Form MD-7

[See sub-rule (1) of rule 21 and sub-rule (2) of rule 21]

Application for Grant of Licence to Manufacture for Sale or for Distribution of Class C or Class D

1. Name of Applicant:		
2. Nature and constitution	on of manufacturer:	
(i.e. proprietorship, p	partnership including Limi	ted
Liability Partnership,	private or public company	y, society,
trust, other to be spec	cified)	
3. (i) Corporate/ register	red office address includin	g
telephone number, m	obile number, fax number	and e-
mail id:		
(ii) Manufacturing si	te address including teleph	one
number, mobile num	ber, fax number and e-mai	id:
(iii) Address for corr	espondence:	
[corporate/ regist	ered office/ manufacturing	site]
4. Details of medical de	vice(s) to be manufactured	[Annexed]:
5. Whether substantial of	equivalence to a predicate of	levice is claimed: (Yes/ No)
6. Fee paid on	Rs	receipt/challan/transaction id
7. I have enclosed the d	ocuments as specified in th	e Fourth Schedule of Medical Devices Rules, 2017.
8. I hereby state and und	lertake that:	
(i) the manufacturing	site is ready for audit or sl	nall be ready for audit onin accordance with
the requirements of	Medical Devices Rules, 20	17.
(ii) I shall comply wi	th all the provisions of th	e Drugs and Cosmetics Act, 1940 (23 of 1940) and the Medical
Devices Rules, 2017	<i>'</i> .	
Place:		Signature
Date:		(Name and designation
		[To be signed digitally]

Annexure:

S.N.	Generic	Model	Intended	Class	Material of	Dimension	Shelf	Sterile or	Brand
	name	No.	use	of	construction	(if any)	life	Non	Name (if
				medical				sterile	registered
				device					under the
									Trade
									Marks
									Act,
									1999)