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Cardiovascular implants and extracorporeal systems - Vascular device-drug combination products - Part 2: Local regulatory information

Contents

Page

Foreword	iv
Introduction	v
1 Scope	1
2 Normative references	1
3 Terms and definitions	1
4 Information on device- and drug-related aspects -- Applicable documents for local guidance	4
4.1 Australia	4
4.1.1 Australia: managing changes	4
4.1.2 Australia: clinical evaluation requirements	5
4.1.3 Australia: audit requirements	5
4.2 Brazil	5
4.2.1 Brazil: managing changes	5
4.2.2 Brazil: clinical evaluation requirements	6
4.2.3 Brazil: audit requirements	6
4.3 Canada	6
4.3.1 Canada: managing changes	6
4.3.2 Canada: clinical evaluation requirements	6
4.3.3 Canada: audit requirements	6
4.4 European Union (EU)	6
4.4.1 EU: managing changes	6
4.4.2 EU: clinical evaluation requirements	7
4.4.3 EU: audit requirements	7
4.5 India	7
4.5.1 India: managing changes	7
4.5.2 India: clinical evaluation requirements	8
4.5.3 India: audit requirements	8
4.6 Japan	8
4.6.1 Japan: managing changes	8
4.6.2 Japan: clinical evaluation requirements	8
4.6.3 Japan: audit requirements	8
4.7 People's Republic of China (PRC)	8
4.7.1 PRC: managing changes	8
4.7.2 PRC: clinical evaluation requirements	9
4.7.3 PRC: audit requirements	10
4.8 Russia	10
4.8.1 Russia: managing changes	10
4.8.2 Russia: clinical evaluation requirements	10
4.8.3 Russia: audit requirements	10
4.9 United States of America (USA)	10
4.9.1 USA: managing changes	10
4.9.2 USA: clinical evaluation requirements	11
4.9.3 USA: audit requirements	12
5 Managing changes that can impact the DCP	12
5.1 Introduction	12

5.2	Change evaluation	12
5.2.1	Identify changes	12
5.2.2	Risk evaluation	13
5.2.3	Guidance for change evaluation	14
5.2.4	Pre-market (during submission)	14
5.3	Interactions with region-specific regulatory authorities - post commercialization	14
Bibliography		23