



27. RE-APPLICATION FOR RENEWAL OF MEDICAL DEVICES AND IVD FOR ALL CLASSIFICATIONS

Center/Office/Division:	CDRRHR-LRD
Classification :	Highly Technical
Type of Transaction :	G2B - Government-to-Businesses
Who May Avail :	Medical Device Manufacturers/Distributors (Importer/Exporter/Wholesaler)/Trader
Fees to be Paid :	Php 1,000.00 + 1% LRF = Php1,010.00

CHECKLIST OF REQUIREMENTS	WHERE TO SECURE
Letter of Intent	Applicant
2. Copy of the Notice of Deficiencies	Applicant
3. Compliance Documents	Applicant / Principal/Manufacturer
4. Payment	FDA Cashier
 NOTES: Submit an electronic/scanned copy (in PDF searchable format of at least 150 dpi) The soft copy should be arranged according to the checklist of requirements. The file name should consist of the name of the requirement. The electronic copy should be contained either in one single continuous file per requirement or single continuous file for all requirements. 	
Submission schedule applies to working days only and excludes national and declared non-working days. In the event of a holiday/non-working day, then the regular schedule shall be followed on the next working and scheduled submission day.	





CLIENT STEPS	AGENCY ACTION	FEES TO BE PAID	PROCESSING TIME	PERSON RESPONSIBLE
 Client sends an email containing the PDF of their compliance to <u>fdac.pacd@fda.gov.ph</u> within the prescribed time period stipulated in the notice of deficiency. * 	Receiving officer sends an acknowledgment email to the client.	None	1 working day	FDAC Officer
	2. FDAC forwards the re-application file to CDRRHR.	None	1 working day	FDAC Officer
	CDRRHR receives the re- application file and decks to the evaluator	None	1 working day	CDRRHR Administrative Staff
	Technical evaluation of application. Recommendation of Approval or Final Disapproval	None	10 working days	Technical Evaluator
	5. Quality Assurance - Checking of recommendation of the Supervisor	None	3 working days	LRD Chief
	Drafting and finalization of certificate or disapproval letter	None	1 working day	Technical Evaluator
	7. Final Approval/Disapproval and signature of the Director	None	1 working day	CDRRHR Director
	Scanning and Transmittal of certificate or disapproval letter to the FDA Records Section.	None	1 working day	CDRRHR Administrative Staff
	Queuing and endorsement to the Releasing Section	None	1 working day	AFS Records Officer / Administrative Officer
	TOTAL	Php1,010.00	20 working days**	

^{*}Submission period is within thirty (30) days from the issuance date of the Letter of Disapproval/Re-application.

**Service is covered under Republic Act No. 3720 Section 21 as amended by Executive Order No. 175 Section 13.