

Urgent Field Safety Notice (Removal) Cordis Vista Brite Tip® & ADROIT™ Guiding Catheters

February 07, 2019

Dear Valued Customer,

The purpose of this communication is to inform you Cordis is recalling 173 lots of Cordis Vista Brite Tip® and ADROIT™ Guiding Catheter product.

Recall Overview:

Cordis has initiated a recall for 173 lots of Vista Brite Tip® and ADROIT™
Guiding Catheter due to frayed pieces of the mounting card being inside the primary packaging. See below image.



Frayed pieces of the mounting card were found in the sterile primary package. While the material may be identified when the product is opened and prepared for use, there is a potential for harm if the material is not discovered or if the device is not prepared properly. Prior to and during preparation of the device, if the frayed material is identified, the user would be prompted to exchange the device for another one, resulting in a preprocedural or intra-procedural delay. However, if undiscovered, the frayed material could enter the patient's vasculature potentially resulting in ischemia, necrosis, or the need for additional intervention. The sterility of the product is not affected. Cordis has not received any reports of patient harm related to this issue.

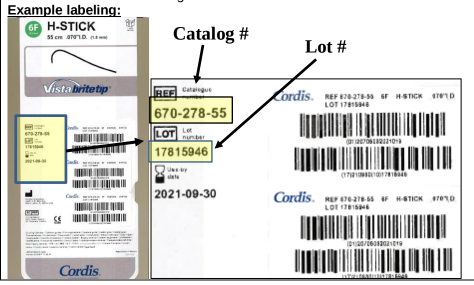
Usage

The Vista Brite Tip® and ADROIT™ Guiding Catheter is intended for use for intravascular introduction of interventional/diagnostic devices into the coronary or peripheral vascular systems.

Details on Affected Devices, to assist in identification of the product involved:

Identification

See Table 1 below for catalog number and lot number list



Why you are being contacted:

You are receiving this letter because our records indicate that you have purchased impacted Vista Brite Tip® and ADROITTM Guiding Catheter lot numbers.

Actions requested on your part:

- 1) Read this Field Safety Notice (Removal) letter.
- 2) Immediately check your inventory to confirm whether or not you have any units from the affected lots in your possession. Identify and set aside any units from the affected lots in a manner that ensures the affected product will not be used. Check all storage and usage locations.
- 3) Review, complete, sign and return the enclosed Acknowledgement Form in accordance with the directions on the form.
- 4) Return all affected product to the Cardinal Health distribution center. Please contact your local sales representative to facilitate return of the affected product, if necessary. Your sales representative will inform you of the product replacement or credit options.
- 5) Share this letter with others in your facility who need to be made aware of this recall and with any other facility that may have been sent the affected units of Vista Brite Tip® and ADROITTM guiding catheter product from your facility. If any units of the affected lots are found to be at the other facility, please arrange the return of the units.
- 6) Maintain awareness of this notice until all affected product has been returned to Cordis and keep a copy of this notice with the affected product.

Description of the problem:

What is the issue?

Cordis became aware of frayed pieces of the mounting card being inside the device primary packaging.

Why are we recalling this product?

Frayed pieces of the mounting card were found in the sterile primary package. While the material may be identified when the product is opened and prepared for use, there is a potential for harm if the material is not discovered or if the device is not prepared properly. Prior to and during preparation of the device, if the frayed material is identified, the user would be prompted to exchange the device for another one, resulting in a pre-procedural or intra-procedural delay. However, if undiscovered, the frayed material could enter the patient's vasculature potentially resulting in ischemia, necrosis, or the need for additional intervention. The sterility of the product is not affected.

What other actions is Cordis taking?

Cordis is investigating the root cause and will take appropriate corrective action. Cordis has not identified any other lots that may be affected.

Available Assistance:

If you have any questions regarding this recall, please contact your local sales representative or local sales office.

Additional Information:

Regulatory Notification

The applicable regulatory agencies and notified body are being notified that Cordis is voluntarily taking this action.

We apologize for any inconvenience this communication may cause. We know that you place high value in our products, and we appreciate your cooperation in this matter. Cordis is committed to maintaining your confidence in the safety and quality of the products that Cordis supplies.

Respectfully yours,

Miguel Ávila

Vice President, Global Quality and Regulatory Affairs

Cordis Corporation

Table 1

Catalog #	Lot Number	Catalog Number	Lot Number
67019055	17744955	588840P	17756076
	17750783		17816499
	17757121		
67021055	17735533	598943P	17746147
0.02.000	17735534		17807995
	17744429		17811028
	17744430		17814511
	17750785		17818155
	17757116		17010133
67021255	17748743	588843P	17777712
07021233	17750476	3000431	17816501
	17755096		17010301
	17757122		
67021455		E0004ED	17720200
	17753002	588845P	17729208
67027055	17779364	588841P	17753948
67027855	17815946	588857P	17756074
		_	17816500
67028055	17805148	588858P	17756075
	17815949		
77821055	17746022	598945P	17755015
	17762925		
77821255	17733016	G780GOND	17724819
	17747583		17742581
	17749800		17756975
	17752698		17771328
	17767925		17794750
	17769354		17814264
	17771053		17816167
	17772929		
77822455	17722360	SM7673	17750494
	17735189		17817649
77827055	17733019	588846P	17733005
	17746026		17736643
	17747584		17737608
	17749793		17742559
	17752703		17740605
	17754875		17743099
	17756762		17743101
	17762928		17745965
	17771058		177-0000
	17771038		
	17816029		
	17817637		
77027055		77828055	1772//00
77827855	17722359	11020000	17724480
	17740611		17767923
	17778655		17771049
			17772926

Table 1 (continued)

Catalog #	Lot Number		
55805400		17806219	
00000100		17809486	
67005400	17811834	17811837 17811839 17813158 17813159 17815025 17815002 17817001 17817002 17817004 17817285 17818563 17818564 17819221 17819222 17821534 17800613 17800614 17800615 17802172 17803537 17803538 17804026 17804222 17804223 17804224 17804765 17804765 17804765 17804766 17804765 17804765 17804765 17805900 17805902 17807627 17811835 17815021 17815022 17815023 17815024 17817003	
67205400	17811836 17819220 17806706 17807595		
	17811677		
	17812321		
	17814278 17814279 17814640 17814641 17814642		
1	17814643		
	1781		