## **Press Release**



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## Dr. Reddy's confirms its voluntary nationwide recall of all Ranitidine products in the U.S. Market

Hyderabad, India, October 23, 2019

For Immediate Release

Hyderabad, India and Princeton, NJ, USA. October 23, 2019 — Dr. Reddy's Laboratories Ltd. (BSE: 500124, NSE: DRREDDY, NYSE: RDY, along with its subsidiaries, together referred to as "Dr. Reddy's") confirms it had initiated a voluntary nationwide recall on October 1, 2019, (at the retail level for over-the-counter products and at the consumer level for prescription products) of all of its ranitidine medications sold in US due to confirmed contamination with N-Nitrosodimethylamine (NDMA) above levels established by the FDA. This recall follows the USFDA's caution note alerting patients and health care professionals that NDMA was found in certain samples of ranitidine. To date, Dr. Reddy's has not received any reports of adverse events related to the recall of Dr. Reddy's Ranitidine products. The recall includes all quantities in the US that are within expiry.

Risk Statement: NDMA is classified as a probable human carcinogen (a substance that could cause cancer) based on results from laboratory tests. NDMA is a known environmental contaminant and found in water and foods, including meats, dairy products, and vegetables.

Ranitidine is available as an over-the-counter (OTC) and prescription drug. Over-the-counter (OTC) ranitidine tablets are used to relieve heartburn associated with acid indigestion and sour stomach. OTC Ranitidine Tablets are also used to prevent heartburn associated with acid indigestion and sour stomach brought on by eating or drinking certain foods and beverages. Prescription ranitidine capsules are prescribed for the short-term treatment of active duodenal ulcer; maintenance therapy for duodenal ulcer patients at reduced dosage after healing of acute ulcers; treatment of pathological hypersecretory conditions (eg, Zollinger-Ellison syndrome and systemic mastocytosis); short-term treatment of active, benign gastric ulcer; maintenance therapy for gastric ulcer patients at reduced dosage after healing of acute ulcers; treatment of GERD (Gastroesophageal reflux disease); treatment of endoscopically diagnosed erosive esophagitis; and for maintenance of healing of erosive esophagitis.

Dr. Reddy's Ranitidine products can be identified by NDC numbers on the product label. All Ranitidine products with expiration dated September 2019 to June 2021 are being recalled:

Description	Strength	Туре	Pack	NDC
Ranitidine Capsules 150mg, 60	150 mg	Rx	60 ct bottle	5511112960
Ranitidine Capsules 150mg, 500	150 mg	Rx	500 ct bottle	5511112905
Ranitidine Capsules 300mg, 30	300 mg	Rx	30 ct bottle	5511113030
Ranitidine Capsules 300mg, 100	300 mg	Rx	100 ct bottle	5511113001
Ranitidine Tablets, USP 150mg,190(2x95)Tray (Sam's Club)	150 mg	ОТС	190 ct (2x95) tray	150062076 (UPC Code 078742089720)
Ranitidine Tablets, USP 150mg, 95 (Walgreens)	150 mg	ОТС	95 ct bottle	0363-0010-62
Ranitidine Tablets, USP 150 mg 220 CT Btl (Walmart)	150 mg	ОТС	220 ct bottle	49035-404-65
Ranitidine Tablets, USP 150mg 50ct Btl (Kroger)	150 mg	OTC	50 ct bottle	30142-505-50
Ranitidine Tablets, USP 150mg 24ct Btl (Kroger)	150 mg	OTC	24 ct bottle	30142-505-34
Ranitidine Tablets, USP 150mg 65 Ct Btl (Walgreens)	150 mg	ОТС	65 ct bottle	0363-0010-61
Ranitidine Tablets, USP 150 TAB 65ct BTL CP32 (Walmart)	150 mg	ОТС	65 ct bottle	49035-404-61
Ranitidine Tablets, USP 150 Tab 200Ct Btl (Walgreens)	150 mg	ОТС	200 ct bottle	0363-0010-01
Ranitidine Tablets, USP 150mg Tabs Btl, 24 (Walgreens)	150 mg	ОТС	24 ct bottle	0363-0010-34
Ranitidine Tablets, USP 75 TAB 30ct Bottle NG (CVS)	75 mg	ОТС	30 ct bottle	69842-871-30
Ranitidine Tablets, USP 75mg Tab 30Ct Btl (Walgreens)	75 mg	ОТС	30 ct bottle	0363-0131-30

75 mg	ОТС	80 ct bottle	0363-0131-80
75 mg	ОТС	80 ct bottle	69842-871-80
75 mg	ОТС	160 ct bottle	69842-871-37
75 mg	OTC	30 ct bottle	30142-131-30
150 mg	ОТС	24 ct bottle	63868-480-24
150 mg	ОТС	130 ct bottle	49035-404-13
150 mg	ОТС	50 ct bottle	63868-480-50
75 mg	ОТС	60 ct bottle	55111-131-60
75 mg	ОТС	60 ct bottle	63868-482-60
75 mg	ОТС	30 ct bottle	63868-482-30
150 mg	ОТС	24 ct bottle	55111-404-34
150 mg	ОТС	95 ct bottle	43598-808-62
150 mg	ОТС	220 ct bottle	43598-808-65
150 mg	ОТС	40 ct bottle	11673-849-40
150 mg	ОТС	24 ct bottle	71713-203-02
150 mg	ОТС	95 ct bottle	71713-203-05
75 mg	ОТС	All counts	57896-715
150 mg	ОТС	All counts	57896-717
	75 mg 75 mg 75 mg 75 mg 150 mg 150 mg 150 mg 75 mg 75 mg 75 mg 75 mg 150 mg	75 mg OTC 75 mg OTC 75 mg OTC 75 mg OTC 150 mg OTC 150 mg OTC 150 mg OTC 75 mg OTC 150 mg OTC	75 mg         OTC         80 ct bottle           75 mg         OTC         160 ct bottle           75 mg         OTC         30 ct bottle           150 mg         OTC         24 ct bottle           150 mg         OTC         130 ct bottle           150 mg         OTC         50 ct bottle           75 mg         OTC         60 ct bottle           75 mg         OTC         60 ct bottle           75 mg         OTC         30 ct bottle           150 mg         OTC         24 ct bottle           150 mg         OTC         220 ct bottle           150 mg         OTC         40 ct bottle           150 mg         OTC         24 ct bottle           150 mg         OTC         All counts

If consumers have questions regarding this recall or to report an adverse event, please contact the Company's Medical Information Call Center at 1-888-375-3784 (1-888-DRL-DRUG) between the hours of 8 a.m. to 8 p.m. ET, Monday through Friday. Patients should contact their healthcare provider if they have experienced any problems that may be related to taking or using this drug product. Adverse reactions or quality concerns experienced with the use of this product may also be reported to the FDA's MedWatch Adverse Event Reporting program either online, by regular mail or by fax. Complete and submit the report online: <a href="www.fda.gov/medwatch/report.htm">www.fda.gov/medwatch/report.htm</a>. Regular Mail or Fax: Download form <a href="www.fda.gov/MedWatch/getforms.htm">www.fda.gov/MedWatch/getforms.htm</a> or call 1-800-332-1088 to request a reporting form, then complete and return to the address on the pre-addressed form, or submit by fax to 1-800-FDA-0178.

This recall is being executed with the knowledge of the U.S. Food and Drug Administration.

RDY-CORP-0919

About Dr. Reddy's: Dr. Reddy's Laboratories Ltd. (BSE: 500124, NSE: DRREDDY, NYSE: RDY) is an integrated pharmaceutical company, committed to providing affordable and innovative medicines for healthier lives. Through its three businesses - Pharmaceutical Services & Active Ingredients, Global Generics and Proprietary Products – Dr. Reddy's offers a portfolio of products and services including APIs, custom pharmaceutical services, generics, biosimilars and differentiated formulations. Our major therapeutic areas of focus are gastrointestinal, cardiovascular, diabetology, oncology, pain management and dermatology. Dr. Reddy's operates in markets across the globe. Our major markets include – USA, India, Russia & CIS countries, and Europe. For more information, log on to: <a href="https://www.drreddys.com">www.drreddys.com</a>

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