



Sun Pharma announces the FDA approval for Ximino

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Sun Pharmaceutical has announced that the US Food and Drug Administration (US FDA) has approved its Supplemental New Drug Application (sNDA) for Ximino (Minocycline HCl) extended-release capsules 45 mg, 90 mg and 135 mg.

Ximino extended-release capsules are indicated for inflammatory lesions of non-nodular moderate to severe acne vulgaris in patients 12 years of age and older.

This approval further strengthens the Company's branded dermatology portfolio in the US. It expects Ximino extended-release capsules to be available for patients during the fourth quarter of 2015.