

U.S. Food and Drug Administration Office of Inspections and Investigations (OII) Office of Medical Devices and Radiological Health Inspectorate (OMDRHI) Division 2–South www.fda.gov

Communication Resources following an FDA Inspection

About Office of Medical Devices and Radiological Health Inspectorate (OMDRHI)

OMDRHI has 4 divisions. Your firm is located within our South Division which covers the states of: AL, AR, FL, GA, LA, MS, NC, OK, PR, SC, TN, TX, and US Virgin Islands. You can learn more about OMDRHI at http://www.fda.gov/OIIDevices.

FDA-483 Responses

Please e-mail your inspection-related correspondence to OII-Devices-South-Firm-Response@fda.hhs.gov

Include your company's name and FEI number found on the form FDA-483 in the subject of the email, and on the cover letter or attachments. We prefer e-mail correspondence for efficiency, fiscal responsibility, and environmental awareness. Be sure that any attachments are readily labeled and/or identified for ease of review and submitted as a single pdf file, with bookmarks to easily identify table of contents, memos, attachments, etc. If a single pdf file exceeds the 100MB size limit, please submit multiple pdf files, with bookmarks, as appropriate. Do not provide multiple folders that contain individual files as this will delay the processing of your response. The agency will acknowledge receipt of your e-mail. Please do not send a back-up hard copy of any correspondence. If hard copy, thumb drive, or compact disc (cd) are the only way you can send a response, please call or email Blake.Bevill@fda.hhs.gov to obtain a mailing address.

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Final Rule: Medical Device Recalls Revised FDA regulation effective in 2026 Submit recall (21 CFR 806) information to On February 2, 2026, the Quality Management OII-Medical-Device-Recalls-SOUTH@fda.hhs.gov System Regulation (OMSR) will go into effect. On For general information on recalls, corrections and the effective date, FDA device inspections will removals, visit: review a manufacturer's compliance with this revised regulation. Links to the final rule and the www.fda.gov/MedicalDevices/DeviceRegulationandG Frequently Asked Questions can be found on uidance/PostmarketRequirements/RecallsCorrection OMDRHO's webpage www.fda.gov/OIIDevices sAndRemovals Additional useful links: ≻ For general information about OMDRHI inspections, including your inspection report, visit What We Do: Inspections at On 02/02/2024 the agency published Quality Management www.fda.gov/OIIDevices System Regulation: Final Rule. Read the rule. For more For general information about **device** \geq information read the Frequently Asked Questions FDA. registration and listing. visit: https://www.fda.gov/medical-devices/how-study-If you have questions about the QMSR, contact the and-market-your-device/device- registration-and-Center for Devices and Radiological Health's listing (CDRH) Division of Industry and Consumer \geq For general information on **mandatory** Education (DICE). reporting requirements, visit: e-mail: DICE@fda.hhs.gov https://www.fda.gov/Mandatory-Reporting-Phone: 1(800) 638-2041 or (301) 796-7100 Requirements-Manufacturers-Importers-andhttps://www.fda.gov/DICE **Device-User-Facilities**