield results comparable to existing manual protocols, with expected differences in purity when comparing size selection only (sEVs qEV) versus size- and affinity-based purification methods (Manual Mag and *EV-Blade*).

Analysis of exosomes derived circulating RNA

We evaluated the *EV-Blade* cartridge as a companion tool for exosomal RNA-based cancer liquid biopsy. In previous work, we analyzed long circulating RNAs found in extracellular

vesicles purified from the peripheral blood of childhood B-ALL patients using total RNA sequencing.²² Given the very limited availability of plasma from childhood B-ALL patients at diagnosis (900 μ L per patient), most extracted RNA was used for sequencing and RT-qPCR validation. To simulate similar conditions for the purpose of validating our technology, we started with a pooled sample of whole blood collected from healthy donors and purified exosomes from three 1 mL aliquots under the three following conditions: (i) healthy donor whole blood, (ii) healthy donor whole blood spiked with sEVs manually purified from REH cell CCM and (iii) healthy donor whole blood spiked with sEVs manually purified from 697 cell CCM. The concentration of spiked sEVs was adjusted to align with C_q values of selected potential biomarker candidates we previously measured in patient samples *via* RT-qPCR.²² These conditions represent healthy individuals, one subtype of low-risk B-ALL (*ETV6::RUNX1* positive) patients and one subtype of high-risk B-ALL (*TCF3::PBX1* positive) patients forming a model for liquid biopsy application in cancer patient risk stratification. Fig. 6a shows RNA yields from *EV-Blade* purified samples, ranging from 3.32 to 5.74 ng. RNA concentrations in spiked plasma samples were similar or slightly lower than those in whole blood samples. The

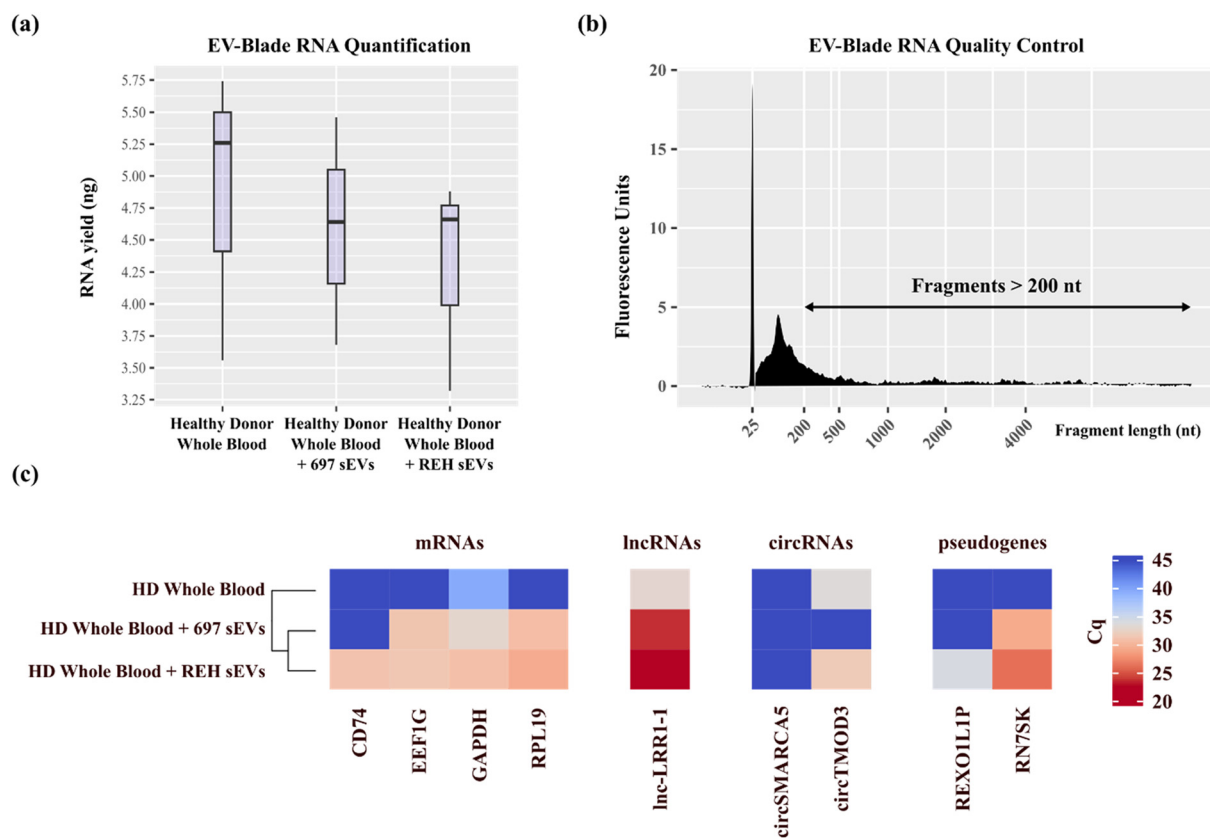


Fig. 6 Exosomes based screening of circulating RNAs. (a) Boxplot showing RNA yields extracted from exosomes purified with the *EV-Blade* device from healthy donors whole blood and whole blood spiked with concentrated sEVs from two different acute lymphoblastic leukemia cellular models (697 and REH). (b) Electropherogram showing the size distribution of RNA fragments in a typical RNA sample extracted from exosomes purified with the *EV-Blade* device from whole-blood. Fragments >200 nt are highlighted. (c) Expression heatmap of selected B-ALL associated genes measured by RT-qPCR in RNA extracted from *EV-Blade* purified exosomes.



collected lysates exhibited a broad distribution of RNA fragment sizes, with the majority measuring <200 nucleotides and larger fragments present at lower abundance, as shown in Fig. 6b. On average, 30% of the total fragments exceeded 200 nucleotides (DV200), with RNA integrity number (RIN) values ranging from 1.7 to 2.4. Previous work by us and others^{22,58–61} have shown the utility of long coding and non-coding RNA transcripts including mRNAs, lncRNAs, circRNAs and pseudogenes biotypes for risk stratification of B-ALL patients. We previously identified potential RNA-based biomarkers in patients' blood-derived exosomes and cellular models that correlate with B-ALL molecular subtypes associated with high- and low-risk conditions.²² We selected a panel of genes in these 4 biotypes and assessed their expression by RT-qPCR (Fig. 6c), revealing significant expression differences for 8 out of 9 genes across the three model conditions, indicating that our exosome-derived RNA profiles can distinguish between different B-ALL subtypes. The *EV-Blade* exosome purification results show that starting from a clinically relevant volume of whole peripheral blood, we were able to recover sufficient RNA to successfully screen a panel of potential B-ALL biomarkers correlating with high- and low-risk conditions in model samples. Further validation with primary patient samples will be critical to confirm these findings and assess the device's sensitivity, specificity, and robustness for routine clinical diagnostics.

Conclusion

This study presents the *EV-Blade*, the first integrated centrifugal-pneumatic microfluidic cartridge to automate size- and affinity-based exosome isolation using cyclic flow-through magnetic capture. This innovative integration enables a complex purification protocol to be fully automated at high flow rates, crucial for processing larger sample volumes (in the milliliter range) compared to previously reported methods that were limited to smaller blood volumes. Automating this process is crucial for clinical adoption as it simplifies and reduces the costs associated with the otherwise complex purification workflow. Handling larger volumes is essential for detecting lowly expressed biomarkers, thereby enhancing disease detection capabilities during the early stages from minimally invasive biofluid collections, such as peripheral blood draws, rather than invasive tissue biopsies.

The *EV-Blade* efficiently isolates specific subsets of sEVs, particularly exosomes, from whole blood in 60 min. Its robust design supports scalable fabrication and consistent capture performance despite manufacturing variations, making it a cost-effective and clinically viable solution for exosome purification. The device removes blood cells, larger EVs, and cellular debris while retaining sEVs within the 60 to 120 nm size range in high quantities. On-chip immunomagnetic capture uses functionalized magnetic nanoparticles assembled in porous magnetic clouds for cyclic flow-through

incubation. This process further purifies exosomes by targeting well-established exosomal surface proteins, effectively eliminating size-similar intracellular organelle contaminants and lipoproteins from recovered samples. We showcase the *EV-Blade's* capabilities in a clinically relevant workflow by processing model samples for liquid biopsies aimed at risk stratification in B-ALL patients. This demonstration highlights the *EV-Blade's* ability to efficiently recover both coding and non-coding RNA transcripts from blood exosomes, facilitating the screening of potential B-ALL biomarkers which can be used for the differentiation of B-ALL subtypes. These results underscore the practical applications of the *EV-Blade* in cancer research and diagnostics, offering a powerful tool for non-invasive biomarker discovery and disease monitoring.

In summary, the *EV-Blade* offers an automated and scalable method to isolate exosomes from whole peripheral blood, efficiently retaining sEVs in plasma through fractionation and cross-flow filtration and providing robust immunomagnetic capture of sEV subsets (*e.g.*, exosomes) for practical omics-based applications in cancer research. Given the simplicity of the device and its ability to fully automate the complex immunomagnetic separation protocol, which can be easily customized for targeting different circulating targets by simply changing the magnetic bead functionalization, this method holds great promise for advancing liquid biopsy-based approaches. It can be easily translated towards multiple applications in cancer research and ultimately in clinical practice, improving the diagnosis and monitoring of various diseases.

Conflicts of interest

The authors declare no conflicts of interest.

Data availability

The data supporting this article have been included in the main body of the article and as part of the supplementary information (SI).

Supplementary information is available. See DOI: <https://doi.org/10.1039/d5lc00977d>.

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