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# **Directing Traffic in Lymph Nodes**

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How do the right cells get to the right place in lymph nodes? It is known that lymphocytes known as B cells (that originate in the bone marrow) migrate to follicles within the nodes, whereas T cells (that originate in the bone marrow and migrate to the thymus gland) reside in an adjacent region known as the paracortex. By combining confocal, electron, and intravital microscopy, Marc Bajénoff, Jackson Egen, Lily Koo, Jean Pierre Laugier, Frédéric Brau, Nicolas Glaichenhaus, and Ronald Germain have demonstrated a role for the stroma of the node in directing these cells to the appropriate location. The stromal cells that are critical in the B cell follicles are follicular dendritic cells (FDCs) and in the paracortex it's the fibroblastic reticular cells (FRCs).

Bajénoff *et al.* employed a variety of strategies to demonstrate the movement of lymphocytes, but their primary model was a mouse chimera. These mice were genetically engineered to express ubiquitin promoter-GFP (green fluorescent protein) then the mice were irradiated to kill all the hemopoietic tissue. These mice were then injected with normal bone marrow cells and the bone marrow was allowed to reconstitute. The cells of the host mice fluoresced when properly illuminated, whereas the "new" bone marrow cells did not. Using this model, they found FRCs provide direction for T cell migration. For example, T cells changed directions along FRC fibers about 93% of the time. Also, the FRCs appeared to be arranged along blood vessels of the lymph node, including the high endothelial venules, and regulated egress of T cells to the paracortex through cell-cell junctions referred to as "exit ramps." Further studies showed that FRCs played a role

in defining the border between T and B cell-occupied areas.

The movement of B cells also appeared to be influenced by FRCs, particularly as they moved through the paracortex. Their movement along the FRC network may enhance B cell stimulation as this would increase the likelihood of their encounters with antigen-laden dendritic cells. Once the B cells arrived in the follicles, their location appeared to be determined by FDCs.

The various imaging studies performed by Bajénoff *et al.* revealed that FRCs do not form an enclosed labyrinth of "corridors" that simply confine the lymphocytes, but rather they form a 3-dimensional meshwork of cell bodies and extended processes that physically interact with the lymphocytes. This provides guidance cues that direct T and B cell movement in the paracortex, and FDCs influence the B cells congregating in follicles.

A few decades ago, the stroma of an organ was generally considered the passive scaffolding that provided the spatial organization for the parenchymal cells that in turn performed the activities of the organ. Evidence has been increasing that the stroma plays more than a passive role. In an elegant series of studies using a variety of microscopic techniques, Bajénoff *et al.* have provided convincing evidence that the stroma of the lymph node plays a key guidance role in facilitating the interaction between rare antigen presenting cells and the corresponding antigen-specific lymphocytes within a densely populated lymph node. Bajénoff *et al.* hypothesized that this in turn promoted a normal immune response.

#### References

- 1 The authors gratefully acknowledge Drs. Ronald Germain and Marc Bajénoff for reviewing this article.
- 2 Bajénoff, M., J.G. Egen, L.Y. Koo, J.P. Laugier, F. Brau, N. Glaichenhaus, and R.N. Germain, Stromal cell networks regulate lymphocyte entry, migration, and territoriality in lymph nodes, *Immunity* 25:1-13, 2006.

Applications of Focused Ion Beam (FIB) on

## INDEX OF ARTICLES

Directing Traffic in Lymph Nodes3
Stephen W. Carmichael and Ellen D. Remstein, Mayo Clinic,
Rochester, MN
Multi-Length Scale Characterization of the Gibeon
Meteorite using Electron Backscatter Diffraction6
Matthew M. Nowell and John O. Carpenter, EDAX-TSL, Draper UT
SEM Provides Critical Process Information in Pharmaceutical Applications12
Ben Lich, FEI Company, Hillsboro, OR
Optimal Noise Filters in High-Resolution
Electron Microscopy16
K. Ishizuka, P. H. C. Eilers* and T. Kogure**, HREM Research
Inc., Higashimatsuyama, Japan, *Utrecht University, Utrecht,
The Netherlands, **University of Tokyo, Tokyo, Japan
Quantification of Contaminant Removal by Evactron
Cleaning Using Quartz Crystal Thickness Monitors22
Christopher G. Morgan, Mark M. Gleason and Ronald Vane
XEI Scientific, Inc., Redwood City, CA
Reconstructing What Was: Software Applied
to Serial Section TEM26
Marcia D. Feinberg and John C. Fiala, Boston University, Boston MA
Overcoming Challenges in Material Science Testing
with the use of Large Specimen SEM Analysis32
Adriana Romero, VisiTec of America LLC, Knoxville, TN
Microscopic analysis of magmatic crystals — Part 2: A SEM study of the stability of accessory zircon under
increasing metamorphic conditions36
Robert Sturm, Department of Materials Engineering and
Physics, University of Salzburg/Austria
Specimen Preparation for Condoms40
Gan Phay Fang, Science & Technology Innovative Centre, Ansell
Shah Alam Sdn Rhd, Selangor, Malaysia

10000 0011 00 01 1100 111 00 111 00 111 011 11 1	4Z
H. L. Hing <sup>1</sup> , C. Burkhardt <sup>2</sup> , P. Gnauck <sup>2</sup> , S. Sally <sup>3</sup> , H. Gelder- bloms <sup>4</sup> , Y. Muranaka <sup>5</sup> , M.A. Kaswandi 1, A.H. A. Aziz 1 & A	
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Reutlingen, Germany, <sup>3</sup> National University, Canberra, Australia, <sup>4</sup>	Rob-
ert Koch Institut, Berlin, Germany, 5 Hamamatsu University School	ol of
Medicine, Hamamatsu, Japan	
Advanced Metallographic Techniques Applied to	
Diesel Particulate Filters	44
Natalio Saenz, Heather Dillon, Shelley Carlson,	
& Gary Maupin, Battelle PNNL, Richland, WA	
New Approaches to Managing, Marketing, and Mone	У
for Maintaining a Core Facility (4Ms)	•
Part 3: Marketing and Managing a	
Research Core/Facility	46
Pankaj Sharma, Purdue University	
The Beginnings of the Southeastern Microscopy Society	52
W. Gray (Jay) Jerome, SEMS Historian	
Industry News	54
NetNotes	
Advertiser's Index	

### **ABOUT THE COYER**

The cover micrograph was obtained from a section of the Gibeon meteorite using Electron Backscatter Diffraction (EBSD) mapping. To analyze this size of sample, a combination beam-stage scanning approach was used. A Widmanstatten structure commonly associated with many iron meteorites is observed. See the article by Nowell and Carpenter starting on page six.