Lab on a Chip



View Article Online

PAPER

Check for updates

Cite this: Lab Chip, 2021, 21, 3378

Received 6th April 2021, Accepted 21st June 2021

DOI: 10.1039/d1lc00292a

rsc.li/loc

Introduction

Elucidating the origins, development and fate of cellular systems is at the forefront of biological enquiry. Increasingly single cell next generation sequencing (NGS) profiling is used to provide a comprehensive map of the cellular population linked to each cell's underlying processes and their role in system biology. In essence, these experiments involve compartmentalising single cells with reporter beads to capture and encode a cell's biological properties prior to delivery to a NGS and bioinformatics pipeline. Cell and bead co-encapsulation requires small volume liquid handling, a central strength of microfluidics.¹ First, elastomeric Quake valves² were used to compartmentalise single cells within an addressable array, marking the beginnings of single-cell diversity screens.^{3,4} Dramatic increases in throughput (*e.g.*

Dual dean entrainment with volume ratio modulation for efficient droplet co-encapsulation: extreme single-cell indexing[†]

Jack Harrington, ^(D)^a Luis Blay Esteban, ^(D)^b Jonathan Butement,^a Andres F. Vallejo, ^(D)^c Simon I. R. Lane,^{bd} Bhavwanti Sheth,^e Maaike S. A. Jongen, ^(D)^a Rachel Parker,^a Patrick S. Stumpf,^f Rosanna C. G. Smith,^a Ben D. MacArthur,^{dg} Matthew J. J. Rose-Zerilli,^{ad} Marta E. Polak, ^(D)^{cd} Tim Underwood^{ad} and Jonathan West ^(D)*^{ad}

The future of single cell diversity screens involves ever-larger sample sizes, dictating the need for higher throughput methods with low analytical noise to accurately describe the nature of the cellular system. Current approaches are limited by the Poisson statistic, requiring dilute cell suspensions and associated losses in throughput. In this contribution, we apply Dean entrainment to both cell and bead inputs, defining different volume packets to effect efficient co-encapsulation. Volume ratio scaling was explored to identify optimal conditions. This enabled the co-encapsulation of single cells with reporter beads at rates of \sim 1 million cells per hour, while increasing assay signal-to-noise with cell multiplet rates of \sim 2.5% and capturing \sim 70% of cells. The method, called Pirouette coupling, extends our capacity to investigate biological systems.

10 000's of cells) emerged from using nanowell arrays^{5–7} and droplet microfluidics.^{8,9}

Following these pivotal technology developments, singlecell analysis is gaining pace with the biology community aiming to decipher ever-larger cellular systems. Cell atlas reference maps, CRISPR and compound library single-cell screening projects typify this trend, with the scalability of the continuous flow droplet microfluidics format suitable for matching such large experiments.¹⁰ However, efforts in this direction face a fundamental problem: cells are randomly encapsulated. The probability of cells being encapsulated in a droplet is described by the Poisson statistic;

$$P_{(x)} = \frac{\lambda^{x} e^{-\lambda}}{x!}$$

where $P_{(x)}$ is the probability of x number of cells being packaged in a droplet and λ the mean number of cells per droplet. Dilute cell suspensions ($\lambda < 0.1$) are used to reduce the probability of co-encapsulating multiple cells and prevent their biology being scrambled during barcoding. As a consequence throughput is greatly limited and cell multiplets cannot be completely excluded, resulting in a trade-off between throughput and analytical noise: for most experiments a signal-to-noise of >20 (<5% multiplet rate,¹¹ see ESI,† Table S1) is acceptable which drastically limits cell concentrations. To compound the problem, solid reporter beads are also delivered as dilute suspensions to avoid clogging. This collides two Poisson statistics as a joint

^a Cancer Sciences, Faculty of Medicine, University of Southampton, UK.

E-mail: J.J.West@soton.ac.uk

^b Faculty of Engineering and Physical Sciences, University of Southampton, UK ^c Clinical and Experimental Sciences, Sir Henry Wellcome Laboratories, Faculty of Medicine, University of Southampton, UK

^d Institute for Life Sciences, University of Southampton, UK

^e School for Biological Sciences, Faculty of Environmental and Life Sciences, University of Southampton, UK

^f Centre for Human Development, Stem Cells and Regeneration, Faculty of Medicine, University of Southampton, UK

^g Mathematical Sciences, University of Southampton, UK

[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/ d1lc00292a

probability distribution (JPD) in which cell and bead coupling rates are necessarily low (<1% of droplets, see ESI,† Fig. S1). Alternatively, hydrogel beads can be delivered to most droplets (50–95%) using packed flows. This reduces the problem to a single Poisson statistic allowing the majority of cells to be captured, albeit necessitating lower droplet generation rates, ultimately producing equivalent throughput to solid bead systems. The general problem of single cell encapsulation has attracted great interest, resulting in a wide variety of passive (trapping,^{12,13} deterministic¹⁴ and inertial ordering^{15–17}) and active (electrical,^{18–20} magnetic,²¹ optical,²² acoustic²³ and mechanical²⁴) droplet generation and encapsulation strategies that have recently been expertly reviewed by Ling *et al.*²⁵

Passive approaches harness flow and channel properties, with the benefit of being easy to operate. In particular, inertial microfluidic formats²⁶⁻²⁸ enable high throughput and can produce entrainment effects to offer the enticing possibility of the periodic delivery of cell and beads, allowing deterministic packaging into droplets to free assays from the limitations of the Poisson statistic. During the high velocity transport (Re > 1) of particle-laden flows where particle diameters approach microchannel dimensions $(a/D_{\rm h} > 0.07)$ particle interaction with the underlying flow field can be predicted by the particle Reynolds number, $\text{Re}_{p} = \text{Re}(a/D_{h})^{2} \ge$ 0.1.^{27,29,30} In this regime the parabolic velocity profile introduces an appreciable shear-gradient lift force that becomes countered by the wall effect lift force to produce an equilibrium position, focusing the particles within the same streamlines. This has the effect to increase the local particle concentration resulting in particle trains³¹ with the interplay between viscous disturbance and inertial lift forces producing an equilibrium defining the inter-particle spacing.³² Using

straight channels these principles have been applied to the formation of *'microfluidic crystals*^{,33} and deterministic cell encapsulation in droplets.^{16,34} With the use of high Reynolds number flows a single equilibrium position can be produced³⁵ to aid entrainment.

Introducing secondary Dean flows (De = $\text{Re}(D_{\rm h}/R)^{1/2}$, where *R* is the channel radius of curvature) created by high velocity transport in curved channels increases migration to attain efficient particle focusing and train formation^{15,17} while operating at slower flow regimes compatible with droplet generation. Curved channels also allow a single, inner-wall, focusing position^{28,36-38} to improve entrainment and avoid additional asynchronous trains in other streamlines which prevent reliable single cell and bead droplet encapsulation. With the benefits of curvature, spiral channels have been used to increase solid bead droplet loading to enhance cell capture efficiencies from \sim 5% to 20%.³⁹ This also introduces gains in throughput, but remains limited to dilute cell suspensions as low multiplet cell encapsulation is governed by the Poisson statistic describing random cell delivery. Entraining both beads and cells has also proved challenging, with one approach requiring dilute cell suspensions to reduce multiplets.40 Consequently, train frequency and length is reduced and many cells are randomly distributed within the flow. New directions are needed to realise the full potential of entrainment for the high throughput coencapsulation of single cells with reporter beads.

In this study, we have implemented the spiral channel approach for the Dean entrainment of both beads and cells. With reporter beads and cells having dissimilar sizes (\emptyset 30 μ m v. ϑ 10–15 μ m), we reasoned that each requires tailored microfluidic dimensions and flow velocities. A two-layer prototype was developed to deliver effective entrainment

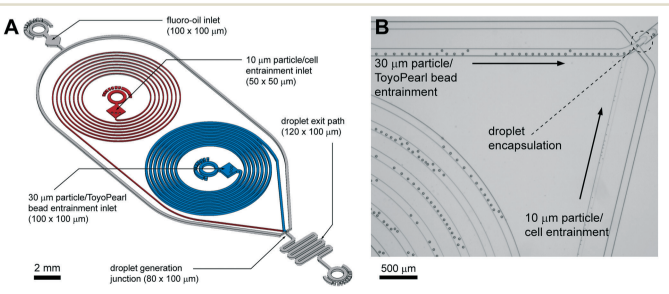


Fig. 1 Pirouette coupling microfluidics circuit for dual Dean entrainment and droplet generation (to scale, (A)). Two-layer fabrication is used to effect entrainment of the dissimilarly-sized reporter beads and cells; spiral channels with a height of 50 µm were used for cell entrainment, and channels 100 µm in height were used for bead entrainment and droplet generation. Example of Dean entrainment of 30 µm and 10 µm polystyrene particles for efficient droplet co-encapsulation (B).

Paper

conditions for the solid reporter beads and mammalian cells (Fig. 1). The volumetric ratio between beads and cells was investigated to identify operating windows for highly efficient co-encapsulation, surpassing the state of the art: we call the approach Pirouette coupling, a technique enabling the large-scale expansion of single-cell experiments.

Materials and methods

Design

Following Dean entrainment principles we designed a twolayer microfluidic prototype that is illustrated in Fig. 1 and provided as a CAD file (SI CAD). The prototype incorporates 50 μ m wide and high spiral channels for cell entrainment, and 100 μ m wide and high spiral channels for ToyoPearl bead entrainment. These high confinement channels ($a/D_h =$ 0.2 and 0.33, respectively) with a minimal aspect ratio promote single equilibrium focusing and single-file particle transport. These channels combine at an 80 μ m-wide droplet generation junction adjoining a 120 μ m-wide droplet exit channel. Each spiral channel has 6 turns with a radius minimum of 1.6 mm and maximum of 3.2 mm to produce an overall length of ~100 mm.

Fabrication and assembly

Pirouette coupling devices were fabricated by standard SU-8 photolithography, followed by replication in PDMS. Inlet and outlet ports were prepared using a 1-mm-diameter biopsy punch (Miltex), and then the device was oxygen plasma bonded (Femto, Deiner) to a glass microscope slide. Surfaces were functionalised by flooding the device with 1% (v/v) trichloro (1*H*,1*H*,2*H*,2*H*-perfluorooctyl)silane (Merck) in HFE-7500TM (3MTM). Plug and play interconnection between 25G needles on the syringes and the device inlets was achieved using polythene tubing (Smiths Medical, ID 0.38 mm; OD 1.09 mm).

Particles, beads and cells

Monodisperse 10-µm-diameter (CV 3.1%) polystyrene particles (Merck) were suspended in PBS, and monodisperse 20- and 30-µm-diameter (CV 3.0% and 3.2%, respectively) polystyrene particles (Merck) and filtered $\leq 40 \ \mu m$ ToyoPearl beads (HW-65S, Tosoh Biosciences, unfunctionalized ChemGene beads) were suspended in filtered, modified DropSeq lysis buffer⁹ (100 mM Tris, pH 7.5, 0.1% sarkosyl, 10 mM EDTA). Human THP-1 and HEK293 cells (ATCC®) were washed and resuspended in filtered PBS with 1% (w/v) BSA. Particle, bead and cell diameter histograms are provided in the ESI†, Fig. S2. Particles, beads and cells were retained in suspension using a vertically orientated syringe with a PTFEcoated samarium cobalt disc magnet rotated at low rpm (\leq 30 a.u., Multi Stirrus™, V&P Scientific) (ESI†, Fig. S3). To avoid particles and beads occluding the channel during high concentration delivery, important instructions are provided in the ESI† Appendix I. This method allows the prolonged

Microfluidics

For the generation of 600 pL (CV < 2%) droplets at ~1800 Hz a QX200 (BioRad) fluoro-oil flow rate of 165 μ L min⁻¹ was used with a total, bead and cell, aqueous flow rate of ~65 μ L min⁻¹. Flow details for volume ratio scaling are provided in ESI† Table S2. High-speed microscopy (Miro Lab310, Vision Research) was used to image entrainment and droplet encapsulation. Video files were pre-processed in ImageJ or directly analysed using custom MATLAB scripts for measuring inter-particle pitch (ESI† Appendix II) and encapsulation (Appendix III). Encapsulation results were verified manually and compared with the joint probability distribution (JPD) based on the random arrival of beads and cells for packaging into droplets as a control metric.

Metric definitions

The signal-to-noise (S:N), multiplet rate (MR), throughput (TP) and capture efficiency (CE) performance metrics describing droplet co-encapsulations are described by:

$$\begin{split} \mathbf{S:N} &= \frac{(\mathbf{BC} + \mathbf{BB} + \mathbf{C})}{(\mathbf{BCC}^+ + \mathbf{BB}^+\mathbf{CC}^+)} \\ \mathbf{MR}(\%) &= 100 \times \frac{(\mathbf{BCC}^+ + \mathbf{BB}^+\mathbf{CC}^+)}{(\mathbf{BC} + \mathbf{BB}^+\mathbf{C} + \mathbf{BCC}^+ + \mathbf{BB}^+\mathbf{CC}^+)} \\ \mathbf{TP} &= \frac{(\mathbf{BC} + \mathbf{BB}^+\mathbf{C})}{\text{interval}} \end{split}$$

$$CE(\%) = 100 \times \frac{(BC + BB^+C)}{C_{\text{total}}}$$

where *B* denotes a 30 μ m polystyrene particle or ToyoPearl bead per droplet, *C* denotes a 10 μ m polystyrene particle or cell, BB⁺ denotes 2 or more particles or beads, CC⁺ denotes 2 or more 10 μ m particles or cells, and *C*_{total} denotes all 10 μ m particles or cells delivered to droplets.

Results and discussion

A first requirement for the high throughput analysis of single cells in droplets is the sustained and homogeneous delivery of high concentration bead and cell suspensions. This is especially the case for the large and dense reporter beads which rapidly sediment yet are also fragile. Initially this challenge was solved using perpetual sedimentation (rotation of a horizontal syringe to produce bead orbits, effectively making them neutrally buoyant).⁴¹ Then, to aid broader uptake in conventional cell biology labs we instead used a vertically orientated syringe with a powerful electromagnet to

gently rotate a disc magnet: the disc magnet disperses beads and cells in all directions, sideways for mixing, upwards to return by gravity and downwards to exit. Both approaches enabled the sustained delivery of cell and particle suspensions suitable for investigating Dean entrainment effects (ESI[†], Fig. S3).

To gain a first understanding of entrainment we employed monodisperse 30 µm polystyrene particles to represent ToyoPearl reporter beads and monodisperse 10 µm polystyrene particles to represent mammalian cells. The emergence of particle entrainment, from disordered to periodic spacing was observed with a 600 000 $mL^{^{-1}}$ 30 μm particle suspension using a 100 mm s⁻¹ mean flow velocity (Rep 0.9, ESI⁺, Fig. S4A). Entrainment requires concentrated suspensions with particle train length increasing and interparticle pitch decreasing with concentration. At 1 million per mL (1.4% volumetric fraction (vf)) a median inter-particle pitch of 75 µm was produced (Fig. 2A). This equates to 2.5D (D = particle diameter) arrangements associated with higher Re_p numbers³² and is attributed to the high volume fraction suspension and prolonged inertial transport (100 mm) supplemented with secondary Dean flows. The concentration could be extended to 1.5 million per mL (2.1% vf), but above this crowding effects reduce periodicity (ESI⁺, Fig. S4B). Entrainment of the 10 μ m polystyrene (\bar{U} = 100 mm s⁻¹, Re_p 0.2) particles was also concentration-dependent, with striking ordering observed at 6 million per mL (0.3% vf, Fig. 2B) producing a median 5D pitch. At these moderate particle concentrations, the increased gaps between trains extends the pitch, and pitch variability. Higher volumetric fractions are feasible. However, in the context of cell processing, such concentrations are unsuitable for maintaining cell viability and promote cell clustering.

The velocity dependence was investigated using flow rates ranging from 1–90 μ L min⁻¹, producing a velocity range of 1.7–150 mm s⁻¹ (Re_p 0.015–1.35, De_{max} 0.04–3.75) for the 30 μ m particles and 6.7–600 mm s⁻¹ (Re_p 0.01–1.2, De_{max} 0.06–5.30) for the 10 μ m particles. 30 μ m particles were tightly entrained with a 2.5*D* pitch throughput the 30–90 μ L min⁻¹ flow range (Fig. 2A), and similarly 10 μ m particles entrained with a 5*D* pitch throughout the 5–30 μ L min⁻¹ flow range (Fig. 2B). At lower flow rates entrainment quality diminishes, ultimately leading to randomly distributed particles with different velocities.

The inter-particle pitch results can be used to predict 30 and 10 μ m particle volume limits for effective single bead and single cell co-encapsulation: using the 25th percentile data, volumes below 650 pL are needed for 30 μ m particles and volumes below 100 pL for 10 μ m particles to obtain efficient co-encapsulations. This requires different bead and cell flow rates, producing a volume ratio (VR) to provide a first approximation to inform flow rates for efficient co-encapsulation and droplet volumes ($V_{\text{bead}} + V_{\text{cell}}$) for optimal throughput:

$$VR = \frac{V_{bead}}{V_{cell}} = \frac{\sigma 2.5 D_{bead} wh}{\sigma 5 D_{cell} wh}$$

where *D* denotes the diameter and σ a length fraction defining bead or cell entrainment statistics, and *w* and *h* are

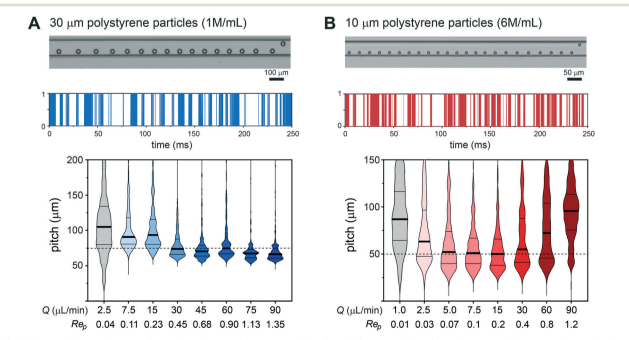


Fig. 2 Scalable particle entrainment. Exemplary frames of 30 μ m particle (A) and 10 μ m particle (B) entrainment with a mean flow velocity of 100 mm s⁻¹. Entrainment intensity profile with distance translated to time of a 250 ms imaging segment illustrating gap and train length variability. Violin plots of the velocity dependence (flow rate and Re_p) of the particle pitch distribution represented as median, 25th and 75th percentiles and data extremities with cut-offs at 200 μ m for 30 μ m particles and 150 μ m for 10 μ m particles (n > 2000 particles per velocity condition). The 2.5*D* and 5*D* inter-particle pitch predictions are indicated with dashed lines. Grey violin plots denote random particle distributions, without entrainment.

Paper

the channel cross-section dimensions. Using these volumes as a guide we introduced flow conditions for the generation of 600 pL droplets. To identify optimal 30 µm particle (referred to as 'beads') and 10 µm particle (referred to as 'cells') input volumes for each droplet, a volume ratio scaling experiment was undertaken: the 'bead' and 'cell' flow rates $(Q_{\text{bead}} \text{ and } Q_{\text{cell}})$ were differentially modulated to produce volume ratios ranging from 1 to 15 (300bead + 300_{cell} pL to 562_{bead} + 38_{cell} pL) while satisfying the requirements for effective Dean entrainment. The coencapsulation results are compared with theoretical results from the joint probability distribution (JPD) in Fig. 3A. With large 'cell' volumes dual Dean entrainment has a substantially reduced single 'cell' and 'bead' coupling frequency due to entrainment increasing the 'cell' multiplet rate. Larger 'bead' volumes also result in 'bead' doublets and multiplets. While cell multiplets are unwanted, bead multiplets are acceptable, only having the effect to (i) randomly oversample cell types and biology without affecting population structure and (ii) use additional beads and read space. At a volume ratio of 10 (545 + 55 pL) a transition occurs in which entrainment becomes beneficial, with the noise ('cell' multiplets) dropping below the theoretical JPD value to produce a signal-to-noise of ~16. As the volume ratio further increases, the 'cell' multiplet rate tends to zero, while the signal (single 'cell' capture rate) remains similar to the JPD results, allowing high throughput and extreme signal-to-noise processing.

To appreciate the different performance metrics the signal-to-noise, throughput and capture efficiency are compared with JPD predictions in Fig. 3B-D. These results demonstrate the merits of dual Dean entrainment; with a volume ratio of 15 the signal-to-noise is 139 (0.7% multiplets), 17-fold higher than random delivery, along with a throughput of 0.5 million per hour and a capture efficiency >50%. The gains in capture efficiency above the JPD prediction result from bead entrainment that produces higher numbers of droplets containing a bead. The 6 million per mL 'cell' concentration has a low volumetric fraction (0.3%), indicating scope for higher cell concentrations. However, higher concentrations introduce localized crowding effects such that gains in throughput are at the expense of the signal-to-noise (Fig. 3E and F). To increase the capture efficiency larger droplet volumes were considered. This allows >70% of droplets to contain a bead, enabling >70%of 'cells' to be captured (see Fig. 3G). However, this would require a higher volume ratio, producing lower 'cell' flow rates that are insufficient for effective entrainment. The ESI[†] video documents dual Dean entrainment for the coencapsulation of periodically spaced 'cell' and 'bead' trains into droplets. Ideal results are shown in Fig. 3H and typical results in ESI[†] Fig. S5.

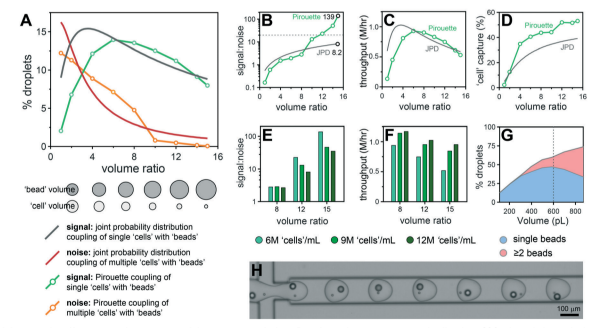


Fig. 3 Volume ratio effects on polystyrene particle co-encapsulation. Droplet volumes were standardized at 600 pL and the ratio between cell and bead volumes modulated between 1:1 and 1:15 (300:300 pL to 38:562 pL). The monodisperse 10 μ m particles are used to approximate cells and denoted as 'cells', and monodisperse 30 μ m particles approximating the ToyoPearl beads denoted as 'beads'. Pirouette coupling results are plotted with the theoretical JPD for given volume ratios with a 'bead' concentration of 1.2 million per mL and a 'cell' concentration of 6 million per mL. The percentage of droplets producing a signal (BC, BB⁺C) and noise (BCC⁺, BB⁺CC⁺) was obtained by analyzing ~3700 droplets per condition (A). The volume ratio effect on signal-to-noise with the S:N 20 threshold indicated using a grey dashed line (B), throughput (C) and capture rate (D). Increasing the 'cell' concentration from 6 million per mL to 9 million per mL and 12 million per mL reduces the signal-to-noise (E) while increasing the throughput (F). Data obtained by analyzing ~1850 droplets per condition. Increasing the droplet volume from 110 to 890 pL results in higher proportions of droplets containing multiple 'beads' (G). Data obtained by analyzing ~3000 droplets for each volume condition. Frame documenting an ideal single 'cell' and single 'bead' co-encapsulation sequence using a volume ratio of 1:12 (H).

Given the promising performance with polystyrene particles we next sought to answer whether dual Dean entrainment can be effectively applied to the co-ToyoPearl beads encapsulation of (unfunctionalised ChemGene beads used in the Drop-seq protocol) with mammalian cells. Both ToyoPearl beads (Ø34.1 µm, CV 9.6%) and human HEK293 cells (Ø14.3 µm, CV 10.6%) were effectively entrained at the same concentrations used for the polystyrene particle experiments demonstrating that the entrainment conditions can manage particles and cells with low monodispersity (ESI[†], Fig. S4C and S6). In addition, the shear flow conditions rapidly disperse cell clusters into trains of single cells, potentially representing a means for sample disaggregation (ESI† Fig. S7). The bead and cell flow rate dependent pitch distributions closely followed those obtained for the 10 and 30 µm polystyrene particles, although the HEK293 cells produced a 3.5D median pitch (10 µm particles = 5D). The results from a volume ratio scaling coencapsulation experiment are documented in Fig. 4A-E. Noise reduction beneath the JPD prediction occurred later with a flow ratio of 16 and stabilized with a flow ratio ≥ 20 , overall resulting in a modest signal-to-noise (24; 4% multiplet rate). Throughput was maintained at ~0.5 million cells per hour, and the capture efficiency was extended to $\sim 70\%$ by using a 1.5 million per mL ToyoPearl bead concentration. Performance was corroborated by repeating the experiment with a THP-1 cells (ESI† Fig. S8). These smaller, polydisperse cells (\emptyset 11.0 μ m, CV 24.7%) eliminate size effects being causative of the reduced signal-to-noise.

We sought to understand the late onset of noise reduction and absence of exponential signal-to-noise scaling with increasing volume ratio. The 10 µm polystyrene particles and HEK293 cells produce equivalent pitch minima (~20 µm), but distinctly different focusing behavior (Fig. 2B and 4F and ESI† Fig. S6 and S9): the solid 10 µm particles are wallfocussed for all flow rates excepting 1 $\mu L\ min^{-1}$ in which focusing collapses, producing random cross-channel positions. In stark contrast, cells are deformable with inner wall focusing only occurring above 30 μ L min⁻¹ (Re_p > 1, ESI† Fig. S7D), below which HEK293 cells become entrained within streamlines towards the channel centre. Migration to equilibrium positions towards the channel center is a likely consequence of a deformation-induced lift force which counters the wall-directed shear-gradient lift force.27,42 However, a full description of cell focusing and entrainment in spiral channels requires further investigation to reliably predict behavior. In addition, operation at these lower flow velocities also reduces the De number, further directing the equilibrium position to the channel center.37 At still lower velocities, required to produce high volume ratios, cells occupy random, unfocussed positions. Cell transport within different streamlines, with different velocities, increases the

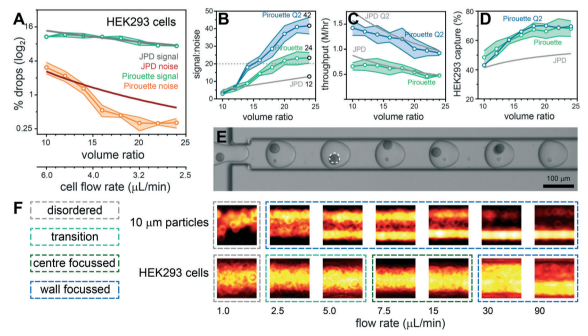


Fig. 4 Volume ratio effects on the co-encapsulation of ToyoPearl beads with HEK293 cells. Droplet volumes were standardized at 600 pL and the ratio between cell and bead volumes modulated between 1:10 and 1:24 (55:545 pL to 24:576 pL). Pirouette coupling results are plotted with the theoretical JPD for given volume ratios with a ToyoPearl bead concentration of 1.5 million per mL and a HEK293 cell concentration of 6 million per mL. The percentage of droplets producing a signal (BC, BB⁺C) and noise (BCC⁺, BB⁺CC⁺) was obtained by analyzing >3500 droplets per condition (n = 3 experiments) (A). The volume ratio effects on signal-to-noise with the S:N 20 threshold indicated using a grey dashed line (B), throughput (C) and capture rate (D) are plotted for standard (Pirouette; green) and double (Pirouette Q2; blue) aqueous flow rates. Frame documenting an efficient single HEK293 cell and single bead co-encapsulation sequence using a volume ratio of 1:16 (E). A dashed circle is used to identify a cell masked by a bead. The 10 μ m polystyrene particles and HEK293 cells have different flow rate-dependent inertial focusing behaviors (F).

Paper

probability of cells arriving together at the droplet generation junction. To achieve higher signal-to-noise sample processing, higher Rep flow conditions are needed for effective cell focusing and entrainment. We investigated the upper limits of the dripping droplet formation regime: aqueous flow rates can be doubled in combination with a 240 µL min⁻¹ oil flow rate while retaining a 600 pL droplet volume. Repeating the volume ratio scaling experiment using these elevated flow conditions improved the signal-to-noise to 42 (2.3% multiplets) with the benefit of doubling the throughput to ~ 1 million cells per hour ($\sim 15\,000$ /minute) while retaining the \sim 70% capture efficiency (Fig. 4B-D). In comparison, entrainment approaches without volume ratio considerations, had throughput limited to 2700 cells per minute³⁹ (ESI[†] Table S1). Higher flow regimes enter a jetting regime with bead-triggering producing higher droplet generation rates for even higher throughput processing.⁴³ However, bead-triggered droplet formation in the jetting regime increases droplet polydispersity, at odds with precisely defined volumes required to effectively coencapsulate single cells by entrainment.

Pirouette coupling out-performed commercial and entrainment-based single cell and reporter bead coencapsulation methods (ESI† Fig. S10, Table S1). Alternative approaches bypass the Poisson-dictated multiplet rate problem by pre-indexing cells by membrane (MULTI-seq⁴⁴) or transcriptome labelling (sci-seq,^{45,46} SPLiT-seq⁴⁷ and scifi-RNA-seq⁴⁸). The scifi-seq method was used to allow droplet 'super-loading', demonstrating a throughput of >150000 nuclei per 10× channel (>500000 nuclei per hour). Each technique has its own deficiencies, such as lengthy procedures, labels being exchanged, cell losses during labelling and volume limitations restricting analyses to nuclei (foregoing the information content from the rest of the cell). Nevertheless, substantial improvements can readily be anticipated in these and other approaches for single cell indexing. Indeed, the current Pirouette coupling prototype represents a blueprint for future iterations incorporating refinements to microfluidic dimensions allowing, for instance, improved focusing (ESI⁺, Fig. S4D) and operation at higher Re_p numbers for enhanced signal-to-noise processing. In addition, bead alternation resulting from overcrowding highlights the need for improved methods for the delivery of high volume fraction particle suspensions with minimal concentration deviations throughout processing. This will reduce crowding events and associated particle alternation, while increasing train length and assay throughput. In general, these technological developments forecast the routine undertaking of large-scale experiments that will become feasible as dramatic cost savings begin to emerge from innovations in sequencing.49

The co-encapsulation efficiencies enabled by Pirouette coupling allow other analytical scenarios to be envisaged, such as experiments requiring cells to be rapidly processed to prevent transcriptome remodeling, those involving different beads reporting different biological dimensions, or a bead to perturb the cell and another bead to report biological outcomes. For example, Pirouette coupling offers the potential for screening genetically-encoded bead-based compound libraries without exhaustive passes to ensure library coverage. Here, the ability of Dean entrainment to process high concentrations of solid beads allows the repertoire of solid phase synthesis methods to be used in library construction. Overall, the co-encapsulation efficiencies lend Pirouette coupling to large-scale experiments that were previously impractical.

Conclusions

Pirouette coupling combines cell and bead entrainment to bypass the limitations of the joint probability distribution during droplet co-encapsulations. This produces profound gains in performance, achieving extreme throughput combined with an enhanced signal-to-noise while capturing the majority of cells. The approach has broad-reaching potential, enabling cellular systems to be comprehensively profiled in health, disease and in response to perturbation.

Author contributions

JH, LBE, JB, AFV, SIRL, BS, MSAJ, RP, PSS, RCGS, MJJRZ and JW undertook the experiments, BDM, MJJRZ, MEP, TU and JW supervised the project, and JW wrote the manuscript, with co-authors reviewing the manuscript.

Conflicts of interest

The authors declare no conflicts of interest.

Acknowledgements

We are indebted to Deborah Mackay for her support. We also thank Gessa Sugiyarto for preparing cells. The project was made possible by financial contributions from the Medical Research Council (JB and MSAJ: MC_PC_15078), a CRUK & Royal College of Surgeons of England Advanced Clinician Scientist Fellowship (JH and TU), an EPSRC CASE studentship (LBE, 1658462), GSK (AFV, ARCP006668), a Sir Henry Dale Wellcome Trust Fellowship (MEP; 109377/Z/15/Z) and a John Goldman Fellowship (MJJRZ; 2016/JGF/0003).

References

- 1 T. A. Duncombe, A. M. Tentori and A. E. Herr, *Nat. Rev. Mol. Cell Biol.*, 2015, **16**, 554.
- 2 M. A. Unger, H.-P. Chou, T. Thorsen, A. Scherer and S. R. Quake, *Science*, 2000, **288**, 113–116.
- 3 J. S. Marcus, W. F. Anderson and S. R. Quake, *Anal. Chem.*, 2006, **78**, 3084–3089.
- 4 A. M. Streets, X. Zhang, C. Cao, Y. Pang, X. Wu, L. Xiong, L. Yang, Y. Fu, L. Zhao, F. Tang and Y. Huang, *Proc. Natl. Acad. Sci. U. S. A.*, 2014, **111**, 7048–7053.
- 5 H. C. Fan, G. K. Fu and S. P. A. Fodor, *Science*, 2015, 347, 1258367.

- 6 T. M. Gierahn, M. H. Wadsworth, T. K. Hughes, B. D. Bryson, A. Butler, R. Satija, S. Fortune, J. C. Love and A. K. Shalek, *Nat. Methods*, 2017, 14, 395–398.
- 7 X. Han, R. Wang, Y. Zhou, L. Fei, H. Sun, S. Lai, A. Saadatpour, Z. Zhou, H. Chen, F. Ye, D. Huang, Y. Xu, W. Huang, M. Jiang, X. Jiang, J. Mao, Y. Chen, C. Lu, J. Xie, Q. Fang, Y. Wang, R. Yue, T. Li, H. Huang, S. H. Orkin, G.-C. Yuan, M. Chen and G. Guo, *Cell*, 2018, 172, 1091–1107. e1017.
- 8 A. M. Klein, L. Mazutis, I. Akartuna, N. Tallapragada, A. Veres, V. Li, L. Peshkin, D. A. Weitz and M. W. Kirschner, *Cell*, 2015, 161, 1187–1201.
- 9 E. Z. Macosko, A. Basu, R. Satija, J. Nemesh, K. Shekhar, M. Goldman, I. Tirosh, A. R. Bialas, N. Kamitaki, E. M. Martersteck, J. J. Trombetta, D. A. Weitz, J. R. Sanes, A. K. Shalek, A. Regev and S. A. McCarroll, *Cell*, 2015, 161, 1202–1214.
- 10 G. X. Y. Zheng, J. M. Terry, P. Belgrader, P. Ryvkin, Z. W. Bent, R. Wilson, S. B. Ziraldo, T. D. Wheeler, G. P. McDermott, J. Zhu, M. T. Gregory, J. Shuga, L. Montesclaros, J. G. Underwood, D. A. Masquelier, S. Y. Nishimura, M. Schnall-Levin, P. W. Wyatt, C. M. Hindson, R. Bharadwaj, A. Wong, K. D. Ness, L. W. Beppu, H. J. Deeg, C. McFarland, K. R. Loeb, W. J. Valente, N. G. Ericson, E. A. Stevens, J. P. Radich, T. S. Mikkelsen, B. J. Hindson and J. H. Bielas, *Nat. Commun.*, 2017, **8**, 14049.
- R. Salomon, D. Kaczorowski, F. Valdes-Mora, R. E. Nordon, A. Neild, N. Farbehi, N. Bartonicek and D. Gallego-Ortega, *Lab Chip*, 2019, **19**, 1706–1727.
- 12 M. Sauzade and E. Brouzes, Lab Chip, 2017, 17, 2186–2192.
- 13 M. Zhang, Y. Zou, X. Xu, X. Zhang, M. Gao, J. Song, P. Huang, Q. Chen, Z. Zhu, W. Lin, R. N. Zare and C. Yang, *Nat. Commun.*, 2020, **11**, 2118.
- 14 A. S. Mao, J.-W. Shin, S. Utech, H. Wang, O. Uzun, W. Li, M. Cooper, Y. Hu, L. Zhang, D. A. Weitz and D. J. Mooney, *Nat. Mater.*, 2017, **16**, 236–243.
- 15 D. Di Carlo, D. Irimia, R. G. Tompkins and M. Toner, Proc. Natl. Acad. Sci. U. S. A., 2007, 104, 18892–18897.
- 16 J. F. Edd, D. Di Carlo, K. J. Humphry, S. Koster, D. Irimia, D. A. Weitz and M. Toner, *Lab Chip*, 2008, 8, 1262–1264.
- 17 E. W. Kemna, R. M. Schoeman, F. Wolbers, I. Vermes, D. A. Weitz and A. van den Berg, *Lab Chip*, 2012, **12**, 2881–2887.
- 18 H. Gu, C. U. Murade, M. H. G. Duits and F. Mugele, Biomicrofluidics, 2011, 5, 011101.
- 19 D. R. Link, E. Grasland-Mongrain, A. Duri, F. Sarrazin, Z. Cheng, G. Cristobal, M. Marquez and D. A. Weitz, *Angew. Chem., Int. Ed.*, 2006, **45**, 2556–2560.
- 20 B. P. Nadappuram, P. Cadinu, A. Barik, A. J. Ainscough, M. J. Devine, M. Kang, J. Gonzalez-Garcia, J. T. Kittler, K. R. Willison, R. Vilar, P. Actis, B. Wojciak-Stothard, S.-H. Oh, A. P. Ivanov and J. B. Edel, *Nat. Nanotechnol.*, 2019, 14, 80–88.
- 21 S. Buryk-Iggers, J. Kieda and S. S. H. Tsai, *AIP Adv.*, 2019, 9, 075106.
- 22 S.-Y. Park, T.-H. Wu, Y. Chen, M. A. Teitell and P.-Y. Chiou, *Lab Chip*, 2011, **11**, 1010–1012.

- 23 D. J. Collins, T. Alan, K. Helmerson and A. Neild, *Lab Chip*, 2013, **13**, 3225–3231.
- 24 J.-H. Choi, S.-K. Lee, J.-M. Lim, S.-M. Yang and G.-R. Yi, *Lab Chip*, 2010, **10**, 456–461.
- 25 S. D. Ling, Y. Geng, A. Chen, Y. Du and J. Xu, *Biomicrofluidics*, 2020, 14, 061508.
- 26 D. Di Carlo, *Lab Chip*, 2009, **9**, 3038–3046.
- 27 H. Amini, W. Lee and D. Di Carlo, *Lab Chip*, 2014, 14, 2739–2761.
- 28 J. M. Martel and M. Toner, Annu. Rev. Biomed. Eng., 2014, 16, 371–396.
- 29 P. M. Holloway, J. Butement, M. Hegde and J. West, *Biomicrofluidics*, 2018, **12**, 044104.
- 30 D. Stoecklein and D. Di Carlo, Anal. Chem., 2019, 91, 296-314.
- 31 J.-P. Matas, V. Glezer, É. Guazzelli and J. F. Morris, *Phys. Fluids*, 2004, 16, 4192–4195.
- 32 S. Kahkeshani, H. Haddadi and D. Di Carlo, *J. Fluid Mech.*, 2016, **786**, R3.
- 33 W. Lee, H. Amini, H. A. Stone and D. Di Carlo, Proc. Natl. Acad. Sci. U. S. A., 2010, 107, 22413–22418.
- 34 T. P. Lagus and J. F. Edd, RSC Adv., 2013, 3, 20512.
- 35 C. Liu, G. Hu, X. Jiang and J. Sun, *Lab Chip*, 2015, 15, 1168–1177.
- 36 S. S. Kuntaegowdanahalli, A. A. S. Bhagat, G. Kumar and I. Papautsky, *Lab Chip*, 2009, 9, 2973–2980.
- 37 J. M. Martel and M. Toner, Sci. Rep., 2013, 3, 3340.
- 38 N. Nivedita, P. Ligrani and I. Papautsky, *Sci. Rep.*, 2017, 7, 44072.
- 39 H. S. Moon, K. Je, J. W. Min, D. Park, K. Y. Han, S. H. Shin, W. Y. Park, C. E. Yoo and S. H. Kim, *Lab Chip*, 2018, 18, 775–784.
- 40 L. Li, P. Wu, Z. Luo, L. Wang, W. Ding, T. Wu, J. Chen, J. He, Y. He, H. Wang, Y. Chen, G. Li, Z. Li and L. He, *ACS Sens.*, 2019, 4, 1299–1305.
- 41 S. I. R. Lane, J. Butement, J. Harrington, T. Underwood, J. Shrimpton and J. West, *Lab Chip*, 2019, **19**, 3771–3775.
- 42 S. C. Hur, N. K. Henderson-MacLennan, E. R. B. McCabe and D. Di Carlo, *Lab Chip*, 2011, **11**, 912–920.
- 43 I. C. Clark and A. R. Abate, *Lab Chip*, 2018, **18**, 3598–3605.
- 44 C. S. McGinnis, D. M. Patterson, J. Winkler, D. N. Conrad, M. Y. Hein, V. Srivastava, J. L. Hu, L. M. Murrow, J. S. Weissman, Z. Werb, E. D. Chow and Z. J. Gartner, *Nat. Methods*, 2019, 16, 619–626.
- J. Cao, J. S. Packer, V. Ramani, D. A. Cusanovich, C. Huynh, R. Daza, X. Qiu, C. Lee, S. N. Furlan, F. J. Steemers, A. Adey, R. H. Waterston, C. Trapnell and J. Shendure, *Science*, 2017, 357, 661.
- 46 J. Cao, M. Spielmann, X. Qiu, X. Huang, D. M. Ibrahim, A. J. Hill, F. Zhang, S. Mundlos, L. Christiansen, F. J. Steemers, C. Trapnell and J. Shendure, *Nature*, 2019, 566, 496–502.
- 47 A. B. Rosenberg, C. M. Roco, R. A. Muscat, A. Kuchina, P. Sample, Z. Yao, L. T. Graybuck, D. J. Peeler, S. Mukherjee, W. Chen, S. H. Pun, D. L. Sellers, B. Tasic and G. Seelig, *Science*, 2018, 360, 176.

View Article Online

- 48 P. Datlinger, A. F. Rendeiro, T. Boenke, T. Krausgruber, D. Barreca and C. Bock, bioRxiv, 2019, DOI: 10.1101/ 2019.12.17.879304.
- 49 Y. Qin, S. Koehler, S. Zhao, R. Mai, Z. Liu, H. Lu and C. Xing, bioRxiv, 2020, DOI: 10.1101/ 2020.12.10.418962.

Paper