

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Item	Manufacturer	link
AquaFlexFlow UF 500 Plus extracorporeal blood circuits	Nuwellis	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-extracorporeal-blood-circuit-issue-nuwellis?utm_medium=email&utm_source=govdelivery
Fluid Delivery Sets	Medline	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-fluid-delivery-set-issue-medline?utm_medium=email&utm_source=govdelivery
Figulla Flex II ASD	Occlutech	https://ansm.sante.fr/informations-de-securite/dm-docclusion-intra-cardiaque-figulla-flex-ii-asd-occluder-occlutech-gmbh
Solution Sets	Baxter Healthcare Corporation	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-solution-set-issue-baxter-healthcare-corporation?utm_medium=email&utm_source=govdelivery
- DLP Pediatric One-Piece Arterial Cannulae - EOPA Arterial Cannula - Select Series Angled Tip Arterial Cannula	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/234
AXIOS Stent and Electrocautery Enhanced Delivery System	Boston Scientific Corp..	https://ade.sfda.gov.sa/Fsca/PublishDetails/235
Dialysis machines.	Fresenius Medical Care.	https://ade.sfda.gov.sa/Fsca/PublishDetails/228
Eclipse Mini Eclipse PRO	Spacelabs Healthcare Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/232
Oxylog 3000 plus	Draeger Medical Systems Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/226

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

PrisMax Systems	Baxter Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/233
draft guidance: Pulse Oximeters for Medical Purposes	FDA	https://www.fda.gov/news-events/press-announcements/fda-proposes-updated-recommendations-help-improve-performance-pulse-oximeters-across-skin-tones?utm_medium=email&utm_source=govdelivery
Medrad Centargo CT Injection System	Imaxeon Pty Ltd	https://ade.sfda.gov.sa/Fsca/PublishDetails/238
draft guidance: Pulse Oximeters for Medical Purposes	U.S. Food and Drug Administration	https://www.fda.gov/news-events/press-announcements/fda-proposes-updated-recommendations-help-improve-performance-pulse-oximeters-across-skin-tones?utm_medium=email&utm_source=govdelivery
VasoView HemoPro 2 (VH-4000 and VH-4001) Endoscopic Vessel Harvesting Systems	Geringe	https://www.fda.gov/medical-devices/medical-device-recalls/endoscopic-vessel-harvesting-evh-system-correction-getinge-and-maquet-cardiovascular-update-use?utm_medium=email&utm_source=govdelivery
	U.S. Food and Drug Administration (FDA)	https://www.fda.gov/medical-devices/medical-devices-news-and-events/medical-device-supply-chain-vulnerabilities-and-public-health-impact-they-have-our-most-vulnerable?utm_medium=email&utm_source=govdelivery
	U.S. Food and Drug Administration (FDA)	https://www.fda.gov/medical-devices/letters-health-care-providers/update-evaluation-airborne-chemicals-neonatal-incubators-letter-health-care-providers?utm_medium=email&utm_source=govdelivery
Ivenix Infusion System	Fresenius Kabi USA	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-infusion-pump-software-issue-fresenius-kabi-usa?utm_medium=email&utm_source=govdelivery
3M™ Prevena™ Plus 125 Therapy Units and Kits and 3M™ V.A.C. ® Via Therapy Units and Kits distributed	KCI USA Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/249

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Disposable insufflation needle	LANDANGER SA	https://ade.sfda.gov.sa/Fsca/PublishDetails/244
Multiple products	Coloplast	https://ade.sfda.gov.sa/Fsca/PublishDetails/245
ROTEM® sigma complete. ROTEM® sigma complete + hep.	Werfen..	https://ade.sfda.gov.sa/Fsca/PublishDetails/250
VINYL EXAMINATION GLOVES	Anhui Intco Medical Products Co., Ltd	https://ade.sfda.gov.sa/Fsca/PublishDetails/246
AD7 and AD7X patient tables which are part of Philips Allura and Azurion systems	Philips Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/265
Cardinal Health Presource Kits:	Cardinal Health 200, LLC..	https://ade.sfda.gov.sa/Fsca/PublishDetails/261
Medima Infusion Pump	ICU Medical, Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/256
Medima P100, P200, and P300 Volumetric infusion Pumps	ICU Medical, Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/255
Multiva 1.5T	Philips Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/262
Optipac Bone Cement	Zimmer Biomet	https://ade.sfda.gov.sa/Fsca/PublishDetails/254

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Philips Allura and Azurion systems	Philips Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/263
Software update for the intensive care ventilator elisa 300/500/600/800/800VIT	Lowenstein Medical GmbH &Co.KG	https://ade.sfda.gov.sa/Fsca/PublishDetails/266
Zenition 50 and Zenition 70 systems	Philips Medical Systems	https://ade.sfda.gov.sa/Fsca/PublishDetails/123
StatStrip Glucose and Glucose/Ketone	Nova Biomedica	https://www.fda.gov/medical-devices/medical-device-recalls/glucose-and-glucoseketone-meter-correction-nova-biomedical-corporation-issues-software-correction?utm_medium=email&utm_source=govdelivery
Neo-Tee T-Piece Resuscitator	Mercury Medical	https://www.fda.gov/medical-devices/medical-device-recalls/gas-powered-emergency-resuscitator-recall-mercury-medical-removes-neo-tee-t-piece-resuscitator-due?utm_medium=email&utm_source=govdelivery
Alinity s System	Abbott GmbH. and Co. KG	https://ade.sfda.gov.sa/Fsca/PublishDetails/269
CARTO® OCTARAY™ Mapping Catheter with TRUEref™ Technology	Biosense Webster Inc. Jo	https://ade.sfda.gov.sa/Fsca/PublishDetails/272
GLUCOSE liquicolor	HUMAN Gesellschaft für Biochemica und Diagnostica	https://ade.sfda.gov.sa/Fsca/PublishDetails/267
Heater-Cooler Unit HCU 40	MAQUET Cardiopulmonary GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/271
Becker and Exacta EDMS	Medtronic Neurosurgery	https://www.fda.gov/medical-devices/medical-device-recalls/pressure-monitoring-device-recall-medtronic-neurosurgery-issues-correction-becker-and-exacta?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Life2000 Ventilator System	Baxter Healthcare	https://www.fda.gov/medical-devices/medical-device-recalls/continuous-ventilator-correction-baxter-healthcare-corporation-issues-correction-life2000-ventilator?utm_medium=email&utm_source=govdelivery
oxygen concentrators.	JIANGSU JUMAO X-CARE MEDICAL EQUIPMENT CO LTD	https://www.fda.gov/medical-devices/medical-device-recalls/oxygen-concentrator-recall-jiangsu-jumao-x-care-medical-equipment-co-ltd-removes-jmc5a-nitruaire-5?utm_medium=email&utm_source=govdelivery
diabetes devices	FDA Alert	https://www.fda.gov/medical-devices/safety-communications/fda-alerts-patients-regularly-check-diabetes-related-smartphone-device-alert-settings-especially?utm_medium=email&utm_source=govdelivery
Rotarex Atherectomy Systems	Vascular, a subsidiary of Becton, Dickinson and Company (BD)	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-atherectomy-catheter-system-issue-bard-peripheral-vascular?utm_medium=email&utm_source=govdelivery
Integrated Arterial Catheters	Medline Industries, LP	https://www.fda.gov/medical-devices/medical-device-recalls/arterial-catheter-recall-medline-industries-lp-removes-integrated-arterial-catheters-due-excess?utm_medium=email&utm_source=govdelivery
Phasitron breathing circuit kits	Sentec/Percussionaire	https://www.fda.gov/medical-devices/medical-device-recalls/breathing-circuit-kit-recall-sentecpercussionaire-removes-vdr4-phasitron-breathing-circuits-due?utm_medium=email&utm_source=govdelivery
LivaNova's Model 1000 SenTiva™ and Model 1000-D SenTiva Duo™ VNS Therapy™ generators	LivaNova PLC	https://ade.sfda.gov.sa/Fsca/PublishDetails/277
TBS iNsight	medimaps group SA	https://ade.sfda.gov.sa/Fsca/PublishDetails/279
Impella RP	Abiomed Inc	https://www.fda.gov/medical-devices/medical-device-recalls/heart-pump-recall-abiomed-inc-updates-use-instructions-impella-rp-smartassist-and-impella-rp-flex?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

AliniQ AMS	Abbott	https://ade.sfda.gov.sa/Fsca/PublishDetails/290
Aortic Root Cannula	Medtronic SA	https://ade.sfda.gov.sa/Fsca/PublishDetails/285
Haemostatic Forceps	Aesculap	https://ade.sfda.gov.sa/Fsca/PublishDetails/287
MiniMed™ Paradigm™, MiniMed™ 600 series, and MiniMed™ 700 series insulin pump systems	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/286
Olympus MAJ-891 Forceps/Irrigation Plug (Isolated Type)	Olympus Corporation of the Americas .	https://ade.sfda.gov.sa/Fsca/PublishDetails/288
SpeedControl Dial component (not implant)	Max Mobility/Permobil	https://www.fda.gov/medical-devices/medical-device-recalls/power-assist-device-recall-max-mobilitypermobil-removes-speedcontrol-dial-component-used-smartdrive?utm_medium=email&utm_source=govdelivery
Allura and Azurion interventional fluoroscopy systems (not implant)	Phillps	https://www.fda.gov/medical-devices/medical-device-recalls/patient-table-correction-philips-updates-use-instructions-allura-and-azurion-systems-due-patient?utm_medium=email&utm_source=govdelivery
Varipulse ablation catheter	Biosense Webster	https://www.fda.gov/medical-devices/medical-device-recalls/ablation-catheter-correction-biosense-webster-updates-use-instructions-varipulse-due-high-rate?utm_medium=email&utm_source=govdelivery
Regard newborn kits (not implant)	ROi CPS, LLC	https://www.fda.gov/medical-devices/medical-device-recalls/regard-newborn-kit-recall-roi-cps-llc-removes-certain-newborn-kits-due-recalled-component-neo-tee-t?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Single Use Guide Sheath Kits (not implant)	Olympus	https://www.fda.gov/medical-devices/medical-device-recalls/endoscope-instrument-recall-olympus-removes-single-use-guide-sheath-kits-due-risk-radiopaque-guide?utm_medium=email&utm_source=govdelivery
MEDRAD Intego 200 PET Infusion Systems (not implant)	Bayer Healthcare LLC..	https://ade.sfda.gov.sa/Fsca/PublishDetails/300
CO2 Filter Line	Covidien LLC....	https://ade.sfda.gov.sa/Fsca/PublishDetails/301
Aisys CS2 ...anesthesia devices (not implant)	GE Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/62
ASSURITY™ AND ENDURITY™ PACEMAKERS	Abbott..	https://ade.sfda.gov.sa/Fsca/PublishDetails/302
HAMILTON-C2/C3	HAMILTON MEDICAL AG	https://ade.sfda.gov.sa/Fsca/PublishDetails/303
Shiley™ Adult Flexible Tracheostomy Tube with TaperGuard™ Cuff Reusable Inner Cannula	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/305
Silvercel Hydro Alginate, (not implant)	Advanced Medical Solutions Limited..	https://ade.sfda.gov.sa/Fsca/PublishDetails/295
VARIPULSE™	Biosense Webster, Inc	https://ansm.sante.fr/informations-de-securite/dispositif-dablation-par-electroporation-rythmologie-catheter-varipulse-biosense-webster
fantômes de cols inclinés 8 M.	Keri Medical	https://ansm.sante.fr/informations-de-securite/orthopedie-fantome-de-col-incline-8-m-keri-medical

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

s adaptateurs pour voies respiratoires(not implant)	Smiths Medical	https://ansm.sante.fr/informations-de-securite/ventilateur-accessoire-adaptateurs-pour-voies-respiratoires-bci-smiths-medical
Vaporizer Sevoflurane(not implant)	Getinge	https://www.fda.gov/medical-devices/medical-device-recalls/vaporizer-recall-getinge-removes-vaporizer-sevoflurane-quick-fill-and-expands-recall-vaporizer?utm_medium=email&utm_source=govdelivery
Tack Endovascular Systems	Philips	https://www.fda.gov/medical-devices/medical-device-recalls/endovascular-system-recall-philips-removes-and-discontinues-distribution-tack-endovascular-system?utm_medium=email&utm_source=govdelivery
Canule de trachéotomie flexible Shiley™	Medtronic	https://ansm.sante.fr/informations-de-securite/canule-de-tracheotomie-flexible-shiley-pour-adultes-avec-chemise-interne-reutilisable-a-ballonnet-taperguard-covidien-llc-medtronic
Spectrum infusion pumps (not implant)	Baxter Healthcare Corporation	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-infusion-pump-issue-baxter-healthcare-corporation?utm_medium=email&utm_source=govdelivery
Delivery cable Flex-Pusher II (51FP100) in combination with Delivery system	Occlutech International AB	https://ade.sfda.gov.sa/Fsca/PublishDetails/310
ProPort™ Plastic Implantable Ports	Smiths Medical ASD, Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/306
générateur thermique HCU 40	Maquet Cardiopulmonary GmbH	https://ansm.sante.fr/informations-de-securite/generateur-thermique-de-cec-heater-cooler-unit-hcu-40-maquet-cardiopulmonary-gmbh-getinge
valves anti-retour « Vaccum Relief Check Valve »	Livanova	https://ansm.sante.fr/informations-de-securite/cec-ecmo-consommable-circuit-set-vacuum-relief-check-valve-livanova

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

SynchroMed	Medtronic	https://ansm.sante.fr/informations-de-securite/pompe-implantable-accessoires-telecommande-patient-mypm-medtronic-inc
BiPAP(not implant)	Philips Respironics	https://ansm.sante.fr/informations-de-securite/appareils-de-ventilation-bipap-a30-bipap-a30-efl-bipap-a30-hybrid-bipap-a40-bipap-a40-efl-bipap-a40-pro-philips-respironics
ProPort Plastic	Smiths Medical	https://www.fda.gov/medical-devices/medical-device-recalls/implantable-port-recall-smiths-medical-removes-proport-plastic-implantable-ports-due-manufacturing?utm_medium=email&utm_source=govdelivery
CVAC Aspiration Systems	Calyxo	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-aspiration-system-issue-calyxo?utm_medium=email&utm_source=govdelivery
BD Alaris Systems (not implant)	Becton, Dickinson	https://www.fda.gov/medical-devices/medical-device-recalls/infusion-pump-software-correction-becton-dickinson-and-company-bd-issues-correction-bd-alaris?utm_medium=email&utm_source=govdelivery
Rocket Thoracentesis Catheter 6Fg & 8Fg	Rocket Medical	https://ade.sfda.gov.sa/Fsca/PublishDetails/324
TruSystem 7500 Surgical Table Systems(not implant)	Baxter Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/322
Vasoview Hemopro 2 (w/Vasoshield) Endoscopic Vessel Harvesting System(not implant)	MAQUET Cardiopulmonary GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/313

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

ORAL/NASAL Endotracheal Tubes	Smiths Medical	https://www.fda.gov/medical-devices/medical-device-recalls/endotracheal-tube-recall-smiths-medical-removes-intubation-oralnasal-endotracheal-tubes-due-smaller?utm_medium=email&utm_source=govdelivery
Aortic Root Cannulas	Medtronic	https://www.fda.gov/medical-devices/medical-device-recalls/vascular-cannula-recall-medtronic-removes-aortic-root-cannula-due-unexpected-loose-material-male?utm_medium=email&utm_source=govdelivery
Origin Data Management software versions 3.1.0, 3.1.1, 3.1.2, 3.2.0, 3.2.1	Brainlab AG	https://ade.sfda.gov.sa/Fsca/PublishDetails/329
Centricity High Acuity Critical Care (CHA CC) and Centricity High Acuity Anesthesia (CHA A) systems (collectively CHA)	GE Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/327
Carestation 620/650/650c and 750/750c Anesthesia Systems	GE Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/328
Minicap Extended Life PD Transfer Set with Twist Clamp	Baxter Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/148
MR systems with 60cm wide bore	Philips Medical Systems	https://ade.sfda.gov.sa/Fsca/PublishDetails/337
ZOLL Powerheart® G5 AED Product Family(it is notimplant)	Cardiac Science Corporation	https://ade.sfda.gov.sa/Fsca/PublishDetails/346
4008 S V10 hemodialysis machine(it is notimplant)	Fresenius Medical Care.	https://ade.sfda.gov.sa/Fsca/PublishDetails/341

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

072 Aspiration System "Hippo" (NON IMPLANT)	Q'Apel Medical, Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/343
Philips CT Big Bore(it is notimplant)	Philips Medical Systems	https://ade.sfda.gov.sa/Fsca/PublishDetails/344
Microstream Advance Intubated CO2 Filter Line, VitaLine Intubated CO2 Filter Line and FilterLine Intubated CO2 Filter Line(it is notimplant)	Philips Medical Systems	https://ade.sfda.gov.sa/Fsca/PublishDetails/347
IntelliSpace Cardiovascular(it is notimplant)	Philips Medical Systems Nederland B.V.	https://ade.sfda.gov.sa/Fsca/PublishDetails/348
PowerPICC Intravascular Catheters	Bard Access Systems	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-intravascular-picc-catheter-issue-bd?utm_medium=email&utm_source=govdelivery
Centricity High Acuity Critical Care (CHA CC)(not implant)	GE Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/358
Ascenda™ Intrathecal Catheter	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/354
HeartMate Mobile Power Unit (MPU)	Abbott	https://www.fda.gov/medical-devices/medical-device-recalls/heart-pump-accessory-removal-abbott-removes-heartmate-mobile-power-unit-due-instances-sudden-power?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Flexi Port Reusable Blood Pressure Cuffs and Two-Piece Reusable Blood Pressure Cuffs kits(its not implant)	Welch Allyn, Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/368
ParaPAC plus™ Model 300 and Model 310 Ventilator(not implant)	Smiths Medical ASD, Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/361
HKH 8820 Wall Holder(not implant)	MAQUET Cardiovascular LLC.	https://ade.sfda.gov.sa/Fsca/PublishDetails/371
FREEGO ENTERAL FEEDING PUMP(not implant)	Zevex International Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/39
MR systems with 60cm wide bore(its not implant)	Philips Medical Systems	https://ade.sfda.gov.sa/Fsca/PublishDetails/337
- Tec 6 Plus - Tec 820 ISO - Tec 820 SEV - Tec 850 ISO - Tec 850 SEV(its not implant)	GE Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/205
CADD-Solis and CADD-Solis VIP Ambulatory Infusion Pumps Thermal Damage(not implant)	Smiths Medical..	https://ade.sfda.gov.sa/Fsca/PublishDetails/374
Hugo™ Robotic-Assisted Surgery (not implant)	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/377

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Centricity High Acuity Critical Care (CHA CC) and Centricity High Acuity Anesthesia (CHA A) systems (collectively CHA)(not implant)	GE Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/379
VentStar Flex 220 (+A126:C139 not implant)	Draeger, Inc	https://www.fda.gov/medical-devices/medical-device-recalls/anesthesia-breathing-circuit-kit-correction-draeger-inc-updates-use-instructions-ventstar-flex-and?utm_medium=email&utm_source=govdelivery
Novum IQ Large Volume Pumps (not implant)	Baxter	https://www.fda.gov/medical-devices/medical-device-recalls/infusion-pump-correction-baxter-updates-instructions-use-novum-iq-large-volume-pump-due-potential?utm_medium=email&utm_source=govdelivery
Shiley Adult Flexible Tracheostomy Tube	Medtronic	https://www.fda.gov/medical-devices/medical-device-recalls/flexible-tracheostomy-tube-recall-medtronic-removes-shiley-adult-flexible-tracheostomy-tube?utm_medium=email&utm_source=govdelivery
MedicaLyte Liquid Bicarbonate Concentrate (not implant)	Nipro	https://www.fda.gov/medical-devices/medical-device-recalls/liquid-bicarbonate-concentrate-recall-nipro-removes-medicalyte-liquid-bicarbonate-concentrate-due?utm_medium=email&utm_source=govdelivery
ACURATE neo2™ Aortic Valve Systems	Boston Scientific Corp..	https://ade.sfda.gov.sa/Fsca/PublishDetails/416
Hippo 072 Aspiration Systems (not implant)	Q'Apel Medical	https://www.fda.gov/medical-devices/medical-device-recalls/aspiration-catheter-recall-qapel-medical-inc-removes-hippo-072-aspiration-system-and-cheetah?utm_medium=email&utm_source=govdelivery
Ivenix LVP Blood Products	Fresenius Kabi	https://www.fda.gov/medical-devices/medical-device-recalls/blood-products-administration-set-recall-fresenius-kabi-removes-large-volume-pump-blood-products?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

OmniLab Advanced + (OLA+) (not implant)	Respironics Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/435
Carestations (not implant)	GE HealthCare	https://www.fda.gov/medical-devices/medical-device-recalls/anesthesia-delivery-systems-recall-ge-healthcare-issues-correction-certain-carestations-due-risk?utm_medium=email&utm_source=govdelivery
Aortic Root Cannulas	Medline Industries LP	https://www.fda.gov/medical-devices/medical-device-recalls/medical-procedure-kits-correction-medline-industries-lp-issues-correction-medline-procedure-kits?utm_medium=email&utm_source=govdelivery
AutoPulse NXT Resuscitation System (not implant)	ZOLL Circulation, Inc	https://www.fda.gov/medical-devices/medical-device-recalls/resuscitation-system-recall-zoll-circulation-inc-recalls-autopulse-nxt-resuscitation-system-due?utm_medium=email&utm_source=govdelivery
Bravo CF Capsule Delivery Devices (not implant)	Medtronic	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-esophageal-ph-monitoring-capsule-issue-medtronic?utm_medium=email&utm_source=govdelivery
Cook Beacon Tip 5.0 Fr Angiographic Catheter	Cook Inc	https://www.fda.gov/medical-devices/medical-device-recalls/angiographic-catheter-recall-cook-removes-beacon-tip-angiographic-catheters-due-tip-separation?utm_medium=email&utm_source=govdelivery
Automated Impella Controllers (AIC) (not implant)	Abiomed	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-blood-pump-controller-issue-abiomed?utm_medium=email&utm_source=govdelivery
Spectrum Infusion Pumps	Baxter	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-infusion-pump-software-issue-baxter?utm_medium=email&utm_source=govdelivery
Infant Heated Wire Circuits	Vyaire	https://www.fda.gov/medical-devices/medical-device-recalls/infant-breathing-system-recall-airlifevyaire-removes-infant-heated-wire-circuits-due-risk?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Ballard Closed Suction Systems	Avanos Medical, Inc.	https://www.fda.gov/medical-devices/medical-device-recalls/closed-suction-catheter-recall-avanos-medical-inc-removes-ballard-closed-suction-systems-due-risk?utm_medium=email&utm_source=govdelivery
Invacare Birdie lifter	Invacare France operations SAS	https://ade.sfda.gov.sa/Fsca/PublishDetails/464
Q-Link 13 components and LikoScale Adapter Kits	Hill Rom Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/457
InPen App	Medtronic MiniMed...	https://ade.sfda.gov.sa/Fsca/PublishDetails/459
Heater Unit HU 35 (NON-IMPLANT)	MAQUET Cardiopulmonary GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/462
HeartSine Samaritan PAD (NON-IMPLANT)	HeartSine Technologies Ltd	https://ade.sfda.gov.sa/Fsca/PublishDetails/454
HAMILTON-C6 (NON-IMPLANT)	HAMILTON MEDICAL AG	https://ade.sfda.gov.sa/Fsca/PublishDetails/449
BiPAP A40 Pro Ventilator, BiPAP A40 EFL Ventilator, and BiPAP A30 EFL Ventilator (NON-IMPLANT)	Respironics Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/138
Philips Allura and Azurion systems(NON-IMPLANT)	Philips Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/263

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Alaris Pump Module model 8100 (NON-IMPLANT)	BD and their subsidiary, CareFusion	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-infusion-set-performance-issue-bd?utm_medium=email&utm_source=govdelivery
Codman Disposable Perforator 14 mm and Codman Craniotomy Kits	Integra LifeSciences	https://www.fda.gov/medical-devices/medical-device-recalls/cranial-drill-recall-integra-lifesciences-recalls-codman-disposable-perforators-due-risk-device?utm_medium=email&utm_source=govdelivery
microbore extension sets	B. Braun Medical Inc	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-microbore-extension-set-issue-b-braun-medical-inc?utm_medium=email&utm_source=govdelivery
Dexcom G6, G7, ONE, and ONE+ (NON-IMPLANT)	Dexcom, Inc	https://www.fda.gov/medical-devices/medical-device-recalls/continuous-glucose-monitor-receiver-recall-dexcom-inc-removes-certain-dexcom-g6-g7-one-and-one?utm_medium=email&utm_source=govdelivery
MicroMyst Applicators	Integra LifeSciences	https://www.fda.gov/medical-devices/medical-device-recalls/applicator-recall-integra-lifesciences-removes-micromyst-applicators-due-potential-sterility?utm_medium=email&utm_source=govdelivery
Craniotomy Kits	Medline Industries, LP	https://www.fda.gov/medical-devices/medical-device-recalls/medical-procedure-kits-correction-medline-industries-lp-issues-correction-medline-craniotomy-kits?utm_medium=email&utm_source=govdelivery
Novum IQ Large Volume Pumps (NON-IMPLANT)	Baxter	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-infusion-pump-issue-baxter?utm_medium=email&utm_source=govdelivery
arterial cannulae	Edwards LifeSciences	https://www.fda.gov/medical-devices/medical-device-recalls/arterial-cannula-recall-edwards-lifesciences-removes-arterial-cannula-due-risk-wire-exposure?utm_medium=email&utm_source=govdelivery
Astral 100 and Astral 150 ventilators (NON-IMPLANT)	ResMed	https://www.bfarm.de/SharedDocs/Kundeninfos/EN/12/2024/08339-24_kundeninfo_en.pdf?__blob=publicationFile

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Nellcor™ Bedside SpO2 Patient Monitoring System (NON-IMPLANT)	Covidien LLC....	https://ade.sfda.gov.sa/Fsca/PublishDetails/465
SureTek Burr Hole Cover Kit	Boston Scientific Corp	https://ade.sfda.gov.sa/Fsca/PublishDetails/466
Vercise Genus™ Deep Brain Stimulation (DBS) Implantable Pulse Generators (IPGs)	Boston Scientific Corp..	https://ade.sfda.gov.sa/Fsca/PublishDetails/467
Corin Operating Tables (NON-IMPLANT)	Getinge Lancer	https://ade.sfda.gov.sa/Fsca/PublishDetails/470
RELIANCE ePTFE defibrillation	Boston Scientific Corp..	https://ade.sfda.gov.sa/Fsca/PublishDetails/487
ESG-410 Electrosurgical Generator	Olympus Corporation of the Americas .	https://ade.sfda.gov.sa/Fsca/PublishDetails/482
PrisMax Systems and TherMax Blood Warmer Units	Vantive Health GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/484
Vasoview Hemopro 2 (w/Vasoshield) Endoscopic Vessel Harvesting System	MAQUET Cardiopulmonary GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/313

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Endopath Echelon Vascular White Reload	Ethicon Endo-Surgery, LLC	https://www.fda.gov/medical-devices/medical-device-recalls/disposable-surgical-stapler-cartridge-correction-ethicon-endo-surgery-llc-issues-correction-endopath?utm_medium=email&utm_source=govdelivery
V30, A30, and A40 ventilators	Philips Respironics,	https://www.fda.gov/medical-devices/medical-device-recalls/continuous-ventilator-correction-philips-respironics-updates-use-instructions-bipap-a30-a40-and-v30?utm_medium=email&utm_source=govdelivery
Catheters	Medline ReNewal	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-electrophysiology-catheter-issue-medline-renewal?utm_medium=email&utm_source=govdelivery
Accez Monthly Color Contact Lenses	MI GWANG CONTACT LENS CO., LTD.	https://ade.sfda.gov.sa/Fsca/PublishDetails/490
WATCHMAN TruSeal™ Access System, WATCHMAN FXD Curve™ Access System	Boston Scientific Corp.. B	https://ade.sfda.gov.sa/Fsca/PublishDetails/492
Surdial X Dialysis Machine	Nipro Corporation.	https://ade.sfda.gov.sa/Fsca/PublishDetails/493
SKULL TRACTION TONG LARGE	Aesculap	https://ade.sfda.gov.sa/Fsca/PublishDetails/494
Allura Xper systems	Philips Medical Systems Nederland B.V.	https://ade.sfda.gov.sa/Fsca/PublishDetails/495

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Left Heart Vent Catheters	Medtronic	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-cardiac-cannula-issue-medtronic?utm_medium=email&utm_source=govdelivery
Carotid WALLSTENT Monorail Endoprosthesis	Boston Scientific	https://www.fda.gov/medical-devices/medical-device-recalls/vascular-stent-recall-boston-scientific-removes-carotid-wallstent-monorail-endoprosthesis-due-risk?utm_medium=email&utm_source=govdelivery
Medin-NC3	medin Medical Innovations GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/511
PhD Hemostasis Valve™	Merit Medical Systems Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/514
t:slim X2 Insulin Pump	Tandem Diabetes Care, Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/508
Multiple MRI systems	Siemens Medical Solutions	https://ade.sfda.gov.sa/Fsca/PublishDetails/505
BeneVision N1 Patient Monitor	BioMedical Electronics Co., Ltd...	https://ade.sfda.gov.sa/Fsca/PublishDetails/506
Heartstring III Proximal Seal System	MAQUET Cardiovascular LLC.	https://ade.sfda.gov.sa/Fsca/PublishDetails/515
Ingenia 1.5T CX, Intera 1.5T, Intera 1.5T Power/Pulsar, SmartPath to dStream for 1.5T	Philips Medical Systems	https://ade.sfda.gov.sa/Fsca/PublishDetails/85

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Automated Impella Controllers	Abiomed	https://www.fda.gov/medical-devices/medical-device-safety/early-alert-automated-impella-controller-issue-abiomed?utm_medium=email&utm_source=govdelivery
Plum Duo Infusion System	ICU Medical	https://www.fda.gov/medical-devices/medical-device-recalls/infusion-pump-correction-icu-medical-inc-issues-correction-plum-duo-infusion-system-due-software?utm_medium=email&utm_source=govdelivery
Extended Tip Applicator	Integra LifeSciences	https://www.fda.gov/medical-devices/medical-device-recalls/applicator-recall-integra-lifesciences-removes-extended-tip-applicator-due-potential-sterility-and?utm_medium=email&utm_source=govdelivery
SPUR II resuscitator	Ambu Inc.	https://www.fda.gov/medical-devices/medical-device-recalls/manual-resuscitator-recall-ambu-inc-removes-spur-ii-resuscitators-due-blocked-manometer-port?utm_medium=email&utm_source=govdelivery
Dexcom G6 iOS App	DexCom Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/522
Vida , MAGNETOM Lumina, MAGNETOM Cima.X , MAGNETOM Seara	Siemens Healthcare GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/520
Tentos 4F Stent System	optimed Medizinische Instrumente GmbH..	https://ade.sfda.gov.sa/Fsca/PublishDetails/521
Optical Lens	Jiangsu Qianyuan New Material Technogy Co.,Ltd.	https://ade.sfda.gov.sa/Fsca/PublishDetails/516

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

Disposable electrodes for defibrillation for use with the ZOLL AED PLUS, AED PRO, AED 3, R Series, X Series Automated External Defibrillator models	FIAB SpA	https://ade.sfda.gov.sa/Fsca/PublishDetails/523
Airvo 2 - myAirvo 2	Fisher & Paykel Healthcare Limited	https://ade.sfda.gov.sa/Fsca/PublishDetails/525
ACCOLADE Family of Pacemakers and CRT-Ps	Boston Scientific Corp..	https://ade.sfda.gov.sa/Fsca/PublishDetails/219
Cartiva Synthetic Cartilage Implant	Cartiva Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/526
DreamStation Auto CPAP and Auto BiPAP	Philips Respironics	https://www.fda.gov/medical-devices/medical-device-recalls/ventilator-recall-philips-respironics-removes-certain-dreamstation-devices-due-programming-errors?utm_medium=email&utm_source=govdelivery
Dexcom G7 Continuous Glucose Monitoring	Dexcom	https://www.fda.gov/medical-devices/medical-device-recalls/continuous-glucose-monitor-apps-correction-dexcom-inc-issues-correction-g7-apps-and-one-apps-due?utm_medium=email&utm_source=govdelivery
t:slim X2 Insulin Pumps	Tandem Diabetes Care	https://www.fda.gov/medical-devices/medical-device-recalls/insulin-pump-correction-tandem-diabetes-care-issues-correction-certain-tslim-x2-insulin-pumps-due?utm_medium=email&utm_source=govdelivery
Medtronic DLP Left Heart Vent Catheters	Medline	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-medline-kits-may-contain-recalled-medtronic-cardiac-cannulas?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

ARGUS PB-3000	Schiller AG	https://ade.sfda.gov.sa/Fsca/PublishDetails/538
Charging unit mounted W/T AC TS7500	Baxter Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/536
Atlan anaesthesia workstations	Draeger Medical Systems Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/537
DEFIGARD HD-7	Schiller AG	https://ade.sfda.gov.sa/Fsca/PublishDetails/539
Vista 120/120 S monitor	Draeger Medical Systems Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/535
RENZAN TM M Peripheral Stent System	Terumo Cardiovascular Systems Corp	https://ade.sfda.gov.sa/Fsca/PublishDetails/550
TactiFlex Ablation Catheter, Sensor Enabled	Abbott Medical	https://ade.sfda.gov.sa/Fsca/PublishDetails/547
Heater Unit HU 35	MAQUET Cardiopulmonary GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/549
Bronchofiberscope, Bronchovideoscope	Olympus Corporation of the Americas .	https://ade.sfda.gov.sa/Fsca/PublishDetails/545

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

SOPHIE 2020 ventilator	Fritz Stephan GmbH.	https://ade.sfda.gov.sa/Fsca/PublishDetails/501
MAQUET CARDIOSAVE Hybrid and MAQUET CARDIOSAVE Rescue	Datascope Corp	https://ade.sfda.gov.sa/Fsca/PublishDetails/553
Datascope Cardiosave Hybrid and Cardiosave Rescue IntraAortic Balloon Pumps (IABP)	Getinge	https://ade.sfda.gov.sa/Fsca/PublishDetails/554
Surdial X Dialysis Machine	Nipro Corporation.	https://ade.sfda.gov.sa/Fsca/PublishDetails/493
Automated Impella Controller.	Abiomed	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-automated-impella-controller-purge-retainer-fixation-issue-abiomed?utm_medium=email&utm_source=govdelivery
TactiFlex Ablation Catheter	Abbott	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-tactiflex-ablation-catheter-issue-abbott?utm_medium=email&utm_source=govdelivery
ViziShot 2 FLEX (19G)	Olympus	https://www.fda.gov/medical-devices/medical-device-recalls/endoscopic-aspiration-needle-recall-olympus-removes-certain-vizishot-2-flex-19g-needles-due?utm_medium=email&utm_source=govdelivery
Ranger Blood/Fluid Warming System	3M Company	https://www.fda.gov/medical-devices/medical-device-recalls/blood-and-plasma-warming-device-correction-3m-company-issues-correction-ranger-bloodfluid-warming?utm_medium=email&utm_source=govdelivery
BD Alaris Pump	BD	https://www.fda.gov/medical-devices/medical-device-recalls/early-alert-infusion-set-performance-issue-bd?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

CONSTELLATION® ULTRAVIT® (10K) and HYPERVIT® (20K) probes	Alcon Laboratories Inc...	https://ade.sfda.gov.sa/Fsca/PublishDetails/557
Catalyft™ PL & PL40 expandable interbody system	Medtronic SA	https://ade.sfda.gov.sa/Fsca/PublishDetails/560
LivaNova's Model 1000 SenTiva™ and Model 1000-D SenTiva Duo™ VNS Therapy™ generators	LivaNova PLC	https://ade.sfda.gov.sa/Fsca/PublishDetails/281
Medistim MiraQ Systems	Medistim ASA	https://ade.sfda.gov.sa/Fsca/PublishDetails/569
AVVIGO+ Multi-Modality Guidance System	Boston Scientific Corp	https://ade.sfda.gov.sa/Fsca/PublishDetails/577
FreeGo Enteral Feeding set	Abbott Ireland Diagnostics Division	https://ade.sfda.gov.sa/Fsca/PublishDetails/578
CODMAN® Disposable Perforator 11mm (26-1222) CODMAN® Disposable Perforator 9mm (26-1223)	Integra LifeSciences Production Corporation	https://ade.sfda.gov.sa/Fsca/PublishDetails/570
IntelliSpace Cardiovascular	Philips Medical Systems Nederland B.V.	https://ade.sfda.gov.sa/Fsca/PublishDetails/348
Salem Sump™	Cardinal Health 200, LLC..	https://ade.sfda.gov.sa/Fsca/PublishDetails/573

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

BD Alaris Syringe Pumps	BD Switzerland Sarl	https://ade.sfda.gov.sa/Fsca/PublishDetails/579
V90 electronic vaporize	Shenzhen Mindray BioMedical Electronics Co.,	https://ade.sfda.gov.sa/Fsca/PublishDetails/581
CentriMagTM Blood Pump	Abbott Medical .	https://ade.sfda.gov.sa/Fsca/PublishDetails/584
HeartMate II® and HeartMate 3™ System Controllers	Abbott Medical ..	https://ade.sfda.gov.sa/Fsca/PublishDetails/585
LGN Tibial Offset Bushing 4MM	Smith & Nephew inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/611
Philips IntelliVue Patient Monitors	Philips Healthcare	https://ade.sfda.gov.sa/Fsca/PublishDetails/613
Zenition 10 & Zenition 30	Philips Medical Systems Nederland B.V.	https://ade.sfda.gov.sa/Fsca/PublishDetails/614
KNEO THREADED FASTENING PINS	Groupe Lépine	https://ade.sfda.gov.sa/Fsca/PublishDetails/618
Rocket Seldinger Chest Drain Kits	Rocket Medical	https://ade.sfda.gov.sa/Fsca/PublishDetails/619

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

ACCOLADE Family of Pacemakers and CRT-Ps	Boston Scientific Corp	https://ade.sfda.gov.sa/Fsca/PublishDetails/219
Kendall SCD 700 Sequential Compression System	Cardinal Health 200, LLC..	https://ade.sfda.gov.sa/Fsca/PublishDetails/599
ErgoStar CM 40, ErgoStar CM 45, ErgoStar CM 55, ErgoStar CM 60	Draeger Medical Systems Inc	https://ade.sfda.gov.sa/Fsca/PublishDetails/601
BD Alaris System Pump (8100) & Bezel Spare Parts	CareFusion 303, Inc..	https://ade.sfda.gov.sa/Fsca/PublishDetails/602
CareLink™ Clinic	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/603
Aurora EV-ICD™ (DVEA3E4), Clinical EV-ICD (DVEX3E4)	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/607
Cardiosave Hybrid	Getinge	https://ade.sfda.gov.sa/Fsca/PublishDetails/608
Vanta Implantable Neurostimulator (INS)	Medtronic Inc.	https://ade.sfda.gov.sa/Fsca/PublishDetails/661
Carestation 600 and 700 series	GE HealthCare	https://www.fda.gov/medical-devices/medical-device-recalls-and-early-alerts/early-alert-anesthesia-system-issue-ge-healthcare?utm_medium=email&utm_source=govdelivery

MEDICAL DEVICES
RECALL AND SAFETY NOTICE 2025
Follow up for implants (by projects /programs department MOH)
Follow up for instruments / non-implants (by engineering department MOH)

HP Flex Pro-C PCs used with the Sensis Vibe application	Siemens Healthcare GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/656
MINOP TROCAR	Aesculap...	https://ade.sfda.gov.sa/Fsca/PublishDetails/650
Riptide™ Aspiration Tubing	Micro Therapeutics Inc (DBA ev3 Neurovascular)	https://ade.sfda.gov.sa/Fsca/PublishDetails/653
Corpuls3 Touch	Elektromedizinische Geräte G. Stemple GmbH	https://ade.sfda.gov.sa/Fsca/PublishDetails/654
FreeStyle Libre 3 and FreeStyle Libre 3 Plus	Abbott Diabetes Care	https://www.fda.gov/medical-devices/medical-device-recalls-and-early-alerts/early-alert-glucose-monitor-sensor-issue-abbott-diabetes-care?utm_medium=email&utm_source=govdelivery
Ligating Devices	Olympus	https://www.fda.gov/medical-devices/medical-device-recalls-and-early-alerts/correction-alert-olympus-updates-use-instructions-ligating-device?utm_medium=email&utm_source=govdelivery
OTESUS OR table column	Getinge Disinfection AB	https://ade.sfda.gov.sa/Fsca/PublishDetails/645
Flexor® Check-Flo® Introducers and Sets Rösch-Uchida Transjugular Liver Access Sets	Cook Incorporated	https://ade.sfda.gov.sa/Fsca/PublishDetails/648

MEDICAL DEVICES RECALL AND SAFETY NOTICE 2025 Follow up for implants (by projects /programs department MOH) Follow up for instruments / non-implants (by engineering department MOH)		
UNIMED Procedure Packs	United Medical Industries Co Ltd	https://ade.sfda.gov.sa/Fsca/PublishDetails/647