Elastomeric microparticles for acoustic mediated bioseparations

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Abstract

Background: Acoustophoresis has been utilized successfully in applications including cell trapping, focusing, and purification. One current limitation of acoustophoresis for cell sorting is the reliance on the inherent physical properties of cells (e.g., compressibility, density) instead of selecting cells based upon biologically relevant surface-presenting antigens. Introducing an acoustophoretic cell sorting approach that allows biochemical specificity may overcome this limitation, thus advancing the value of acoustophoresis approaches for both the basic research and clinical fields.

Results: The results presented herein demonstrate the ability for negative acoustic contrast particles (NACPs) to specifically capture and transport positive acoustic contrast particles (PACPs) to the antinode of an ultrasound standing wave. Emulsification and post curing of pre-polymers, either polydimethylsiloxane (PDMS) or polyvinylmethylsiloxane (PVMS), within aqueous surfactant solution results in the formation of stable NACPs that focus onto pressure antinodes. We used either photochemical reactions with biotin-tetrafluorophenyl azide (biotin-TFPA) or end-functionalization of Pluronic F108 surfactant to biofunctionalize NACPs. These biotinylated NACPs bind specifically to streptavidin polystyrene microparticles (as cell surrogates) and transport them to the pressure antinode within an acoustofluidic chip.

Conclusion: To the best of our knowledge, this is the first demonstration of using NACPs as carriers for transport of PACPs in an ultrasound standing wave. By using different silicones (i.e., PDMS, PVMS) and curing chemistries, we demonstrate versatility of silicone materials for NACPs and advance the understanding of useful approaches for preparing NACPs. This bioseparation scheme holds potential for applications requiring rapid, continuous separations such as sorting and analysis of cells and biomolecules.

Keywords: Cell separation, Continuous cell sorting, Acoustofluidics, Particle synthesis, Ultrasound standing wave
within aqueous media, whereas particles with a negative $\phi$ (i.e., NACPs) transport to the acoustic pressure antinodes within aqueous media.

$$F = \left(\frac{\pi P_o^2 V^2 \rho_w^2}{2\lambda}\right) \cdot \phi(\beta, \rho) \cdot \sin(2kx)$$

$$\phi(\beta, \rho) = \frac{5\rho_p - 2\rho_w}{2\rho_p + \rho_w} \frac{\beta_p}{\beta_w}$$

The capacity to relocate PACPs to pressure nodes has been used in various approaches for focusing and separation of mammalian cells [5-11]. For example, the recently commercialized Attune® flow cytometer (Life Technologies) substitutes traditional hydrodynamic focusing with ultrasonic standing wave fields to focus cells into a single flowing stream prior to laser interrogation [5]. To increase the high-throughput capacity of flow cytometry, Piyasena et al. recently developed multi-node acoustic focusing and demonstrated up to 37 parallel flow streams [6]. Peterson et al. exploited the inherent contrast factor of constituents from whole blood to separate and sort positive contrast erythrocytes from negative contrast lipids within an acoustofluidic device [7,8]. Strategies for separating two particle populations with contrast factors of the same sign can exploit differences in the magnitude of the acoustic force [9,10]. In certain cases, the contrast factor can be adjusted by changing the density of the solution, as shown in a report separating polystyrene and PMMA microparticles by increasing the salt concentration of the media [11].

We seek to augment acoustophoretic particle sorting capabilities by introducing newly designed negative acoustic contrast particles (NACPs) with the capacity for specific biomolecular recognition and relocation of PACPs to antinodes of ultrasound standing waves. Since NACPs move in the direction opposite to the majority of mammalian cells, we hypothesized that biofunctional NACPs can capture and specifically transport targeted cells (or other PACPs) to the pressure antinodes, provided the total acoustic force of the NACPs is greater than the total acoustic force of the PACPs.

Other NACP carriers for the transport of streptavidin PACPs to pressure antinodes 

**Results and discussion**

**Silicone microparticles as biofunctional NACPs**

Silicone elastomers offer properties suitable for NACPs such as compressibility at mild temperature (e.g., Young’s modulus ~1MPa for typical PDMS formulations) [14]. Here, all NACPs were prepared by emulsifying silicone pre-polymers in aqueous surfactant solutions and subsequently curing to produce solid microparticles (Figure 2A). Because homogenization produces polydisperse particles, filtration or centrifugation was employed to narrow the breadth of particle size distributions. In one example,
filtration of NACPs with a 12 μm polycarbonate filter resulted in an average particle diameter of 6 ±3 μm (Additional file 1). Although a variety of surfactants enabled formation of silicone-in-water emulsions, the importance of surfactant type became evident when attempting to re-suspend cured NACPs in surfactant-free buffer, which often resulted in irreversible particle aggregation. Here, we found that the block copolymer surfactant, Pluronic® F108, stabilizes silicone microparticles likely due to the strong association of the hydrophobic polypropylene oxide block with silicone [15]. We further exploited this stable association by end-functionalizing Pluronic® F108 with biotin (Figure 2B,C). Biotin-Pluronic F108 enables use of the streptavidin protein as a linker between NACPs and any biotinylated analyte (e.g., cells labelled with biotinylated antibodies).

We also sought to evaluate the feasibility of direct modification of NACPs. Typically, surface modification of PDMS is accomplished by employing modification methods such as ultraviolet (UV)/ozone irradiation [16], UV graft polymerization [13], oxygen plasma treatment [17], and adsorption [18]. These modification approaches are usually performed on macroscopic silicone surfaces not held to the unique stringencies required to functionalize NACPS. For NACPs, conditions must be avoided that cause significant change in modulus or irreversible microparticle aggregation. For instance, modification of PDMS surfaces via plasma treatment results in the formation of brittle silica layers [19] which could affect the negative acoustic contrast property. Here, to evaluate direct, covalent modification of particles, we used PVMS which contains vinyl groups and can be functionalized chemically without forming a silica-like crust [16]. To first evaluate and compare chemical groups in both PDMS and PVMS, bulk samples were prepared and characterized using ATR-FTIR (Figure 3). PVMS material displays characteristic vinyl peaks at 958 cm⁻¹ (C=C twist, =CH₂ wagging), 1,408 cm⁻¹ (=CH₂ scissors), and 1,597 cm⁻¹ (C=C stretch). While vinyl groups are versatile for various chemical reactions (e.g., thiolene or metathesis coupling), our studies revealed that relatively simple photochemical reaction with biotin-TFPA results in biofunctionalization of PVMS particles (Figure 4A,B). Photoreacting biotin-TFPA with PVMS microparticles and subsequently adding fluorescent streptavidin resulted in significant differences in fluorescent signal between positive and negative samples (Additional file 2). For example, signal to background values (S/B) of fluorescent images of PVMS microparticles functionalized with biotin-TFPA and fluorescent streptavidin was 22 ± 2, whereas the negative control reaction without light irradiation was 9.0 ± 0.3, suggesting a biotinylation reaction of NACPs occurred. Notably, these studies...
cannot discern the exact location of biotinylation (e.g., vinyl groups or associated surfactant), as TFPA may react with C-H, N-H, or C=C groups [20]. Initial attempts at using biotin-TFPA to functionalize PDMS microparticles resulted in similar trends with a S/B values of 22 ± 3, supporting the non-specificity of biotin-TFPA. Overall, these studies demonstrate the utility of using biotin-TFPA for bio-functionalization of silicone microparticles.

Next, we sought to evaluate the acoustic responsiveness of these silicone microparticles. Our results show that microparticles prepared from either PVMS or PDMS function as NACPs within aqueous media (Figures 4 and 5). For example, a mixture of biotinylated PVMS NACPs and non-biotinylated polystyrene microparticles randomly distribute within an acoustofluidic channel in the absence of a standing wave field (Figure 4C). Upon application of an operating frequency of 2.98 MHz to generate an ultrasound standing wave within the microchannel (wavelength = 2 × channel width), polystyrene and PVMS microparticles separate (Figure 4D). Here, particle separation occurred in less than one second as determined under this experimental setup. Determining the precise rate of particle separation would require further measurements with a high speed camera to track trajectories of individual particles. Incompressible positive acoustic contrast polystyrene particles transport to the center of channel, corresponding to the pressure node, whereas compressible PVMS NACPs transport to the channel sidewalls, corresponding to the pressure antinodes. The capacity for both PDMS and PVMS to function as NACPs (Figures 4 and 5) illustrates the versatility of using silicone elastomers with different chemical compositions. Although only two silicone materials were tested here, we envision schemes to enhance the repertoire of available functional groups by employing a range of functional silicones that could be used for bioconjugation reactions. It is important to note is that the “PDMS” microparticles studied here (i.e., Sylgard 184, which is a blend of PDMS, silica, and resin fillers) exhibited negative acoustic contrast despite containing silica fillers.

![Figure 4](image)

**Figure 4 Acoustic response of silicone NACPs.** Brightfield image (A) and corresponding fluorescence image (B) of PVMS microparticles functionalized with biotin-TFPA and subsequently labelled with streptavidin Alexa Fluor® 488. The fluorescent image was acquired during a 250 ms exposure. The scale bars represent 50 μm. (C, D) Fluorescence images show a mixture of PVMS microparticles (red, functionalized with biotin-TFPA and streptavidin Alexa Fluor® 546) and polystyrene microparticles (green, non-biotinylated, Spherotech, 10–13 μm diameter) within a channel of an acoustofluidic device with (C) and without (D) activation of the PZT. Mixture contained a 1:7 ratio of polystyrene/PVMS microparticles. Images acquired in the absence of flow. Dashed lines are included to demarcate the channel boundaries.
NACPs as carriers for acoustic-mediated separations

The separation of silicone NACPs from polystyrene microparticles demonstrated in Figure 4 encouraged further investigations aimed at evaluating potential for the use of NACPs in cell separations. We hypothesized that NACP-PACP complexes within aqueous media will transport to pressure antinodes, provided the total radiation force from NACPs in the complex is greater than the total radiation force from PACPs in the complex. To this end, we employed polystyrene microparticles as surrogates for mammalian cells and investigated separation characteristics using NACPs prepared from PDMS. The brightfield image (Figure 2B) and accompanying fluorescent image (Figure 2C) show association between streptavidin coated polystyrene and PDMS microparticles functionalized with biotin-Pluronic F108. Notably, within the acoustofluidic device, the NACP-polystyrene microparticle complexes transport in unison to the pressure antinode (Figure 5B). This suggests the feasibility of a new bioseparation technique where transport of targeted PACPs (e.g., cells) will rely on specific, well-defined interactions with the NACPs. Figure 5 shows all PACP-NACP complexes transported to the antinode at the acoustofluidic wall (e.g., ~14 NACPs and ~12 PACPs in four separate complexes). However, additional studies are required to further understand the effects of parameters, such as particle ratios, flow rates, and applied voltages on efficiency of separation.

As expected, in the absence of fluid flow, NACPs accumulate at the pressure antinodes along the acoustofluidic channel walls during activation of the PZT (Figures 4 and 5). Secondary acoustic forces contribute to the aggregation of NACPs, as previously described for lipids in milk emulsions and whole blood [7,8,21]. This NAPC aggregation may be reduced by introducing flow to the channel. As recently demonstrated, laminar flow within the channel enables NACPs to maintain their position at the pressure antinode while simultaneously...
moving along laminar streamlines to the downstream trifurcation [12]. This capacity to couple relocation with downstream sample collection facilitates continuous sorting applications.

To the best of our knowledge, this is the first report documenting the use of NACPs as carriers for active transport of PACPs in acoustofluidic systems. Although polystyrene microparticles were used as cell surrogates in this preliminary investigation to demonstrate separation, this approach should be suitable for cell sorting based on binding of NACPs to specific cell surface antigens. Because the positive acoustic contrast factor value of cells is less than polystyrene beads [22], we anticipate that cell-NACP complexes should readily transport to pressure antinodes. Thus, this method holds potential as a complement to current cell sorting techniques (e.g., fluorescence-activated or magnetic-activated cell sorting). In contrast to these conventional methods, the present technique offers the possibility of enhanced selectivity and separation efficiency since ultrasound wave fields exert forces on both NACPs and PACPs in opposing directions. Given this promise, it is necessary to further examine several aspects of using NACPs in cellular separations. For example, the role of bioaffinity bond strength between particles that are being subjected to force in opposite directions may need to be studied in detail. Likewise, the features that enable the primary radiation force of NACPs to dominate that of PACPs requires further investigation. The transport of PACPs to the pressure antinodes will only occur when a complex of PACP bound to NACPs exhibits an overall negative acoustic contrast factor, which can be adjusted through the volume, density, and bulk modulus of the NACPs. In the current study, these properties have converged to favor the relocation of PACP-NACP complexes to the antinode. We anticipate that future experimental and computational investigations will reveal the optimal parameters that support efficient cell separation.

Conclusions

This report communicates a new approach for bioseparation that employs polysiloxane-based microparticles with a negative acoustic contrast property. Emulsifying and post-curing pre-polymers within aqueous surfactant results in stable microparticles that transport to the pressure antinode of an ultrasonic standing wave field in aqueous media. By using polysiloxanes with different chemical compositions and curing chemistries (i.e., PDMS, PVMS), we demonstrate versatility and general utility of silicone materials as negative acoustic contrast agents. Both photochemical and physical adsorption approaches are used to biofunctionalize NACPs, ultimately enabling the specific capture and transport of PACPs to an acoustic pressure antinode. These results encourage further pursuits aimed at using NACPs for cell separation, owing to potential advantages of this system such as high sensitivity, selectivity, portability and low cost.

Methods

Preparation & functionalization of NACPs

Preparing PVMS particles: A mixture of 1.0 g of hydroxyl-terminated PVMS [14], 0.07 g vinylmethoxysiloxane homopolymer (Gelest), and between 0.02 g and 0.03g tin octoate catalyst (Gelest) was thoroughly stirred and combined with a solution of 0.5 or 0.7 wt% Pluronic® F108 (Aldrich) in ultrapure water (Mill-Q, 18MΩ resistivity). The mixture was briefly vortexted, homogenized using a PT 1200E homogenizer (Polytron) with a 3mm rotor for 5 min at 18,750 rpm, and stirred for at least 2 hr at ~50°C. The polydisperse emulsion was permitted to cure via alkoxy condensation of silanol-terminated PVMS with vinylmethoxysiloxane. Particles were left at ambient conditions for approximately one week, then filtered through a 12 μm polycarbonate membrane (Whatman, Cyclopore), and stored at ambient conditions until use. Preparing PDMS particles: A mixture comprising a 1:10 weight ratio of curing agent: base of Sylgard® 184 (Dow Chemical) was thoroughly mixed and 1 gram of the mixture was subsequently combined with 1 wt% of Pluronic F108. The mixture was homogenized as previously described. The emulsion was incubated at 45°C, stirring for at least 1.5 hrs and subsequently left at ambient conditions for at least 12 hrs to permit curing. Functionalization: For reactions with biotin-TFPA (Quanta Biodesign), ~5 × 107 PVMS microparticles were washed with 1× PBS by centrifuging and resuspending the pellet in a final volume of 2 mL of 1× PBS. The microparticles were transferred to a cylindrical glass vial (2.5 cm diameter) and 3 mg biotin-TFPA in 100 μL of dimethyacetamide was added. Light irradiation occurred using an Omnircure S1000 equipped with a high pressure mercury lamp and an internal 320–500 nm filter. The associated light guide was placed ~5 mm above the stirring solution for 30 min at a light intensity of ~100 mW/cm2 at a wavelength of 365 nm, (as measured by Powermax USB sensor, Coherent). The resultant yellow solution was stored at 4°C until use. Biotinylation of Pluronic F108 surfactant followed a similarly reported protocol [20]. Briefly, hydroxyl end groups on F108 were modified to succinimidyl carbonate using N,N-disuccinimidyl carbonate (Aldrich) and 4-(dimethylamino)pyridine (Aldrich) and subsequently reacted with biotin-hydrazide (Aldrich). Once biotinylated, Pluronic F108 was used to prepare silicone emulsions as previously described. Subsequent addition of streptavidin (AlexaFluor® 488 or AlexaFluor® 546) to NACPs occurred by washing particles at least three times by centrifuging and resuspending...
the pellet in 1× PBS, and incubating with either 1 μM or 1.7 μM of streptavidin for 30 min at room temperature.

Characterization of negative acoustic contrast materials and microparticles

Attenuated total reflection-Fourier transform infrared (ATR-FTIR) spectra were acquired using a Thermo Electron Nicolet 8700 spectrometer (Ge crystal, 32 scans, 4 cm⁻² resolution). Scanning electron microscopy (SEM) images were obtained using model FEI XL 30 SEM under ultra-high resolution mode after sputter coating the samples with approximately 6 nm of gold. Optical microscopy images were obtained using an upright Zeiss Axio Imager A2 microscope with appropriate filter set (ex 470/40, em 525/50 or ex 545/25, em 605/70 or ex 365, em 445/50).

Bioseparation studies

Binding between streptavidin polystyrene microparticles (Polysciences, YG microspheres, 6 μm) and PDMS NACPs (encapsulated with rhodamine B, functionalized with biotin-F108) occurred by combining ~10⁶ polystyrene particles and ~10⁷ PDMS particles and incubating for 30 minutes at room temperature with end-over-end rotation. Before combining with the polystyrene microparticles, ~10⁷ PDMS NACPs were washed three times with 1× PBS. Polystyrene particles were added directly from the manufacturer’s stock without washing. Bioseparation events within the channel were monitored through the glass lid of the acoustofluidic device using fluorescent microscopy.

Fabrication of acoustofluidic device

The acoustofluidic device (Additional file 3) was prepared using standard photolithography, deep reactive-ion etching, anodic bonding and plasma bonding. The device contained a downstream collection module and an acoustic (piezoelectric) actuation element (i.e., lead zirconate titanate, PZT, 841 WFB, d₃₃ = 0.3 nm/V, APC International). The channel width was designed to operate at a half wavelength resonant mode (e.g., 252 μm and frequency of 2.94 MHz or 272 μm and frequency of 2.72MHz) resulting in an antinode at both channel walls and a single node in the channel center line. For the experiments, an electric signal with peak-to-peak voltage of 31 V was applied to the PZT. Prior to running experiments, the acoustofluidic channels were treated with a solution of Pluronic F108.

Additional files

Additional file 2: Analysis of PVMS microparticles after functionalization with biotin-TFFA and fluorescent streptavidin.

Histogram showing the signal to background (S/B) fluorescent values of: (A) PVMS microparticles (~5 × 10⁴ particles/ml) combined with biotin-TFFA and irradiated with a 320–500 nm light source (~10 mW/cm²) for 30 minutes. The particles were subsequently labeled with streptavidin AlexaFluor® 488 and washed with 1× PBS. The negative control reactions were performed identical except (B) without light irradiation or (C) without biotin-TFFA. All fluorescent values were taken from images acquired using a 40x objective and 25 ms exposure. Three separate fluorescent images from the same sample were taken and used to calculate standard deviations.

Additional file 3: Acoustofluidic device. Digital camera images showing (A) the glass top and (B) the silicon underside of an exemplary acoustofluidic device. To collect downstream sorted particles, a trifurcation arrangement was designed with two side outlets and a single middle outlet, where negative and positive acoustic contrast particles would exit, respectively. The PZT is attached to the silicon underside.

Abbreviations

PACPs: Positive acoustic contrast particles; NACPs: Negative acoustic contrast particles; PBS: Phosphate buffered saline; PDMS: Polydimethylsiloxane; PVMS: Polvynölymethylsiloxane; TFFA: Tetrafluorophenyl azide.

Competing interests

The authors declare no competing interests.

Authors’ contributions

LMJ, LG, CWS, and MS performed the experiments including the microparticle preparation and characterization, acoustofluidic chip fabrication, and bioseparation experiments. KE prepared and characterized PVMS material and contributed to PVMS particle formulation. KC helped design preparation procedures for PDMS particles. All authors offered scientific input during these studies and assisted in preparation of the manuscript. All authors approved the manuscript.

Acknowledgements

This work was supported by the National Science Foundation (NSF, through the Research Triangle MRSEC, DMR-1121107 and CBET-10-50116). LMJ thanks The Hartwell Foundation (Biomedical Research Fellowship) and CWS is grateful for a NSF Graduate Research Fellowship (1106401). MS thanks the Pratt Research Fellows program at Duke University. KC thanks the National Institutes of Health (NH RR020064, NIH RR001315). We thank Zijian Zhou at Duke University for the images of the acoustofluidic chip.

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Received: 14 June 2013 Accepted: 14 June 2013

Published: 28 June 2013

References

1. Gor’kov LP: On the forces acting on a small particle in an acoustical field in an ideal fluid. Soviet Physics-Doklady 1962, 6:773–775.
2. Bruus H: Acoustofluidics 7: the acoustic radiation force on small particles. Lab Chip 2012, 12:1014–1021.
3. Laurell T, Peterson F, Nilsson A: Chip integrated strategies for acoustic separation and manipulation of cells and particles. Chem Soc Rev 2007, 36:492–506.
4. Lenshof A, Magnusson C, Laurell T: Acoustofluidics 8: applications of acoustophoresis in continuous flow microsystems. Lab Chip 2012, 12:1210–1222.
5. Ward M, Turner P, Delohn M, Kaduchak G: Unit 1.22 fundamentals of acoustic cytometry. Current Protocols in Cytometry 2009, 491:22.1–1.22.12.

6. Piyasena ME, Austin Suthanthirapj PP, Applegate RW Jr, Goumas AM, Woods TA, López GP, Graves SW: Multinode acoustic focusing for parallel flow cytometry. Anal Chem 2012, 84:1831–1839.

7. Petersson F, Nilsson A, Holm C, Jönsson H, Laurell T: Separation of lipids from blood utilizing ultrasonic standing waves in microfluidic channels. Analyst 2004, 129:938–943.

8. Petersson F, Nilsson A, Holm C, Jönsson H, Laurell T: Continuous separation of lipid particles from erythrocytes by means of laminar flow and acoustic standing wave forces. Lab Chip 2005, 5:20–22.

9. Thévoz P, Adams JD, Shea H, Bruus H, Soh T: Acoustophoretic synchronization of mammalian cells in microchannels. Anal Chem 2010, 82:3094–3098.

10. Dykes J, Lenshof A, Åstrand-Grundström I, Laurell T, Scheding S: Efficient removal of platelets from peripheral blood progenitor cell products using a novel micro-chip based acoustophoretic platform. PLoS One 2011, 6:e23074.

11. Petersson F, Åberg L, Swärd-Nilsson AM, Laurell T: Free flow acoustophoresis: microfluidic-based mode of particle and cell separation. Anal Chem 2007, 79:5117–5123.

12. Cushing KW, Piyasena ME, Carroll NJ, Maestas GC, López BA, Edwards BS, Graves SW, López GP: Elastomeric negative acoustic contrast particles for affinity capture assays. Anal Chem 2013, 85:2208–2215.

13. Hu S, Ren X, Bachman M, Sims CE, Li GP, Allbritton N: Surface modification of poly(dimethylsiloxane) microfluidic devices by ultraviolet polymer grafting. Anal Chem 2002, 74:4117–4123.

14. Fuard D, Tzvetkova-Chevolleau T, Decossas S, Tracqui P, Schiavone P: Optimization of poly-di-methyl-siloxane (PDMS) substrates for studying cellular adhesion and motility. Microelectron Eng 2008, 85:1289–1293.

15. Hellmich W, Regtmeier J, Duong TT, Rost R, Anselmetti D, Ros A: Poly (oxyethylene) based surface coatings for poly(dimethylsiloxane) microchannels. Langmuir 2005, 21:7551–7557.

16. Efimenko K, Crowe JA, Manias E, Schwark DW, Fischer DA, Genzer J: Rapid formation of soft hydrophilic silicone elastomer surfaces. Polymer 2005, 46:9329–9341.

17. Ferguson GS, Chaudhury MK, Biebuyck H, Whitesides GM: Monolayers on disordered substrates: self-assembly of alkyltrichlorosilanes on a surface-modified polyethylene and poly(dimethylsiloxane). Macromol 1993, 26:5870–5875.

18. Linder V, Verpoorte E, Thorman W, de Rooij NF, Sigrist H: Surface biopassivation of replicated poly(dimethylsiloxane) microfluidic channels and application to heterogeneous immunoreaction with on-chip fluorescence detection. Anal Chem 2001, 73:4181–4189.

19. Owen MJ, Smith PJ: Plasma treatment of polydimethylsiloxane. J Adhes Sci Technol 1994, 8:1063–1075.

20. Hermanson GT: From Bioconjugate Techniques. 2nd edition. Oxford, UK: Elsevier; 2008: 204, 215, 940–945.

21. Grevval C, Augustsson P, Folkenberg JR, Laurell T: Harmonic microchip acoustophoresis: a route to online raw milk sample precondition in protein and lipid content quality control. Anal Chem 2009, 81:6195–6200.

22. Hartono D, Liu Y, Tan PL, Then XYS, Yung LL, Lim K: On-chip measurements of cell compressibility via acoustic radiation. Lab Chip 2011, 11:4072–4080.