High-speed particle detection and tracking in microfluidic devices using event-based sensing

Jessie Howell1, Tansy Hammarton2, Yoann Altmann3*, Melanie Jimenez1*

1 Biomedical Engineering Division, James Watt School of Engineering, University of Glasgow, Glasgow, G12 8LT, UK
2 Institute of Infection, Immunity and Inflammation, University of Glasgow, Glasgow, G12 8TA, UK
3 School of Engineering and Physical Sciences, Heriot-Watt University, Edinburgh Campus, Edinburgh, EH14 4AS, UK

Corresponding author: Melanie.Jimenez@glasgow.ac.uk

Abstract
Visualising fluids and particles within channels is a key element of microfluidic work. Current imaging methods for particle image velocimetry often require expensive high-speed cameras with powerful illuminating sources, thus potentially limiting accessibility. This paper explores for the first time the potential of an event-based camera for particle and fluid behaviours characterisation in a microfluidic system. Event-based cameras have the unique capacity to detect light intensity changes asynchronously and to record spatial and temporal information with low latency, low power and high dynamic range. Event-based cameras could consequently be relevant for detecting light intensity changes due to moving particles, chemical reactions or intake of fluorescent dyes by cells to mention a few. As a proof-of-principle, event-based sensing was tested in this work to detect and track 1 µm and 10 µm-diameter particles flowing in a microfluidic channel. Importantly, experiments were performed by directly connecting the camera to a standard fluorescence microscope, only relying on the microscope arc lamp for illumination. We propose a data processing strategy that allowed particle tracking in both bright-field and fluorescence imaging. Tracking was achieved for particle velocities up to 0.4 m.s⁻¹ demonstrating that event-based cameras could be a new paradigm shift in microscopic imaging.

Introduction
Event-based cameras emerged in the 1990s as neuromorphic vision sensors mimicking biological retinas (1). Unlike frame-based cameras, event-based cameras respond to, and only record, brightness changes (log intensity depicted as log(n) in Figure 1-A), asynchronously and independently for each pixel. When a change in brightness is detected at a given pixel, the event information is transmitted, that is, its (x,y) location on the pixel array, its time stamp and the sign/polarity of the change (increase (+1 in Figure 1-B) or decrease (-1 in Figure 1-B)).

Using current event-based cameras (available under 5k USD), events are detected with microsecond resolution. Since only events are transmitted, event-based cameras offer low latency, low power (ca. 10 mW) and high dynamic range (>120 dB) (2). These “silicon retinas” have become increasingly popular for high-speed robotic vision, e.g., for ball detection
(3)(4)(5), gesture recognition (6)(7), 3D mapping (8) or for unmanned aerial vehicles (9) and predator robots (10). They have also been used for tracking macroscopic objects such as vehicles (11) or stars/satellites (12)(13) (see the recent survey (28) which discusses applications and challenges of event-based sensing). Importantly, application to the "micro-world" is limited. The work proposed in (14) uses event-based cameras for micro-robotics, demonstrating tracking of microparticles in a petri dish while (15) focuses on neural activity imaging. In (16), particle tracking in a fluid-solid system has also been tested in a 5 cm ID pipe with 950 µm particles. The potential of event-based sensing for microfluidic applications remains consequently untapped.

Figure 1. Event-based detection of a particle over time. **A.** represents the light flux reaching a given pixel when a fluorescent particle passes through the pixel field of view. **B.** When the light intensity change (with respect to the last recorded event) exceeds a user-defined threshold, the camera records a new event whose polarity encodes the sign of the intensity change. **C.** Particle passing through the pixel field of view, generating light intensity changes depicted in A.

Due to their characteristic microscopic scale, microfluidic systems primarily rely on imaging technologies (such as microscopes and cameras) to monitor fluids and particles inside a channel. Imaging modalities offer considerable flexibility and find applications in quality control (e.g., detection of dust/bubble), performance evaluation (e.g., mixing, separation, detection) or in better understanding localised phenomena but limitations remain. Let's consider inertial microfluidics for example.

Inertial focusing devices have been widely used by the community for their unique capabilities to focus and separate particles based on size, shape and/or deformability (17). Despite advances in the field of computational inertial microfluidics (18), prototypes are often tailored to a targeted application following long design/test/optimise iterations to empirically explore the capabilities of new channel designs. Accurately imaging fluid and particle behaviour has become essential to assess the underlying physical phenomena and inform further design.
changes. Imaging inside inertial focusing devices (and microfluidics channels in general) typically relies on either long-exposure fluorescence or high-speed imaging.

Long-exposure fluorescence consists of imaging fluorescent particles over an extended period of time (long enough to be representative of the particle/fluid behaviour) and of building corresponding composite images (by stacking/integrating several images). This approach is particularly well suited to inertial devices with clear visualisation of "streaks" representing fluorescent particles that are focused at specific locations inside the channel. By mapping the intensity profile, an estimation of the focusing efficiency can be obtained. Long-exposure has been used with a wide range of particles (e.g., beads (19)-(20)(21)-(22)(23), cancer cells (24), bacteria (25)-(26)(27) or fungal cells (28) to mention a few) and has enhanced our understanding of the impact of channel geometry on focusing (20)-(21)-(29)-(30). Advantages of this imaging approach include a compatibility with standard fluorescent microscopes equipped with conventional cameras and relatively little data processing. Importantly however, the information extracted is reduced to global (i.e., population) behaviours. Moreover, high particle concentrations are usually required to ensure detectable fluorescent signals, hindering an in-depth understanding of single particle behaviour.

On the other hand, high-speed imaging unveils other aspects of inertial focusing devices such as the formation of trains of particles (31), the measurement of migration velocities (32) or the number of focusing positions (29). In contrast to long-fluorescence imaging, studies exploiting high-speed imaging for in-depth quantification are scarcer. One reason might be the requirement of bespoke and expensive imaging systems to limit motion blur. Typically, imaging can be performed with a high-speed camera synchronised with a high-power, pulsed illumination source to reach exposure times in the order of 1 – 10 µs (32)-(33)-(34)-(35). Although products have been commercialised e.g., by Dolomite, Fluigent or PreciGenome to offer plug-and-play solutions to the community, such products are often limited to brightfield imaging. In the presence of mixed populations, as often occurs in microfluidic systems, detecting and differentiating particles without fluorescent signals can be a challenge. New approaches are emerging to unlock the potential of high-speed fluorescence and/or 3D imaging (36)-(37)-(38); however, micro-particle image velocimetry (μPIV) remains the most widely accessible technique. In μPIV, the illumination is provided by a high-power, pulsed laser to record pairs of images with a short time delay. μPIV set-ups or similar have been used for inertial focusing systems to access particle or fluid velocities (31)-(32)-(39)-(40)-(41)-(42)-(43). Access to μPIV can be challenging due to high capital cost; commercialised μPIV set-ups are also often limited to one wavelength, thus requiring iterative measurements for mixed populations (one measurement per fluorescent population).

In this work, we investigate the potential of event-based cameras as a cost-effective alternative to particle detection and tracking in microfluidic devices that 1) is compatible with standard microscopes, 2) does not rely on high-power pulsed illumination sources, 3) is significantly less data-consuming and less expensive than traditional, frame-based cameras and 4) is attractive for both bright-field and fluorescence imaging. As proof-of-principle, particle detection and tracking were performed in an inertial focusing device in both fluorescence and bright-field modes for 1 µm and 10 µm-diameter beads. For the first time, this work reveals the unique capabilities of event-based sensing for overcoming some of the commonly encountered challenges in microfluidics imaging.
Materials and Methods

Bead preparation
Red fluorescent polystyrene beads, 10 µm or 1 µm in diameter (Magsphere) were diluted in filtered phosphate-buffered saline (PBS) supplemented with 0.1% v/v Triton X-100 to a final concentration of 1 x 10^4 – 5 x 10^5 beads mL⁻¹. Beads concentrations were determined using a haemocytometer.

Microfluidic setup
A spiral device was fabricated by lithography (Epigem, UK) and consisted of a channel in an Archimedes spiral, with a rectangular cross section of 360 µm (width) x 60 µm (height). The channel had one inlet, four outlets and 6 loops (Figure 2). Samples were injected into the spiral channel via the inlet with a mid-pressure syringe pump (neMESYS 1000N, Cetoni, Germany) and 1/16” polytetrafluoroethylene tubing with an internal diameter of 0.5 mm. Applied flowrates in this work ranged from 0.05 to 2 mL min⁻¹ corresponding to Reynolds numbers in the interval 4 – 159; the Reynolds number is defined as Re=ρUDh/µ, where ρ is the fluid density, µ is the fluid viscosity, U is the velocity of the fluid and Dh is the hydraulic diameter of the channel.

Prior to any measurement, and in between samples, 5 mL filtered PBS was flushed through the syringe three times, and then through the spiral at 1.5 mL min⁻¹, to clean the system. Experiments were performed in triplicate.

Imaging setup
Particles flowing in the spiral were characterised by capturing video footage (15 seconds duration, from triplicate experiments) using two separate cameras. An event-based camera (CSD3SHCD, Prophesee) consisting of a 480 x 360 pixels CMOS vision sensor, 20 µm x 20 µm event-based pixels and >10k frames per second (fps) typical equivalent frame rate, was mounted on a Zeiss Axioskop 2 fluorescence microscope (Zeiss, Germany) to visualise the spiral at 10X magnification. Prophesee player software (version 1.4.1-1935316) was used to adjust the camera settings and to record videos. To validate data from the event-based camera, a DinoLite camera (Dino-Lite) with a resolution of 1280 x 1024 pixels and a frame rate of 30 frames per second (fps) was mounted on a clamp stand and positioned above the spiral channel, and images recorded at ~50X magnification, 30 fps using DinoCapture 2.0 software.

Data analysis was performed using a bespoke data processing pipeline implemented in Matlab, which is detailed in the following section.

Event-data collection and pre-processing
As mentioned above, event-based cameras do not provide series of frames but, instead, lists of time-tagged events. Thus, pre-processing steps are required prior to information extraction. As explained in Figure 1, events are recorded when the intensity change exceeds a user-defined threshold. If this threshold is set too small, a large number of events are recorded, including signal events but also spurious events considered as “nuisance events”. For extremely low threshold values, this can lead to read-out issues whereby not all the events can be properly recorded and transmitted. Conversely, using a large threshold value reduces the number of background events but also the number of signal events, potentially hindering the detection of particles generating intensity changes. Consequently, this threshold has to be
set carefully. Here, the trade-off between accurate particle detection and low background noise was found via visual inspection using the camera software for each Reynolds number and illumination mode. Note that, in the bright-field mode, the intensity changes are weaker than in the fluorescence mode, leading to lower thresholds in practice.

In principle, particle detection can be performed directly from the stream of detection events. However, since event-based cameras are still an emerging technology, it is currently easier to map the recorded events onto temporal frames and use mature image processing tools to perform the detection task. This is the approach adopted here. One of the main advantages of event-based cameras is that this frame-based representation can be obtained after the data acquisition, and the frame rate, as well as the exposure, can be user-defined. Each pixel is subject to a dead time, i.e., a period of time after each detection, during which that pixel is not able to record subsequent events. This dead time varies depending on the overall number of events recorded in the array but is typically lower than 50 µs with the camera used here. Thus, integration periods smaller than this value are not recommended. In the context of event-based cameras, we identify the integration period as the temporal window used to aggregate events and build a frame. Conversely, the longer this integration period, the blurrier the reconstructed frames. Here exposures in the 100 µs – 1 ms range were used, depending on the expected particle velocity and illumination mode, to find the best visual trade-off between satisfactory particle detection and motion blur. Note that within a frame or integration period, several events can be recorded by the camera at a given pixel. In that case, only the most recent event is considered to generate the frame. The frame rate is also user-defined and can be set independently from the per frame integration time, e.g., it is possible to consider overlapping integration windows. Again, setting this parameter can depend on the expected speed of particles to be tracked. Low frame rates can lead to large distances travelled by the particles between two frames and can jeopardize the tracking task, especially when multiple particles are present in the field of view simultaneously (a challenging data association problem). High frame rates make the tracking task easier, but might unnecessarily increase the number of frames to be processed. In this work, the frame rate has been set to 20k fps, to ensure the particles are visible in a sufficient number of frames to estimate their velocity (> 40 frames).

Results and discussion

As presented in Figure 2, the passage of spherical particles through a spiral channel with a rectangular cross-section (360 µm x 60 µm) was analysed with an event-based camera. Particle detection and tracking were performed near the outlet of the spiral (1200 µm x 1500 µm region of interest (ROI) in Figure 2-B).
Figure 2. Imaging in inertial focusing devices. A. Standard approaches for imaging particles in inertial focusing devices include bright-field (or phase contrast) images of moving particles, composite images from long-exposure fluorescent signals or single fluorescent particle visualisation (e.g., using particle image velocimetry). B. Schematic of the microfluidic device tested in this work; ROI: region of interest. C. An event-based camera was used for particle detection and tracking in the ROI. Negative (blue) events correspond to pixels detecting a decrease in brightness and positive (red-brown) ones to an increase.

Based on previous work with similar designs (44)(45)(46), 10 µm rigid spherical beads were expected to focus towards the inner wall of the spiral channel for Reynolds numbers above ~50 in the region of interest of the channel, while 1 µm particles were expected to remain unfocused. It can also be noted that according to (47), no secondary Dean vortices are expected for the Reynolds numbers in the range of interest (Re<160). With an average fluid velocity in the spiral of 0.5 m.s⁻¹ at Re = 50, a light source with a very short pulse duration (~10 µs) would normally be necessary for single particle detection to circumvent motion blur. In this work, only a standard lighting source (Zeiss HBO100 Mercury vapor short-arc lamp) from a fluorescent microscope was used with the event-based camera.

Visualisation and detection of microparticles

As visible in Figures 3-A and -B (and video in the supplementary information), 10 µm fluorescent particles can be clearly seen at a low Reynolds number (Re = 4) using the event-based camera. The motion of particles in bright-field mode (Figure 3-C) was also demonstrated, although the contrast is generally lower than in the fluorescence mode. Figures 3 A-C highlight one specific characteristic of event-based cameras, namely the possibility to define, after the data recording, the integration period and frame rate for visualisation/analysis purposes, as discussed above. Although not all event-based cameras provide this feature, the camera used in this work also records information allowing the reconstruction of grey scale images (but at a much lower time resolution than the event-data processed here). This feature was used to create a long-exposure intensity image (in bright-field mode) for calibration purposes and to identify the location of the channel in the field of view. An example is depicted in Figure 3-D.
Figure 3. Raw data for 10 µm particle visualisation inside the spiral channel at Re = 4 in fluorescence mode for an integration time of 100 µs (A) and 750 µs (B). C. Example of raw data for bright-field imaging with the event camera and an integration time of 750 µs. Red arrows denote the position of particles in the channel, red pixels correspond to pixels detecting an increase in brightness and blue ones to a decrease. D. Reconstructed long-exposure intensity image. The scale bars (in the bottom right corner of each subplot) correspond to 200 µm, the inner and outer walls are defined in A.

It is interesting to note from Figure 3 that the trails produced by moving particles (e.g., the negative (blue) events for fluorescent particles in Figures 3 A-B) are usually longer than the events created when the particles first become visible in a given pixel. As presented in Figure 4-A, this phenomenon is due to the fact that only the most recent event is kept at each pixel when constructing the frames. Depending on the selected integration period, the reconstructed image consequently has more negative (blue) events than positive (red) ones for fluorescent particles (and more positive events for bright-field reconstructed images).

Figure 3 also shows that moving fluorescent particles induce a local increase of light flux, while particles in bright-field mode are characterized by a local reduction of the light flux. Consequently, fluorescent particles first produce positive events and then negative events, depicted in red and blue, respectively, in Figures 3-A and -B. Conversely, particles in bright-field first produce negative events and then positive events (Figure 3-C). This detail is essential to process data appropriately and differences between bright-field and fluorescence imaging are further highlighted via the simulated data presented in Figure 4-B.

In bright-field mode, static particles typically appear as dark rings (48). The darker edges and lighter centre can create a characteristic pattern on the reconstructed images (see simulated images in Figure 4-B, left panels), especially for slowly moving particles. Despite not being visible in Figure 3-C due to the high velocity of the particles, this pattern was observed at higher magnification (20x) and lower particle velocity (~0.0003 m.s⁻¹), as reported in Figure 4-C. It can also be noted that for slow-moving objects, the length of the tail could be used to estimate the particle velocity from a single image, i.e., without the need for advanced tracking.
algorithms. However, Figure 3 shows that the length of the tail of high-velocity particles cannot be accurately measured due to the low signal-to-noise ratio in that case.

**Figure 4.** A. Example of image reconstruction (single pixel presented here) from recorded asynchronous events. The events are mapped onto temporal frames based on a user-defined integration window. This gives access to a sequence of reconstructed images that can then be analysed using image processing tools. The event camera only keeps information corresponding to the last event, leading to “trails” (higher number of negative (blue) events in this example) on the reconstructed image. B. Simulation of moving particle between times t1 and t2 in bright-field and fluorescence modes and corresponding events. The third row from the top illustrates how the (simulated) reconstructed event frames are expected to look like (for a sufficiently short time delay t2-t1) and short integration time. The displacement of the particles is further highlighted by the dashed yellow lines. C. Reconstructed images obtained from measurements of 10 µm particles slowly moving in the spiral channel for an integration time of 10 ms and 100 ms. As in Figure 3 A-B, the longer the integration time, the longer the trails.

Consequently, in the following sections, only the positive events are used to detect particles in fluorescence, and respectively, negative events to detect particles in bright-field mode. More precisely, the frames of positive or negative events are first denoised using morphological transforms (e.g., erosion and dilation) to remove isolated events. The particles are then extracted by identifying groups of spatially connected events whose size falls into a predefined range. The position of the particle is then computed as the position of the centroid of each region. Note that here, all the particles used had the same apparent size, but the size/shape of the connected regions could be used in the future to classify particles and potentially enhance the tracking results.
Quantification of focusing behaviours in fluorescence and bright-field modes

In addition to demonstrating that 10 µm particles can be detected at low Reynolds numbers, we investigated whether detection was possible for the range of Reynolds numbers which are usually considered for focusing experiments with such microfluidic designs. The spatial distribution (expressed as the distance to the spiral channel inner wall) of particles detected next to the outlet of the channel, in a region of constant cross-section (before the opening), was recorded at different flow rates (Figure 5-A). For visualisation purposes, violin plots were used as a comprehensive representation of the spatial distribution of particles. A wide horizontal spread of these plots (normalised by the number of detected particles in each case) corresponds to a large number of particles detected within a narrow region of the channel (characteristics of focusing).

Figure 5. A. Distribution of particles detected in one video recorded with an event-based camera using fluorescence microscopy for Reynolds numbers Re = 4, 8, 16, 40 and 159, corresponding to applied flow rates in the interval 0.02 – 2 mL/min and averaged fluid velocities in the interval 0.04 – 1.54 m.s⁻¹. B. Distribution of particles detected with an event-based camera using bright-field microscopy for Reynolds numbers Re = 4, 8, 16. Inset. For Re = 16 (blue plots), three replicates (blue, red and black) are superimposed in the inset for both fluorescence and bright-field imaging. All the distributions depicted in this figure are normalized such that they do not depend on the particle concentration. Thus, all the violin plots (expect in the inset which has presents a different vertical scale) have the same area.
Particles were successfully detected in fluorescence mode for \( \text{Re} \in [4,159] \). The focusing of beads close to the inner wall is clearly visible when the Reynolds number increases, as expected for this microfluidic design (45). Note that each distribution has been derived from at least 1000 particles and that similar distribution profiles can be plotted at any x-location along the channel imaged (cf. Figure 3-D for definition of x axis), or mapped onto the entire section of the channel imaged (the channel length that is imaged with the current set-up is circa 1.5 mm).

For comparison purposes, similar experiments were performed in bright-field mode. Due to a lower contrast between beads and background, reliable detection was only possible up to \( \text{Re} = 16 \) (Figure 5-B). To measure the similarity between the distributions obtained in Figures 5-A and -B, we computed their percentage overlap. For \( \text{Re} = 4, 8 \) and 16, the overlap between the distributions plotted in Figures 5-A and -B was 88%, 79% and 77%, respectively. The distributions of fluorescence and bright-field modes are generally in good agreement and the increasing discrepancy with increasing \( \text{Re} \) can be partly explained by a degradation of the detection performance in bright-field mode.

With an average fluid velocity of 0.2 m.s\(^{-1}\) at \( \text{Re} = 16 \), being able to detect particles without a pulsed light confirms the potential of event-based cameras for particle detection in microfluidic channels in both fluorescence and bright-field modes. Note that 1) results for both fluorescence and bright-field modes were highly reproducible as demonstrated for three replicates at \( \text{Re} = 16 \) (cf. blue inset, Figure 5-B; overlap \( > 94\% \) for replicates in fluorescence mode and \( > 89\% \) in bright-field mode) and 2) detection at higher Reynolds number might possibly be achieved upon further optimisation of the hardware/software (see discussion in conclusion).

As a control, long-exposure fluorescence imaging was used for all the Reynolds numbers considered (cf. Figure 6-A) – in this case, images were recorded at 30 fps and stacked over a 15 second period. Fluorescence intensity distributions were then estimated in a measurement window similar to the one used in Figure 5. As visible in Figure 6-B, the intensity distributions follow patterns and trends similar to those obtained using the event-based camera, with focusing occurring at the inner wall of the spiral channel with increasing Reynolds numbers. The horizontal spread of the fluorescence distribution at \( \text{Re} = 159 \) does not appear as sharp as the particle distribution, which might be due to pixel saturation and the likely non-linear relationship between the particle and light intensity densities. Only one main mode is also observed in Figure 6-B for long-fluorescence imaging data, while two streams of focusing seem to be detected at lower Reynolds numbers with the event-camera. Multiple focusing streamlines in inertial devices have been previously reported in the literature, especially for higher volume fractions (44,49). However, the particle concentration used here has been kept the same for event-based and long fluorescence imaging experiments. Although the exact nature of this observed second mode remains unclear; its observation for both event-based fluorescence and bright-field modes seems to confirm that this is not an artefact from the cameras or from the data processing pipeline.
Figure 6. A. Composite images of 10 μm fluorescent beads flowing near the outlet of a spiral channel at Re = 16, 40 and 159. The dashed lines correspond to the channel walls, the yellow rectangles are regions where the intensity distributions have been estimated and the arrow highlights the focused stream of beads at Re = 159. Scale bars: 200 μm. B. Intensity distributions estimated in the measurement windows (yellow rectangles in A) and depicted as distances to the channel inner wall in blue. Particle densities obtained from the event-camera are superimposed in red. All the distributions depicted in this figure are normalised such that all the violin plots have the same area.

Particle tracking and velocity mapping

In this section, the potential of event-based imaging for particle tracking is investigated. Only results in fluorescence mode are reported here but tracking was also achieved in bright-field mode for all the Reynolds numbers reported in Figure 5-B.

Depending on the concentration of particles, a varying number of particles can be observed simultaneously in the field of view. Consequently, algorithms for multiple target tracking (MTT) can be used. Although it might be possible to identify particles from the complete set of generated frames, in practice, it is computationally intractable, given the high frame rates considered (20M frames for a 10 s experiment). In all experiments performed, the number of particles simultaneously present was relatively low (less than 10) and the particles presented similar trajectories and velocities. Thus, a standard online approach to MTT was adopted,
which updates the particle tracks sequentially as each frame is processed. Following the particle detection steps described above, the data association problem was solved using a variant of the Munkres algorithm (50)(51). This problem consists of deciding which detected particles are associated with existing tracks (from previous frames) and which are new. Once the data association is performed, the actual tracking of each particle is performed using a standard Kalman filter (52) assuming a near-constant velocity model (for each track). The algorithm also includes a track ending mechanism, which terminates tracks for particles that have not been seen over a given period. This makes the algorithm more robust against missed particles (which might not be detected in a few frames during the detection process).

Figure 7-A presents an example of 4 tracks of 10 µm particles identified in the channel at Re = 4 (one colour per track), leading to the estimated particle velocities plotted in Figure 7-B. An example of tracking is also visible in the video (cf. supplementary information). The estimated average particle velocity of 0.05 m.s⁻¹ is in accordance with an average fluid velocity of 0.04 m.s⁻¹. A slight decrease in particle velocity can be observed when approaching the outlets (x > 1000 µm), as expected due to the opening of the channel. Particle velocity profiles were then plotted as a function of the distance to the inner wall for different increasing Reynolds numbers. As presented in Figure 7-C, successful tracking was achieved up to Re = 40; the colormap in this figure highlights in purple, regions where many detected particles present similar velocity and distance to the inner wall, for a given Reynolds number. For Re < 40, an approximately constant velocity is measured for all the particles tracked (>1000 for each experiment). For particles detected closer to the inner wall, especially at Re = 40 with a high number of particles tightly focused in the region near y = 45 µm, a decrease in particle velocity is observed. This behaviour in particle velocity is further confirmed by the probability density functions displayed in Figure 7-D.
Figure 7. A. Example of tracks for four randomly picked 10 µm particles flowing in the spiral at Re = 4 (one colour per track). Dashed lines correspond to the microfluidic channel. Each track contains approximately 800 positions. B. Corresponding estimated particle velocity (raw data) as a function of the x-position in the channel. C. Particle velocity profile as a function of the distance to the inner wall for Re = 4-40 and D. corresponding probability density functions (P.D.F.). Plots in C are based on particles tracked in the blue region of interest highlighted in A. Colormap highlights particle density, with dense regions in purple (linear colour map L17 in (53)).

The results in Figure 7 confirm that event-based cameras can be used to track individual particle behaviours in the size range of commonly used biological cells. In order to map fluid patterns, however, smaller sizes would typically be used for µPIV experiments. Experiments were consequently also conducted with 1 µm fluorescent particles at Re = 4. With the current set-up, one pixel of the 480 x 360 pixels CMOS vision sensor corresponds to a ~3.3 µm x 3.3 µm field of view. As presented in Figure 8-A, a subpixel detection was possible in fluorescence mode, with 1 µm particles successfully detected across the channel. The estimated probability density function of particle velocity is displayed in Figure 8-B. With an average fluid velocity of 0.04 m.s⁻¹ it seems that only particles with lower velocities (<0.03 m.s⁻¹) were successfully detected and tracked. Due to the small size of the particles, the fluorescence-induced intensity change was lower than with the previously tested 10 µm particles, causing a lower signal-to-noise ratio while the particle passing time was shorter. Importantly, the fact that 1 µm beads could still be detected and tracked illustrates further the potential of event-based sensing for
μPIV experiments in microfluidic devices; a higher detection performance could be reached by either increasing the magnification or working with sensors with a higher number of pixels (cf. (2) for a descriptive review of existing cameras available and characteristics).

Figure 8. Tracking results of 1 µm fluorescent particles at Re = 4. Probability density functions (P.D.F.) of A, tracked particles as a function of the distance to the spiral inner wall and B, corresponding particle velocities. Plots A and B are based on particles tracked in the blue region of interest highlighted in Figure 7-A.

For both 10 and 1 µm particles, no information was gained on the z-position of particles detected at this stage; a similar approach to (32) using velocimetric reconstruction for reconstructing the z-position could potentially be used with event-based data too.

Conclusions

Event-based cameras offer unique advantages to track high-speed phenomena thanks to their sensors acting as silicon retinas. Although the benefits of this technology have been already demonstrated for robotics, its potential for biological/microscopic applications remains largely untapped. In this work, the performance of an event-based camera for detecting and tracking micrometric particles in a microfluidic channel was evaluated. Inertial focusing devices, due to their high working Reynolds numbers, are often recognised as challenging systems for individual particle tracking. Current approaches typically rely on high power, pulsed illumination sources and expensive micro-PIV setups to track fluorescent particles. The present work demonstrates that event-based cameras can offer an alternative to such state-of-the-art imaging setups. Particle detection was possible for a wide range of Reynolds numbers, up to Re = 159, simply by using a standard fluorescence microscope (and lighting), both in bright-field and fluorescence modes. As opposed to micro-PIV setups, the event-based camera is not limited to one wavelength; any particles that are excitable in the visible spectrum with the microscope can potentially be detected. Although more challenging to accurately track, the velocity profile of particles down to 1 µm was also measured with the tested set-up. Since the application of event-based cameras to the microfluidic world is still new, specific challenges/limitations also need to be considered. The field of event-based cameras is fast evolving with always faster and more sensitive sensors being developed. For instance, efforts are currently made to increase the fill-factor and reduce the pitch of event-based detectors, and at the same time to produce larger arrays to improve the spatial resolution. However, it is important to mention that as opposed to the camera tested here, most products do not directly offer grey scale “reconstructed image”. This might cause significant difficulties to set up the system (e.g., for the focus) since only moving/blinking objects are visible on display. This could be tackled by engineering new tools to help with the calibration either in the setup itself or computationally.
Finally, the drastic changes in data—from images to events—imposes the development of a new framework for processing. Importantly, it has been demonstrated here that data can be analysed to extract relevant information (e.g., particle focusing position, particle velocity) and can also be directly compared to images (e.g., comparison with composite image from fluorescent imaging). This, in addition to their high sensitivity to intensity changes, compatibility with standard microscopes, high speed capabilities, low consumption and lower costs compared to standard high-speed cameras, makes event-based cameras unique candidates to change our way of characterising the microscopic world.

Conflicts of interest: There are no conflicts to declare.

Acknowledgements: MJ and YA are supported by the Royal Academy of Engineering under the Research Fellowship scheme (RF/201718/1741 and RF201617/16/31). MJ would also like to thank the Engineering and Physical Sciences Research Council (EPSRC) and the Royal Society for their support (EP/R006482/1 and RGS/R1:191188).

Authors contribution: JH, TH and MJ designed and performed the experiments. YA designed the data processing approach and analysed the experimental results with MJ. JH, YA and MJ wrote the manuscript; all authors discussed the results and commented on the manuscript. YA and MJ equally contributed to this work.

Bibliographic references

rofluidics: a review. Lab Chip


33. Holloway PM, Butement J, Hegde M, West J. Serial integration of Dean-structured sample cores with linear inertial focussing for enhanced particle and cell sorting.


