

## **Important Correction of Drug Information**

May 23, 2018

## SUBJECT: Notice to Healthcare Professionals of Error in Labeling of the Folic Acid Content in Baxter's Premier ProRx INFUVITE® ADULT Multiple Vitamins Injection

Dear Healthcare Professional:

The purpose of this letter is to inform you of an labeling error of the folic acid content on the label on Vial # 2 of Baxter's Premier ProRx INFUVITE® ADULT Multiple Vitamins Injection.

## Summary of Folic Acid Labeling Error on the 5 mL Vial #2 label

The strength of Folic Acid is erroneously stated as 600 mg on the Vial #2 label. The correct strength is 600 micrograms, as stated on the package insert and the carton.

The product has the appropriate quantity of Folic Acid and is used as a single dose with no calculation when preparing an infusion for patient administration. The product itself is considered safe for the patient as its quality and efficacy is not affected.

Baxter's Infuvite Adult Multiple Vitamins Injection is manufactured by Sandoz Canada Inc., and distributed by Baxter. The 5 mL, 10 pack contains five (5) of Vial #1 and five (5) of Vial #2. One of each vial is required for a single dose.

Sandoz Inc. has evaluated all the approved INFUVITE® ADULT Multiple Vitamins Injection labels and has determined that this error was only on the Premier ProRx product Vial #2 label.

The following lots of ProRX INFUVITE Adult Single Dose 10 pack (5) vials are impacted by this error.

Material Description	Lot	Expiration Date	NDC Code
ProRX INFUVITE Adult Single Dose 10LIVI US	GY5919	08. 2018	54643-7862-9
ProRX INFUVITE Adult Single Dose 10LIVI US	GZ9290	08. 2018	54643-7862-9
ProRX INFUVITE Adult Single Dose 10LIVI US	HG7655	12. 2018	54643-7862-9

If you have any questions or comments on the information provided in this letter, please contact: Sandoz Quality Compliance Call Center at 1-800-525-2492, Email: <a href="mailto:qa.drugsafety@sandoz.com">qa.drugsafety@sandoz.com</a>.

Adverse events or quality problems experienced with the use of this product may also be reported to the FDA's MedWatch Adverse Event Reporting Program either online, or regular mail, or by fax:

• Complete and submit the report **Online**: www.fda.gov/medwatch/report.htm

## Sandoz Inc.

100 College Road West Princeton, NJ 08540 USA www.us.sandoz.com



• 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 preaddressed form, or submit by fax to 1-800-FDA-0178.

Thank you again for your support of INFUVITE® ADULT Multiple Vitamins Injection.

Sincerely, **SANDOZ INC.** 

Anthony Maffia III, Vice President, Regulatory Affairs