Visualizing medium and biodistribution in complex cell culture bioreactors using in vivo imaging

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Visualizing medium and bio-distribution in complex cell culture bioreactors using in-vivo imaging

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Abstract. There is a dearth of technology and methods to aid process characterization, control and scale-up of complex culture platforms that provide niche micro-environments for some stem cell-based products. We have demonstrated a novel use of 3d in-vivo imaging systems to visualize medium flow and cell distribution within a complex culture platform (hollow fiber bioreactor) to aid characterization of potential spatial heterogeneity and identify potential routes of bioreactor failure or sources of variability. This can then aid process characterization and control of such systems with a view to scale-up. Two potential sources of variation were observed with multiple bioreactors repeatedly imaged using two different imaging systems: shortcutting of medium between adjacent inlet and outlet ports with the potential to create medium gradients within the bioreactor, and localization of bioluminescent murine 4T1-luc2 cells upon inoculation with the potential to create variable seeding densities at different points within the cell growth chamber. The ability of the imaging technique to identify these key operational bioreactor characteristics demonstrates an emerging technique in troubleshooting and engineering optimization of bioreactor performance.
Stem cell-based products always incorporate some heterogeneity due to variability in biological input materials, different micro-environments in the bioprocess and cell sensitivity to extrinsic factors. Improving the understanding, characterization and control of a bioprocess, is essential in ensuring process reproducibility and product quality\(^1\). In-process measurements aid process understanding and control, and one of the most commonly used methods of in-process measurement is cell number and cell viability determination, but many tests are invasive and subjective\(^1, 5\). There is a requirement for improved process control equipment and techniques, and the use of quantitative non-invasive and non-destructive automated imaging techniques are expected to become more prevalent for on-line monitoring\(^6\). However current techniques are primarily suited to 2-dimensional adherent cell culture formats, and for many applications 3d culture is preferable for scalable manufacture\(^7\). Although many 3d culture systems have on-line monitoring and culture maintenance capabilities, and traditional 3d stirred tank production technology is known to perform and be readily scalable for suspension cultures\(^2, 8, 9\), more complex 3d platforms are required for some cell-based bioprocesses in order to provide a more appropriate micro-environment\(^9, 10\). For example, perfused hollow fiber bioreactors have been used for primary human hepatocytes\(^11\) and hematopoietic progenitor cells\(^12\), where the stem cell niche is considered of critical importance to successful expansion, differentiation and maturation. An advantage of these more complex perfused bioreactors is a higher surface area to volume ratio leading to potentially higher product yields, improved homogeneity relative to fed-batch or static methods, and reduced waste accumulation. Yet the disadvantages associated with such systems are variable harvest yield, increased risk of heterogeneous culture conditions or cell distribution with complex internal structure/flow (primarily rigid systems), increased risk of problematic cell dissociation, and unknown or negative effects of continuous removal of paracrine factors\(^13, 6, 10\). There is a requirement to evaluate methods of in-process bioreactor
monitoring and bioreactor failure modes in such systems that are inherently difficult to characterize in order to assess what drives variability in such systems. Here we demonstrate the proof of principle that 3d in-vivo imaging systems can be utilized to image medium and bio-distribution in a hollow fiber cell culture bioreactor to aid characterization of potential spatial heterogeneity and sources of variability.

Two potential sources of heterogeneity were investigated: media distribution and cell seed distribution within the bioreactor. Equipment for in vivo bio-monitoring were considered as good candidates to allow monitoring of flow patterns and cell distribution in the bioreactor. The laboratory scale bioreactor has an 8 mL cell chamber whereas the clinical scale bioreactor has an 800 mL cell chamber (Stemcell Systems GmbH, Berlin, Germany). The cell chamber comprises layers of counter-currently perfused medium capillaries with integrated gas capillaries. Cells are cultured outside the fibers in the extra-capillary space and medium / gas is circulated through semi-permeable fibers through the intra-capillary space. The internal structure of the bioreactor, the perfusion set up, perfusion and inoculation conditions have been described in detail elsewhere. Two imaging systems were used; the Pearl Impulse system (LI-COR Biosciences, Cambridge, UK) and the Ivis Kinetic system (Caliper Life Sciences, Cheshire, UK). As the Pearl Impulse system does not have the chamber capacity to analyze an 800 mL bioreactor, a further high capacity system, the Ivis Kinetic, was also investigated as this has the potential for downstream use with a scaled bioreactor.

Medium distribution. The Pearl Impulse Imager was shown to be capable of detecting artificially induced fluorescence and flow patterns of fluorescent liquid (IR Dye® 800CW probe, LI-COR Biosciences) within the bioreactor and these signals were distinguishable from background auto-fluorescence of the plastic. Fluorescent dye-spiked buffer was perfused under counter-current flow through the bioreactor to capture video of flow characteristics. Figure 1 shows a cross-section of still images from a representative video
sequence. It identifies shortcutting of medium flow from the inlet port to the adjacent exit port, rather than crossing the bioreactor to the appropriate opposite exit port for each medium capillary network. The medium shortcutting was observed at the higher flow 20 mL/min medium recirculation rate\textsuperscript{13} and could potentially decrease the homogeneity of the bioreactor and linearity of process scalability. As perfusion continued for a few minutes medium was subsequently capable of circulating to all parts of the chamber at this flow rate (as shown in the latter images from the video sequence). Prior to using the Ivis Kinetic imager, spectral analysis of the hollow fiber bioreactor was conducted using an Ivis Spectrum system (Caliper Life Sciences) to image the bioreactor with a large number of filters (green fluorescent protein, GFP, range to near-infrared, NIR, range) to identify excitation/emission combinations for the lowest background noise signal. The 710/780 excitation/emission filter combination provided the lowest background fluorescence, indicating the Xenolight CF750 dye (Caliper Life Sciences) or similar alternative and the 700 nanometer (nm) filter wheel in the Ivis Kinetic would achieve the best fluorescent imaging data (Figure 2A). Additionally, Figure 2B, with blocked excitation, showed that the bioreactor had no detectable phosphorescence suggesting the possibility of tracking a bioluminescent cell population. The shortcutting of fluorescent dye-spiked media between adjacent medium in-flow and out-flow ports observed previously was also observed when bioreactors were imaged using the Ivis Kinetic under identical counter-current medium perfusion conditions (Figure 2C shows a still image from a representative video sequence). The effect of medium short-cutting was observed in multiple bioreactors with two different imaging systems suggesting that this could be a source of potential process variation. Additionally, as medium short-cutting was observed at the higher circulation flow rate, there is potential for differences or gradients to arise at lower flow rates such as the 4 mL/min cell feeding rate\textsuperscript{13}. 
Cell Seeding distribution. A mock cell seeding exercise using the Pearl Impulse Imager and inoculating fluorescent dye-spiked buffer (IR Dye® 800CW, 800 nm fluorescence channel) into the cell inoculation port appeared to show uneven dye distribution after seeding, thereby suggesting the potential for non-homogenous cell density within the cell chamber as another possible candidate for introducing process variability. To further investigate cell distribution after seeding into the cell chamber of the bioreactor, a population of bioluminescent murine cells (4T1-luc2, Caliper Life Sciences) were inoculated into the bioreactor under conditions described previously. The Ivis Kinetic imager was able to detect the bioluminescent population within the hollow fiber bioreactor, and showed that the cells appeared to localize in the center of the cell chamber (Figure 3), suggesting that distribution in the compartment may not be homogenous at inoculation.

Shortcutting of medium between adjacent inlet and outlet ports has the potential to create medium gradients, and heterogeneous distribution of cells upon inoculation has the potential to create variable seeding densities within the bioreactor. Such sources of variation could cause considerable process performance heterogeneity problems with scale-up. Small scale bioreactor systems using low cell densities tend to function well as there is an excess of nutrients available. However, as systems are scaled and higher cell densities achieved, early identification of heterogeneity issues can be critical as cell and medium gradients may result in some cells being starved of both nutrients and oxygen with others being overfed. Variation in harvest cell yield from the bioreactors has been observed previously and the ability of the imaging technique to identify these key operational bioreactor characteristics that may contribute to such variation demonstrates a powerful emerging role in troubleshooting and engineering optimization of bioreactor performance. Defining procedures to maximize cell seeding homogeneity (e.g. optimized liquid chasing or bioreactor rotation procedures) could
improve this situation and be further explored with imaging as an initial output, with subsequent population distribution tracking, cell yield, viability and characterization measurements. Imaging could also be used as an initial output alongside cell yield and viability to further inform modeling experiments and explore the effect of optimizing medium perfusion rates and distribution on cell expansion or other bioreactor phenomena such as Starling flows (secondary flow in the cell chamber driven by elevated medium recirculation levels in the capillaries). Technique limitations would also need to be further defined, for example the qualitative nature of the 2d heat-map data may be addressed by region of interest (ROI) definition to attain quantitative fluorescence and phosphorescence data from the Imaging system software. As well as further definition of the potential for obtaining complete 3d images and quantitative data. More work is also required to define the achievable resolution of the technique, for example whether single cell imaging is possible.

The technique may also be useful for other hollow fiber technology applications. For example, Pinzon et al. (2013) recently demonstrated a microbe-based bio-surfactant production process using hollow fiber technology under denitrifying conditions in order to eliminate the common oxygen transfer limitations and foaming difficulties that are inherent to the current aerobic process. Cell entrapment within the extra-capillary space offered the advantage of easy soluble product recovery from perfused medium and the authors suggest bioreactor scale-up should be explored. The imaging technique described may offer advantages to the scaling process to assess labeled cell distribution or tracing labeled feedstock compounds for localization or limitation. Additionally as continuous product harvesting via multivalent cation addition and extraction is an attractive concept the technique could be used to assess residues alongside cell tracking and measures of cell viability. Another area where the technique may offer unique advantages is for those
processes that require further cell immobilization, for example alginate immobilization to promote stem cell differentiation or increased levels of insulin production in pancreatic/islet-like cell clusters. A novel bioprocess for alginate immobilization of mammalian cells within a hollow-fiber bioreactor was recently demonstrated by Hoesli and colleagues (2009) and key processing issues included the obstruction of medium flow due to alginate crossing the membranes from the extra-capillary space into the intra-capillary space as well as cell recovery from the extra-capillary space. The method used to assess effective alginate gelling during method development was visual inspection of sawed hollow fiber cartridges. The imaging technique described would eliminate the need for destruction of hollow fiber cartridges and could be used to trace labeled alginate for effective gelling or labeled medium flows through alginate bioreactors. Additionally, labeled cells could be used for method development of cell recovery or to trace effective removal and residues of temperature sensitive alginate gel for more effective recovery.

The work described demonstrates that as a proof-of-principle medium and bio-distribution imaging in complex hollow fiber cell culture bioreactors is possible using 3d in-vivo imaging techniques. Additionally, imaging of medium distribution and bioluminescent cell populations in this manner could be particularly useful for early detection of heterogeneity within complex 3d cell culture platforms and aid engineering of operational parameters to improve process performance and control prior to scale-up.

**Materials and Methods.** The Pearl Impulse system is a fluorescent imaging platform with laser excitation and near-infrared (NIR) fluorescence detection (see http://www.licor.com/bio/products/imaging_systems/pearl/pearl_imager.jsp). The Ivis Kinetic system is a filter-based platform for fluorescence (green fluorescent protein, GFP-NIR range) or bio-luminescence (see...
The hollow fiber bioreactors were provided courtesy of Celgene Cellular Therapeutics (New Jersey, USA); they are produced by Stemcell Systems GmbH (Berlin, Germany)\textsuperscript{13,18}. Each laboratory scale bioreactor consists of an 8 mL cell compartment and multiple layers of three independent capillary networks that are interwoven and provide counter-current medium perfusion and integrated gas supply. The internal structure of the bioreactor, the perfusion set up, perfusion and inoculation conditions have been described in detail elsewhere\textsuperscript{13}. Bioreactors were perfused at the medium recirculation rate of 20 mL/min\textsuperscript{13}. Fluorescent imaging of liquid flow patterns with the Pearl Impulse imager (calibrated to manufacturer’s standards) was performed using the IR Dye\textsuperscript{®} 800CW probe (1µg/mL in PBS, LI-COR Biosciences), 800 nanometers (nm) fluorescence channel detection and 0.5 s image acquisition. Spectral analysis of the bioreactor was performed using an Ivis Spectrum imaging platform (Caliper Life Sciences, calibrated to manufacturer’s standards) to image the bioreactor rapidly with GFP range to NIR range filters for background fluorescence as well as phosphorescence. Fluorescent imaging of liquid flow patterns with the Ivis Kinetic imager (calibrated to manufacturer’s standards) was performed using the Xenolight CF750 dye (1 mM in dH\textsubscript{2}O), 700 nm filter wheel detection and 0.5 s image acquisition. Cell localization imaging was performed using the luciferase expressing 4T1-\textit{luc2} Bioware Ultra murine cell line, cultured according to manufacturer’s conditions in RPMI-1640 medium (ATCC, #30-2001) supplemented with 10% FBS (Fisher Scientific, Loughborough, UK). 4T1-\textit{luc2} cells were prepared to $5 \times 10^4$/mL density and exposed to Xenolight D-luciferin potassium salt substrate (150 µg/mL in RPMI-1640) at 37ºC for 5 min. Cells were inoculated into the 8 mL bioreactor cell chamber port at a density of $8 \times 10^5$/mL (6.4×10\textsuperscript{6} cells in total), as described previously\textsuperscript{13}, for subsequent imaging (37 ºC imaging chamber). All reagents, cell lines and substrates for Ivis Kinetic Imaging were provided courtesy of Caliper Life Sciences. Imaging was
performed in duplicate (minimum) with duplicate (minimum) bioreactors unless otherwise stated, a total of 6 bioreactors were used in this study.

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References


**Figure Captions**

**Figure 1.** Image A shows the hollow fiber bioreactor structure: (Port 1) gas outlet, (2) medium outlet, (3) counter-current flow medium inlet, (4) cell inoculation port, (5) gas inlet, (6) medium inlet and (7) medium outlet. Inset: illustrated hollow fiber layers of gassing (grey), medium (pink), and counter-current flow medium (blue) fibers. Image B shows a cross-section of still images from a representative video sequence of counter-current flow medium perfusion. The images were taken using the Pearl Impulse Imaging system, IR Dye® 800CW and the 800 nm fluorescence channel. Numbered bioreactor ports (1-7) are clearly visible at the beginning of the sequence (B1) and the bioreactor remains in the same orientation throughout imaging. White arrows show counter-current medium perfusion (B1) and black arrows show medium short-cutting to adjacent rather than opposite exit ports (B4).
Figure 2. Spectral analysis of the hollow fiber bioreactor using an Ivis Spectrum imaging system and GFP to NIR range filters. Bioreactor ports are numbered from 1-7 (in white) to show orientation (black port numbers painted on the bioreactor casing have been obscured by signal in some places). Image A shows the minimal background noise signal can be achieved with the 710/780 excitation/emission filter combination. Image B shows blocked excitation and no detectable phosphorescence for bioluminescence experimentation. Image C is taken from a video sequence using the Ivis Kinetic Imaging system (710/780 excitation/emission filter combination) and shows Xenolight CF750 fluorescent dye-spiked counter-current medium perfusion (white arrows) and medium short cutting to adjacent rather than opposite exit ports (black arrows).

Figure 3. Ivis Kinetic Imager photographs of the bioreactor from two angles after inoculation of luciferase expressing 4T1-luc2 Bioware Ultra murine cells (6.4×10⁶ cells at 8×10⁵/mL density) into the cell chamber to assess inoculated cell distribution. Bioreactor ports are numbered from 1-7 (in white) to show orientation (black port numbers painted on the bioreactor casing have been obscured by signal in some places). Port 4 is used for cell inoculation into the cell chamber.
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