RIQAS

COAGULATION PROGRAMME

RQ9135

	Lab. Reference Number	
Please tick the correct option:	agulation	
	This is an update to an existing Co	agulation registration
If you wish to register multiple instruments	s, please complete separate enrolment	documents for each instrument
On each document please state an instrur	ment identification name here	
Instrument Group Reports Instrument group reports can be provided on request. Pleas Inter-Laboratory Group Reports To receive inter-laboratory group reports, please contact RI		ibutor for more details.
Please indicate cycles required in boxes Cycle 17 January 2025 - December 2025	s below Cycle 18	January 2026 - December 2026
Please indicate kit required		12 months
RQ9135/a - Coagulation (5 Selected parameter	rs only: PT, aPTT, TT, Fibrinogen, Anti-thro	ombin III)
RQ9135/b - Coagulation Programme (all analyt	tes, choose from 16)	
Primary Contact Details: (CAPITAL LET	TERS ONLY)	
QA Officer		
Laboratory / Hospital Name		
Department		
Postal Address		
City	State	
Postal / Zip Code Cou	ntry	
Telephone Number		
Randox Office / Distributor		1

Lab. Reference Number	
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RIQAS COAGULATION PROGRAMME

RIQASNet - ELECTRONIC CORRESPONDENCE

Participation on RIQAS requires access to RIQASNet, a web-based online method for result entry, viewing of released reports and addition or change of assay details. In addition, PDF reports can be e-mailed to up to 3 e-mail addresses. Internet access and login details are required for RIQASNet. A login will be supplied by RIQAS based on "e-mail address 1" below. It is also possible to receive a csy file containing the information found on the summary page of the routine report.

I wish to receive a summary csv file (csv files must be sent to the same email addresses as the PDF reports)	FOR RIQAS USE ONLY RIQASNet No
	Date added: By:
	PDF copies set to
	csv copies set to

Primary Contact email for RIQASNet/PDF reports/summary csv files (Please write in capital letters only)					
E-mail address 1:					
E-mail addresses for	additional PDF reports/summary csv files				
E-mail address 2:					
E-mail address 3:					

Customer Declaration: By submitting this enrolment document to RIQAS, either directly or via my local Randox representative, I, (the customer of RIQAS) confirm that:

- 1) I have read and understood the RIQAS policies stated in the most recent Method Questionnaire associated with this programme.
- 2) I understand that the submission of this enrolment document to RIQAS marks the beginning of an on-going agreement, and I will be automatically enrolled in subsequent cycles of this programme until RIQAS receives written confirmation of my cancellation. This should be received by RIQAS 12 weeks prior to the month in which the cycle starts.
- 3) I understand that I must inform RIQAS of any changes to my contact details, assay details or contract status
- 4) I authorise Randox Laboratories Ltd. to send communication related to the products and service provided to the e-mail or postal addresses stated on this document
- 5) I understand that I am permitted to request disclosure of, change or erase personal details held by Randox Laboratories Ltd. at any time.

REGISTRATION OF ASSAY DETAILS

It is possible to inform RIQAS of your chosen parameters and assay details by

- 1) Completing the 'REGISTRATION OF ASSAY DETAILS' on the following pages OR
- 2) Adding your own assay details using RIQASNet

Please select one of the following options

	I wish to add my own assay details via RIQASNet once I have received my username, password and Lab Reference Number from RIQAS (You do not need to complete the 'REGISTRATION OF ASSAY DETAILS' section of this document)
П	I wish to inform RIQAS of my assay details using this enrolment document
	(please complete all remaining pages of the 'REGISTRATION OF ASSAY DETAILS' section)

For any further queries, please contact your local Randox office, Sales Representative or RIQAS directly.

Please contact RIQAS at

Tel: +44 (0) 28 9445 4399
E-Mail: mail@riqas.com
RIQAS Scheme Co-ordinator: Sarah Fleck

 $RANDOX\,LABORATORIES\,LTD.,\,55\,Diamond\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,BT29\,4QY,\,United\,Kingdom\,Road,\,Crumlin,\,County\,Antrim,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,County,\,Count$

This programme is accredited by UKAS TO ISO/IEC 17043:2010 via Fixed Scope



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RIQAS COAGULATION PROGRAMME

REGISTRATION OF ASSAY DETAILS

ONLY COMPLETE THIS SECTION IF YOU DO NOT INTEND TO REGISTER YOUR METHODS VIA RIQASNET

Please indicate your requirements by ✓ or by writing in the boxes below. Current participants should complete the document only for method changes.

This programme is not suitable for use with the ITC Hemochron Signature or Roche Coaguchek Instruments.

ANALYTE	METHOD CODE	INSTRUMENT	REAGENT	SI UNITS 🗸	OTHER UNITS
FACTOR II ACTIVITY				%	N/A
FACTOR V ACTIVITY				%	N/A
FACTOR VII ACTIVITY				%	N/A
FACTOR VIII ACTIVITY				%	N/A
FACTOR IX ACTIVITY				%	N/A
FACTOR X ACTIVITY				%	N/A
FACTOR XI ACTIVITY				%	N/A
FACTOR XII ACTIVITY				%	N/A
D-DIMER (PILOT)				ug/l	
KIT NAME/ CATALOGUE NUMBER		KIT LOT NUMBER		CONVERSION FACTOR	
FIBRINOGEN				g/l	
PLASMINOGEN ACTIVITY				%	N/A
ANTITHROMBIN III ACTIVITY				%	N/A
PROTEIN C ACTIVITY				%	N/A
PROTEIN S ACTIVITY				%	N/A
FREE PROTEIN S				%	N/A
aPTT AS A RATIO				Ratio	N/A
aPTT IN SECONDS				s	N/A

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Lau.	Reiei	ence	NUII	ıber

Eub. Reference Rumber						

RIQAS COAGULATION PROGRAMME REGISTRATION OF ASSAY DETAILS

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ANALYTE	METHOD CODE	INSTRUMENT	REAGENT	SI UNITS 🗸	OTHER UNITS
PT AS AN INR				INR	N/A
PT ACTIVITY				%	N/A
PT AS A RATIO				Ratio	N/A
PT IN SECONDS				s	N/A
THROMBIN TIME				s	N/A
Please use this space to de	escribe "other" metho	ods, instruments	s and reagents.		

PLEASE NOTE: PT may be registered as seconds, % activity, ratio or INR. Registration of all of these will count as one analyte. aPTT may be registered as seconds or ratio. Registration of both will count as one analyte PLEASE NOTE Some users of ACL TOP instruments when used in combination with Hemosil Synthasil reagents may be unable to attain a clot with extended read time samples.