



July 2016

### URGENT PRODUCT NOTIFICATION

Product Name	Catalog No.	Lot No.	Expiration Date
BD Tritest™ CD3 FITC/CD4 PE/CD45 PerCP (IVD)	340383	49983	31 July 2016
		01810	30 September 2016
		82527	30 September 2016
		54689	30 November 2016
		36715	31 December 2016
		15614	31 January 2017
		57797	28 February 2017

Dear Customer,

BD has recently confirmed that the vial labels for the above listed lot numbers of this product state that the product contains gelatin. The product does not contain gelatin, and the label should state "in buffer." See Figure 1.

Figure 1: Vial Label



The product may continue to be used. Product performance is not impacted, since the buffer contains a stabilizer equivalent to gelatin. No action is required.

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Our records indicate you may have been shipped one of the above referenced lots. Per the label, the product may have been ordered using Catalog No. 340402 for the reagents listed above with BD Trucount™ tubes.

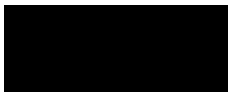
**YOU NEED TO TAKE THE FOLLOWING ACTIONS:**

1. Please fill in the attached form, sign and fax the completed form to your local distributor contact confirming that you have read and understood this notice.
2. We also request that you forward this notice to any other person or groups within your organization for whom this information may be relevant.

For all other inquiries please contact your local BD representative and they will ensure that you are put in contact with the most appropriate individual to address your concerns.

Please accept our apology for any inconvenience this may cause. BD is committed to providing you with the highest quality products. Thank you for your continued support.

Sincerely,



Rose Kwong  
Senior Regulatory Affairs & Compliance Executive

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### URGENT PRODUCT NOTIFICATION (CUSTOMER ACKNOWLEDGEMENT)

Please fill in the information below so that we may acknowledge your receipt of this notification. Simply complete and return the completed form to your local distributor contact.

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BD Tritest™ CD3 FITC/CD4 PE/CD45 PerCP (IVD)	340383	49983	31 July 2016
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Please tick as appropriate.

I confirmed that I have read and understood the Product Notification dated July 2016, regarding the above BD products.

Other comments:

\_\_\_\_\_  
\_\_\_\_\_

**Completed by:**

Name: \_\_\_\_\_ Signature/Date: \_\_\_\_\_

Facility: \_\_\_\_\_

Please use full, current facility name. Do not use initials.

Street Address: \_\_\_\_\_

Telephone No.: \_\_\_\_\_ Fax No.: \_\_\_\_\_

Company stamp:

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