

LETTER OF AUTHORIZATION

We, LumiQuick Diagnostics, Inc., having a registered office at 2946 Scott Blvd, Santa Clara, CA 95054, USA assign SRL SANMEDICO, having a registered office at A. Corobceanu street 7A, apt. 9, Chişinău MD-2012, Moldova, as our authorized representative in correspondence with the conditions of directive 98/79/EEC.

We declare that the company mentioned above is authorized to register, notify, renew or modify the registration of medical devices on the territory of the Republic of Moldova.

This letter is valid through December 31, 2023 and will automatically renewed upon the agreement of both companies. Should you have questions, please contact us.

Best regards,

Charles Yu

President

Date: January 19, 2022



Australia | Canada | China | Japan | The Netherlands | United States

E M E R G O 🥑 E U R O P E

24 March 2009

Mr. Jeff Wang LumiQuick Diagnostics, Inc. 2946 Scott Blvd. Santa Clara, CA 95054

Dear Mr.Jeff Wang:

I am writing to inform you that today, we have notified by registered mail the Competent Authority in the following countries:

Austria	Bulgaria	Cyprus	Czech Republic	Denmark	Estonia			
Finland	France	Germany	Greece	Hungary	Iceland			
Ireland	Italy	Latvia	Liechtenstein	Lithuania	Luxembourg			
Malta	The Netherlands		Norway	Poland	Portugal			
Romania	Slovakia	Slovenia	Spain	Sweden	Switzerland			
United Kingdom								

With this notification, LumiQuick Diagnostics, Inc. has met the requirements of the In-vitro Diagnostics Directive, 98/79/EC for the following devices:

- Adeno/Rota Virus
- Cardiac Marker
- Dengue IgG/IgM Combo (registered only in Italy and The Netherlands)
- Drugs of Abuse
- Fecal Occult Blood (registered only in Italy and The Netherlands)
- H. Pylori Ab/Ag
- HCG
- Legionella (registered only in Italy and The Netherlands)
- LH (registered only in Italy and The Netherlands)
- Strep A (registered only in Italy and The Netherlands)

As of today and without any further notice from the respective Competent Authorities, LumiQuick Diagnostics, Inc. can consider the respective devices and Authorized Representative as officially registered.

If you have any questions, please do not hesitate to contact me.

Yours sincerely,

Rene van de Zande President & CEO Emergo Europe



EmergoEurope.com



Declaration of Conformity

Product name	Mode//number		
Fecal Occult Blood Test Device	ces		
QuickProfile Fecal Occult Blo QuickProfile Fecal Occult Blo		72001 72006	
MANUFACTURER			
Name of company	Address	Representative	
LumiQuick Diagnostics, Inc.	2946 Scott Blvd. Santa Clara, CA 95054 USA	Jeff Wang	
AUTHORIZED REPRESENTATIVE			
Name of company	Address	Telephone/email	
Emergo Europe	Prinsessegracht 20 2514 AP The Hague, Netherlands	+31.70.345.8570 - phone +31.70.346.7299 - fax europe@emergogroup.com	
CONFORMITY			
ASSESSMENT			
Device classification	Route to compliance	Standards applied	
Class: Self-Certify	Annex III of IVDD 98/79/EC	ISO 13485:2003	

LumiQuick Diagnostics, Inc. declares that the above mentioned products meet the provision of the Council Directive 98/79/EC for In Vitro Diagnostic Medical Devices and Directive 98/79/EC as transposed in the national laws of the Member States.

COMPANY REPRESENTATIVE: Jeff Wang

TITLE: Quality Systems Manager

SIGNATURE:

nico

DATE: 28/04/2017





Certificate of Registration

QUALITY MANAGEMENT SYSTEM - ISO 13485:2016

This is to certify that:

LumiQuick Diagnostics, Inc. 2946 Scott Blvd Santa Clara California 95054 USA

Holds Certificate No:

FM 574919

and operates a Quality Management System which complies with the requirements of ISO 13485:2016 for the following scope:

The design, development, manufacture and distribution of in vitro diagnostics test kits and reagents used in the diagnosis and management of disease status, including Infectious Diseases tests, Drugs of Abuse tests, Cardiac Monitor tests, Cancer Marker tests, Fertility Hormone tests, ELISA tests & Urine Chemistry tests.

jan CS

For and on behalf of BSI:

Gary E Slack, Senior Vice President - Medical Devices

Original Registration Date: 2011-10-20 Latest Revision Date: 2020-08-31





Effective Date: 2020-10-20 Expiry Date: 2023-10-19

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...making excellence a habit."

This certificate remains the property of BSI and shall be returned immediately upon request. An electronic certificate can be authenticated <u>online</u>. Printed copies can be validated at www.bsigroup.com/ClientDirectory To be read in conjunction with the scope above or the attached appendix.

Americas Headquarters: BSI Group America Inc., 12950 Worldgate Drive, Suite 800, Herndon, VA 20170-6007 USA A Member of the BSI Group of Companies.



Quick PROFILE[™] FECAL OCCULT BLOOD TEST CARD

FOR THE QUALITATIVE ASSESSMENT OF HUMAN HEMOGLOBIN IN FECES



Catalog No.: 72001

For In-Vitro Diagnostic Use

INTENDED USE

QuickProfile[™] Fecal Occult Blood test is a qualitative test that detects human hemoglobin in human fecal specimens. The test is a visual one step, in-vitro assay. It is intended for professional use to help diagnose gastrointestinal bleeding.

SUMMARY AND EXPLANATION

Colorectal cancer is the third most common cancer in the world. "Fecal occult blood" is generally defined as a blood loss of less than 50 mL/d. The appearance of occult blood in human fecal specimen is often associated with gastrointestinal diseases which might cause colorectal cancer if not treated promptly and properly. The traditional guaiac-based method lacks sensitivity and specificity, and has diet restrictions prior to the testing.

QuickProfile[™] Fecal Occult Blood Test uses the technology of immunochromatographic sandwich assay. The test is more sensitive and more specific than the traditional guaiac assay. It is easier to interpret the result. In addition, unlike the guaiac assays, the accuracy of the test is not affected by the diet of the patients.

TEST PRINCIPLE

QuickProfile[™] Fecal Occult Blood Test is composed of two units, a fecal collection tube and a test device. A fecal specimen is collected in the collection tube containing sample extraction buffer, and then added to the test device. When sample is added to sample pad, it moves through the conjugate pad and mobilizes the gold anti-h hemoglobin antibody conjugate that is coated on the conjugate pad. The mixture moves along the membrane by capillary action and reacts with anti-h hemoglobin antibody that is coated on the test region. If h hemoglobin is present at levels of 50 ng/mL or greater, the result is the formation of a colored band in the test region. If there is no h hemoglobin in the sample, the area will remain colorless. The sample continues to move to the control area where goat anti-mouse IgG antibody will capture gold-antibody conjugate to form a pink to purple color, indicating the test is working and the result is valid.

MATERIAL PROVIDED

- QuickProfile[™] Fecal Occult Blood Test device Test zone: contains mice monoclonal anti-hemoglobin antibody. Control zone: contains goat anti-mouse IgG antibody. Conjugate pad: contains gold-mice monoclonal anti-hemoglobin antibody conjugate.
- 2. Fecal specimen collection tube
- The collection tube contains 2 ml of buffer.
- 3. Instructions for use

MATERIALS REQUIRED BUT NOT SUPPLIED

Timer or clock.

STORAGE

- 1. Store the test device in the original sealed pouch and the fecal specimen collection tube at 4 to 30°C. Do Not Freeze.
- 2. The expiration date given was established under these storage conditions.
- 3. The test device should remain in its original sealed pouch until ready for us.
- 4. The device is designed for single use. Once the pouch is opened, the device must be tested as soon as possible and cannot be reused.

72001

9-16-2015

PRECAUTIONS

- 1. For in-vitro diagnostic use only.
- 2. Do not use product beyond the expiration date.

DCR 15-052 5044 E3R2 3. Handle all specimens as potentially infectious.

PATIENT PREPARATION

- 1. Specimen should not be collected during or within three days of a menstrual period, or if the patient suffers from bleeding hemorrhoids or blood in the urine.
- 2. Alcohol, aspirin and other medications, taken in excess, may cause gastrointestinal irritation resulting in occult bleeding. Such substances should be discontinued at least 48 hours prior to testing.
- 3. Dietary restrictions are not necessary.

SPECIMEN COLLECTION WITH SAMPLE TUBE TYPE I

- 1. Stool specimens can be collected at any time of the day.
- 2. Collect a random sample of feces in a clean, dry receptacle.
- 3. Unscrew the bottom cap (red end) of the collection tube and remove the applicator stick.
- 4. Insert the stick into the fecal specimen at several different sites.
- 5. Insert the sampled applicator back to the tube and tighten the bottom (red end) securely. The hold that only allows the stick goes through will prevent the access sample from getting into the tube.
- Shake the tubes with bottom cap (red end) vigorously for about 5 seconds to release and disperse the stool sample into the collection buffer.





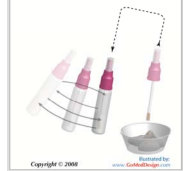
Specimen collection Steps 3 and 4

Specimen collection Steps 5 and 6

SPECIMEN COLLECTION WITH SAMPLE TUBE TYPE II

- 1. Stool specimens can be collected at any time of the day.
- 2. Collect a random sample of feces in a clean, dry receptacle.
- 3. Unscrew the cap (red end) of the collection tube and remove the applicator stick.
- Insert the stick into the fecal specimen at several different sites.
- Insert the sampled applicator back to the tube and tighten the cap securely.
- Shake the tubes vigorously for about 5 seconds to release and disperse the stool sample into the collection buffer.





Specimen collection Steps 3 and 4

Specimen collection Steps 5 and 6

SPECIMEN STABILITY

The sample can be stored at room temperature (8 - 30°C) up to seven days if not immediately tested. If the condition allowed, the sample can also be refrigerated ($2 - 8^{\circ}$ C) for better storage.

QUALITY CONTROL

- It is recommended that a positive control, with a level between 50-200 ng/mL h hemoglobin and a negative 1. control, 0 ng/mL h hemoglobin, be used. Control materials, which are not provided with this test kit, are commercially available.
- 2. The control band is an internal reagent and procedural control. It will appear if the test has been performed correctly and the reagents are reactive.

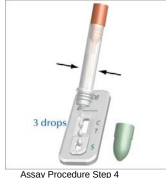
You should always follow local, state and federal guidelines for running QC.

PROCEDURE

- 1. Bring all materials and specimens to room temperature (8-30°C).
- 2. Remove the test card from the sealed foil pouch.
- 3. 3.1 If collection tube Type I is used, remove the tip protection cap (green). Holding the tube upright with tip pointed toward the direction away from the test performer, Snap off the tip.
- 3.2 If collection tube Type II is used, hold the tube upright with tip pointed toward the direction away from the test performer, Snap off the tip.
- Hold the tube in a vertical position over the sample well of the test card and deliver 3 drops (120-150 µL) of 4. sample into the sample well marked as "S" on the cassette.
- 5. Read the results between 3 and 10 minutes.

Note: Results read after 10 minutes may not be accurate.





Assay Procedure Step 3.1







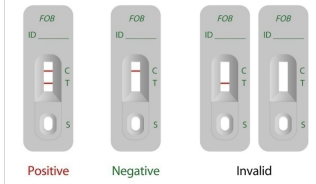
If two colored bands are visible within 3 minutes, the test result is positive and valid. Note: Specimens containing very low levels of h hemoglobin may develop two colored bands over 10 minutes.

DCR 15-052 5044 E3R2

Negative:

If test area has no colored band and the control area displays a colored band, the result is negative and valid. Invalid result:

The test result is invalid if a colored band does not form in the control region. The sample must be retested using a new test device.



LIMITATIONS OF THE PROCEDURE

- A number of conditions, as mentioned in "Patient Preparation", can cause false positive results. 1.
- As with all diagnostic tests, a definitive clinical diagnosis should not be based on the result of a single test, 2. but should only be made by the physician after all clinical and laboratory findings have been evaluated.

PERFORMANCE CHARACTERISTICS

A. Sensitivity:

The analytical sensitivity of the test is 50 ng/mL h hemoglobin or 12.5 µg h hemoglobin/g feces.

B. Specificity:

The test is specific to human hemoglobin. Samples containing the following substances were tested on both positive and negative controls with no effect on test results.

Substances	Concentrations
Beef hemoglobin	1 mg/mL
Chicken hemoglobin	1 mg/mL
Goat hemoglobin	1 mg/mL
Horse hemoglobin	1 mg/mL
Pork hemoglobin	1 mg/mL
Rabbit hemoglobin	1 mg/mL
Duck hemoglobin	1 mg/mL
Dog hemoglobin	1 mg/mL
Horseredish peroxidase	1 mg/mL

C. Interference testing:

The following substances were added to h hemoglobin free and 50 ng/mL controls. No interference was found with any of the substances at the following concentrations:

Acetaminophen	20 mg/dL
Acetylsalicyclic acid	20 mg/dL
Ampicillin	40 mg/dL
Ascorbic acid	40 mg/dL
Atropine	40 mg/dL
Caffeine	40 mg/dL
Gentisic acid	40 mg/dL
Glucose	2000 mg/dL
Human albumin	2000 mg/dL
Urea	4000 mg/dL
Uric acid	10 mg/dL

REFERENCES

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- 5. Taranen, M.J., et al. "Immunological detection of fecal occult blood in colorectal cancer", Br. J. Cancer, Vol. 49 141, (1984).

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~ 30°C

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