



February 8, 2017

ArQule to Present at The Leerink Partners 6th Annual Global Healthcare Conference on February 15, 2017

BURLINGTON, Mass.--(BUSINESS WIRE)-- ArQule, Inc. (Nasdaq: ARQL) today announced that Paolo Pucci, Chief Executive Officer, and Dr. Brian Schwartz, Chief Medical Officer and Head of Research and Development, will present at The Leerink Partners 6th Annual Global Healthcare Conference on February 15th, 2017, at 10:00 a.m. ET at the Lotte New York Palace in New York, New York.

You can access the live webcast of the presentation via the "Investors & Media" section of our website, www.arqule.com, under "Events & Presentations". A replay of the webcast will be available shortly after the conclusion of the presentation.

About ArQule

[ArQule](http://www.arqule.com) is a biopharmaceutical company engaged in the research and development of targeted therapeutics to treat cancers and rare diseases. Our mission is to discover, develop and commercialize novel small molecule drugs in areas of high unmet need that will dramatically extend and improve the lives of our patients. Our clinical-stage pipeline consists of five drug candidates, all of which are in targeted, biomarker-defined patient populations, making [ArQule](http://www.arqule.com) a leader among companies our size in precision medicine. ArQule's lead product, in phase 3 clinical development, is tivantinib (ARQ 197), an oral, selective inhibitor of the c-MET receptor tyrosine kinase, for second-line treatment of hepatocellular carcinoma in partnership with Daiichi Sankyo in the West and Kyowa Hakko Kirin in Asia. ArQule's proprietary pipeline includes: ARQ 087, a multi-kinase inhibitor designed to preferentially inhibit the fibroblast growth factor receptor (FGFR) family, in phase 2 for iCCA and in phase 1b for multiple oncology indications; ARQ 092, a selective inhibitor of the AKT serine/threonine kinase, in phase 1 for multiple oncology indications as well as ultra-rare Proteus syndrome, in partnership with the National Institutes of Health (NIH); ARQ 751, a next generation AKT inhibitor, in phase 1 for patients with AKT1 and PI3K mutations; and ARQ 761, a β -lapachone analog being evaluated as a promoter of NQO1-mediated programmed cancer cell necrosis, in phase 1/2 in multiple oncology indications in partnership with the University of Texas Southwestern Medical Center. In addition, we have advanced ARQ 531, an investigational, orally bioavailable, potent and reversible inhibitor of both wild type and C481S-mutant BTK, into toxicology testing and plan to file an Investigational New Drug Application in early 2017. ArQule's current discovery efforts are focused on the identification and development of novel kinase inhibitors, leveraging the Company's proprietary library of compounds. You can follow us on [Twitter](#) and [LinkedIn](#).

View source version on [businesswire.com](http://www.businesswire.com): <http://www.businesswire.com/news/home/20170208005029/en/>

ArQule, Inc.
Dawn Schottlandt, 781-994-0300
Sr. Director, Investor Relations/ Corp. Communications
www.arqule.com

Source: ArQule, Inc.

News Provided by Acquire Media