



May 17, 2012

## **Astrom Biosciences to Present at World Stem Cells & Regenerative Medicine Congress**

ANN ARBOR, Mich., May 17, 2012 (GLOBE NEWSWIRE) -- Astrom Biosciences, Inc. (Nasdaq:ASTM), the leading developer of patient-specific expanded multicellular therapies for the treatment of severe chronic cardiovascular diseases, today announced that company president and CEO Tim Mayleben will be presenting at the World Stem Cells & Regenerative Medicine Congress at the Park Plaza hotel in London, UK. The presentation entitled "Phase 3 Development of a Cellular Therapy Product" will take place on Monday, May 21, 2012 at 4:40 pm (BST).

The Astrom presentation will cover the benefits of a special protocol assessment and offer insights on achieving manufacturing readiness. The presentation will also address the role of clinicians and patients in the Phase 3 development process.

### **About Astrom Biosciences**

Astrom Biosciences is the leader in developing patient-specific, expanded multicellular therapies for use in the treatment of patients with severe, chronic cardiovascular diseases. The company's proprietary cell-processing technology enables the manufacture of ixmyelocel-T, a patient-specific multicellular therapy expanded from a patient's own bone marrow and delivered directly to damaged tissues. Astrom has advanced ixmyelocel-T into late-stage clinical development, including a Phase 3 clinical program to study patients with critical limb ischemia and a planned Phase 2b clinical trial in patients with ischemic dilated cardiomyopathy. For more information, please visit Astrom's website at [www.astrom.com](http://www.astrom.com). For more information on the pivotal REVIVE Phase 3 clinical trial, please visit the trial website at [www.revivecli.com](http://www.revivecli.com).

The Astrom Biosciences, Inc. logo is available at <http://www.globenewswire.com/newsroom/prs/?pkgid=3663>

*This document contains forward-looking statements, including, without limitation, statements concerning clinical trial plans and progress, objectives and expectations, clinical activity timing, intended product development, the performance and contribution of certain individuals and expected timing of collecting and analyzing treatment data, all of which involve certain risks and uncertainties. These statements are often, but are not always, made through the use of words or phrases such as "anticipates," "intends," "estimates," "plans," "expects," "we believe," "we intend," and similar words or phrases, or future or conditional verbs such as "will," "would," "should," "potential," "could," "may," or similar expressions. Actual results may differ significantly from the expectations contained in the forward-looking statements. Among the factors that may result in differences are the inherent uncertainties associated with clinical trial and product development activities, regulatory approval requirements, competitive developments, and the availability of resources and the allocation of resources among different potential uses. These and other significant factors are discussed in greater detail in Astrom's Annual or Transition Report on Form 10-K or 10-K/T, Quarterly Reports on Form 10-Q and other filings with the Securities and Exchange Commission. These forward-looking statements reflect management's current views and Astrom does not undertake to update any of these forward-looking statements to reflect a change in its views or events or circumstances that occur after the date of this release except as required by law.*

CONTACT: Media contact

Andrea Coan

Berry & Company

[acoan@berrypr.com](mailto:acoan@berrypr.com)

(212) 253-8881

Investor contact

Danielle Spangler

The Trout Group

[dspangler@troutgroup.com](mailto:dspangler@troutgroup.com)

(646) 378-2924