



Press Release

Wednesday, September 14, 2011

Strides Arcolab Limited,
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BSE: 532531
NSE: STAR

CONTINUED US FDA APPROVAL STATUS FOR ORAL DOSAGE FACILITY

“Zero Inspectional observation”

September 14, 2011, Bangalore: Strides Arcolab is pleased to announce that its Oral Dosage Forms manufacturing site (KRS Gardens) in Bangalore was recently inspected by the USFDA as part of GMP compliance audit and the facility continues to be approved with “Zero Inspectional observation” reported in FDA 483.

The last US FDA inspection and approval for this facility was in the year 2008.

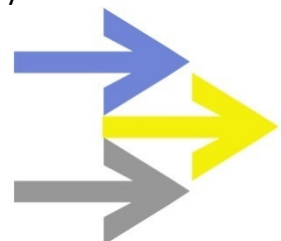
The KRS Gardens facility in Bangalore manufactures oral dosage forms such as tablets and capsules (both hard gelatine and soft gelatine). The manufacturing plant supports important current and future submissions for the US Market.

Manish Gupta, CEO – Pharma of Strides said “this continued approval of the site with Zero observation in FDA 483 augurs very well for the Pharma division of the Company in this challenging regulatory and business environment”.

About Strides Arcolab Limited

Strides Arcolab, listed on the Bombay Stock Exchange Limited (532531) and National Stock Exchange of India Limited (STAR), is a global pharmaceutical company headquartered in Bangalore, India that develops and manufactures a wide range of IP-led niche pharmaceutical products with an emphasis on sterile injectables.

The Company has 14 manufacturing facilities across 6 countries with presence in more than 75 countries in developed and emerging markets. Manufacturing is ably supported by a 350-scientist strong global R&D Centre located in Bangalore.



Additional information is available at the company's website at www.stridesarco.com.

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