

Mylan Receives Approval for Generic Version of Procardia XL(R) Tablets

PITTSBURGH, June 24, 2010 /PRNewswire via COMTEX News Network/ -- Mylan Inc. (Nasdaq: MYL) today announced that its subsidiary Mylan Pharmaceuticals Inc. has received final approval from the U.S. Food and Drug Administration (FDA) for its Abbreviated New Drug Application (ANDA) for Nifedipine Extended-release Tablets USP, 30 mg, 60 mg and 90 mg, the generic version of Pfizer's Procardia XL(R) Tablets, a treatment for angina and/or hypertension.

Nifedipine Tablets had U.S. sales of approximately \$234 million for the 12 months ending March 31, 2010, according to IMS Health.

Currently, Mylan has 137 ANDAs pending FDA approval representing \$94.9 billion in annual brand sales, according to IMS Health. Forty-one of these pending ANDAs are potential first-to-file opportunities, representing \$20.4 billion in annual brand sales, for the 12 months ending Dec. 31, 2009 according to IMS Health.

Mylan Inc. ranks among the leading generic and specialty pharmaceutical companies in the world and provides products to customers in more than 140 countries and territories. The company maintains one of the industry's broadest and highest quality product portfolios supported by a robust product pipeline; operates one of the world's largest active pharmaceutical ingredient manufacturers; and runs a specialty business focused on respiratory, allergy and psychiatric therapies. For more information, please visit www.mylan.com.

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