



Auditing

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	ОСТ	NOV	DEC
Active Pharmaceutical Ingredients (API) Auditing	1						5 Virtual						
Auditing to Pharmaceutical Standards	1			12 Virtual									2 Virtual
GMP Lead Auditor – Pharmaceutical Quality Systems (IRCA Ref: PR325)	5	15-19 Virtual	Cook	o-1 Mar erson kham, IK		20-24 In person Crewe, UK		1-5 In person Cookham, UK			14-18 Virtual		
QMS Lead Auditor – Pharmaceutical Supply Chain (IRCA Ref: PR330)	5						17-21 In person Cookham, UK			9-13 Virtual		18-22 In person Cookham, UK	
Internal Auditor	2		7-8 Virtual				5-6 In person Reading, UK					4-5 In person Liverpool, UK	
GDP/RP Auditor	2	10-11 Virtual		26-27 In person Cookham, UK				2-3 In person Reading, UK		3-4 In person Reading, UK			4-5 In person Reading, UK
Virtual Auditing Skills	1			14 Virtual			12 Virtual				2 Virtual		

Biopharmaceutical

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	OCT	NOV	DEC
Introduction to ATMPs and route to market	1				4 Virtual								
Good clinical practice for ATMPs	1			11 Virtual				9 Virtual					
Biotechnology Issues	1					20 Virtual						4 Virtual	

Information is correct at the time of going to print.
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Good Manufacturing Practice

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	OCT	NOV	DEC
Good Manufacturing Practice Advanced RSC Approved	3			25-27 Virtual		13-15 In person Liverpool, UK		15-17 In person Reading, UK		23-25 Virtual		12-14 Virtual	
Good Manufacturing Practice – The Essentials	1			21 Virtual		8 Virtual		1 Virtual				5 Virtual	
Good Manufacturing Practice in the Laboratory	1				4 Virtual					2 Virtual			
Interpretation of Statistical Values Used in GMP Applications	1		12 Virtual				19 Virtual						
Management of GMP Inspections	2			7-8 Virtual									
Medical Device Regulations (MDR) 2017/745	1					8 Virtual						6 Virtual	
Pharmaceutical Product Development – GMP Requirements	2			4-5 Virtual						9-10 In person Reading, UK			
Pharmaceutical Quality Management System	3				23-25 Virtual							6-8 In person Liverpool, UK	

Microbiology

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	OCT	NOV	DEC
Microbiology Environmental Monitoring in Partnership with Cherwell Laboratories	1		6 In person Oxford, UK								31 In person Oxford, UK		
Introduction to Pharmaceutical Microbiology	1				17 Virtual					12 Virtual			
Manufacturing Sterile Products	2				25-26 In person Reading, UK					19-20 Virtual		26-27 Virtual	
Pharmaceutical Microbiology Advanced	3				8-10 In person Wokingham, UK			29-31 In person Wokingham, UK			15-17 In person Wokingham, UK		
Water Systems and Microbiological Control	1				23 Virtual					5 Virtual			



Qualified Person



APPROVED TRAINING

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COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	ОСТ	NOV	DEC
Active Pharmaceutical Ingredients	2				15-16 In person Reading, UK		5-6 In person Liverpool, UK					18-19 Virtual	
Analysis and Testing	3				22-24 In person Reading, UK			8-10 In person Reading, UK			21-23 In person Reading, UK		
Biotechnology Issues	1					20 Virtual						4 Virtual	
Investigational Medicinal Products	2	10-11 Virtual				16-17 In person Liverpool, UK							2-3 In person Reading, UK
Mathematics and Statistics	3		20-22 Virtual					3-5 In person Liverpool, UK				5-7 In person Reading, UK	
Medicinal Chemistry and Therapeutics	3			11-13 Virtual						2-4 Virtual			
Pharmaceutical Formulation and Processing Part 1*	2*	29-30 Virtual				28-29 In person Reading, UK				9-10 Virtual			
Pharmaceutical Formulation and Processing Part 1* – Optional Practical Day	1*					30 In person Reading, UK				11 In person Reading, UK			
Pharmaceutical Formulation and Processing Part 2*	3*			5-7 In person Reading, UK				22-24 In person Liverpool, UK				26-28 Virtual	
Pharmaceutical Law and Administration	2	22-23 Virtual					3-4 In person Reading, UK				1-2 In person Liverpool, UK		
Pharmaceutical Microbiology	3				8–10 In person Wokingham, UK			29-31 In person Wokingham, UK			15-17 In person Wokingham, UK		
Pharmaceutical Packaging	3		5-7 Virtual							16-18 In person Barnstaple, UK			10-12 In person Barnstaple, UK
Pharmaceutical Quality Systems	3	24-26 Virtual					25-27 In person Liverpool, UK				8-10 In person Reading, UK		
Role and Professional Duties	2	15-16 Virtual				7-8 In person Reading, UK					30-31 In person Liverpool, UK		

^{*}Please note to cover the criteria set out in the QP study guide, both Part 1 and Part 2 should be attended.





Supply Chain

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	OCT	NOV	DEC
Good Distribution Practice	1			20 Virtual			4 Virtual				14 Virtual		
Responsible Person and GDP (Cogent Gold Standard Approved)	3		12-14 In person Reading, UK			13-15 In person Reading, UK		15-17 Virtual		16-18 In person Liverpool, UK		6-8 Virtual	
Responsible Person Refresher	1				18 Virtual			25 Virtual				28 Virtual	
Responsible Person Forum	2 hours	19 Virtual		1 Virtual		10 Virtual		19 Virtual		13 Virtual		15 Virtual	

Validation

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	OCT	NOV	DEC
Cleaning Validation	1			14 Virtual							15 Virtual		
Introduction to Validation	1					1 Virtual					2 Virtual		
GxP Computerised Systems Validation and Compliance	2		7-8 Virtual			1-2 Virtual					23-24 Virtual		
Process Validation and Qualification, including Validation Methods	2			26-27 In person Reading, UK						17-18 In person Reading, UK			
GAMP 5 Second Edition Workshop	1					14 In person Reading, UK						5 In person Reading, UK	

Group booking discount

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Contact our team for details.

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Other Courses

COURSE	DAYS	JAN	FEB	MAR	APR	MAY	JUN	JUL	AUG	SEPT	OCT	NOV	DEC
Data Integrity, Electronic Records and Signatures	1			7 Virtual							22 Virtual		
Introduction to Pharmacovigilance	1						27 Virtual						
Leadership and Influencing Skills for QPs & New Managers	1		22 Virtual								29 Virtual		
Quality Risk Management – How to apply ICH Q9 in Practice	1				29 In person Reading, UK			9 In person Reading, UK				20 In person Reading, UK	
QC Chemistry Crash Course	1			25 Virtual						24 Virtual			
Root Cause Analysis and CAPA	2			20-21 Virtual									5-6 Virtual
Technical Report Writing	1		15 Virtual					11 Virtual					
Technology Transfer of Pharmaceutical Products	1		5 In person Reading, UK			7 In person Reading, UK							

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- Preparing for a Regulatory Inspection
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- Annex 1
- Data Integrity
- Good Manufacturing Practice
- Responsible Person
- GMP Lead Auditor
- Internal Auditor
- Cleaning Validation





About RSSL Training & Consultancy

We've been training and developing pharmaceutical, biopharmaceutical, medical device, and healthcare professionals across the complete life cycle of pharmaceutical products for over 30 years. We know what it takes to succeed and employ leading industry experts to deliver the most up-to-date training and CPD so individuals can achieve their career aspirations and companies have the skills they need to excel on the global stage and make a positive difference in the world.



RSSL Training & Consultancy

The Reading Science Centre Whiteknights Campus Pepper Lane Reading Berkshire RG6 6LA

Tel: **+44 (0)118 918 4076**

Email: trainingsales@rssl.com

Web: www.rssl.com

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