

TISSUE ENGINEERING INTELLIGENCE UNIT 5

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Biomaterials in the Design and Reliability of Medical Devices



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CONTENTS

Preface	viii
1. Overview and Introduction: Unique Aspects of Biomaterials in the Safety and Efficacy of Medical Implant Devices	1
<i>Michael N. Helmus</i>	
Materials Selection	1
Functional Requirements	2
Biomaterials and Regulatory Guidelines	53
2. Standards and Guidelines for Biocompatibility of Medical Devices	74
<i>Sharon J. Northup</i>	
Product Registration Requirements	74
Standards and Guidelines for Biocompatibility Testing	106
Chemical Characterization	107
Risk Assessment	113
Case Studies of Materials Toxicity Risk Assessment	117
Conclusion	125
3. Regulation of Medical Devices	127
<i>Barry Sall</i>	
U.S. Food and Drug Administration Regulation of Medical Devices	127
4. Nonclinical Medical Device Testing.....	144
<i>Sharon J. Northup</i>	
Overview of Biocompatibility Procedures	144
Sample Preparation	146
Detailed Description of Biocompatibility Procedures	148
Cytotoxicity	149
Sensitization	150
Irritation Assays	152
Systemic Toxicity (Single and Repeated Dose Toxicity Including Pyrogenicity)	152
Pyrogenicity	156
Genotoxicity	157
Implantation	160
Hemocompatibility	162
Chronic Toxicity and Carcinogenicity	164
Reproductive and Developmental Toxicity	165
Biodegradation	165
Toxicokinetic Studies	166
Effectiveness Testing	167
Externally Communicating Devices	169
Implanted Devices	169
Conclusion	170

5. Failure Analysis: Learning for the Future from the Past	172
<i>Michael N. Helmus</i>	
6. Product Development in a Small Company Environment	178
<i>Roger W. Snyder</i>	
Records and Record Keeping	179
Testing	181
Materials and Components	182
Prototyping	184
Vendor Relationships	185
Sterilization and Shelf Life	186
Production Facilities	187
Conclusion	189
7. Tissue Engineering Constructs and Commercialization	191
<i>Kelvin G.M. Brockbank</i>	
Case Studies: Development of Effective Transport Solutions and Devices to Enable Product Distribution	194
Development of Methods to Increase Product Shelf-life	195
8. Testing of Biomaterials Modified with Bioactive Molecules: A Case Study	198
<i>Katherine S. Tweden</i>	
Characterization of the Nature and Uniformity of the Modification	199
Quantitation of the Modification	202
Assessment of the Biological Activity In Vitro	202
In Vitro Challenges	204
Manufacturing Ruggedness	204
Assessment of the Biological Activity In Vivo	205
Immobilization of an RGD-Containing Peptide: A Case Study	206
Materials	206
Results	209
Appendix: Selecting Contract Labs	223
<i>Barry Sall</i>	
Index	224