

## LIMING BIO

Diagnostics are ASSURED StrongStep®

## **Neisseria Gonorrhoeae Antigen Rapid Test**

(Dyed latex immunochromatography)

REF	500020	Specimen: Swab
Language: English		Version: 02
Effective Date: 2011-12		

For professional in vitro diagnostic use only.

#### **INTENDED USE**

StrongStep® Neisseria Gonorrhoeae Antigen Rapid Test is an immunochromatographic assay for the qualitative presumptive detection of *Neisseria gonorrhoeae* in female endocervical swab and male urethral swab specimens. This kit is intended for use as an aid in the diagnosis of Gonorrhea.

#### INTRODUCTION

Gonorrhea is a sexually transmitted disease caused by the bacterium Neisseria gonorrhoeae. Gonorrhea is one of the most common infectious bacterial diseases and is most frequently transmitted during sexual intercourse, including vaginal, oral and anal sex. The causative organism can infect the throat, producing a severe sore throat. It can infect the anus and rectum, producing a condition called proctitis. With females, it can infect the vagina, causing irritation with drainage (vaginitis). Infection of the urethra may cause urethritis with burning, painful urination, and a discharge. When women have symptoms, they often note vaginal discharge, increased urinary frequency, and urinary discomfort. But there are  $5\% \sim 20\%$  of men and 60% of women patient that do not show any symptoms. Spread of the organism to the fallopian tubes and abdomen may cause severe lower-abdominal pain and fever. The average incubation for Gonorrhea is approximately 2 to 5 days following sexual contact with an infected partner. However, symptoms may appear as late as 2 weeks. A preliminary diagnosis of Gonorrhea can be made at the time of examination. In women, Gonorrhea is a common cause of pelvic inflammatory disease (PID), PID can lead to internal abscesses and long-lasting, chronic pelvic pain. PID can damage the fallopian tubes enough to cause infertility or increase the risk of ectopic pregnancy. The swab of urethral or endocervical discharge can be taken and tested using StrongStep® Neisseria Gonorrhoeae Antigen Rapid Test.

## PRINCIPLE

StrongStep® Neisseria Gonorrhoeae Antigen Rapid Test detects Neisseria gonorrhoeae through visual interpretation of color development on the internal strip. Gonococcal Antigen-specific polyclonal antibody is immobilized on the test region of the membrane. During testing, the specimen reacts with monoclonal anti-Gonococcus antibodies conjugated to colored particles and precoated onto the sample pad of the test. The mixture then migrates through the membrane by capillary action and interacts with reagents on the membrane. If there are sufficient Gonococcal antigens in the specimen, a colored band will form at the test region of the membrane. The presence of this colored band indicates a positive result, while its absence indicates a negative result. The appearance of a colored band at the control region serves as a procedural control, indicating that the proper volume of specimen has been added and membrane wicking has occurred.

## KIT COMPONENTS

20 Individually packed test devices	Each test contains colored conjugates and reactive reagents precoated at the corresponding regions.		
1 bottle of Extraction Buffer A - Buffer solution containing 0.2 M hydroxide with yellow cap.			
1 bottle of Extraction Buffer B - 10ml	Buffer solution containing 0.2 M hydrochloric acid with white cap.		
20 Extraction tubes	For specimens preparation use.		
2 Workstation	Place for holding buffer vials and tubes.		
1 Package insert	For operation instruction.		
1 Positive control swab	ntrol swab Contain inactived Gonococci and sodium		
(on request only)	azide. For external control.		
1 Negative control swab	Not contain Gonorrhea. For external		
(on request only)	control.		

#### MATERIALS REQUIRED BUT NOT PROVIDED

Timer	For timing use

#### **PRECAUTIONS**

- . For professional in vitro diagnostic use only.
- Do not use after the expiration date indicated on the package. Do not use the test
  if the foil pouch is damaged. Do not reuse tests.
- This kit contains products of animal origin. Certified knowledge of the origin and/or sanitary state of the animals does not completely guarantee the absence of transmissible pathogenic agents. It is therefore recommended that these products be treated as potentially infectious, and handled by observing usual safety precautions (e.g., do not ingest or inhale).
- Avoid cross-contamination of specimens by using a new extraction tube for each specimen obtained.
- Read the entire procedure carefully prior to testing.
- Do not eat, drink or smoke in any area where specimens and kits are handled. Handle all specimens as if they contain infectious agents. Observe established precautions against microbiological hazards throughout the procedure and follow standard procedures for the proper disposal of specimens. Wear protective clothing such as laboratory coats, disposable gloves and eye protection when specimens are assayed.
- Do not interchange or mix reagents from different lots. Do not mix solution bottle caps.
- Humidity and temperature can adversely affect results.
- When the assay procedure is completed, dispose swabs carefully after autoclaving them at 121°C for at least 20 minutes. Alternatively, swabs can be treated with 0.5% sodium hypochlorite (i.e., household bleach) for one hour before disposal.
- Used testing materials should be discarded according to local regulations.
- Do not use cytology brushes with pregnant patients.

#### STORAGE AND STABILITY

- The kit should be stored at 2-30°C until the expiry date printed on the sealed pouch.
- The test must remain in the sealed pouch until use.
- Do not freeze.
- Care should be taken to protect the components of this kit from contamination. Do
  not use if there is evidence of microbial contamination or precipitation. Biological
  contamination of dispensing equipments, containers or reagents can lead to false
  results.

## **SPECIMEN COLLECTION AND STORAGE**

- The quality of specimen obtained is of extreme importance. Detection of gonococci requires a rigorous and thorough collection technique which provides cellular material rather than just body fluids. Do not use 0.9% sodium chloride to treat swabs before collecting specimens.
- For female endocervical specimens:
  - Use only Dacron or Rayon tipped sterile swabs with plastic shafts. It is recommend to use the swab supplied by the kits manufacturer(The swabs are not contained in this kit, for the ordering information, please contact the manufacturer or local distributor, the catalog numbers are 207000female. 208000-male). Swabs from other suppliers have not been validated. Swabs with cotton tips or wooden shafts are not recommended.
  - Before specimen collection, remove excess mucus from the endocervical area
    with a separate swab or cotton ball and discard. The swab should be inserted
    into the endocervical canal, past the squamocolumnar junction, until most of
    the tip is no longer visible. This will permit acquisition of columnar or cuboidal
    epithelial cells which are the main reservoir of gonorrhea organisms. Firmly
    rotate the swab for 15 20 seconds without contamination with exocervical or
    vaginal cells.
  - If the swab may be tested immediately, replace the swab into the extraction tube.

#### • For male urethral specimens:

- Standard wire-shafted fiber-tipped swabs should be used for urethral specimen collection. Instruct the patients not to urinate at least two hours prior to specimen collection.
- Insert the swab 2-4 cm into the urea, rotate for 3-5 seconds and withdraw it. If
  the swab may be tested immediately, replace the swab into the extraction
  tube.
- Do not place the swab in any transport device containing medium. Transport
  medium interferes with the assay, and viability of organisms is not required for the
  assay. If immediate testing is not possible, patient samples should be placed in a
  dry transport tube for storage or transport. The swabs may be stored for 24 hours
  at room temperature (15-30°C) or 1 week at 4°C or no more than 6 month
  at -20°C. All specimens should be allowed to reach a room temperature of
  15-30°C before testing.

#### **PROCEDURE**

Bring tests, specimens, reagents and/or controls to room temperature (15-30°C) before use.

#### 1. Prepare endocervical or urethral swab specimens:

- Place a clean extraction tube in the workstation. Add 8 drops of Extraction Buffer A into the extraction tube.
- Immerse the patient swab into the extraction tube and wait 2 minutes. While
  waiting, use a circular motion to roll the swab against the side of the extraction
  tube so that the liquid is expressed from the swab and can reabsorb.
- At the end of the extraction time, add 8 drops of Extraction Buffer B to the
  tube and extract for another 1 minute in the same way. Then squeeze the
  swab firmly against the tube to expel as much liquid as possible from the
  swab. Discard the swab following guidelines for handling infectious agents.
- The extracted specimen can remain at room temperature for 60 minutes without affecting the test result.
- Remove the test from the sealed pouch and place it on a clean, level surface. Label the device with patient or control identification. For best results, the assay should be performed within one hour.
- 3. Add 3 drops (approximately 100  $\mu$ L) of extracted specimen from the extraction tube to the specimen well (S) of the test cassette.

Avoid trapping air bubbles in the specimen well (S), and do not add any solution to the result window.

As the test begins to work, color will migrate across the membrane.

Wait for the colored band(s) to appear. The result should be read at 15 minutes. Do not interpret the result after 20 minutes.

#### **INTERPRETATION OF RESULTS**

POSITIVE RESULT:	Two colored bands appear on the membrane. One band appears in the control region (C) and another band appears in the test region (T).
NEGATIVE RESULT:	Only one colored band appears, in the control region (C). No colored band appears in the test region (T).
INVALID RESULT:	Control band fails to appear. Results from any test which has not produced a control band in the specified read time must be discarded. Please review the procedure and repeat with a new test. If the problem persists, discontinue using the kit immediately and contact your local distributor.

#### NOTE:

- The intensity of color in the test region (T) may vary depending on the concentration of analytes present in the specimen. Note that this is a qualitative test, and cannot determine the concentrations of analytes in specimens.
- 2. Insufficient specimen volume, incorrect operation procedure or expired tests are the most likely reasons for control band failure.

## QUALITY CONTROL

- Internal procedural controls are included in the test. A colored band appearing in the control region (C) is considered as an internal positive procedural control, confirming sufficient specimen volume and correct procedural technique.
- External procedural controls may provided(on request only) in the kits to
  ensure that the tests function properly. Also, the Controls may be used to
  demonstrate proper performance by the test operator. To perform a positive
  or negative control test, complete the steps in the Test Procedure section
  treating the control swab in the same manner as a specimen swab.

## LIMITATIONS OF THE TEST

- StrongStep® Neisseria Gonorrhoeae Antigen Rapid Test is for professional in vitro diagnostic use, and should only be used for the qualitative detection of Neisseria gonorrhoeae. No meaning should be inferred from the color intensity or width of any apparent bands.
- 2. This test will only indicate the presence of Gonococcal antigen in specimens from both viable and non-viable Neisseria gonorrhoeae.
- Detection of Gonorrhea is dependent on the number of organisms present in the specimen. This may be affected by specimen collection methods and patient factors such as age, history of STD, presence of symptoms, etc. The minimum

- detection level of this test may vary according to serovar.
- 4. Women with vaginal discharge should be evaluated for risk factors of cervicitis and pelvic imflammatory diseases caused by other organisms including Candida albicans, Trichomonas vaginalis or Bacterial Vaginosis(These can be also diagnosed by LimingBio's other products: 500030 Candida albicans antigen rapid test; 500040 Trichomonas vaginalis antigen rapid test;500060 Candida albicans/Trichomonas vaginalis combo antigen rapid test; 500080 Bacterial vaginosis rapid test)
- 5. As with all diagnostic tests, a definitive clinical diagnosis should not be based on the results of a single test, but should only be made by the physician after all clinical and laboratory findings have been evaluated.
- Therapeutic failure or success cannot be determined as antigen may persist following appropriate antimicrobial therapy.
- Excessive blood (>50 µL in case of female swabs and >20 µL in case of male swabs) may cause false positive results. Endocervical samples from female patients should not be collected during menstrual period.

## PERFORMANCE CHARACTERISTICS

# Table: StrongStep® Neisseria Gonorrhoeae Antigen Rapid Test vs. Culture

#### **Female Endocervical Specimens**

Relative Sensitivity:			Cult	ture	
96.9% (89.2%-99.6%)* Relative Specificity:			+	-	Total
95.8% (93.6%-97.4%)*	StrongStep®		62	21	83
Overall Agreement: 95.9% (93.9%-97.4%)*	Gonorroheae Test	-	2	473	475
*95% Confidence Interval			64	494	558

#### **Male Urethral Specimens**

Relative Sensitivity:			Culture		
97.8% (93.7%-99.6%)* Relative Specificity:			+	ı	Total
99.5% (98.2%-99.9%)*	StrongStep®		134	2	136
Overall Agreement:         Gonorrohea           99.1% (97.8%-99.7%)*         Test	Gonorroheae Test	-	3	389	392
*95% Confidence Interval			137	391	528

#### Specificity:

Cross reactivity with organisms has been studied using suspensions of  $10^7$  CFU/ml. The following organisms produced negative results with the test:

Acinetobacter calcoaceticus	Pseudomona aeruginosa	
Acinetobacter spp	Gardnerella vaginalis	
Enterococcus faecalis	Salmonella choleraesius	
Enterococcus faecium	Candida albicans	
Staphylococcus aureus	Proteus vulgaris	
Proteus mirabilis	Hemophilus influenzae	
Chlamydia trachomatis	Klebsiella pneumoniae	
Group B/C Streptococcus	Ureaplasma Urealyticum	
Mycoplasma hominis	Trichomonas vaginalis	

## LITERATURE REFERENCES

- Knapp, J.S. et al. Neisseria gonorrhoeae. Manual of Clinical Microbiology, Sixth Edition, ASM Press, Washington DC., 324-325 (1995).
- Centers for Disease Control and Prevention. Sexually Transmitted Diseases Treatment Guidelines 2002. Morbidity and Mortality Weekly Report (2002), 51(RR-

## **GLOSSARY OF SYMBOLS**

REF	Catalog number	1	Temperature limitation		
	Consult instructions for use	LOT	Batch code		
IVD	In vitro diagnostic medical device	8	Use by		
***	Manufacturer	Σ	Contains sufficient for <n> tests</n>		
2	Do not reuse	EC REP	Authorized representative in the European Community		
CE	CE marked according to IVD Medical Devices Directive 98/79/EC				

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