

Blood Glucose Test Strips (Glucose Dehydrogenase FAD-Dependent) Package Insert

ΕN

> Specification

Model: SM211

Catalog: SM2111011, SM2111012, SM2111014, SM2111015, SM2111019

Intended Use

ACCUGENCE® Blood Glucose Test Strips (Glucose Dehydrogenase FAD-dependent) are used with ACCUGENCE® Multi-Monitoring Meter to quantitatively measure the glucose concentration in fresh capillary whole blood samples. ACCUGENCE® Blood Glucose Test Strips and applicable meter are intended to be used only outside the body (in vitro diagnostic use) by people with diabetes at home, as an aid to monitor the effectiveness of diabetes control. They are only for self-testing, not for near-patient testing.

The test strips shall not be used for screening, diagnosis, or aid to diagnosis of diabetes. They are not automated.

> Test Principle

When blood sample is applied to the end tip of the test strip, the sample is then automatically absorbed into the reaction cell where the reaction takes place. A transient electrical current is formed during the reaction and measured by the meter. The glucose results is then calculated based on this current and is shown on the meter display.

> Composition

Each test strip contains reactive and non-reactive chemicals. These chemicals are: Glucose Dehydrogenase FAD-Dependent <15 IU, Mediator <100 μg , Non-reactive Ingredients (Buffer, etc.).

Each test strip vial contains a drying agent.

> Storage and Handling

- Store test strips in a cool dry place at 2-35 °C (36-95 °F) and 10-90% relative humidity.
 Do not freeze. Keep away from heat and direct sunlight. Exposure to temperatures and/or humidity outside the storage limits may cause inaccurate readings.
- The unopened expiration date is printed on the vial.
- **Note:** All expiration dates are printed in Year-Month-Day format. 2020-01-01 means January 1th, 2020.
- A new vial of test strips may be used for 6 months after first opening. Write the opened expiration date on the vial label when you open a new vial.
- Do not use your test strips beyond the unopened expiration date or the opened expiration date whichever comes first. Discard any unused test strips beyond the expiration date, because they may cause inaccurate results.
- Store unused test strips only in the original vial with the cap closed tightly. Do not transfer the test strips to any other container.
- Do not store the meter, the test strips or control solutions near bleach or cleaners with bleach
- . Open the vial only when taking out a test strip for use.
- Close the vial cap tightly immediately after removing a test strip. Use each test strip as soon as you take it out of the vial.
- Use the test strips at temperatures between 5-45 °C (41-113 °F).
- Use the test strips at 10%-90% relative humidity. Do not store or use the test strips in high heat and moisture areas such as the bathroom or kitchen.
- Make sure your meter and test strips are about the same temperature before you test.
- Do not use test strips that are torn, bent, damaged, altered, or contaminated.
- Do not use test strips from a vial that is damaged or left open to air.
- Test strips are for single use only. Do not reuse test strips.
- Repeated insertion and removal of a test strip into the meter strip port may result in reading errors.

Precautions

- Do not use new test strips if the vial is open or damaged in any way. This could lead to
 error messages or inaccurate results.
- Matching the code number on the meter to the code number on the test strip vial is
 essential to obtain accurate results. Refer to the User's Manual for the detailed
 instructions about coding.
- Use universal blood precautions when handling, and disposing of, blood glucose
 monitoring materials. All patient samples and materials with which they come in
 contact are considered biohazards and should be handled as if capable of transmitting
 infection even after you have performed cleaning and disinfection. Follow proper
 precautions in accordance with all local regulations when disposing of all materials.
- . Do not use a lancet that has been used by others.
- Wash your hands thoroughly with soap and water after handling the meter, lancing device or test strips.
- Keep your meter and lancing device clean.
- Apply sample only to the tip of the test strip. Do not apply to the top of the test strip.
 This may result in a false reading.
- $\bullet \hspace{0.4cm}$ Do not put the used test strips back in the vial after taking a test.
- Keep the test strip vial away from children and animals.
- Always consult your doctor before making any changes to your treatment plan.
- Any serious incident shall be reported to manufacturer and competent authority of the Member State in which user and/or patient is established.

Materials Provided

Test Strips
 Calibration Chip
 Package Insert

Materials Required but Not Provided

• Multi-monitoring (Model: PM900) meter • Sterile Lancets • Lancing Device • Control Solution

➤ Instructions for Use

See your User's Manual for complete instructions for blood sample collection before use.

- 1. Open the cap of the test strip vial, remove a test strip. Reclose the vial cap immediately to protect the unused test strips from humidity.
- 2. Run the test following the instructions in your User's Manual.
- The test result will be shown on the meter display window. This result should fall within the target range. Your doctor should recommend your target range. If your results are higher or lower, ask your doctor what to do.

Explanation of Test Results

- The ACCUGENCE® Blood Glucose Test Strips (Glucose Dehydrogenase FAD-Dependent) gives accurate blood glucose readings within the range of 0.6-33.3 mmol/L (10-600 mg/dl).
- "LO" means that your meter has determined that you blood glucose result is lower than 0.6 mmol/L (10 mg/dL).
- "HI" means that your meter has determined that you blood glucose result is higher than 33.3 mmol/L (600 mg/dL).

> Range of Expected Values

Blood glucose monitoring requires the help of a doctor. Together you can set your own range of expected blood glucose values, arrange your testing times, and discuss the meaning of your blood glucose results.

Expected blood glucose levels for people without diabetes1:

Time	Range, mg/dL	Range, mmol/L
Fasting and Before Meals	70 - 100	3.9 - 5.6
2 Hours After Meal	Less than 140	Less than 7.8

Checking the System

See your user's manual for how to take good care of your meter.

Do a quality control test to make sure that the meter and test strips are working well together. Follow the control test procedure in your User's Manual. Use only ACCUGENCE® Glucose Control Solutions. Contact Customer Support for information on purchasing control solution.

There are 3 levels for ACCUGENCE® Glucose Control Solutions. When a control test completed, determine whether the test result is within the range printed on the test vial. If the obtained results fail outside this range, repeat the control test.

CAUTION: If your quality control test result falls outside the control range shown on the test strip vial, DO NOT use the system to test your blood. The system may not be working properly. If you cannot correct the problem, contact your local distributor for help.

Limitation

- Do not use the meter in any way that is not specified by the manufacturer. Otherwise, the system might not work the way it is supposed to.
- The test strips are for testing fresh capillary whole blood. Do not use with serum or plasma samples.
- The test strips should not use for the testing of newborns.
- Very high (above 70%) and very low (below 10%) hematocrit levels can cause false results. Talk to your doctor to find out your hematocrit level.
- Fatty substances (triglycerides up to 33.9 mmol/L (3000 mg/dL) or cholesterol up to 12.9 mmol/L (500 mg/dL)) have no significant effect on test results.
- Ascorbic acid (vitamin C), acetaminophen, uric acid, salicylates, and other reducing substances when occurring in normal blood or normal therapeutic concentrations do not significantly affect results. However, abnormally high concentrations in blood may cause inaccurately high results.
- The test strips may be used at altitudes up to 10,000 feet (3,048 meters).
- If peripheral circulation is impaired, collection of capillary blood is not advise as the
 results might not be a true reflection of the physiological blood glucose level. This may
 apply in the following circumstances: Severe dehydration as a result of diabetic
 ketoacidosis or due to hyperglycaemic hyperosmolar non-ketotic syndrome,
 hypotension, shock, decompensated heart failure NYHA Class IV, or peripheral arterial
 occlusive disease.
- The test strips are not recommended for use in critically ill patients.

Performance Characteristics

Calibration and Traceability: The ACCUGENCE® Blood Glucose Test Strips (Glucose Dehydrogenase FAD-Dependent) are calibrated by using YSI Model 2300 Glucose Analyzer, a laboratory reference instrument. It is traceable to NIST SRM 917c. The test results are calibrated to be plasma equivalent.

System measurement range: 0.6-33.3 mmol/L (10-600 mg/dL) Required sample size: $0.7 \mu L$

Tested time: 5 seconds
Measurement Repeatability

The evaluation was conducted with 10 meters, 3 strip lots, and 5 samples with different glucose concentrations, and 10 measurements were performed with each combination of meter, strip lot and sample. The results are shown below.

	Diood Julipic 1	2 3 4	<u> </u>
N 300 300 300 300	N 300	300 300 300	300

Grand Mean	2.11 mmol/L	4.15 mmol/L	7.03 mmol/L	10.61 mmol/L	17.70 mmol/L
Grand Weari	(37.9 mg/dL)	(74.6 mg/dL)	(126.6 mg/dL)	(190.9 mg/dL)	(318.6 mg/dL)
Pooled SD	0.05 mmol/L	0.07 mmol/L	0.11 mmol/L	0.16 mmol/L	0.26 mmol/L
Pooled SD	(0.98 mg/dL)	(1.25 mg/dL)	(2.06 mg/dL)	(2.83 mg/dL)	(4.69 mg/dL)
Pooled CV (%)	2.6	1.7	1.6	1.5	1.5

Intermediate Measurement Precision

The evaluation was conducted with one measurement of each sample per day and was conducted with 10 meters, 3 strip lots and control solutions at 3 glucose concentrations over 10 days. The results are shown below.

Control Solutions	1	2	3	
N	300	300	300	
Grand Mean	2.05 mmol/L (37.0 mg/dL)	6.24 mmol/L (112.3 mg/dL)	17.72 mmol/L (318.9 mg/dL)	
Pooled SD	0.05 mmol/L (0.88 mg/dL)	0.10 mmol/L (1.72 mg/dL)	0.22 mmol/L (3.97 mg/dL)	
Pooled CV (%)	2.4	1.5	1.2	

System Accuracy

System accuracy was evaluated with fresh fingertip capillary blood samples by comparing glucose measurements from the ACCUGENCE® Multi-Monitoring Meters tested with ACCUGENCE® Blood Glucose Test Strips (Glucose Dehydrogenase FAD-Dependent) to the reference glucose values obtained by YSI Model 2300 Glucose Analyzer, and the evaluation was conducted with 100 different subjects taking duplicate measurements from each of 3 strip lots by trained operators. The following results were obtained.

Data across the entire reportable range meets the acceptance criteria of \geq 95 % of the measured glucose values within either \pm 0.83 mmol/L (\pm 15 mg/dL) of the reference values at glucose concentrations <5.55 mmol/L (<100 mg/dL) or within \pm 15 % at glucose concentrations \geq 5.55 mmol/L (\pm 100 mg/dL).

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System accuracy results for glucose concentration <5.55 mmol/L (<100 mg/dL)						
Within ±0.28 mmol/L (Within ±5 mg/dL)	Within ±0.56 mmol/L (Within ±10 mg/dL)	Within ±0.83 mmol/L (Within ±15 mg/dL)				
98/144 (68.1%)	133/144 (92.4%)	144/144 (100.0%)				
System accuracy results for glucose concentration ≥5.55 mmol/L (≥100 mg/dL)						
Within ± 5%	Within ± 10%	Within ± 15%				
223/456 (48.9%)	401/456 (87.9%)	456/456 (100.0%)				
	System accuracy results for glucose concentrations between 1.4 mmol/L (25.6 mg/dL) and 32.9 mmol/L (592.7 mg/dL)					
Within ±0.83 mmol/L (± 15 mg/dL) or ±15%						
600/600 (100.0%)						

User Performance Evaluation

User performance was evaluated by comparing the accuracy of capillary blood glucose values measured by lay persons to the capillary blood glucose values measured by YSI Model 2300 Glucose Analyzer, a laboratory instrument.

The study evaluating glucose values from fingertip capillary blood samples obtained by 100 lay persons showed the following results: 99.0% within ±0.83 mmol/L (±15 mg/dL) of the reference values at glucose concentrations <5.55 mmol/L (<100 mg/dL) or within ±15% of the reference values at glucose concentrations ≥5.55 mmol/L (≥100 mg/dL).

Reference

 ${\bf 1.\ Standards\ of\ Medical\ Care\ in\ Diabetes-2017.\ American\ Diabetes\ Association.}$

➢ Index of System

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i	Consult instructions for use	X	Use by	CODE	Code Number
IVD	For <i>in vitro</i> diagnostic use only	LOT	Lot Number	REF	Catalog #
+210-43510	Store between 2-35°C (36-95 °F)	1	Manufacturer	Ø	Do not reuse
Σ	Contains sufficient for <n> tests</n>		Authorized Representative		6 months expiry date from the date of first open vial
CE	CE Marking		Unique Device Identifier	Į,	For self-testing

> The summary of safety and performance

Intended users can log in to the European database on medical devices (Eudamed) to request the summary of safety and performance (SSP) of the device or contact the manufacturer to obtain it.



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