

MC2 Therapeutics's wynzora cream gets FDA nod to treat Plaque Psoriasis

21 November 2019 | News

The pivotal phase 3 trial demonstrated that Wynzora™ Cream treatment has a substantial and statistically significantly greater efficacy compared to Taclonex® Topical Suspension



MC2 Therapeutics, an emerging pharmaceutical company focused on novel PAD™ Technology-based topical therapies for chronic inflammatory conditions, has announced that the U.S. Food and Drug Administration (FDA) has accepted for review the New Drug Application (NDA) for Wynzora™ Cream. MC2 Therapeutics is seeking marketing approval for Wynzora™ Cream for the treatment of plaque psoriasis. The FDA has set July 20th, 2020 as the Prescription Drug User Fee Act (PDUFA) action date.

MC2 Therapeutics' NDA for WynzoraTM Cream is comprised of extensive quality, non-clinical and clinical data. Specifically, data from the pivotal phase 3 trial demonstrated that WynzoraTM Cream treatment has a substantial and statistically significantly greater efficacy compared to Taclonex® Topical Suspension ("Taclonex®") based on treatment success defined as a minimum two-point decrease in the Physician Global Assessment (PGA) score to clear or almost clear disease (40.1% versus 24.0%, p < 0.0001).

"We are very proud of the overall clinical profile of Wynzora™ Cream and look forward to continuing our interaction with the FDA during the NDA review", said Jesper J. Lange, President & CEO of MC2 Therapeutics and added "In parallel we will continue our efforts to ensure widespread patient access to Wynzora™ Cream pending approval".