

【Product Name】

Preanesthetic Profile (7+3)

【Packing Specification】

Type B: 1 Test / Disc, 10 Discs / Box.

Type B with diluent container.

【Testing Instrument】

CareSign-C chemistry analyzer

【Intended Use】

Preanesthetic Profile (7+3) used with the CareSign-C chemistry analyzer, is intended to be used for the in vitro quantitative determination of total Protein (TP), albumin (ALB), alanine aminotransferase (ALT), alkaline phosphatase (ALP), creatinine (CRE), urea nitrogen(BUN), glucose (GLU) in heparinized whole blood, heparinized plasma, or serum in a clinical laboratory setting or point-of-care location.

The Preanesthetic Profile (7+3) measurements are used in the diagnosis of liver and gallbladder diseases, urinary system diseases, glucose metabolism disorder.

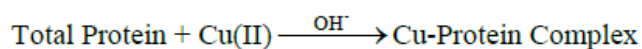
【Principles of Testing】

The Preanesthetic Profile (7+3) is used to quantitatively test the concentration of the 7 biochemical indicators in the sample, which is based on the spectrophotometry. The principles are as follows:

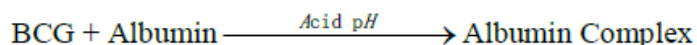
1. Total Protein (TP)

The total protein method is a Biuret reaction, the protein solution is treated with cupric [Cu(II)] ions in a strong alkaline medium. The Cu(II) ions react with peptide bonds between the carbonyl oxygen and amide nitrogen atoms to form a colored Cu-protein complex.

The amount of total protein present in the sample is directly proportional to the absorbance of the Cu-protein complex. The total protein test is an endpoint reaction and the absorbance is measured as the difference in absorbance between 546 nm and 800 nm.

**2. Albumin (ALB)**

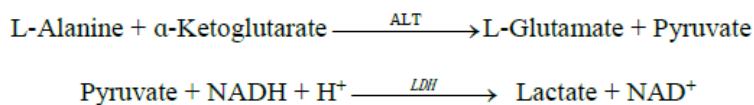
Bromcresol green (BCG), when bound with albumin, changes color from a yellow to green color. The absorbance maximum changes with the color shift.



Bound albumin is proportional to the concentration of albumin in the sample. This is an endpoint reaction that is measured as the difference in absorbance between 600 nm and 700 nm.

3. Alanine Aminotransferase (ALT)

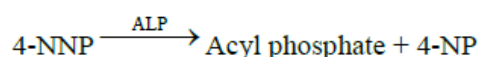
ALT catalyzes the transfer of an amino group from L-alanine to α -ketoglutarate to form L-glutamate and pyruvate. Lactate dehydrogenase catalyzes the conversion of pyruvate to lactate. Concomitantly, NADH is oxidized to NAD^+ , as illustrated in the following reaction scheme.



The rate of change of the absorbance difference between 340 nm and 405 nm is due to the conversion of NADH to NAD^+ and is directly proportional to the amount of ALT present in the sample.

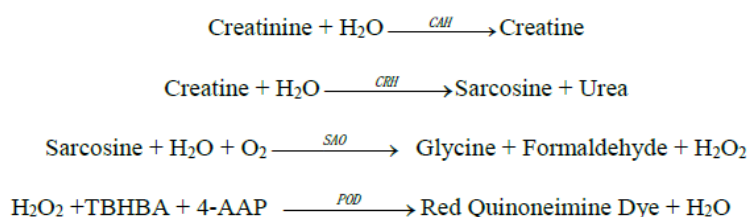
4. Alkaline Phosphatase (ALP)

Under the catalysis of ALP, the Phosphoric acid on nitrobenzene (4-NNP) was turned into Para nitro phenol (4-NP). 4-NP shows a yellow color in alkaline solution. At the wavelength of 405/505nm, the ALP activity can be calculated by monitoring the absorbance change rate.



5. Creatinine (CRE)

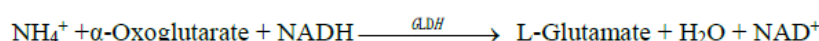
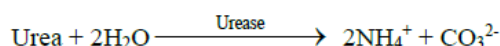
In the coupled enzyme reactions, creatinineamidohydrolase (CAH) hydrolyzes creatinine to creatine. A second enzyme, creatineamidinohydrolase (CRH), catalyzes the formation of sarcosine from creatine. Sarcosine oxidase (SAO) causes the oxidation of sarcosine to glycine, formaldehyde and hydrogen peroxide (H_2O_2). In a Trinder finish, peroxidase (POD) catalyzes the reaction among the hydrogen peroxide, 2, 4, 6-tribromo-3-hydroxybenzoic acid (TBHBA) and 4-aminoantipyrine (4-AAP) into a red quinoneimine dye. Potassium ferrocyanide and ascorbate oxidase are added to the reaction mixture to minimize the potential interference of bilirubin and ascorbic acid respectively.



Two cuvettes are used to determine the concentration of creatinine in the sample. Endogenous creatine is measured in the blank cuvette, which is subtracted from the combined endogenous creatine and the creatine formed from the enzyme reactions in the test cuvette. Once the endogenous creatine is eliminated from the calculations, the concentration of creatinine is proportional to the intensity of the red color produced. The endpoint reaction is measured as the difference in absorbance at 546 nm and 700 nm.

6. Urea Nitrogen (BUN)

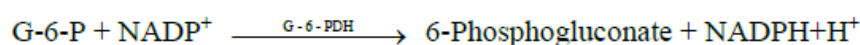
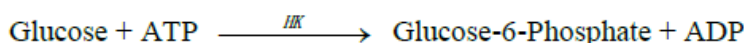
In the coupled-enzyme reaction, urease hydrolyzes urea into ammonia and carbon dioxide. Upon combining ammonia with α -oxoglutarate and reduced nicotinamide adenine dinucleotide (NADH), the enzyme glutamate dehydrogenase (GLDH) oxidizes NADH to NAD^+ .



The rate of change of the absorbance difference between 340 nm and 405 nm is caused by the conversion of NADH to NAD⁺ and is directly proportional to the amount of urea present in the sample.

7. Glucose (GLU)

The reaction of glucose with adenosine triphosphate (ATP) catalyzed by hexokinase (HK), produces glucose-6-phosphate (G-6-P) and adenosine diphosphate (ADP). Glucose-6-phosphate dehydrogenase (G-6-PDH) catalyzes the reaction of G-6-P into 6-phosphogluconate and the reduction of nicotinamide adenine dinucleotide phosphate (NADP⁺) to NADPH.



The absorbance is measured bichromatically at 340 nm and 405 nm. The production of NADPH is directly proportional to the amount of glucose present in the sample.

【Principle of Operation】

Refer to the CareSign-C chemistry analyzer Operator's Manual, for the Principles and Limitations of the Procedure.

【Description of Reagents】

Each Preanesthetic Profile (7+3) contains lyophilized test-specific reagent beads. A lyophilized blank reagent bead includes in each disc for a judgment of error 0233.

Type B is the reagent disc with diluent container.

Calibration information is included in barcode code. Please check it on the label.

The component of each Preanesthetic Profile (7+3) is as follows (after redissolution):

Component	Quantity
Total protein assay reagent	13.5 µL
Albumin assay reagent	13.5 µL
Alanine Aminotransferase assay reagent	13.5 µL
Alkaline Phosphatase assay reagent	13.5 µL
Creatinine assay reagent	13.5 µL
Urea assay reagent	13.5 µL
Glucose assay reagent	6.6 µL
Stabilizer	Appropriate amount

【Storage】

Store reagent discs in their sealed pouches at a temperature of 2-8°C (36-46°F). Do not expose opened

or unopened discs to direct sunlight or temperatures exceeding 32°C (90°F). Reagent discs may be used until the expiration date indicated on the package, which is also encoded in the unique code printed on the sealing pouch.

A torn or damaged pouch may allow moisture to reach the unused disc, adversely affecting its performance. Therefore, do not use any disc from a damaged pouch.

【Sample Requirements】

Sample collection techniques are described in the “Sample requirement” section of the CareSing-C chemistry analyzer Operator’s Manual.

The required sample usage is 100 µL of lithium heparin whole blood, lithium heparin plasma, serum or quality controls.

Whole blood samples collected by venipuncture must be homogeneous before transferring the sample to a reagent disc.

At the same time, it is necessary to carry out the test within 60 minutes. Before taking the test, shake the lithium heparin blood collection tube gently upside down several times.

The glucose concentration is affected by the patient's feeding time and the storage environment after the sample is collected. In order to accurately measure glucose, a sample of the patient should be taken after at least 12 hours of fasting. For uncentrifuged samples stored at room temperature, the glucose concentration is reduced by about 5-12 mg / dL in 1 hour.

Light may cause total bilirubin to decompose, causing deviations in the test results. Whole blood samples that are not tested immediately should be stored in a dark environment.

Use only lithium heparin evacuated specimen collection tubes for whole blood or plasma samples.

The test was started within 10 minutes after transferring the sample to the reagent disc.

【Interfering Substances】

Studies on known drugs or chemicals have found that when the interfering substances contained in the sample exceed the contents in the table below, the final test results are affected.

Analyte	Interfering substances concentration (≤)						
	Bilirubin mg/dL	Intralipid mg/dL	Hemoglobin mg/dL	Vitamin C mg/dL	Pyruvate mmol/L	NH ₄ Cl mmol/L	Creatine µmol/L
TP	25	1050	200	—	—	—	—
ALB	40	600	1000	—	—	—	—
ALT	40	600	50	50	1	—	—
ALP	40	1050	400	—	—	—	—
CRE	40	1050	500	25	—	—	600
BUN	25	600	1000	—	—	1	—
GLU	40	600	1000	50	—	—	—

【Procedure】

■ Materials Provided

Preanesthetic Profile (7+3)

CareSign-C chemistry analyzer

Please tear off the aluminum strip before using Type B.

Transfer pipettes (fixed volume 100 µL for sample) and tips

■ Test Procedure

The complete sample collection and step-by-step operating procedures are detailed in the Operator's Manual for the CareSign-C chemistry analyzer.

■ Calibration

Each batch of reagent is calibrated using Randox standard serum to obtain the disc-specific calibration parameters before shipment.

The calibration parameters stored in the two-dimensional code printed on the sealed pouch are provided to analyzer at the time of scanning the code.

Refer to the Operator's Manual for specific information.

■ Quality Control

Refer to Operator's Manual of the CareSign-C chemistry analyzer. Performance of the CareSign-C chemistry analyzer can be verified by running controls. For a list of approved quality control materials with acceptance ranges, please consult the manual.

If control results are out of range, repeat one time. If still out of range, call i-SENS customer service or local distributors for technical support. Do not report the results if controls are outside their labeled limits.

■ Results

The CareSign-C chemistry analyzer automatically calculates and prints the analyte concentrations in the sample. Details regarding endpoint and rate reaction calculations can be found in the CareSign-C chemistry analyzer Operator's Manual.

【Normal Reference Ranges】

These ranges are provided as a guideline only. It is recommended that your office or institution establish normal ranges for your particular patient population.

Analyte	SI Units	Common Units
TP	Dog: 52 ~ 82g/L;	Dog: 5.2 ~ 8.2g/dL;
	Cat: 54 ~ 89g/L	Cat: 5.4 ~ 8.9g/dL
ALB	Dog: 22 ~ 44g/L;	Dog: 2.2 ~ 4.4 g/dL;
	Cat: 22 ~ 45g/L	Cat: 2.2 ~ 4.5 g/dL
ALT	Dog: 10 ~ 140U/L;	Dog: 10 ~ 140U/L;
	Cat: 8.2 ~ 123U/L	Cat: 8.2 ~ 123U/L
ALP	Dog: 20 ~ 150U/L;	Dog: 20 ~ 150U/L;
	Cat: 10 ~ 90U/L	Cat: 10 ~ 90U/L
CRE	Dog: 27 ~ 149μmol/L;	Dog: 0.3 ~ 1.7mg/dL;
	Cat: 27 ~ 223μmol/L	Cat: 0.3 ~ 2.5mg/dL
BUN	Dog: 2.5 ~ 11.5mmol/L	Dog: 7 ~ 32mg/dL
	Cat: 3.6 ~ 15.5mmol/L	Cat: 10 ~ 43mg/dL
GLU	Dog: 3.89 ~ 7.95mmol/L	Dog: 70 ~ 143mg/dL
	Cat: 4.11 ~ 8.84mmol/L	Cat: 74 ~ 159mg/dL

【Interpretation of Results】

Physiological interferents, such as hemolysis, icterus, and lipemia, can cause changes in the reported concentrations of certain analytes. Sample indices are printed at the bottom of each printout to inform

the operator about any abnormalities in the sample. The operator should take care to avoid hemolysis caused by improper blood collection techniques.

The CareSign-C chemistry analyzer suppresses any results that are affected by >10% interference from hemolysis, lipemia or icterus. “HEM”, “LIP”, or “ICT” respectively, is printed on the printout in place of the result.

Any result for a particular test that exceeds the assay range should be analyzed by another approved test method or sent to a referral laboratory. Do not dilute the sample and run it again on the CareSign-C chemistry analyzer.

【Limitations of Procedure】

The Preanesthetic Profile (7+3) should be used with the CareSign-C chemistry analyzer, and is just used for in vitro diagnosis (IVD).

As with any diagnostic test procedure, all other test procedures including the clinical status of the patient, should be considered prior to final diagnosis.

【Performance Characteristics】

Accuracy

Analyte	The relative deviation or absolute deviation should meet the following requirements
TP	B% ≤ 5.0%
ALB	B% ≤ 6.0%
ALT	B% ≤ 15.0%
ALP	B% ≤ 10.0%
CRE	B% ≤ 10.0%
BUN	B% ≤ 15.0%
GLU	B% ≤ 20.0%

Batch precision

Analyte	Coefficient of variation (≤ *)
TP	2.0%
ALB	2.0%
ALT	5.0%
ALP	5.0%
CRE	5.0%
BUN	5.0%
GLU	5.0%

Inter batch precision

Analyte	Relative Range (≤ *)
TP	5.0%
ALB	5.0%

ALT	10.0%
ALP	10.0%
CRE	10.0%
BUN	10.0%
GLU	10.0%

Dynamic Ranges

Analyte	Dynamic Ranges
TP	20~100g/L
ALB	10 ~ 60g/L
ALT	5 ~ 1500U/L
ALP	5 ~ 2000U/L
CRE	20 ~ 2000 μ mol/L
BUN	0.9~35.7mmol/L
GLU	1 ~ 35 mmol/L

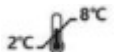
【Notes】

Used reagent discs contain animal body fluids. It is essential to follow good laboratory safety practices when handling and disposing of these used discs. For instructions on cleaning biohazardous spills, refer to the CareSign-C chemistry analyzer Operator's Manual.

The reagent discs are made of plastic and may crack or chip if dropped. Never use a disc that has been dropped, as it may spray biohazardous material throughout the interior of the analyzer.

Reagent beads may contain acids or caustic substances. Operators do not come into contact with the reagent beads when following the recommended procedures. It is important to avoid ingestion, skin contact, or inhalation of the reagent beads.

【Symbols Used in Labelling】

Symbol	Explanation
	Veterinary use only
	Manufacturer
	Use-by date
	Batch code
	Date of manufacture
	Consult instructions for use
	Limit of temperature
	Do not re-use

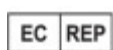
【Manufactured for i-SENS by MNCHIP】



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