# RANDOX

# RANDOX INTERNATIONAL QUALITY ASSESSMENT SCHEME





# RIQAS

# THE LARGEST INTERNATIONAL EQA SCHEME WITH OVER 45,000 LAB PARTICIPANTS



01	BENEFITS
02	EQA
03	RIQAS REPORTS
04	Web-based data transfer
05	PARTICIPATION IN RIQAS
06	STANDARD REPORT
07	TEXT SECTION
08	HISTOGRAM
09	LEVEY-JENNINGS CHART
10	TARGET SCORE CHART
П	%DEVIATION CHARTS
12	MULTI-METHOD STAT SECTION
13	SUMMARY PAGE
14	MULTI-INSTRUMENT REPORT
15	URINE TOXICOLOGY REPORT
16	URINE TOXICOLOGY REPORT SCREENING SECTION
17	URINE TOXICOLOGY REPORT QUANTITATIVE SECTION
18	URINALYSIS REPORT
19	SEROLOGY: SCREENING (QUALITATIVE) REPORT
20	SEROLOGY: SCREENING (QUANTITATIVE) REPORT
21	QUANTITATIVE END-OF-CYCLE REPORT
22	CHART SECTION (END-OF-CYCLE REPORT)
23	TEXT SECTION (END-OF-CYCLE REPORT)
25	CURRENT & PREVIOUS CYCLE ABSOLUTE SDIs (END-OF-CYCLE REPORT)
26	CERTIFICATE OF PERFORMANCE (END-OF-CYCLE REPORT)
27	MONITORING EQA PERFORMANCE
30	RIQAS PROGRAMMES
35	PARAMETER INDEX
49	RELATED PRODUCTS
51	RANDOX - A GLOBAL DIAGNOSTIC SOLUTIONS PROVIDER
52	CONTACT US

#### **BENEFITS**

Delivering a comprehensive yet cost effective EQA solution, RIQAS will help meet regulatory requirements and increase confidence in test system accuracy.



#### Large Database of Users

• A high level of participation means peer group numbers are maximised whilst ensuring availability of data for a wide range of instruments and methods.



#### User-friendly Reports

- Simple, one page per parameter format, enables at-a-glance performance assessment, saving valuable laboratory time.
- Complimentary multi-instrument and interlaboratory reports allow comparative performance assessment of all laboratory systems and multiple connected laboratories.
- End-of-Cycle reports, summarising performance compared to the previous cycle, allow you to identify improvements in quality over time.



#### Cost Effective

- Our extensive range of multi-analyte programmes will reduce the number of individual programmes required to cover your test menu, saving both time and money.
- Reduced parameter options for selected programmes offer greater flexibility, ensuring suitability for laboratories of all sizes and budgets.
- Register up to five instruments per programme (volume permitting) at no extra cost for comparative performance assessment.



#### Frequency

- Frequent reporting allows early identification of system errors and implementation of any necessary corrective actions with minimum disruption to the lab.
- With a turnaround of less than 72 hours for most reports, corrective action can be implemented earlier, potentially reducing costly errors with patient results.



#### High Quality Samples

- Samples spanning clinically relevant levels allow identification of concentration related biases, helping to ensure accurate instrument performance.
- Human samples free from interfering preservatives increase confidence that EQA performance mirrors the performance of patient samples.
- Reference method values are provided in the Clinical Chemistry programme for selected parameters and lots while for the Immunosuppressant programme they are provided for all parameters and lots.



#### Highly Accredited

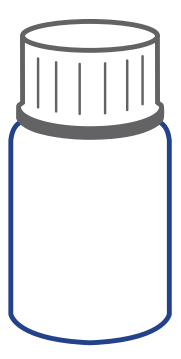
- Programmes accepted by National and International accreditation bodies worldwide.
- Participant certificates provide evidence of participation in a reputable EQA scheme.

RIQAS is the largest international EQA scheme in the world. It is used by more than 45,000 laboratory participants in 133 countries. 33 programmes are currently available.

#### **RIQAS Programmes**

- Ammonia/Ethanol
- Anti-TSH Receptor
- Blood Gas
- BNP
- Cardiac
- Cerebrospinal Fluid (CSF)
- Clinical Chemistry
- Coagulation
- CO-Oximetry
- CYFRA 21-1
- ESR
- Glycated Haemoglobin (HbA1c)
- Haematology
- Human Urine
- Immunoassay
- Immunoassay Speciality 1
- Immunoassay Speciality 2

- Immunosuppressant Drugs
- Lipid
- Liquid Cardiac
- Maternal Screening
- Serology Epstein Barr Virus (EBV)
- Serology (HIV/Hepatitis)
- Serology (Syphilis)
- Serology (ToRCH)
- Specific Proteins
- Sweat Testing
- Therapeutic Drugs
- Trace Elements in Blood
- Trace Elements in Serum
- Trace Elements in Urine
- Urinalysis
- Urine Toxicology



#### Accreditation

- RIQAS provides certificates as proof of EQA participation and performance for laboratory accreditation purposes.
- RIQAS is a UKAS accredited Proficiency Testing Provider, No. 0010, and is accredited to ISO/IEC 17043:2010, 'Conformity Assessment- General Requirements for Proficiency Testing'.
- Accreditation to ISO/IEC 17043:2010 highlights the superior quality and excellence of RIQAS.

#### **UK Performance Surveillance**

- Recognised by the Joint Working Group on Quality Assurance (JWG QA).
- Recognised by various National Quality Assurance Advisory Panels (NQAAP).

#### Independent Advisory Panel

RIQAS participants have access to an independent advisory panel consisting of scientific and clinical experts. This ensures professional and ethical conduct of the scheme and participant confidentiality.

RIQAS support staff are on hand to offer advice and troubleshoot technical queries.

# **RIQAS REPORTS**

RIQAS reports are presented in a user-friendly, one page per parameter format. This allows easy interpretation of your analytical performance.

#### **RIQAS** Reports

- Statistical breakdown by all methods, your method and, where applicable, your instrument, including running means for the last 10 samples.
- Compare your instrument group, method group and all methods using the histogram.
- Identify trends, biases and precision problems using the visual charts.
- The Target Score chart grades your performance in a moving window over the last 20 samples, including the previous cycle.
- At-a-glance summary page for all parameters in the programme.
- Compare your result with statistically robust consensus means.
- Identify acceptable and poor performance using fit-for-purpose performance indicators:
  - SDI
  - %Deviation
  - Target Score



#### **Summary CSV Files**

It is possible to receive an additional summary of your report statistics, acceptable limits and performance indicators as a .csv file for every sample.

#### Multi-Instrument Reports

Laboratories can register up to five instruments at no extra cost. Individual reports for each instrument plus a unique multi-instrument report are provided. The multi-instrument report plots the performance of each individual instrument on a single, colour coded Levey-Jennings chart, ensuring instant identification of any differences in instrument performance. Additional sample packs may be ordered as required if volume supplied is insufficient for the registered instruments.

#### Laboratory Group Reports

The Group Reporting facility enables laboratory groups or chains to monitor the performance of satellite sites. Each affiliated laboratory will receive their individual reports with the group supervisor also receiving a summary report comparing each laboratory in the network.

### WEB-BASED DATA TRANSFER

# RIQAS.Net offers easy, direct access for the submission of results and retrieval of reports direct from the RIQAS host server.

- Available in multiple languages.
- Confidentiality and security is maintained through the use of password protected access.
- Submit current, corrected and future results (normal policies apply), directly into the RIQAS database. Receipt of results is confirmed by e-mail.
- Multi-lingual registration identifier provides simple identification of multiple registrations.
- Additions and changes to assay details can be made quickly and easily online.
- Requests for new method, instrument and reagent codes can be made online.
- Reports are emailed in PDF format as soon as they are prepared.
- Reports for the previous two cycles can be downloaded from the website.
- View, print, store or distribute reports as you wish.
- Update your laboratory's certificate of participation details in multiple languages.
- All that is required is web access, Adobe Reader (for viewing reports) and a valid password to access the system.
- No additional software required.



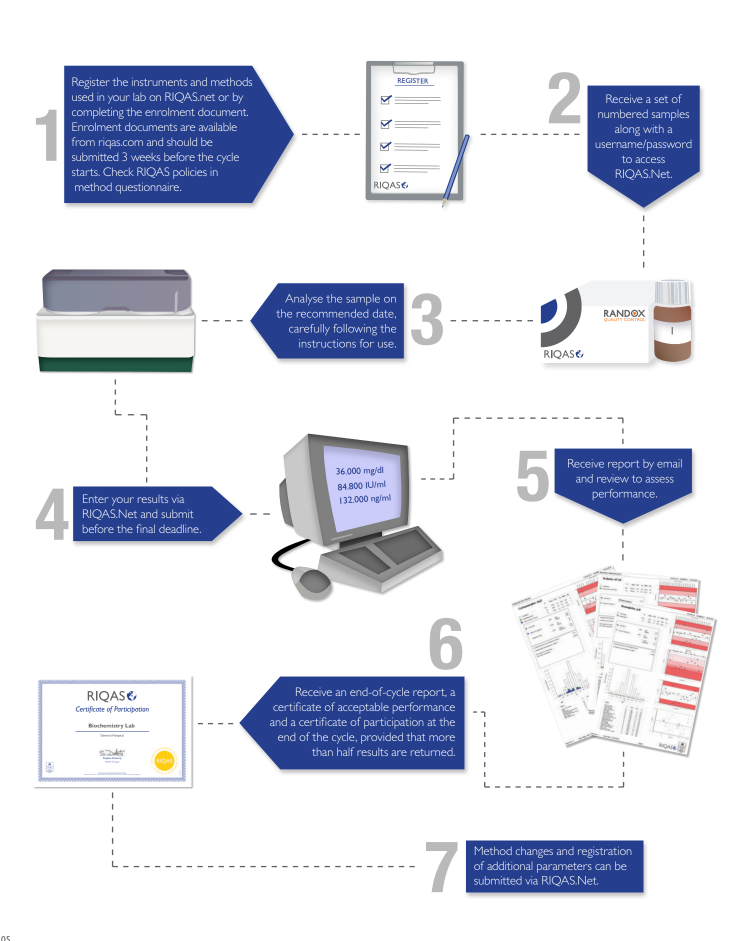






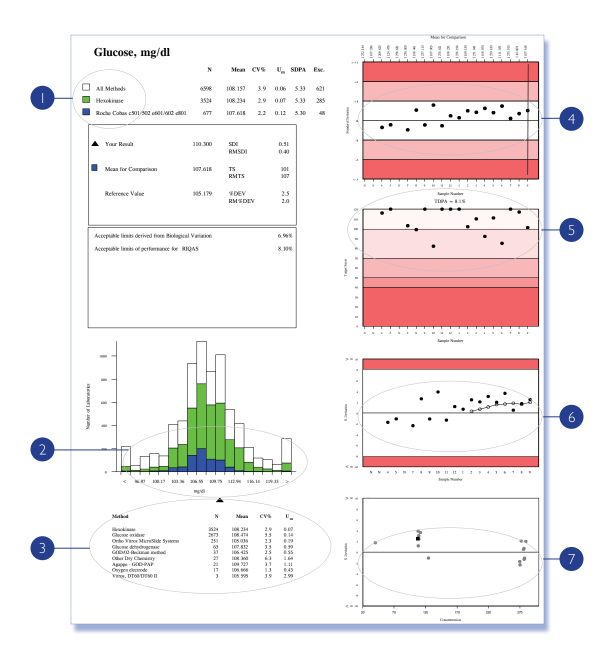
# PARTICIPATION IN RIQAS

#### Participation in RIQAS follows these simple steps:



# STANDARD REPORT

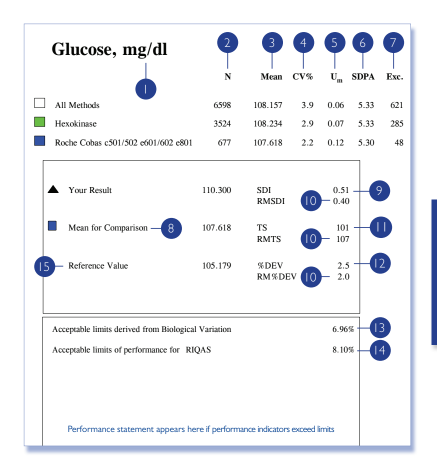
Performance data is presented in a one page format with up to seven sub-reports.



Text Section:	Statistics for all methods, your method and instrument group (programme specific).
Histogram:	Method and instrument comparison.
Multi-Method Stat Section:	Enables assessment of the performance of each method.
Levey-Jennings Chart:	Details features of your laboratory's performance.
Target Score:	This unique chart provides a numerical index of performance, allowing at-a-glance assessment.
%Deviation by Sample:	Helps to identify trends and shifts in performance.
%Deviation by Concentration:	Rapid assessment of concentration related biases.

### **TEXT SECTION**

#### The text section summarises the statistical information for each parameter.



RIQAS performance indicators include SDI, Target Score and %Deviation.

Acceptable performance criteria:

SDI < 2

Target score ≥ 50

%Deviation < defined acceptable limits

- Report is presented in your chosen unit.
- Number of returned results used to generate Mean for Comparison.
- 3 Average value of all laboratories' results.
- 4 Coefficient of Variation.
- Uncertainty associated with the Mean for Comparison.

$$U_{m} = 1.25 \times SD$$

SDPA = Standard Deviation for Performance Assessment, calculated from the Target Deviation for Performance Assessment (TDPA) and the Mean for Comparison.

$$SDPA = \frac{TDPA \times Mean for Comparison}{t-value \times 100}$$

t-value = factor which represents the % of poor performers reflected in the TDPA (t-value  $\sim$  1.645 when  $\sim$ 10% laboratories achieve poor performance), SDPA is combined with U  $_{\rm m}$ , where appropriate.

If  $U_m > (0.3 \times SDPA)$  then SDPA adjusted =  $\sqrt{(U_m^2 + SDPA^2)}$  and the reported value is suffixed with "a"

If  $U_m$  is less than (0.3 x SDPA) then SDPA adjusted = SDPA

- After statistical reduction, some results are excluded.
- 8 Ideally this will be your instrument group mean. If N<5 for instrument group, your method group Mean is selected as Mean for Comparison.
- Running Mean average of the last 10 performance indicators is used to monitor performance over time and concentration range.
- Target Score The closer a value is to 120, the better the performance.

$$TS = \log_{10} \left( 3.16 \times \frac{TDPA}{|\%Dev|} \right) \times 100$$

%Deviation from the Mean for Comparison -

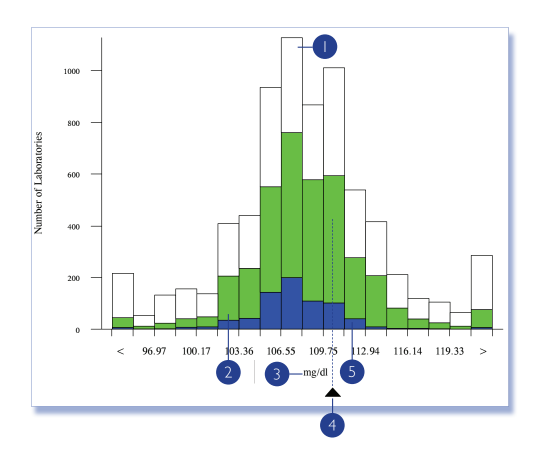
 $\text{\%Dev} = \frac{\text{Your Result - Mean for Comparison}}{\text{Mean for Comparison}} \times 100$ 

The closer the value is to zero, the better the performance.

- Biological Variation stated for information purposes only.
- Performance limit set for this parameter.
- Reference values quoted for information purposes, where applicable.

The Bar Graph is intended as a quick visualisation of how your lab's result compares to the method mean, instrument mean and all method mean.







200 laboratories reported values between 101.77 and 103.36 in your method group.

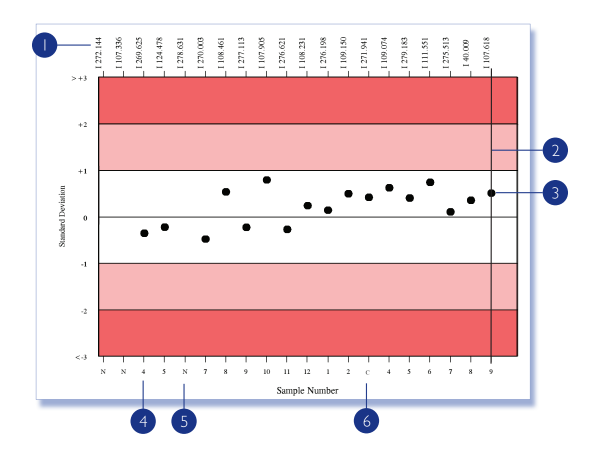
3 RIQAS reports show your unit of measurement.

4 Your result is indicated by the black triangle.

41 laboratories reported values between 111.35 and 112.94 in your instrument group.

# LEVEY-JENNINGS CHART

SDIs reflect laboratory performance in relation to fit-for-purpose SDPAs and are useful to monitor performance over time. Acceptable performance is SDI < 2.



- The Mean for Comparison for each sample is indicated at the top of the chart. This allows easy assessment of concentration related bias:
  - I: Instrument mean M: Method mean
  - A: All method mean
- This line indicates a change in registration details for this parameter.
- 3 Your SDI (Standard Deviation Index).

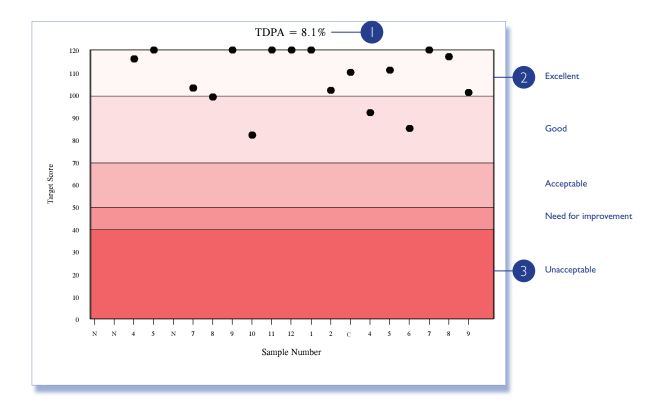
- 4 Sample number.
- N = No result returned from your laboratory.
  - C = Corrected results will be accepted for non-analytical errors.

    Corrected results will be accepted up to 4 weeks after the final submission deadline, on application, with evidence of analysis. Late results are only accepted if there has been a Randox error.

R = Incorrect results can be removed retrospectively on request.

# TARGET SCORE CHART

The Target Score (TS) allows you to assess your performance at a glance. The TS relates the %Deviation of your result from the Mean to a Target Deviation for Performance Assessment (TDPA). TDPAs are set to encourage participants to achieve and maintain acceptable performance. TDPAs are fit-for-purpose performance criteria which are set taking guidance from ISO/IEC17043, ISO13528 and IUPAC. Target Deviations for Performance Assessment are also used to calculate the Standard Deviation for Performance Assessment (SDPA).



This is the upper deviation limit of performance for this parameter. TDPAs are reviewed regularly and deemed fit for purpose by the RIQAS Advisory Panel.

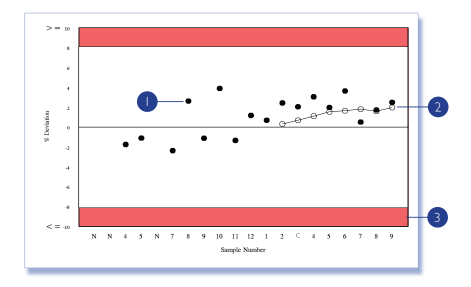
High scores ≥50 in the lighter shaded area represent acceptable, good or excellent performance.

Heavy shading for values 10 to 50 signifies poor performance.

# **%DEVIATION CHARTS**

The %Deviation by sample chart helps to identify trends and shifts in performance.

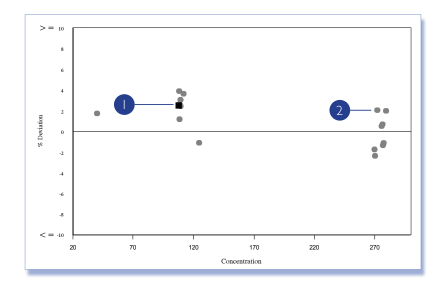
%Deviation = 
$$\frac{\text{Your Result - Consensus Mean}}{\text{Consensus Mean}} \times 100\%$$



- %Deviation from Mean for Comparison.
- Plot of Running Mean %Deviations (average of the last 10 %Deviations for the sample indicated).

Acceptable limits of performance. These are defaulted to RIQAS TDPAs but can be set to e.g. biological variation or regulatory requirement on request.

The %Deviation by concentration chart enables rapid assessment of concentration related biases. Biases at low or high concentrations can be easily determined.



Current sample indicated by square.

2

%Deviation at specific concentration.

# **MULTI-METHOD STAT SECTION**

This section provides an easy way of assessing the performance of other methods used to analyse the parameter in question.

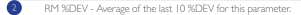
Method	N	Mean	CV%	$\mathbf{U}_{\mathbf{m}}$
Hexokinase	3524	108.234	2.9	0.07
Glucose oxidase	2673	108.474	5.5	0.14
Ortho Vitros MicroSlide Systems	251	105.036	2.3	0.19
Glucose dehydrogenase	63	107.832	3.5	0.59
GOD/02-Beckman method	37	106.425	2.5	0.55
Other Dry Chemistry	27	108.360	6.3	1.64
Agappe - GOD-PAP	21	109.727	3.7	1.11
Oxygen electrode	17	106.666	1.3	0.43
Vitros, DT60/DT60 II	3	105.595	3.9	2.99

# **SUMMARY PAGE**

Located at the back of the RIQAS Report, the Summary Page collates the key information, allowing participants to review the performance of all parameters at-a-glance.

	Mean for	Your							
Analyte	Comparison	Result	SDI	RMSDI	%DEV	RM%DEV	TS	RMTS	Performance
Albumin	2.120	2.230	1.00	0.37 —	5.2	2.0	72	107	
Alkaline Phosphatase	17.705	19.000	0.61	-0.27	7.3	-2.9	93	105	
ALT (GPT)	12.387	12.000	-0.33	-0.47	-3.1	-3.8	119	103	
Amylase, Total	20.454	22.000	0.72	-0.29	7.6	-2.5	86	103	_
AST (GOT)	11.976	11.000	-0.86	-0.03	-8.2	-0.4 —	78	100 —	3
Bicarbonate	8.203	6.900	-1.48	0.15	-15.9	1.5	54	98	
Bilirubin, Direct	0.251	0.380	2.57	2.64	51.3	47.2	31	29	<b>A</b> – 4
Bilirubin, Total	0.701	0.640	-0.91	-0.29	-8.8	-2.9	76	101	
Calcium	6.074	6.020	-0.19	-0.40	-0.9	-1.8	120	92	
Chloride	76.353	77.000	0.30	-0.28	0.8	-0.8	120	98	
Cholesterol	112.696	110.000	-0.55	0.05	2.4	0.2	97	115	
CK, Total	111.659	111.000	-0.08	0.35	-0.6	2.5	120	107	
Creatinine	0.607	0.620	0.27	0.06	2.1	0.5	120	117	
Glucose	36.429	36.000	-0.26	-0.84	-1.2	-3.7	120	82	
HDL-Cholesterol	98.836	102.000	0.21	-0.04	3.2	-0.4	120	113	
Iron	97.374	99.000	0.28	0.01	1.7	0.1	120	114	
Lactate		No Result		Too Few		Too Few	N/A	N/A	
LD (LDH)	85.894	87.000	0.11	-0.70	1.3	-6.3	120	89	
Magnesium	1.313	1.390	0.79	-0.07	5.8	-0.5	82	107	
Phosphate, Inorganic	1.451	1.540	1.02	0.02	6.1	0.1	71	112	
Potassium	1.770	1.840	1.10	-0.25	3.9	-0.7	67	99	
Protein, Total	3.850	3.830	-0.11	0.07	-0.5	0.3	120	114	
Sodium	112.537	114.000	0.58	-0.01	1.3	-0.0	95	104	
TIBC	133.143	133.000	-0.01	-0.01	-0.1	-0.1	120	117	
Trig Total	23.626	24.000	0.18	-0.09	1.6	-0.6	120	114	
Urea	5.872	5.000	-2.02	-0.57	<u>-14.9</u>	-4.0	41	95	<b>A</b>
Uric Acid (Urate)	3.135	3.100	-0.20	-0.44	-1.1	-2.4	120	107	
			ORM	ISDI -0.05	OR	M%DEV 0.8	OPM'	TS 102	
			OKW	13D1 -0.03	OK	VI WDE V U.O	OKM	13 102	





RMTS - Average of the last 10 Target Scores for this parameter.

SDI > 2

TS < 50

%DEV > acceptable limits set



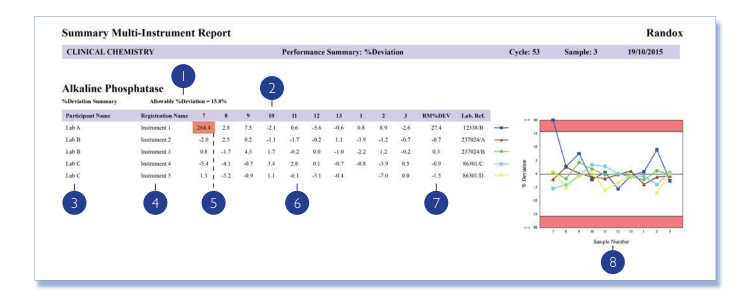
6 Overall RM%DEV = average RM%DEV for this sample distribution.

Overall RMTS = average RMTS for this sample distribution.

Red triangle appears when all performance indicators (SDI, %DEV and TS) exceed acceptable performance, i.e: when

# **MULTI-INSTRUMENT REPORT**

Register up to five instruments per programme at no extra cost. In addition to a standard report for each instrument, a multi-instrument report is also provided allowing comparitive performance assessment.



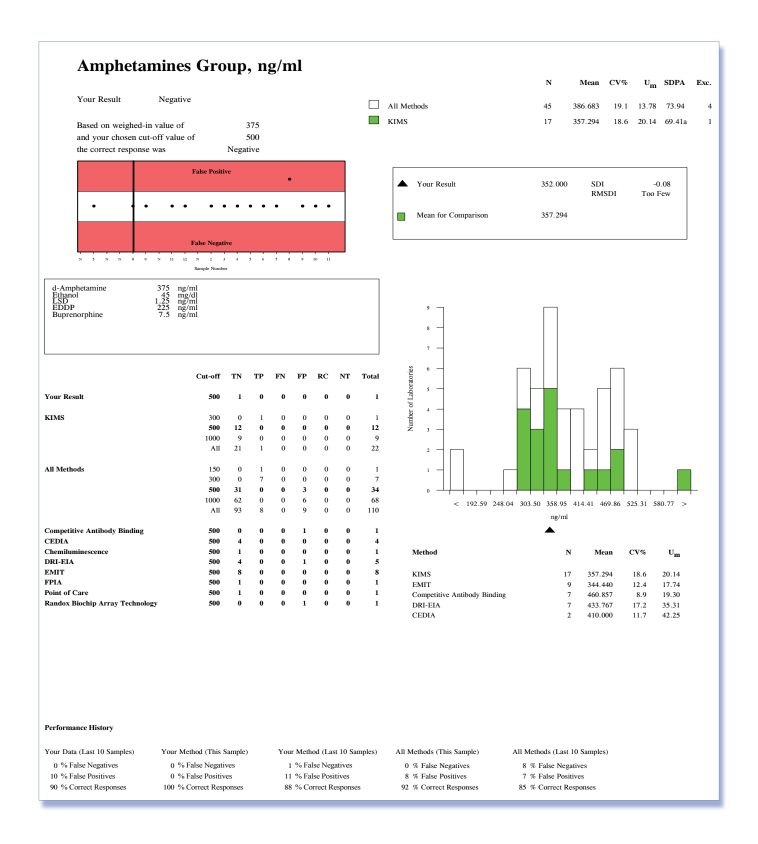


# URINE TOXICOLOGY REPORT

Laboratory performance is presented in both quantitative and qualitative screening formats, allowing for easy interpretation at-a-glance.

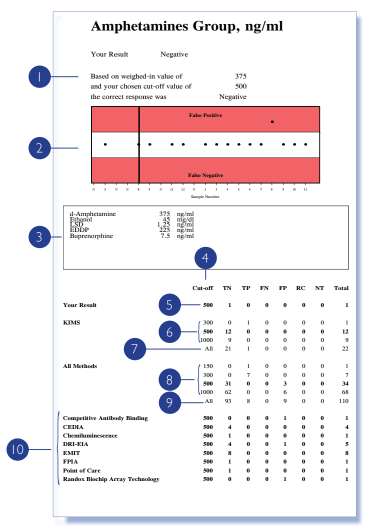
#### **Screening Section**

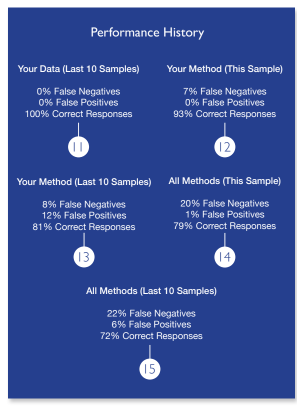
#### **Quantitative Section**



# URINE TOXICOLOGY REPORT SCREENING SECTION

#### Qualitative comparison of screening results available for each parameter.





- Text section shows the correct response for the lab based on a comparison between the weighed in value and the lab's cut off value.
- Screening Results: This chart is a quick visualisation of your performance over the last 20 samples. A result in the white section indicates a correct response. A result in the upper red section indicates a False Positive response, and a result in the lower red section indicates a False Negative response.
- 3 Comment section for RIQAS to provide your laboratory with additional relevant information regarding this sample, such as spiked metabolite concentration
- Screening result response categories. All abbreviations indicated at the bottom of the report page.

Key

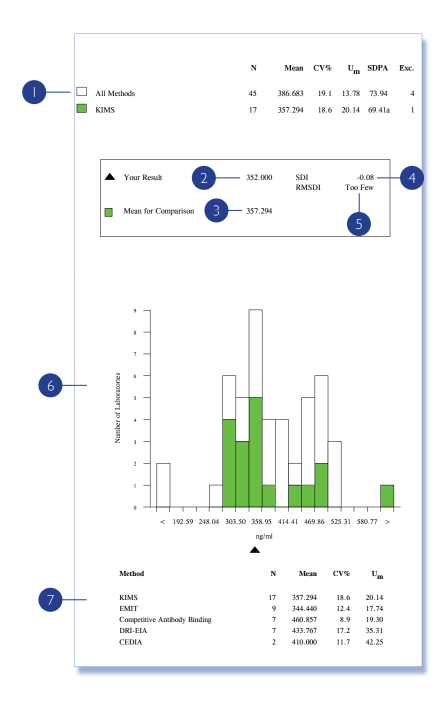
TN - true negative TP - true positive FN - false negative FP - false positive RC - sent for confirmation NT - not tested

- Screening Summary: Your screening result shown in the appropriate response category and your cut off for this sample.
- 6 Screening results for all cut-offs returned for this sample within your method group.

- 7 Total screening results over all your cut-offs for your laboratory's method.
- 8 Screening results for all cut-offs returned for this sample over all methods.
- Total screening results over all cut-offs for all methods.
- Screening results for other methods using same cut-off as your laboratory.
- Performance history for this parameter, based on previous 10 samples.
- Performance of your method over all cut-offs for this sample.
- Performance history of your method over all cut-offs, based on the previous 10 samples.
- Performance of all methods over all cut-offs for this sample.
- Performance history of all methods over all cut-offs, based on the previous 10 samples.

# URINE TOXICOLOGY REPORT QUANTITATIVE SECTION

#### Quantitative statistical comparison available for each parameter.



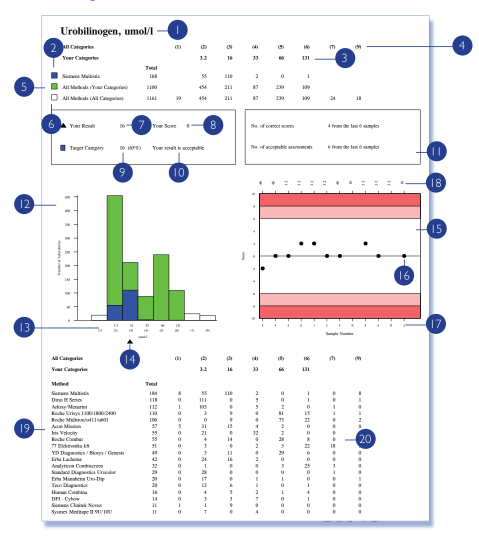


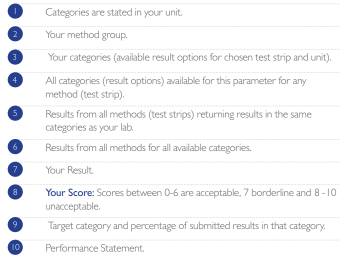
- Your Result.
- Your Mean for Comparison.
- Standard Deviation Index = (Your Result Mean for Comparison)
   SD of Mean for comparison
- Sunning mean SDI = average of last 10 SDIs for this parameter (If fewer than 10 results, "Too Few" is printed).
- Quantitative Results Histogram: This graph provides a quick visualisation of how your quantitative result falls into the overall picture for all methods and your method group.
- 7 All available method statistics for this sample.

# **URINALYSIS REPORT**

Your performance for each parameter is presented in a simple, convenient report.

#### Screening Results





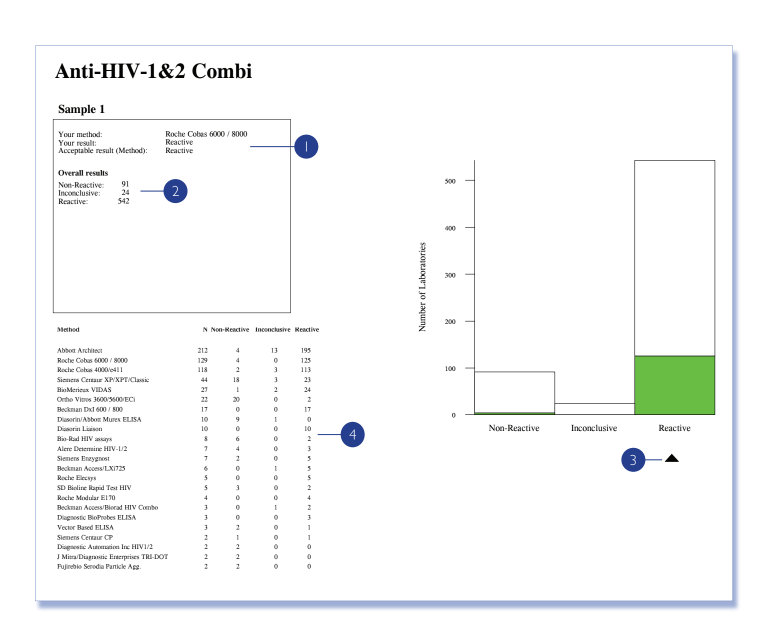
0	Comments Box: Provides number of correct scores and acceptable assessments for the last 6 samples.
12	<b>Categories Histogram:</b> A quick visualisation of how your lab's result falls into the overall picture for your categories.
13	Possible reporting categories for your method.
14	Your result is indicated by the black triangle.
15	Levey-Jennings Chart: Acceptable scores (0-6) have no shading, borderline scores (7) have light red shading, unacceptable scores (8-10) have dark red shading.
16	Score for each sample number.
17	Sample Number:
18	Target Categories.
19	All methods reported for this parameter.

Detailed summary of results: This table enables you to see how you

compare to all other results.

# SEROLOGY: SCREENING (QUALITATIVE) REPORT

Your performance for multiple samples is presented in a convenient single report per quarterly distribution.





Overall Summary shows the number of results for this parameter and sample which are non-reactive, inconclusive or reactive.

3	Your Result is shown as a black to other laboratories in groups:	triangle on the category chart compared
	All Methods	Your Method
4	Summary shows performance of the parameter:	of all the methods used to analyse

# SEROLOGY: SCREENING (QUANTITATIVE) REPORT

Your performance for multiple samples is presented in a convenient single report per quarterly distribution.





Your result is presented on the bar graph as a black triangle, showing how you compare to:

All Methods

Your Method

Multi Method Statistics section provides an easy way of assessing the performance of the methods used to analyse the parameter.

4 Levey-Jennings chart - Your SDIs for previous 20 samples.

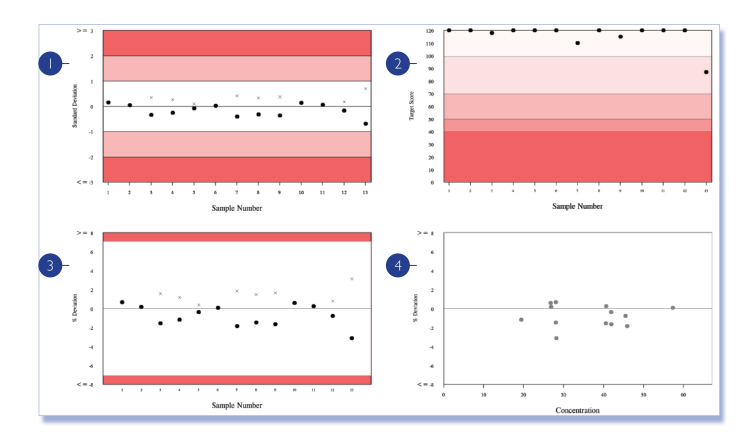
# QUANTITATIVE (END-OF-CYCLE REPORT)

The End-of-Cycle Report is sent to labs receiving standard reports at the end of each cycle and provides a complete summary of statistics. Results can also be compared to the previous cycle.

#### Albumin, g/l Method: Bromocresol Purple Instrument: Siemens/Dade Dimension RxL/Max/Xpand Reagent: Siemens/Dade Behring RIQAS TDPA: 7.1% **Biological Variation:** Mean for Comparison CV% SDPA SDI TS Sample Result Unit N Um % Deviation 28.200 0.10 28.013 2.4 1.26 0.15 120 0.67 2 26.900 g/l 87 26.853 2.7 0.101.21 0.04 120 0.17 3 39.900 g/l 71 40.531 2.5 0.15 1.82 -0.35 118 -1.56 4 19.200 81 19.429 2.5 0.07 0.87 -0.26 120 -1.18 g/l 41.700 67 41.859 2.0 0.13 1.88 -0.08 120 -0.38 g/l 6 57.300 g/l 57.257 2.7 0.21 2.58 0.02 120 0.08 45.000 2.1 2.06 g/l 72 45.850 0.14 -0.41 110 -1.85 8 27.600 g/1 87 28.013 2.5 0.09 1.26 -0.33 120 -1.47 g/l 41.200 70 41.891 2.2 0.14 1.88 -0.37 115 -1.65 10 0.59 26.900 26.742 3.3 0.12 1.20 120 g/l 83 0.13 40.700 0.14 11 g/l 71 40.601 2.2 1.83 0.05 120 0.24 12 45.100 g/l 80 Ι 45.456 2.2 0.14 2.04 -0.17120 -0.7828.179 13 27.300 g/1 63 0.09 1.27 -0.6987 -3.12Cycle 45 Cycle 46 Cycle Average SDI -0.23-0.18 Cycle Average TS 110 116 Cycle Average %DEV -1.05 -0.79 0.36 0.24 Cycle Average Absolute SDI Cycle Average Absolute %DEV 1.06 1.63 110 50 Sample Number Sample Number Sample Number Concentration

# CHART SECTION (END-OF-CYCLE REPORT)

Your results for current cycle shown in various diagrams.



•	Levey-Jennings chart	Shows your SDIs for a full cycle.
		Shows SDI (positive and negative)
		× Shows absolute SDI
2	Target Score chart	Shows your Target Scores for a full cycle.
3	%Deviation by sample chart	Shows your %Deviations for a full cycle.
		Acceptable limits equal to TDPA unless alternative limits are registered by the lab.
		Shows %Deviation (positive and negative)
		x Shows absolute %Deviation
4	%Deviation by Concentration chart	Shows your results for a full cycle.

# TEXT SECTION (END-OF-CYCLE REPORT)

The text section summarises the statistical information for all samples.

Albumin, g/l

Method: Bromocresol Purple

Instrument: Siemens/Dade Dimension RxL/Max/Xpand

Reagent: Siemens/Dade Behring

RIQAS TDPA: 7.1% Biological Variation: 3.99

Your assay details at the end of the cycle.

The RIQAS TDPA and biological variation for the parameter are shown if available.



Sample	Result	Unit	N	Mean	SDPA	Um	CV%	SDI	TS	% Deviation
1	28.200	g/l	68	I 28.013	1.26	0.10	2.4	0.15	120	0.7
2	26.900	g/l	87	I 26.853	1.21	0.10	2.7	0.04	120	0.2
3	39.900	g/l	71	M 40.531	1.82	0.15	2.5	-0.36	116	-1.5
4	19.200	g/l	81	I 19.429	0.87	0.07	2.5	-0.27	120	-1.2
5	41.700	g/l	67	I 41.942	1.88	0.13	2.0	-0.09	120	-0.4
6	57.300	g/l	87	I 57.257	2.58	0.21	2.7	0.02	120	0.1
7	45.000	g/l	72	I 45.850	2.06	0.14	2.1	-0.43	108	-1.8
8	27.600	g/l	87	I 28.011	1.26	0.09	2.5	-0.34	118	-1.5
9	41.200	g/l	70	I 41.823	1.88	0.14	2.2	-0.38	113	-1.6
10	26.900	g/l	83	I 26.742	1.20	0.12	3.3	0.14	120	0.6
11	40.700	g/l	71	I 40.601	1.83	0.13	2.2	0.06	120	0.2
12	45.100	g/l	80	I 45.119	2.05	0.14	2.2	-0.18	120	-0.8
13	27.300	g/l	63	I 28.454	1.27	0.09	2.0	-0.72	86	-3.1

Summary of your results and statistics are shown, including Mean for Comparison, SDPA, %CV, U<sub>m</sub>, SDI, Target Score, %Deviation.

		Cycle 45	Cycle 46
	Cycle Average SDI	-0.23	-0.18
15	Cycle Average TS	110	116
	Cycle Average %DEV	-1.05	-0.79
16	Cycle Average Absolute SDI	0.36	0.24
16	Cycle Average Absolute %DEV	1.63	1.06

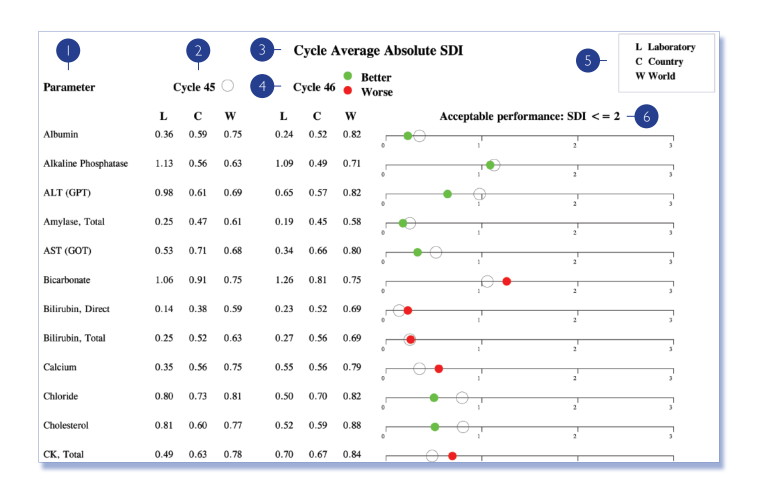
Table containing a summary of your performance for previous cycle and current cycle, including Average Absolute SDIs and %Deviations.

# TEXT SECTION (END-OF-CYCLE REPORT)

0	Report presented in your chosen unit	Cycle average of your performation of the Cycle average of your performation of the Cycle average of your performation of the Cycle average of your performance of	ormance indicators – Standard Deviation Deviation.	
2	Your assay details as of the last sample		(Sum of SDIs returned for the completed cycle)	
3	RIQAS TDPA and Biological variation	Cycle Average SDI =	(Number of samples returned in cycle)	
4	Sample number	Cycle Average	(Sum of your Target Scores returned for the completed cycle)	
5	Your results for each sample	Target Score =		
			(Number of samples returned in cycle)	
6	Unit your result was returned in		(Sum of your %Deviations returned	
7	umber of results used for statistical analysis	Cycle Average	for the completed cycle)	
		%Deviation =	(Number of samples returned in cycle)	
8	Mean for Comparison		(· · · · · · · · · · · · · · · · · · ·	
9	SDPA = Standard Deviation for performance assessment		values of your SDI and %Deviation.  far a value is from zero regardless of the	
10	Uncertainty of Mean for Comparison	Sign. This is an indication of	the magnitude of accuracy.	
		Cools Assessed	(Sum of your Absolute SDIs returned for the completed cycle)	
w	Coefficient of Variation (%)	Cycle Average Absolute SDI =	(Number of samples returned in cycle)	
12	Your Standard Deviation Index			
13	Your Target Score	Cycle Average Absolute %Deviation =	(Sum of your Absolute %Deviations returned for the completed cycle)	
14	Your %Deviation		(Number of samples returned in cycle)	

# CURRENT & PREVIOUS CYCLE ABSOLUTE SDIs (END-OF-CYCLE REPORT)

Based on the cycle average absolute SDI, this chart provides a visual representation of your laboratory's performance compared to the previous cycle.



Parameter list	List of all parameters registered.
2 Results for previous cycle	Indicated by open circle on the chart.
Report title - Cycle Average Absolute SDI	This shows your performance this cycle compared to the previous cycle.
4 Results for current cycle	Indicated by a closed circle on the chart.
5 Legend	Cycle Average Absolute SDIs are shown for:
	<ul> <li>L Your results throughout the cycle</li> <li>C All labs within your own country</li> <li>W All labs Worldwide</li> </ul>
6 Graphical representation of Absolute SDIs	Acceptable performance is $\leq 2$ .
	If Absolute SDI for current cycle is less than that for the previous cycle, this is indicated by a green circle.
	If Absolute SDI for current cycle is greater than that for the previous cycle,

The closer the circle is to zero, the better the performance.

this is indicated by a red circle.

# CERTIFICATE OF PERFORMANCE (END-OF-CYCLE REPORT)

An End-of-Cycle report will be issued for all registrations. However, the Certificate of Performance will only be available for parameters where results for at least 50% of samples in the cycle have been returned. Labs joining after the beginning of the cycle will only receive the Certificate of Performance if they meet this criterion. Any parameters not included on the Certificate of Acceptable Performance will be listed on the Notification of Unacceptable Performance.

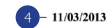


# CERTIFICATE OF ACCEPTABLE PERFORMANCE

RIQAS Department
Randox Laboratories
CRUMLIN
COUNTY ANTRIM
BT29 4QY
UNITED KINGDOM







This is to certify that the above participant took part in a cycle of external quality assessment and achieved an acceptable level of performance (Cycle Average Absolute SDI <=2) for the following parameters:

5	- Cycle Average Absolute SDI
Albumin - Bromocresol Purple - Siemens/Dade Dimension RxL/Max/Xpand	0.50
Alkaline Phosphatase - Dade Dimension, AMP buffer - Siemens/Dade Dimension RxL/Max/Xpa	and 1.22
ALT (GPT) - Tris buffer with P5P - Siemens/Dade Dimension RxL/Max/Xpand	0.53
A TOTAL DELIVER A STATE OF THE	0.04

Amylase, Total - Dade Behring 2-chloro-pNPG3 - Siemens/Dade Dimension RxL/Max/Xpand 0.34 AST (GOT) - Tris buffer with P5P - Siemens/Dade Dimension RxL/Max/Xpand 0.55 Bicarbonate - Enzymatic - Siemens/Dade Dimension RxL/Max/Xpand 1.08 Bilirubin, Direct - Diazo with Sulphanilic Acid - Siemens/Dade Dimension RxL/Max/Xpand 0.19 Bilirubin, Total - Diazo with Sulphanilic Acid - Siemens/Dade Dimension RxL/Max/Xpand 0.26 Calcium - Cresolphthalein complexone - Siemens/Dade Dimension RxL/Max/Xpand 0.49 Chloride - ISE, indirect - Siemens/Dade Dimension RxL/Max/Xpand 0.70 Cholesterol - Dimension-Dade Behring reagents - Siemens/Dade Dimension RxL/Max/Xpand 0.54 CK, Total - CK-NAC (IFCC) - Siemens/Dade Dimension RxL/Max/Xpand 0.26 Creatinine - Alkaline picrate no deprot. - Siemens/Dade Dimension RxL/Max/Xpand 0.44 GGT - Gamma glut'3-carb'4-nitro (IFCC) - Siemens/Dade Dimension RxL/Max/Xpand 0.25 Glucose - Hexokinase - Siemens/Dade Dimension RxL/Max/Xpand 0.70

•	Full registration address	Your full registration address details.
2	Your lab reference number	Used to identify each lab.
3	Programme / cycle number	Programme and current, completed cycle number.
4	Date	Date End-of-Cycle report is issued.
5	Parameters	List of parameters including the assay details for which cycle absolute SDI is $\leq 2$ .
6	Average Absolute SDI	Your Cycle Average Absolute SDI.

# MONITORING EQA PERFORMANCE

Each EQA report should be evaluated and any poor performance investigated. A step by step approach should be adopted consisting of the following three steps:

#### 1. Investigate the source of the problem

In order to identify the source of the problem, it is useful to be aware of the most common causes of poor EQA performance. Errors can occur at any stage of the testing process; however, EQA is most concerned with detecting analytical errors i.e. errors that occur during the analysis of the sample.

Most analytical errors can be easily divided into three main areas; clerical errors, systematic errors and random errors. Systematic errors result in inaccurate results that consistently show a positive or negative bias. Random errors, on the other hand, affect precision and result in fluctuations in either direction.

It may be possible that, after extensive investigations, the root cause of the poor performance cannot be established. Poor performance for a single sample could be attributed to random error. If poor performance has been noted for several samples, a systematic error is the most likely cause and the analytical process should be reviewed.



#### Clerical errors

- Transcription errors
- Incorrect units used
- Incorrect sample tested
- Incorrect method classification
- Calculation/conversion error

#### Systematic errors

- Sample/Reagent prep/handling
- Reagent/calibrator/standardisation change
- Instrument/reagent/calibrator fault
- Inexperienced operators
- Reagent deterioration
- · Inappropriate method

#### Random errors

- · Bubbles in reagent
- Bubbles in reagent/sample pipette
- Temperature fluctuations
- Poor pipetting technique
- Poor operator technique

The flowchart (page 29) is designed to help you investigate any apparent poor performance.

#### 2. Implement corrective actions

Some errors can be readily recognised as simple clerical errors and easily corrected. If there is evidence of systematic or random error however more detailed corrective actions must be taken.

#### Systematic Error

In the event of a systematic error, the following suggested actions may help to resolve the problem:

- Perform instrument maintenance
- Recalibrate instrument
- Review reagent/sample storage
- Check pipettes

- Prepare fresh reagents & re-run sample
- · Perform staff training

#### Random Error

If all possible causes have been excluded, a single unacceptable result is most likely due to random error. Re-run the sample; if the result of repeat analysis is acceptable then corrective action is not required. If the issue persists, investigate possible sources of systematic error.

#### 3. Check the effectiveness of corrective actions

The effectiveness or impact of any corrective actions taken can be assessed by continuing to monitor analytical performance over time.

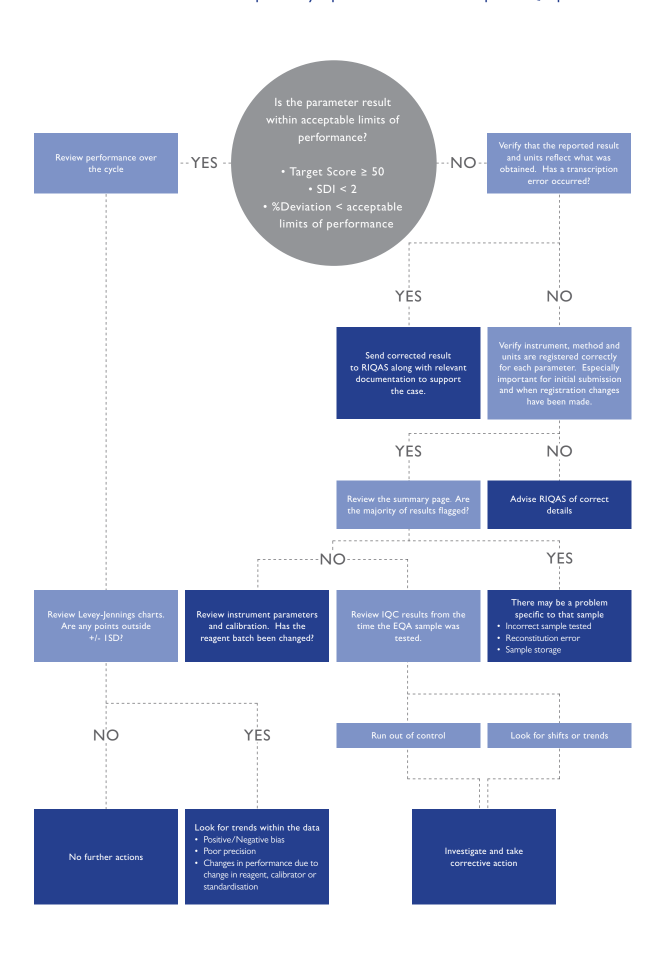
# MONITORING EQA PERFORMANCE

A checklist similar to the one below is extremely useful when investigating poor EQA performance and may help you to determine the root cause of the problem and initiate corrective actions.

Laboratory:				
Cycle Number:	Sample Number:			
Analysis Date:	Analyte:			
Mean for Comparison:		Lab Result: SDI: %Dev:		
I. Specimen Handling		e. Error due to imprecision; check IQC in terms of		
a. Samples received in good condition		%Deviation compared to deviation observed in EQA		
b. Samples stored/prepared appropriately	N	f. IQC target correctly assigned		
c. Integrity of the sample is acceptable	N			
		5. Calibration		
2. Clerical		a. Date of last calibration		
a. Correct result entered		b. Calibration frequency acceptable	<b>M M</b>	
b. Correct use of decimal point and units		c. Last calibration acceptable		
c. Calculations, if any, performed correctly				
(even if automated)		6. Instrument		
d. Conversion factors applied to results before submission		a. Daily maintenance performed on date of sample analysis		
		b. Special maintenance performed prior to sample analysis		
3. Registration and Mean for Comparison		c. Instrument operated correctly		
a. Registered in the correct method/instrument group		d. Operator fully trained		
b. Changed method or instrument without advising RIQAS		7. D		
c. Peer Group changed due to the number of participants		7. Reagents		
returning results e.g. from method to instrument		a. Reagents prepared and stored correctly		
d. An obvious bias between method and instrument means		b. Reagents within open vial stability		
(check histogram and stats sections)		8. EQA sample		
4. Internal Quality Control		a. Initial value		
a. %Deviation of IQC (at similar conc to that of EQA) on		b. Re-run value		
sample analysis date acceptable		c. Issue observed in previous EQA samples at a similar		
b. Shift in IQC in the periods just before and after EQA		concentration (check %Deviation by concentration and		
sample analysis		Levey Jennings charts)		
c. Trends in IQC in the periods before and after EQA		d. All parameters affected (to the same extent) - possible		
sample analysis		reconstitution error (check %Deviation on summary pages)		
d. Random IQC variation on sample analysis date				
Conclusion:		Remedial Action:		
			•••••	
Lab Manager: Date:		Lab Director: Date:		

# MONITORING EQA PERFORMANCE

The flow chart below can be used to help identify a possible root cause for poor EQA performance.



Lactate

# Ammonia/Ethanol Programme+ With target scoring



RQ9164 (2 ml)

2 Parameters Samples every month,  $1 \times 12$  month cycle, 12 month subscription

Ethanol

### Anti-TSH Receptor Programme+ With target scoring



I Parameter

Samples every month,  $1 \times 12$  month cycle, 12 month subscription

Anti-TSH Receptor (TRAb)

### Blood Gas Programme With target scoring



RQ9134 (1.8 ml) RQ9134/A (1.8 ml) First registered instrument Subsequent instruments 10 Parameters 10 Parameters Samples every month, 1 x 12 month cycle, 12 month subscription

pCO, CO<sub>2</sub>(Total)\* pН Ca++ Na+ рО, Glucose

#### BNP Programme+ With target scoring



RQ9165 (1 ml)

I Parameter

Samples every month, 1 x 12 month cycle, 12 month subscription

#### Cardiac Programme With target scoring



RQ9127/a (1 ml) RQ9127/b (1 ml) 2 Parameters only (choose from 7) Full 7 Parameters Samples every 2 weeks, 2  $\times$  6 monthly cycles, 12 month subscription

CK, Total CK-MB (Mass) Myoglobin Troponin T CK-MB (Activity) Homocysteine Troponin I

#### Cerebrospinal Fluid Programme+ With target scoring



RQ9168 (3 ml) 7 Parameters

Samples every month,  $1 \times 12$  month cycle, 12 month subscription

Albumin Glucose Lactate Sodium Protein (Total) Chloride lgG

#### Coagulation Programme With target scoring



RQ9135/b (1 ml) RQ9135/a (1 ml) 5 Selected parameters only (aPTT, PT, TT, Fibrinogen, Antithrombin III) Samples every month,  $1\times12$  month cycle, 12 month subscription

Plasminogen Factor VII  $PT \ (including \ INR)$ Protein C Factor VIII Factor IX Protein S Factor II Factor X

Fibrinogen Antithrombin III Factor XI Factor V





PURPLE = The only parameters available on RQ9135/a

Factor XII

D-dimer\*

\* = Pilot study ongoing

# **RIQAS PROGRAMMES**

# CO-Oximetry Programme+



Samples every month, 1 x 12 month cycle, 12 month subscription Carboxyhaemoglobin (COHb / HbCO) Methaemoglobin (MetHb)

Oxygen Saturation (sO2 / Vol O2) Oxyhaemoglobin (O2Hb / HbO2)

Total Haemoglobin (tHb)

# CYFRA 21-1 Programme+



Oxygen Content (O2CT)

RQ9175 (1 ml) I Parameter

Deoxyhaemoglobin (HHb)

Samples every month, 1 x 12 month cycle, 12 month subscription

CYFRA 21-1 (Cytokeratin 19)

# ESR Programme+



RQ9163 (4.5 ml)

I Parameter

RQ9112/a (5 ml)

2 samples per quarterly distribution, 1  $\times$  12 month cycle, 12 month subcription

ESR (Erythrocyte Sedimentation Rate)

# General Clinical Chemistry Programme With target scoring

RQ9112/b (5 ml)



10 Parameters only 17 Parameters only Full 52 Parameters Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription, reference method values ACE (Angiotensin Converting Enzyme) Calcium Acid Phosphatase (Prostatic) Calcium (Ionised) HDL-Cholesterol Sodium Acid Phosphatase (Total) Chloride TIBC T<sub>3</sub> (Free) T<sub>3</sub> (Total) T<sub>4</sub> (Free) T<sub>4</sub> (Total) Adjusted Calcium Cholesterol Lactate Albumin Cholinesterase LD (LDH) Alkaline Phosphatase CK, Total (CPK) Lipase ALT (ALAT) Copper Lithium Amylase (Pancreatic) Creatinine Magnesium Triglycerides Amylase (Total) D-3-Hydroxybutyrate NEFA TSH AST (ASAT) **EGFR** Non-HDL Cholesterol UIBC Osmolality Bicarbonate Fructosamine Urea Phosphate (Inorganic) Uric Acid Bile Acids γGT Bilirubin (Direct) . GLDH Potassium Zinc Bilirubin (Total) Glucose Protein (Total)

# Glycated Haemoglobin Programme (HbAIc) With target scoring



RQ9129 (0.5ml)

2 Parameters

Samples every month,  $1 \times 12$  month cycle, 12 month subscription

HbAlc Total Haemoglobin

# Haematology Programme With target scoring



RQ9118 (2 ml) 11 Parameters

Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription

Haemoglobin (Hb) Mean Cell Haemoglobin (MCH)

Mean Cell Haemoglobin Concentration (MCHC) Mean Cell Volume (MCV) Mean Platelet Volume (MPV)

Platelets (PLT) Plateletcrit (PCT) Red Blood Cell Count (RBC) Red Cell Distribution Width (RDW) Total White Blood Cell Count (WBC)





# Human Urine Programme With target scoring



# RQ9115 (10 ml) 25 Parameters Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription

Protein (Total) Creatinine Normetanephrine Albumin/Microalbumin Magnesium Sodium Dopamine Osmolality Amylase **Epinephrine** Urea Calcium Glucose Oxalate Uric Acid Chloride Metanephrine Phosphate (Inorganic) VMA 5-HIAA Copper Norepinephrine Potassium Cortisol

### Immunoassay Programme With target scoring



RQ9125/a (5 ml) 4 Parameters only (choose from 55) Samples every two weeks, 2 x 6 monthly cy Samples every month, 1 x 12 month cycle,	RQ9125/b (5 ml) 13 Parameters only (choose from 55) cles, 12 month subscription (RQ9125/a, RQ917 12 month subscription (RQ9130)	RQ9125/c (5 ml) Full 55 Parameters 25/b, RQ9125/c)	RQ9130 (5 ml) Full 55 Parameters
ACTH	DHEA Unconjugated	17-OH-Progesterone	T <sub>4</sub> (Free)
AFP	Digoxin	Paracetamol	T <sub>4</sub> (Total)
Aldosterone	Estriol Total*	Phenobarbital	Testosterone (Free)*
Amikacin	Ethosuximide*	Phenytoin	Testosterone (Total)

Theophylline Androstenedione Ferritin Primidone\* β-2-Microglobulin Progesterone Folate Thyroglobulin CA125 FSH Prolactin Tobramycin\* CA15-3 Gentamicin PSA (Free) TSH CA19-9 PSA (Total) Valproic Acid Carbamazepine hCG PTH Vancomycin Salicylate CEA ΙgΕ Vitamin B12 1-25-(OH)<sub>2</sub>-Vitamin D\* Cortisol Insulin SHBG T<sub>3</sub> (Free) T<sub>3</sub> (Total) C-Peptide 25-OH-Vitamin D LH DHEA-Sulphate Oestradiol

# Immunoassay Speciality I Programme+ With target scoring



RQ9141 (2 ml)
10 Parameters
Samples every month, 1 x 12 month cycle, 12 month subscription

 I-25-(OH)<sub>2</sub>-Vitamin D\*
 Anti-TG
 Osteocalcin
 Insulin

 25-OH-Vitamin D
 Anti-TPO
 Procalcitonin
 FTH

 C-Peptide
 IGF-I
 PTH

# Immunoassay Speciality 2 Programme+ With target scoring



# RQ9142 (1 ml) 5 Parameters Samples every month, 1 $\times$ 12 month cycle, 12 month subscription

Calcitonin Procalcitonin Plasma Renin Activity Renin (Direct Concentration)
Gastrin

# Immunosuppressant Programme+



RQ9159 (2 ml) 4 Parameters Samples every month,  $1\times12$  month cycle, 12 month subscription, reference method values

Ciclosporin Everolimus Sirolimus Tacrolimus

# Lipid Programme With target scoring



RQ9126/a (3 ml)  RQ9126/b (3 ml)  Parameters only (choose from 7)  Full 7 Parameters  Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription	1 0			
3 Parameters only (choose from 7) Full 7 Parameters	RO9126/a (3 ml)	RO9126/b (3 ml)		
Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription	3 Parameters only (choose from 7)	Full / Parameters		
	Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription			

Apolipoprotein A ICholesterol (Total)LDL-CholesterolTriglyceridesApolipoprotein BHDL-CholesterolLipoprotein (a)





PURPLE = The only parameters available on RQ9135/a

+ = Not accredited

\* = Pilot study ongoing

# **RIQAS PROGRAMMES**

# Liquid Cardiac Programme With target scoring



#### RQ9136 (3 ml)

9 Parameters

Samples every month, 1 x 12 month cycle, 12 month subscription

CK-MR Mass Homocysteine Myoglobin Troponin I NT proBNP D-dimer hsCRP Troponin T Digoxin

#### Maternal Screening Programme With target scoring



#### RQ9137 (1 ml)

6 Parameters

Samples every month,  $1 \times 12$  month cycle, 12 month subscription

Total hCG PAPP-A Unconjugated Oestriol free β-hCG Inhibin A

# Serology (EBV) Programme+



#### RQ9153 (1 ml)

3 Parameters

3 samples per quarterly distribution, 1 x 12 month cycle, 12 month subscription, Quantitative and Qualitative results

Anti-EBV VCA IgG Anti-EBNA IgG Anti-EBV VCA IgM

# Serology (HIV-Hepatitis) Programme+



#### RQ9151 (1.8 ml)

10 Parameters

 $5 \ samples \ per \ quarterly \ distribution, \ 1 \times 12 \ month \ cycle, \ 12 \ month \ subscription, \ Quantitative \ and \ Qualitative \ results$ 

Anti-HIV-I Anti-HCV Anti-HTLV-II HBsAg Anti-HIV-2 Anti-HBc Anti-HTLV-1&2 Combined Anti-HIV-1&2 Combined Anti-HTLV-I Anti-CMV

# Serology (Syphilis) Programme+



I Parameter

3 samples per quarterly distribution, 1 x 12 month cycle, 12 month subscription, Quantitative and Qualitative results

Syphilis (Methods available include immunoassay RPR, VDRL and TPHA)

# Serology (ToRCH) Programme+



#### RQ9152 (1 ml)

12 Parameters

5 samples per quarterly distribution, 1 x 12 month cycle, 12 month subscription, Quantitative and Qualitative results

Anti-Rubella IgM Anti-HSV I IgM Anti-Toxoplasma IgG Anti-HSV2 lgG Anti-Toxoplasma IgM Anti-CMV lgG Anti-HSV 2 IgM Anti-Rubella IgG Anti-CMV lgM Anti-HSV-1&2 lgG Combined Anti-HSV I + 2 IgM Combined

# Specific Proteins Programme With target scoring



RQ9114 (3 ml) 26 Parameters	RQ9160 (2 ml)	RQ9161 (1 ml)		
Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription				
AFP Albumin $\alpha$ -1-Acid glycoprotein $\alpha$ -1-Artitrypsin $\alpha$ -2-Macroglobulin Anti Streptolysin O	β-2-Microglobulin Ceruloplasmin Complement C <sub>3</sub> Complement C <sub>4</sub> C-Reactive Protein Ferritin	IgA IgE IgG IgM Kappa Light Chain (Free) Kappa Light Chain (Total)	Lambda Light Chain (Total) Prealbumin (Transthyretin) Retinol Binding Protein Rheumatoid Factor Transferrin	
Antithrombin III	Haptoglobin	Lambda Light Chain (Free)		





PURPLE = The only parameters available on RQ9135/a

+ = Not accredited

\* = Pilot study ongoing

## Sweat Testing Programme+



RQ9173 (2 ml)

2 Parameters  $^{'}$  Samples every month, 1 x 12 month cycle, 12 month subscription

Chloride

Conductivity

### Therapeutic Drugs Programme With target scoring



RQ9111 (5 ml) 18 Parameters

Samples every 2 weeks, 2 x 6 monthly cycles, 12 month subscription, Weighed-in values

Amikacin Ethosuximide Caffeine Gentamicin Carbamazenine Lithium Ciclosporin Methotrexate Digoxin

Paracetamol (Acetaminophen)

Phenobarbital Phenytoin Primidone Salicylic Acid Theophylline

Tobramycin Valproic Acid Vancomycin

### Trace Elements In Blood Programme+



RQ9172 (3 ml)

Samples every month, 1 x 12 month cycle, 12 month subscription

Lead Magnesium lodine

Manganese Selenium

7inc

7inc

### Trace Elements In Serum Programme+



RQ9170 (3 ml) 10 Parameters

Samples every month,  $1 \times 12$  month cycle, 12 month subscription

Aluminium Copper Manganese Chromium Iodine Nickel Cobalt Lead Selenium

### Trace Elements In Urine Programme+



RQ9171 (3 ml)

Samples every month,  $1 \times 12$  month cycle, 12 month subscription

Nickel Copper Magnesium Manganese Thallium Chromium lodine Cobalt Molybdenum

### Urinalysis Programme+ With scoring



RQ9138 (12 ml)

14 Parameters

Samples every 2 months, 1 x 12 month cycle, 12 month subscription

Galactose Specific Gravity Albumin Leukocytes Bilirubin Glucose Nitrite . Urobilinogen Blood Creatinine Ketones Protein

### Urine Toxicology Programme+



RQ9139 (5 ml) 20 Parameters

Samples every month, 1 x 12 month cycle, 12 month subscription

Benzoylecgonine Buprenorphine Cannabinoids (THC) Cotinine Creatinine d-Amphetamine

d-Methamphetamine FDDP Ethanol Free Morphine Lorazepam LSD

MDMA Methadone Nortriptyline Norpropoxyphene Oxazepam Phencyclidine

Phenobarbital Secobarbital

= Liquid ready-to-use samples



PURPLE = The only parameters available on RQ9135/a

\* = Pilot study ongoing

+ = No	t accredited	+	+				+					emistry					llity 1 +	llity 2 +
* = Pilo	t study ongoing	Ammonia / Ethanol +	Anti-TSH Receptor +	ias			Cerebrospinal Fluid +	tion	CO-Oximetry +	21-1 +		General Clinical Chemistry		ology	Urine	assay	Immunoassay Speciality I +	Immunoassay Speciality 2 +
PURPLE	E = The only parameters available on RQ9135/a	Ammon	Anti-TSI	Blood Gas	BNP +	Cardiac	Cerebro	Coagulation	CO-OX	CYFRA 21-1 +	ESR +	General	HbAIc	Haematology	Human Urine	Immunoassay	Immuno	Immuno
#	I-25-(OH) <sub>2</sub> -Vitamin D*															Χ	Χ	
	17-OH-Progesterone															X		
	25-OH-Vitamin D															X	X	
	5-HIAA														X			
Α	lpha-I-Acid Glycoprotein																	
	α-I-Antitryspin																	
	α-2-Macroglobulin																	
	ACE (Angiotensin Converting Enzyme)											X						
	Acid Phosphatase (Prostatic)											X						
	Acid Phosphatase (Total)											X						
	ACR														X			
	ACTH															X		
	Adjusted Calcium*											X						
	AFP															X		
	Albumin						X					X			X			
	Aldosterone															X		
	Alkaline Phosphatase											X						
	ALT (ALAT)											X						
	Aluminium																	
	Amikacin															X		
	Ammonia	X																
	Amylase (Pancreatic)											X						
	Amylase (Total)											X			X			
	Androstenedione															X		
	Anti Streptolysin O (ASO)																	
	Anti-CMV																	
	Anti-CMV IgG																	
	Anti-CMV IgM																	
	Anti-EBNA IgG																	
	Anti-EBV VCA IgG																	
	Anti-EBV VCA IgM																	
	Anti-HBc																	
	Anti-HCV																	
	Anti-HIV-I																	
	Anti-HIV-I & 2 Combined																	
	Anti-HIV-2																	
	Anti-HSV- I & 2 IgG Combined																	
	Anti-HSV- I & 2 IgM Combined																	
	Anti-HSVI IgG																	
	Anti-HSVI IgM																	
	Anti-HSV2 IgG																	
	<del></del>																	

Immunosuppressant +		ardiac	Maternal Screening	Serology (EBV) +	Serology (HIV / Hepatitis) +	Serology (Syphilis) +	Serology (ToRCH) +	Proteins	esting +	Therapeutic Drug	Trace Elements in Blood +	Trace Elements in Serum +	Trace Elements in Urine +	+ s	Urine Toxicology +	+ = Not accredited  * = Pilot study ongoing	
Immunos	Lipid	Liquid Cardiac	Materna	Serology	Serology	Serology	Serology	Specific Proteins	Sweat Testing +	Therape	Trace Ele	Trace Eld	Trace Ele	Urinalysis +	Urine To	PURPLE = The only parameters available on RQ	9135/a
																I-25-(OH) <sub>2</sub> -Vitamin D*	#
																17-OH-Progesterone	
																25-OH-Vitamin D	
																5-HIAA	
								X								lpha-I-Acid Glycoprotein	Α
								X								lpha-I-Antitryspin	
								X								lpha-2-Macroglobulin	
																ACE (Angiotensin Converting Enzyme)	
																Acid Phosphatase (Prostatic)	
																Acid Phosphatase (Total)	
																ACR	
																ACTH	
																Adjusted Calcium*	
			X					X								AFP	
								X						X		Albumin	
																Aldosterone	
																Alkaline Phosphatase	
																ALT (ALAT)	
												X				Aluminium	
										X						Amikacin	
																Ammonia	
																Amylase (Pancreatic)	
																Amylase (Total)	
																Androstenedione	
								X								Anti Streptolysin O (ASO)	
					V			^								Anti-CMV	
					X		V										
							×									Anti-CMV IgG	
				V			X									Anti-CMV IgM	
				X												Anti-EBNA IgG	
				X												Anti-EBV VCA I M	
				X												Anti-EBV VCA IgM	
					X											Anti-HBc	
					X											Anti-HCV	
					X											Anti-HIV-I	
					X											Anti-HIV-I & 2 Combined	
					X											Anti-HIV-2	
							X									Anti-HSV- I & 2 IgG Combined	
							X									Anti-HSV- I & 2 IgM Combined	
							X									Anti-HSVI IgG	
							X									Anti-HSVI IgM	
							X									Anti-HSV2 IgG	

+ = No	t accredited											mistry					ty   +	ty 2 +
* = Pilo	t study ongoing	Ammonia / Ethanol +	Anti-TSH Receptor +	18			Cerebrospinal Fluid +	ion	netry +	+ 1-17		General Clinical Chemistry		logy	Jrine	ıssay	Immunoassay Speciality I +	Immunoassay Speciality 2 +
PURPLE	E = The only parameters available on RQ9135/a	Ammoni	Anti-TSH	Blood Gas	BNP +	Cardiac	Cerebro	Coagulation	CO-Oximetry +	CYFRA 21-1 +	ESR +	General	HbAIc	Haematology	Human Urine	Immunoassay	Immunos	Immunos
Α	Anti-HSV2 IgM																	
	Anti-HTLV-I & 2 Combined																	
	Anti-HTLV-I																	
	Anti-HTLV-II																	
	Anti-Rubella IgG																	
	Anti-Rubella IgM																	
	Anti-TG																X	
	Antithrombin III							X										
	Anti-Toxoplasma IgG																	
	Anti-Toxoplasma IgM																	
	Anti-TPO																X	
	Anti-TSH Receptor (TRAb)		X															
	Apolipoprotein Al																	
	Apolipoprotein B																	
	aPTT							X										
	AST (ASAT)											X						
В	β-2-Microglobulin															X		
	Benzoylecgonine																	
	Bicarbonate			X								X						
	Bile Acids											X						
	Bilirubin (Direct)											X						
	Bilirubin (Total)											X						
												^						
	Blood BNP				X													
					^													
C	Buprenorphine CA15-3															V		
																X		
	CA19-9															X		
	CA125															X		
	Cadmium																	
	Caffeine																	
	Calcitonin																	X
	Calcium											X			X			
	Calcium (Ionised)			X								X						
	Cannabinoids (THC)																	
	Carbamazepine															Χ		
	Carboxyhaemoglobin (COHb / HbCO)								X									
	CEA															Χ		
	Ceruloplasmin																	
	Chloride			X			X					X			X			
	Cholesterol (Total)											Χ						

Immunosuppressant +	Lipid	Liquid Cardiac	Maternal Screening	Serology (EBV) +	Serology (HIV / Hepatitis) +	Serology (Syphilis) +	Serology (ToRCH) +	Specific Proteins	Sweat Testing +	Therapeutic Drug	Trace Elements in Blood +	Trace Elements in Serum +	Trace Elements in Urine +	Urinalysis +	Urine Toxicology +	+ = Not accredited  * = Pilot study ongoing  PURPLE = The only parameters available on RQ	9135/a
트	<u> </u>	Ĕ	Σ̈́	Se	Se	Se		Sp	Š	È	Ė	Ĕ	Ĕ	Š	Ď		
					.,		X									Anti-HSV2 IgM	Α
					X											Anti-HTLV-I & 2 Combined	
					X											Anti-HTLV-I	
					X											Anti-HTLV-II	
							X									Anti-Rubella IgG	
							X									Anti-Rubella IgM	
																Anti-TG	
								X								Antithrombin III	
							X									Anti-Toxoplasma IgG	
							X									Anti-Toxoplasma IgM	
																Anti-TPO	
																Anti-TSH Receptor (TRAb)	
	X															Apolipoprotein Al	
	X															Apolipoprotein B	
																аРТТ	
																AST (ASAT)	
								X								β-2-Microglobulin	В
															X	Benzoylecgonine	
																Bicarbonate	
																Bile Acids	
																Bilirubin (Direct)	
														X		Bilirubin (Total)	
														Χ		Blood	
																BNP	
															X	Buprenorphine	
																CA15-3	С
																CA19-9	
																CA125	
													X			Cadmium	
										V			^			Caffeine Caffeine	
										X							
																Calcitonin	
																Calcium	
																Calcium (Ionised)	
															X	Cannabinoids (THC)	
										X						Carbamazepine	
																Carboxyhaemoglobin (COHb / HbCO)	
																CEA	
								X								Ceruloplasmin	
									X							Chloride	
	X															Cholesterol (Total)	

+ = No	t accredited	_	+				+					mistry					ity I +	ity 2 +
* = Pilo	t study ongoing	Ammonia / Ethanol +	Anti-TSH Receptor +	as			Cerebrospinal Fluid +	tion	CO-Oximetry +	21-1 +		General Clinical Chemistry		ology	Urine	assay	Immunoassay Speciality 1 +	Immunoassay Speciality 2 +
PURPLI	E = The only parameters available on RQ9135/a	Ammoni	Anti-TS	Blood Gas	BNP +	Cardiac	Cerebro	Coagulation	CO-0xi	CYFRA 21-1 +	ESR +	General	HbAIc	Haematology	Human Urine	Immunoassay	Immuno	Immuno
С	Cholinesterase											Х						
	Chromium																	
	Ciclosporin																	
	CK, Total					X						X						
	CK-MB (Activity)					X												
	CK-MB (Mass)					X												
	Cobalt																	
	Complement C <sub>3</sub>																	
	Complement C₄																	
	Conductivity																	
	Copper											X			X			
	Cortisol														X	X		
	Cotinine																	
	C-Peptide															X	X	
	C-Reactive Protein (CRP)																	
	Creatinine											X			X			
	CYFRA 21-1 (Cytokeratin 19)									X								
D	D-3-Hydroxybutyrate											X						
	d-Amphetamine																	
	D-Dimer* <sup>∆</sup>							X										
	Deoxyhaemoglobin (HHb)								X									
	DHEA Unconjugated															X		
	DHEA-Sulphate															X		
	Digoxin															X		
	d-Methamphetamine																	
	Dopamine														X			
Е	EDDP																	
	EGFR*											X						
	Epinephrine Epinephrine														X			
	ESR										X							
	Estriol Total*															X		
	Ethanol	X																
	Ethosuximide* <sup>Δ</sup>															X		
	Everolimus																	
F	Factor II							X										
	Factor IX							X										
	Factor V							X										
	Factor VII							X										
	Factor VIII							X										
	Factor X							X										
	Factor XI							X										
	120001711																	

Immunosuppressant +	Lipid	Liquid Cardiac	Maternal Screening	Serology (EBV) +	Serology (HIV / Hepatitis) +	Serology (Syphilis) +	Serology (ToRCH) +	Specific Proteins	Sweat Testing +	Therapeutic Drug	Trace Elements in Blood +	Trace Elements in Serum +	Trace Elements in Urine +	Urinalysis +	Urine Toxicology +	+ = Not accredited  * = Pilot study ongoing  PURPLE = The only parameters available on RQ91	35/a
																Cholinesterase	С
												X	X			Chromium	
X										X						Ciclosporin	
																CK, Total	
																CK-MB (Activity)	
		X														CK-MB (Mass)	
												X	X			Cobalt	
								X								Complement C <sub>3</sub>	
								X								Complement C <sub>4</sub>	
									X							Conductivity	
											X	X	X			Copper	
																Cortisol	
															X	Cotinine	
																C-Peptide	
								X								C-Reactive Protein (CRP)	
														Χ	Χ	Creatinine	
																CYFRA 21-1 (Cytokeratin 19)	
																D-3-Hydroxybutyrate	D
															X	d-Amphetamine	
		X														D-Dimer* <sup>∆</sup>	
																Deoxyhaemoglobin (HHb)	
																DHEA Unconjugated	
																DHEA-Sulphate	
		X								X						Digoxin	
															X	d-Methamphetamine	
																Dopamine	
															Χ	EDDP	Е
																EGFR*	
																Epinephrine	
																ESR	
																Estriol Total*	
															X	Ethanol	
										Χ						Ethosuximide* <sup>Δ</sup>	
×																Everolimus	
																Factor II	F
																Factor IX	
																Factor V	
																Factor VII	
																Factor VIII	
																Factor X	
																Factor XI	

 $<sup>^{\</sup>vartriangle}$  Pilot status only in certain programmes. Please check pages 32 - 36 for more information.

# = Flot study ongoing  * = Flot study ongoing  PURPLE = The only parameters available on RQ21135/a  *	+ = No	t accredited											mistry					ity   +	ity 2 +
F	* = Pilo	t study ongoing	a / Ethanol +	Receptor 4	as			spinal Fluid	ion	netry +	+  - 2		Clinical Che		logy	Jrine	ıssay	ıssay Speciali	Immunoassay Speciality 2 +
Ferritin	PURPLE	= The only parameters available on RQ9135/a	Ammoni	Anti-TSF	Blood G	BNP +	Cardiac	Cerebro	Coagulat	CO-Oxi	CYFRA 3	ESR +	General	HbAIc	Haemato	Human L	Immunos	Immunos	Immunos
Politice	F	Factor XII							X										
Foliate   Free Morphine   Free Morphine   Free β-NCG		Ferritin															Χ		
Free Morphine   Free Morphi		Fibrinogen							Χ										
Free β-hCG		Folate															X		
Fructosamine		Free Morphine																	
FSH		free β-hCG																	
G   Galactose		Fructosamine											X						
Galactose   Gastrin   Ga		FSH															X		
Gastrin   Gentamicin   Growth Hormone (GH)   X   X   X   X   X   X   X   X   X	G	γ-GT											X						
Gentamicin		Galactose																	
Growth Hormone (GH)		Gastrin																	X
GLDH		Gentamicin															X		
Glucose		Growth Hormone (GH)															X		
Haematocrit (HCT)		GLDH											X						
Haemoglobin (Hb)		Glucose			X			X					X			X			
Total Haemoglobin (Hb)	Н	Haematocrit (HCT)													X				
Haptoglobin		Haemoglobin (Hb)													X				
HbA1c		Total Haemoglobin (tHb)								X				X					
HBSAG		Haptoglobin																	
HBDH		HbAlc												X					
hCG		HBsAG																	
HDL-Cholesterol		НВДН											X						
Homocysteine		hCG															X		
hsCRP		HDL-Cholesterol											X						
IgA		Homocysteine					X												
IgE		hsCRP																	
IGF-I	- 1	lgA																	
IgG		lgE															X		
IgM         Inhibin A           Insulin         X           Iodine         X           Iron         X           K         Kappa Light Chain (Free)           Kappa Light Chain (Total)         X           Ketones         X		IGF-I																X	
Inhibin A   Insulin		IgG						X											
Insulin		IgM																	
Iodine   Iron   X   X   X   X   X   X   X   X   X		Inhibin A																	
Iron X X X X X X X X X X X X X X X X X X X		Insulin															Χ	X	
K Kappa Light Chain (Free) Kappa Light Chain (Total) Ketones		lodine																	
Kappa Light Chain (Total)  Ketones		Iron											X						
Ketones Edward E	K	Kappa Light Chain (Free)																	
		Kappa Light Chain (Total)																	
Lactate X X X		Ketones																	
	L	Lactate			X			X					X						
Lambda Light Chain (Free)		Lambda Light Chain (Free)																	

Factor XII	Immunosuppressant +	Lipid	Liquid Cardiac	Maternal Screening	Serology (EBV) +	Serology (HIV / Hepatitis) +	Serology (Syphilis) +	Serology (ToRCH) +	Specific Proteins	Sweat Testing +	Therapeutic Drug	Trace Elements in Blood +	Trace Elements in Serum +	Trace Elements in Urine +	Urinalysis +	Urine Toxicology +	+ = Not accredited  * = Pilot study ongoing  PURPLE = The only parameters available on Ro	Q9135/a
Ferritin   Filtering   Filte	드			Σ	Ň	Š	Š	Ň	S	Ń	-	-	-	<b>—</b>	$\supset$	$\supset$		
Fibrological   Foliate																		F
									X									
Fructosamine																X		
FSH				X													free β-hCG	
																	FSH	
Castrin   Captamicin   Captam																	γ-GT	G
															X		Galactose	
																	Gastrin	
											Χ						Gentamicin	
																	Growth Hormone (GH)	
Haematocrit (HCT)																	GLDH	
Haemoglobin (Hb)															X		Glucose	
Total Haemoglobin (tHb)																	Haematocrit (HCT)	Н
																	Haemoglobin (Hb)	
HbA1c																	Total Haemoglobin (tHb)	
HBSAG									X								Haptoglobin	
HBDH																	HbAIc	
						X											HBsAG	
X																	НВОН	
X															X		hCG	
X		X															HDL-Cholesterol	
			X														Homocysteine	
									X								IgA	1
									X									
X																		
Insulin				X														
X X X I Iodine Iron  X X X X X I Iodine Iron  X X X X X X X X X I Iodine Iron  X X X X X X X X X X X X X X X X X X X																		
Iron   Kappa Light Chain (Free)   K   X   Kappa Light Chain (Total)   X   Ketones   Lactate   L												X	X	X				
X Kappa Light Chain (Free)  X Kappa Light Chain (Total)  X Ketones  Lactate  L																		
X Kappa Light Chain (Total) X Ketones Lactate L									X									К
X Ketones  Lactate  Lactate																		
Lactate L									, (						X			
															,(			
X Lambda Light Chain (Free)									X									

+ = Not accredited  * = Pilot study ongoing  PURPLE = The only parameters available on RQ9135/a  Lambda Light Chain (Total)  LD (LDH)  LDL-Cholesterol  Lead	Human Urine	Immunoassay	Immunoassay Speciality 1 +	Immunoassay Speciality 2 +
L Lambda Light Chain (Total)  LD (LDH)  LDL-Cholesterol	Human	Immuno	Immuno	Immuno
LD (LDH)  LDL-Cholesterol				
LDL-Cholesterol LDL-Cholesterol				
Lead				
Leukocytes Leukocytes				
Lipase				
Lipoprotein (a)				
Lithium				
Lorazepam				
LSD				
Luteinising Hormone (LH)		X		
M Magnesium X	X			
Manganese Manganese				
MDMA				
Mean Cell Haemoglobin (MCH)				
Mean Cell Haemoglobin Concentration (MCHC)				
Mean Cell Volume (MCV)				
Mean Platelet Volume (MPV)				
Metanephrine Metanephrine	X			
Methadone Methadone				
Methaemoglobin (MetHb)				
Methotrexate Methotrexate				
Molybdenum Molybdenum				
Myoglobin X				
N NEFA X				
Nickel Nickel				
Nitrite Nitrite				
Non-HDL Cholesterol*				
Norepinephrine	X			
Normetanephrine	X			
Norpropoxyphene Norpropoxyphene				
Nortriptyline Nortriptyline				
NTproBNP				
O Oestradiol		Х		
Osmolality X	X			
Osteocalcin			X	
Oxalate Oxalate	X			
Oxazepam Oxazepam				
Oxygen Content (O2CT)				
Oxygen Saturation (sO2 / Vol O2)				
Oxyhaemoglobin (O2Hb / HbO2)				

Immunosuppressant +		Jiac	creening	:BV) +	Serology (HIV / Hepatitis) +	yphilis) +	-oRCH) +	oteins	ing +	c Drug	Trace Elements in Blood +	Trace Elements in Serum +	Trace Elements in Urine +		cology +	+ = Not accredited  * = Pilot study ongoing	
Immunosnb	Lipid	Liquid Cardiac	Maternal Screening	Serology (EBV) +	Serology (F	Serology (Syphilis) +	Serology (ToRCH) +	Specific Proteins	Sweat Testing +	Therapeutic Drug	Trace Elem	Trace Elem	Trace Elem	Urinalysis +	Urine Toxicology +	PURPLE = The only parameters available on Re	Q9135/a
								Χ								Lambda Light Chain (Total)	L
																LD (LDH)	
	X															LDL-Cholesterol	
											X	X	X			Lead	
														X		Leukocytes	
																Lipase	
	X															Lipoprotein (a)	
										X						Lithium	
															X	Lorazepam	
															X	LSD	
																Luteinising Hormone (LH)	
											X		X			Magnesium	М
											X	X	X			Manganese	
															X	MDMA	
																Mean Cell Haemoglobin (MCH)	
																Mean Cell Haemoglobin Concentration (MCHC)	
																Mean Cell Volume (MCV)	
																Mean Platelet Volume (MPV)	
																Metanephrine	
															X	Methadone	
																Methaemoglobin (MetHb)	
										X						Methotrexate	
													X			Molybdenum	
		X														Myoglobin	
																NEFA	N
												X	X			Nickel	
														X		Nitrite	
																Non-HDL Cholesterol*	
																Norepinephrine	
																Normetanephrine	
															X	Norpropoxyphene	
															X	Nortriptyline	
		X														NTproBNP	
																Oestradiol	0
																Osmolality	
																Osteocalcin	
															V	Oxalate	
															X	Oxazepam (O3CT)	
																Oxygen Content (O2CT)	
																Oxygen Saturation (sO2 / Vol O2)	
																Oxyhaemoglobin (O2Hb / HbO2)	

+ = No	t accredited		_									mistry					ity   +	ity 2 +
* = Pilo	t study ongoing	Ammonia / Ethanol +	Anti-TSH Receptor +	as			Cerebrospinal Fluid +	tion	CO-Oximetry +	21-1 +		General Clinical Chemistry		ology	Jrine	assay	Immunoassay Speciality I +	Immunoassay Speciality 2
PURPLE	E = The only parameters available on RQ9135/a	Ammoni	Anti-TS	Blood Gas	BNP +	Cardiac	Cerebro	Coagulation	CO-0xi	CYFRA 21-1 +	ESR +	General	HbAIc	Haematology	Human Urine	Immunoassay	Immuno	Immuno
Р	PAPP-A																	
	Paracetamol (Acetaminophen)															X		
	pCO <sub>2</sub>			Χ														
	pH			X														
	Phencyclidine																	
	Phenobarbital															X		
	Phenytoin															X		
	Phosphate (Inorganic)											X			X			
	Plasma Renin Activity																	Х
	Plasminogen							X										
	Plateletcrit (PCT)													X				
	Platelets (PLT)													X				
	$pO_2$			X														
	Potassium			X								X			X			
	Prealbumin (Transthyretin)																	
	Primidone* <sup>Δ</sup>															X		
	Procalcitonin																X	X
	Progesterone															X		
	Prolactin															X		
	Protein (Total)						X					X			X			
	Protein C							X										
	Protein S							X										
	PSA (Free)															X		
	PSA (Total)											X				X		
	PT (Including INR)							X										
	PTH															X	X	
R	Red Blood Bell Count (RBC)													X				
	Red Cell Distribution Width (RDW)													X				
	Renin (Direct Concentration)																	X
	Retinol Binding Protein																	
	Rheumatoid Factor																	
S	Salicylic Acid															X		
	Secobarbital																	
	Selenium																	
	SHBG															X		
	Sirolimus																	
	Sodium			Х			X					X			X			
	Specific Gravity																	
	Syphilis																	
Т	T <sub>3</sub> (Free)											Χ				Χ		

Blood + + + Curine + + + + + + + + + + + + + + + + + + +	+ = Not accredited
Immunosuppressant + Lipid Liquid Cardiac Maternal Screening Serology (EBV) + Serology (HIV / Hepatitis) + Serology (Syphilis) + Serology (ToRCH) + Therapeutic Drug Trace Elements in Blood + Trace Elements in Serum + Trace Elements in Urine + Urinalysis +	* = Pilot study ongoing
X	PAPP-A P
X	Paracetamol (Acetaminophen)
	pCO <sub>2</sub>
X	pH
X X X	Phencyclidine
× ×	
	Phenytoin Phosphate (Inorganic)
	Plasma Renin Activity
	Plasminogen
	Plateletcrit (PCT)
	Platelets (PLT)
	pO,
	Potassium
X	Prealbumin (Transthyretin)
X	Primidone* <sup>A</sup>
	Procalcitonin
	Progesterone
	Prolactin
X	Protein (Total)
	Protein C
	Protein S
	PSA (Free)
	PSA (Total)
	PT (Including INR)
	PTH
	Red Blood Bell Count (RBC)
	Red Cell Distribution Width (RDW)
	Renin (Direct Concentration)
X	Retinol Binding Protein
X	Rheumatoid Factor
X	Salicylic Acid S
X	Secobarbital
X X	Selenium
	SHBG
X	Sirolimus
	Sodium
X	Specific Gravity
X	Syphilis
	T <sub>3</sub> (Free)

 $<sup>^{\</sup>vartriangle}$  Pilot status only in certain programmes. Please check pages 32 - 36 for more information.

+ = No	ot accredited											nistry					+ - 6:	.y 2 +
* = Pilo	ot study ongoing	Ammonia / Ethanol +	Anti-TSH Receptor +	Blood Gas			Cerebrospinal Fluid +	tion	CO-Oximetry +	21-1 +		General Clinical Chemistry		ology	Urine	assay	Immunoassay Speciality I +	Immunoassay Speciality 2 +
PURPLI	PURPLE = The only parameters available on RQ9135/a				BNP +	Cardiac	Cerebro	Coagulation	00-0X	CYFRA 21-1 +	ESR +	General	HbAIc	Haematology	Human Urine	Immunoassay	Immuno	Immuno
Т	T <sub>3</sub> (Total)											X				X		
	T <sub>4</sub> (Free)											Χ				Χ		
	T <sub>4</sub> (Total)											X				X		
	Tacrolimus																	
	Testosterone (Free)*															X		
	Testosterone (Total)															X		
	Thallium																	
	Theophylline															X		
	Thyroglobulin															X		
	TIBC											X						
	Tobramycin* <sup>∆</sup>															X		
	Total hCG																	
	Transferrin																	
	Triglycerides											X						
	Troponin I					X												
	Troponin T					X												
	TSH											X				X		
	П							X										
U	UIBC											X						
	Unconjugated Oestriol																	
	Urea											X			X			
	Uric Acid											X			X			
	Urobilinogen																	
٧	Valproic Acid															X		
	Vancomycin															X		
	Vitamin B12															X		
	VMA														X			
W	Total White Blood Cell Count (WBC)													X				
Z	Zinc											Χ						

Immunosuppressant +	Lipid	Liquid Cardiac	Maternal Screening	Serology (EBV) +	Serology (HIV / Hepatitis) +	Serology (Syphilis) +	Serology (ToRCH) +	Specific Proteins	Sweat Testing +	Therapeutic Drug	Trace Elements in Blood +	Trace Elements in Serum +	Trace Elements in Urine +	Urinalysis +	Urine Toxicology +	+ = Not accredited  * = Pilot study ongoing  PURPLE = The only parameters available on RQ9135		
_				S	S	S	S	S	S							T <sub>3</sub> (Total)	Т	
																T <sub>4</sub> (Free)		
																T <sub>4</sub> (Total)		
X																Tacrolimus		
																Testosterone (Free)*		
																Testosterone (Total)		
													X			Thallium		
										X						Theophylline		
																Thyroglobulin		
																TIBC		
										X						Tobramycin* <sup>Δ</sup>		
			X													Total hCG		
								X								Transferrin		
	X															Triglycerides		
		X														Troponin I		
		X														Troponin T		
																TSH		
																тт		
																UIBC	U	
			X													Unconjugated Oestriol		
																Urea		
																Uric Acid		
														X		Urobilinogen		
										X						Valproic Acid	٧	
										X						Vancomycin		
																Vitamin B12		
																VMA		
																Total White Blood Cell Count (WBC)	W	

 $<sup>^{\</sup>vartriangle}$  Pilot status only in certain programmes. Please check pages 32 - 36 for more information.

### **RELATED PRODUCTS**

# ACUSERA True Third Party Quality Controls

As a world leading manufacturer of multi-analyte true third party controls, thousands of laboratories rely on Randox to accurately assess test system performance and ultimately empower them with the confidence required to release patient test results. With more than 400 analytes available, the number of individual controls required to cover your test menu is significantly reduced while simultaneously reducing costs, time and storage space. A choice of formats is available, including liquid or lyophilised, which ensures flexibility and suitability for laboratories of all sizes and budgets. Many features of the Acusera range can help you to meet ISO 15189:2012 requirements:

- Designed to react to the test system in the same manner as a patient sample, helping to reduce inconvenient shifts in QC results when reagent batch is changed and ultimately providing a true indication of laboratory performance.
- The presence of analytes at key decision levels ensures accurate instrument performance and eliminates the need for additional low/high controls at extra expense.
- Manufactured independently from any instrument, the Acusera range delivers unbiased performance assessment with any instrument or method, while eliminating the need for multiple instrument specific controls.

#### **Product Portfolio**

Antioxidants | Blood Gas | Cardiac Markers | Routine Chemistry | Coagulation | Haematology | Diabetes | Immunoassay | Immunology | Infectious Diseases (Serology) | Lipids | POCT | Therapeutic Drugs | Toxicology | Urine Chemistry



Uniquely combining more than 100 analytes conveniently in a single control, laboratories can significantly reduce costs and consolidate without compromising on quality. As true third party controls, unbiased performance assessment with any instrument or method is guaranteed.

### **RELATED PRODUCTS**

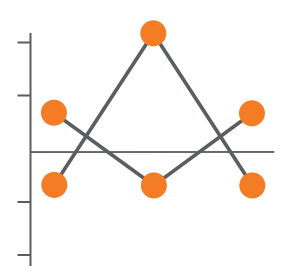
# ACUSERA 24.7 Interlaboratory Data Management

Designed for use with the Acusera range of third party controls, the Acusera 24•7 software helps laboratories monitor and interpret their QC data. Access to an impressive range of features, including interactive charts, the automatic calculation of Measurement Uncertainty & Sigma Metrics and live peer group data generated from our extensive database of laboratory participants, ensures Acusera 24•7 is the most comprehensive package available.

- Advanced statistical analysis with automatic calculation of performance metrics including; Sigma, UM, TE & %Bias.
- Instantly discover how you compare to your peers with peer group statistics updated live in real-time reducing time and money spent troubleshooting.
- Interactive charts allowing you to add events and multiple data sets for quick and easy performance monitoring.
- · Automated data import with bi-directional connection to LIMS (eliminating manual data entry).

### Software Features

Dashboard | Result History | Interactive Levey-Jennings Charts | Interactive Histogram Charts | Performance Summary Charts | Statistical Analysis Report | Statistical Metrics Report | Audit Trail Report Uncertainty of Measurement Report | Exception Report | Peer Group Statistics | Acusera Advisor



'The laboratory shall have a procedure to prevent the release of patient results in the event of quality control failure. When the quality controls rules are violated and indicate that examination results are likely to contain significant errors the results shall be rejected... Quality Control data shall be reviewed at regular intervals to detect trends in examination performance'.

ISO 15189:2012

### RANDOX - A GLOBAL DIAGNOSTIC SOLUTIONS PROVIDER

Randox has been supplying laboratories worldwide with revolutionary diagnostic solutions for over 35 years. Our experience and expertise allow us to create a leading product portfolio of high quality diagnostic tools which offer reliable and rapid diagnosis. We believe that by providing laboratories with the right tools, we can improve healthcare worldwide.

#### **RX SERIES**



Renowned for quality and reliability, the RX series combines robust hardware and intuitive software with the world leading RX series test menu comprising an extensive range of high quality reagents including routine chemistries, specific proteins, lipids, therapeutic drugs, drugs of abuse, antioxidants and diabetes testing. The RX series offers excellence in patient care delivering unrivalled precision and accuracy for results you can trust, guaranteeing real cost savings through consolidation of routine and specialised tests onto one single platform.

### **REAGENTS**



Randox offers an extensive range of third-party diagnostic reagents which are internationally recognised as being of the highest quality; producing accurate and precise results. At Randox, we re-invest significantly in R&D to ensure we meet the ever-changing needs of the laboratory. Consequently, Randox offer a range of novel and superior performance assays, including: sdLDL-C, Lipoprotein (a), H-FABP, Adiponectin, Copper and Zinc. Applications are available detailing instrument-specific settings for the convenient use of Randox Reagents on numerous clinical chemistry analysers.

### **EVIDENCE SERIES**



In 2002, Randox invented the world's first, Biochip Array Technology, offering highly specific tests, coupled to the highly sensitive chemiluminescent detection, providing quantitative results instantly changing the landscape of diagnostic testing forever. The Randox Evidence Series of multi-analyte immunoanalysers provide an unrivalled increase in patient information per sample offering diagnostic, prognostic and predictive solutions across a variety of disease areas with a highly advanced clinical and toxicology immunoassay test menu including cardiac, diabetes, drugs of abuse, metabolic and renal markers.

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