

Consumer Product Safety Improvement Act (CPSIA) General Conformity Certificate

As required by section 102 of the Consumer Product Safety Improvement Act of 2008 , codified at 15 U.S.C. §2063(a), the following certifies the product referenced meets all applicable testing standards

Amgen NDC Number: 55513-0073-07
Amgen Drug Product: 038 FDP US 30mg 7Tbl Sample
Dosage: 30mg
Finished Drug Product Lot Number: 1051383

Consumer Product Safety Commission (CPSC) safety requirements to which the above-referenced product is being certified: Child-resistant packaging requirements under the Poison Prevention Packaging Act. The applicable CPSC regulations are codified at 16 CFR Parts 1700 and 1701.

Date of (month/year) and place of Manufacture of Finished Drug Product:

Date of Manufacture: July 2014

Place of Manufacture: Patheon, Inc., Toronto Region Operations
2100 Syntex Court
Mississauga, Ontario L5N7K9
Canada

Drug Product Package Packaging Testing [Per Amgen Technical Assessment # TA-004171](#)

Date of testing: [5/21/2001](#)

Location of testing (incl. city/county): [Perritt Laboratories, Hightstown, NJ 08520](#)

U.S Importer of Record	Amgen U.S.A 12000 Plantside Drive Louisville, KY 40299 502-266-2706
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For additional information, please contact the following party responsible for maintaining the test results:

Amgen Inc Attn: Amgen Medical Information One Amgen Center Drive Thousand Oaks, CA 91320 Phone Number: 800-77-AMGEN Contact fax Number: 866-29-AMGEN Email: www.amgenmedinfo.com
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