Solvent-selective routing for centrifugally automated solid-phase purification of RNA

Nikolay Dimov · Eoin Clancy · Jennifer Gaughran · David Boyle · Darren Mc Auley · Macdara T. Glynn · Róisín M. Dwyer · Helena Coughlan · Thomas Barry · Louise M. Barrett · Terry J. Smith · Jens Ducrée

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Abstract We present a disc-based module for rotationally controlled solid-phase purification of RNA from cell lysate. To this end, multi-stage routing of a sequence of aqueous and organic liquids into designated waste and elution reservoirs is implemented by a network of strategically placed, solvent-selective composite valves. Using a bead-based stationary phase at the entrance of the router, we show that total RNA is purified with high integrity from cultured MCF7 and T47D cell lines, human leucocytes and Haemophilus influenzae cell lysates. Furthermore, we demonstrate the broad applicability of the device through the in vitro amplification of RNA purified on-disc using RT-PCR and NASBA. Our novel router will be at the pivot of a forthcoming, fully integrated and automated sample preparation system for RNA-based analysis.

Keywords Lab-on-a-disc · RNA purification · Solvent-selective valves · Molecular diagnostics

1 Introduction

Microfluidic lab-on-a-chip technologies have been shown to automate and reduce the time-to-result of a wealth of bioanalytical assays, and ultimately enable their deployment at the point of need. Whilst a range of detection techniques remains implemented on-chip, the integration of sample preparation with detection has been a bottleneck in the development of microfluidic devices for nucleic acid testing (Foudeh et al. 2012; McCalla and Tripathi 2011). In many instances, RNA is a preferable diagnostics target, such as the detection of retroviruses or in the expression analysis of genes. Also, in assays for the detection of pathogens, RNA can be used to obtain information on pathogen viability as RNA is less stable than the DNA. Furthermore, since RNA molecules are typically present in multi-copy (100–1,000 s/per cell), the potential for detection without in vitro enzymatic amplification is possible. One such class of RNA species that have recently received attention for their diagnostic potential is micro-RNAs (miRNAs). miRNAs are small non-coding RNAs that are involved in modulating gene expression at the transcriptional and post-transcriptional level. Their dysregulation has been shown to be associated with a wide variety of human diseases, including cancer (Mitchell et al. 2008; Heneghan et al. 2010), diabetes and cardiovascular diseases. In terms of bacterial RNA diagnostics, ribosomal RNA (rRNA) and in particular 16S...
rRNA remains the gold standard. Other functional high-
copy number bacterial RNA molecules such as transfer
messenger RNA (tmRNA) encoded by the ssrA gene has
been demonstrated to be a useful marker in bacterial diag-
nostics (O’Grady et al. 2009; Clancy et al. 2012).

The automation of microfluidic platforms often involves
the use of costly peripheral equipment, such as syringe
pumps, external valves and pressure controllers. These
chip in a laboratory devices require complex, multi-stage
off-chip liquid handling steps, thereby severely limiting
their widespread adoption in clinical testing. Centrifugal
microfluidic Lab-on-a-Disc (LoaD) systems bear the poten-
tial to integrate sample preparation with detection to create
full-fledged sample-to-answer devices (Madou et al. 2006;
Ducrée et al. 2007). A simple rotary microsystem devel-
oped by Park et al. (2012a) enabled the purification of RNA
from viral lysate with frequency-controlled release of rea-
gents. Using centrifugal microfluidics, Cho et al. presented
a device capable of one-step DNA extraction of pathogen-
specific DNA from whole blood (Cho et al. 2007). Also
multiplexing of immunoassays on-disc (Lee et al. 2009;
Park et al. 2012) and parallelized biochemical analysis (Lee
et al. 2011) have been demonstrated.

As all liquids resident on the disc are simultaneously
exposed to the same centrifugal field, valving technolo-
gies are pivotal for establishing a sequence of liquid han-
dling steps. Over the years, the scientific community has
pioneered a repertoire of valving schemes, which are commensurate with the rotational nature of the lab-on-a-
disc platform. Amongst the central aspects governing the
choice of the valving scheme are the upper limit of toler-
able spin frequencies (e.g. during multi-stage sample
prep), the open-state hydrodynamic resistance and vapour
barrier properties (e.g. for storage and release of liquid
reagents) on behalf of flow control and the complexities
involved in fabrication and actuation on the hardware side.
Valve actuation on LoaD platforms can be categorized into
two schemes: the first type of valve is controlled by the
system-innate spindle motor such as capillary burst valves
(Ducrée et al. 2007; Zoval and Madou 2004) or siphons
(Schembri et al. 1995; Steigert et al. 2007; Nwankire et al.
2013); the alternative, externally actuated schemes often
involve the manipulation of a sacrificial material by an
external stimulus like heat, for instance, thermally actu-
ated wax valves (Park et al. 2007). Recent reports have
utilized such wax valves for the on-disc integration of
biochemical assays and immunoassays (Lee et al. 2009,
2011). To avoid the manufacturing complexity of incor-
porating on-disc heat-induced valves, Mark et al. (2008)
introduced a pneumatic microvalve using a thin burstable
foil. Zehnle et al. (2012) swiftly balance the pressures on a
spinning disc and use the advancing liquid as a valve to
achieve inward pumping. In another study, Gorkin et al. III
(2012) integrated water-dissolvable films as once-off valve
seals. By trapping air between this dissolvable film and the
liquid, they implemented centrifugopneumatic valving
and gating of on-board-stored reagents for a wide range of
burst frequencies. We expand upon these dissolvable film
valves and develop automated, solvent-selective routing on
a LoaD platform.

For solid-phase purification (SPP) of nucleic acids, we
here exploit the well-established Boom chemistry (Boom
et al. 1990). Fluidic routing to direct flow to a selected out-
put at a bifurcation between a waste and an elution outlet
is critical for centrifugally implemented automation. Kim
et al. (2008) developed a flow switch by using a capillary
valve upstream of an open chamber and a unique 3D junc-
tion geometry. A similar router, solely controlled by the
rotationally actuated hydrodynamic Coriolis pseudo-force,
was reported by Brenner et al. (2005). This virtual routing
concept was further sophisticated by Haeberle et al. (2007)
who successfully extracted DNA from calf thymus using
silica beads by alternating the sense of rotation. Based on a
droplet deflection Coriolis-force driven router, they recov-
ered 16 % of the initial nucleic acid. A recent study by
Jung reports 81 % capture efficiency of RNA from lysed
influenza A H3N2 virus using silane, i.e. TEOS-treated
glass beads on-disc (Jung et al. 2013). Alternatively, 100 %
efficient automated extraction of human genomic
DNA is demonstrated by Kloke et al. (2014) who imple-
ment novel ball-pen pierceable seals to route the sample
lysate through an integrated silica membrane in a Lab
Tube platform. Whilst these systems have contributed
greatly to the development of SPP, to date, reliable, high-
efficiency and low-complexity routing of flows (e.g. aque-
ous or organic flows) on LoaD platforms still remains a
challenge.

We report a routing scheme which utilizes solvent-
selective valving with unidirectional rotational actuation
for SPP of RNA, thereby obviating the need for external
actuation other than the system-inherent spinning rates.
Previously, Kinahan et al. (2014) successfully achieved
solid-phase purification of total RNA from MCF7 cell
lysates. Multiple dissolvable film valving strategies were
presented, which allowed inbuilt control of reagent release
for automated sample preparation and integration of labor-
atory unit operations on LoaD at constant angular veloc-
ity. However, the article briefly touches on the quality of
the purified samples and the mechanisms of routing. In
contrast, here we investigate solid-phase purification of
RNA from various cells, study the suitability of the method
for further downstream amplifications, and expand on the
selective routing through integrated hydrophobic mem-
brane and dissolvable film valves. Our findings will be at
the core of a future, fully automated centrifugal microflu-
idic nucleic acid analyser.
2 Principle of operation

This paper addresses the missing link between upstream, chemically induced cell lysis and 3-phase RNA extraction and downstream nucleic acid analysis techniques which are both well-established bench-top methods that have also been demonstrated microfluidically using lab-on-a-chip/lab-on-a-disc systems (Linares et al. 2011). The single-step acid guanidinium thiocyanate-phenol-chloroform method, also known as 3-phase RNA extraction, is based on the discovery that RNA (other than DNA and proteins) remains soluble in the acidic aqueous phase (Brenner et al. 2005; Chomczynski and Sacchi 1987). Formulations of acidic guanidinium thiocyanate and phenol are commercially available under various brand names (TRIzol from LifeTechnologies, TRI Reagent from Sigma). In general, the reagent contains phenol that ruptures the cells and guanidinium thiocyanate, a chaotropic salt that strips protein complexes from RNA. The addition of chloroform (or an alternative reagent such as 4-bromoanisole or 1-bromo-3-chloropropane), prior to centrifugation, permits the separation of the non-polar (organic) and polar (aqueous) phases. Due to differences in their solubility, DNA is retained in the organic phase whilst the RNA is concentrated in the aqueous phase, which also contains salt contamination. For accurate downstream analysis of RNA, it is essential that extracted RNA is free of contaminants such as chaotropic salts and phenol (Bustin and Nolan 2004; Tan and Yiap 2009).

Here, we convey the centrifugal microfluidic automation of a 4-stage purification of RNA from the aqueous phase of TRI Reagent-lysed human and bacterial cells. Using human MCF7 breast cancer cells, we validate the device. Subsequently, we demonstrate the applicability of the device for nucleic acid diagnostics with the in vitro enzymatic amplification of miRNAs from cultured T47D cells and human leucocytes using RT-PCR. Furthermore, using nucleic acid based amplification (NASBA) we demonstrate the utility of the device for bacterial molecular diagnostics using H. influenzae as a model organism.

First, we demonstrate the underlying principle of solvent-selective routing (Fig. 1) by strategically placing two solvent-selective valve types (Fig. 1a, b) in a simple microfluidic network, inverse-Y bifurcation connecting an inner loading chamber to two outlets (Fig. 1c, d). Within the three-dimensional (3D) disc architecture similar to an earlier reported system (Gorkin et al. III 2012), we placed the two outgoing channels in separate layers. These channels
communicate via a vertical through hole which is initially sealed by a solvent-selective film, in this case a hydrophobic, PTFE-supported membrane valve (HMV) (Fig. 1a, c) which becomes permeable upon exposure to organic solvents. As the HMV is impermeable to aqueous solutions, the first aqueous flow is routed through a narrow channel to the right collection chamber (Fig. 1c, 1–2). However, the HMV is permeable for the subsequent organic solution (Fig. 1c, 3–4), which has thus two options for exiting. The outflow into the left reservoir can be biased by a lower flow resistance, e.g. through a shorter connection channel with a larger cross section compared to the other outlet.

In addition, the centrifugally stabilized liquid volume already resident in the right outlet blocks all vents, thus acting akin to a solid plug to practically create a dead-end channel. The system can be simplified because both liquids, e.g. aqueous solutions, have identical contact angles with the channel walls and the channel has uniform cross section on both sides of the air pocket. For the sake of clarity, we hence neglect capillary and inertial effects, so the flow into the right outlet will hence stop once the pressure in the interspersed air pocket balances the centrifugal pressure exerted by the incoming liquid.

The air pocket thus acts as a fluidic capacitance (Kim et al. 2008) where the final pressure of the air pocket \( P_{\text{ini}} \cdot (V_{\text{ini}}/V) \) depends on the initial (typically atmospheric) pressure \( P_{\text{ini}} \) and the ratio of the initial and final volumes \( V_{\text{ini}} / V \), respectively (Fig. 1d). The flow in the lateral channel stops once the pressure in the air pocket equilibrates the centrifugally induced pressure \( \rho \omega^2 \bar{r} \Delta r \) with the liquid density \( \rho \), the angular frequency \( \omega \), the radial length \( \bar{r} \) and the mean position \( \Delta r \) of the incoming liquid plug. So essentially, the here considered routing function is tightly linked to the sequence of liquids and sealing the air pocket formed between the advancing and the stationary liquids.

We integrate a bench-top method starting with (1) loading the aqueous lysate, (2) isopropanol (IPA)-mediated RNA precipitation from the aqueous phase, followed by (3) washing with ethanol (EtOH) and eventual (4) resuspension of the purified RNA in aqueous buffer. A glass bead solid-phase support is utilized to enhance the purification efficiency of the system. To implement the automated routing of two organic solvents between a first and last aqueous phases, we established a sequence of a HMV and a selective router is based on the following protocol: first, the centrifugal microfluidic SPP with this solvent-adelated, 100-µm-wide radial grooves (Fig. 2a) to geometrically retain the glass beads (≤106 µm) under the impact of the centrifugal flow. With its microporous (pore size 0.45 µm) PTFE barrier, the first HMV blocks the aqueous sample and eluate (Fig. 2b, e) while providing passage of the organic solvents (Fig. 2c, d). These organic phases are both directed to their designated waste (\( W_{\text{org}} \)), which is the hydrodynamically preferred, axial outlet of the open-channel situation on the centrifugal platform.

The hydrophilic siphon, downstream from the loading chamber, lets the aqueous fraction pass at reduced spin rates, while it holds back subsequent IPA and EtOH before the crest point at increased rates until they are fully diverted into \( W_{\text{org}} \). Organic solutions are therefore effectively restricted to the loading chamber, solid phase and ascending siphon arm. The siphon thus minimizes the risk of critical cross-contamination of the elution reservoir \( E_{\text{aq}} \) with organic solvents. Following this intermediate, organic routing phase, we intersperse a drying period for the previously primed channels and membranes, which could be enhanced by centrifugation (Garcia-Cordero et al. 2010). Again the most crucial factor for a subsequent molecular amplification step is that \( E_{\text{aq}} \) remains free of potential contaminants from preceding solutions.

In the final step, the aqueous elution buffer desorbs the RNA bound to the solid phase. After passing the hydrophilic siphon during slow spinning, the RNA-depleted sample (Fig. 2b) and the elution buffer (Fig. 2e, f) are diverted in a binary fashion to the aqueous waste (\( W_{\text{aq}} \)) and the elution chamber (\( E_{\text{aq}} \)), respectively. To this end, a normally closed (water) dissolvable film valve (DFV) initially seals the radial outlet to route the liquid into \( W_{\text{aq}} \). Once in contact with the valve, the first liquid initiates the timed dissolution of the thin film so the DFV is opened to direct the subsequent elution buffer to its destination \( E_{\text{aq}} \). Additionally, the hydrophobic membrane (HM) covering the outlet of the aqueous waste (\( W_{\text{aq}} \)), denoted in Fig. 2b, prevents leakage while venting the chamber. Thus, the HM retains the RNA-depleted fraction and maintains the air pocket with the advancing aqueous elution buffer. Contact between the two liquids is averted because of the fluid capacitance, i.e. pressure equilibrium on the two sides of the air pocket (Fig. 2e, f). In total, the router utilizes four phase-selective valves (HMV, hydrophilic siphon, DFV and HM) with different functionalities that allow automated sample purification without exposure of the eluted RNA to preceding flows.

The centrifugal microfluidic SPP with this solvent-selective router is based on the following protocol: first, the crude aqueous extract from a homogenized biological sample is introduced onto the beads inside the loading chamber (Fig. 2b). The RNA from the solution binds to the acid-washed glass beads as a result of charge-charge interactions under chaotropic conditions (Boom et al. 1990).
Using a specific spin frequency, the RNA-depleted fraction is delivered through the siphon into $W_{aq}$. The flow disintegrates the thin film so the DFV is open to route the subsequent elution buffer into $E_{aq}$. Next, the disc is stopped and IPA is pipetted into the loading chamber ($L$) to precipitate any remaining unbound RNA (Fig. 2c). A small volume of the RNA-depleted aqueous solution remains trapped in the channel prior to the siphon crest. Both, its flow resistance and the wettability of the PTFE-supported membrane make the HMV impermeable for water, even at high rotational frequencies (75 Hz). As long as the pressure drop across the membrane is lower than the minimum pressure required to drive all of the permeate phase through the membrane (Adamo et al. 2013), the HMV remains closed. Once the IPA is introduced onto disc and is in contact with the HMV pores, it passes through the PTFE-supported membrane. According to the supplier, organic solvents, IPA in this case, changes the membrane permeability and thus gates the RNA-depleted aqueous solution through the pores of the HMV into the organic waste. The high angular velocity (75 Hz) guarantees that IPA does not shoot over the crest of the siphon, which is a low-frequency pass valve. Such HMV permeability facilitates the emptying of the channel, prior to the siphon crest, into the $W_{org}$ and prevents transfer of the high-salt concentration solution to the elution chamber ($E_{aq}$). The lost volume does not influence the extraction efficiency as the RNA has been retained on the solid phase prior to the treatment with IPA. It takes approximately 3.5 min at spin frequency 75 Hz for the membrane to open as the RNA-depleted solution prevents the IPA from direct contact with the HMV. This is the time required for the IPA to reach the PTFE membrane through the channel at that given spin rate. Using a solvent with higher miscibility would accelerate the processing time. Therefore, the time required to open the HMV is solvent specific, and it also depends on the geometry of the upstream channel, spin rate and intrinsic properties (pore size, contact angle) of the integrated PTFE filter. EtOH is consecutively loaded that rinses away salts from the beads and the precipitated polynucleotides (Fig. 2d), and also residues from the pre-crest channel region. During the drying period, as the EtOH evaporates from the pores of the PTFE-supported membrane, the valve returns to its normally closed state, hence allowing the elution buffer to pass by the HMV undisturbed, at low rotational frequencies over the siphon crest into $E_{aq}$. Finally, 100 µL of aqueous elution buffer is introduced and driven at lower frequencies (7.5 Hz) through the solid phase where it retrieves the purified RNA, then pass
the HMV over the siphon crest and through the now open DFV and vertical channel into the $E_{aq}$ reservoir (Fig. 2e).

3 Materials and methods

The following materials were used for the study: acid-washed glass beads ($\leq 106 \mu m$), absolute ethanol, isopropanol, phosphate buffered saline (PBS), pressure-sensitive adhesive (PSA, Adhesive Research Inc., Ireland), polymethyl methacrylate (PMMA) sheets (Evonik Industries AG, UK), dissolvable film (FA36, Harke PackPro, Germany), hydrophobic membrane (PTFE membrane circles, 0.45-µm pore diameter, Whatman), DI water (TKA, Germany), nuclease-free water (VWR, UK), RNAse ZAP (Biosciences, Ireland), TRI Reagent, 4-bromoanisole (Bio-Science). All reagents listed above were obtained from Sigma-Aldrich unless otherwise stated.

3.1 Valve assembly

These valves were assembled into a tab structure using two pieces of PSA (Fig. 1a, b). First, the flow-through region in a tab was cut out from the PSA using a precision knife cutter (Graphtec, Yokohama, Japan). The protective foil was peeled off the PSA and either a hydrophobic membrane (HM) or a water-dissolvable film (DF) was stacked on the surface (Fig. 1a, b). The excess material was cut off and removed without penetrating the DF layer. A final cut through the PSA defined the tab size. To improve the mechanical stability of the fluid exposed area, a second PSA layer covered the HM/DF at the top.

3.2 Fabrication and assembly of the disc

The device consisted of three PMMA discs ($\phi 120 \text{ mm} \times \phi 15 \text{ mm} \times 1.5 \text{ mm}$) and two binding PSA layers (Fig. 3). Each disc was cut to size and processed on a CO$_2$ laser (Epilog Zing, US): the bottom disc had its draining channels ablated; in the middle disc, liquid loading channels and collection chambers were cut out. Additionally in the middle disc connecting channels, siphons and valve grooves were CNC milled (MDX-40 Rolland, UK) into the backside. Inlets and outlet via-wholes were ablated in the top disc. All three discs were sonicated in 2 % aqueous solution Micro 90 (International Products Corp., USA) for 30 min at 50 °C. The discs were then transferred to DI water and sonicated under the same conditions for another 30 min. Finally, the discs were heat-dried at 80 °C for 45 min.

Prior to disc assembly, the work area on the bench was decontaminated using RNase ZAP and cleaned with 70 % IPA. Valves were placed in their designated grooves in the middle disc. Intermittent layers of PSA with the cut out silhouettes (Graphtec, Yokohama, Japan) of all chambers and channels were aligned with their contour parts on the PMMA discs using a bespoke assembly rig. The assembled device was passed multiple times through a roller press (Hot Roll Laminator, Chemsultant Int., USA) to reinforce the bonds of the five-layer structure. Finally, 30 mg of acid-washed, dried glass beads was introduced to the loading chamber $L$.

3.3 MCF7 cell culture

All cell culture reagents were obtained from Sigma-Aldrich (MO, USA) unless otherwise stated. MCF7 cells (DSMZ, Braunschweig, Germany) were cultured in 75 cm$^2$ flasks in DMEM media, supplemented with 10 % foetal bovine serum (FBS), 100 U mL$^{-1}$ penicillin and 100 µg mL$^{-1}$ streptomycin. The cultures were maintained at 37 °C with 5 % CO$_2$. Cells were harvested by incubation in 5 mL 0.25 % trypsin/0.1 % EDTA at 37 °C for 5 min followed by neutralization with 5 mL culture medium. Cells were centrifuged at 300×g for 4 min and resuspended in culture medium. The cells used in this study were collected between their 17 and 25 passages.
3.4 T47D cell culture

The ductal breast cancer (T47D) cells were cultured in RPMI-1640 supplemented with 0.2 U mL⁻¹ bovine insulin and FBS to a final concentration of 10 % at 37 °C in 5 % CO₂. Cells were harvested by incubation in trypsin–EDTA solution at 37 °C for 5 min followed by neutralization with 5 mL culture medium. Cells were counted and centrifuged at 300×g for 4 min to pellet the cells.

3.5 Haemophilus influenzae growth conditions

H. influenzae (DSMZ 4690) was cultured overnight in haemophilus test medium (HTM, Oxoid, UK). The following morning, a new culture (10 mL) was inoculated with a 100 µL overnight culture aliquot and allowed to grow to exponential phase (4 hrs). Cells were harvested, and the optical density of the culture was measured at 600 nm and compared to a previously generated growth curve. Cells were diluted in HTM to 1 × 10⁷ cells/mL and pelleted.

3.6 Leucocyte preparation

Human whole blood was obtained from a healthy male volunteer. The leucocyte fraction was isolated by gradient density centrifugation using Ficoll Paque Plus (GE Life Sciences) according to the manufacturers instructions. The cells were washed twice in 1× PBS and pelleted.

3.7 Bench-top cell lysis and RNA extraction with TRI Reagent

Cells were lysed by mixing a 25 µL aliquot of harvested cells in culture media (DMEM, FBS, penicillin and streptomycin) with 80 µL TRI Reagent. The aliquots contained 2.96 × 10⁵–7.4 × 10⁴ MCF7 cells. The cells were enumerated by an automated cell counter (Scepter Handheld, Milipore, USA) and in a haematocrit chamber (depth 0.1 mm, Marienfeld, Germany). For all other cell types (T47D, H. influenzae and leucocytes), the cell pellet was resuspended in a mixture containing 80 µL TRI Reagent and 25 µL nuclease-free water. To each sample, 5 µL of 4-bromoanisole was added. The mixture was then vortexed for 1 min and incubated for 9 min at room temperature prior to purification. All samples were then centrifuged at 14,000×g for 5 min at room temperature, and the aqueous phase was removed for on-disc purification.

3.8 Measuring the contact angle between PMMA and aqueous cell homogenate

Following lysis, 60 µL of the aqueous phase was collected, and an equal volume of water was added. From this, a 1 µL aliquot was placed on a clean, dry PMMA surface (n = 5) and images of the sitting droplets were acquired. The contact angles were calculated using the tangential method from the built-in software of the goniometer (DataPhysics Instruments GmbH, Germany). The hydrophilic siphon design (Online Resource 2) was calculated based on these data.

3.9 Bead-based total RNA purification on-disc

The use of silica-based substrates in combination with chaotropic salts to purify nucleic acids has been widely implemented (Boom et al. 1990). Under basic and near neutral pH, the silanol groups on glass or silica surfaces are negatively charged (Wen et al. 2008) due to their pKa of 5–7. In spite of the pKa, charge–charge interactions can take place in high-salt concentration solution. Also, strong ionic conditions (NaCl) and lower pH (pH = 5) can mask native hydroxyl negative charges and permit binding of the RNA. In the protocol we implement here, the chaotropic salt from the TRI reagent compensates the anionic charges by removing water molecules from both glass and nucleotides. To strip the beads of the bound RNA, a low ionic strength buffer at near neutral pH 8.1 is used.

On disc, 30 µg of dry acid-washed glass beads were introduced into the main chamber. Next, 120 µL of cell lysate mixed with water (1:1, aqueous phase/water) was loaded. The mixture was incubated for 5 min on a stationary disc to enable RNA binding to the beads. The beads were then subjected to two sequential washing steps with 60 µL of 100 % IPA and 60 µL of 75 % ethanol aqueous solution (EtOH). Finally, the purified RNA was retrieved from the beads with the addition of 100 µL elution buffer (50 mM Tris-HCl, 1 × EDTA at pH 8.1). The liquids were automatically collected in designated chambers (Fig. 2 and ESM-1). The entire volume of the eluted fraction (100 µL) was recovered from the collection chamber (Eₐq) and analysed.

3.10 Measuring the concentration, purity and integrity of extracted RNA

The concentration and integrity of the purified total RNA was determined by capillary electrophoresis (RNA 6000 Pico kit, Bioanalyzer 2100, Agilent technologies, USA) according to the manufacturers’ instructions. The quality of the purified RNA was assessed by the RIN (RNA Integrity Number) provided by a built-in algorithm of the Agilent Expert Software. The RIN algorithm analyses the entire electrophoretic trace originating from an RNA sample. First, to determine whether RIN could be calculated, depending on important elements of the electrophogram such as the Pre region, 5S region, Fast region, Inter
region and Past region are evaluated. If a critical anomaly is detected in any of these regions, the RIN is not computed. Baseline correction and normalization are automatically applied to each electropherogram prior to the feature extraction. Then, based on Bayesian learning technique, the algorithm builds regression models using indicated peak positions, heights, areas, area ratios, S/N ratio, maximum and minimum values, and waviness of the electropherogram trace to assign a 1–10 score (Schroeder et al. 2006). RIN score of 1 means the sample is degraded, and 10 scores for completely intact RNA. According to the description of Agilent, the RIN algorithm is developed to utilize neural networks and adaptive learning in conjunction with a large database of eukaryote total RNA samples, and the RIN score is largely independent of the amount of RNA used and the origin of the sample.

3.11 RT-PCR amplification

The microRNAs, miR-16 and miR-195 were reverse transcribed using miRNA-specific stem loop primers (Life Technologies) and the TaqMan® MicroRNA reverse transcription kit (Life technologies); 1.33 µL of RT product was subsequently amplified by PCR in a 20 µL reaction using the TaqMan® universal PCR master mix II kit and miRNA-specific primers and probes (Life Technologies) on a LightCycler 480 thermocycler (Roche).

3.12 NASBA amplification

The tmRNA transcript in RNA purified from *H. influenzae* was amplified in a real-time NASBA reaction using the NucliSENS EasyQ basic kit version 2 (bioMérieux, Lyon, France) on a LightCycler 2.0 thermocycler (Roche) using the primers (5-3) P1; AATCTC- AATACGACTCACTATAGGGAGAAGGCTTCGATCCTCAAACGGT, P2; GCAGCTTAATAACCT and a molecular beacon 5’FAM-CCCGAGTTGGGGATAACGC-GGAGTCAACTCGG 3’DAB (MWG Eurofins, Germany). Each sample was amplified in a 20 µL reaction consisting of 10 µL of NASBA reagent-primer mix, 5 µL of RNA template and 5 µL of the enzyme mixture (avian myeloblastosis virus reverse transcriptase, RNase H, and T7 RNA polymerase).

4 Results and discussion

4.1 Fluidic analysis

Proper operation with liquid volumes, conditions and spin frequencies for each stage of the purification procedure was experimentally validated. The final spin frequency protocol for the purification is outlined in Fig. 4. For total RNA, a 5-min incubation period was necessary to enable the beads to bind nucleic acids present in the sample. Another 5-min incubation period was required to provide sufficient time for the RNA to elute from the beads into the aqueous elution buffer in the reverse process. These incubation times and elution conditions relate directly to the properties of the solid phase and may vary for different bead materials, e.g. silica (Wen et al. 2008; Duarte et al. 2010), polystyrene (Duarte et al. 2011) or chitosan (Kim et al. 2009b). One advantage of the router presented in this work is that the solid phase inside the loading chamber can be easily substituted, e.g. to reduce the incubation times and/or increase the extraction efficiencies.

It was determined that increasing the period of incubation prior to elution further increased the extraction efficiency. The 5-min drying period (Fig. 4) facilitated the evaporation of any remaining EtOH from the siphon. Dry- ing of the siphon prior to the final elution step stabilized the contact angle, thereby preventing elution buffer from seeping over the siphon crest, thus preventing the elution buffer from mixing with aqueous waste fraction.

4.2 On-disc RNA purification

To validate proper functioning of the on-disc SPP with the solvent-selective router, we performed lysis and phase separation on the bench of MCF7 cells with TRI Reagent, and subsequent on-disc RNA purification from the aqueous phase obtained from the cell lysate. The electropherograms obtained for different numbers of MCF7 cells in Fig. 5 show that high-quality RNA was recovered from the eluted fraction.

For the MCF7 RNA studied (n = 8), RIN values were in the range of 7.2–9.2. An average of 16.8 ng was recovered from the 2.96 × 10^5 cells, which was less than the 23.1
ng from $1.48 \times 10^5$ MCF7 cells. Evidently, the cell number and the amount of ribosomal RNA (18S and 28S) are not correlated as indicated by the peak height variations in Fig. 5. According to previous studies by Hiorns et al. (2004), random fluctuations of ribosomal RNA are not uncommon; in the authors’ opinion, fluctuations in RNA levels can also be due to low recovery after purification. However, the RIN is the criterion used here to evaluate solid-phase extraction and RNA quality on LoaD, which is in the scope of the current article. Based on qualitative analysis of the rRNA, we conclude that the purified RNA has preserved high integrity after on-disc purification with the solvent-specific router.

Further analysis indicates that the quantity of recovered small RNAs is proportional to the number of cells in a starting sample. From the integrated area of the peaks with migration time between 24 and 28 s, the small RNA fraction was quantified for four different cell concentrations. We focused on this region as many potential biomarkers have been identified as small RNAs (Iorio and Croce 2012; Kosaka et al. 2010). Figure 5 compares the purified small RNA concentrations to cell content. In spite of the significant lost of RNA, which is discussed in the following paragraphs, the results suggest that our LoaD platform is applicable for small RNA purification from acid guanidinium isothiocyanate phenol-lysed samples.

The purification efficiency was further investigated by comparing the RNA content from the purified ($E_{aq}$) to the unbound ($W_{aq}$) fractions. Our quantitative measurements show that the RNA concentration is higher in the purified fraction in comparison with the non-bound, waste fraction. A concentration of 180 pg $\mu$L$^{-1}$ was measured in the purified fraction (100 $\mu$L), against 37 pg $\mu$L$^{-1}$ (120 $\mu$L) in the waste fraction (Fig. 6). These were also compared with the control sample, containing culture media only, where no RNA was detected, demonstrating that the recovered RNA originated from the cells and not from the growth media.

Literature reports on-chip SPP efficiency for DNA of 42.5 % using static silica beads (Duarte et al. 2010). Higher extraction efficiencies of 63.9 % are achieved for DNA using dynamic methods with magnetically induced enhanced mixing (Duarte et al. 2011) and 71.0 % for RNA utilizing chitosan-coated beads on-chip (Hagan et al. 2009) and recently 81.0 % on-disc with (TEOS)-treated glass beads (Jung et al. 2013) to capture RNA from influenza (A H3N2) lysates with known concentrations. The Lab Tube system developed by Kloke et al. (2014) is capable of 100 % automated extraction of human genomic DNA using integrated silica membrane.

As part of our study, bench-top extraction was performed lysing $9.5 \times 10^4$ MCF7 cells, using IPA precipitation and two consecutive washes with EtOH. We measured total RNA of 32.3 ng from that sample. The RNA recovered from an identical sample after on-disc extraction resulted in 3.9 ng of total RNA. Assuming that the bench-top extraction was 100 % efficient, our purification efficiency was
12.1 % (weight) of extracted total RNA from an identical sample utilizing a purification protocol without beads. The percentage of purified RNA on-disc varied between samples with different cell counts. For the $1.48 \times 10^5$ cells, 43 % of the total RNA was retained on the beads, from which 7.2 % was recovered from the elution fraction. Variations in the packaging of the solid phase would inevitably lead to fluctuations of the amount of recovered total RNA. Even in a tightly packed monolith, diffusion of the RNA molecules would act as a limiting factor. According to the Einstein-Smoluchowski relation for one-dimensional diffusion ($x^2 = 2D \cdot t$) during the incubation time (5 min), an RNA molecule with diffusion constant of $10^{-7}$ cm$^2$/s, for prokaryotic (16S) ribosomal RNA (Tam et al. 1981), would only travel 7.7 µm. Introducing mixing (Duarte et al. 2011) in the L chamber of the LoaD system can increase the capture efficiency at the loading stage. Further investigation and optimization of the solid-phase extraction protocol should follow in order to further raise both the capture and elution efficiency of total RNA.

A closer look at the electropherogram illustrated in Fig. 6 reveals that the run-time of the sample from $W_{aq}$ is increased in relation to the typical migration times. We attribute this behaviour to residual salts in the $W_{aq}$ chamber (Copois et al. 2007).

4.3 Assessment of on-disc RNA purity

To further investigate the presence of contaminants in the eluted fraction, we sought to determine whether salts and/or other contaminants would inhibit enzymatic amplification of RNA species present in the $E_{aq}$. Nucleic acid modification enzymes and polymerases are inhibited by organic species such as phenol (Katcher and Schwartz 1994), ethanol (Huggett et al. 2008) and the presence of guanidinium salts. We therefore sought to determine whether on-disc-purified RNA was amplifiable by RT-PCR and NASBA. For RT-PCR, RNA was purified from the lysate of $1 \times 10^5$ T47D cells (human ductal breast epithelial tumour cell line) and the leucocyte fraction from 1 mL of human whole blood. The microRNAs, miR-195 and miR-16 were first reverse transcribed using a sequence specific primer and then amplified by real-time PCR. The comparison study between the breast cancer-related miR-195 and the housekeeping miR-16 after RT-PCR is plotted in Fig. 7.

In the RT-PCR, the number of cycles needed for the amplification-associated fluorescence to reach a specific threshold level of detection, called crossing point (Cp) value, is inversely correlated to the amount of nucleic acid found in the original sample (Valasek and Repa 2005). For the T47D cells, Cp values of 30.6 and 29.1 were obtained for miR-16 with on-disc and bench-top purified RNA, respectively. For miR-195, the Cp values were 27.4 and 25.3 for on-disc and bench-top purified RNA, respectively (Fig. 7a). This result indicates an approximately threefold reduction in the quantity of both miRNAs detected from disc-purified RNA compared to bench-top purified RNA. In the case of RNA purified from leucocytes, the difference was even greater. For miR-16, there is a 600-fold reduction in the quantity detected (9.2 cycles) and a ninefold reduction (3.1 cycles) for miR-195. Nonetheless, these results demonstrate that our LoaD is capable of purifying small RNAs that are amplifiable from two different biological sample types.

One of the main questions in this study was whether the system was capable of purifying RNA from different organism types (human cells and bacteria) that could be later used in various downstream analysis processes. To further address this question, $H. influenzae$ cells were
lysed, RNA purified on disc, and tested for quality using NASBA amplification. The isothermal nature of the reaction lends it to point of care diagnostics. Whilst the reaction is isothermal, it does require the simultaneous action of three enzymes to work. Figure 8 indicates that the amount of RNA purified on-disc and amplified is comparable to the bench-top purified RNA. Decrease of fluorescence signal from NTC (Fig. 8) is attributed to photobleaching of the molecular beacon, FAM/DABSYL as the fluorophore/quencher (Summerer and Marx 2002). It should also be noted that the tmRNA transcript is 360 bases in length, considerably longer than miRNAs.

Preferential retention of tmRNA on disc should correlate to the intrinsic properties of the system and/or the extraction protocol. Prior studies have noted that non-specific adsorption is minimal for RNA or DNA on PMMA (Zhao et al. 2012). Therefore, we believe that it is highly unlikely that non-specific adsorption is a factor in RNA retention within the system. The literature described (Balladur et al. 1997) adsorption as a three-step mechanism starting with (1) diffusion, in our case with convection, from the solution to the surface, (2) lateral diffusion on the surface involving rearrangement of adsorbed entities and (3) adsorption/desorption equilibrium of the molecules at the interface. The observed selectivity during on-disc SPP could be explained in part by the centrifugally induced flux, which forced the liquid through the bead bed in the L chamber. At this initial stage of the protocol, time is another factor. Though velocity and contact time contribute to the selective adsorption of tmRNA; in the authors opinion, selectivity was mainly governed by desorption. Our system, however, is limited by the frequencies and accelerations during rotation due to the region of operation of the siphon valve; at low frequency and acceleration, the liquid would pass over the crest (ESM-2, Sup. Fig. 3). For instance, constant angular velocity under similar experimental conditions resulted in higher recovery of total RNA from MCF7 cells (Kinahan et al. 2014). Other known factors are the solid phase and binding conditions, which can significantly influence the efficiency of extraction as demonstrated by Kloke et al. (2014), who used silica membrane, and binding from ethanol for 2 min, for ample capture of DNA. In the context of reversible binding (3), it would be more difficult for larger RNA species (18S and 28S), due to their larger hydrodynamic radii, to remain adsorbed on the solid surface under the shear stress resulting from the disc acceleration (Kim and Hee Jang 2004) during washing with IPA and EtOH. This finding has important implications for developing LoaD systems, which target size-specific purification of molecules by solid-phase purification (SPP).

Despite the limitations in capture and elution during the solid-phase purification of total RNA, sufficient quantities and quality of RNA for downstream analysis are retained and recovered utilizing the solvent-selective LoaD router. These findings further support the feasibility of the router for sample preparations from both prokaryotic and eukaryotic samples.

5 Conclusions and outlook

In summary, we have purified total RNA with high integrity from cell lysates through an automated, merely rotationally actuated flow control strategy involving a network of solvent-selective valves. Purifying mammalian and bacterial RNA, we demonstrated that the applicability of the platform did not depend on the source but on the extraction conditions. Under the studied conditions, the shear stress...
induced by acceleration resulted in an overall low recovery of total RNA. In the future, we seek to improve the total RNA extraction efficiency by using alternative solid-phase materials and by improved packing of the solid phase. Based on the developed extraction platform, the fully automated system is planned to integrate pre-storage and release of reagents (Oordt et al. 2013), an upstream chemical cell lysis and the first stage 3-phase liquid–liquid RNA extraction. A downstream microarray will facilitate a sample-to-answer, point-of-care molecular diagnostic for early diagnostics of breast cancer measuring the expression levels of circulating miRNA molecules, including miR-16.

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