

IMDRF Membership Application Form

Applications must be submitted at least two (2) months before an IMDRF Management Committee meeting, which are usually held two times each year (for example, March and September (variable each year)).

If the application is for a Regional Harmonization Initiative, the application must be submitted by the Chair of the RHI. Any questions should be directed to the Chair of the IMDRF Management Committee which is listed on the IMDRF website.

Type of Membership

If other, please specify:
Contact Details for Applicant: Name of Applicant Organization:
Contact Person(s):
Title:
Address:
Phone:
Email:
Depending on the type of application, please fill out the corresponding section:

Management Committee Member

Have you been an Official Observer for at least the past 3 years?
 Yes
 No

2. List the IMDRF meetings (including teleconferences) your organization has attended in at least the past 3 consecutive years:

3.	List the IMDRF Working Groups to which you have appointed experts and have been actively involved in:			
OF	FICIAL OBSERVER			
Auth	nority			
1.	Are you a Regulatory Author			
	Yes	No		
2.	Do you have laws and regulations in place for medical devices that build on GHTF and IMDRF foundations and principles?			
		No		
	If yes, please provide the relevant description of related enforcemen	law or regulation, a comprehensive description of its contents and a t activities:		
3.	Please describe any activiti	es or initiatives you undertook or are currently undertaking		
J .	to bring scientific or regulate	ory innovation in the field of medical devices, including any lerging and technical regulatory issues:		
4.	IMDRF guidance document	onformity assessment of devices building on GHTF and ts? No		
		n of your conformity assessment program:		

Contribution to IMDRF

lmp	lementation of IMDRF Gu	idelines				
6.		egy regarding the implementation of IMDRF guidelines:				
7.		RF documents were implemented and provide relevant				
	documentation to support	evidence of implementation:				
REGIONAL HARMONIZATION INITIATIVE						
1.	Are you an association/initiative comprising medical device regulatory authorities representing the majority of countries in a certain region/area of the world?					
	Yes	No .				
	If yes, please describe the count	ries/region you are representing:				
2.	Do you have a mandate of Yes	regional harmonization amongst your members?				
	100					

Describe how your organization contributes or can contribute resources and expertise to the objectives of IMDRF and how its membership would be a benefit to IMDRF:

3.	Please provide a brief description of the activities you goals of fostering global regulatory convergence, leve available safe and effective medical devices globally:	eraging resources and making
4.	Please state who would be representing your RHI (e.	g. Member, Chair, Secretariat, etc):
 Signa	ture	Date