Report Submitted 4/23/2021 to the Vermont Office of the Attorney General for Introduction of a New Prescription Drug to Market Janssen Pharmaceuticals, Inc.

Information required pursuant to 18 VSA § 4637(c),(d)

PONVORYTM

Requirement	Submission - PONVORY TM
18 VSA § 4637(c)(1) A description of the marketing and pricing plans used in the launch of the new drug in the United States and internationally	While specific marketing and pricing plans are not available in the public domain, generally we plan to market in the US and promote to appropriate healthcare professionals who treat individuals diagnosed with Relapsing forms of Multiple Sclerosis (RMS). The pricing plan has WAC set for a 30 day supply of the 20 mg maintenance dose at \$8,083.50 and WAC for the Multiple Strength, Titration Pack 14 set at \$3,772.30 . The list price of PONVORY TM is not reflective of discounts and rebates which may be available through Medicaid, Medicare, and commercial insurance. PONVORY TM will also be discounted as required under the 340B program, Federal Supply Schedule, and other government programs. International approvals are pending.
18 VSA § 4637(c)(2) The estimated volume of patients who may be prescribed the drug	Janssen's estimated volume of patients who may be prescribed PONVORY is not available in the public domain.
18 VSA § 4637(c)(3) Whether the drug was granted breakthrough therapy designation or priority review by the FDA prior to final approval	The drug was not granted breakthrough therapy designation nor priority review
18 VSA § 4637(c)(4) The date and price of acquisition if the drug was not developed by the manufacturer.	Not an acquisition