



Medical Devices Safety Notice

The National Health Regulatory Authority would like to alert all governmental and private healthcare facilities, local agents and distributors that the below medical device:

Device Details	
Device Name	Original Perfusor® Line
affected devices	8723060
Lot No.	22D16E8SC6
	22C11E8SC6
Manufacturer	B. Braun Melsungen AG
Country of Origin	Germany
Reference	https://ncmdr.sfda.gov.sa/FileDownLoad.ashx?f=ca&fid=11908
	NHRA initiates this FSN due to risk for holes and leakages in the device. The deviation might
Reason of Recall	harbour the risk for the patient of microbial contamination, under supply, open
	patient access, or air infusion.
Action should be taken	Please stop using the above mentioned medical device and contact the authorized
	representative Wael Pharmacy Co. W.L. L at vincent@waelpharmacy.com to take the
	necessary action for recall.

Your cooperation is highly appreciated in improving health services in the Kingdom of Bahrain.

Recall 2023 0003 25/Jan/2023