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Prefilled syringes - Part 6: Plastic barrels for injectables and sterilized subassembled syringes ready for filling (ISO 11040-6:2019)

Contents	Page
National foreword	4
National Annex NA (informative) Bibliography	5
Foreword	7
Introduction	8
1 Scope	9
2 Normative references	9
3 Terms and definitions	10
4 General requirements	11
4.1 Quality systems	11
4.2 Testing	11
4.3 Documentation	11
5 Dimensions and designation	12
5.1 Design including dimensions	12
5.2 Design requirements	15
5.2.1 Head design	15
5.2.2 Dead space	15
5.2.3 Functional testing of Luer cone/Luer lock connection	15
5.2.4 Flange breakage resistance	15
5.2.5 Syringe tip breakage resistance	15
6 Requirements	15
6.1 General	15
6.2 Material	16
6.2.1 General	16
6.2.2 Duty of notification concerning modifications to polymers	16
6.2.3 Needle	16
6.2.4 Closure system	17
6.2.5 Closure system integrity	18
6.3 Physical requirements	18
6.3.1 Sterilization	18
6.3.2 Clarity and transparency	18
6.3.3 Particulate contamination	18
6.3.4 Lubricants	19
6.4 Chemical requirements	19
6.5 Endotoxins and biological requirements	19
7 Graduation	20
8 Packaging and labelling	20
Annex A (informative) Examples of types of sterilized subassembled syringes ready for filling	21
Annex B (informative) Head designs	24

Annex C (normative) Test methods for syringe barrels	26
Annex D (informative) Sample preparation for endotoxin and particulate determination	32
Annex E (informative) Evaluation of syringe lubrication by glide force test method	36
Annex F (informative) Needle penetration test	39
Annex G (normative) Test methods for syringe closure systems	42
Annex H (informative) Dye solution tightness test	57
Annex I (informative) Guidance on materials	59
Bibliography	60