§830.20

§830.20 Requirements for a unique device identifier.

- A unique device identifier (UDI) must:
- (a) Be issued under a system operated by FDA or an FDA-accredited issuing agency;
- (b) Conform to each of the following international standards:
- (1) ISO/IEC 15459-2, which is incorporated by reference at §830.10;
- (2) ISO/IEC 15459–4, which is incorporated by reference at $\$\,830.10;$ and
- (3) ISO/IEC 15459-6, which is incorporated by reference at §830.10.
- (c) Use only characters and numbers from the invariant character set of ISO/IEC 646, which is incorporated by reference at §830.10.

[78 FR 58825, Sept. 24, 2013]

§830.40 Use and discontinuation of a device identifier.

- (a) Only one device identifier from any particular system for the issuance of unique device identifiers (UDIs) may be used to identify a particular version or model of a device. A particular version or model may be identified by UDIs from two or more systems for the issuance of UDIs.
- (b) A device identifier shall be used to identify only one version or model.
- (c) In the event that a version or model of a device is discontinued, its device identifier may not be reassigned to another device. If a discontinued version or model is re-introduced and no changes have been made that would require the use of a new device identifier, the device identifier that was previously in use may be used to identify the device.
- (d) In the event that an issuing agency relinquishes or does not renew its accreditation, you may continue to use a previously issued UDI until such time as §830.50 requires you to assign a new device identifier.

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§830.50 Changes that require use of a new device identifier.

(a) Whenever you make a change to a device that is required to bear a unique device identifier (UDI) on its label, and the change results in a new version or

model, you must assign a new device identifier to the new version or model.

(b) Whenever you create a new device package, you must assign a new device identifier to the new device package.

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§830.60 Relabeling of a device that is required to bear a unique device identifier.

If you relabel a device that is required to bear a unique device identifier (UDI), you must:

- (a) Assign a new device identifier to the device, and
- (b) Keep a record showing the relationship of the prior device identifier to your new device identifier.

[78 FR 58825, Sept. 24, 2013]

Subpart C—FDA Accreditation of an Issuing Agency

§ 830.100 FDA accreditation of an issuing agency.

- (a) Eligibility. A private organization may apply for accreditation as an issuing agency.
- (b) Accreditation criteria. FDA may accredit an organization as an issuing agency, if the system it will operate:
- (1) Will employ unique device identifiers (UDIs) that meet the requirements of this part to adequately identify a device through its distribution and use;
- (2) Conforms to each of the following international standards:
- (i) ISO/IEC 15459-2, which is incorporated by reference at §830.10;
- (ii) ISO/IEC 15459-4, which is incorporated by reference at §830.10;
- (iii) ISO/IEC 15459-6, which is incorporated by reference at §830.10.
- (3) Uses only characters and numbers from the invariant character set of ISO/IEC 646, which is incorporated by reference at §830.10.
- (4) Will be available to all users according to a single set of consistent, fair, and reasonable terms and conditions.
- (5) Will protect against conflicts of interest between the issuing agency (and its officers, employees, and other agents) and labelers (and their officers, employees, and other agents) seeking