Council Directive of 7 June 1988 on the inspection and verification of Good Laboratory Practice (GLP) (88/320/EEC) (repealed)

Article 1	
Article 2	
Article 3	
Article 4	
Article 5	
Article 6	
Article 7	
Article 8	
Article 9	
Article 10	
	ANNEX

PART A

REVISED GUIDES FOR COMPLIANCE MONITORING PROCEDURES FOR GOOD LABORATORY PRACTICE

Definitions of terms

Components of good laboratory practice compliance monitoring procedures

Administration
Confidentiality
Personnel and training
(National) GLP compliance programmes
Follow-up to test facility inspections and study audits
Appeals procedures

PART B

REVISED GUIDANCE FOR THE CONDUCT OF TEST FACILITY INSPECTIONS AND STUDY AUDITS

Introduction

Definitions of terms

Test facility inspections

Inspection procedures

Status: EU Directives are being published on this site to aid cross referencing from UK legislation. After IP completion day (31 December 2020 11pm) no further amendments will be applied to this version.

Pre-inspection Starting conference Organisation and personnel Quality assurance programme **Facilities** Care, housing and containment of biological test systems Apparatus, materials, reagents and specimens Test systems

Physical and chemical systems Biological test systems Test and reference substances Standard operating procedures Performance of the study Reporting of study results Storage and retention of records

Study audits

Completion of inspection or study audit