Field Name brand_name	<b>Type</b> string	Description The Proprietary/Trade/Brand name of the medical device as used in device labeling or in the catalog. This information may 1) be on a label attached to a durable device, 2) be on a package of a disposable device, or 3) appear in labeling materials of an implantable device. The brand name is the name that is typically registered with USPTO and have the <sup>®</sup> and/or TM symbol.
		This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
catalog_number	string	The catalog, reference, or product number found on the device label or accompanying packaging to identify a particular product.
		This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
commercial_distribution_end_date	date	Indicates the date the device is no longer held or offered for sale. See 21 CFR 807.3(b) for exceptions. The device may or may not still be available for purchase in the marketplace.
commercial_distribution_status	string	Indicates whether the device is in commercial distribution as defined under 21 CFR 807.3(b).
		This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
		Value is one of the following
		In Commercial Distribution = In Commercial Distribution
		Not in Commercial Distribution = Not in Commercial Distribution
company_name	string	Company name associated with the labeler DUNS Number entered in the DI Record.
		This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
device count in base package	integer	Number of medical devices in the base package.
device description	string	Additional relevant information about the device that is not already captured as a distinct GUDID data attribute.
has donation id number	boolean	The Donation Identification Number is applicable to devices that are also regulated as HCT/Ps and is a number that is assigned to each
		donation. This number/code is required to be part of the UDI when included on the label in order to provide the means to track the device back to its manufacturing source or otherwise allow the history of the device manufacturing, packaging, labeling, distribution and use to be determined.
		Value is one of the following
		true = true
		false = false
has_expiration_date	boolean	The date by which the label of a device states the device must or should be used. This date is required to be part of the UDI when included on
		the label in order to provide the means to track the device back to its manufacturing source or otherwise allow the history of the device manufacturing, packaging, labeling, distribution and use to be determined.
		Value is one of the following
		true = true
		false = false
has_lot_or_batch_number	boolean	The number assigned to one or more device(s) that consist of a single type, model, class, size, composition, or software version that are
		manufactured under essentially the same conditions and that are intended to have uniform characteristics and quality within specified limits.
		This number is required to be part of the UDI when included on the label in order to provide the means to track the device back to its
		manufacturing source or otherwise allow the history of the device manufacturing, packaging, labeling, distribution and use to be determined.
		Value is one of the following
		true = true
		false = false
has_manufacturing_date	boolean	The date on which a device is manufactured. This date is required to be part of the UDI when included on the label in order to provide the means
		to track the device back to its manufacturing source or otherwise allow the history of the device manufacturing, packaging, labeling, distribution and use to be determined.
		Value is one of the following
		true = true
		false = false

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	has_serial_number	boolean	The number that allows for the identification of a device, indicating its position within a series. This number is required to be part of the UDI
			when included on the label in order to provide the means to track the device back to its manufacturing source or otherwise allow the history of the device manufacturing packaging labeling distribution and use to be determined
			Value is one of the following
			false = false
	is_combination_product	boolean	Indicates that the product is comprised of two or more regulated products that are physically, chemically, or otherwise combined or mixed and produced as a single entity; packaged together as a single package; or packaged separately for the intended use together as defined under 21
			CFR 3.2(e). At least one of the products in the combination product must be a device in this case.
			Value is one of the following
			true = true false = false
	is_direct_marking_exempt	boolean	The device is exempt from Direct Marking requirements under 21 CFR 801.45.
			Value is one of the following
			true = true
		h a sha a s	false = false
	is_hct_p	boolean	Indicates that the product contains or consists of human cells or tissues that are intended for implantation, infusion, or transfer intended for implantation, infusion, or transfer
			Value is one of the following true = true
		h a a l a a a	false = false
	IS_KIL	nearood	including medical devices, that are packaged together to achieve a common intended use and are being distributed as a medical device.
			Value is one of the following
			true = true
	is labeled as no nrl	boolean	false = false
		boolean	contains this information. Only applicable to devices not subject to the requirements under 21 CFR 801.437. Not all medical products that are
			NOT made with natural rubber latex will be marked.
			Value is one of the following
			true = true false = false
	is_labeled_as_nrl	boolean	Indicates that the device or packaging contains natural rubber that contacts humans as described under 21 CFR 801.437. The value true
			indicates that the device label or packaging contains one of the following statements: (1) 'Caution: This Product Contains Natural Rubber Latex Which May Cause Allergic Reactions'. (2) 'This Product Contains Dry Natural Rubber'. (3) 'Caution: The Packaging of This Product Contains
			Natural Rubber Latex Which May Cause Allergic Reactions' or (4) 'The Packaging of This Product Contains Dry Natural Rubber'.
			Value is one of the following
			true = true
	is_otc	boolean	faise = faise Indicates that the device does not require a prescription to use and can be purchased over the counter.
			Value is one of the following
			true = true
	is nm exempt	hooloon	false = false Indicates whether the device is exempt from premarket notification requirements
	ι3_μιτ_ενειτιμι	boolean	
			Value is one of the following
			false = false
	is_rx	boolean	Indicates whether the device requires a prescription.
			Value is one of the following
			true = true
	is_single_use	boolean	Indicates that the device is intended for one use or on a single patient during a single procedure.
			Value is one of the following
			true = true
	labelar duna number	string	false = false
	Tabeler_duns_number	string	by lenders and potential business partners to help predict the reliability and/or financial stability of the company in question.
	mri_safety	string	Indicates the MRI Safety Information, if any, that is present in the device labeling. Please see the ASTM F2503-13 standard for more information.
			This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
			Value is one of the following
			MR Safe = MR Safe
			MR Unsafe = MR Unsafe
			Labeling does not contain MRI Safety Information = Labeling does not contain MRI Safety Information
	public version data	data	Auto accigned the day file is generated with Time Stemp. All evicting records will have first data accigned the day developed file is generated
	public_version_date	uate	with this data element.
	public version number	string	Auto accigned version number, accigned just before file generation. All evisting records will have version 1 accigned
	public_version_number	String	Auto assigned version number, assigned just before me generation, an existing records will have version 1 assigned.
	public_version_status	string	Definition forthcoming.
	publish_date	date	Indicates the date the DI Record gets published and is available via Public Search.
	record key	string	Current ophancoments will allow the Drimany DL to change after the DL record has been released to the public. To ensure records can be linked
	record_key	string	and managed, a record key will be provided; Unique alphanumeric value, auto generated.
	record status	string	Indicates the status of the DI Record.
		0.000	
			Value is one of the following Published = Published
			Unpublished = Unpublished
	sterilization.is sterile	boolean	Deactivated = Deactivated Indicates the medical device is free from viable microorganisms. See ISO/TS 11139.
			Value is one of the following true = true
			false = false
	sterilization.is_sterilization_prior_use	boolean	indicates that the device requires sterilization prior to use.
			Value is one of the following
			rue = rue false = false
	sterilization.sterilization_methods	string	Indicates the method(s) of sterilization that can be used for this device.
			Value is one of the following
			Chlorine Dioxide = Chlorine Dioxide
			שיא Heat Sterilization = שיא Heat Sterilization Ethylene Oxide = Ethylene Oxide
			High Intensity Light or Pulse Light = High Intensity Light or Pulse Light
			High-level Disintectant = High-level Disintectant Hydrogen Peroxide = Hydrogen Peroxide
			Liquid Chemical = Liquid Chemical
			Moist Heat or Steam Sterilization = Moist Heat or Steam Sterilization
			Nitrogen Dioxide = Nitrogen Dioxide
			Peracetic Acid = Peracetic Acid
			Radiation Sterilization = Radiation Sterilization
			Supercritical Carbon Dioxide = Supercritical Carbon Dioxide
	product codes code	ctring	Ultraviolet Light = Ultraviolet Light A three-letter identifier assigned to a device category
		วนเมย	
	product codes name	string	This is an .exact field. It has been indexed both as its exact string content, and also tokenized. Name associated with the three-letter Product Code
	version_or_model_number	string	The version or model found on the device label or accompanying packaging used to identify a category or design of a device. The version or
			model identifies all devices that have specifications, performance, size, and composition within limits set by the labeler.
			This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
Device identifiers	identifiers.id	string	Registration status code.
			Value is one of the following
			<ul> <li> <i>μ</i> = Active <i>μ</i> = Active avaiting assignment of registration number <i>μ</i> = Active <i>μ</i> = Active <i>μ</i> = Active <i>μ</i> = Active <i>μ</i> = Active <i>μ</i> = Active</li></ul>
Device identifiers	identifiers.issuing_agency	string	Identifies whether facility is an initial importer.
			Value is one of the following
			Y = Yes
Device identifiers	identifiers.package_discontinue date	date	N = NO Year that registration expires (expires 12/31 of that year).
Device identifiers	identifiers.package_status	string	Facility or US agent address line 1.
Device identifiers Device identifiers	identifiers.package_type identifiers.quantity_per_package	string integer	Facility or US agent address line 2. Facility or US agent city.
		J	
Device identifiers	identifiers.type	string	Facility or US agent US state or foreign state or province.
Device identifiers	identifiers.unit_of_use_id	string	Number of devices noted in the adverse event report. Almost always 1. May be empty if report_source_code contains Voluntary report.
			This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
Customer Contact	customer_contacts.email	string	Facility or US agent Zip code.
			This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
Customer Contact	customer_contacts.phone	string	Name associated with the facility or US agent.
			This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
Device Size	device_sizes.text	string	Facility foreign postal code.
			This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
	dovice sizes type	string	Facility identifier assigned to facility by the FDA Office of Regulatory Affairs.
Device Size Device Size	device_sizes.value	string	Facility identifier assigned to facility by the FDA Office of Regulatory Affairs.

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2 = Class II (moderate to high risk): general controls and special controls 3 = Class III (high risk): general controls and Premarket Approval (PMA) U = Unclassified N = Not classified F = HDE
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OpenFDA fields device name device name device.
This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
OpenFDA fields regulation Medical Specialty is assigned to facility by the FDA Office of Regulation (e.g. 21 CER Dart 898 is Orthonadia Davisos) which is why Class 2 devices lack
the "Pogulation Medical Specialty" field
the Regulation Medical Specialty field.
This is an .exact field. It has been indexed both as its exact string content, and also tokenized.
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This is an .exact field. It has been indexed both as its exact string content, and also tokenized. OpenFDA fields regulation_number (Code of Federal regulations Title 21, 862.00 through 892.00). The classification regulation covers various aspects of design, clinical evaluation,