

**ACUTA announces the release of
ACUTA Regulatory Information Management (ARIM™)
version 3.2**

Marlborough, MA – February 8, 2018 - ACUTA is pleased to announce the release of ARIM version 3.2.

Highlights in v3.2:

- All modules in ARIM 3.2 support Australia eCTD submissions version 3.1.
- Support for Single Sign-On using OKTA.
- The 'My Favorites' feature has been added to the Dashboard.
- A Reports widget has been added to the Dashboard, providing access to the Study Status and Application Status reports.
- New enhancements have been added to the Veeva Vault integration.
- Users are able launch documents from previous sequences from the life-cycle.
- Users are able to add a Reason and enter the Location values for their signature when signing the PDF document.
- Registration Tracking users can now view the Formulation hierarchy in a Tree View.
- Replying to and resolving comments in a comment thread will now notify all involved users.
- The efficiency of loading the Sequence, Current and Cumulative Views has been improved.

Note that ARIM v2.3.3 will be supported in parallel to ARIM v3.2 until further notice. Our policy is to provide support for the current release plus two minor releases. We will communicate our plans to migrate multi-tenant clients to ARIM v3.2 during the coming weeks.

If you would like more information about this release, please contact ACUTA Technical Support team members at support@acutallc.com.

Sincerely,

ACUTA Product Management

www.acutallc.com

1-508-466-7799

Check out the following promotional videos.

[ARIM Viewer](#)

[ARIM Correspondence and Commitments](#)

The ARIM Cloud offering is powered by Microsoft Azure and advances the Life Science industry's urgent drive for an affordable solution using technology tailored to small and medium size company needs, with increased efficiencies and cost savings. For more information on ARIM, please visit <http://www.acutallc.com/arim>

About ACUTA, LLC

ACUTA was founded in 2012 to assist Life Science companies in complying with ever changing regulatory requirements that guide the product lifecycle through approval and maintenance. ACUTA's founder Shy Kumar and his team members are well known in the industry, with over 20 years of experience. They have successfully deployed state-of-the-art solutions that were well received by industry and users. ACUTA's vision is to develop innovative solutions to assist Life Sciences and related companies with their regulatory information management, which ultimately benefit everyone and specifically the needs of patients around the world. ACUTA is headquartered in Marlborough, MA, USA.