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Aerie Pharmaceuticals Announces Appointment of Huan Sheng, M.D., Ph.D., as Director Clinical Research and Drug Safety

IRVINE, Calif.--(BUSINESS WIRE)-- Aerie Pharmaceuticals, Inc. (NASDAQ:AERI), a clinical-stage pharmaceutical company focused on the discovery, development, and commercialization of first-in-class therapies for the treatment of glaucoma and other diseases of the eye, today announced the appointment of Huan Sheng, M.D., Ph.D., as Director Clinical Research and Drug Safety, reporting to Kristine Erickson, O.D., Ph.D., Aerie's Vice President Clinical Research. Dr. Sheng will be responsible for the development and implementation of Aerie's drug safety surveillance and clinical research strategy, and the management and oversight of Aerie's clinical development and drug safety programs. Dr. Sheng was most recently Senior Clinical Lead in Alcon's Global Clinical and Regulatory Affairs organization. In addition to previous experience at the Vistakon® Division of Johnson & Johnson Vision Care, Inc., as a Senior Clinical Research Scientist/Clinical Project Lead, Dr. Sheng also has extensive academic ophthalmology experience. Dr. Sheng is a licensed ophthalmologist and also holds a Ph.D. in Vision Science.

In connection with her acceptance of the position as Director Clinical Research and Drug Safety, Dr. Sheng will receive an award of 16,000 stock options. The stock options will vest over 4 years, with 25% vesting on the first anniversary of the hire date and the remainder vesting ratably on each of the subsequent 36 monthly anniversaries of the hire date. This award was made outside of Aerie's stockholder-approved equity incentive plan and was approved by the Company's independent directors as an inducement material to Dr. Sheng's entering into employment with the Company in reliance on NASDAQ Listing Rule 5635(c)(4), which requires this public announcement.

About Aerie Pharmaceuticals, Inc.

Aerie is a clinical-stage pharmaceutical company focused on the discovery, development and commercialization of first-in-class therapies for the treatment of patients with glaucoma and other diseases of the eye. Aerie's two lead product candidates are once-daily IOP-lowering therapies with novel mechanisms of action to treat patients with glaucoma or ocular hypertension. The NDA filing for Rhopressa™ (netarsudil ophthalmic solution) 0.02% was originally submitted in the third quarter of 2016 and is expected to be resubmitted near the end of the first quarter of 2017. The second product candidate, Roclatan™ (netarsudil/latanoprost ophthalmic solution) 0.02%/0.005%, which is a fixed dose combination of Rhopressa™ and widely prescribed PGA latanoprost, currently has two Phase 3 registration trials underway, named Mercury 1 and Mercury 2. If these trials are successful, a Roclatan™ NDA filing is expected to take place near year-end 2017. Aerie is also focused on the development of additional product candidates and technologies in ophthalmology.

Forward-Looking Statements

This press release contains forward-looking statements for purposes of the safe harbor provisions of the Private Securities Litigation Reform Act of 1995. We may, in some cases, use terms such as "predicts," "believes," "potential," "proposed," "continue," "estimates," "anticipates," "expects," "plans," "intends," "may," "could," "might," "will," "should," "exploring," "pursuing" or other words that convey uncertainty of future events or outcomes to identify these forward-looking statements. Forward-looking statements include statements regarding our intentions, beliefs, projections, outlook, analyses or current expectations concerning, among other things: the success, timing and cost of our ongoing and anticipated preclinical studies and clinical trials for our current product candidates, including statements regarding the timing of initiation and completion of the studies and trials; our expectations regarding the clinical effectiveness of our product candidates and results of our clinical trials; the timing of and our ability to obtain and maintain U.S. Food and Drug Administration or other regulatory authority approval of, or other action with respect to, our product candidates, including the expected timing of NDA filings for our product candidates; our expectations regarding the commercialization of our product candidates; the potential advantages of our product candidates; our plans to pursue development of our product candidates for additional indications and other therapeutic opportunities; our plans to explore possible uses of our existing proprietary compounds beyond glaucoma; our ability to protect our proprietary technology and enforce our intellectual property rights; and our expectations regarding strategic operations, including our ability to in-license or acquire additional ophthalmic products or product candidates. By their nature, forward-looking statements involve risks and uncertainties because they relate to

events, competitive dynamics and industry change, and depend on regulatory approvals and economic circumstances that may or may not occur in the future or may occur on longer or shorter timelines than anticipated. We discuss many of these risks in greater detail under the heading "Risk Factors" in the quarterly and annual reports that we file with the Securities and Exchange Commission (SEC). Forward-looking statements are not guarantees of future performance and our actual results of operations, financial condition and liquidity, and the development of the industry in which we operate may differ materially from the forward-looking statements contained in this press release. Any forward-looking statements that we make in this press release speak only as of the date of this press release. We assume no obligation to update our forward-looking statements whether as a result of new information, future events or otherwise, after the date of this press release.

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