Automated Identification and Localization of Hematopoietic Stem Cells in 3D Intravital Microscopy Data

Reema A. Khorshed,1,* Edwin D. Hawkins,1 Delfim Duarte,1 Mark K. Scott,1,2,3 Olufolake A. Akinduro,1 Narges M. Rashidi,1,4 Martin Spitaler,2,5 and Cristina Lo Celso1,*

1Department of Life Sciences, Imperial College London, London SW7 2AZ, UK
2Facility for Imaging by Light Microscopy, Imperial College London, London SW7 2AZ, UK
3Present address: Centre for Dynamic Imaging, The Walter & Eliza Hall Institute, 1G Royal Parade, Parkville, VIC 3052, Australia
4Present address: The Ragon Institute of MGH, MIT and Harvard, 400 Technology Square, Cambridge, MA 02139, USA
5Present address: Max Planck Institute for Biochemistry, Am Klopferspitz 18, 82152 Martinsried, Germany

*Correspondence: r.khorshed@imperial.ac.uk (R.A.K.), c.lo-celso@imperial.ac.uk (C.L.C.)
http://dx.doi.org/10.1016/j.stemcr.2015.05.017
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SUMMARY

Measuring three-dimensional (3D) localization of hematopoietic stem cells (HSCs) within the bone marrow microenvironment using intravital microscopy is a rapidly expanding research theme. This approach holds the key to understanding the detail of HSC-niche interactions, which are critical for appropriate stem cell function. Due to the complex tissue architecture of the bone marrow and to the progressive introduction of scattering and signal loss at increasing imaging depths, there is no ready-made software to handle efficient segmentation and unbiased analysis of the data. To address this, we developed an automated image analysis tool that simplifies and standardizes the biological interpretation of 3D HSC microenvironment images. The algorithm identifies HSCs and measures their localization relative to surrounding osteoblast cells and bone collagen. We demonstrate how this tool is effective, consistent, and accurate in the analysis of the proposed approach compared to current manual analysis and its wider applicability to analyze other 3D bone marrow components.

INTRODUCTION

Precise regulation of somatic stem cell function is essential for the survival of multicellular living organisms ranging from C. elegans to humans. Somatic stem cells maintain themselves while their progeny turn over and differentiate to maintain the tissue they reside in throughout life. This process is deregulated during disease and aging; therefore, increasing attention has been dedicated to understanding somatic stem cells with an aim to improve both prevention and treatment of disease. The correct functioning of somatic stem cells depends on complex and dynamic interactions with specific cellular and molecular components of the microenvironment that surrounds them (together called “niche”) (Scadden, 2014), and in vivo imaging of stem cells is expanding and promising field that provides a unique perspective of their behavior in situ. To date, this approach has been directly responsible for generating new hypotheses on the crucial role of the stem cell niche (Ritsema et al., 2014; Rompolas et al., 2012).

Hematopoietic stem cells (HSCs) maintain the turnover of red blood cells, platelets, and immune cells. They reside in the bone marrow, where several cell types contribute to their regulation (Lo Celso et al., 2011; Morrison and Scadden, 2014). How the concerted action of multiple niche components regulates HSC fate is not clear, and understanding the localization of HSCs relative to multiple surrounding cellular and structural constituents of the bone marrow microenvironment is the first step toward solving this puzzle. Single-cell resolution intravital microscopy of fluorescently labeled HSCs and niche components allows direct observation of HSCs in mouse bone marrow. We, and others, have successfully used fluorescent dyes to label HSCs prior to transplantation to achieve their detection in vivo through the bone of anesthetized recipient mice (Köhler et al., 2009; Lo Celso et al., 2009).

The endosteal niche is a bone marrow microenvironment proximal to trabecular/cortical bone, lined by cells of the osteoblast lineage, and associated with effective HSC engraftment as well as maintenance of their long-term survival and function (Calvi et al., 2003; Kunisaki et al., 2013; Lane et al., 2011). Two-photon microscopy is essential to detect second harmonic generation (SHG) signal emitted by bone collagen and to recognize the location of bone marrow cavities and their endosteal surface (Lo Celso et al., 2011). Transgenic reporter mice expressing GFP under the control of an osteoblast-specific promoter (herein referred to as Col2.3GFP mice) (Kalajzic et al., 2002) are a powerful tool for visualizing the HSC endosteal niche. Using these and other reporter strategies, manual analysis of HSC in vivo images has provided the indication that normal HSCs localize near vasculature, endosteum, osteoblastic cells, and nestin-positive mesenchymal progenitor cells (Lane et al., 2011; Lo Celso et al., 2009; Méndez-Ferrer et al., 2010; Sanchez-Aguilera et al., 2011). However, such analysis has the following limitations: (1) it is time-consuming and, therefore, limited to measuring a few parameters; and (2) it is subject to human error, leading to...
intra- and inter-researcher inconsistencies. We reasoned that specialized image analysis tools would simplify the biological interpretation of 3D HSC microenvironment images, and they not only would provide unbiased data analysis but also expand the number of measurable parameters with the potential to uncover new aspects of HSC biology. Although several segmentation and classification methods have been developed for 2D (Chung and Vese, 2009; Saikumar et al., 2012; Yeo et al., 2011) and 3D datasets (Lou et al., 2014; Nandy et al., 2014; Pop et al., 2013), their application to in vivo bone marrow images is not straightforward due to the complexity of the structures of interest. In conjunction, light scattering caused by the surrounding tissue, especially by overlaying bone, limits the resolution of in vivo microscopy of bone marrow compared to that of other tissues or ex vivo techniques. To overcome these issues, we propose a local heterogeneity-based image segmentation (LH-SEG) approach that utilizes multi-resolution segmentation (Mallinis et al., 2008) and mean intensity difference to neighbor thresholding. This approach measures local morphological and intensity homogeneity and combines these values with neighborhood distance features to segment and threshold the objects of interest. To ensure reliable edge detection across objects with high-intensity heterogeneity, LH-SEG is applied on each 2D slice prior to 3D rendering.

HSCs labeled ex vivo using lipophilic membrane dyes such as 1,1'-Dioctadecyl-3,3,3',3'-Tetramethylindodicarbocyanine (DiD) generate a bright fluorescent signal (Lo Celso et al., 2009, 2011). However, these dyes lead to background signal from cell debris and aggregates (Progatzky et al., 2013), posing the extra challenge of filtering the signal from cell debris and aggregates (Progatzky et al., 2009, 2011). However, these dyes lead to back-}

RESULTS

Variability of Intravital Microscopy Datasets

Combined confocal/two-photon microscopy allows detection of HSCs by intravital microscopy of mouse calvarium bone marrow; however, a variety of acquisition settings can be used on different days and by different users, leading to overall brighter/dimmer images and a range of field-of-view sizes and depths. We aimed to develop an image analysis protocol sufficiently flexible to deal with such variations, as it is essential to eliminate the bottleneck of data processing.

To test the applicability of our analysis protocol to a wide range of HSC in vivo datasets, we worked with ten independently acquired image datasets, exhibiting variability at both the biological and image acquisition levels. Each dataset contained 6–12 3D stacks (fields of view), including one or multiple DiD signals, originated from multiple experiments (summarized in Table S1) as follows: control HSCs (from wild-type, untreated mice) injected into Col2.3GFP recipients; control HSCs injected into Col2.3GFP × Col2.3Dkk1 double-transgenic mice (in which the same promoter drives expression of GFP and the Wnt inhibitor Dkk1, causing HSCs to lose self-renewal ability through a still unknown mechanism (Fleming et al., 2008); and infection-exposed HSCs injected into Col2.3GFP recipients (these HSCs, having become motile, interact with the bone marrow microenvironment differently than controls [Rashidi et al., 2014]).

Of the ten datasets analyzed, five (datasets 5, 6, 8, 9, and 10) contained SHG bone signal in addition to DiD and GFP signals. Movie S1 shows one example of raw data (dataset 10, field of view 4). All images were encoded at 8 bits per pixel, at a resolution of 512 x 512 pixels, but z stacks had varying sizes, with 512 pixels in the x and y dimensions corresponding to 213–620 μm and 12–102 slices acquired with a step size of 1–5 μm, with each slice corresponding to 1 voxel in the z dimension. Moreover, each dataset was generated using unique acquisition settings, from exciting laser power to detectors’ gain and offset. The datasets used reflected the range of settings selected by individual researchers performing intravitual microscopy experiments, and they allowed us to immediately test whether our image analysis protocol would provide reliable results in independently acquired datasets.

LH-SEG

The two main challenges faced when segmenting bone marrow in vivo images are the following: (1) the unpredictable shapes and sizes of stroma components, as osteoblasts have asymmetrical contours and cluster into irregular groups, and bone cavities vary in sizes, from micropores to large areas occupied by bone marrow; and (2) the variable levels of fluorescence intensity, due to loss of signal with increasing depth and, in the case of osteoblasts, to the co-existence of cells expressing higher and lower levels of GFP within the same field of view and sometimes even within the same cluster. Similar challenges apply to the...
segmentation of HSCs, because, even though their shape is more homogeneous, they vary from one to another in terms of signal intensity due to their varying depth and DiD being diluted upon cell division (Figure 1).

To handle fluorescence intensity heterogeneity, we applied a convolution Gaussian blur filter with $(3 \times 3)$ kernel size (Moon, 2001), using a small kernel size to reduce intensity heterogeneity while conserving object structures.

To segment the objects of interest, we developed a two-step method that recapitulates how humans identify objects based on how their intensities compare to other objects in the surrounding neighborhood. We named this segmentation approach LH-SEG. To minimize artifacts due to depth-dependent loss of signal, this method was applied to each 2D slice.

Multi-resolution segmentation is the first step of LH-SEG. It breaks highly heterogeneous images into a number of smaller segments, each of them more homogeneous in terms of both texture and shape (Baatz and Schäpe, 2000). It starts with a single pixel and iteratively merges further pixels in a number of loops as long as a threshold of homogeneity is not exceeded locally, within the segment.

The homogeneity threshold depends on the value of the scale parameter $\alpha$, which reflects a combination of shape and texture homogeneity (Baatz and Schäpe, 2000). The selection of the value of this parameter depends on the physical structure of the objects of interest as well as their textural characteristics. Selecting high scale parameters results in fewer, larger segments that can be bigger than the object observed, while lower scale parameter values result in smaller segments. However, the smaller the $\alpha$, the longer the processing time (Table S2). The selection of the value of this parameter depends on the physical structure of objects as well as their textural characteristics; therefore, we selected the largest value that would provide efficient segmentation of each category of objects (HSCs, osteoblasts, and bone) throughout our datasets (Figure 2B).

To reconstruct the objects of interest from the segments obtained, we merged adjacent image segments based on the mean intensity difference to neighborhood (MDN) threshold. MDN threshold describes the difference between an image segment and its neighboring image segments, in terms of mean intensity values, and is defined as follows:

$$T_{\alpha}(v) = \frac{1}{w} \sum_{u \in N_V(d)} w_u \left[ c_k(v) - c_k(u) \right], \quad \text{(Equation 1)}$$

where $w$ is the image channel weight. Images are weighted by the distance between the segmented image objects, defined as follows:

$$w = \sum_{u \in N_V(d)} w_u \alpha^d \quad \text{(Equation 2)}$$

where $v$ and $u$ are two segmented image objects, $N_V$ is the direct neighbors to the segmented image object $v$, $u$ is defined as a direct neighbor to $v$ if the minimum distance between them is less than or equal to $d$, $d$ is the distance...
between neighboring segments and defined as the radius of
the segmented image object perimeter in pixel, \( w_u \) is the
weight of the segmented image object defined by the differ-
ence of the mean intensity value between \( v \) and \( u \) in a given
distance \( d \), and \( \tau_k \) is the mean intensity value of channel \( k \).

The appropriate MDN threshold for \( T_{\psi} \) and distance
feature \( d \) need to be selected for effective segmentation of
each image object category (HSC, osteoblast, and bone).

Progressive rounds of merges are repeated until the MDN
of the resulting object is equal to or greater than a selected
MDN threshold within a particular neighborhood dis-
tance. Given the different nature of HSCs, bone, and oste-
oblasts in terms of shape and intensity, we optimized
different values for the MDN threshold and neighborhood
distance parameters for each object category. DiD objects
showed low-intensity heterogeneity and required a higher

Figure 2. The LH-SEG Method
(A) Representative raw data from intravital
microscopy experiments. (Left) Typical
signal from a DiD-labeled HSC is shown.
(Middle) GFP signal from osteoblast cells in
a Col2.3GFP reporter mouse is shown.
(Right) SHG signal obtained from 840-nm
two-photon excitation of collagen to
resolve bone structures is shown.
(B) Results of multi-resolution segmenta-
tion of each cellular component in (A) are
shown.
(C) Results of MDN thresholding of each
cellular component in (B) are shown.
(D) Results of automated detection of the
components in (A) following segmentation
and thresholding are shown.
compared to bone and osteoblast clusters, which presented higher intensity heterogeneity. Conversely, the parameter distance $d$ depends on the object size. DiD objects covered a smaller neighborhood compared to osteoblasts and bone, which are much larger; therefore, we selected a smaller $d$ for DiD objects and a larger $d$ for bone and osteoblasts (Table S3). The protocol for parameter selection and optimization for these and other types of objects is available in the Supplemental Experimental Procedures.

Once the objects were segmented in each 2D slice (Figures 2C and 2D), we linked them across the z dimension, according to the original z step size, to form the final 3D objects (Figure 3). Once the parameters were optimized on one initial dataset, we could apply them to the remaining nine datasets without any further alteration.

**Machine Learning Classification of HSCs**

A further challenge posed by the analysis of intravital microscopy images of chemically labeled HSCs is the discrimination between genuine HSCs and similar signal generated as a consequence of cell death and dye aggregation, or shedding of the dye onto neighboring cells and structures (Progatzky et al., 2013). While LH-SEG eliminated...
the smallest DiD debris, aggregates and non-specifically labeled cells and structures remained in the segmented, 3D-reconstructed images and constituted false-positive signals that needed to be eliminated. As these objects share common characteristics with real HSCs, their elimination based solely on morphological and textural thresholds is a challenging task, and one often impossible to solve manually. To tackle this problem, we used a supervised 3D object-based classification approach to identify DiD signal as either positive (genuine) or negative (false) HSC objects. We selected the decision tree classifier (Agarwal and Sharma, 2011; Aydemir and Kayikcioglu, 2014) for this task due to its computational simplicity and illustrative attributes. The decision tree classifier weights boundaries between different classes based on their discriminative power. This type of classifier does not require a feature optimization task prior to classification; therefore, we could use a vast number of textural, intensity, and morphological features (Table S4) to train the classifier. It then automatically selected the discriminative features and the boundaries that defined different classes of DiD objects.

Due to the highly variable appearance of DiD objects, we manually prepared a training set containing three classes of DiD objects, based on the morphological characteristics (Figure 4). The first class, HSC-Class-1, comprises HSCs showing rounder shapes and smoother surfaces, as we would expect for quiescent, non-motile cells. The second class, HSC-Class-2, contains HSCs that have less rounded shapes and present small uropod protrusions (Krummel and Macara, 2006), as we previously observed in time-lapse images of migratory HSCs (Rashidi et al., 2014). The third class, False-HSCs, are DiD objects characterized by highly irregular morphology, for example, very pronounced protrusions, longer than the main diameter of the object itself, and objects that could represent cell doublets or clusters (Figure 4).

For the training task, two DiD objects were annotated as HSC-Class-1, five for HSC-Class-2, and 11 for False-HSCs (Figure 4). The number of samples was optimized for each class to avoid under-fitting and over-fitting the classifier. In fact, larger training samples resulted in increased false negatives. The size of each training set correlated with the heterogeneity of the objects within each class; thus, DiD objects that exhibited low variation, such as HSC-Class-1, required fewer samples compared to those with higher variation (HSC-Class-2 and False-HSCs).
To test whether machine learning could deal with our non-standardized datasets, all training samples were taken from one image dataset only (dataset 1, Table S1). We then tested the classifier on the nine remaining datasets. A 3-fold cross-validation was sufficient for training the classifier, as increasing the folds did not show any evidence of improving classification accuracy.

3D Localization: Distance Measurements
To minimize the complexity of the minimal distance search, we first extracted all the points that belonged to the surface of the HSC, osteoblast, and bone objects, and then we measured the distances between surface points of each object pair by computing their Euclidian distance (Ye, 1988), taking into account the anisotropic resolution of the image value, as follows:

\[
dist = \sqrt{(x_1 - x_2)^2 + (y_1 - y_2)^2 + (z_1 - z_2)^2},
\]

(Equation 3)

where dist is the distance between two points \((x_1, y_1, z_1)\) and \((x_2, y_2, z_2)\), and \(sd\) is the distance between slices.

To calculate the minimum distance from each HSC to its neighboring osteoblast, we selected the closest point of the nearest osteoblast and the closest point of that particular HSC. Closest points of each pair were pseudo colored for visual identification of the nearest edges of associated objects. All analysis up to this point was performed using pixel units, and the results obtained here were then converted to micrometers (Figures 5A–5D). Of note, when we extracted positional data for control and infection-exposed HSCs, their distributions relative to osteoblasts were equivalent. However, when in a further experiment we injected, imaged, and analyzed myeloid progenitor (MP) cells (Lin− c-Kit+ Sca-1−), they could be found further away from osteoblasts than HSCs. The position relative to endostal surface was slightly but significantly different for HSCs and exp.HSCs (Figures 5E and S5).

Performance Evaluation
Segmentation Accuracy
We evaluated the performance of the LH-SEG method by comparing its results to manually annotated data taken from each dataset. We also compared our proposed automated segmentation method to the following three commonly used segmentation approaches: automated local thresholding “Bernsen” (Sales et al., 2011), automated global thresholding (Otsu, 1979), and level-set segmentation, all available through the Fiji image adjustments and segmentation plugin (Schindelin et al., 2012). We evaluated each of these methods after optimization against the benchmarking manual segmentation and, therefore, could assess the improvement achieved by LH-SEG. We selected ten examples (one from each dataset) of DiD objects and GFP+ osteoblasts that exhibited different intensity, edge morphology, and neighborhood characteristics. Bone signal was excluded from the quantitative segmentation evaluation, because the size of this structure makes it difficult to correctly select the bone region for the manual benchmarking. A user selected a 2D slice and segmented the objects manually (benchmarking); the manual segment was then converted into a mask and placed over the automated segmentation mask for comparison. Regions of the automated segmentation that matched the manual segmentation regions were considered true positives (TP), regions of the automated segmentation that did not match with the manual segmentation regions were false positives (FP), and regions of the manual segmentation missed by the automated segmentations were false negatives (FN)(Figures S1 and S2).

In the case of the DiD objects (Figure S1), automated local thresholding parameters were optimized for the first image and provided good results; however, when the same parameters were used to segment DiD objects from other images, they led to poor segmentation. Thus, this method would require optimization for each individual image and would only give high TP values for DiD objects that exhibit high-intensity levels and result in high FN values when DiD objects exhibited medium to low intensities.

Global thresholding yielded poor results and was difficult to optimize due to the nature of DiD signal, which exhibits low contrast with the background and a high level of noise.

Level-set segmentation provided high TP in cases where HSCs exhibited high contrast between the cell edges and the background and increased FP in cases where a high level of background noise around the cell edges was found. Our proposed LH-SEG method maintained consistency in segmenting HSCs regardless of the DiD objects intensity, contrast level, and background noise and provided high TP and low FN and FP in all tested data using only one set of optimized parameters.

Automated local thresholding and level-set segmentation yielded poor results for osteoblasts, which exhibit high-intensity heterogeneity (Figure S2). Both methods were only able to detect osteoblastic regions that exhibited high-intensity levels compared to their surrounding neighborhood, making it impossible to optimize the parameters to work across all our image datasets. Osteoblastic regions of intermediate intensity were not detected using these methods and resulted in increased FP. Global thresholding resulted in increased FP when images included osteoblastic regions of heterogeneous intensities. However, when osteoblasts had homogeneous intensity and good contrast
compared to the background, global thresholding provided high TP and low FN and FP. Our proposed LH-SEG method maintained consistency in segmenting osteoblastic cell regions regardless of their intensity, contrast level, and background noise and provided high TP and low FN and FP across all ten tested datasets, despite using only one single set of optimized parameters.

To measure the accuracy of each automated segmentation method, we used the Jaccard similarity index defined as follows:

\[ J(M, A) = \frac{|M \cap A|}{|M \cup A|} \]  

(Equation 4)

where \( A \) represents the automated segmentation results and \( M \) is the manual benchmarking data. \( M \cap A \) is the TP regions, and \( M \cup A \) is the sum of TP, FP, and FN regions. LH-SEG achieved the highest and most reproducible \( J \) values for both HSC and osteoblast segmentation (Figure 6A). Qualitative evaluation of LH-SEG of bone showed no evidence of it being
Machine Learning Classification Accuracy

Manual classification (benchmark) was performed on unsegmented images and included three classes of objects as follows: HSC-Class-1, HSC-Class-2, and False-HSCs, as we used to train the classifier. We did not manually classify all False-HSCs as it was unfeasible due to their large number; therefore, any DiD object that was not classified in any of the three manual classes was automatically considered a benchmark False-HSC.

The classifier assessed all DiD objects present in the ten datasets (Table S1) and classified them as HSC-Class-1, HSC-Class-2, and False-HSCs according to the training set. This identified 14 HSC-Class-1 and 94 HSC-Class-2, including seven bona fide HSCs that were manually missed, however, correctly identified and classified by the classifier. All remaining signal was classified as False-HSCs. This was consistent with the fact that only few HSCs are observed in each imaged mouse (i.e., in each dataset) (Lo Celso et al., 2011). We then evaluated the classification results by calculating the Precision and Recall values defined in Equations 5 and 6.

\[
\text{Precision} = \frac{TP}{TP + FP} \quad \text{(Equation 5)}
\]

\[
\text{Recall} = \frac{TP}{TP + FN} \quad \text{(Equation 6)}
\]
Figure 7. Overview of Our Automated Pipeline and Its Applicability to Large Areas of Bone Marrow and a Wide Range of Niche Components

(A) Raw data are subjected to an initial pre-processing smoothing. LH-SEG identifies DiD-labeled objects, osteoblastic cells, and bone collagen; the 2D objects resulting from the segmentation are linked in 3D according to the raw data step size, then the decision tree classifier separates HSCs from DiD non-specific signal and HSC-to-osteoblast and HSC-to-bone shortest 3D distances are measured.

(B) (Left) 3D rendering of a calvarium tile scan including DiD-labeled HSC signal in red, GFP+ osteoblasts in green, and SHG bone collagen in blue. (Right) Higher magnification image of a detail of the tile scan is shown.

(legend continued on next page)
where \( TP \) represents the number of correctly classified bona fide HSCs, \( FP \) represents the number of false HSCs classified as bona fide HSCs, and \( FN \) represents the number of bona fide HSCs classified as false HSCs. The Precision value represents the fraction of correctly classified bona fide HSCs, while the Recall value represents the fraction of HSCs selected by the classifier that are bona fide HSCs. Precision and Recall are \( 1.0 \) if the classifier does not report any errors. Based on all tested 102 bona fide HSCs, only three bona fide HSCs were misclassified by machine learning and were \( FN \) and two classified HSCs were \( FP \) (Figure 6B), giving a Precision value of 0.98 and a Recall value of 0.97.

### 3D Localization Accuracy

Although segmentation accuracy plays a vital part in determining the efficacy of the minimum distance measurements, it is also important to evaluate the performance of the automated distance measurement algorithm. Distance measurement benchmarking was done manually (Lo Celso et al., 2009) using the commercial software Volocity, with users drawing lines between two points manually selected to be the closest edges of the selected DiD object and neighboring osteoblast/bone within unsegmented 3D images. Manual 3D distance measurements were set to 0 when DiD-labeled objects touched osteoblast or bone. All manual and automated measures are presented in Figure S3.

To evaluate the results of the automated 3D distance measurements, we calculated the error percentage for all ten datasets combined as defined in Equation 7.

\[
\text{% Error} = \left( \frac{\overline{d}(M) - \overline{d}(A)}{\overline{d}(M)} \right) \times 100, \quad \text{(Equation 7)}
\]

where \( \overline{d}(M) \) is the number of DiD objects used for the manual 3D distance measurement benchmarking, and \( \overline{d}(A) \) is the number of DiD objects where the automated 3D distance measurement matched the benchmarking measurements. Distance measurements \( \overline{d}_m \) and \( \overline{d}_a \) (Equation 8) were a match \( M \) if their difference was \( \leq 5 \) \( \mu m \), which is the upper margin of the error expected when measuring 3D distances manually (Figure 6C).

\[
|\overline{d}_m - \overline{d}_a| \leq 5 \mu m \times |M|. \quad \text{(Equation 8)}
\]

where \( \overline{d}_m \) is the manual distance measurement of HSCs to nearest osteoblast/bone, and \( \overline{d}_a \) is the automated one.

Automated HSC to osteoblast localization measurements proved to be accurate when compared to the manual 3D distance measurements (with 6.8% error based on the 102 HSCs measured). Furthermore, automated HSC-to-bone localization measurements resulted in no error based on all 43 HSCs in the datasets that had bone signal.

### Wide Applicability of LH-SEG and 3D Measurements

Given the size and complexity of bone marrow tissue, we examined whether our method would allow analysis of areas larger than a single field of view and of hematopoietic cells and niche components other than DiD-labeled HSCs, osteoblasts, and bone (Figure 7A). We injected DiD-labeled HSCs in a Col2.3GFP recipient, and, instead of searching for HSCs and acquiring fields of view that contained promising DiD signal, we set up a tiled acquisition of a calvarium area of approximately 4,413 \( \times \) 3,272 \( \times \) 125 \( \mu m \). Our method successfully segmented osteoblasts, bone, and DiD signal; classified DiD signal; and identified six bona fide HSCs in locations equivalent to those normally acquired as single fields of view (Figure 7B). Also, using the same approach, we were able to identify and measure tomato+ committed MPs (Figures 7C, 5E, and 5F). Machine learning classification was not applied to this type of cell, as it is only needed to distinguish chemically labeled cells from cell debris and aggregates.

Next we tested whether other bone marrow components would be analyzable using our method. We imaged wild-type recipients reconstituted with MacBlue bone marrow, in which cells of the monocytic-macrophage lineage express CFP (Hume, 2011), and injected with TRITC-labeled dextran to highlight all vasculature immediately prior to imaging. By identifying appropriate LH-SEG parameters for macrophages and vessels, we were able to segment and 3D render macrophages adjacent to bone marrow sinusoids (Figure 7D). Similarly, we identified tomato-expressing MPs injected in a nestin GFP recipient (Méndez-Ferrer et al., 2010) and measured their position relative to GFP-expressing nestin+ mesenchymal progenitors (Figure 7E; data not shown).

(C) (Left) 3D rendering of a calvarium tile scan including tomato+ progenitor cells in magenta, GFP+ osteoblasts in green, and SHG bone collagen in blue. (Right) Higher magnification image of a detail of the tile scan is shown.

(D) (Left) Representative example of automated detection of TRITC dextran-labeled blood vessels (purple) and CFP+ macrophages (yellow) following LH-SEG segmentation on a 2D slice. (Right) 3D rendering after segmentation and linking of each 2D slice in the z stack containing the image shown on the left. Inset shows higher magnification of a detail of the 3D rendering.

(E) (Left) Representative example of automated detection of nestin cells (green), tomato+ progenitors (red), and SHG bone signal (blue) following LH-SEG segmentation on a 2D slice. (Right) 3D rendering after segmentation and linking of each 2D slice in the z stack containing the image shown on the left.
DISCUSSION

We have developed an automated segmentation, classification, and localization measurement algorithm (Figure 7A) for analysis of confocal and two-photon microscopy images of HSCs, osteoblasts, and bone obtained through intravital microscopy of mouse bone marrow. In particular, we addressed the following two main challenges posed by HSC intravital microscopy image analysis: (1) non-standardized datasets, which vary based on each specific experiment and user; and (2) highly variable image quality and signal intensity, due to the complexity of the structures observed and the increasing signal loss and scatter with depth of imaging. The use of non-standardized datasets with varying acquisition settings (i.e., magnification, excitation power, detector gain, and z step size), reflecting the non-standardized nature of intravital microscopy experiments, allowed us to test and demonstrate the robustness of the proposed approach.

While the majority of segmentation algorithms work best with objects exhibiting a narrow range of signal intensity, LH-SEG could deal with highly heterogeneous objects because it recognizes the intensity differences between each object and its neighborhood, independently of the size of the object. This method, therefore, did not require any assumption on the size or shape of the objects segmented, and was particularly suited to segment bone and osteoblast clusters as they exhibit variable morphology.

Furthermore, machine learning classification of DiD objects separated genuine HSCs from DiD debris and was sufficiently flexible to recognize the variable appearance of HSCs. In our experience, this is the most subjective step of manual analysis of HSC images, leading to some HSCs not being included in the analysis (false negative) and some debris being further analyzed instead. The decision tree machine learning approach, which automates the feature optimization task, efficiently selected the few discriminative features that best identified HSCs from false signal. Of note, this method proved slightly more effective than manual analysis of 3D stacks, as it allowed detection and analysis of a higher number of HSCs (seven cells had not been recognized by the human eye). Most importantly, it sped up the analysis process and increased its objectivity, therefore improving current manual analysis. Of note, 3D stacks are acquired by definition in areas where the user has identified HSCs being present, and, therefore, the margin left for detection improvement is small. Our analysis method is critically helpful when large, non-pre-selected areas of bone marrow are acquired and analyzed, such as the 3D tiles we tested, which are extremely time consuming to be analyzed manually.

Our analysis algorithm segments, classifies, and localizes HSCs in their natural niche based on their morphology, texture, and surroundings, modeling the processes that the human brain uses to identify and understand relational features of the HSCs within an image. As a result, we obtained a high level of accuracy for segmentation, classification, and localization measurements, and only few errors, discussed in detail below, were reported for the classification of DiD objects (5/111 total cells identified by machine learning or manual user) and HSC localization (7/102 HSC-to-osteoblast and 0/43 HSC-to-bone measures). Classification errors occurred between HSC-Class-2 and False-HSCs, likely due to the common characteristics that both types of DiD objects share (less rounded shapes and uropod/irregular protrusions), which can mislead the classifier. Given their low frequency, these errors are unlikely to affect further analysis and, therefore, do not impinge on the validity of the automated results obtained.

Localization error was reported in circumstances where the automated approach measured the distance to a brighter osteoblast while a dimmer osteoblast was detected manually (2/102), or vice versa (2/102). In both cases, this type of error could be further rectified by increasing or decreasing the MDN threshold for the specific 3D stack; however, the resulting values may not be applicable to the majority of the other datasets. It was more efficient to have values that yielded rare errors across highly variable datasets than to have to manually select new values for each dataset generated. Another source of localization error was the position of HSC in the 3D stack (3/102): HSCs positioned at the edge of the xy field may be closer to osteoblasts located outside the field of view, and HSCs positioned too deep in the bone marrow yield unreliable measures because the signal is too distorted (and they could be closer to osteoblasts/endosteal surface on the cavity bottom [Lo Celso et al., 2009]). These HSCs are easily identifiable when collecting the data and simply should be eliminated from the analysis, as even manually it would be impossible to provide a reliable measurement.

Of note, the median of the differences between all manual and automated distance measurements was zero, indicating that the error in our method is unbiased relative to manual measurements and, therefore, our approach is not skewing the results obtained. Thus, the overall distribution of HSC positions observed with our automated algorithm was equivalent to that obtained manually from previously published datasets (Lane et al., 2011; Lo Celso et al., 2009), with the advantage that they were obtained in a fraction of the time.

As little is known about the spatial organization of hematopoietic and stroma cells in the bone marrow, our method will be useful to further elucidate the relative position of multiple cell types. Here we identified that
exp. HSCs, despite being motile, are found to have the same positional distribution relative to osteoblasts as steady-state HSCs, while MP distribution is wider. The observed differing distributions relative to osteoblasts and endosteme may reflect biological differences.

Our image analysis tool detects and analyzes single DiD-labeled HSCs. Methods to distinguish doublets, clusters, and dividing cells will require further refinement. However, the workflow described here allows automated segmentation, classification, and localization of HSCs during the initial stage of engraftment (homing), a critical step for the success of bone marrow transplantation therapeutic protocols, and one for which evidence has been accumulated of the correlation between HSC position relative to niche cells/components and their long-term function (Lo Celso et al., 2009; Lane et al., 2011). HSC proliferation may be better studied using different experimental approaches altogether, based on endogenous expression of fluorescent reporters rather than chemical dyes that dilute upon cell division (Progatzky et al., 2013).

Our approach was developed to detect and analyze DiD-labeled HSCs, osteoblasts, and bone collagen, but it was immediately expandable to the analysis of further bone marrow niche components, including ones with highly variable appearance, such as nestin cells and blood vessels. While analysis of large areas of bone marrow did not increase the number of HSCs detected per mouse, this approach considerably simplified and sped up acquisition and analysis of images, especially those with large numbers of cells, such as MPs. Based on our data, we can conclude that our approach is applicable to the analysis of a broad range of 3D and intravital microscopy images from other tissues and organs, leading to a faster pace of discovery.

**EXPERIMENTAL PROCEDURES**

**Generation of Raw Data by Intravital Microscopy**

All animal work was performed according to the UK Home Office Animals (Scientific Procedures) Act regulations and was approved by the Imperial College Ethics committee and by the Home Office. Intravital microscopy of HSCs was performed as described previously (Lo Celso et al., 2009; Rashidi et al., 2014), with the donor and recipient mice listed in Table S1. The lipophilic dye DiD was used to label all HSCs (LKS CD150⁺ CD48⁻ or LKS CD34⁺ Flk2⁻). In some experiments, Lin⁻ c-Kit⁺ Sca-1⁻ MPs were purified from the bone marrow of mT/mG donor mice (Muzumdar et al., 2002) and Nestin-GFP (Mendez-Ferrer et al., 2010) reporter mice, and identified as tomato⁺ cells. MacBlue bone marrow was a kind gift of Professor D. Hume (Edinburgh University) and was injected into irradiated wild-type recipients. Once fully reconstituted, mice were injected with 50 μl of 40 mg/ml TRITC-dextran to label blood vessels. All imaged mice were anesthetized and their scalps were removed and replaced with imaging windows (Rashidi et al., 2014). All microscopy was performed at Imperial College with a Leica SP5 (Facility for Imaging by Light Microscopy) and a Zeiss LSM 780 upright confocal/two-photon combined microscopes (Scott et al., 2014).

**Automated Method Implementation and Availability**

The presented method was developed in Definiens Developer XD 64. The source code is available in the Supplemental Information available online (Segment-Classify-Measure-distance-3D.dcp).

Computations were conducted on Intel-Core i5-3427U processor at 1.80 GHz and 4.0 GB RAM, running the 64-bit Windows 7 operating system. The total processing time per 3D stack is 5 min 26 s for a typical 512 × 512 × 18 (x,y,z)-pixel image size with three channels for DiD signal, GFP⁺ osteoblast cells, and SHG bone collagen signal, and where two real HSCs were identified. Analysis of large datasets can be carried out without manual supervision. The processing time varies depending on the 3D stack size/depth and the number of HSCs found.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, three figures, four tables, and one movie and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2015.05.017.

**ACKNOWLEDGMENTS**

The Kay Kendall Leukaemia Fund, Human Frontiers Science Program, Cancer Research UK, European Research Council, and Biotechnology and Biological Sciences Research Council supported this work. E.D.H. was supported by a European Hematology Association Fellowship. We thank Dr. D. Keller (Imperial College Facility for Imaging by Light Microscopy) for technical support; Professor D. Hume and Dr. C. Pridans for kindly providing MacBlue bone marrow; and Professor K. Duffy, Dr. L. Pizarro, and Dr. A. Dufour for comments on the manuscript.

Received: December 12, 2014
Revised: May 29, 2015
Accepted: May 29, 2015
Published: June 25, 2015

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Author/s:
Khorshed, RA; Hawkins, ED; Duarte, D; Scott, MK; Akinduro, OA; Rashidi, NM; Spitaler, M; Lo Celso, C

Title:
Automated Identification and Localization of Hematopoietic Stem Cells in 3D Intravital Microscopy Data

Date:
2015-07-14

Citation:
Khorshed, R. A., Hawkins, E. D., Duarte, D., Scott, M. K., Akinduro, O. A., Rashidi, N. M., Spitaler, M. & Lo Celso, C. (2015). Automated Identification and Localization of Hematopoietic Stem Cells in 3D Intravital Microscopy Data. STEM CELL REPORTS, 5 (1), pp.139-153. https://doi.org/10.1016/j.stemcr.2015.05.017.

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