

Buruburu Business Complex Suite No.26, Mumias South Road, Nairobi. P.O Box 4963-00100, Nairobi, Kenya.

ADVANCED CURRENT GOOD MANUFACTURING PRACTICES(cGMP) 3 DAYS 8th – 10th MAY 2024

This course is presented over 3 days and provides a real depth of information on the main aspects of pharmaceutical GMP. During the course the **main GMP requirements Quality Control and batch release** are covered. In addition, the course also covers the main elements of the **Quality Management System** needed to provide medicines of the highest quality, including the requirements for **documentation**, **training and system monitoring and review**. The course is full of **interactive exercises** and workshops throughout the programme.

DAY 1 (8 th May 2024)				
9.00-9.30	Principles & Practices in cGMP			
	Introduction and Benefits of cGMP			
0.20.10.20				
9.30-10.30	GMP –Rules & guidelines;			
	• European Union (EU) GMP and EU Guide to GMP			
10.30 - 11.00	GMP in the United States Other GMP from Around the world Tea Break			
11.00-13.00	Premise & Facility Design			
	Suitable premises and Facility design			
	 Heating Ventilation and Air conditioning Access ,Security and Pest Control 			
13.00- 14.00	Lunch Break			
14.00 - 16.30	Equipment, Maintenance and Calibration			
14.00 - 10.30	Selection of equipment and Installation			
Day 2 (9 th May 2024)	Planned Preventative Maintenance Calibration of measuring equipment's			
9.00 - 10.30				
9.00 - 10.30	 Good Manufacturing Practices(GMP) Regulations; CRF role in cGMP Regulation 			
	 21 CFR Part 210: Processing, Packing, or Holding 			
10.30 - 11.00	Tea Break			
11.00 – 13.00	21 CFR Part 211: Finished Pharmaceuticals			
	21 CFR Part 600: Biological Products			
13.00 – 14.00	Lunch Break			
14.00 -16.30	• 21 CFR Part 600: Biological Products:			
4	21 CFR Part 11: Electronic Records and Signatures			
DAY 3 (10 th May 2024)				
09.00 - 10.30	Good Manufacturing Practices (GMP) and Quality Management			
	System(QMS)			
	People & Training			
	Organization charts, Job description and training records			
	• GMP and job specific training			
	Training design and evaluation			

10.30 - 11.00	Tea Break				
11.00 – 13.00	Key Personnel in GMP				
	• The Heads of Production, QC and Qualified personnel				
	The role of Quality and Quality Assurance				
	The importance of Senior management				
	Documentation, Records and Data integrity				
	Control and approval of documents and records				
	Data integrity and regulatory concerns				
13.00 - 14.00	Lunch Break				
14.00 – 15.30	Quality Risk Management				
	Decision making based on risk				
	• ICH Q9 and its requirements				
	Reactive & Proactive risk assessments				
	The Quality Management Systems				
	Batch review, and release, Product quality review, Internal, Auditing,				
	Management review.				
15.30 - 16.00	Directors remarks and issue of certificates				
Date		Cost	Venue		
8 th – 10 th May 2024		Ksh. 63,800.00 or	NAIROBI		
		USD 638.00			
Reg: Deadline: Asap					