While many tools exist to study immune-cell chemotaxis in vitro, current methods often lack desirable features. Using fluid-walled microfluidics, circuits are built around primary murine macrophages deposited in pre-defined patterns on Petri dishes or microplates. Concentration gradients of complement component 5α (C5α) are established in flow-free or flowing environments, cell migration imaged, and cell directionality and velocity correlated to calculated local C5α concentrations. In flow-free circuits built around patterned macrophages, only cells nearest the C5α source migrate regardless of local attractant concentration. Conversely, in flowing circuits free from intercellular signaling and attractant degradation, only cells distant from the source migrate. In both systems, cells respond to lower C5α concentrations than previously reported (~0.1 pM). Finally, macrophages follow instantly-shifted gradients better than slowly-shifting ones, suggesting that migration depends on both spatial and temporal responses to concentration.

1. Introduction

Cells often experience environments containing heterogeneous chemical signals. These signals can induce directed cell migration (i.e., chemotaxis) and play crucial roles in regulating tissue homeostasis and inflammation.[1,2] For example, chemotactic signals are essential for immunity, as they direct neutrophils and monocytes to inflammation sites. Macrophages also respond to chemotactic signals to discover and repair tissue damage, destroy infections, and maintain homeostasis. However, chemotaxis also contributes to pathological processes such as atherosclerosis.[3,4] Various chemotaxis assays have been used since the 1960s but all have significant limitations. Transwell assays are technically simple and widely-used; a porous insert containing cells is placed inside a well loaded with attractant, and (once a concentration gradient is established by diffusion) cells migrate through micrometer-sized pores into the well, and chemotaxis is quantified by removing the insert and counting cells in the well.[5] The xCELLigence assay provided a major technical advance; impedance changes are measured in real-time as cells travel through pores in a modified Boyden chamber.[6] Alternatives to the transwell assay were introduced to address some of its limitations, including tracking and monitoring single cells (as in Dunn chambers),[7] and detecting cell reversibility or fugetaxis (as in under-agarose migration assays).[8] More recently, microfluidic systems have been developed[9] that enable control of stable gradients,[10] distinction between different types of movement (e.g., chemotaxis, chemokinesis—non-directional cell migration, and fugetaxis[11]), tracking individual cells in real-time,[12] and increased throughput[13]—sometimes achieved with less reliance on specialized equipment.[14] While microfluidic approaches show great promise, their uptake in biomedical research has been impeded by the technical complexity required to operate devices, long fabrication and prototyping times, the problematic biocompatibility of the plastic often used (i.e., polydimethylsiloxane,
2. Results

2.1. Workflow and Operation

2.1.1. Printing Circuits Around Living Cells

Circuits are printed on standard 35 mm Petri dishes using a custom-built ‘printer’ as described previously.[18] Briefly, the printer consists of a 3-axis traverse that holds a dispensing needle used to add/remove liquids. We first print BMDMs in any desired 2D pattern by simply infusing culture medium containing suspended cells through the dispensing needle and onto the surface of a dish (Figure 1A(ii)). The dish is incubated (5 min) so cells attach, and the bottom of the dish is then gently and completely covered with medium; as macrophages are strongly adherent, they remain firmly attached to the dish where originally deposited (Figure 1A(iii)). Most medium is now removed to leave a thin layer, which is immediately covered with FC40—a bioinert and liquid fluorocarbon that prevents evaporation. After returning the dish to the printer in the original orientation, another dispensing needle jets a stream of FC40 through the thin layer of medium onto the bottom of the dish (Figure 1A(iii)); this jet pushes medium aside so FC40 gently and completely covers the bottom (Figure 1A(iv)). If originally deposited (Figure 1A(i)). The dish is incubated (5 min) so cells attach, and the bottom of the dish is then gently and completely covered with medium; as macrophages are strongly adherent, they remain firmly attached to the dish where originally deposited (Figure 1A(iii)). Most medium is now removed to leave a thin layer, which is immediately covered with FC40—a bioinert and liquid fluorocarbon that prevents evaporation. After returning the dish to the printer in the original orientation, another dispensing needle jets a stream of FC40 through the thin layer of medium onto the bottom of the dish (Figure 1A(iii)); this jet pushes medium aside so FC40 gently and completely covers the bottom (Figure 1A(iv)).

2.1.2. Loading and Generating Stable Gradients in Passive Circuits

In passive circuits, reagents are pipetted directly through the fluid walls into the circuit (Figure 1B(i–iii))—either manually or robotically on the printer—so that no attractant can flow out of the bottom chamber into the conduit (Figure 1B(iv)). Instead, upon reaching equilibrium, attractant simply diffuses out of the bottom chamber to create a gradient from the bottom to the top of the conduit.

2.1.3. Determining Steady-State Times and Gradient Stability

The rate of change in concentration of a substance by diffusion is described by Fick’s second law as:

\[
\frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial x^2}
\]

(c = concentration, t = time, x = distance, D = diffusion coefficient). During an initial transient phase, the concentration gradient in the conduit is defined using the 1D solution to the diffusion equation in a semi-infinite medium, where originally deposited (Figure 1A(i)). Most medium is now removed to leave a thin layer, which is immediately covered with FC40—a bioinert and liquid fluorocarbon that prevents evaporation. After returning the dish to the printer in the original orientation, another dispensing needle jets a stream of FC40 through the thin layer of medium onto the bottom of the dish (Figure 1A(iii)); this jet pushes medium aside so FC40 gently and completely covers the bottom (Figure 1A(iv)).

After reaching steady-state, the gradient in the conduit is linear and set to vary from 0 to 10 nM (\(c_{\text{max}} = 10 \text{ nM}\)). As \(t_{\text{SS}} \propto x^2\), doubling conduit length quadruples \(t_{\text{SS}}\) (e.g., for C5a in our system, if \(x = 2 \text{ mm}\), \(t_{\text{SS}} = 1.5 \text{ h}\); if \(x = 4 \text{ mm}\), \(t_{\text{SS}} = 6 \text{ h}\); Table S1, Supporting Information).

We next determine the rate of mass transfer through a conduit from Fick’s first law of diffusion:

\[
J = -D \frac{\partial c}{\partial x} \left( \frac{m}{A} \right)
\]

\(m = \text{mass flow rate, } A = \text{area}\) to assess the duration of these stable gradients (and predict maximum experimental runtimes). Assuming the concentration gradient remains approximately constant until 5% of the analyte’s initial concentration \(c_{\text{max}}\) reaches the top chamber where it becomes evenly distributed (\(c_{\text{end}} = 0.05c_{\text{max}}\)), we define the maximum experimental...
runtime as $t_{\text{end}} = 1.05 \frac{2m_{\text{CsA}}}{DAc_{\text{max}}}$ ($A =$ conduit cross-sectional area, $m_{\text{CsA}} =$ mass of CsA). The cross-sectional area of a conduit is expressed as $A = \frac{a^2}{2 \sin \theta} \left( \frac{\theta \pi}{90} - \sin 2\theta \right)$, where $a =$ conduit half-width and $\theta =$ contact angle. In this study, conduits have fixed widths so $t_{\text{end}} = x$, thus $t_{\text{end}}$ varies proportionally with conduit length (e.g., for $x = 2$ mm, $t_{\text{end}} = 15$ h; for $x = 4$ mm, $t_{\text{end}} = 30$ h; Figure S1 and Table S1, Supporting Information).

### 2.1.4. Moving Dishes Containing Passive Circuits Does Not Impair Gradient Formation

As passive circuits are prepared on the printer and moved to a microscope for imaging, movement-induced advective flows could perturb gradients; therefore, we monitor gradient uniformity as follows. Twelve circuits are printed and loaded with fluorescein on the printer (in a biosafety cabinet to ensure sterility), moved to an imaging system (an IncuCyte ZOOM in an
incubator), and scanned using fluorescence; fluorescent fronts in all conduits are located in essentially the same places, consistent with little advective perturbation (Figure 1C).

2.1.5. Setting Up Gradients in Active Circuits

The passive circuits described previously have the advantage of being simple to set up and do not require external pumps; however, their gradients change over time as diffusion drives the system to equilibrium. In contrast, gradients generated by diffusion between two laminar streams (one containing attractant) flowing continuously through one conduit remain stable and unchanging for as long as flow continues. Stable gradients in ‘m’-shaped circuits—consisting of two circuit arms leading into a single central conduit—are thus established. Briefly, two needles are inserted separately into left and right arms of the circuit; needles are filled with medium ± attractant

Figure 2. Tracking chemotactic bias in dumbbells jetted around BMDMs. C5a (10 nM) is generally added to the bottom chamber; time-lapse movies (phase contrast) of the conduit made over 48 h, trajectories of individual cells extracted, and rose plots generated (3 biological repeats for each condition). A) Migration of cells initially presents only in the top chamber. i) Cartoon showing starting positions of cells (grey with orange nuclei). ii,iii) Images of conduit. Cells migrate into an empty conduit; many are elongated with characteristic pseudopodia. Scale bar = 250 µm. iv) Cell trajectories. Downward tracks towards C5a (blue) outnumber upward ones (red). Black tracks: stationary cells. v) Rose plot with the circles indicating number of trajectory points falling into each bin and measurements of $v_{taxis}$ confirming this bias. B) Migration of cells initially present in top chamber and conduit. i) Cartoon showing starting positions of cells. ii,iii) Images of conduit; many cells become elongated. Scale bar = 250 µm. iv) Trajectories are again biased towards C5a. v) Rose plot with the circles indicating number of trajectory points falling into each bin and measurements of $v_{taxis}$ confirming this bias. C) Controls. i) Cells initially present in the top chamber + conduit and rose plots with ii) medium alone or iii) with C5a in both chambers show lack of bias. D) Mean NGDR (+SD, 3 biological replicates) for C5a gradients from conditions in panels A, B, C(ii), and C(iii); one-way ANOVA statistical significance *$p<0.033$, **$p<0.0021$, ***$p<0.0002$. 
and are connected via polytetrafluoroethylene (PTFE) tubing to two syringes on one syringe pump. The syringes now start infusing at a constant flow rate, so medium (red dye added in Figure 1D(ii)) and medium + attractant (represented by blue dye in Figure 1D(ii)) meet as laminar streams in the central conduit. Diffusion then creates a stable concentration gradient across the width of the conduit. The steepness of this gradient is controlled by varying the overall flow rate in the conduit (lower rates yield shallower gradients, Figure 1D(iii)). A third needle connected to another syringe on a separate pump is inserted into a sink at the end of the central conduit and withdraws fluid at an equivalent flow rate.

2.2. Chemotaxis of BMDMs in Passive (Flow-Free) Circuits

2.2.1. Exposing BMDMs to Gradients of C5a

We demonstrate the approach by monitoring chemotaxis of BMDMs in response to a gradient of C5a that varies from 0 to 10 nM—the concentration eliciting the strongest chemotactic response in transwell dose-response experiments (Figure S2, Supporting Information). Initially, all cells are in the top chamber of a dumbbell-shaped circuit (Figure 2A(ii)). C5a (10 nM) is pipetted into the bottom chamber, and a time-lapse movie is made of the conduit over the next 48 h (Movie S1, Supporting Information; 1 frame every 20 min). Almost no cells are seen in the first movie frame (Figure 2A(iii)). As the gradient becomes established (tSS = 0.9 h), cells enter the conduit and by the last frame some even reach the bottom chamber (Figure 2A(iii)). Tracks of individual cell trajectories are now derived by analyzing successive images in the movie (Experimental Section); these tracks show a biased movement of cells towards C5a (Figure 2A(iv)); blue tracks indicate movement towards C5a, red ones away from it, and black dots show static cells). Trajectories are also binned according to angle and displayed as rose plots (Figure 2A(v)); blue segments indicate tracks moving towards C5a, and red ones away from it, with segment length reflecting the number of trajectory points per bin). We next build circuits around cells initially located in the conduit and examine the response of these cells to the same gradient (Figure 2B; Movie S2, Supporting Information). Many cells are visible in the first frame (Figure 2B(ii)), and only some trajectories are biased towards C5a (Figure 2B(iv)); in other words, only some cells chemotax, and rose plots confirm this (Figure 2B(v)).

2.2.2. Distinguishing Between Chemotaxis and Chemokinesis

In the absence of a gradient, BMDMs typically move randomly. Whilst rose plots clearly indicate that some cells exhibit directed movement in a C5a gradient (and so chemotaxis), rose plots from controls performed in circuits without C5a in either chamber (Movie S3, Supporting Information), or with the attractant initially in both chambers (Movie S4, Supporting Information), show no bias (Figure 2C(ii,iii)). We also quantify cells’ average drift velocity in the direction of the attractant ($v_{\text{taxis}}$, average y-velocity) and net-to-gross displacement ratio (NGDR) for each condition. $v_{\text{taxis}}$s calculated by averaging the y-velocities of each trajectory (positive velocities towards the top chamber, negative velocities towards the bottom chamber). Thus, in a gradient, negative values of $v_{\text{taxis}}$ indicate biased movement towards C5a (Figure 2A(v),B(v)). In controls, $v_{\text{taxis}} = 0 \mu$m min$^{-1}$, confirming the absence of chemotactic bias (Figure 2C(ii,iii)). The NGDR is defined as the straight-line distance between start and end points of a trajectory divided by the total distance traveled along the path of the trajectory (see for instance,[21]). Therefore, an NGDR of 1 indicates a cell travels in a straight line from start to finish irrespective of whether it is towards or away from attractant, while progressively lower values point to decreased directedness (Figure 2D). All these results confirm previous ones showing that C5a is a chemoattractant for BMDMs.[19]

2.3. Assessing Population Dynamics in BMDM Chemotaxis Assays

2.3.1. Patternning Cells in Conduits Reveals Unique Population Dynamics

Thus far, we have imaged short conduits (length 1.5 mm) initially containing no cells, or those where cells are uniformly distributed everywhere. We now compare responses of cells distributed in 4 different patterns in longer 6 mm conduits (chosen to improve resolution, but note that now $t_{\text{SS}} = 13.6$ h). Initially, cells occupy either the whole conduit (Figure 3A, purple), or just the top (Figure 3B, green), middle (Figure 3C, red), or bottom third (Figure 3D, blue). Since all 4 conduits have the same dimensions and as bottom chambers are filled with the same C5a concentration, diffusion will establish identical gradients in all conduits that develop similarly over time. Remarkably, trajectories of cells at equivalent points along the y axis in the 4 conduits often differ. Thus, consider the conduit initially filled uniformly with cells: only cells in the bottom third yield long blue trajectories and a biased rose plot, whilst those above them do not (Figure 3A). This might be expected: responders lie close to the highest attractant concentration, and non-responders, far from it. Now consider the other three panels (Figure 3B–D). The front closest to C5a is always rich in responders, but this is found at different distances from the lower chamber. In other words, migratory responses of macrophages seem to depend more on the position of the cells at the front of the population, and not on the absolute C5a concentration expected at that point in the conduit. This result highlights one advantage of the approach: printing circuits around cells in precise user-defined patterns enables identification of new behaviors.

2.3.2. Determining the Lowest C5a Concentrations Eliciting Chemotaxis

We next determine the lowest C5a concentration eliciting chemotaxis ($c_{\text{min}}$) in each of the four circuits described in Figure 3. Macrophages move by extending pseudopodia that pull the cell over the substrate, and pseudopodia extending towards an
attractant are early visible markers of chemotaxis.[22,23] Therefore, we use our movies to trace cells yielding the long blue trajectories back in time, identify when and where they first extend pseudopodia towards attractant (Figure S3, Supporting Information), and calculate the local C5a concentration present at that time and place (using the solution to the 1D diffusion equation). Because these movies are collected using a framerate of 20 min, temporal resolution is limited (higher frame rates can be acquired by reducing the imaged area or increasing scan rate, but then overheating can alter cell behavior and even kill cells). Therefore, we recognize the concentration, \( c_{\text{min,obs}} \), present in the earliest frame in which such pseudopodia are seen; this is an upper limit of \( c_{\text{min}} \). Then, in movies used for Figure 3A,D (Movies S5 and S8, Supporting Information), marker pseudopodia are seen at the bottom of the conduit in the first frame, and \( c_{\text{min}} \) is \( \approx 1 \) nM (Table 1). For movies used for Figure 3B,C (Movies S6 and S7, Supporting Information), markers are seen in the second frame, and values for \( c_{\text{min}} \) are 0.1 \( \mu \)M and 79 \( \mu \)M respectively (Table 1; comparable to the minimum concentration eliciting a response in xCELLigence transwell assays, Figure S2, Supporting Information). This gives a 1000-fold range in the lowest C5a concentration eliciting a response. Values for the concentration gradients at the points where cells respond are also given in Table 1. As these gradients are so shallow, and if a macrophage is 0.1 mm in length, this makes it likely that the first responders in Figure 3B experience a C5a concentration of \( c_{\text{min}} \) at one end of their body and zero at the other. Such low concentrations of \( c_{\text{min}} \) suggest high sensitivity of this assay, which combined with the theoretical calculations provides new ways of confidently studying chemoattractant gradients in ranges lower than previously reported.[24,25]

### 2.4. Comparing Multiple Cell Types and Higher-Throughput Chemotaxis Studies in Microplates

#### 2.4.1. Label-Free Comparison of Different Cell Populations in a Single Circuit

We next design and implement circuits with 3 chambers that enable comparison of chemotactic responses of two distinct cell populations without requiring distinguishing cell labels (Figure 4A). The middle chamber serves as a C5a reservoir, while chambers and conduits above and below contain different cell types. To establish gradients, medium is first added

![Figure 3](image-url)
to the top chamber (1 µL, \( P_{\text{top}} = 20 \text{ Pa} \)), a smaller volume than the bottom one (0.9 µL, \( P_{\text{bottom}} = 18.5 \text{ Pa} \)), and an even smaller volume of C5a to the middle one (0.6 µL, \( P_{\text{middle}} = 13 \text{ Pa} \)). This creates pressure gradients that drive medium into the C5a chamber (Figure 4A(i)), red dye represents medium, blue represents medium + C5a until the system reaches equilibrium; then, diffusion creates a steady gradient of C5a along both conduits (Figure 4A(iii)). To demonstrate the approach, C5a-receptor 1 knock-out (KO) BMDMs are initially at the top, wild-type (WT) BMDMs at the bottom, and 10 nM C5a in the middle. As expected, KO cells exhibit mostly unbiased movement in the upper conduit (Figure 4A(iv), rose plot in top inset), as WT cells chemotax towards C5a (Figure 4A(iv), rose plot in bottom inset; Movies S9 and S10, Supporting Information). Although a small proportion of KO cells migrate directionally towards C5a, this may be due to incomplete KO of all C5a receptors in some cells or potential contribution of another receptor (C5aR2) to migratory mechanisms. This demonstrates how phenotypic differences of various cell populations can be observed and analyzed in a single microfluidic system without the need for additional labeling to distinguish subpopulations.

### 2.4.2. Increasing Throughput Using Microplates

We now demonstrate a method to print and operate circuits in conventional microplates to perform higher-throughput assays. The technique uses a custom-built three-axis traverse that is larger than the printer used thus far and can accommodate bigger substrates such as microplates; however, it is operated much like the printer. Then, for example, 12 identical dumbbell circuits are printed in each well of a 6-well microplate (i.e., 72 circuits in total; Figure 4B(i)). As before, gradients of chemoattractant are generated and circuits imaged to analyze cellular response. We thus compare the response of BMDMs cultured in the top chamber of circuits and exposed to gradients of C5a, to that of BMDMs plated everywhere in circuits with uniform distributions of C5a or in the absence of C5a. Results show a clear increase in the mean NGDR for cells experiencing gradients compared to controls (Figure 4B(ii), as in Figure 2D). The flexibility of the approach allows circuits with various shapes to be shrunk to fit inside wells of 12- to 96-well plates (Figure S4, Supporting Information).
2.5. Chemotaxis of BMDMs in Active (Flowing) Circuits

2.5.1. Generating an Unshifting Stable Gradient Circuits

Here, we use an ‘m’-shaped circuit and infuse medium ± 10 nM C5a into left and right arms, respectively (Figure 1D). These two streams converge as laminar streams in the central arm to flow over BMDMs, and diffusion establishes a C5a gradient (from 0–10 nM) across the width of the conduit. Concentrations are determined using the solution to the 1D time-dependent diffusion equation in finite media, where

$$q(x,t) = \frac{\epsilon_{\text{max}}}{2} \text{erfc} \left( \frac{x}{\sqrt{4D\mu t}} \right)$$

and $t =$ time a molecule spends in the conduit.\(^{[20]}\) Flow velocities at any point in fluid-walled conduits can be predicted using a simplified power law\(^{[26]}\) if the geometry of the conduit footprint, flow rate, and fluid properties of perfusing medium are known. We thus determine flow velocities to transform the time-dependent diffusion solution into a distance along the device substitution using $t = \frac{x}{\mu}$, where $\mu$ is the mean flow velocity at a given $x$-location.

2.5.2. BMDMs Respond to Low Concentrations of C5a in Unshifting Stable Gradients

As before, we make a movie over 48 h to follow chemotaxis of BMDMs across the central channel in the ‘m’. Many migrate from the left to the center, leaving behind a cell-depleted zone (Figure 5A(ii,iii); Movie S11, Supporting Information); individual trajectories and a rose plot (segment length indicates the number of trajectories per bin) confirm chemotaxis towards C5a (Figure 5A(iii)). Note that the cell band just to the right of this depleted zone is enriched in taxing cells superimposed on non-responsive ones that were initially present. A theoretical prediction of the normalized concentration gradient is now mapped onto the image of cells at 48 h (Figure 5A(iv)); colors depict local concentrations from 0 to 1 (corresponding to 0–10 nM C5a), and curves the gradient contours (red line—5 nM, with black lines correspond to decreasing concentrations). The leftmost black line is chosen to represent the minimum concentration triggering a response, and maps onto the leftmost edge of the depleted region (equivalent to $\epsilon_{\text{max}} \times 10^{-5} = 0.1 \text{ pM}$). While the diameter of a murine macrophage is typically $\approx 20 \mu$m,\(^{[27]}\) their length can be $\approx 100 \mu$m when extending pseudopodia and moving (Figure 5A(ii)). As a result, cells in this region of the conduit easily sense one to two orders of magnitude difference in C5a concentration across their body. Remarkably, and in contrast to flow-free circuits, it is now cells far from the front that respond (as those at the front do not). We will argue in the Section, “Discussion” that this difference results from the effects of flow continually maintaining attractant levels. The predicted C5a concentrations around responsive cells in both flow-free and flowing circuits are comparable to the lowest concentrations eliciting taxis in the xCELLigence assay (Figure S2, Supporting Information), thus validating our findings. To our knowledge, the response of macrophages to such low concentrations of C5a has not been shown before.

2.5.3. Controlling Cell Movement by Shifting the Gradient

We now attempt to control cell movement by shifting the gradient a known distance to the right in order to recruit more responsive cells in specified regions of the conduit. First, a 10 nM gradient of C5a is generated over cells cultured in the conduit of an ‘m’-shaped circuit as before, and cells are imaged on a microscope (Figure 5B(ii)). As expected, after 24 h, a band of BMDMs has migrated into the central region of the conduit (Figure 5B(ii)). The gradient is now abruptly shifted 200 μm to the right by increasing the flow rate of the left stream (7.15 μL h$^{-1}$) and decreasing the flow rate of the right one (2.85 μL h$^{-1}$) (Figure S5, Supporting Information) as the overall flow rate in the conduit remains unchanged (10 μL h$^{-1}$). Cells are then imaged for an additional 24 h (Movie S12, Supporting Information). Analysis of cell trajectories and a rose plot reveals that the band of cells distributed along the center of the conduit shifts in the direction of C5a, indicating a clear bias towards the newly-positioned gradient (Figure 5B(iii,iv); note the large proportion of blue tracks towards C5a, and enlarged blue segment in the rose plot).

Since BMDMs follow a gradient of C5a when abruptly shifted, we investigate whether cells can follow a continuously-shifting gradient. As before, BMDMs are exposed to a 10 nM gradient of C5a and imaged over 24 h (as in Figure 5A), during which a sub-population of cells migrate from the left to the center of the conduit where the concentration of C5a is higher (Figure 5C(i,iii)). Over the following 24 h, the gradient is shifted continuously from the center of the conduit to 150 μm from the right edge, by gradually increasing the flow rate through the left arm (from 5 to 9.98 μL h$^{-1}$) and decreasing the flow rate through the right one (from 5 to 0.02 μL h$^{-1}$) which is performed using the ‘ramp’ function of the syringe pump. Analysis of cell trajectories during this time reveals that BMDMs do not follow the gradient as efficiently, and the chemotactic bias observed previously reduces (rose plots in Figure 5C(iii)). However, when the gradient remains stationary for an additional 24 h, chemotaxis becomes stronger, and a new cell band appears on the right (Figure 5C(iv); Movie S13, Supporting Information). This confirms that cells remain competent to chemotax throughout the 72 h, and that sweeping the gradient across more cells—at least at the pace chosen in this experiment—does not significantly recruit more cells into the enriched band. To our best knowledge, the difference in macrophage response to abrupt and continuously shifting gradients has not been reported before and is a feature easily detected using our approach. This behavior is consistent with cells responding poorly when their leading edges sense falling attractant concentrations despite the presence of a gradient across their whole bodies; then, they only migrate actively once the gradient stops moving. In vivo, gradients of chemokines are likely evolving over time and therefore this new platform has the potential to facilitate our understanding of macrophage response to changing inflammatory landscape in tissues.

3. Discussion

We describe a suite of chemotaxis assays designed using micro-fluidic circuits built around living cells on standard Petri dishes.
A stationary

i  t = 0

5 μL/h
media

5 μL/h
C5a

ii  t = 48 h

iii tracks

iv normalised gradient

B abrupt shift

i  t = 0

5 μL/h

5 μL/h

ii  t = 24 h

5 μL/h

5 μL/h

7.15 μL/h

2.85 μL/h

iii  t = 48 h

IV tracks 24 – 48 h

C static, gradual shift

i  t = 0

5 μL/h

5 μL/h

ii  t = 24 h

5 μL/h

5 μL/h

media ↑

C5a ↓

iii  t = 48 h

4.98 μL/h

0.02 μL/h

gradually shift gradient

iv  t = 72 h

10^5 0.5

10^5 0.5

10^5 0.5

60

60

60
Table 2. Comparing traditional and fluid-walled chemotaxis assays.

| Assay                              | Transwell assays | Direct observation and cell tracking chambers | Under agarose assays | Traditional microfluidic assays | Fluid-walled microfluidics |
|-----------------------------------|------------------|----------------------------------------------|----------------------|--------------------------------|----------------------------|
| Stable gradients                  | ±[38]            | ±[38,43]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Single-cell tracking              | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Fugetaxis detection               | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Distinction between chemotaxis and chemokinesis | ±[31,40] | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| High throughput                   | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Specialized equipment             | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Real-time observation             | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Multiple cell types               | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Patterned cells                   | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Multiple chemoattractants         | ±[31,40]         | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |
| Cell isolation/circuit reconfiguration | ±[31,40] | ±[31,40]                                     | ±[31,40]            | ±[31,40]                      | ±[31,40]                   |

(Figure 1) [18] and apply them to investigate the complex interactions involved in the chemotaxis of primary murine macrophages to C5a—a small peptide released at inflammation sites. Taking advantage of properties of circuits with fluid walls, this method addresses many limitations of traditional chemotaxis assays (Table 2). Gradients of C5a are established in flow-free (passive) dumbbell-shaped circuits, and theoretical models are developed to predict local C5a concentrations; as fluid walls are transparent, [28] time-lapse movies are made, and individual cell trajectories are analyzed (Figure 2). Remarkable results are obtained by comparing responses of cells initially occupying different parts of 4 identical conduits—either the whole conduit, just the top, middle, or bottom third. One might expect cells to respond according to position and so to predicted C5a concentration gradient; instead, responders are the ones closest to the highest C5a concentration irrespective of position (Figure 3). A proof-of-concept control in a modified 3-chamber circuit then compares the chemotactic response of WT and KO cells (without any cell labels) and shows that WT cells migrate towards C5a more efficiently than KO ones (Figure 4A). Passive circuits are also built on well plates to increase throughput (e.g., 12 circuits in each well of a 6-well plate; Figure 4B). Finally, tenfold steeper gradients than the ones in flow-free circuits are generated with external pumps driving input flows through ‘m’-shaped circuits; now, responders are distant from the front (Figure 5A). Cell movement is also controlled by shifting gradients from the center of a conduit to the right-hand wall (by 200 or 750 μm) either abruptly or continuously (Figure 5B,C). Interestingly, cells do not efficiently follow a continuously-shifting gradient, although they do so once the gradient stops moving. In other words, cells seem to sense the presence of C5a in both time and space. In both passive and active circuits, cells respond to far lower concentrations than previously reported (=0.1 pM), and this is validated using the real-time xCELLigence transwell assay (Figure S2, Supporting Information).

Figure 5. Chemotaxis in steep C5a gradients held stationary, shifted abruptly to the right, or shifted progressively to the right. BMDMs in the central conduit of ‘m’-shaped circuits are exposed to laminar streams of medium and medium + 10 nM C5a, so diffusion creates a concentration gradient across the conduit. Movies are collected as the position of this gradient is changed (either abruptly or steadily) at times indicated by varying input flows. As fluid walls are transparent, [28] time-lapse movies are made, and individual cell trajectories are analyzed (Figure 2). Remarkable results are obtained by comparing responses of cells initially occupying different parts of four identical conduits—either the whole conduit, just the top, middle, or bottom third. One might expect cells to respond according to position and so to predicted C5a concentration gradient; instead, responders are the ones closest to the highest C5a concentration irrespective of position (Figure 3). A proof-of-concept control in a modified 3-chamber circuit then compares the chemotactic response of WT and KO cells (without any cell labels) and shows that WT cells migrate towards C5a more efficiently than KO ones (Figure 4A). Passive circuits are also built on well plates to increase throughput (e.g., 12 circuits in each well of a 6-well plate; Figure 4B). Finally, tenfold steeper gradients than the ones in flow-free circuits are generated with external pumps driving input flows through ‘m’-shaped circuits; now, responders are distant from the front (Figure 5A). Cell movement is also controlled by shifting gradients from the center of a conduit to the right-hand wall (by 200 or 750 μm) either abruptly or continuously (Figure 5B,C). Interestingly, cells do not efficiently follow a continuously-shifting gradient, although they do so once the gradient stops moving. In other words, cells seem to sense the presence of C5a in both time and space. In both passive and active circuits, cells respond to far lower concentrations than previously reported (=0.1 pM), and this is validated using the real-time xCELLigence transwell assay (Figure S2, Supporting Information).
We discuss these complex results in the light of three inter-related observations. First, consider the gradients in the 4 identical passive circuits. In the absence of cells, and since all circuits have the same dimensions and contents, gradients will evolve in them identically. However, cells at equivalent points along the 4 conduits (and so experiencing similar C5a concentrations and gradients) respond in some cases, but not in others (Figure 3). Then, C5a concentration gradient cannot be the only trigger of taxis. Second, there is clear evidence that cells can distort pre-existing gradients, and even generate ones from a uniform field of attractant by binding, internalizing, or degrading attractant around them.[29] Macrophages also change their migration when other cells further up a gradient distort the concentrations behind them.[2,30] Moreover, self-generated gradients can guide cells robustly over longer distances than pre-existing ones, with responding cells always being the ones at the front.[2,29] This is consistent with what we observe in flow-free gradients; taxing cells are at the front, irrespective of predicted C5a concentration (Figure 3). Third, responding cells can emit signals to near neighbors that amplify an initial response, and reduce a late one (e.g., neutrophils initially signal to each other to swarm towards infecting bacteria, and later they reduce excessive swarming and so reduce inflammation[31]). Although other responses to concentration gradients by spatial sensing. [33]

It is widely accepted that eukaryotic cells regulate their responses to concentration gradients by spatial sensing.[31] Additionally, migrating myeloid cells (dendritic cells and neutrophils) sense temporal dynamics of chemoattractant concentrations and respond to rising concentrations of CCL19 and CXCL12—a behavior that contrasts with C5a.[34] Moreover, neutrophils migrate further towards shifting gradients of interleukin-8 than static ones.[35] Note that there is ample time for signals to diffuse between cells in our flow-free circuits, but not in our flowing ones—as any secreted signals are quickly flushed away (e.g., in Figure 5A, medium in the conduit is wholly replaced every minute). Then, it is no longer surprising that responders are at the front in flow-free gradients (Figure 3), but not in flowing ones (Figure 5) where inter-cellular signaling becomes impossible. Therefore, our findings suggest that macrophages can sense C5a both spatially and temporally, and that shifting gradients modulate what is already a complex response.

A technical limitation of the current study results from the low temporal resolution of recordings; estimation of the minimum concentration ($c_{min}$) eliciting chemotaxis in passive circuits is limited by framerate. However, our approach has various advantages; it allows users to generate stable gradients over cells freely-patterned in 2D, track individual cells, distinguish between types of movement (chemotaxis/chemokinesis), compare cells of different types without requiring additional cell labels, and—in the future—increase throughput, and isolate cells of interest in real-time for downstream analysis, as cells can be selectively retrieved through fluid walls.[36,37] We thus hope this study provides new ways of evaluating chemotaxis and spatio-temporal sensing in adhesive mammalian cells. For example, this approach should allow comparison of transcriptomes and proteomes of responding and non-responding cells for downstream analysis.

4. Experimental Section

Cells, Media, Culture Conditions, Reagents: C57BL/6, Balb/c or 5ar1tm1Cge/J on Balb/c background (C5aR KO) were used in this study. C57BL/6 were bred in Oxford, the Balb/c WT plus the C5aR KO were bought from the Jackson Laboratory. To obtain BMDMs, bone marrow was extracted from femurs and tibia of mice and frozen for up to 6 months in –80 °C prior to use. Upon defrosting the bone marrow was cultured for 7 days (37 °C, 5% CO2) in high-glucose Dulbecco’s Modified Eagle’s Medium (DMEM, Sigma) enriched with 10% L929-conditioned media (containing macrophage colony-stimulating factor), 10% Fetal Bovine Serum (FBS, Sigma) and 1% penicillin plus streptomycin (P/S, Gibco). As this combination was used throughout the study, the mixture was simply referred to as ‘medium’. 8 mL medium was used for the first 4 d; then, 3 mL was removed and replaced with 5 mL fresh medium. Cells were plated on dishes on day 6, circuits built around them on the same or next day for passive and active circuits, respectively, and chemoattractants added on day 7. Most circuits were built on 35 mm Petri dishes (Corning, 430165) or 6-well plates (Corning, 3516). 12- to 96-well microplates were additionally used (Corning) for higher-throughput applications. In all chemotaxis assays, recombinant mouse complement component 5a was used (C5a, 10 nM; R&D Systems).

Imaging: BMDMs were imaged at a rate of 3 frames h⁻¹ (for experiments in dishes) or 2 frames h⁻¹ (for experiments in plates) using an IncuCyte ZOOM, a live-cell imaging system (Sartorius, Gottingen, Germany). The IncuCyte was itself contained in an incubator (37 °C, 5% CO₂).

Printing Circuits: In all experiments, cells were first plated on either Petri dishes or microplates, and—after attaching to the substrate—the circuit was jetted around them. Cell plating and circuit printing were performed on custom-made three-axis traverses (iotaSciences Ltd, Oxford, UK) referred to as a ‘printer’ (for circuits on Petri dishes), and larger ‘pro-printer’ (for circuits on microplates).

Loading and Patterning Cells: Cells were first plated on the dish using the printer.[39] Briefly, the three-axis of the printer holds a dispensing needle (25G, Adhesive Dispensing Ltd) connected by PTFE tubing (Cole-Parmer) to a 250 μL glass syringe (Hamilton) controlled by a syringe pump (iotaSciences Ltd). The needle was brought 300 μm above the dish surface and infuses BMDMs in medium + 2% fibronectin (Sigma, F1141; 1 million cells mL⁻¹). The needle was then moved over the surface of the dish to deposit cells in the desired pattern. After printing, the dish was incubated (5 min) to allow cells to attach.

Jetting Circuits: The dish was now removed from the incubator and 1 mL medium was gently added to wet the entire bottom. Most medium was removed, and 1 mL FC40 (iotaSciences Ltd) was added to cover the remaining thin film of medium. The dish was placed back on the printer in the same orientation and a second needle loaded with FC40 jets a stream of FC40 vertically downwards from 500 μm above the surface of the dish. When this jet contacts the surface, it displaces the underlying medium, and—as FC40 wets polystyrene better than water—it replaces medium to form a fluid wall (a medium:FC40 interface). The needle was then moved over the dish and around cells to create fluid walls that form the footprint of the circuit.[39] For passive circuits, 1 μL medium was initially added to each chamber after jetting the circuit.

Circuit Design: Two types of circuit were used, static (passive) circuits without flow, and dynamic (active) ones with flow. The first passive circuits used were dumbbell-shaped and consist of two square chambers (footprints 2.5 x 2.5 mm) connected by a thin conduit (600 μm wide, 1.5 mm long). Circuits were built around cells so cells end up in the top chamber, in the top chamber + conduit, or in bands at different positions in the conduit. C5a was then added to the bottom chamber. For these experiments, 12 circuits were printed in each dish (Figure 1C). Cells in different locations of the circuit grow under different heights of medium due to the curved profile of the overlying fluid interface,
and this might affect cell behavior; therefore, cells growing at different positions were tracked throughout static circuits in medium or a uniform distribution of C5a (i.e., without a gradient), and find that average cell velocities were unaffected by position in the circuit (Figure S8, Supporting Information).

For cell-comparison studies (Figure 4A), a slightly modified version of the passive circuit was used by connecting three chambers (also 2.5 mm wide) to two conduits (600 µm wide, 1.5 µm long). Due to larger circuit sizes, 6 circuits were printed in each dish.

For flow-generated gradients, ‘m’-shaped circuits were used. These circuits consist of two arms that join into a conduit that ends in a large circular sink (7 mm wide). Cells grow in the central conduit (6 mm long, 1.8 mm wide), and side arms were used to perfuse medium or medium + C5a.

Passive Circuit: As described previously, reagents were loaded using the printer, and dishes subsequently moved to the IncuCyte for imaging. Loading was performed by matching reagent volumes to ensure that C5a remains in the bottom chamber and does not flow into the conduit. To achieve this, pressure in the C5a chamber was kept lower than the medium chamber during loading, so medium flows into the C5a chamber until the system reaches equilibrium and then diffusion can create a steady gradient down the conduit. This was done by first infusing 1 µL medium into the top chamber, and then 0.5 µL C5a in the bottom chamber. After loading all 12 circuits, the dish was gently moved from printer to incubator (to not cause sudden flows of C5a) and placed inside the IncuCyte for imaging.

Active Circuit: Gradients in active circuits were generated as in previous studies.[37] Briefly, two 500 µL glass syringes (Hamilton) were loaded with either medium or medium + 10 nM C5a, placed on a syringe pump (Harvard Apparatus), and connected to needles via PTFE tubing. Using a 3D printed adaptor that fits around the edge of the dish and holds the needles, needles were inserted into each arm of the circuit (left—medium, right—C5a). A third needle—also connected to a 1 mL glass syringe (Hamilton) on a separate syringe pump and held in the adaptor—was inserted into the sink. The first two needles were set to infuse (at rates indicated in Figure 5; total flow rate in the conduit is 600 µL h⁻¹). Cells grow in the central conduit (6 mm long, 1.8 mm wide), and side arms were used to perfuse medium or medium + C5a.

Determining the Diffusion Coefficient of C5a: In both types of circuits, theoretical models were used to predict local concentrations and gradient steepness across conduits. The models use well-known solutions to the 1D diffusion equation (Supporting Information) that require the diffusion coefficient of C5a to be known. D_C5a was thus determined using the Stokes–Einstein equation for the diffusion of a spherical particle in water of radius r as

\[
D_C5a = \frac{\kappa B T}{6 \pi \eta r}
\]

Note: The presentation of the third sentence in the abstract was changed on September 9, 2022, after initial publication online.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

P.R.C. and E.J.W. each hold equity in and receive emoluments from iotaSciences Ltd., a company exploiting the fluid-shaping technology; iotaSciences Ltd. also provided the printers, the FC40STAR, and a scholarship for C.D. All other authors declare they have no competing interests.

Author contributions

C.D. and A.N.R. contributed equally. C.D., A.N.R., P. R.C., D.R.G., and E.J.W. designed the research; C.D., A.N.R., and F.N. performed experiments; C.D., A.N.R., and J.H.R.W. analyzed data; and all authors worked on the paper.

Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.

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Author contributions

C.D. and A.N.R. contributed equally. C.D., A.N.R., P. R.C., D.R.G., and E.J.W. designed the research; C.D., A.N.R., and F.N. performed experiments; C.D., A.N.R., and J.H.R.W. analyzed data; and all authors worked on the paper.

Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.
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