



URGENT MEDICAL DEVICE RECALL

Access CEA Reagent Kit

For use with the Access Family of Immunoassay Systems*

REF	LOT	
33200	595027 595029	15-FEB-16 28-FEB-16

^{*} The Access Family of Immunoassay Systems includes the Access 2, UniCel DxI 800 and UniCel DxI 600, UniCel DxC 600i, and the UniCel DxC 880i, UniCel DxC 860i, UniCel DxC 680i, and UniCel DxC 660i systems.

Attention Beckman Coulter Customer:

Beckman Coulter is initiating a field action for the product listed above. This letter contains important information that needs your immediate attention.

ISSUE:	Beckman Coulter has determined that the Access CEA reagent packs (P/N 33200) of the lots listed above were filled incorrectly. These packs contain insufficient quantity of reagents in one of the pack wells.
IMPACT:	The impact is dependent upon the instrument and software version installed at the time the reagent lot was in use:
	Access 2 systems running software version 3.3.1 or lower, and Access 2i systems running software version 6.1 or lower with these reagent lots:
	 The instrument may have generated incorrect results of 0.0 ng/mL when using an affected pack.
	 All other results greater than 0.0 ng/mL are not affected by this issue and are correct.
	Access 2 systems running software version 3.4.2 and Access 2i systems running software version 6.2.2 or higher with these lots:
	 The affected packs would have been detected by process monitoring with a result flag QSD (indicates reagent dispense is insufficient), and the pack would have been disabled by the instrument.
	 No patient result would have been generated.
	UniCel DxI systems running any software version:
	 The packs would have been detected by reagent pack monitoring with a result flag QSD, and the pack would have been disabled by the instrument.
	 No patient result would have been generated.

Telephone: (800) 854-3633

www.beckmancoulter.com

Internet:



ACTION:	To all customers that received the affected lots listed above: • Discard the Access CEA reagent pack lots listed above. To Access 2 customers with software version 3.3.1 or lower and Access 2i custome with software version 6.1 or lower ONLY:	
	Review your patient results that were reported as 0.0 ng/mL and did not match the clinical status of the patients.	
	• At the discretion of your Laboratory Director, notify clinicians that it is possible the Access CEA sample results reported by your laboratory were affected by this issue.	
RESOLUTION:	The affected Access CEA reagent pack lots are no longer being distributed.	

Please share this information with your laboratory staff and retain this notification as part of your laboratory Quality System documentation. If you have forwarded any of the affected products listed above to another laboratory, please provide them a copy of this letter.

Please complete and return the enclosed Response Form within 10 days so we are assured you have received this important communication.

If you have any questions regarding this notice, please contact our Customer Technical Support:

- From our website: http://www.beckmancoulter.com
- By phone: call 1-800-854-3633 in the United States and Canada.
- Outside the United States and Canada, contact your local Beckman Coulter representative.

To request replacement material:

- In the United States or Canada, complete the attached Replacement Order Form. Fax the completed form to Beckman Coulter at 1-800-232-3828, or email it to AskBeckman@Beckman.com.
- To request replacement material in other geographies, please contact your local Beckman Coulter representative.

We apologize for the inconvenience that this caused your laboratory.

Sincerely,

Noreen Galvin, Ph.D. Vice President Quality & Regulatory Affairs

Enclosure: Response Form

Replacement Order Form

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