Aerie Pharmaceuticals to Present at the Cowen 40th Annual Healthcare Conference

February 24, 2020

DURHAM, N.C.--(BUSINESS WIRE)--Feb. 24, 2020-- Aerie Pharmaceuticals, Inc. (NASDAQ:AERI), an ophthalmic pharmaceutical company focused on the discovery, development and commercialization of first-in-class therapies for the treatment of patients with open-angle glaucoma, dry eye, retinal diseases and other diseases of the eye today announced that Vicente Anido, Jr., Ph.D., Chairman and Chief Executive Officer, will present in a fireside discussion at the Cowen 40th Annual Healthcare Conference on Monday, March 2, 2020 at 12:00 p.m. Eastern Time in Boston, MA. Dr. Anido will provide an Aerie overview and provide a business update.

The fireside discussion will be webcast live and may be accessed by visiting Aerie’s website at http://investors.aeriepharma.com. A replay of the webcast will be available for 10 business days.

About Aerie Pharmaceuticals, Inc.

Aerie is an ophthalmic pharmaceutical company focused on the discovery, development and commercialization of first-in-class therapies for the treatment of patients with open-angle glaucoma, dry eye, retinal diseases and other diseases of the eye. Aerie’s first product, Rhopressa® (netarsudil ophthalmic solution) 0.02%, a once-daily eye drop approved by the U.S. Food and Drug Administration (FDA) for the reduction of elevated intraocular pressure (IOP) in patients with open-angle glaucoma or ocular hypertension, was launched in the United States in April 2018. In clinical trials of Rhopressa®, the most common adverse reactions were conjunctival hyperemia, corneal verticillata, instillation site pain, and conjunctival hemorrhage. More information about Rhopressa®, including the product label, is available at www.rhopressa.com. Aerie’s second product for the reduction of elevated IOP in patients with open-angle glaucoma or ocular hypertension, Rocklatan® (netarsudil and latanoprost ophthalmic solution) 0.02%/0.005%, the first and only fixed-dose combination of Rhopressa® and the widely-prescribed PGA (prostaglandin analog) latanoprost, has been approved by the FDA and was launched in the United States in the second quarter of 2019. In clinical trials of Rocklatan®, the most common adverse reactions were conjunctival hyperemia, corneal verticillata, instillation site pain, and conjunctival hemorrhage. More information about Rocklatan®, including the product label, is available at www.rocklatan.com. Aerie continues to focus on global expansion and the development of additional product candidates and technologies in ophthalmology, including for dry eye, wet age-related macular degeneration and diabetic macular edema. More information is available at www.aeriepharma.com.

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Media: Tad Heitmann 949-526-8747; theitmann@aeriepharma.com
Investors: Ami Bavishi 908-947-3949; abavishi@aeriepharma.com

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