EXCELLENT MEASUREMENT OF SPECIFIC PROTEINS
SPECIFIC PROTEINS

Alpha-1-Acid Glycoprotein | Alpha-1-Antitrypsin | Anti-Streptolysin O
Apolipoproteins | Ceruloplasmin | Complement C3 | Complement C4
C-Reactive Protein | Cystatin C | Ferritin | Haptoglobin
IgA | IgE | IgG | IgM | Lipoprotein (a) | Microalbumin | Myoglobin
Rheumatoid Factor | Soluble Transferrin Receptor (sTfR) | Transferrin | Transthyretin |
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KEY

NP
Niche Product
When you see this symbol you will know that Randox have one of the only automated biochemistry assays available on the market

UF
Unique Feature
When you see this symbol you will know that this feature is unique to the Randox product
BENEFITS OF RANDOX 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.

We have the largest test menu of 115 assays, covering over 100 disease markers including specific proteins, lipids, therapeutic drug monitoring, drugs of abuse, antioxidants, coagulation, diabetes and veterinary testing.

A wide range of formats and methods are available providing greater flexibility and choice for any laboratory size.

In addition to flexible pack sizes and a comprehensive list of analyser applications, we can also provide dedicated reagent packs (Randox Easy Read and Easy Fit reagents) for a wide range of chemistry analysers providing you with freedom of choice from an independent manufacturer.

EXPAND ROUTINE TESTING
With speciality assays for 195 of the most common clinical chemistry analysers; assays which usually require dedicated equipment such as nephelometers (or was previously only available as an ELISA) can now be run on automated biochemistry analysers, allowing your laboratory to expand its routine test menu. E.g. TxBCardio™, cystatin C, adiponectin, CRP, Lp(a) and many more.

EXPAND YOUR TEST MENU WITHOUT EXPANDING YOUR LAB
There is no need to buy any extra equipment in order to expand your test menu. Our reagents can be programmed onto the majority of the most common biochemistry analysers.

REDUCE LABOUR
Reduce time with liquid ready-to-use reagents, automated methods (compared to the traditional laborious ELISA methods used for some tests such as cystatin C or adiponectin); and our easy-fit options.

REDUCE THE RISK OF ERRORS AND HAVE CONFIDENCE IN PATIENT RESULTS
Our traceability of material and extremely tight manufacturing tolerances ensure uniformity across reagent batches reducing lot-to-lot variability. Our assays are validated against gold-standard methods; increasing confidence in patient test results.

REDUCE CE COSTS
We can help create cost-savings for your laboratory through excellent reagent stability; by eliminating the need for costly re-runs through the excellent quality of products; and by offering a range of kit sizes (including smaller kit sizes for niche tests to reduce waste).

BRING TESTING IN-HOUSE
With smaller kit sizes and excellent reagent stability (most are stable for 28 days on-board the analyser), you don’t have to worry about reagent wastage, allowing testing to be brought in-house.
Alpha-1-Acid Glycoprotein

WHAT IS ALPHA-1-ACID GLYCOPROTEIN?

Alpha-1-Acid Glycoprotein (AAG), also known as orosomucoid, is an acute-phase reactant synthesised in the liver in response to inflammation and tissue damage. The normal range for healthy individuals is 50-120 mg/dl.

CLINICAL SIGNIFICANCE

Markedly higher AAG levels are observed in a number of conditions such as inflammatory diseases and acute myocardial infarction, trauma. High AAG levels have also been observed in pregnancy and surgery. AAG concentrations rise rapidly until 48 hours after surgery followed by little change until about 120 hours, regardless of the severity of the injury. Serum AAG levels also provide a useful diagnostic tool in neonates with bacterial infections.

BENEFITS OF THE Randox AAG ASSAY

- Wide measuring range of 24.6-453 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox AAG assay on a wide range of biochemistry analysers
- Complementary controls and calibrator offering a complete testing package
- Traceable to ERM DA470k/IFCC

Alpha-1-Antitrypsin

WHAT IS ALPHA-1-ANTITRYPSIN?

Alpha-1-Antitrypsin (AAT) primarily serves to protect the elastin fibres of the lungs by inhibiting neutrophil elastase. The normal range for healthy individuals is 90-200 mg/dl.

CLINICAL SIGNIFICANCE

Low levels of AAT (<80 mg/dl) have major clinical importance in association with emphysema and liver disease. Increased levels (3-4 times normal) can occur as a result of trauma, pregnancy, administration of oestrogens or typhoid vaccine.

BENEFITS OF THE Randox ALT ASSAY

- Wide measuring range of 38.7-660 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox ALT assay on a wide range of biochemistry analysers
- Complementary controls and calibrator offering a complete testing package
- Traceable to ERM DA470k/IFCC
Anti-Streptolysin O

WHAT IS ANTI-STREPTOLYSIN O?

Streptolysin O (SLO) is a lethal, exocellular protein produced by Group A Streptococcal bacteria. The SLO toxin lyases erythrocytes and many other animal cells by disruption of the cytoplasmic and similar membranes. Consequently, anti-streptolysin O antibodies (ASO) are produced by the host to neutralise the haemolytic action of the SLO. Levels of ASO in serum are dependent on the age of the patient, geographical location and the local incidence of streptococcal infection. The upper limit of the normal range, internationally, is 200IU/ml as this value is rarely exceeded without symptoms indicative of streptococcal infection.

CLINICAL SIGNIFICANCE

ASO measurement can provide information on the extent and severity of infection. A streptococcal infection is considered to be one where there is a two-dilution rise in titre between acute and convalescent stage serum. Hence, the test should be repeated after 1 to 2 weeks. Raised ASO levels may also be present in other conditions such as scarlet fever, acute rheumatic arthritis, tonsillitis and various other streptococcal infections as well as in healthy carriers.

The World Health Organisation (WHO) recommends the use of ASO to aid the diagnosis of streptococcal infections such as rheumatic fever and glomerulonephritis.

BENEFITS OF THE RANDOX ASO ASSAY

- Wide measuring range of 28-1314 IU/ml for the accurate detection of clinically important results
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox ASO assay on a wide range of biochemistry analysers
- Complementary controls and calibrator offering a complete testing package
- Single point calibration, directly calibrated against NIBSC 97/662
- No interference from C1q complement
**Apolipoprotein A-I**

**WHAT IS APO A-I?**
Apo A-I is the major protein component of HDL particles in plasma and plays a role in lipid metabolism. Chylomicrons secreted from the intestinal enterocyte also contain Apo A-I, but it is quickly transferred to HDL in the bloodstream.

**CLINICAL SIGNIFICANCE**
The chief role of Apo A-I is the activation of lecithin cholesterol acyl transferase (LCAT) and the capture and removal of free cholesterol from hepatic tissues, also known as reverse cholesterol transport. Apo A-I is therefore non-atherogenic, showing an inverse relationship to cardiovascular risk.

**BENEFITS OF THE RANDOX A-I ASSAY**
- Wide measuring range of 5.27 - 251 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox Apo A-I assay on a wide range of biochemistry analysers
- Calibrators supplied with kits simplifying the ordering process
- Complementary controls offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results

**Apolipoprotein A-II**

**WHAT IS APO A-II?**
Apo A-II is a major constituent of High Density Lipoprotein (HDL) particles and plays an important role in the reverse cholesterol transport and lipid metabolism. The increased production of Apo A-II promotes atherosclerosis by decreasing the proportion of anti-atherogenic HDL containing Apo A-I.

**CLINICAL SIGNIFICANCE**
Increased production of Apo A-II promotes atherosclerosis by decreasing the proportion of anti-atherogenic HDL containing Apo A-I.

**BENEFITS OF THE RANDOX APO A-II ASSAY**
- Wide measuring range of 6.75 - 61.1 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox Apo A-II assay on a wide range of biochemistry analysers
- Calibrators supplied with kits simplifying the ordering process
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results
Apolipoprotein B

WHAT IS APO B?
Apo B is a component of LDL cholesterol and enables tissue cells to take up cholesterol. Apo B is considered atherogenic making it a useful risk marker for CVD.

CLINICAL SIGNIFICANCE
Elevated levels of Apo B correlates with an increased risk of CVD, even when total and LDL cholesterol levels are within the normal range, making Apo B an important risk marker.

BENEFITS OF THE RANDOX APO B ASSAY
- Wide measuring range of 9.37 - 233 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox Apo B assay on a wide range of biochemistry analysers
- Calibrator supplied with some kits simplifying the ordering process
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results

Apolipoprotein C-II

WHAT IS APO C-II?
Apo C-II acts as a co-factor for lipoprotein lipase, an enzyme that hydrolyses triglycerides in chylomicrons and vLDL. Lipoprotein lipase cannot function without Apo C-II; it is essential for the normal triglyceride breakdown. If an individual is deficient in Apo C-II, they would experience a build up of triglycerides and subsequent hypertriglyceridemia.

CLINICAL SIGNIFICANCE
Patients have been identified with excessive hypertriglyceridemia due to a deficiency in Apo C-II. Conditions associated with Apo C-II and hypertriglyceridemia include: chylomicronemia, xanthomas, and recurrent pancreatitis.

BENEFITS OF THE RANDOX APO C-II ASSAY
- Excellent sensitivity of 1.48 mg/dl, ensuring depleted levels are detected
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox C-II assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results
Apolipoprotein C-III

WHAT IS APO C-III?
Apo C-III inhibits the action of lipoprotein lipase and as a result modulates the uptake of triglyceride-rich lipoproteins.

CLINICAL SIGNIFICANCE
Elevated levels of Apo C-III are associated with both primary and secondary hypertriglyceridemia. Genetically determined Apo C-III deficiency in humans has shown to increase the rate of triglyceride clearance from the plasma by 6 to 7-fold. Elevated Apo C-III levels have been reported in many pathological conditions including type 2 diabetes, hyperbilirubinemia, kidney deficiency and decreased thyroid function. Several factors influence Apo C-III levels including: gender, age, menopause and genetic polymorphisms in the Apo C-III gene.

BENEFITS OF THE RANDOX APO C-III ASSAY
- Excellent linearity of 2.17 mg/dl for the comfortable detection of elevated levels
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox Apo C-III assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results

Apolipoprotein E

WHAT IS APO E?
Apo E is a protein synthesised mainly in the brain and to a lesser extent in the spleen, lungs, adrenal glands, ovaries, kidneys, muscle cells and macrophages. Apo E's functions include the transport of triglycerides to the liver tissues and the distribution of cholesterol between cells.

CLINICAL SIGNIFICANCE
Apo E deficiency gives rise to high cholesterol and triglyceride levels, promoting atherosclerosis. The polymorphism has been associated with diseases other than cardiovascular disease, for example E4 is implicated in Alzheimer's disease.

BENEFITS OF THE RANDOX APO E ASSAY
- Wide measuring range of 1.04-12.3 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox Apo E assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results
Ceruloplasmin

WHAT IS CERULOPLASMIN?

Ceruloplasmin is an alpha 2 globulin synthesised primarily in the liver. It binds to copper after it is absorbed from the gastrointestinal (GI) tract and is responsible for transporting more than 90% of all copper to various tissues within the body. Ceruloplasmin has several important functions including ferroxidase activity, amine oxidase activity and superoxidase activity. It is also involved in homeostasis.

CLINICAL SIGNIFICANCE

The main use of ceruloplasmin is in the diagnosis of Wilson disease, a rare inherited disorder characterised by liver damage and neurological deterioration. Individuals with Wilson disease often have decreased levels of ceruloplasmin which subsequently leads to a build-up of copper in the liver, brain and other organs. If Wilson disease is identified before significant copper deposits affect the function of the major organs, serious damage can be avoided. If left undiagnosed and untreated, Wilson disease can be fatal. Early detection and treatment is therefore crucial in order to prevent permanent damage and halt disease progression.

As an acute-phase reactant, ceruloplasmin concentrations are also elevated in cases of bacterial infection, stress, pregnancy, leukemia, Hodgkin’s disease, systemic lupus erythematosis and rheumatoid arthritis.

Deficiency of ceruloplasmin is strongly associated with copper toxicity. Unbound copper may result in an increase in oxidative stress, damaging cells and tissues and accelerating neurological degeneration.

BENEFITS OF THE RANDOX CERULOPLASMIN ASSAY

- Wide measuring range of 6.29-73.8 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox ceruloplasmin assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results
Complement C3

**WHAT IS COMPLEMENT C3?**
Complement C3 is the most important and prevalent protein within the complement system. Complement C3 works in conjunction with antibodies and other factors to protect the body from invasion by pathogens. When activated by either the classical or alternative pathway, complement C3 acts on biological membranes and may cause cell death.

**CLINICAL SIGNIFICANCE**
Decreased complement C3 levels are important in determining inherited or acquired deficiencies and may increase the susceptibility to infections or developing autoimmune disorders. Conversely, levels may rise in a variety of inflammatory and necrotic disorders as part of the acute-phase plasma protein response.

**BENEFITS OF THE RANDOX COMPLEMENT C3 ASSAY**
- Wide measuring range of 15.5-502 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox complement C3 assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results
- Traceable to ERM DA470k/IFCC

Complement C4

**WHAT IS COMPLEMENT C4?**
Complement C4 is a complex biological system which works in conjunction with antibodies and other factors to protect the body from invasion by pathogens. When activated by either the classical or alternative pathway, complement C4 acts on biological membranes and may cause cell death.

**CLINICAL SIGNIFICANCE**
Decreased complement C4 levels are important in determining inherited or acquired deficiencies and may increase the susceptibility to infections or developing autoimmune disorders. Conversely, levels may rise in a variety of inflammatory and necrotic disorders as part of the acute-phase plasma protein response.

**BENEFITS OF THE RANDOX COMPLEMENT C4 ASSAY**
- Wide measuring range of 3.41-152 mg/dl for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox complement C4 assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Limited interference from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results
- Traceable to ERM DA470k/IFCC
C-Reactive Protein (CRP)

WHAT IS C-REACTIVE PROTEIN?
C-Reactive Protein or CRP is an acute phase reactant found at low concentrations in the serum of normal patients. During the acute phase response of an inflammatory reaction, CRP levels are elevated up to 1000-fold making it an excellent early indicator of infection.

WHY MEASURE CRP?
Elevated CRP levels indicate an inflammatory response has been activated. CRP is more sensitive compared to other acute phase reactants which can take 24 hours to produce a detectable signal. Levels above the normal range (>6 mg/L) are used to diagnose and monitor rheumatic diseases and other inflammatory conditions. Recent research has widened the clinical significance of CRP and measurement within the normal range can be used for risk assessment of cardiovascular disease, detection of infection in neonates and in early detection of renal allograft rejection.

Inflammation
CRP is useful in monitoring recovery from surgery and rheumatic disorders. CRP levels can rise to 30-100 mg/l after surgery.

Cardiovascular Disease
Inflammation is being linked to the development of atherosclerosis. CRP levels at the upper end of the normal range may indicate that a low-level inflammatory response has been initiated and there is an increased risk of cardiovascular disease and stroke.

Neonate Infection
Neonatal susceptibility to bacterial infection is a well-established complication. Early detection of infection is important to enable rapid and effective antibiotic treatment and to minimise antibiotic dose. This will lead to a reduction in risk of antibiotic-related kidney damage and enable earlier neonatal recovery from infection.

Renal Allograft Rejection
Monitoring postoperative CRP levels in renal transplant patients enables early detection of allograft rejection. CRP levels peak around 3 days earlier than the creatinine levels, a standard biochemical marker for allograft assessment. Recent studies have indicated significant elevations in CRP levels in all cases of rejection indicating 100% sensitivity.

Infection
During viral infections such as upper respiratory tract infections, influenza, pneumonia and meningitis, CRP levels will be slightly elevated to around 10-20 mg/l. Elevated CRP levels from 30->200 mg/l are found in bacterial infections such as septic arthritis, meningitis, pneumonia and also in rheumatoid arthritis.
CRP LEVELS SHOWING CARDIAC RISK AND DISEASES

- High cardiac risk
- Intermediate cardiac risk
- Low cardiac risk
- Normal range
- Viral infection
- Myocardial infarction
- After surgery
- Rheumatoid arthritis

CRP KITS FROM RANDOX

- **Full Range CRP**: Test for measurement of CRP levels within and outside the normal range (0.3 - 161 mg/l)
- **CRP**: Test for measurement of CRP levels outside the normal range (2.88 - 220 mg/l)
- **High Sensitivity CRP**: Test for measurement of low CRP levels within the normal range (0.270 - 10 mg/l)
### CRP

**WHY USE THE RANDOX CRP ASSAY?**

CRP measurements are indicative of infection, tissue injury, inflammatory disorders and associated diseases. Recent research highlights that CRP measurements can also be used to assess the risk of cardiovascular disease, infection in neonates and for the early detection of renal allograft rejection. Elevated levels of CRP are associated with a 3-fold risk of myocardial infarction.

**BENEFITS OF THE RANDOX CRP ASSAY**

- **Wide measuring range** of 3.57-219 mg/L for the accurate detection of clinically important results
- **Automated latex enhanced immunoturbidimetric assay** eliminating the need for any dedicated equipment
- **Liquid ready-to-use reagents** for convenience and ease-of-use
- **Stable to expiry** when stored at +2 to +8°C
- **Applications available** detailing instrument-specific settings for the convenient use of the Randox full range CRP assay on a wide range of biochemistry analysers
- **Complementary controls and calibrators** offering a complete testing package
- **Limited interference** – from Bilirubin, Haemoglobin, Intralipid® and Triglycerides, producing more accurate results

### Full-Range CRP

**WHY USE THE RANDOX FULL-RANGE CRP ASSAY?**

The Randox Full-Range CRP assay combines exceptional sensitivity in the low CRP range with excellent linearity at elevated levels for an all-inclusive test. The Randox Full-Range CRP assay enables the precise and accurate measurement of CRP levels both within and above the normal range, all within one kit, eliminating the need for a dedicated high sensitivity CRP test. With instrument-specific applications, the Randox CRP assay can be automated on a range of biochemistry analysers for the rapid routine assessment of CRP levels.

**BENEFITS OF THE RANDOX FULL-RANGE CRP ASSAY**

- **Wide measuring range** of 0.18-165 mg/L for the accurate detection of clinically important results
- **Automated latex enhanced immunoturbidimetric assay** eliminating the need for any dedicated equipment
- **Liquid ready-to-use reagents** for convenience and ease-of-use
- **Stable to expiry** when stored at +2 to +8°C
- **Applications available** detailing instrument-specific settings for the convenient use of the Randox full range CRP assay on a wide range of biochemistry analysers
- **Complementary controls and calibrators** offering a complete testing package
- **Traceable to ERM DA474k/IFCC**
High-Sensitivity CRP

WHY USE THE RANDOX HIGH-SENSITIVITY CRP ASSAY?

High sensitivity CRP, or hsCRP, in addition to lipid evaluation and risk scoring systems helps in the assessment of cardiovascular disease (CVD) risk. The American Heart Association (AHA) and Centre for Disease Control and Prevention (CDC) now recommend the use of hsCRP as a more sensitive marker of CVD risk compared to traditional CRP assays. The hsCRP assay is particularly useful in predicting future cardiac events in individuals with no previous history of CVD. Healthy individuals with CRP levels higher than 3mg/l are 2 to 4 times more likely to have a heart attack or stroke. Approximately half of all heart attacks occur in patients who have a normal lipid profile and are classified as low risk based on traditional methods of risk estimation. The measurement of hsCRP can therefore help clinicians to identify these individuals earlier; it can also be used to evaluate the risk of a recurrent cardiac event. In high risk groups there have even been indications that hsCRP could be used as a prognostic tool.

AHA/CDC RISK ASSESSMENT GUIDELINES:

<table>
<thead>
<tr>
<th>Risk Level</th>
<th>hs-CRP (mg/L)</th>
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<tbody>
<tr>
<td>Low</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Average</td>
<td>1-3</td>
</tr>
<tr>
<td>High</td>
<td>&gt;3</td>
</tr>
</tbody>
</table>

BENEFITS OF THE RANDOX CRP ASSAY

- Wide measuring range of 0.477-10 mg/l for the accurate detection of clinically important results
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox hsCRP assay on a wide range of biochemistry analysers
- High sensitivity CRP control and calibrator offering a complete testing package
Cystatin C

WHAT IS CYSTATIN C?

Cystatin C is a small (13 kDa) cysteine proteinase inhibitor that is steadily produced by all nucleated cells. The normal range for healthy individuals is 0.57-1.05 mg/l.

CLINICAL SIGNIFICANCE

The small molecular weight of cystatin C means it can be freely filtered by the glomerular membrane; cystatin C levels in the blood are therefore indicative of a normal or impaired Glomerular Filtration Rate (GFR). Creatinine is perhaps the most widely used marker of GFR however unlike cystatin C, creatinine levels are affected by non-renal factors including: age, gender, race, muscle mass, diet and disease. As creatinine is secreted by tubular cells into the bloodstream a 24 hour urine sample is required, furthermore creatinine is unable to detect small decreases in GFR. Cystatin C on the other hand is degraded in renal tubular cells and is not secreted by the kidneys meaning plasma and serum levels are dependent on the GFR.

Cancer therapeutics can damage renal function; early indication of such damage through the measurement of cystatin C levels is therefore important as it allows the oncologist to adjust drug dosage.

Poor diabetes control can also affect renal function, therefore monitoring cystatin C levels is also useful in diabetic patients.

BENEFITS OF THE RANDOX CYSTATIN C ASSAY

- Wide measuring range of 0.4-10 mg/L for the accurate detection of clinically important results
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox cystatin C assay on a wide range of biochemistry analysers
- Cystatin C control and calibrators offering a complete testing package
Ferritin

WHAT IS FERRITIN?
Iron is essential for respiration as it forms part of the haemoglobin structure. It is the iron atom which binds to oxygen and is responsible for oxygen transport to bodily cells and tissues. Iron is also found in myoglobin and other proteins and enzymes. Ferritin is a metalloprotein that stores iron intracellularly, mainly in the liver, spleen and bone marrow. Ferritin is an accurate and indirect indicator of the body’s iron stores and levels are measured in iron deficiency and iron toxicity.

CLINICAL SIGNIFICANCE
Anaemia is a shortage of red blood cells with a wide range of causes including excessive bleeding, haemolysis and reduced red blood cell production. Anaemia symptoms tend to be vague, for example fatigue and shortness of breath, therefore anaemia is diagnosed using a full blood count.
Tests such as ferritin, glucose and renal function are used to confirm the diagnosis and determine the cause of anaemia. In iron toxicity, iron accumulates in the body causing organ dysfunction. High levels of stored iron may be an indicator of infection, liver disease, alcohol abuse, overactive thyroid, rheumatoid arthritis, certain cancers or Wilson Disease.

Early diagnosis of these conditions is critical as treatment is relatively simple: periodic phlebotomies and minor dietary changes. Serum iron, total iron-binding capacity and transferrin tests are also used in diagnosis. CRP may be used to rule out elevated ferritin levels due to inflammation.

BENEFITS OF THE RANDOX FERRITIN ASSAY
• Wide measuring range of 5.08 - 443 ng/ml for the accurate detection of clinically important results
• Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
• Liquid ready-to-use reagents for convenience and ease-of-use
• Stable to expiry when stored at +2 to +8°C
• Applications available detailing instrument-specific settings for the convenient use of the Randox ferritin assay on a wide range of biochemistry analysers
• Complementary controls and calibrators offering a complete testing package
Haptoglobin

WHAT IS HAPTOGLOBIN?
Haptoglobin is an acute phase protein made primarily in the liver. Its main function is to bind free oxyhaemoglobin and remove it from circulation in order to prevent renal injury and iron loss following haemolysis. Haptoglobin is also thought to have both anti-inflammatory and antioxidant properties.

CLINICAL SIGNIFICANCE
Haptoglobin measurements are used in the diagnosis of haemolytic anaemia and to distinguish it from other types of anaemia. In haemolytic anaemia haptoglobin levels in the blood decrease significantly, low levels however may also indicate red blood cell destruction due to sickle cell anaemia or thalassemia. In certain cases of liver disease haptoglobin levels may also be low as the liver cannot manufacture normal levels of the protein.

As an acute phase reactant, haptoglobin levels in the blood are significantly increased in response to infection, inflammation, acute tissue necrosis, malignant tumours, burns, surgery and trauma.

BENEFITS OF THE RANDOX HAPTOGLOBIN ASSAY
• Wide measuring range of 0.1-3.68 g/l for the accurate detection of clinically important results
• Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
• Liquid ready-to-use reagents for convenience and ease-of-use
• Stable to expiry when stored at +2 to +8°C
• Applications available detailing instrument-specific settings for the convenient use of the Randox haptoglobin assay on a wide range of biochemistry analysers
• Complementary controls and calibrators offering a complete testing package
• Traceable to ERM DA470k/IFCC
Immunoglobulin A

WHAT IS IgA?
IgA makes up 15% to 20% of the body’s immunoglobulin pool. Its main role is to provide antibody activity at or near mucosal surfaces, such as in the gastrointestinal system, genitourinary system and respiratory system. For this reason, secretions from these areas contain high levels of IgA (in a specialised secretory IgA form). Examples of secretions with high IgA levels include saliva, bile, airway secretions, genitourinary secretions and milk. IgA fixes complement via the alternative pathway.

CLINICAL SIGNIFICANCE
Measurement of IgA is used to diagnose diseases of the respiratory tract e.g. tuberculosis, Chron’s disease and early cirrhosis of the liver. It is also useful in monitoring therapy of IgA myeloma and evaluating IgA immunity. IgA in colostrum and milk is important in neonatal defence against infection.

BENEFITS OF THE RANDOX IgA ASSAY
- Wide measuring range of 0.21-5.9 g/l for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox IgA assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Traceable to ERM DA470k/IFCC

Immunoglobulin E

WHAT IS IgE?
IgE accounts for 0.002% of immunoglobulins in the body. IgE antibodies are the chief immunoglobulin responsible for immediate hypersensitivity reactions in humans.

CLINICAL SIGNIFICANCE
Continual production of IgE antibodies in response to common naturally occurring allergens and the production of excessive amounts of histamine by the IgE bound mast cells results in the development of such clinically important allergic reactions such as asthma, hay fever, dermatitis and food allergies. Elevated IgE levels are also seen in parasitic diseases, IgE myeloma and in hepatitis.

BENEFITS OF THE RANDOX IgE ASSAY
- Wide measuring range of 19.6-1007 IU/ml for the accurate measurement of clinically important results
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox IgE assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package

ALL ORDERING INFORMATION CAN BE FOUND ON PAGE 23–24
RANDOX REAGENTS
**Immunoglobulin G**

WHAT IS IgG?

This is the principle immunoglobulin in normal human serum, and makes up 80% of the total serum immunoglobulin in adult humans. IgG is the main form of antibody produced in response to secondary infections. This immunoglobulin provides one of the body’s major defences against bacterial infection in adults, unborn children and newborn babies as it is the only immunoglobulin that can cross the placenta.

CLINICAL SIGNIFICANCE

Measurement of IgG is the basis of the serological diagnosis of several infectious diseases. Uses of IgG measurement include diagnosis of infectious and inflammatory diseases, diagnosis of malignancies, to detect the presence of soluble antigens and monitoring therapy in myeloma.

BENEFITS OF THE RANDOX IgG ASSAY

- Wide measuring range of 2.8-27.6 g/l for the accurate detection of clinically important results
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox IgG assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Traceable to ERM DA470k/IFCC

**Immunoglobulin M**

WHAT IS IgM?

IgM is the major antibody (accounting for approximately 10% of the body’s immunoglobulins) found in serum in the first week following infection. This antibody is particularly effective in combating bacterial infections because of its high binding affinity for proteins responsible for destroying bacterial cells (complement proteins). IgM is usually present in humans as groups of five molecules. IgM does not cross the placenta.

CLINICAL SIGNIFICANCE

IgM measurement has the following uses: to establish diagnosis and monitor therapy in Waldenström’s macroglobulinemia and plasma cell myeloma, to detect intra-uterine infection by measuring levels in newborn babies, diagnosis of primary biliary cirrhosis, viral hepatitis, rheumatoid arthritis and parasitic infections.

BENEFITS OF THE RANDOX IgM ASSAY

- Wide measuring range of 0.23-3.77 g/l for the accurate detection of clinically important results
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox IgM assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
- Traceable to ERM DA470k/IFCC
Lipoprotein (a)

WHAT IS LIPOPROTEIN (A)?

Lp(a) is an LDL-like particle with a cholesterol-rich core and a molecule of apo B-100 linked by a disulphide bridge to the glycoprotein apo(a). Apo(a) is unique in that it is extremely heterogeneous in size (which is genetically predetermined) due to the Kringle 4 type 2 domain which can be present in up to 40 copies. The size heterogeneity of apo(a) affects to varying degrees the outcome of many commercially available Lp(a) kits, resulting in overestimation of samples containing large apo(a) molecules and an underestimation of samples containing small apo(a) molecules. Research has documented and shown the Randox method to be one of only a few to exhibit minimum size-related bias.

CLINICAL SIGNIFICANCE

Lipoprotein (a) in combination with other lipid tests can provide clinicians with much needed additional information on an individual's risk of CVD. High levels of Lp(a) are known to occur in individuals with an otherwise normal lipid profile and as such it is thought to contribute to an increased risk of cardiovascular disease independent of other lipids. It is also of particular use in assessing the risk of coronary heart disease in specific populations as Lp(a) concentrations are genetically determined and vary with ethnic population.

Although not a routinely requested test the National Cholesterol Education Programme and the National Academy of Clinical Biochemistry recognise the usefulness of Lp(a) and recommend testing patients with a family history of premature CVD or those classified as moderate/high risk. In June 2010, the European Atherosclerosis Society (EAS) also published a consensus paper recommending the widespread use of Lp(a) as a screening tool in those at intermediate or high risk of cardiovascular disease.

BENEFITS OF THE RANDOX Lp(a) ASSAY

- The Randox Lp(a) assay is one of the only methodologies on the market that detects the non-variable part of the Lp(a) molecule and therefore suffers minimal size-related bias, providing more accurate and consistent results. The Randox Lp(a) kit is standardised to the WHO/IFCC reference material SRM 2B and is closest in terms of agreement to the ELISA reference method.
- Five-point calibrators with accuracy-based assigned target values are provided which accurately reflect the heterogeneity of isoforms present in the general population.
- Wide measuring range of 3-90 mg/dl for accurate detection of clinically important results.
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment.
- Measuring units available in nmol/L upon request.
- Highly sensitive and specific method for the detection of Lp(a) in serum and plasma.
- Applications available detailing instrument-specific settings for the convenient use of the Randox Lp(a) assay on a wide range of biochemistry analysers.
- Liquid ready-to-use reagents for convenience and ease-of-use.
- Lp(a) controls and calibrator offering a complete testing package.
Microalbumin

WHAT IS MICROALBUMIN?

Kidney function may be assessed through measurement of albumin levels in the urine. Normal kidney function entails filtering of waste products from the blood across tiny capillaries in the glomerulus. Kidney malfunction results when the capillaries become blocked, resulting in a build-up of waste products in the blood and a loss of important proteins. Kidney deterioration is progressive and begins with small amounts of albumin leaking into the urine. This is known as microalbuminuria and indicates early signs of nephropathy. The term ‘micro-’ refers to low concentrations of urinary albumin.

Progression of kidney disease will lead to larger amounts of albumin leaking into the urine which may develop further to end stage renal disease. Kidney disease is a major concern in diabetic patients and early detection and treatment may slow the onset and progression of the condition.

CONDITION ALBUMIN LEVELS

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CLINICAL SIGNIFICANCE

Albumin is one of the major plasma proteins. In normal circumstances, albumin molecules are too large to cross the glomerular basement membrane, therefore, albumin is usually present in very low concentrations in urine. Damage to the glomerular basement membrane can alter its permeability allowing albumin to enter the urine. Sustained elevations of urinary albumin concentrations are called microalbuminuria.

Microalbumin testing can identify individuals with diabetic nephropathy approximately 5-10 years earlier than proteinuria tests helping reduce the incidence of end-stage renal disease. It is recommended that patients at a high-risk of renal disease (hypertension and diabetes) are tested for microalbumin.

BENEFITS OF THE RANDOX MICROALBUMIN ASSAY

- Excellent sensitivity of 5.11 mg/l ensuring depleted albumin concentrations are detected
- Calibrator supplied with the kit simplifying the ordering process
- Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox microalbumin assay on a wide range of biochemistry analysers
- Microalbumin control and calibrator offering a complete testing package
Myoglobin

WHAT IS MYOGLOBIN?
Myoglobin is an oxygen binding haem-protein found in cardiac and skeletal muscle. It is released into the circulatory system when muscle cells are damaged.

CLINICAL SIGNIFICANCE
Elevated levels of myoglobin is a strong indicator of muscle tissue damage and muscular inflammation. Myoglobin determination can be useful in the diagnosis of myocardial infarction, myositis and in the treatment of myopathy and muscular dystrophy.

BENEFITS OF THE RANDOX MYOGLOBIN ASSAY
- Wide measuring range of 20.1-725 ng/ml for the accurate detection of clinically important results
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox myoglobin assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package

Rheumatoid Factor

WHAT IS RHEUMATOID FACTOR?
The agglutination reaction between antibody coated red blood cells and rheumatoid arthritis (RA) sera, was first demonstrated by Waaler and Rose in 1940. The reaction has since been shown to be caused by certain factors in the RA sera. These rheumatoid factors (RF) are a heterogeneous group of high molecular weight auto-antibodies directed against the body’s own immunoglobulins. They are produced by plasma cells present at sites of tissue injury. The initiating antigen is thought to be one or more viruses or viral antigens that persist in the joint tissues.

CLINICAL SIGNIFICANCE
Research has shown that both environmental and genetic factors can affect the production of RF with various biological properties. Although they may be found in all immunoglobulin classes, the RF most frequently detected is the IgM type; present in about 75% of adult patients with RA and about 10% of children with juvenile RA. RF has also been observed in the serum of patients with lupus erythematosus, hepatitis, liver cirrhosis, syphilis and various other conditions; but the RF titre is much lower than in RA.

BENEFITS OF THE RANDOX RF ASSAY
- Wide measuring range of 6.72-104 IU/ml for the accurate detection of clinically important results
- Accurate assessment of RF titre (calibrant standardised against primary WHO material; 1st British Standard 64/2)
- No interference from C1q complement
- Automated latex enhanced immunoturbidimetric assay eliminating the need for any dedicated equipment
- Liquid ready-to-use reagents for convenience and ease-of-use
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox RF assay on a wide range of biochemistry analysers
- Complementary controls and calibrators offering a complete testing package
WHAT IS sTfR?

Transferrin transports iron around the body, donating it to bodily cells by interacting with a specific membrane receptor, the transferrin receptor (TfR). The soluble transferrin receptor (sTfR) is a truncated form (shortened) of the transferrin receptor, formed as a result of proteolysis (the breakdown of proteins into smaller polypeptides or amino acids). A soluble form of the TfR (sTfR) has been identified in animal and human serum, circulating freely in the blood. The serum concentration of sTfR is directly proportional to the concentration of the membrane bound transferrin receptors.

CLINICAL SIGNIFICANCE

sTfR is a marker of iron status. In iron deficiency anaemia, sTfR levels are significantly increased, however, remain normal in acute phase conditions including: chronic diseases and inflammation. As such, sTfR measurement is useful in the differential diagnosis of anaemia: anaemia of chronic disease (ACD) or iron deficiency anaemia (IDA).

In IDA, increased sTfR levels have been observed in haemolytic anaemia, sickle cell anaemia, B12 deficiency and functional iron deficiency in pregnancy.

In ACD, sTfR levels do not correlate with iron status. This was observed in patients with chronic illnesses (cystic fibrosis and cancer), certain infections, autoimmune diseases (insulin-dependent diabetics) and inflammatory diseases.

sTfR is a useful test where ferritin tests may be inconclusive. As ferritin is an acute phase reactant, levels will increase in response to inflammation. As sTfR is not an acute phase reactant, sTfR levels are not impacted by inflammation.

BENEFITS OF THE RANDOX sTfR ASSAY

- Liquid ready-to-use reagent for convenience and ease-of-use
- Latex Enhanced Immunoturbidimetric method facilitating testing on biochemistry analysers and eliminating the need for dedicated equipment
- Excellent measuring range of 0.5 – 11.77 mg/L, for the comfortable detection of clinically important results
- Excellent correlation coefficient of r=0.977 when compared against another commercially available methods
- Stable to expiry when stored at +2 to +8°C
- Applications available detailing instrument-specific settings for the convenient use of the Randox sTfR assay on a wide range of biochemistry analysers
Transferrin

WHAT IS TRANSFERRIN?
Transferrin (siderophilin) is the principal iron binding and transport protein in human plasma and can bind two molecules of iron. The normal range for healthy adults is 200-380 mg/dl. Iron availability in the plasma regulates transferrin levels which increase when plasma iron is low.

CLINICAL SIGNIFICANCE
Transferrin levels increase during pregnancy and oestrogen administration and correlate closely with Total Iron Binding Capacity of serum. Plasma transferrin levels are associated with a range of conditions including anaemia, iron deficiency, inflammation or malignancy, liver disease, malnutrition and protein loss.

BENEFITS OF THE RANDOX TRANSFERRIN ASSAY
• Wide measuring range of 7.60-497 mg/dl for the accurate detection of clinically important results
• Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
• Liquid ready-to-use reagents for convenience and ease-of-use
• Stable to expiry when stored at +2 to +8°C
• Applications available detailing instrument-specific settings for the convenient use of the Randox transferrin assay on a wide range of biochemistry analysers
• Complementary controls and calibrators offering a complete testing package

Transthyretin (Prealbumin)

WHAT IS TRANSTHYRETIN?
Transthyretin, or prealbumin, may be referred to as thyroxin-binding prealbumin as it binds thyroxine and triiodothyronine. Transthyretin has a molecular mass of 55kDa and the prealbumin name is derived from its electrophoretic mobility, as it migrates faster than albumin on electrophoresis. It is manufactured in the liver hepatocytes and forms a complex with retinol-binding protein, to aid the transport of vitamin A.

CLINICAL SIGNIFICANCE
Transthyretin is a negative acute phase reactant and serum levels fall in inflammation, malignancy, cirrhosis of the liver and protein-wasting diseases of the gut or kidneys, owing to decreased synthesis. Elevated levels have been reported in Hodgkinson’s disease. Transthyretin is a good marker of visceral protein status and positive nitrogen balance.

Early Detection of Malnutrition
Transthyretin is a specific clinical indicator of nutritional risk in the management of diseases such as HIV/AIDS, renal disease, diabetes, pneumonia and cancer. Nutritional assessment using transthyretin has also been effective in surgical cases, pre-surgical screening, fractures and wound healing.

Transthyretin Screening
Transthyretin screening can lead to the early identification of patients with Protein Calorie Malnutrition (PCM) and implementation of diet therapy. Transthyretin screening is more specific than albumin for the identification of PCM risk.

BENEFITS OF THE RANDOX TRANSTHYRETIN ASSAY
• Wide measuring range of 2.3-65 mg/dl for the accurate detection of clinically important results
• Automated immunoturbidimetric assay eliminating the need for any dedicated equipment
• Liquid ready-to-use reagents for convenience and ease-of-use
• Stable to expiry when stored at +2 to +8°C
• Applications available detailing instrument-specific settings for the convenient use of the Randox transthyretin assay on a wide range of biochemistry analysers
• Complementary controls and calibrators offering a complete testing package
### ORDERING INFORMATION

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<td>R1 4 x 8.7ml, R2 4 x 3ml</td>
<td>IM8045</td>
</tr>
<tr>
<td>IgM</td>
<td>Immunoturbidimetric</td>
<td>3 x 20ml</td>
<td>IM7977</td>
</tr>
<tr>
<td>IgM</td>
<td>Immunoturbidimetric</td>
<td>R1 4 x 8.7ml, R2 4 x 3.9 ml</td>
<td>IM3834</td>
</tr>
<tr>
<td>Lipoprotein (a)</td>
<td>LEI</td>
<td>R1 1 x 30 ml, R2 1 x 15ml</td>
<td>LP2757</td>
</tr>
<tr>
<td>Lipoprotein (a)</td>
<td>LEI</td>
<td>R1 1 x 10 ml, R2 1 x 6ml</td>
<td>LP3403</td>
</tr>
<tr>
<td>Lipoprotein (a)</td>
<td>LEI</td>
<td>R1 1 x 10ml, R2 1 x 6.5ml</td>
<td>LP8324</td>
</tr>
<tr>
<td>Microalbumin</td>
<td>Peg Enhanced Immunoturbidimetric</td>
<td>R1 3 x 100ml, R2 2.5 x 7ml</td>
<td>MA2423</td>
</tr>
<tr>
<td>Microalbumin</td>
<td>Peg Enhanced Immunoturbidimetric</td>
<td>R1 1 x 60ml, R2 1 x 7ml</td>
<td>MA2426</td>
</tr>
<tr>
<td>Microalbumin</td>
<td>Peg Enhanced Immunoturbidimetric</td>
<td>R1 6 x 20ml, R2 3 x 8ml</td>
<td>MA3828</td>
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<tr>
<td>Microalbumin</td>
<td>Peg Enhanced Immunoturbidimetric</td>
<td>R1 2 x 20ml, R2 2 x 6.6ml</td>
<td>MA8056</td>
</tr>
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<td>Microalbumin</td>
<td>Peg Enhanced Immunoturbidimetric</td>
<td>R1 1 x 20ml, R2 1 x 4.6ml</td>
<td>MA8325</td>
</tr>
<tr>
<td>Myoglobin</td>
<td>LEI</td>
<td>R1 1 x 9.5ml, R2 1 x 4.5ml</td>
<td>MY2127</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>LEI</td>
<td>R1 2 x 20ml, R2 2 x 8ml</td>
<td>RF3836</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>LEI</td>
<td>R1 2 x 15ml, R2 1 x 10ml</td>
<td>RF7980</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>Disposable Latex Slide Test</td>
<td>100T</td>
<td>RF1436</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>Glass Latex Slide Test</td>
<td>100T</td>
<td>RF2716</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>LEI</td>
<td>R1 2 x 8.7ml, R2 2 x 4.7ml</td>
<td>RF8063</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>LEI</td>
<td>R1 1 x 11.7ml, R2 1 x 5.7ml</td>
<td>RF8345</td>
</tr>
<tr>
<td>Soluble Transferrin Receptor (sTR)</td>
<td>LEI</td>
<td>R1 1 x 9.0ml, R2 1 x 5.8ml</td>
<td>TF10159</td>
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<td>Transferrin</td>
<td>Immunoturbidimetric</td>
<td>R1 6 x 20ml, R2 2 x 15ml</td>
<td>TF7197</td>
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<tr>
<td>Transferrin</td>
<td>Immunoturbidimetric</td>
<td>R1 6 x 20ml, R2 3 x 14ml</td>
<td>TF3831</td>
</tr>
<tr>
<td>Transthyretin (Prealbumin)</td>
<td>Immunoturbidimetric</td>
<td>R1 6 x 20ml, R2 3 x 11ml</td>
<td>PA3843</td>
</tr>
</tbody>
</table>

**KEY**

- * For use with RX analysers only
- ♦ Indicates liquid option
- (C) Indicates calibrator included in kit
- (S) Indicates standard included in kit, and is for manual and semi-automated use only
- LEI Latex Enhanced Immunoturbidimetric
- T Number of tests

To receive a quotation or for further enquiries, please contact reagents@randox.com

To place an order, please contact order.entry@randox.com
Immunoturbidimetry methods have become the main technique for performing protein tests. The transition from nephelometry has been cautious but is increasing as laboratories enjoy the comparability and flexibility of immunoturbidimetry.

Immunoturbidimetry and nephelometry both measure the turbidity of a sample to determine the level of an analyte. Upon addition of the assay reagent, antibodies and antigen cluster to form an immune complex that precipitates, increasing the turbidity of the sample. When light is passed through the reaction solution, some light is scattered by the sample, some light is absorbed by the sample and the rest passes through the sample.

Immunoturbidimetry measures the absorbance of the light by the sample, nephelometry measures the light scattered at a fixed angle. The level of analyte is determined by comparison with a calibrator of known concentration.

Immunoturbidimetry is ideal for the detection of proteins, where the analyte concentration is inversely proportional to the transmitted light signal. Historically nephelometry has been more sensitive than conventional immunoturbidimetry. In latex-enhanced immunoturbidimetry, inert microscopic particles enlarge the immune complexes, amplifying the reaction and significantly increasing the sensitivity of the reaction.

Nephelometers are dedicated analysers only capable of performing this type of assay. In addition, they:
- have high consumable costs
- require highly trained personnel

Immunoturbidimetric tests are carried out on routine biochemistry analysers that are:
- versatile
- fast
- cost-effective
- offer longer reagent stability
- sensitive

The main advantage of nephelometry was its sensitivity; however latex-enhanced immunoturbidimetry has closed this gap. Immunoturbidimetric tests are an increasingly accepted alternative to nephelometry for specific protein assays, and studies have shown a close correlation between Randox immunoturbidimetric tests and nephelometry.

Correlation between nephelometric and immunoturbidimetric methods (n=38)

Correlation between nephelometric and immunoturbidimetric methods (n=38)

R² = 0.99
A-Z PORTFOLIO OF REAGENTS

Acetaminophen (Paracetamol) | CO₂ Total
Acid Phosphatase | Cocaine Metabolite
Adiponectin | Complement C3
Albumin | Complement C4
Aldolase | Copper
Alkaline Phosphatase | Creatinine
Alpha-1-Acid Glycoprotein | CRP
Alpha-1-Antitrypsin | CRP (Canine)
Alanine Aminotransferase (ALT) | CRP (Full Range)
Ammonia | CRP (High Sensitivity)
Amylase | Cystatin C
Amylase (Pancreatic) | Digoxin
Anti-Streptolysin O (ASO) | Ecstasy
Apolipoprotein A-I | EDDP
Apolipoprotein A-II | Ethanol
Apolipoprotein B | Ferritin
Apolipoprotein C-II | Fructosamine
Apolipoprotein C-III | G-6-PDH
Apolipoprotein E | Gamma GT
Aspartate Aminotransferase (AST) | Gentamicin
Barbiturates | GLDH
Benzodiazepines | Glucose
β₂-Microglobulin | Glutamate
Bile Acids | Glutamine
Bilirubin Direct | Glutathione Peroxidase (Ransel)
Bilirubin Total | Glutathione Reductase
Calcium | Glycerol
Cannabinoids | Haemoglobin
Carbamazepine | Haptoglobin
Ceruloplasmin | HbA1c
Chloride | HbA1c II
Cholesterol (Total) | Heart-type Fatty Acid Binding Protein
Cholesterol (HDL) | (H-FABP)
Cholesterol (HDL3) | Homocysteine
Cholesterol (LDL) | D-3-Hydroxybutyrate (Ranbut)
Cholesterol (sdLDL-C) | IgA
Cholinesterase | IgE
CK-MB | IgG
CK-NAC | IgM

Iron
Iron/UIBC
L-Lactate
Lactate Dehydrogenase (L-P)
Lactate Dehydrogenase (P-L)
Leucine Aminopeptidase (LAP)
Lipase
Lipoprotein (a)
Lithium
Magnesium
Methadone
Methamphetamine
Microalbumin
Myoglobin
NEFA (Non-Esterified Fatty Acids)
Opiates
Phenobarbital
Phenytoin
Phosphorus
Potassium
Pregnancy Test
Rheumatoid Factor (RF)
Sodium
Soluble Transferrin Receptor (sTfR)
sPLA₂-11A
Superoxide Dismutase (Ransod)
Syphilis
Total Iron Binding Capacity (TIBC)
Total Antioxidant Status (TAS)
Total Protein
Transferrin
Transthyretin (Prealbumin)
Triglycerides
TxB2Cardio™
Urea
Uric Acid
Urinary Protein
Valproic Acid
Zinc
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 health care 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.

**INTERNAL QUALITY CONTROL**

Acusera third party quality controls are made using the highest quality material of human origin, ensuring they react like a real patient sample. With more than 390 analytes available across the Acusera range we can uniquely reduce the number of controls required while reducing costs and time. Our product range includes clinical chemistry, immunoassay, urine, immunology and more. Qnostics molecular controls for infectious disease testing are designed to meet the demands of today’s molecular diagnostics laboratory while effectively monitoring the entire testing process. Our whole pathogen molecular controls comprise hundreds of characterised viral, bacterial and fungal targets.

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RIQAS is the world’s largest international EQA scheme with more than 45,000 participants worldwide. 33 comprehensive, yet flexible programmes cover a wide range of clinical diagnostic testing including chemistry, immunoassay, cardiac, urine, serology and more. Our programmes benefit from a wide range of concentrations, frequent reporting, rapid feedback and user-friendly reports. The QCMD range of molecular infectious disease EQA programmes feature a whole pathogen matrix ensuring a true test of patient sample analysis. With access to over 90 programmes including blood borne viruses, respiratory diseases, multi-pathogen infections and more, there is something for every laboratory.

**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 immunoanalyser’s 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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