

pSivida Corp. Announces Commercial Launch of ILUVIEN® in Germany with First Patient Treated

WATERTOWN, MA. May 7, 2013 -- pSivida Corp. (NASDAQ:<u>PSDV</u> - <u>News</u>), a leader in developing sustained release, drug delivery products for treatment of back-of-the-eye diseases, today announced that ILUVIEN[®], the first sustained release pharmaceutical product for the treatment of chronic diabetic macular edema (DME), is now commercially available in Germany. pSivida's licensee Alimera Sciences reported that the first patient has been treated following the commercial launch in Germany.

"We are very pleased ILUVIEN is now available in Germany as well as for privately insured and private pay patients in the U.K.," said Dr. Paul Ashton, President and Chief Executive Officer. "We are also very pleased by Alimera's resubmission of the NDA for ILUVIEN to the FDA and the recently announced PDUFA goal date of October 17, 2013. We will be entitled to 20% of net profits, as defined, in each of Germany and the U.K. from sales of ILUVIEN by Alimera. If the FDA approves ILUVIEN, we would also be entitled to an additional \$25 million milestone payment from Alimera as well as 20% of net profits on any sales in the U.S. by Alimera."

About pSivida Corp.

pSivida Corp., headquartered in Watertown, MA, develops tiny, sustained release, drug delivery products designed to deliver drugs at a controlled and steady rate for months or years. pSivida is currently focused on treatment of chronic diseases of the back of the eye utilizing its core technology systems, Durasert[™] and BioSilicon[™]. The injectable, sustained release micro-insert ILUVIEN® for the treatment of chronic DME considered insufficiently responsive to available therapies, licensed to Alimera Sciences, Inc., has received marketing authorization in Austria, France, Germany, Portugal, Spain and the U.K. and is awaiting authorization in Italy. Alimera has resubmitted the New Drug Application for ILUVIEN for DME to the U.S. Food and Drug Administration. pSivida plans to institute pivotal Phase III clinical trials for the treatment of posterior uveitis, a chronic back-of-the-eye disease, with the same micro-insert as ILUVIEN for DME. An investigator-sponsored clinical trial is ongoing for an injectable, bioerodible micro-insert to treat glaucoma and ocular hypertension. pSivida's FDA-approved Retisert® licensed to Bausch & Lomb Incorporated provides long-term, sustained drug delivery to posterior uveitis.

SAFE HARBOR STATEMENTS UNDER THE PRIVATE SECURITIES LITIGATION REFORM ACT OF 1995: Various statements made in this release are forward-looking, and are inherently subject to risks, uncertainties and potentially inaccurate assumptions. All statements that address activities, events or developments that we intend, expect or believe may occur in the future are forward-looking statements. The following are some of the factors that could cause actual results to differ materially from the anticipated results or other expectations expressed, anticipated or implied in our forward-looking statements: uncertainties with respect to: Alimera's ability to obtain regulatory approval for, and if approved, to finance, successfully commercialize and achieve market acceptance of, and generate revenues to pSivida from, ILUVIEN for DME in the U.S.; Alimera's ability to finance, achieve additional marketing approvals, successfully commercialize and achieve market acceptance of, and generate revenues to pSivida from, ILUVIEN for DME in the EU; outcome of reimbursement for ILUVIEN in the U.K.; financing and success of Phase III posterior uveitis trials including efficacy, side effects and

risk/benefit profile of the posterior uveitis micro-insert; initiation, financing and success of Latanoprost Product Phase II trials and exercise by Pfizer of its option; development of products using Tethadur and BioSilicon; initiation and completion of clinical trials and obtaining regulatory approval of product candidates; adverse side effects; ability to attain profitability; ability to obtain additional capital; further impairment of intangible assets; fluctuations in operating results; decline in royalty revenues; ability to, and to find partners to, develop and market products; termination of license agreements; competition and other developments affecting sales of products; market acceptance; protection of intellectual property and avoiding intellectual property infringement; retention of key personnel; product liability; consolidation in the pharmaceutical and biotechnology industries; compliance with environmental laws; manufacturing risks; risks and costs of international business operations; credit and financial market conditions; legislative or regulatory changes; volatility of stock price; possible dilution; possible influence by Pfizer; absence of dividends; and other factors described in our filings with the SEC. Given these uncertainties, readers are cautioned not to place undue reliance on such forward-looking statements. Our forwardlooking statements speak only as of the dates on which they are made. We do not undertake any obligation to publicly update or revise our forward-looking statements even if experience or future changes makes it clear that any projected results expressed or implied in such statements will not be realized.

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