Medincell’s Partner Teva presented New Data Providing Schizophrenia Treatment Insights into Switching to UZEDY® (risperidone) from Invega Sustenna® (paliperidone palmitate)

Teva presented seven studies from its LAI schizophrenia research program at the Psych Congress Elevate 2024 Annual Meeting, which took place from May 30 to June 2 in Las Vegas, Nevada.

Presentations included data informing clinical strategies for switching patients to UZEDY, an extended-release injectable suspension of risperidone for subcutaneous use every one or two months for the treatment of schizophrenia in adults, from a once-monthly intramuscular injection of Invega Sustenna.

Eric Hughes, MD, PhD, Executive Vice President of Global R&D and Chief Medical Officer at Teva said: “UZEDY is a long-acting formulation of risperidone with an innovative delivery system that requires no loading dose or oral supplementation and can be dosed at one- or two-month intervals to help with prevention of relapse. It can also be administered subcutaneously under the skin instead of intramuscularly, which is an important feature to discuss with patients. We are proud to share these clinical insights that may help healthcare providers understand more about switching between long-acting options for their schizophrenia patients.”

Read here Teva’s press release

About Medincell

Medincell is a clinical- and commercial-stage biopharmaceutical licensing company developing long-acting injectable drugs in many therapeutic areas. Our innovative treatments aim to guarantee compliance with medical prescriptions, to improve the effectiveness and accessibility of medicines, and to reduce their environmental footprint. They combine active pharmaceutical ingredients with our proprietary BEPO® technology which controls the delivery of a drug at a therapeutic level for several days, weeks or months from the subcutaneous or local injection of a simple deposit of a few millimeters, entirely bioresorbable. The first treatment based on BEPO® technology, intended for the treatment of schizophrenia, was approved by the FDA in April 2023, and is now distributed in the United States by Teva under the name UZEDY® (BEPO® technology is licensed to Teva under the name SteadyTeq™). We collaborate with leading pharmaceutical companies and foundations to improve global health through new treatment options. Based in Montpellier, Medincell currently employs more than 140 people representing more than 25 different nationalities.

UZEDY® and SteadyTeq™ are trademarks of Teva Pharmaceuticals

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This press release contains forward-looking statements, including statements regarding Company’s expectations for (i) the timing, progress and outcome of its clinical trials; (ii) its clinical benefits and competitive positioning of its product candidates; (iii) its ability to obtain regulatory approvals, commence commercial production and achieve market penetration and sales; (iv) its future product portfolio; (v) its future partnering arrangements; (vi) its future capital needs, capital expenditure plans and ability to obtain funding; and (vii) prospective financial matters regarding its business. Although the Company believes that its expectations are based on reasonable assumptions, any statements other than statements of historical facts that may be contained in this press release relating to future events are forward-looking statements and subject to change without notice, factors beyond the Company’s control and the Company’s financial capabilities.

These statements may include, but are not limited to, any statement beginning with, followed by or including words or phrases such as “objective”, “believe”, “anticipate”, “expect”, “foresee”, “aim”, “intend”, “may”, “anticipate”, “estimate”, “plan”, “project”, “will”, “may”, “probably”, “potential”, “should”, “could” and other words and phrases of the same meaning or used in negative form. Forward-looking statements are subject to inherent risks and uncertainties beyond the Company’s control that may, if any, cause actual results, performance, or achievements to differ materially from those anticipated or expressed explicitly or implicitly by such forward-looking statements. A list and description of these risks, contingencies and uncertainties can be found in the documents filed by the Company with the Autorité des Marchés Financiers (the “AMF”) pursuant to its regulatory obligations, including the Company’s registration document, registered with the AMF on September 4, 2018, under number I. 18-062 (the “Registration Document”), as well as in the documents and reports to be published subsequently by the Company. In particular, readers’ attention is drawn to the section entitled “Facteurs de Risques” on page 26 of the Registration Document.

Any forward-looking statements made by or on behalf of the Company speak only as of the date they are made. Except as required by law, the Company does not undertake any obligation to publicly update these forward-looking statements or to update the reasons why actual results could differ materially from those anticipated by the forward-looking statements, including in the event that new information becomes available. The Company’s update of one or more forward-looking statements does not imply that the Company will make any further updates to such forward-looking statements or other forward-looking statements. Readers are cautioned not to place undue reliance on these forward-looking statements.

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