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Product Details for NDA 202155

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ELIQUIS (APIXABAN) 2.5MG	Marketing Status: Prescription
Active Ingredient: APIXABAN Proprietary Name: ELIQUIS Dosage Form; Route of Administration: TABLET; ORAL Strength: 2.5MG Reference Listed Drug: Yes Reference Standard: No TE Code: Application Number: N202155 Product Number: 001 Approval Date: Dec 28, 2012	

Applicant Holder Full Name: BRISTOL MYERS SQUIBB CO PHARMACEUTICAL RESEARCH INSTITUTE
Marketing Status: Prescription
[Patent and Exclusivity Information](#)

ELIQUIS (APIXABAN)
5MG
Marketing Status: Prescription

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