

Alembic Pharma gets USFDA permission for anti-depression drug

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"The company has received approval from the United States Food and Drug Administration (USFDA) for its abbreviated new drug application (ANDA) for Bupropion Hydrochloride tablets," Alembic Pharmaceuticals said in a BSE filing.

The approval is for strengths of 75 mg and 100 mg.

The approved product is therapeutically equivalent to the reference listed drug product (RLD) Wellbutrin tablets of GlaxoSmithKline LLC.

Quoting IMS December 2017 data, the company said Bupropion Hydrochloride tablets had an estimated market size of USD 37 million.