

Scilex Holding Company announces the State of Indiana Medicaid will add Elyxyb as a preferred agent to its preferred drug list (PDL) effective October 1, 2023

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PALO ALTO, Calif., Sept. 25, 2023 (GLOBE NEWSWIRE) -- Scilex Holding Company (Nasdaq: SCLX, "Scilex" or "Company"), an innovative revenuegenerating company focused on acquiring, developing and commercializing non-opioid pain management products for the treatment of acute and chronic pain, announced the State of Indiana Medicaid will add Elyxyb as a preferred agent to its Preferred Drug List ("PDL") effective October 1, 2023.

As Indiana is a Single PDL State, where the five Indiana Managed Medicaid plans (Anthem, CareSource, MDwise, Managed Health Services (MHS), and UnitedHealthcare) must follow the State PDL, this change improves access to Elyxyb for all eligible patients under the Indiana Medicaid umbrella, the combined traditional Medicaid and Managed Medicaid populations, totaling approximately 1.2M lives, between the ages of 18-64.

Under the new terms, Elyxyb will be Preferred with the following prior authorization (PA) criteria:

- Migraine Dx, 18 & Older
- Trial and failure of 1 preferred triptan or contraindication to triptans
- Elyxyb will have a QL of 6 bottles.

"We are pleased to announce our first Elyxyb managed health care win with the Medicaid plan in Indiana, which we believe is an important step to expand utilization of Elyxyb throughout the U.S. Importantly, our diversified portfolio of unique, leading programs will increasingly allow us to offer patients complementary and standalone opioid-sparing outpatient pain management solutions," said Jaisim Shah, Chief Executive Officer of Scilex Holding Company.

About Scilex Holding Company

Scilex Holding Company is an innovative revenue-generating company focused on acquiring, developing and commercializing non-opioid pain management products for the treatment of acute and chronic pain. Scilex is uncompromising in its focus to become the global pain management leader committed to social, environmental, economic, and ethical principles to responsibly develop pharmaceutical products to maximize quality of life. Results from the Phase III Pivotal Trial C.L.E.A.R. Program for SEMDEXATM, its novel, non-opioid product for the treatment of lumbosacral radicular pain (sciatica), were announced in March 2022. Scilex participated in the type C meeting for purposes of pre-NDA discussion with the FDA and is pending official minutes in writing from the FDA. Scilex targets indications with high unmet needs and large market opportunities with non-opioid therapies for the treatment of patients with moderate to severe pain. Scilex launched its first commercial product ZTIido® in October 2018, in-licensed a commercial product Gloperba® in June 2022, and launched its third FDA-approved product ElyxybTM in April 2023. It is also developing its late-stage pipeline, which includes a pivotal Phase 3 candidate, and one Phase 2 and one Phase 1 candidate. Its commercial product, ZTIido® (Idocaine topical system) 1.8%, or ZTIido®, is a prescription lidocaine topical product approved by the U.S. Food and Drug Administration for the relief of pain associated with post-herpetic neuralgia, which is a form of post-shingles nerve pain. Scilex in-licensed the exclusive right to commercialize Gloperba® (colchicine USP) oral solution, an FDA-approved prophylactic treatment for painful gout flares in adults, in the U.S. Scilex in-licensed the exclusive rights to commercialize Gloperba® by 2024, and is well-positioned to market and distribute those product. Scilex's three product candidates are SP-102 (injectable dexamethasone sodium phosphate viscous gel product containing 10 mg dexamethasone), or SEMDEXATM in Phase 3, novel,

sodium phosphate viscous gel product containing 10 mg dexamethasone), or SEMDEXA[™], a Phase 3, novel, viscous gel formulation of a widely used corticosteroid for epidural injections to treat lumbosacral radicular pain, or sciatica, with FDA Fast Track status; SP-103 (lidocaine topical system) 5.4%, a Phase 2 study, triple-strength formulation of ZTlido®, for the treatment of acute low back pain, with FDA Fast Track status; and SP-104, 4.5 mg Delayed Burst Release Low Dose Naltrexone Hydrochloride (DBR-LDN) Capsule, for the treatment of chronic pain, fibromyalgia that has completed multiple Phase 1 trial programs and is expected to initiate Phase 2 trials in 2024.

Scilex Holding Company is headquartered in Palo Alto, California.

Forward-Looking Statements

This press release and any statements made for and during any presentation or meeting concerning the matters discussed in this press release contain forward-looking statements related to Scilex and its subsidiaries under the safe harbor provisions of Section 21E of the Private Securities Litigation Reform Act of 1995 and are subject to risks and uncertainties that could cause actual results to differ materially from those projected. Forward-looking statements include statements regarding the potential growth of Elyxyb in managed care plans, the potential benefits related to the addition of Elyxyb to the State of Indiana Medicaid PDL, Scilex's belief that it is well positioned to continue its growth over the next several years, Scilex's long-term objectives and commercialization plans, Scilex's potential to attract new capital, future opportunities for Scilex, Scilex's future business strategies, the expected cash resources of Scilex and the expected uses thereof; Scilex's current and prospective product candidates, planned clinical trials and preclinical activities and potential provals, as well as the potential for market acceptance of any approved products and the related market opportunity; statements regarding ZTIIdo®, Gloperba®, ELYXYB®, SP-102 (SEMDEXA[™]), SP-103 or SP-104, if approved by the FDA; Scilex's development and commercialization plans; and Scilex's products, technologies and prospects.

Risks and uncertainties that could cause Scilex's actual results to differ materially and adversely from those expressed in our forward-looking statements, include, but are not limited to: risks associated with the unpredictability of trading markets and whether a market will be established for Scilex's common stock; general economic, political and business conditions; risks related to the ongoing COVID-19 pandemic; the risk that the potential product candidates that Scilex develops may not progress through clinical development or receive required regulatory approvals within expected timelines or at all; risks relating to uncertainty regarding the regulatory pathway for Scilex's product candidates; the risk that Scilex will be unable to successfully market or gain market acceptance of its product candidates; the risk that Scilex's product candidates may not be beneficial to

patients or successfully commercialized; the risk that Scilex has overestimated the size of the target patient population, their willingness to try new therapies and the willingness of physicians to prescribe these therapies; risks that the outcome of the trials for SP-103 or SP-104 may not be successful; risks that the prior results of the clinical trials of SP-102 (SEMDEXA[™]), SP-103 or SP-104 may not be replicated; regulatory and intellectual property risks; and other risks and uncertainties indicated from time to time and other risks set forth in Scilex's filings with the Securities and Exchange Commission. Investors are cautioned not to place undue reliance on these forward-looking statements, which speak only as of the date of this release, and Scilex undertakes no obligation to update any forward-looking statement in this press release except as may be required by law.

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SEMDEXA[™] (SP-102) is a trademark owned by Semnur Pharmaceuticals, Inc., a wholly-owned subsidiary of Scilex Holding Company. A proprietary name review by the FDA is planned.

ZTlido® is a registered trademark owned by Scilex Pharmaceuticals Inc., a wholly-owned subsidiary of Scilex Holding Company.

Gloperba® is the subject of an exclusive, transferable license to use the registered trademark by Scilex Holding Company.

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