

**ALL VALUE ASSIGNMENT RANGES CALCULATED USING
ABAXIS REAGENTS ON ABAXIS ANALYZERS**

Product Number ALCV-G14033-050
Lot # 464019002
Containing Vial Lots S1 #464119002, S2 #464219002
& S3 #464319002
Expiration Date: **05 2021**
3 Levels x 3 vials x 0.5mL

Intended Use

NOD® Verification Samples are human Liquid assayed serum samples used to validate analyzer performance according to CLIA guidelines. The laboratory testing procedures are listed in the following text. Verification Samples must be run at least every six months in Labs using non-waived Reagent discs 400-0026 Hepatic Panel; 400-0031 Basic Metabolic Panel Plus; and 400-0034 MetLyte Plus CRP.

Summary and Principles

The use of independent quality verification sample materials is indicated as an objective assessment of the precision of methods and techniques in use and is an integral part of good laboratory practices. Three levels of verification samples are available to allow performance monitoring within the clinical range.

Reagent Composition

This product is prepared from purified human serum to which biochemical material (human and animal tissue extracts); drugs, chemicals, stabilizers and preservatives have been added. The verification samples are in a prepackaged liquid form to avoid potential error or contaminate being introduced during reconstitution.

Storage and Stability

To achieve maximum shelf life for the Verification kit store unopened at ≤ -20°C. Store vials away from the light. Bilirubin may decrease over the product shelf life.

Thawed and Unopened: The Verification Samples can be used for up to **14 days** when stored unopened 2-8°C. For optimum Bilirubin and CO₂ stability avoid

prolonged exposure of the Verification Sample vials to ambient air / room temperature / light.

Procedure

The verification samples should be treated the same as a patient sample and run according to the instructions accompanying the instrument, kit, or reagent being used. Before sampling the verification samples should be mixed thoroughly but gently.

Thaw **NOD®** Verification Samples at room temperature (18-25° C) for 30 minutes or until completely thawed. Mix the vial thoroughly by inverting several times, before sampling gently swirls until homogeneous with no visible signs of precipitate. Avoid vigorous shaking. After sampling the Verification samples should be promptly re-capped and stored a 2-8°C. Dispose of at the end of day or upon completion of data collection.

Limitation of Procedure

(a) This product should not be used past the expiration date (b) if there is evidence of microbial contamination in the verification samples or excessive turbidity discard the vial (c) This Product is not intended for use as a standard.

The assay values recovered in the laboratory are method dependent and reflect reagent, method and technique and instrument variations. If methods and / or reagents are changed or modified the resulting assay value may be different.

Assignment of Values

The mean values printed on the circular were derived from replicate analyses on the Piccolo Clinical Chemistry Analyzer and are specific for this lot of Liquid Assayed Chemistry Verification Samples. .
NOVA-ONE Tech Support will provide Data Reduction analysis assistance for control and verification results.

Specific Performance Characteristics

To ensure the reliability and usefulness of the verification samples, the product must be properly handled and stored as described.

Individual donor units used in the preparation of this product have been tested by FDA approved methods for anti-HIV 1 & 2, HBsAg, anti-HCV, HIV-1 antigens and Syphilis and found non-reactive. No test method can offer complete assurance that products derived from human source material will not transmit infectious diseases. Therefore, this product should be considered potentially infective and be treated in the same manner as a patient specimen.

Re-Ordering Information: Verification P/N ALCV-G14033-050 or Control P/N ALPC-G14026-100
By Ordering On Line at **NOVA-ONE.NET**; or **Fax 818-348-9696**



ALL VALUE ASSIGNMENT RANGES CALCULATED USING ABAXIS REAGENTS ON ABAXIS ANALYZERS**Assigned Values and Ranges (Representative Values)****Lot # 464019002 (Containing Vial Lots S1 # 464119002, S2 # 464219002 & S3 # 464319002)**

Expiration Date: 05 - 2021

METHOD: Abaxis Piccolo	Sample 1	Sample 2	Sample 3
Analyte, Units	Mean	Mean	Mean
Albumin g/dL	2.0	3.9	5.4
Alkaline Phosphatase (ALP) U/L	23	1058	1847
ALT/SGPT U/L	27	777	1544
Amylase, Pancreatic U/L	31	1296	2349
Aspartate Aminotransferase (AST/SGOT) U/L	21	863	1705
Bilirubin – Direct mg/dL	0.2	2.3	4.4
Bilirubin – Total mg/dL	0.4	2.9	5.5
BUN (Urea Nitrogen) mg/dL (**Use with low recovery reagent rotors P/N 400-0023, 400-0028 & 400-0031)	6	63	123
BUN (Urea Nitrogen) mg/dL	7	64	123
Calcium Total mg/dL	4.4	11.1	14.9
Carbon Dioxide (CO ₂) mmol/L	30	22	12
Chloride mmol/L	89	111	135
HDL Cholesterol mg/dL	24	49	85
Total Cholesterol mg/dL	76	243	378
Creatine Kinase (CK) U/L	34	1791	3396
Creatinine mg/dL	0.7	8.6	16.7
GGT U/L	14	1273	2363
Glucose mg/dL	29	342	639
Lactate mmol/L	0.31	4.69	9.63
Lactate Dehydrogenase (LDH) U/L	67	409	816
Magnesium mg/dL	0.8	3.8	7.8
Phosphorus mg/dL	16.5	9.6	2.6
Potassium mmol/L	2.1	5.1	7.7
Protein, Total g/dL	3.3	6.4	9.6
Sodium mmol/L	119	141	165
Triglycerides mg/dL	50	202	350
Uric Acid mg/dL	1.1	7.6	15.4

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