



Biocon Limited
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CIN : L24234KA1978PLC003417

www.biocon.com

April 26, 2021

To The Manager, BSE Limited Department of Corporate Services Phiroze Jeejeebhoy Towers, Dalal Street, Mumbai – 400 001 Scrip Code - 532523	To The Manager, National Stock Exchange of India Limited Corporate Communication Department Exchange Plaza, Bandra Kurla Complex Mumbai – 400 050 Scrip Symbol - Biocon
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Subject: Company Statement

Dear Sir/Madam,

Please find attached a Company Statement titled “**Biocon Biologics and Viatrix Receive European Commission Approval for Biosimilar Bevacizumab.**”

The above information will also be available on the website of the Company at www.biocon.com.

Kindly take the same on record and acknowledge.

Thanking You,

Yours faithfully,

For **Biocon Limited**

Mayank Verma
Company Secretary and Compliance Officer



NOTIFICATION TO STOCK EXCHANGE

COMPANY STATEMENT

Biocon Biologics and Viatris Receive European Commission Approval for Biosimilar Bevacizumab

Bengaluru, India; April 26, 2021:

“This is to inform that **Biocon Biologics Ltd.**, a subsidiary of **Biocon Ltd. (BSE code: 532523, NSE: BIOCON)**, has announced that **Abevmy® 100 & 400 mg**, a **biosimilar of Bevacizumab co-developed with Viatris Inc. (NASDAQ: VTRS)** has **received marketing authorization approval** from the **European Commission** following the positive recommendation by the Committee for Medicinal Products for Human Use (CHMP) of the European Medicines Agency.

Abevmy® 100 & 400 mg, a **biosimilar Bevacizumab**, is approved for the treatment in **metastatic colorectal carcinoma, metastatic breast cancer, non-small-cell lung carcinoma, glioblastoma, ovarian, cervical and renal cancer** as part of a specific regimen.

The centralized marketing authorization granted by the EC is valid in all EU Member States as well as in the European Economic Area (EEA) countries Iceland, Liechtenstein and Norway.

“The European Commission’s approval of our biosimilar Bevacizumab will enable us to offer this biologic therapy to cancer patients in the EU along with our partner Viatris. The addition of biosimilar Bevacizumab will strengthen our portfolio of biosimilars for cancer in the EU, which include biosimilar Trastuzumab and biosimilar Pegfilgrastim. This approval is an outcome of a great team effort and years of hard work and underlines our commitment to expand affordable access to life-saving biosimilars and make an enduring impact on global health.”

-- Company Spokesperson, Biocon Biologics.

For more information

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