

(2) Any personal, medical, and similar information (including the serial number of implanted devices), which would constitute an invasion of personal privacy under §20.63 of this chapter. FDA will disclose to a patient who requests a report, all the information in the report concerning that patient, as provided in §20.61 of this chapter; and

(3) Any names and other identifying information of a third party voluntarily submitting an adverse event report.

(c) FDA may not disclose the identity of a device user facility which makes a report under this part except in connection with:

(1) An action brought to enforce section 301(q) of the act, including the failure or refusal to furnish material or information required by section 519 of the act;

(2) A communication to a manufacturer of a device which is the subject of a report required by a user facility under §803.30;

(3) A disclosure relating to a manufacturer or distributor adverse event report that is required under section 519(a) of the act; or

(4) A disclosure to employees of the Department of Health and Human Services, to the Department of Justice, or to the duly authorized committees and subcommittees of the Congress.

§803.10 General description of reports required from user facilities and manufacturers.

(a) *Device user facilities.* User facilities must submit the following reports, which are described more fully in subpart C of this part.

(1) User facilities must submit MDR reports of individual adverse events within 10 days after the user facility becomes aware of an MDR reportable event as described in §§803.30 and 803.32.

(i) User facilities must submit reports of device-related deaths to FDA and to the manufacturer, if known.

(ii) User facilities must submit reports of device-related serious injuries to manufacturers, or to FDA, if the manufacturer is unknown.

(2) User facilities must submit semi-annual reports as described in §803.33.

(b) [Reserved]

(c) *Device manufacturers.* Manufacturers must submit the following reports as described more fully in subpart E of this part:

(1) MDR reports of individual adverse events within 30 days after the manufacturer becomes aware of a reportable death, serious injury, or malfunction as described in §§803.50 and 803.52.

(2) MDR reports of individual adverse events within 5 days of:

(i) Becoming aware that a reportable MDR event requires remedial action to prevent an unreasonable risk of substantial harm to the public health or,

(ii) Becoming aware of an MDR reportable event for which FDA has made a written request, as described in §803.53.

(3) Annual baseline reports as described in §803.55.

(4) Supplemental reports if they obtain information that was not provided in an initial report as described in §803.56.

(5) Annual certification to FDA of the number of MDR reports filed during the preceding year as described in §803.57.

§803.11 Obtaining the forms.

User facilities and manufacturers must submit all reports of individual adverse events on FDA Form 3500A (MEDWATCH form) or in an electronic equivalent as approved under §803.14. This form and all other forms referenced in this section can also be obtained from the Consolidated Forms and Publications Office, Washington Commerce Center, 3222 Hubbard Rd., Landover, MD 20785, or from the Division of Small Manufacturers Assistance, Office of Health and Industry Programs, Center for Devices and Radiological Health, 1350 Piccard Dr. (HFZ-220), Rockville, MD 20850, telephone facsimile (FAX) 301-443-8818. FDA Form 3500A may also be obtained from the Food and Drug Administration, MEDWATCH (HF-2), 5600 Fishers Lane, rm. 9-57, Rockville, MD 20850, 301-443-0117.

§803.12 Where to submit reports.

(a) Any written report or additional information required under this part shall be submitted to: Food and Drug Administration, Center for Devices and