

ADVANCED CURRENT GOOD MANUFACTURING PRACTICES(ACGMP)

25th - 27th
MARCH 2026

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	25-03-26	EVENTS
		Registration and Climate Setting
9:00 – 09:30 am		Principles & Practices in cGMP • Introduction and Benefits of cGMP
09:30 – 10:30 am		TEA- BREAK
11:00 – 13:00 p.m		GMP –Rules & guidelines; • European Union (EU) GMP and EU Guide to GMP • GMP in the United States Other GMP from Around the world
13:00 – 14:00 p.m		LUNCH - BREAK
14:00 – 16:30 p.m		Equipment, Maintenance and Calibration • Selection of equipment and Installation • Planned Preventative Maintenance Calibration of measuring equipment's

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Day 2	26-03-26	EVENTS
9:00 – 10:30 am	Good Manufacturing Practices(GMP) Regulations; <ul style="list-style-type: none"> • CRF role in cGMP Regulation • 21 CFR Part 210: Processing, Packing, or Holding 	
10:30 – 11:00 am	TEA- BREAK	
11:00 – 13:00 p.m	<ul style="list-style-type: none"> • 21 CFR Part 211: Finished Pharmaceuticals • 21 CFR Part 600: Biological Products 	
13:00 – 14:00 p.m	LUNCH - BREAK	
14:00 – 16:30 p.m	<ul style="list-style-type: none"> • 21 CFR Part 600: Biological Products: • 21 CFR Part 11: Electronic Records and Signatures 	

Day 3	27-03-26	EVENTS
9:00 – 10:30 am	Good Manufacturing Practices (GMP) and Quality Management System(QMS) People & Training <ul style="list-style-type: none"> • Organization charts, Job description and training records • GMP and job specific training • Training design and evaluation 	
10:30 – 11:00 am	TEA- BREAK	
11:00 – 13:00 p.m	Key Personnel in GMP <ul style="list-style-type: none"> • 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 p.m	LUNCH - BREAK	
14:00 – 15:30 p.m	Quality Risk Management <ul style="list-style-type: none"> • 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 p.m	Directors remarks and issue of certificates	

Deadline: 11th March 2026

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Cost Kes. 90,000.00
or USD 850.00
exclusive of taxes

NAIROBI

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Online Training available at: www.chromafrica.com