#### Teva Pharmaceuticals USA, Inc.

## URGENT DRUG RECALL – Retail LEVEL - INITIATED 2/5/16 PARICALCITOL Capsules, 1 mcg

**RECALLED BY:** 

Teva Pharmaceuticals USA, Inc.

Horsham, PA 19044

Lot#	Exp. Date 2/2016	Strength 1 mcg	Bottle Size 30 count	NDC 0093-7656-56
13013.009A				

#### Dear Customer:

Teva Pharmaceuticals USA, Inc. is voluntarily recalling the above lot of **PARICALCITOL Capsules**, **1 mcg** distributed under the **Teva Pharmaceuticals label**. This recall is being carried out to the **Retail LEVEL** due to out of specification test results for impurities during stability testing. The use of or exposure to the product may cause temporary or medically reversible adverse health consequences, and the probability of serious adverse health consequences is likely remote.

Wholesalers / Distributors / Retailers - Please perform the following activities:

- Examine your inventory immediately for the specified lot of PARICALCITOL Capsules, 1 mcg
- Our records indicate we shipped this product between May 21, 2014 and November 3, 2014.
- Immediately discontinue distribution of the specific lot being recalled.
- Please perform a SUB-RECALL to your retail accounts using this Recall Notification and Stock Response Form.
- Promptly complete the attached recall stock response and reply via fax number 817-868-5362 or mail, even if you have **no** product to return.

Completed Recall Stock Response forms can be mailed, emailed, or sent via FAX to Inmar Attn: Recall Coordinator, 4332 Empire Road Suite 200, Fort Worth, TX 76155. Inmar Email address: <a href="mailto:recallnotice@inmar.com">recallnotice@inmar.com</a>. Inmar FAX: 817-868-5362. Inmar will send a <a href="mailto:Return Goods Authorization">Return Goods Authorization</a> label and shipping label. Appropriate credit for product returns, plus handling and shipping expenses, will be issued upon receipt of said product with the Return Goods Authorization form. All recalled product returned without a Return Goods Authorization label may delay the issuance of your credit.

This recall is being made with the knowledge of the Food & Drug Administration. Your cooperation and prompt response to this notice is appreciated. If you have Customer Service related questions, please contact Teva Customer Service at 800-545-8800. For medical-related questions please contact Medical Information at 888-838-2872, option 3, then, option 4. For product complaint-related questions please contact Quality Assurance Services at 888-838-2872, option 3, then, option 3. If you need a Recall Stock Response form, contact Inmar at 800-967-5952 or acquire it from clsnetlink.com.

Sincerely,

James S. Young

Director, Quality Systems & Regulatory Compliance

Teva Pharmaceuticals USA, Inc.

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### **STOCK RESPONSE FORM**

Please fill out compl	<u>etely</u>			Date	
DIRECT CUSTOM	ERS ONLY: [	Ooes this respons	se include <u>all</u> DC loca	tions? TYES [	□NO
Customer/Store Name	e:			DEA	#:
DEA # is required; i	f not provided	the processing o	f your form will be de	layed	
Address:					
					Zip:
Contact Name (please print):			Telepho		
Lot #	Exp. Date	Strength	Bottle Size	NDC	Quantity to Return (count partial as 1)
13013.009A	2/2016	1 mcg	30 count	0093-7656-56	
	TOMERS ON	LY: Please con	aplete the following:	DEA #:	
City:			State:		
	<u>Inqu</u>	iries regarding	this recall are to be	directed to the follow	ving:
Recall Sto	-	•	turn kit is not received en Option 3. Please <u>d</u>		ss days contact Inmar at onse form.
	lical-related que	estions - contact	stions - contact Teva C Medical Information tact Quality Assurance	888-838-2872, option	
•	•	-	17-868-5362 or E-ma		•
Inmar/MedTurn Use C	only:				
Scan	Labels		Store	Kit	D.B