

Publicly Accessible Databases for MDSAP Audits

	Main page	Market Authorizations	Site Licenses/ Registrations	Adverse Events	Advisory Notices	Other topics
TGA	TGA	eBusiness	NA	Adverse Event Notifications (DAEN)	Recalls (SARA)	TGA Act & Regulations Standards Orders and Medical Devices Clinical Evidence Guidelines IVD Guidelines Other guidance
ANVISA	ANVISA	Product Registration	International Manufacturers ID	Adverse Events and Quality Issues* <small>*computerized system developed by ANVISA for the news of incidents, adverse events (AE) and technical complaints (QT) related to the use of products and services</small>	Recalls, Counterfeit, Suspended Products	ANVISA Website Guidance

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				under sanitary surveillance		
Health Canada	Health Canada Medical Devices	Medical Device Active License Listing (MDALL)	Medical Device Establishment Licence Listing (MDEL)	Medical Device Incidents	Recalls and Safety Alerts	NA
MHLW/ PMDA	PMDA	NA	Foreign Manufacturing Sites	NA	Recalls	PMD Act MHLW MO169 Standards for Medical Devices in Japan
FDA	FDA	Device Registration and Listing (online search) Establishment Registration & Device Listing	Device Registration and Listing (online search) Establishment Registration & Device Listing	MDR (online search) MAUDE	List of Device Recalls Medical Device Safety (online search) Recalls	Inspection Classification 510(K) Premarket Notification Premarket Approval (PMA) Total Product Life Cycle

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						Product Code Builder Unique Device Identifier CFR Code of Federal Regulations Title 21 eCFR ICH Guidelines Warning Letters