Global Misoprostol Registration by Indication

Registration is the process by which a drug is approved by a regulatory agency for importation, distribution and marketing for a specific medical indication after a thorough review of its effectiveness, safety and manufacturing process. It is a key strategy and critical first step to improving access to any drug or device.

This registration map is prepared by Venture Strategies Innovations (VSI) and reflects the organization’s most current knowledge of regulatory approvals of misoprostol globally.

For further information visit www.vsinnovations.org, or to share updated misoprostol registration information with VSI, contact comm@vsinnovations.org.

*Misoprostol may or may not be registered for gastric ulcers