CE IVD MICROPOINT

qLabs[®] APTT Test Strips

REF QS-9 Pro Contains: 12 test strips V REF Q-2 Plus qLabs[®] ElectroMeter Plus

For Health Care Professional Use Only

INTENDED USE

The gLabs[®] APTT test strip is designed t provide quantitative determination of Activated Partial Thromboplastin Time (APTT).

qLabs® ElectroMeter Plus instrument using good decisions regarding the need for such fresh capillary whole blood.

for in vitro diagnostic use. It is suitable for and accurate measurement of clotting capacity professional use only.

INTRODUCTION

of the blood in response to the damage of of the intrinsic coagulation pathway, which blood vessels, in order to prevent excessive involves the coagulation factor XII, XI, IX, VIII, bleeding. A clot formed inappropriately in the X, V, II and fibrinogen. It is also used to monitor areas of heart, lung and brain, however, can the effectiveness of heparin therapy. The APTT hinder normal blood flow and may result in is a modification of the Partial Thromboplastin life-threatening events such as stroke and Time (PTT); it can provide a more precise and heart attack. For patients who lose their ability sensitive assay. to properly reabsorb clots and patients who

have low tolerance to clots, anticoagulation medicine (blood thinner) is prescribed. Because these medications may have narrow therapeutic windows and be sensitive to diet and lifestyle, it may be necessary for patients to The aLabs® APTT test is performed on the adjust their dosage regularly. In order to make an adjustment, it is important for the patient to NOTE: The gLabs® APTT test strip is intended know the clotting status of their blood. A quick is critical for the safety and effectiveness of the anticoagulation therapy.

The APTT is a general coagulation test used Blood normally clots to slow down the flow for screening and measuring the functionality

The glabs[®] APTT test strip measures the **REAGENTS** blood's ability to clot which determines the Activated Partial Thromboplastin Time (APTT) on whole blood.

TEST PRINCIPLE

aLabs[®] APTT test strips are used together with gLabs[®] ElectroMeter Plus. The meter automatically detects the insertion of a 1. For in vitro diagnostic use only. Do not aLabs[®] APTT test strip and heats the strip up to a preset operating temperature. After a drop of blood is applied to the strip, the capillary channels carry the blood to the reaction zones where the blood mixes with pre-printed reagents and starts to coagulate. 3. Use only fresh capillary whole blood. Each strip contains two reaction zones: 4. Never add blood to a test strip after the test one for APTT testing, another for on-board QC testing. Each reaction zone contains one pair of metallic electrodes, to which a constant voltage is applied by the meter. As the coagulation of the blood proceeds, the 6. Do not move the meter during a test. current monitored across the two electrodes changes. The meter detects the change of the current in the reaction zone and identifies a clot endpoint. Based on an analysis of the test data, the resultant clot endpoint is then The health status of the patient may affect converted to a value that is more familiar to the test. Please take this into consideration the clinician.

Each test strip contains a standardized amount of phospholipid, particulate activator, stabilizers and buffers on the test zone. Individual test strip is packaged in a pouch with one desiccant bag.

ECAUTIONS & WARNINGS

- take internally.
- 2. Follow proper infection control guidelines for handling all blood specimens and related items
- has begur
- 5. Do not use strong repetitive pressure to collect the sample.

7. Do not use test strip that past their marked expiration date, or which have been improperly stored

before making a therapeutic judgment based on the test results. Failure to do so may cause serious consequences.

STORAGE & HANDLING

- aLabs[®] APTT test strips can be stored i the refrigerator at 2°C to 8°C or at room temperature (below 32 °C) until the expiration date. Do not freeze.
- Store strips in their original foil pouch until ready to use.
- If refrigerated, allow the sealed pouch t equilibrate to room temperature for 5 minutes before opening it for testing.
- Use the test strip within 10 minutes after opening the foil pouch.

TEST PROCEDURE

- When powered on, the meter will automatically enter the Test Mode and prompt you to insert a test strip.
- I Insert a test strip into the test strip guide on the meter. Remove a fresh test strip from foil pouch. Insert it into the test strip guide so that the electrode end goes in first On the end of the strip you should be able to read the word "APTT" appearing from left to riaht.
- 2. Enter the patient ID. The patient ID car be entered either via user's manual input or the built-in barcode scanner. Please refer to meter's user manual to correctly enter a patient ID.
- Enter the strip code and the lot numbe When you insert a fresh test strip, the meter will prompt you to enter a strip code and lot

number. If the code and lot number already match with the code and lot on the test strip pouch, simply press the "OK" button. If it does not match, you can either use the left arrow button to select the "PEN" symbol to manually input the required information or use the right arrow button to select the "BarCode" symbol to input the required information with the built-in barcode scanner. Please follow the user's manual to ensure the code and lot number are entered correctly.

- 4. Confirm the strip code and the lot number. Once you have input the strip code and the lot number, the meter will automatically check whether the strip is expired. If it is expired, an error will be reported. You will not be able to test with an expired strip.
- Always match the strip code. lot number and test item name on the display with those on the strip pouch. Failure to do so may cause inaccurate results.
- 5. Wait for the meter to warm up. Once the strip code and lot number have been confirmed, the meter will automatically warm up for the test. When it is ready to perform a test, the meter will beep and prompt the user to apply a blood sample.
- 6. Obtain a fingerstick blood sample. It is important that you use the correct technique to obtain the right type and amount of blood

sample. If the procedure is not followed, it can cause inaccurate results.

- 6.1 Increase blood circulation by
- Warming the hand by soaking in warm water
- Warming the hand with a heating pad or hand warmer
- Massaging the finger gently
- Holding the hand below the heart
- 6.2 Identify a site on the finger to puncture:
- On one of the middle fingers of either hand
- Near the top of the finger on either side
- Away from any calluses or scars



6.3 Clean the selected area with 70% clean isopropyl alcohol, or an alcohol pad. Dry thoroughly with cotton or gauze.



6.4 Puncture the finger following the instructions for the lancet that you are using.



6.5 Apply gentle, continuous pressure until a large, hanging drop of blood forms.

7. Add blood sample. Apply the blood directly on the sample well of the strip. The minimum sample volume is 10 µL.



- and time.
- 9. Finish the test. Discard the used lancet and test strip into an puncture resistant waste container. All blood samples should be regarded as potentially hazardous. The meter will turn itself off after 5 minutes if no buttons are pressed

Material Provided

• gLabs[®] APTT test strips

Material Required, but Not Provided

- aLabs[®] ElectroMeter Plus
- Alcohol pads and gauze
- Lancet device
- Puncture resistant waste container

SAMPLE COLLECTION AND HANDLIN

H47-A) guidelines to obtain blood samples for outside of this range may give unusual APTT testing.

RESULTS

the test method, it is recommended that the blood to clot through the normal means. same method be used whenever doing routine patient monitoring.

Interfering metabolites: The gLabs[®] system is validated to work in the presence of unusually Normal Range: high concentrations of hemoglobin, bilirubin, Results for normal blood were determined or triglycerides (see LIMITATIONS section below). Presence of these metabolites at by testing 20 subjects who were not taking anticoagulant medication. The ranges found concentrations above these limits may lead to prolonged clot times.



8. Perform APTT test. After adding were 31 – 42 sec. Due to many variables that blood sample, the system will start test affect clotting times, each individual laboratory automatically. The test results will appear as should establish relevant normal range for its APTT value, QC value, along with the date respective patient population.

Therapeutic Range:

Therapeutic heparin levels of 0.2 - 0.4 U/mL should give 1.5 - 2.5 times the mean normal APTT values. Due to many variables that affect clotting times, each individual laboratory should establish relevant APTT therapeutic ranges for its respective patient population.

Unexpected Results:

When the ElectoMeter displays an APTT result outside of the expected therapeutic range, it may or may not be due to an unusual clinical situation.

What causes unexpected results:

Hematocrit: The gLabs[®] system is validated to work reliably with blood having hematocrit Follow the institutional and NCCLS (H21-A3, values between 30% and 55%, Blood samples values.

Interfering antibodies: Conditions (such as Lupus) that produce antiphospholipid Since APTT results are expected to vary with antibodies may interfere with the ability of

Medications: Certain medications, including Precision: both prescription and over the counter, may interfere with oral anticoagulants, and may lead to an anomalous APTT result.

may interfere with anticoagulant therapy.

Diet: Oral anticoagulants may be sensitive to food, alcohol, and nutritional supplements. APTT: normal donor

What to do:

Whenever you encounter an unexpected result, please repeat the test with a fresh aLabs[®] test strip. If the result is seen a second time, please consult immediately with vour local distributor.

PERFORMANCE CHARACTERISTICS

Normal Range:

According to CLSI C28-A2, the normal range of aLabs[®] APTT tests were evaluated using fresh fingerstick whole blood from normal volunteer donors which is 31-42 seconds (n=20).

NOTE: Each institution should establish its own normal range and target range of therapeutic anticoagulation based on its patient population.

NOTE: gLabs[®] APTT values greater than 420 will be reported as ">420" and may indicate excessive blood coagulation activation, possibly due to specimen contamination upon sample collection or processing and should be repeated.

The precision of the APTT test was evaluated using fresh fingerstick whole blood from normal volunteer donor and heparinized fresh Disease state: Certain medical conditions venous whole blood from normal volunteer donor.

		Ν	Mean	S.D.	CV
			(sec)	(sec)	(%)
Day 1	Lot 1	6	33.3	1.4	4.1
Day 2	Lot 2	6	35.5	1.2	3.5
Day 3	Lot 3	6	35.2	0.7	2.1

APTT: heparinized normal donor

		Ν	Mean	S.D.	CV
			(sec)	(sec)	(%)
Day 1	Lot 1	6	87.9	6.1	6.9
Day 2	Lot 2	6	92.2	1.6	1.7
Day 3	Lot 3	6	78.8	3.8	4.8

Accuracy:

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Regression analysis of the qLabs® APTT test compared to a central laboratory analyzer (n=203).



Heparin sensitivity:

qLabs[®] APTT test strips are sensitive to the presence of therapeutic levels (0.2 - 0.4 U/mL by protamine titration) of heparin in the sample. The sensitivity curves below are obtained by adding increasing quantities of unfractionated porcine heparin to aliquots of a normal donor blood.



NOTE: The heparin sensitivity curve is unique to each patient and can vary due to many variables (e.g. different source of heparin being used). The curves are intended to serve as examples only.

LIMITATIONS

- 1. The gLabs[®] system is designed to use fresh capillary whole blood. Plasma or anticoagulated whole blood should not be used.
- 2. The drop of blood must be a minimum of 10 µL.
- 3. Hematocrit ranges between 30% and 55% will not affect test results.
- 4. In vitro studies show no significant effect in blood samples containing up to 10 mg/ dL of bilirubin, 100 mg/dL of hemoglobin.
- 5. The gLabs[®] APTT test strips are validated to perform at temperatures in the range 10 to 35°C, and 10 to 90% RH (relative humidity). This includes a 10 minute out of pouch exposure of the strips at these conditions.
- 6. As with all diagnostic tests, gLabs[®] APTT test results should be scrutinized in light of a specific patient's condition and anticoagulant therapy. Any results exhibiting inconsistency with the patient's clinical status should be repeated or supplemented with additional test data or repeated with other testing methods.

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PERFORMANCE SPECIFICATIONS

Category	Performance Specification		
Intended sample	Fresh fingerstick		
Operating temperature range	10 to 35°C		
Operating humidity range	10 to 90% RH		
Out-of-pouch stability	10 minutes		
Shelf life	12 months (2 - 32°C, in pouch with desiccant)		
Measurable Range	20-420 sec		
Accuracy	Regression analysis vs. central lab: Correlation: $r \ge 0.90$		
Precision	$CV \le 7\%$		
Hematocrit range	30% to 55%		
Time to result	7 minutes		
Sensitivity to heparin	APTT test is sensitive up to 0.6 U/mL for unfractionated heparin		

ADDITIONAL INFORMATION

If you have any questions regarding the use of this product, please call your local representative/distributor, or our customer service at +86 755 86296766.

SYMBOLS

Symbols	Explanation		
IVD	In vitro diagnostics		
***	Name and Address of Manufacturer		
CE	CE Mark		
EC REP	European Authorized Representative		
X	Temperature limitation		
LOT	Lot number		
\Box	Expiry Date		
2	Do not reuse		
REF	Catalogue number		
Σ $\Sigma = n$	Contains sufficient for n tests		
	Caution. Read carefully		



MICROPOINT



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