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# Optical feedback control loop for the precise and robust acoustic focusing of cells, micro- and nanoparticles†

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Despite a long history and the vast number of applications demonstrated, very few market products incorporate acoustophoresis. Because a human operator must run and control a device during an experiment, most devices are limited to proof of concepts. On top of a possible detuning due to temperature changes, the human operator introduces a bias which reduces the reproducibility, performance and reliability of devices. To mitigate some of these problems, we propose an optical feedback control loop that optimizes the excitation frequency. We investigate the improvements that can be expected when a human operator is replaced for acoustic micro- and nanometer particle focusing experiments. Three experiments previously conducted in our group were taken as a benchmark. In addition to being automatic, this resulted in the feedback control loop displaying a superior performance compared to an experienced scientist in 1) improving the particle focusing by at least a factor of two for 5 µm diameter PS particles, 2) increasing the range of flow rates in which 1 µm diameter PS particles could be focused and 3) was even capable of focusing 600 nm diameter PS particles at a frequency of 1.72075 MHz. Furthermore, the feedback control loop is capable of focusing biological cells in one and two pressure nodes. The requirements for the feedback control loop are: an optical setup, a run-of-the-mill computer and a computer controllable function generator. Thus resulting in a cost-effective, high-throughput and automated method to rapidly increase the efficiency of established systems. The code for the feedback control loop is openly accessible and the authors explicitly wish that the community uses and modifies the feedback control loop to their own needs.

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#### Introduction

The Kundt's tube described in 1866 (ref. 1) is based on acoustophoresis, a contactless, label-free, and non-invasive method to actively control the position of particles over a wide range of sizes using acoustic pressure waves.<sup>2</sup>

In 1938 Hillary W. St. Clair filed, to our knowledge, the first patent related to acoustophoresis "Sonic flocculator and method of flocculating smoke or the like". Continued

research on both the theoretical as well as the experimental aspect of acoustophoresis has led to a vast number of applications such as acoustic particle levitation,<sup>4</sup> acoustic holograms<sup>5</sup> and tweezers,<sup>6</sup> aiding in 3D mechanical characterization<sup>7</sup> and isolating cancer cells from blood<sup>8</sup> to name only a few.

Despite a long history, a plethora of publications in the field and interesting applications demonstrated in research, only a small number of companies offer products based on acoustic particle manipulation. There are many reasons for this such as the currently imprecise manufacturing which could be tackled by improved cleanroom processes or by improving the predictability or reliability of devices using more complex production methods or designs. In the enhanced understanding of the fundamental principles of acoustophoresis, aiding in device design, various computational models have been created. Adding these computational models, deep learning can be used to find a device design that creates a desired acoustic potential.

The beforehand mentioned approaches to improve the performance of acoustic devices place an emphasis on

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improving the theoretical device performance. When considering a prevalent application of acoustophoresis, focusing micro- and nanometer particles, 20-26 additional methods have sought to further increase the experimental outcome. A successful outcome is achieved by reducing the influence of acoustic streaming vs. the acoustic radiation force. Due to the size dependent scaling, seen in eqn (2) and (6), this becomes increasingly difficult if the particles' size is decreased to below 1 micron. In addition, temperature changes must be taken into account, which result in a detuning of the device and therefore lower pressure amplitudes. An approach is to control the excitation frequency of the piezoelectric transducer which is enabling the focusing via a feedback control loop (FCL). This is accomplished by measuring the resonance response of the system and thus indirectly effects the outcome of the experiment. Two parameters are currently being used to measure the resonance, which are the impedance of the system<sup>27–29</sup> or the electric current.<sup>30</sup> The measured resonance behavior of the system is then used to tune the excitation frequency of the driving piezoelectric transducer. These state of the art methods of which parameters can be used to implement a FCL are summarized in Table 1.

It however has been noted that these approaches, which are based on measuring electric signals, might not be able to correctly identify the optimal excitation frequency of a coupled system consisting of the piezoelectric transducer and the microfluidic channel.31 To try and elevate this shortcoming, the system can be characterized a priori to an experiment to try and find the corresponding resonance frequencies<sup>32</sup> or an FCL can be combined with a lookup table.<sup>29</sup> However, all methods require additional electrical equipment and while increasing the performance of a system, they only indirectly influence the performance, especially for particles close to the critical radius as actuating the system at resonance does not guarantee an optimal streaming suppression. An optical FCL has been reported only in the context of particle positioning<sup>33</sup> which uses a visual input to control the position of one particle in two dimensions. An optical FCL, which directly takes the desired outcome as the control signal, has not been presented before. Our goal was therefore to expand on the idea of an optical FCL, schematically represented in Fig. 1, which directly effects the performance of a system classified by the quality of the particle focusing. The shift from a resonance control to an optical control can also help in finding the optimal excitation frequency when acoustic streaming is present. Thus being able to focus particles that otherwise would be

Table 1 Summary of state of the art methods

	Measured parameter	Year published
Goddard et al. <sup>27</sup>	Impedance	2006
Suthanthiraraj et al. <sup>28</sup>	Impedance	2012
Kalb et al. <sup>29</sup>	Impedance	2018
Farmehini <i>et al.</i> <sup>30</sup>	Electric current	2021

dominated by acoustic streaming. The FCL uses the particle linewidth (LW) as the parameter to optimize, where the LW is the spread of the particles in the microfluidic channel of a device, Fig. 2. The FCL can easily be integrated into any preexisting setup that allows for an optical readout during the experiment, increasing the sample throughput whilst not increasing the cost.

Optically automatizing the determination of an optimal excitation frequency leads to an increase of the acoustic force acting on the particles of interest. This lead to an improved performance of acoustophoretic devices when compared to three benchmark tests<sup>10</sup> by 1) narrowing the focusing of the particles by at least a factor of two for 5 µm PS particles, 2) increasing the range of flow rates in which 1 µm diameter PS particles could be focused and 3) focusing particles with a diameter as small as 600 nm (Fig. 3). The manipulation of such small particles is a significant step, since in state of the art publications, a very complicated setup needs to be utilised to manipulate particles in this size range,<sup>34</sup> or no fluid flow could be applied.35 Finally, we demonstrate the versatility of the setup by showing two different types of devices and modes of operation (Fig. 4) using biological cells. The above mentioned augmented performance, based on automatising the focusing, is needed for applications to be successful outside of laboratory settings. We therefore believe that our novel optical FCL can be used as a foundation to further the development and integration of acoustophoresis into market products.<sup>9,36</sup> This is in line with the increasing need of simpler and more adaptable lab on a chip devices.<sup>37</sup>

## Theoretical background

#### Acoustic radiation force (ARF)

The FCL demonstrated in this paper allows to precisely focus particles of interest. This focusing exploits the ARF which, given a spherical particle with a radius much smaller than the acoustic wavelength in an inviscid fluid, is commonly approximated as the negative Gradient of the Gor'kov potential<sup>38</sup>

$$F_{\rm rad} = -\nabla U,\tag{1}$$

with the Gor'kov potential

$$U = \frac{4\pi}{3} r^3 \left[ f_1(\tilde{\kappa}) \frac{1}{2\rho_0 c_0^2} \langle p_1^2 \rangle - f_2(\tilde{\rho}) \frac{3}{4} \rho_0 \langle v_1^2 \rangle \right]$$
 (2)

where r is the radius of the particle,  $\rho_0$  is the density of the fluid,  $c_0$  the speed of sound of the fluid,  $\langle p_1^2 \rangle$  the first order time averaged square of the incident acoustic pressure and  $\langle \mathbf{v}_1^2 \rangle$  the first order time averaged square of the incident acoustic velocity. Furthermore, the monopole coefficient  $f_1(\tilde{\kappa})$  which is related to the relative compressibility and the dipole coefficient  $f_2(\tilde{\rho})$  which is related to the relative density

$$f_1(\tilde{\kappa}) = 1 - \frac{\kappa_{\rm p}}{\kappa_0} \tag{3}$$

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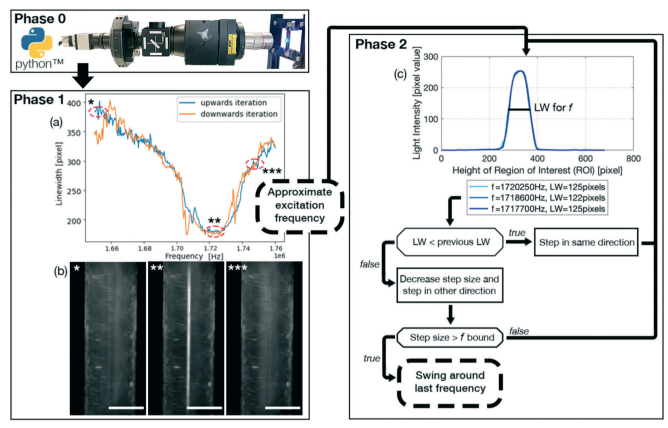


Fig. 1 Schematic overview of the feedback control loop (FCL). Phase 0 initializes the FCL and the video stream from the device being filmed (blue dot) by the uEye camera (grey metal housing above the black arrow on the left) is taken by a screen capture function and fed into the FCL. Phase 1 (a) recorded particle linewidths (LW), a measure for particle focusing, for phase 1 of the FCL using 5  $\mu$ m PS particles focused in the center of the device by an acoustic standing wave for the frequency range 1.66–1.78 MHz with a 600  $\mu$ L min<sup>-1</sup> flow rate. The function generator was controlled by the FCL, sweeping through the frequency spectrum with 400 Hz steps to find the minimal linewidth. (b) \*t = 0 s, \*\*t = 68 s, \*\*\*t = 110 s are images, where the asterisk (\*) correlates the graphical readout of the frequency sweep to the red dashed circles of (a) of the upwards iteration (blue data entries). Phase 2 on the right indicates how the approximate excitation frequency of phase 1 is used to initialize phase 2, which ultimately leads to the determination of an optimal excitation frequency. For clarity, a safeguard against continuously stepping in the wrong frequency direction is left out of the schematic. (c) 3 different frequencies obtained from phase 2 are plotted against each other and corresponds to the phase 2 of (a), indicating the precision of phase 1 and illustrates that for simple systems, such as a glass capillary and 5  $\mu$ m PS particles, phase 1 might be sufficient and that a more precise excitation frequency does not result in a significantly narrower linewidth. White scale bar (b): 500  $\mu$ m.

$$f_2(\tilde{\rho}) = 2\frac{\rho_{\rm p} - \rho_0}{2\rho_{\rm p} + \rho_0}$$
 (4)

where  $\kappa_{\rm p}$  is the compressibility of a particle,  $\kappa_{\rm 0}$  is the compressibility of the fluid and  $\rho_{\rm p}$  is the density of the particle.

A standing wave is required for a large ARF magnitude. The resonance frequency for the *n*th ultrasonic resonance mode of a one-dimensional standing wave with hard wall boundary conditions is<sup>39</sup>

$$f_{\rm res}^{\rm 1D} = \frac{c_0 n}{2w} \tag{5}$$

where w is the width of the channel.

#### Stokes' drag force

In addition to the ARF, a second force, the Stokes' drag force produced by the acoustic streaming velocity  $\nu_{str}$ , acts on the particle in the fluid<sup>40,41</sup>

$$F_{\rm str} = 6\pi \eta r (\mathbf{v}_{\rm str} - \mathbf{v}_{\rm p}), \tag{6}$$

where  $\eta$  is the dynamic viscosity of the fluid and  $v_p$  the velocity of the particle. In addition to the Stokes' drag force resulting from the acoustic streaming velocity  $v_{str}$ , the Stokes' drag force results from the viscous resistance for particles moving through a fluid and is therefore always present for  $v_p \neq 0$ . The Stokes' drag force, no matter the origin, scales with r (eqn (6)) and the ARF scales with  $r^3$ (eqn (1)). This means that overall the Stokes' drag force always dictates the motion of the particle, but the influence of the Stokes' drag force coming from the acoustic streaming velocity is diminished compared to the ARF as particles get larger. Therefore, there is a radius, termed the critical radius, above which the ARF cannot overcome the Stokes' drag force stemming from the particle motion, but can overcome the Stokes' drag force resulting from the acoustic streaming velocity. This critical radius can be analytically approximated in 1D<sup>39</sup>

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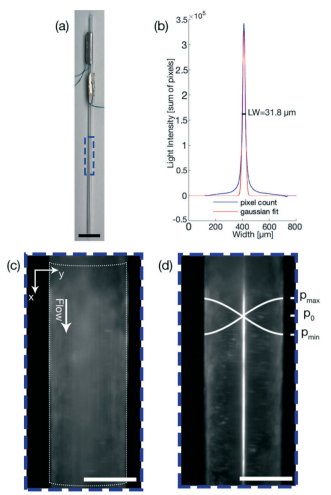


Fig. 2 (a) Glass capillary with two piezoelectric transducers and the ROI for all results generated using the glass capillary indicated by a blue dashed box. (b) Evaluation of the linewidth of the experiment shown in (d) using a self written MATLAB code, where a Gaussian fit is used to include ~70% of all particles. (c) 5 μm PS particles when the FCL is inactive at t=0. (d) Focused 5 μm PS particles at f=1.7204 MHz, V=20 V<sub>pp</sub> and a flow rate of 100 μL min<sup>-1</sup> at t=1.12 s. Black (a) and white (c) and (d) scale bars: 5000 μm respectively 500 μm.

$$r_{\rm c}^{\rm 1D} = \sqrt{\frac{3\psi}{2\Phi}}\delta\tag{7}$$

where  $\psi$  is a geometry dependent factor and the acoustic contrast factor  $\Phi$ 

$$\Phi = \frac{1}{3}f_1(\tilde{\kappa}) + \frac{1}{2}f_2(\tilde{\rho}),\tag{8}$$

and the viscous boundary layer  $\delta$ 

$$\delta = \sqrt{\frac{\eta}{\pi \rho_0 f}}. (9)$$

*Exempli gratia*, the 1D analytical critical radius  $r_{\rm c}$  of a polystyrene particle in water,  $\Phi\approx 0.165$  at f=1.75 MHz in a rectangular microfluidic channel, therefore  $\psi=\frac{3}{8},^{42}$  is  $r_{{\rm c},a}^{\rm 1D}=0.78\,\mu{\rm m}$ . The numerical model for our specific geometry which describes the diminished influence of the acoustic

streaming compared to the influence of the ARF described by Gerlt *et al.*,  $^{43}$  resulted in a numerical critical radius  $r_{c,n}$  = 0.35  $\mu$ m. This still does not match our experimental findings as will be seen in Table 2 and Fig. 4, hence providing further evidence that the critical radius for a given system is not known prior to an experiment.  $^{44}$ 

#### Materials & methods

#### Setup

The setup employed is analogous to the one used in Gerlt et al.43 the fluid flow through a circular glass capillary (76 mm length, 1 mm outer diameter, 0.75 mm inner diameter) and a glass - silicon - glass device (device: 50 mm length, 12 mm width, 1.4 mm thickness; focusing channel: 15 mm length, 1 mm width, 0.2 mm height) was controlled by volumetric syringe pumps (Nemesys, Cetoni). A function generator, connected to the computer running the FCL, (AFG-2225, GW Instek) powered the piezoelectric transducers. The signal from the function generator was amplified (325LA Linear Power Amplifier, Electronics & Innovation) and monitored using an oscilloscope (UTD2025CL, UNI-T). For optical data acquisition a custom made setup from THORLABS consisting of specifically selected parts from the Cerna® series was used. The video feed generated with a 5× objective (M Plan Apo 5×/0.14, Mitutoyo) and a uEye camera (UI-3160CP Rev. 2.1, iDS, 1920 × 1200 pixels, 60 fps), as shown in Fig. 1 phase 0, was captured from one of the two computer screens and fed into the FCL running on PyCharm 2021.1.3 (Edu) with Python 3.8.

#### Glass capillary

In order to compare the device performance as best as possible, the tests were performed with the same glass capillaries, piezoelectric transducers and production procedure that was used by Gerlt *et al.*<sup>43</sup> the glass capillary (TW100-4, World Precision Instruments) has two piezoelectric transducers (10 mm length, 2 mm width, 1 mm thickness, Pz26, Meggitt Ferroperm) glued on using an electrically conductive Epoxy glue (H20E, Epoxy Technology). Copper cables (0.15 mm diameter) were attached to the piezoelectric transducers and the electrical connection was established with an electrically conductive silver paste.

#### Glass-silicon-glass device

The glass–silicon–glass device, seen in in Fig. 4(c), was produced by bonding a 500  $\mu m$  thick glass wafer to a 200  $\mu m$  thick silicon wafer. Channels were patterned onto the exposed silicon wafer using photolithography (resist: S1828, Shipley, 4'000 rpm; developer: AZ351B, Microchemicals). The full thickness of the silicon wafer was etched away with an inductively coupled plasma (ICP) deep reactive ion etching (DRIE) machine (Estrellas, Oxford instruments). Following the etching, a 700  $\mu m$  thick glass wafer was anodically bonded onto the exposed silicon wafer. The wafer was then

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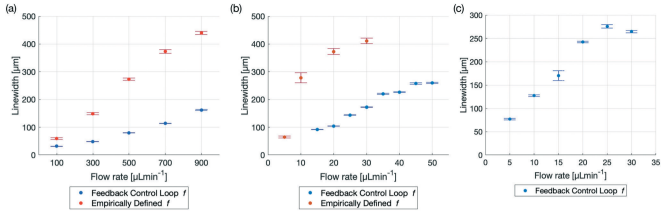


Fig. 3 Comparison of particle focusing in a glass capillary with a empirically defined excitation frequency  $^{43}$  vs. an optimal excitation frequency found by the FCL for different particle sizes and flow rates. All results were generated with the same setup and capillary driving both piezoelectric transducers at  $V = 20 \text{ V}_{pp}$  and the precise linewidth can be found in Table 2. (a) Comparison of 5 μm diameter PS particles for the flow rates: 100, 300 500, 700 and 900 μL min<sup>-1</sup>. The excitation frequency defined by the FCL is f = 1.7204 MHz. The improved linewidth is not only seen in the smaller linewidths at higher flow rates, but also in the smaller error bars, indicating a more precise and robust focusing. (b) The focusing of 1 μm diameter PS particles using an empirically defined frequency was possible for the flow rate of 5 μL min<sup>-1</sup>. At 10 μL min<sup>-1</sup> the linewidth was already close to half of the diameter of the capillary and at higher flow rates, the focusing was nearly non existent. Employing the FCL, which found the frequency at f = 1.7204 MHz, focusing was possible even with flows up to 50 μL min<sup>-1</sup>. (c) 600 nm diameter PS particles could be focused by the FCL, when utilizing phase 1 and phase 2. The frequency defined by the FCL from phase 1 was 1.7202 MHz and phase 2 further iterated and found an improved excitation frequency at 1.72075 MHz. Even at high flow rates, e.g. 25 μL min<sup>-1</sup> and 30 μL min<sup>-1</sup> the linewidth was still smaller than half of the capillary diameter.

diced into individual chips with a wafer saw (DAD3221, Disco corporation). Fused silica capillaries (164  $\pm$  6  $\mu m$  outer diameter, 100  $\pm$  6  $\mu m$  inner diameter, Molex) were inserted into the inlets and outlets of the chips and fixed with a two-component glue (5 Minute Epoxy, Devcon). The piezoelectric transducers, Epoxy glue, wiring and an electrically conductive silver pasted are the same ones used for the glass capillaries.

#### Polystyrene particles

Green fluorescent polystyrene (PS) particles (microParticles GmbH, Germany) with diameters of 5.19  $\pm$  0.14  $\mu m,~1.08~\pm$  0.04  $\mu m$  and 600  $\pm$  20 nm were used for all experiments in Fig. 3 and all experiments were conducted with 0.5% v/v.

#### Cell culture

The bone cancer cells used is the SaOs-2 cell line. The cell line was kept at the standard 37 °C and 5% CO<sub>2</sub> and 95% air. The cell media used is DMEM – F12 Ham (D8437, Sigma) supplemented with 10% fetal bovine serum (10270106, Thermo) and 1% P/S. The cells were passaged when 60% confluency was reached. The diameter of the cells is 15.73  $\pm$  0.42  $\mu m$  (CellDrop BF, DeNovix).

#### Feedback control loop (FCL)

The FCL, shown schematically in Fig. 1 and available on Gitlab (https://gitlab.com/acoustofluidics/feedback-controlloop), is implemented in the widely used programming language Python and is split into 3 phases,

- Phase 0: initialization of the system.
- Phase 1: approximate determination of an optimal excitation frequency.

• Phase 2: precise determination and stabilisation of the optimal excitation frequency.

A highlight of the code is the ImageGrab function. This allows the user to continuously capture the image directly from the computer screen of the computer controlling the imaging system in order to use the image for the FCL to determine the linewidth. This ensures that any preexisting optical setup can be utilized. Therefore, the imaging system does not need to be adapted to the FCL, thus being an augmentation without any additional costs. It is recommended to use two monitors, such that the plotting does not interfere with the screen capture.

In the following description, this typeset was used to highlight variables.

Phase 0. The connection to the function generator is established and all necessary Python libraries are loaded. Libraries required are pyserial, opency-python, keyboard, termcolor, numpy, and matplotlib. For setups that deviate from the setup demonstrated here, the script needs to be altered in order to accommodate for different function generators. In addition, the parameters for the experiment need to be set, such as where the frequency sweep starts and where it stops, defined as freq\_min, freq\_max. freq\_min and freq\_max can either be defined manually starting at a frequency defined by experience or calculated using a user defined percentage interval. In this phase, the region of interest (ROI) needs to be chosen and the code creates a background image, which is used for a background subtraction later on, if needed. In addition to the listed parameters and working directories, parameter sets for phase 1 and phase 2 need to be defined. These parameters depend on the flow rate, input voltage, particle concentration and

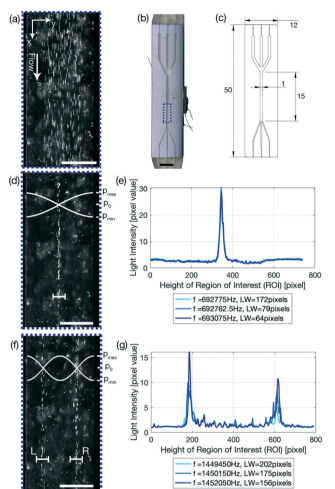


Fig. 4 Glass-silicon-glass device used for the manipulation of biological cells. (a) SaOs-2 cells when the FCL is inactive and a cell concentration of  $0.361 \times 10^5$  cells per ml. (b) Image of the device and the ROI indicated by the dashed blue box. The top and the bottom of the device have a beige coloring due to the epoxy glue used to glue the capillary connections. A piezoelectric transducer is glued onto the side of the device at half of the height on the right hand side. (c) Sketch of the device with the dimensions given in mm. Device: 50 mm length, 12 mm width, 1.4 mm thickness; focusing channel: 15 mm length, 1 mm width, 0.2 mm height (d) SaOs-2 cells focusing in the center of the channel of the device with a linewidth of 31.28  $\pm$  0.98  $\mu$ m at f = 0.7101 MHz and a flow rate of 10  $\mu$ L min<sup>-1</sup>. The white crossbar indicating the focusing is not to scale. (e) Plot of 3 frequencies and the corresponding linewidths for phase 2. (f) Cells focusing in two lines. The left-hand-side (L) linewidth is 33.01  $\pm$  5.27  $\mu m$  and the righthand-side (R) linewidth is 36.15  $\pm$  1.00  $\mu$ m at f = 1.453 MHz and a flow rate of 10 µL min<sup>-1</sup>. (g) Plot of 3 frequencies and the corresponding linewidths for phase 2. (e) & (g) were re-plotted with altered colors and a reduced amount (three) of plotted frequencies. The voltage was 30  $V_{pp}$ . The lighting angle and the video settings, mainly a high exposure time set new at the beginning of each experiment, result in the bright cells. Black and white scale bar: 5000  $\mu m$  respectively 500  $\mu m$ .

size to name a few. Exemplary values that were used for the experiments can be found in the code on Gitlab (https://gitlab.com/acoustofluidics/feedback-control-loop). As to be expected, the time required for each phase highly depends on the set of parameters and the frequency range chosen. Phase 1 of Fig. 1 for instance took ~4 minutes.

**Table 2** Comparison of the LW between the empirically defined excitation frequency and the frequency found by the FCL (FCL). This data corresponds to the data from Fig. 3. \*No data available from previous experiments. \*\*Data was excluded due to particle trapping

Flow rate [μL min <sup>-1</sup> ]	LW for an empirical $f$ [ $\mu$ m]	LW for FCL defined $f$ [ $\mu$ m]
5 μm diameter	PS particles	
100	59.54 (±2.67)	$31.80 (\pm 0.58)$
300	149.02 (±4.26)	$48.51 (\pm 0.28)$
500	273.17 (±3.92)	80.27 (±0.71)
700	373.19 (±6.65)	114.27 (±0.57)
900	439.65 (±5.67)	162.55 (±1.31)
1 μm diameter	PS particles	
5	65.02 (±3.92)	**
10	277.62 (±18.20)	**
15	*	92.21 (±0.72)
20	372.93 (±10.42)	104.61 (±0.26)
25	*	144.14 (±0.63)
30	411.29 (±9.94)	172.41 (±0.89)
35	*	216.66 (±1.91)
40	*	$225.83(\pm 1.14)$
45	*	257.22 (±2.22)
50	*	259.76 (±1.88)
600 nm diamet	er PS particles	
5	*	77.23 (±1.50)
10	*	127.42 (±1.57)
15	*	170.37 (±10.48)
20	*	242.52 (±1.80)
25	*	275.95 (±3.74)
30	*	264.81 (±2.61)

The set of parameters that need to be defined for phase 1 is comprised of: initial\_time\_delay\_1 which is the initial time delay for system to settle, frequency\_step\_size\_1 which is the frequency step size, time\_delay\_1 which is the time delay between frequency steps.

The set of parameters that need to be defined for phase 2 is comprised of: wait\_time\_transition\_2 which is the time delay for the system to settle between phase 1 and phase 2. frequency\_step\_size\_2 which is the frequency step size. time\_delay\_2 which is the time delay between the frequency steps. Threshold which is the threshold value for a change of direction, if the linewidth has not improved in the past |threshold| iterations. swing\_value which is the value with which the step size is multiplied with after changing the direction of iteration. The swing\_value is therefore  $\in$  (0, 1). freq\_bound is the lower frequency boundary for minimal step size. In our experience, a |freq\_bound| = 20 [Hz] leads to optimal results.

Phase 1. The FCL steps through from the lowest to the highest set frequency and once the highest frequency is reached, starts to step down again, as shown in Fig. 1(a), which is the plot generated after phase 1 is completed. The FCL then automatically sets the frequency to the approximate excitation frequency determined by the best average linewidth from the up- and downward iteration. The linewidth calculation is demonstrated in Fig. 2, for the up- and downward iteration. The up- and downward iteration

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allows for an averaging of the best linewidth to double-check the optimal frequency. The up- and downward iteration shown in Fig. 1 is not necessarily needed, as the curves look similar enough, but the lower the flow rate and the smaller the particle size, the more important the up- and downward iteration becomes, as the ARF decreases with decreasing particle size. The theoretical resonance frequency  $f_{\rm res}^{\rm 1D}$  calculated using eqn (5) can differ from both the empirically chosen frequency and the frequency found by the FCL. This highlights how precarious finding an approximately optimal excitation frequency is, and that even experienced users can introduce human error, especially as many devices show focusing over a multiple hundred kHz range, exemplary seen in Fig. 1(a), therefore making it is easy to pick an excitation

frequency which is far off from an optimal excitation frequency.

Phase 2. In phase 1, the functionality of the FCL is limited to finding an approximate starting value for the optimal excitation frequency. Once the approximate excitation frequency is found, phase 2 narrows down the range of the optimal excitation frequency, as illustrated in Fig. 1(c). To initialize phase 2, the FCL takes the linewidth of the approximate excitation frequency determined in phase 1 and sets the linewidth as the best previous linewidth. Then the FCL starts to iterate downwards with the step size frequency\_step\_size\_2. The linewidth for each (cur\_linewidth) is compared to the best previous linewidth (best linewidth). If best\_linewidth cur\_linewidth, best\_linewidth is set to cur\_linewidth. If best\_linewidth is narrower than cur\_linewidth an internal counter (count) is increased by 1. This is done until count = threshold. Once count = threshold is reached, continuing to step in the same direction does not yield a narrower linewidth. Therefore frequency\_step\_size\_2 is then multiplied by swing\_value, count is set to zero and the FCL steps in the opposite direction with the decreased step size. This process is continued until frequency\_step\_size\_2 is smaller than freq\_bound. When freq\_bound is reached, the step size is kept constant for the remainder of the experiment and the FCL continuously oscillates around the frequency. This can be used to balance out any fluctuations in concentration, flow rate or temperature, to name a few common disturbances.

It is recommended that for a first rough characterization of the system phase 1 is run at high flow rates, e.g. 100 µL min<sup>-1</sup> and above, with particle diameters of at least a few micrometer diameter and in a large frequency range, e.g. ±10% of the expected resonance frequency. In a second round of characterisation, the frequency step sizes and frequency range can be decreased and shifted symmetrically around the approximately optimal excitation frequency and time\_delays can be increased for an even more precise approximate excitation frequency. The thus found approximate excitation frequency can be constant over multiple days and experiments when using multi micrometer diameter particles. For particles around one micrometer in diameter or smaller it is recommended to use phase 2.

#### Linewidth calculation

The linewidth was chosen as the parameter that the FCL optimizes, and in our case is defined as the spread of particles over the microfluidic channel, thus being a measure for the focusing of the particles. The linewidth is determined differently in phase 1 and phase 2.

In phase 1 the linewidth is calculated by taking the sum of the pixel values of all the pixels in the direction of flow (indicated by the *x*-axis in Fig. 2(c)) and is plotted over the width of the channel (indicated by the *y*-axis in Fig. 2(c)). The linewidth is then calculated using a Gaussian fit as the region in which 2 standard deviations, or  $\sim$ 70% of the total sum of the pixels, are located. This linewidth readout is then plotted over the frequency range as shown in Fig. 1(a).

In phase 2, as seen in Fig. 1(c), 4(e) and (g), the pixel distribution over the whole channel is displayed for a given frequency, where the pixel values range from 0 (pure black) to 255 (pure white). The resulting pixel value is the average pixel value of all the pixels in the direction of flow (indicated by the *x*-axis in Fig. 4(e) and (g)) and is plotted over the width of the channel (indicated by the *y*-axis in Fig. 4(e) and (g)). The linewidth is calculated the same way as for phase 1, but multiple frequencies are plotted, providing live output of how the particle distribution is evolving. The linewidth is given in the legend, with the corresponding frequency.

#### Results

The data presented provides a brief overview of some of the use cases where the FCL can be employed to increase device performance and throughput, whilst being independent of a human operator. Fig. 3 and Table 2 compare the relative performance between an excitation frequency determined empirically<sup>10</sup> and the excitation frequency defined by the FCL. All experiments were conducted with 0.5% v/v.

#### Improved focusing of 5 micrometer diameter PS particles

There is significant demand for narrow LWs at high flow rates to increase the throughput while minimizing clogging and it was recently demonstrated that flow rates of up to 1000  $\mu L \ min^{-1}$  are relevant and achievable for cell separation.45 Flow rates of up to 900 µL min-1 were demonstrated, in ref. 43 albeit the focusing drastically decreased at high flow rates, Fig. 3. The FCL presented here is capable of focusing the PS particles down to a narrower LW, with an increased stability, where the stability increase is defined as the reduced size of the error of the measurements, as illustrated in Fig. 3 and Table 2. The FCL resulted in a LW which was half as narrow as the LW that resulted from determining the excitation frequency empirically for a flow rate of 100 μL min<sup>-1</sup> (94 mms<sup>-1</sup>). For higher flow rates, the LW remained narrower by even larger margins and at a 900 μL min<sup>-1</sup> (850 mms<sup>-1</sup>) flow rate the LW resulting from the FCL was about the same width as the LW for a flow rate of 300 μL min<sup>-1</sup> for the empirically determined frequency. The

frequency chosen for this comparison is f=1.7204 MHz, found using only phase 1 of the FCL as in Fig. 1. Technically, phase 2 could have been run as well, but as the LW was already half of the LW when using the empirically determined excitation frequency (f=1.74 MHz), phase 2 was

#### Higher flow rate for 1 micrometer diameter PS particles

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Decreasing the particle size increases the difficulty to focus the particles as the ARF decreases and therefore more precise excitation frequencies are required. Focusing of 1 µm diameter PS particles for instance is possible however, above 10 μL min<sup>-1</sup> Gerlt et al. 43 were not able to focus 1 μm PS particles, as the excitation frequency determined empirically has a limited precision. The FCL demonstrates that 1 µm particles can be focused beyond 10  $\mu L \, min^{-1}$ . Even at 5 times the flow rate (50 µL min<sup>-1</sup> instead of 10 µL min<sup>-1</sup>) the particle LW was still narrower with the frequency chosen by the FCL, compared to the empirically chosen frequency in Fig. 3. The lack of focusing of the FCL of 5 and 10 μL min<sup>-1</sup> flow rates can be attributed to PS particles being trapped in the device whereas the trapping can only be overcome at higher flow rates. As for the 5 µm PS particles, only phase 1 was used to generate this data.

#### 600 nanometer diameter PS particle focusing

Particle sizes greater than one micrometer in diameter are important, but with improving device designs and manufacturing precision, focusing of nanometer particles is a realistic aim. Building on the corpus of nanometer particle trapping46 and separation,47 the FCL demonstrated the capability to use bulk acoustic wave (BAW) devices to focus nanometer particles. When the particle radius is around the critical radius  $r_c$  the acoustic streaming and thus the Stokes' drag force resulting from the acoustic streaming can dominate the movement of the particles in the fluid, therefore phase 1 and phase 2 of the FCL were run in order to find an excitation frequency at which the dominating force was still the ARF. eqn (7) indicates that the analytical and numerical critical radius where the AS dominates and no focusing is possible is  $r_{\mathrm{c},a}^{\mathrm{1D}} = 0.78\,\mathrm{\mu m}$  respectively  $r_{\mathrm{c},n} =$ 0.35 µm. The focusing of 600 nm diameter PS particles however show that both the analytical and the numerical solution both provide critical radii above which focusing was demonstrated in Fig. 3. While<sup>43</sup> was not able to find any focusing of 600 nm diameter particles, the FCL presented here could identify excitation frequencies at which the 600 nm diameter particles could be focused. Even at higher flow rates, such as 30 µL min<sup>-1</sup>, the excitation frequency determined by the FCL proved to be good enough for focusing particles.

#### Glass-silicon-glass device and n = 1, 2 resonance mode

The glass capillary was used to demonstrate the focusing of PS particles using the FCL and to quantify the performance increase of the system. To further elucidate how powerful the FCL is, a glass-silicon-glass device was used. The glasssilicon-glass device demonstrates that the FCL is capable of not only finding an improved excitation frequency for other setups than glass capillaries and PS particles, but can furthermore focus biological cells in an n = 1 resonance mode as detailed in eqn (5). n > 1 resonance modes are however also important in many processes. By altering the frequency range, and only the frequency range, the FCL is able to define the optimal frequency for resonance modes n> 1, such as the n = 2 resonance mode shown in Fig. 1(f) and (g). This is made possible, as the FCL is searching for the narrowest LW, and only chooses to look at one line, thus enabling this feature of the FCL. Therefore it is not surprising that the LWs for the resonance modes n = 1and n = 2 are approximately the same, Fig. 4(d) 31.28  $\pm$  0.98  $\mu$ m compared to (f) (L) 33.01 ± 5.27  $\mu$ m and (R) 36.15 ± 1.00 μm, which also corresponds to the line width found for the 5  $\mu$ m PS particles with a flow rate of 100  $\mu$ L min<sup>-1</sup> (31.80  $\pm$  0.58 μm). Another interesting readout to point out from Fig. 4(e) and (g) is that, by keeping the total cell number constant, the pixel value of the light intensity drops by half if the mode is increased from n = 1 (e) to n = 2 (g) which is as expected as there are half as many cells per modal line, and indicates that the FCL could be used to read out sample concentration. Another value to note is that in Fig. 1(c) the pixel value is 255, indicating a pure white line, which is expected as the video shows a stream of white in the middle. The cells, which are too few in number to form a continuous line in the channel even when focused, and therefore leave black spots in between the individual cells, have a much smaller average pixel value. Thus indicating that using higher concentrations is beneficial as this suppresses background noise, while at the same time allows for a higher sample concentrations and thus results in a higher sample throughput.

#### Conclusions

The FCL was compared against 3 experiments previously conducted in our lab  $^{43}$  and demonstrates a superior performance compared to experienced scientists in 1) reducing the linewidth by at least a factor of two for 5  $\mu$ m PS particles, 2) increasing the range of flow rates in which 1  $\mu$ m diameter PS particles could be focused and 3) identifying an excitation frequency for which 600 nm diameter PS particles could be focused. Furthermore, the FCL is capable of manipulating biological cells and resonance modes of n greater than 1.

These results indicate that by replacing the human operator by an optical FCL, *ceteris paribus*, the performance of a device can be augmented by automating otherwise laborious tasks. This replacement increases the device reliability without increasing the production complexity, increases the throughput, overall yields narrower focusing and focusing of 600 nm diameter PS particles can be

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achieved. A limitation, was that for small particles and low flow rates, such as 1 μm diameter PS particles at 5 μL min<sup>-1</sup> and 10 µL min<sup>-1</sup>, particles were trapped at the piezoelectric transducers. This could be prevented by adapting the code to search for an optimal excitation frequency and then placing a frequency offset on the found frequency as the ARF decreases with sub-optimal frequency selection. This idea could also be used to find an excitation frequency in which all particles are focused, and then placing a small offset in order to separate the particles by size. A further limitation was that particles below 600 nm diameter could not be focused. Provided an advanced imaging system with a higher resolution, we believe that even smaller particles could be focused. Research groups with devices already mounted on high resolution optical systems could for instance use the screen capture functionality function as an adapter for their optical system and test this hypothesis.

The FCL relies on an optical input to minimize the particle linewidth. The frequency thus chosen and continuously adapted by the FCL is therefore not necessarily the exact resonance frequency of the system, which sets the presented FCL apart from previously reported FCLs. As the FCL requires no additional equipment, provided that the setup has an optical readout and the function generator can be controlled by a computer, we thus present a cost-effective, high-throughput and automated method to rapidly increase the efficiency of established systems used for particle focusing. The source code for the FCL is openly accessible; ideas and augmentations from the community are welcomed.

#### Conflicts of interest

There are no conflicts to declare.

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