

# **EyePoint Pharmaceuticals to Present at Upcoming Investor Conferences**

August 28, 2018

WATERTOWN, Mass., Aug. 28, 2018 (GLOBE NEWSWIRE) -- EyePoint Pharmaceuticals, Inc. (NASDAQ:EYPT), a specialty biopharmaceutical company committed to developing and commercializing innovative ophthalmic products, today announced that Nancy Lurker, President and Chief Executive Officer, is scheduled to participate in the following investor conferences:

- B. Riley FBR Healthcare Conference on Tuesday, September 4 at 12:25 PM ET
- H.C. Wainwright & Co. 20<sup>th</sup> Annual Global Investment Conference on Wednesday, September 5 at 10:00 AM ET

A live audio webcast and subsequent archived replay of the H.C. Wainwright & Co. 20<sup>th</sup> Annual Global Investment Conference presentation may be accessed via the Investors section of the Company's website at <a href="https://www.eyepoint.com">www.eyepoint.com</a>. The replay will be available for 90 days after the event.

## **About EyePoint Pharmaceuticals**

EyePoint Pharmaceuticals, Inc. (formerly pSivida Corp.) (www.eyepointpharma.com), headquartered in Watertown, MA, is a specialty biopharmaceutical company committed to developing and commercializing innovative ophthalmic products in indications with high unmet medical need to help improve the lives of patients with serious eye disorders. The Company has developed three of only four FDA-approved sustained-release treatments for back-of-the-eye diseases. In addition, DEXYCU™ was approved by the FDA on February 9, 2018. DEXYCU, administered as a single intraocular dose at the end of ocular surgery for the treatment of postoperative inflammation, is the first and only FDA-approved intraocular product with this indication. DEXYCU employs the Verisome® extended-release drug delivery technology, which encompasses a broad number of related but distinct drug delivery systems capable of incorporating an extensive range of active agents, including small molecules, proteins and monoclonal antibodies. ILUVIEN® (fluocinolone acetonide intravitreal implant), a micro-insert for diabetic macular edema, licensed to Alimera Sciences, Inc., is currently sold directly in the U.S. and several EU countries. Retisert® (fluocinolone acetonide intravitreal implant), for posterior uveitis, is licensed to and sold by Bausch & Lomb, Inc. The New Drug Application (NDA) for EyePoint's lead product candidate, YUTIQ™ three-year treatment of non-infectious uveitis affecting the posterior segment of the eye, has been accepted for filing by the FDA and is currently under standard review with a Prescription Drug User Fee Act (PDUFA) date of November 5, 2018. The Company's pre-clinical development program is focused on using its core Durasert™ and the Verisome platform technologies to deliver drugs to treat wet age-related macular degeneration, glaucoma, and other diseases. To learn more about the Company, please visit www.eyepointpharma.com and connect on Twitter, LinkedIn, Facebook and Google+.

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Source: EyePoint Pharmaceuticals, Inc.