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in vivo

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Abstract. Accurate quantification of circulating cell populations in mice is important in many areas of preclinical biomedical research. Normally, this is done either by extraction and analysis of small blood samples or, more recently, by using microscopy-based \textit{in vivo} fluorescence flow cytometry. We describe a new technological approach to this problem using detection of diffuse fluorescent light from relatively large blood vessels \textit{in vivo}. The diffuse fluorescence flow cytometer (DFFC) uses a laser to illuminate a mouse limb and an array of optical fibers coupled to a high-sensitivity photomultiplier tube array operating in photon counting mode to detect weak fluorescence signals from cells. We first demonstrate that the DFFC instrument is capable of detecting fluorescent microspheres and Vybrant-DiD-labeled cells in a custom-made optical flow phantom with similar size, optical properties, linear flow rates, and autofluorescence as a mouse limb. We also present preliminary data demonstrating that the DFFC is capable of detecting circulating cells in nude mice \textit{in vivo}. In principle, this device would allow interrogation of the whole blood volume of a mouse in minutes, with sensitivity improvement by several orders of magnitude compared to current approaches.

Keywords: road image; diffuse; \textit{in vivo}; flow cytometry; fluorescence.

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1 Introduction

There are many applications in preclinical biomedical research where it is desirable to measure the number of cells circulating in the bloodstream in small animals. For example, the number of circulating tumor cells is known to be an important predictor of progression of cancer metastasis.\textsuperscript{1-3} In current practice, circulating cell populations are normally quantified in mice by extraction of small blood samples and \textit{ex-vivo} analysis using, for example, flow cytometry,\textsuperscript{5} hemocytometry, or microfluidic devices.\textsuperscript{6-10} However, typical sampling volumes for nonterminal experiments are less than 1% of the total blood volume of the animal, so very rare cell types are difficult to detect.\textsuperscript{11} Measurement of changes in cell populations over hours or minutes is also challenging because blood samples are typically drawn only once per day.\textsuperscript{11}

These limitations have motivated the development of noninvasive techniques that can be operated continuously without drawing blood samples.\textsuperscript{12-18} Microscopy-based \textit{in vivo} flow cytometry of fluorescently labeled circulating cells, wherein a laser beam is focused through a microscope objective across a small blood vessel in the ear or retina of a mouse, is one such approach. As cells cross the laser path, a fluorescent pulse is generated that can be detected with a photomultiplier tube (PMT).\textsuperscript{12,13} While \textit{in vivo} flow cytometry has proven useful for many applications—including enumeration of circulating red blood cells, T-lymphocytes, and multiple myeloma (MM) cells,\textsuperscript{12,13,20,21} the limitation is that relatively small blood volumes are sampled, specifically on the order of 1–5 $\mu$L per minute. Therefore, monitoring of about 1% of the blood volume of a mouse requires approximately 30 min. This sets a practical lower limit of sensitivity of about $10^4$ to $10^5$ cells in circulation. As such, very rare circulating cell types may escape detection entirely.

Here we describe a new technological approach to this problem using high-sensitivity detection of diffuse fluorescent light from relatively large blood vessels \textit{in vivo}. Specifically, our goal is to develop an instrument capable of detecting fluorescently labeled circulating cells in mice in the range of less than $10^4$ cells in circulation, i.e., below the sensitivity range of microscopy-based instant-varying fading channel (IVFC; \textit{in vivo} flow cytometry). Our strategy was to develop a fluorescence sensing ring that would be placed around a relatively narrow, 2 to 3 mm limb of a mouse, where total circulating blood flow rates are approximately 0.2 to 0.5 mL per minute.\textsuperscript{22} Therefore, in principle, the entire $2.5$ mL blood volume of a mouse could be interrogated in minutes. As we discuss, the principal engineering challenges in developing this instrument were i. high-sensitivity detection of very low signal levels from individual cells, ii. rejection of interfering background autofluorescence, and iii. minimizing movement (breathing) artifacts in mice.

In this paper, we first describe our diffuse fluorescence flow cytometer (DFFC) instrument and validate its operation with a custom-made optical flow phantom model. We demonstrate that the instrument can successfully detect single fluorescent microspheres and fluorescently labeled cells passing through flow phantoms with similar size, optical properties, autofluorescence, and flow speeds of a mouse limb with excellent counting accuracy. We also present preliminary \textit{in vivo} validation of our design by detecting fluorescently labeled circulating MM cells in a nude mouse limb.
cells in the tails of nude mice. To our knowledge, the concept of enumerating circulating cells with diffuse fluorescence light has not been described previously. We anticipate that our DFFC instrument will have many potential applications in biomedical research, including the detection of cancer metastasis at earlier stages and counting circulating hematopoietic stem cells in vivo.

2 Methods and Materials

2.1 System Description

A schematic diagram and photograph of the DFFC instrument is shown in Figs. 1(a) and 1(b), respectively. The sample—either a limb mimicking flow phantom or a mouse limb—was placed in the center of the custom-made ring, and could be illuminated by two solid-state lasers emitting at 642 nm (DL640-050-O, CrystaLaser, Inc., Reno, NV). The output of each laser was passed through 640-nm excitation clean-up filters with a 10-nm bandpass (Z640/10x, Chroma Technology, Rockingham, VT), and the power of the sample was 15 mW with a spot size of 1 mm diameter. For the work described in this article, only a single laser was operated in continuous wave (CW) mode, but as we discuss, in the future we plan to modulate both lasers so as to allow coarse tomographic localization of fluorescent cells in the cross-section of the phantom or mouse limb.

Emitted fluorescence signal from the sample was detected with six optical fibers that were arranged around the ring holder as shown in Fig. 1. Six detector fibers were used so as to allow close to full-angle collection of the emitted fluorescent light. As we demonstrate, the signal from individual fluorescent microspheres and fluorescently labeled cells was detectable on each of the six detection channels, but in this work, these were summed to improve the overall signal-to-noise ratio (SNR). Special cut 2 × 3 mm filters centered at 700 nm with a 50-nm bandpass (ET700/50, Chroma Technology, Rockingham, VT) were placed in front of collection fibers; this blocked diffusely and specularly reflected light from directly entering the fiber and generating secondary autofluorescence. On the opposite end, the fibers were terminated on a custom-designed filter housing with hardening. During phantom characterization experiments, the output from each detection optical fiber was simultaneously (the two additional PMT channels were unused). The photon-counting threshold was set to −100 mV for each channel, and the sampling rate was set to a rate of 100 samples/s. The maximum number of photon counts per time sample was 65,535 (hardware limited) on each of the six detection channels. For each experiment, the MCS card was configured to acquire continuously for 7500 samples, which was equivalent to 75 s. The 75 s measurement "run" could be repeated an arbitrary number of times with approximately a 0.1-s time delay between cycles to allow for writing of the data to the hard drive. Experimental automation was performed using the analog outputs of a multifunction data acquisition card (DAQ; NI-USB-6251, National Instruments, Austin, TX) controlled with the same personal computer.

2.2 Limb-Mimicking Optical Flow Phantoms

To test and characterize our instrument, we first developed an optical flow phantom with similar size, optical properties, and flow speeds as a mouse limb or tail [Fig. 1(b), inset]. The phantoms were constructed from polyester resin material (Casting Craft, Fields Landing, CA) with titanium oxide (TiO2; Sigma-Aldrich, Inc., St. Louis, MO) and India ink (Higgins Ink, Bellwood, IL) added to adjust the optical properties.23 Phantoms were first made with final optical properties close to that of biological tissue at near-infrared wavelengths,24 specifically with the reduced scattering coefficient μs = 15 cm−1 and absorption coefficient μa = 0.1 cm−1 (these baseline optical properties were used for all experiments in this paper unless otherwise specified). The liquid resin material was placed in a 3-mm diameter by a 1-cm cylindrical mold with a length of 250 μm internal diameter Tygon tubing (TGY-010-C, Small Parts, Inc., Seattle, WA) passed through the center before hardening. During phantom characterization experiments, the Tygon tubing was connected to a 30-gauge insulin syringe (Easy Touch, Loveland, OH) that contained a solution of either

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Fig. 1 (a) A schematic of the diffuse fluorescence flow cytometry instrument, and (b) a photograph of the device during in vivo operation with a limb-mimicking optical flow phantom (inset). The position of the six detection fibers (D1 to D6) are shown.
fluorescent microspheres or fluorescently labeled cells as described below. The syringe was placed in a microsyringe pump (70-2209, Harvard Apparatus, Holliston, MA) that could be configured to produce linear flow rates in the range of 600 μm/s to 15 cm/s. A new flow phantom was used for each of the experiments described herein.

Second, to investigate the effect of the flow phantom optical attenuation on the measured fluorescence signal, we constructed a set of phantoms with increasing concentrations of India ink. Specifically, the final absorption coefficients of the phantoms were μ_a = 0.1, 0.25, 0.4, and 0.55 cm^-1. This was repeated four times for each ink concentration.

2.3 Fluorescent Microspheres

Characterization of the DFFC instrument was first performed using commercially available flow cytometry calibration microspheres (6-μm PeakFlow Claret, P-24670, Invitrogen, Carlsbad, CA) with an absorption peak near 645 nm and an emission peak near 695 nm. These microspheres were chosen since they have similar absorption and emission characteristics, such as the Cy5.5, Alexafluor-680, and Vybrant-DiD NIR fluorescent dyes that we anticipate using with our system. Further, according to the manufacturer’s specifications, these microspheres have similar size and fluorescence emission intensity as fluorescently labeled cells. Before each experiment, the microspheres were diluted and suspended in phosphate-buffered saline (PBS) solution at final concentrations between 10^2 and 10^4 spheres/mL. Before dilution, and before each experiment, the microsphere suspensions were sonicated for 5 min (2510, Branson Ultrasonics Corp., Danbury, CT) to prevent clumping.

2.4 Characterization of DFFC Detection Sensitivity

To determine the counting accuracy of our DFFC instrument, microspheres were suspended in PBS and 250 μL samples were passed through a flow phantom at a constant linear flow speed of 1 cm/s. For these experiments, a total of eight samples in the range of less than 10^3 microspheres/mL were prepared; i.e., in the target sensitivity range of our instrument. Accurate dilution of microsphere solutions at very low concentrations is difficult. Therefore, to obtain “true” microsphere concentrations, the samples were collected in a microcentrifuge tube after analysis. These were subsequently counted with a commercial flow cytometer (Cell Lab Quanta SC, 771917, Beckman Coulter, Brea, CA), which has ±5% accuracy in the range of 3 × 10^4 to 2 × 10^6 spheres/mL.

To obtain a total microsphere count with our DFFC instrument, we applied a thresholding algorithm to our measured data as follows. As will be shown, when a fluorescent microsphere passed through the instrument detection ring, a transient fluorescence signal (i.e., a “spike”) was recorded. The signal from all six detection channels were summed, and any fluorescence spike exceeding one standard deviation above the maximum measured background—determined by first running a blank sample of PBS through the flow phantom—was counted as a single microsphere. In practice, this threshold was approximately 300 photon counts. The count total for the 250 μL sample was then multiplied by 4 to arrive at a concentration in microspheres per mL for direct comparison to the commercial flow cytometer measurements. We note that we did not attempt to account for possible simultaneous arrival of more than one microsphere (i.e., “double spikes”) in this work. The implications of this are discussed below.

2.5 Characterization of the Useful DFFC Flow Speed Range

Since we anticipate that a large range of blood flow rates will be present in the instrument field of view in a mouse limb in vivo (i.e., in different-sized blood vessels), we next performed a series of experiments to determine the range of flow speeds for which our system could detect single microspheres. Microsphere suspensions were prepared in the range of 10^2 to 10^3 spheres/mL. Then 1-mL samples were run through the flow phantom at varying linear flow speeds between 600 μm/s and 15 cm/s. The latter value reflects a higher flow speed than we anticipate in limb blood vessels in mice in vivo, and the former is the lowest flow speed that the microsyringe pump could produce. Experiments were repeated three times for each flow speed. As above, the signals from all six channels were summed, and measured fluorescent spikes were analyzed for each to obtain the full width half maximum (FWHM), defined as the duration of time between data points with magnitude halfway between the maximum value for a given spike and the mean background. All spikes were analyzed for each experimental condition and the mean and standard deviation FHWMs were recorded.

2.6 Characterization with Fluorescently Labeled Cells in Flow Phantoms In Vitro

We next tested the ability of the DFFC instrument to detect fluorescently labeled cells in our phantom model using either Jurkat T-lymphocyte cells (American Type Culture Collection, Manassas, VA) or MM cells (Northwestern University, Chicago, IL). Jurkat T-lymphocytes were cultured in 75-cm² tissue culture flasks at 37 °C in a humidified atmosphere with 5% CO_2 and 95% air. The cells were cultured in RPMI 1640 supplemented with 100 U/mL penicillin, 100 μg/mL streptomycin, and 10% fetal bovine serum (FBS). Cells were grown to confluence (approximately 1 × 10^6 cells/mL) and then dyed with 10 μmol/L of Vybrant DiD (V-22887, Invitrogen) cell-labeling solution that had been incubated for 30 min. Vybrant-DiD is a nonspecific lipophilic dye that brightly labels cell membranes without loss of viability. The cell suspension was centrifuged and washed repeatedly in PBS and then resuspended in PBS at final concentrations of approximately 10^3 cells/mL.

MM cells were harvested using trypsin, spun down at 400 g, and then resuspended in RPMI with 0.1% bovine serum albumin (BSA) at a concentration of 1 × 10^6 cells/mL. Cells were dyed using a final concentration of 1 μmol/L of Vybrant-DiD and incubated for 30 min at 37 °C. At the end of the incubation process, PBS was added (2% of total volume) to prevent cell clumping during centrifuging. Cells were centrifuged as before and washed, once with RPMI with BSA to remove any free DiD in suspension, and again with RPMI only. They were then resuspended at approximately 10^6 cells/mL.

For these experiments, the syringe pump was configured to produce a linear flow speed of 1 cm/s. Following data collection, the signal from each of the six detection channels was summed and the fluorescence spike amplitude from each cell was analyzed to determine the relative fluorescence intensity compared to fluorescent microspheres. The fluorescence signal from the cells and microspheres were also quantified with the commercial flow cytometer for comparison with our DFFC instrument.
2.7 Experiments in Mice In Vivo

As an initial in vivo test of the DFFC instrument, we performed a limited number of experiments in mice with injected MM cells. All mice were handled in accordance with Northeastern University’s Division of Laboratory Animal Medicine policies on animal care. MM cells were used since they have been used previously for microscopy-based in vivo flow cytometry experiments\(^\text{12,13}\) and circulate with known kinetics. Further, as we demonstrate, MM cells exhibited better Vybrant-DiD labeling (measured intensity) than Jurkat cells. Nude (nu/nu) mice first were anesthetized using a cocktail of ketamine (100 mg/kg) and xylazine (5 mg/kg), injected i.p. The mice were then placed on an adjustable platform with a warming pad and their tails were passed through the detection ring. Each tail was gently secured at each end with medical tape so that it would remain taut (but not tight enough to restrict circulation) in order to minimize breathing movement artifacts.

A total of \(10^6\) Vybrant-DiD-labeled MM cells were suspended in 100 μL in RPMI. Cells were injected retro-orbitally into the mice \((N = 3)\) while the instrument was operating so that changes in the measured fluorescence signal could be observed immediately after injection. Measurements were made until approximately 15 min after injection. We also performed control (sham) injections with unlabeled MM cells on a separate set of mice \((N = 3)\).

3 Results

3.1 Fluorescent Microspheres

Figure 2 shows a sample data set collected from each of the six detection channels of the DFFC instrument over a 30-s interval when fluorescent microspheres were run through a limb-mimicking optical flow phantom. For this experiment, a concentration of 500 microspheres/mL with a flow speed of 1 cm/s was used. As each fluorescent microsphere passed through the instrument field of view, a transient fluorescent signal (i.e., a “spike”) was observed. As shown, the amplitudes of these spikes were typically in the range of 1000 to 5000 photon counts above the background, depending on the detection channel. We also note that we observed intraspike variability in amplitude and width even within a single detection channel. This was primarily due to variations in speed across the flow profile inside the Tygon tubing; i.e., because microspheres near the center of the tubing move faster than those near the walls of the tubing. This was confirmed by observing the movement of fluorescent microspheres in bare Tygon tubing with a fluorescence microscope. For all subsequent analysis in this work, the signal from the six channels was summed; i.e., integrated over all detection angles. This was not a necessary step since microspheres were detectable on individual channels, but it improved the instrument SNR. The exact improvement varied somewhat between experiments and detector channels but was generally by approximately 5 to 7 dB (e.g., for the data in Fig. 2, the average channel SNR was approximately 19 dB, whereas the SNR of the summed signal was approximately 26 dB). As we discuss, we plan in future work to treat the signal from each channel independently, specifically to approximately localize the fluorescent microsphere or cell tomographically in the cross-section of the sample.

3.2 Effect of the Optical Phantom Absorption Coefficient on Measured Signals

To investigate the effect of varying the optical absorption of our flow phantoms on the measured fluorescence signal, we increased the quantity of India ink so that \(\mu_a\) increased from 0.1 to 0.55 cm\(^{-1}\). These data are summarized in Fig. 3. Unsurprisingly, increasing the absorption coefficient reduced the amplitude of measured spikes; specifically, increasing \(\mu_a\) by a factor of 5.5 decreased the amplitude of the measured spikes by a factor of 4. However, microspheres were easily detectable above the background in all cases. This range of absorption coefficients covers reported literature values in the red and near-infrared region for biological tissues\(^\text{24}\) and therefore shows the feasibility of this technique in a phantom model. Further, we added ink to the PBS media in which the microspheres were suspended so that \(\mu_a\) of the media increased from 0 to 0.6 cm\(^{-1}\). Increasing the absorption coefficient here had only negligible effect on the measured spike height in this range (data not shown) since the volume of PBS in the 250-μm diameter tubing was very small compared to the bulk phantom material.

3.3 Analysis of DFFC Detection Sensitivity

We next characterized the detection sensitivity of our DFFC instrument in our flow phantom model. Using the thresholding

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Fig. 2 Sample fluorescence signals from detector channels D1 to 6 (a–f) as microspheres passed through a limb-mimicking flow phantom at a concentration of 500 spheres/mL and a linear flow speed of 1 cm/s. Transient “spikes” (shown as arrows, panel a) were detected as each microsphere passed through the instrument field of view.
algorithm described above, we counted the number of microspheres measurable with our system in 250 μL samples and then compared this concentration to that obtained with the same sample using a commercial flow cytometer. Figure 4 shows a cell count comparison between microspheres on our DFFC instrument and the flow cytometer for a set of eight experiments, all of which were in the target concentration range of less than 10⁵ cells per mL. The dashed line indicates the ideal 1:1 correspondence between the two systems. Generally, we observed very good correlation between the two instruments, with a mean error of less than 20%. We note that the microsphere concentrations used here were at least an order of magnitude below the recommended operating range of the commercial flow cytometer of 3 × 10⁴ cells/mL. Therefore, deviations between the two measurements are at least partly due to the accuracy of the commercial system at low concentrations (as opposed to our DFFC prototype). From these data, we conclude that our DFFC instrument could detect individual microspheres with very high sensitivity.

3.4 Analysis of DFFC Flow Speed Range

To ensure that the DFFC instrument was capable of detecting single microspheres over a large range of blood vessel flow speeds, we investigated the effect of varying the speed from 600 μm/s to 15 cm/s on the measured fluorescence spike width. For these experiments, microsphere concentrations between 10² and 10⁵ spheres/mL were used. These data are summarized in Fig. 5. Each data point represents the mean and standard deviation of at least 100 fluorescent spikes at each flow speed. The highest flow speed (15 cm/s) represents a faster speed than occurs normally in blood vessels in mice *in vivo*. The slowest flow speed used here (600 μm/s) was the lower limit that could be produced with our microsyringe pump rather than a detection limit of our instrument. We note that the relatively large standard deviations for each point shown do not reflect error in the measurements, but real observed variations in flow speeds of the microspheres across the flow profile as discussed above. Finally, it is interesting to note that the linearity of the curve plotted on a log-log scale is consistent with the anticipated 1/x relationship, since the product of each FWHM and flow speed combination must equal the fixed instrument field of view. Therefore, from this analysis, we were able to estimate that the field of view of the DFFC instrument is approximately 0.7 mm.

3.5 Detection of Fluorescently Labeled Cells in Flow Phantoms

We next tested the ability of our DFFC instrument to detect fluorescently labeled cells (as opposed to microspheres) through the diffusive optical flow phantom. Vybrant-DiD-labeled Jurkat T-Lymphocyte cells and MM cells were used. Sample data summed from all six detection channels are shown in Fig. 6. Here, a constant flow speed of 1 cm/s and cell concentrations of approximately 10⁴ cells/mL were used. Sample data from Vybrant-DiD-labeled Jurkat cells [Fig. 6(a)] and MM cells [Fig. 6(b)], as well as data measured from fluorescent microspheres [Fig. 6(c)], are shown for comparison. The relative mean and standard deviation of measured fluorescent spike heights for each case are shown in Fig. 6(d), averaged over 1000 individual spikes. From these data, it can be seen that microspheres had the highest fluorescence intensity on average, followed by MM cells and Jurkat cells, which exhibited about 48% and 10% of the fluorescence intensity of the microspheres, respectively. These relative intensities also generally agreed well with conventional flow cytometry analysis [Fig. 6(d)], which showed that MM cells were on average about 50% as bright as the microspheres.
as fluorescent microspheres, and Jurkat cells were approximately 6% as bright. In summary, this series of experiments demonstrated that, although fluorescently labeled cells exhibited a lower level of emitted fluorescence than the fluorescent microspheres, our DFFC prototype could robustly detect diffuse fluorescent light from individual cells passing through an optical flow phantom.

3.6 Experiments in Mice In Vivo

Finally, we conducted preliminary in vivo testing of our DFFC instrument in mice, with results shown in Fig. 7. After being anesthetized, nu/nu mice were placed on the translation platform and their tails placed inside the detection ring of our instrument. While the DFFC instrument was in operation, a total of $10^6$ MM cells suspended in 100 μL of media were injected retro-orbitally. Sample data for mice injected with fluorescently labeled MM cells and unlabeled control cells are shown in Figs. 7(a) and 7(b), respectively. We note here that the time scale is approximately 15 min for both experiments (i.e., significantly longer than that shown in previous figures), and the data from multiple time bins has been summed to 0.1-s increments. Magnified (120-s) portions of the curves are shown in the inset figures. Arrows indicate the time of the retro-orbital injection, and the signal fluctuations immediately prior to these were caused by unavoidable movement of the mouse during the injection. When fluorescently labeled cells were injected, an increase in the measured fluorescence signal due to the bolus of cells entering the bloodstream was observed within seconds. This increase began to decay over the course of several minutes, which we

![Graphs and diagrams](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics-037001-6-March-2012-Vol.17(3))

Fig. 7 The measured fluorescence signal from the tail of a mouse over approximately 15 min. when (a) $10^6$ Vybrant-DiD-labeled MM cells and (b) unlabeled control cells were injected retro-orbitally. Inset: magnified sections of the curves.
interpret to be due to the dilution of the bolus of injected cells in the mouse blood volume. On top of the larger bolus, individual fluorescent spikes were detected, which we interpret to be due to brightly labeled fluorescent cells passing through the instrument field of view. In additional experiments that were performed (data not shown), these spikes were observable up to 1 hour after injection. The average FWHM of the measured fluorescent spikes was about 0.23 s, which corresponds to a linear flow speed inside the instrument field of view of approximately 4.5 mm/s (Fig. 5). This speed agrees well with reported literature values of average flow speeds in mouse tail arteries of less than 8 cm/s.22,23,26 We note that relatively small numbers of fluorescent spikes were observed here, paradoxically because large numbers of cells were injected in these tests. This resulted primarily in a DC fluorescence increase as a result of the bolus, as opposed to many individual spikes as in our phantom experiments.

In contrast, when unlabeled control cells were injected [Fig. 7(b)], neither the bolus nor individual fluorescent spikes were observed. The measured average background autofluorescence signal was similar in amplitude to that observed in the phantom. However, inspection of the data revealed that there was a small but consistent 1 to 2 Hz component in the signal, which we attribute to artifacts from breathing movements of the mouse (this component was not present in the detected signal in our phantom studies). While the amplitude of this signal component was significantly lower than the fluorescent signals from circulating cells, minimization of these motion artifacts by carefully securing the tail was nonetheless critical in these experiments.

Finally, we note that a steady decrease in the background signal of about 1% per minute was observed over the course of all the experiments. The cause for this decrease is unclear, but we hypothesize that it may be due to a decrease in the core temperature of the mouse during the experiment or photo-bleaching of native tissue chromophores.

4 Discussion and Conclusions

In this work, we described and characterized a new instrument for detecting circulating cells in mice with diffuse fluorescent light. We demonstrated that the DFFC instrument is capable of robustly detecting individual fluorescent microspheres and fluourescently labeled cells in an optical flow phantom with similar size as a mouse forelimb, hindlimb, or tail and with optical properties in the range of reported literature values for biological tissue in the red and near-infrared region ($\mu'_s = 15 \text{ cm}^{-1}$; $\mu_a = 0.1$ to 0.55 cm$^{-1}$). While our experiments with Vybrant DiD-labeled cells indicated that the signal from fluorescent microspheres was larger than labeled cells, MM cells were nonetheless brightly labeled (at approximately 50% of the intensity of microspheres) and these were easily detectable with our DFFC instrument. Labeled Jurkat T-Lymphocyte cells had an average of about 6% of the brightness of the microspheres, and our analysis indicates that we detected about 50% of these cells with our instrument in our in vitro flow phantom studies.

As we discussed, the most difficult engineering challenge we faced was rejection of interfering background autofluorescence, which was found to originate from the optical flow phantom or mouse limb in the instrument field of view, but also the experimental apparatus itself (e.g., the optical fibers). Since even modest amounts of autofluorescence could obscure the very weak fluorescence signal from individual cells, significant effort was required to minimize this background signal in the development phase of this work, specifically with respect to instrument geometry and selection of appropriate excitation and emission filters.

With respect to the detection sensitivity and count accuracy, our data indicated (Fig. 3) that the measured count rate of our DFFC prototype correlated very well with conventional flow cytometry in the target sensitivity range of $<10^3$ spheres/mL. As discussed, we did not attempt to account for the simultaneous arrival of multiple microspheres in the instrument field of view with our cell-counting algorithm. However, at higher concentrations, our threshold approach could lead to undercounting errors; i.e., since simultaneous arriving cells would be single-counted. This issue could be addressed in the future by using more sophisticated counting algorithms (for example, by analyzing the measured pulse width and height). However, we note that this issue is not of immediate concern since our goal is to detect cells at very low concentrations (of $<10^3$ cells total).

An obvious challenge in applying our DFFC instrument to accurate cell counting (as opposed to cell detection) in mice in vivo is the variation in the number, direction, and flow speed ranges of blood vessels in a tail or limb. With respect to flow speed, our analysis (Fig. 4) demonstrates that the DFFC is capable of robustly detecting cells over several orders of magnitude of flow speeds. As we have noted, the highest flow speed used (15 cm/s) was higher than the anticipated arterial blood flow speeds in mice. On the other hand, the lower limit of detection will be defined practically by the ability to identify a fluorescence signal as a “spike.” In particular, very slow moving cells will yield very long fluorescent signals; for example, a cell moving at 50 $\mu$m/s would produce a transient signal with a FWHM of approximately 14 s (assuming a 0.7-mm field of view). Careful analysis of measured data sets would therefore be required to distinguish this effect from instrument DC drift.

Second, the presence of multiple blood vessels within the DFFC field of view in vivo presents a challenge in accurate quantification of the number of circulating cells. We have conducted a series of experiments in phantoms with multiple embedded flow channels (not shown) and demonstrated that the DFFC instrument can detect cells from multiple vessels simultaneously. However, it is conceivable (or even likely) that a single cell in vivo may pass through the instrument more than once on its return trip through the vasculature, so that cells may be double-counted. To address this issue, we are developing algorithms to localize the fluorescent cell in the cross-section of the instrument field of view tomographically using differences in the measured signal amplitude on the six detector channels (although beyond the scope of the current work, we have recently shown that this is possible using two modulated light sources and plan to report this in a forthcoming article). We will use this data in combination with a priori anatomical information to identify approximately the location of the blood vessel in which the cell is moving to avoid double-counting.

We also note that ultimately our goal is not to provide an absolute count of circulating cells, since it is entirely conceivable that a given circulating cell may happen not to pass through a blood vessel in the DFFC field of view during an acquisition (or, as we have already noted, a cell may pass...
through the field of view multiple times). Rather, our intention is to use the count rate (detected cells per minute) as a metric to quantify relative changes in circulating cell populations in response to, for example, disease progression\(^\text{30}\) or disruption of the homing process,\(^\text{30}\) as is often done for microscopy-based IVFC applications but at significantly lower cell concentrations. Our hypothesis is that, given sufficiently long acquisition times—on the order of 15 to 30 min—mixing of the blood volume will be sufficient so that count rates will be statistically accurate. Testing of this hypothesis is the subject of ongoing research.

Finally, we performed a feasibility test of the DFFC instrument in detecting fluorescently labeled MM cells in the tails of nude mice \textit{in vivo}. For this initial set of experiments, relatively large numbers of cells were injected (10\(^6\) cells in 100 \(\mu\)L), which resulted primarily in a DC increase in the detected fluorescent signal with a relatively small number of measured fluorescent spikes. We note that the use of retro-orbital injections most likely resulted in significant loss of cells at the injection site, so the actual number of circulating cells in these experiments is not known. More accurate quantification, using concurrent measurements with microscopy-based IVFC, is the subject of ongoing work. In future work, we will reduce the concentration of injected cells by several orders of magnitude until the target instrument sensitivity range of less than 10\(^3\) cells in circulation is reached.

We also tried placing the DFFC ring around the forelimb of a mouse when fluorescently labeled MM cells were in circulation, which resulted in measurement of slightly larger fluorescent spikes than in the tail (presumably because of lower optical absorption), but movement artifacts were significantly larger since the forelimb was more difficult to immobilize. As we discussed, movement artifacts in the range of 1 to 2 Hz were present in the measured signal in the tail even in control mice, and special care was taken to minimize tail movement from breathing. In the future, we plan to add additional “out-of-band” detection wavelengths to the DFFC to allow better subtraction of interfering background signals.

In summary, in this work we described a new instrument to detect fluorescently labeled cells using diffuse light in mice \textit{in vivo}. Translation of this approach to larger clinical scales, while of potential interest in the longer term, is beyond the immediate intent of this work. This will allow interrogation of large blood volumes (up to 0.2 to 0.5 mL of blood per minute), which in principle would allow sampling of the entire blood volume of a mouse in minutes. This would not only increase the sampling rate but would allow the detection of circulating cells at several orders of magnitude lower concentration than is currently possible. We anticipate that this instrument would have many applications in preclinical biomedical research, including enumeration of circulating tumor cells at early stages, as well as tracking of hematopoietic stem cells \textit{in vivo}.

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\section*{References}


