

 **BD BACTEC™ Peds Plus™/F Culture Vials**
Soybean-Casein Digest Broth with Resins in a Plastic Vial

IVD Rx Only  500008334(05)
2019-09
English

INTENDED USE

BD BACTEC™ Peds Plus™/F culture vials (enriched Soybean-Casein Digest broth with CO₂) are for aerobic blood cultures. Principal use is with the BD BACTEC fluorescent series instruments for the qualitative culture and recovery of aerobic microorganisms (mainly bacteria and yeast) from pediatric and non-pediatric blood specimens which are generally less than 3 mL in volume.

SUMMARY AND EXPLANATION

The sample to be tested is inoculated into one or more vials which are inserted into the BD BACTEC fluorescent series instrument for incubation and periodic reading. Each vial contains a chemical sensor which can detect increases in CO₂ produced by the growth of microorganisms. The sensor is monitored by the instrument every ten minutes for an increase in its fluorescence, which is proportional to the amount of CO₂ present. A positive reading indicates the presumptive presence of viable microorganisms in the vial. Detection is limited to microorganisms that will grow in a particular type of medium.

Resins have been described for the treatment of blood specimens both prior to and after their inoculation into culture media. Resins have been incorporated into BD BACTEC culture media to enhance recovery of organisms without a need for special processing.^{1-3,8}

PRINCIPLES OF THE PROCEDURE

If microorganisms are present in the test sample inoculated into the BD BACTEC vial, CO₂ will be produced when the organisms metabolize the substrates present in the vial. Increases in the fluorescence of the vial sensor caused by the higher amount of CO₂ are monitored by the BD BACTEC fluorescent series instrument. Analysis of the rate and amount of CO₂ increase enables the BD BACTEC fluorescent series instrument to determine if the vial is positive, i.e., that the test sample contains viable organisms.

REAGENTS

The BD BACTEC culture vials contain the following reactive ingredients prior to processing:

List of Ingredients	BD BACTEC Peds Plus/F	List of Ingredients	BD BACTEC Peds Plus/F
Processed Water	40 mL	Hemin	0.0005% w/v
Soybean-Casein Digest Broth	2.75% w/v	Menadione	0.00005% w/v
Yeast Extract	0.25% w/v	Sodium Polyanetholsulfonate (SPS)	0.02% w/v
Animal Tissue Digest	0.10% w/v	Pyridoxal HCl (Vitamin B ₆)	0.001% w/v
Sodium Pyruvate	0.10% w/v	Nonionic Adsorbing Resin	10.0% w/v
Dextrose	0.06% w/v	Cationic Exchange Resin	0.6% w/v
Sucrose	0.08% w/v		

All BD BACTEC media are dispensed with added CO₂.

Warnings and Precautions:

The prepared culture vials are for *in vitro* diagnostic use. This Product Contains Dry Natural Rubber.

Pathogenic microorganisms, including hepatitis viruses and Human Immunodeficiency Virus, may be present in clinical specimens. "Standard Precautions"⁴⁻⁷ and institutional guidelines should be followed in handling all items contaminated with blood and other body fluids.

Prior to use, each vial should be examined for evidence of contamination such as cloudiness, bulging or depressed septum, or leakage. DO NOT USE any vial showing evidence of contamination. A contaminated vial could contain positive pressure. If a contaminated vial is used for direct draw, gas or contaminated culture media could be refluxed into the patient's vein. Vial contamination may not be readily apparent. If a direct draw procedure is used, monitor the process closely to avoid refluxing materials into the patient.

Prior to use, the user should examine the vials for evidence of damage or deterioration. Vials displaying turbidity, contamination, or discoloration (darkening) should not be used. On rare occasions a vial may not be sealed sufficiently. The contents of the vials may leak or spill, especially if the vial is inverted. If the vial has been inoculated, treat the leak or spill with caution, as pathogenic organisms/agents may be present. Before discarding, sterilize all inoculated vials by autoclaving.

Positive culture vials for subculturing or staining, etc.: before sampling it is necessary to release gas which often builds up due to microbial metabolism. Sampling should be performed in a biological safety cabinet if possible, and appropriate protective clothing, including gloves and masks, should be worn. See Procedure section for more information on subculturing.

To minimize the potential of leakage during inoculation of specimen into culture vials, use syringes with permanently attached needles or BD Luer-Lok™ tips.

Molecular tests performed on positive blood cultures will detect both viable and non-viable organisms commonly found in culture media. Therefore, Molecular test results should be evaluated in conjunction with Gram Stain results in accordance with standard-of-care practices as well as manufacturer's instructions for use.

Storage Instructions

The BD BACTEC vials are ready for use as received and require no reconstitution or dilution. Store in a cool, dry place (2–25 °C), **out of direct light.**

SPECIMEN COLLECTION

The specimen must be collected using sterile techniques to reduce the chance of contamination. The range of blood volume which can be cultured is 0.5 to 5.0 mL. If the volume of blood cultured is less than 0.5 mL, recovery of some fastidious organisms, such as *Haemophilus* species, may require the use of an appropriate supplement, as described later in this Package Insert. It is recommended that the specimen be inoculated into the BD BACTEC vials at bedside. Most commonly, a syringe with a BD Luer-Lok brand tip is used to draw the sample. If appropriate, a BD Vacutainer® brand Needle Holder and a BD Vacutainer brand Blood Collection Set, Vacutainer Safety-Lok™ Blood Collection Set or other tubing "butterfly" set may be used. If using a needle and tubing set (direct draw), carefully observe the direction of blood flow when starting sample collection. The vacuum in the vial will usually exceed 5 mL, so the user should monitor the volume collected by means of the 5 mL graduation marks on the vial label. When the recommended 1–3 mL has been drawn, the flow should be stopped by crimping the tubing and removing the tubing set from the BD BACTEC vial. **The inoculated BD BACTEC vial should be transported as quickly as possible to the laboratory.**

PROCEDURE

Remove the flip-off cap from the BD BACTEC vial top and inspect the vial for cracks, contamination, excessive cloudiness, and bulging or indented septum. DO NOT USE if any defect is noted. Before inoculating, swab the septum with alcohol (iodine is not recommended). Aseptically inject or draw directly a maximum of 5 mL of specimen per vial (see Limitations of the Procedure). Inoculated vials should be placed in the BD BACTEC fluorescent series instrument as soon as possible for incubation and monitoring. If placement of an inoculated vial into the instrument has been delayed and visible growth is apparent, it should not be tested in the BD BACTEC fluorescent series instrument, but rather it should be subcultured, Gram-stained and treated as a presumptively positive vial.

Vials entered into the instrument will be automatically tested every ten minutes for the duration of the testing protocol period. Positive vials will be determined by the BD BACTEC fluorescent series instrument and identified as such (see the appropriate BD BACTEC Fluorescent Series Instrument User's Manual). The sensor inside the vial will not appear visibly different in positive and negative vials, however, the BD BACTEC fluorescent series instrument can determine a difference in fluorescence.

If at the end of the testing period a negative vial appears visually positive (i.e., chocolatized blood, bulging septum, and/or lysed), it should be subcultured and Gram-stained and treated as a presumptively positive.

Positive vials should be subcultured and Gram-stained. In a great majority of cases, organisms will be seen and a preliminary report can be made to the physician. Subcultures to solid media and a preliminary direct antimicrobial susceptibility test may be prepared from fluid in the BD BACTEC vials.

Subculturing: Prior to subculturing, put the vial in an upright position, and place an alcohol wipe over the septum. To release pressure in the vial, use an appropriate venting unit (BD Cat. No. 249560, or equivalent). The needle should be removed after the pressure is released and before sampling the vial for subculture. The insertion and withdrawal of the needle should be done in a straight-line motion, avoiding any twisting motions.

QUALITY CONTROL

Quality control requirements must be performed in accordance with applicable local, state and/or federal regulations or accreditation requirements and your laboratory's standard Quality Control procedures. It is recommended that the user refer to pertinent CLSI guidance and CLIA regulations for appropriate Quality Control practices.

DO NOT USE culture vials past their expiration date.

DO NOT USE culture vials that exhibit any cracks or defects; discard the vial in the appropriate manner.

Quality Control Certificates are provided with each carton of media. Quality Control Certificates list test organisms, including ATCC® cultures specified in the CLSI Standard M22, *Quality Control for Commercially Prepared Microbiological Culture Media*. The range of time-to-detection in hours was ≤ 72 hours for each of the organisms listed on the Quality Control Certificate for this medium:

Peds Plus Medium Organisms

<i>Streptococcus pyogenes</i> ATCC 19615	<i>Neisseria meningitidis</i> ATCC 13090
<i>Escherichia coli</i> ATCC 25922	<i>Alcaligenes faecalis</i> ATCC 8750
<i>Streptococcus pneumoniae*</i> ATCC 6305	<i>Haemophilus influenzae</i> ATCC 19418
<i>Pseudomonas aeruginosa</i> ATCC 27853	<i>Staphylococcus aureus</i> ATCC 25923
<i>Candida albicans</i> ATCC 18804	

* CLSI-recommended strain

For information on Quality Control for the BD BACTEC fluorescent series instrument, refer to the appropriate BD BACTEC Fluorescent Series Instrument User's Manual.

RESULTS

A positive sample is determined by the BD BACTEC fluorescent series instrument and indicates the presumptive presence of viable microorganisms in the vial.

LIMITATIONS OF THE PROCEDURE

Contamination

Care must be taken to prevent contamination of the sample during collection and inoculation into the BD BACTEC vial. A contaminated sample will give a positive reading, but will not indicate a relevant clinical sample. Such a determination must be made by the user based on such factors as type of organism recovered, occurrence of the same organism in multiple cultures, patient history, etc.

Recovery of SPS Sensitive Organisms From Blood Samples

Because blood can neutralize the toxicity of SPS toward organisms sensitive to SPS (such as some *Neisseria* species), the presence of recommended volumes of blood (1–3 mL) can help to optimize recovery of these organisms.

Some organisms may be dependent on having a minimum amount of blood in the medium for optimal growth. Fastidious organisms, such as certain *Haemophilus* species, require growth factors from the blood specimen, such as NAD, or factor V. Optimal growth of these organisms is dependent on having greater than a minimum of 0.5 mL blood in the specimen. If the blood specimen volume is very small (0.5 mL or less), an appropriate supplement may be required for recovery of these organisms. BD BACTEC FOS™ Fastidious Organism Supplement (catalog number 442153) may be used as a nutritional supplement.

Nonviable Organisms

A Gram-stained smear from a culture medium may contain small numbers of nonviable organisms derived from media constituents, staining reagents, immersion oil, glass slides, and specimens used for inoculation. In addition, the patient specimen may contain organisms that will not grow in the culture medium or in media used for subculture. Such specimens should be subcultured to special media as appropriate.

Antibiotic Activity

Neutralization of the antibiotic activity by resins varies depending on dosage level and timing of specimen collection.

Studies have demonstrated that the resins present in this medium do not adequately neutralize meropenem preparations.

Studies have demonstrated that the resins present in this medium adequately neutralize the antifungal agent fluconazole with *Candida albicans*. However, other antifungal agents/yeast combinations have not been tested/evaluated.

Recovery of *Streptococcus pneumoniae*

In aerobic media, *S. pneumoniae* will typically be visually and instrument positive, but in some cases no organism will be seen on Gram stain or recovered on routine subculture. If an anaerobic vial was also inoculated, the organism can usually be recovered by performing an aerobic subculture of the anaerobic vial, since this organism has been reported to grow well under anaerobic conditions.⁹

General Considerations

Recovery of isolates will be achieved by adding the recommended volume 1–3 mL of blood. Use of lower or higher volumes may adversely affect recovery and/or detection. Blood may contain antimicrobials or other inhibitors which may slow or prevent the growth of microorganisms. False negative readings may result when certain organisms are present which do not produce enough CO₂ to be detected by the system or significant growth has occurred before placing the vial into the system. False positivity may occur when the white blood cell count is high. The default 5-day (120 hours) protocol was utilized for all analytical testing with the BD BACTEC Peds Plus/F culture media and protocol lengths of >5 days have not been evaluated.

EXPECTED VALUES AND SPECIFIC PERFORMANCE CHARACTERISTICS

Internal studies have demonstrated that antibiotics are effectively neutralized by the resins used in BD BACTEC resin media. In these tests, antibiotics were added in clinically relevant concentrations directly to resin media prior to inoculation with susceptible strains. These tests showed equivalent performance in BD BACTEC Peds Plus in plastic vial compared to BD BACTEC Peds Plus in glass vial.

A total of 984 paired sets inoculated with 0.5 mL and 5.0 mL of blood at 10–100 CFU per vial were evaluated across the four instruments comprising the BD BACTEC fluorescent-series instrument family: BD BACTEC 9050, BD BACTEC 9240, BD BACTEC FX and the BD BACTEC FX40. Of the 984 paired sets 953 sets recovered organisms within the instrument series. There were 18 sets with no detection of organisms in either the plastic or glass vial that included *Candida albicans* (4 sets) *Haemophilus influenzae* (9 sets) and *Haemophilus parainfluenzae* (5 sets). There were 4 sets with no detection in the plastic vial that included *Candida albicans* (2 sets), *Enterococcus faecalis* (1 set) and *Haemophilus influenzae* (1 set). There were 9 sets with no detection in the glass vial that included *Candida albicans* (3 sets), *Haemophilus influenzae* (1 set) *Haemophilus parainfluenzae* (4 sets) and *Pediococcus acidilactici* (1 set) with no detection in the glass vial. The detection rate of *Candida albicans*, *Enterococcus faecalis* and *Pediococcus acidilactici* was 73%, 98% and 98% respectively under these test conditions. The *Haemophilus* species detection rates were 69% with 0.5 mL blood, and 100% with 5.0 mL, due to the quality (freshness) and volume of blood used in the test. There were five organisms with false negative results (i.e., end of protocol, instrument negative vials with a positive terminal subculture) observed with the BD BACTEC Peds Plus/F medium contained in a plastic vial using 0.5 mL of bagged blood: *H. influenzae* inoculated at 54, 65 CFU, *Haemophilus parainfluenzae* inoculated at 4, 58 CFU, *Candida glabrata*, inoculated at 1 CFU, *Micrococcus luteus* inoculated at 0 CFU, and *Cryptococcus neoformans* inoculated at 0 CFU. Three *Haemophilus influenzae* strains were species that were retested using 0.5 and 1 mL fresh instead of bagged blood and were detected in both glass and plastic vials.

An additional study of 492 paired sets inoculated with 3mL of blood at 10–100 CFU per vial were evaluated across the four instruments comprising the BD BACTEC fluorescent-series instrument family: BD BACTEC 9050, BD BACTEC 9240, BD BACTEC FX and the BD BACTEC FX40. All organisms recovered from the 492 paired sets across the four BACTEC instruments. The *Haemophilus* species detection rate was 100% with 3.0 mL blood due to the volume of blood used in the test. There were 4 sets that favored the glass vial, the mean time to detection was <10%; these vials included *Candida glabrata*, *Stenotrophomonas maltophilia*, *Candida albicans* and *Haemophilus parainfluenzae*.

The following organisms were evaluated in the analytical studies: *Abiotrophia defective*, *Acinetobacter Iwoffii*, *Actinobacillus actinomycetemcomitans*, *Aerococcus viridans*, *Alcaligenes faecalis*, *Bacillus subtilis*, *Candida albicans*, *Candida glabrata*, *Candida tropicalis*, *Candida parapsilosis*, *Cardiobacterium hominis*, *Corynebacterium jeikeium*, *Cryptococcus neoformans*, *Eikenella corrodens*, *Enterobacter cloacae*, *Enterococcus faecalis*, *Escherichia coli*, *Granulicatella adiacens*, *Haemophilus influenzae*, *Haemophilus influenzae*, type a, *Haemophilus influenzae*, type b, *Haemophilus parainfluenzae*, *Kingella kingae*, *Klebsiella pneumoniae*, *Leuconostoc mesenteroides*, *Micrococcus luteus*, *Neisseria gonorrhoeae*, *Neisseria meningitidis*, *Pediococcus acidilactici*, *Proteus mirabilis*, *Providencia stuartii*, *Pseudomonas aeruginosa*, *Staphylococcus aureus*, *Staphylococcus epidermidis*, *Stenotrophomonas maltophilia*, *Rothia mucilaginosa* (formerly *Stomatococcus mucilaginosus*), *Streptococcus agalactiae*, four strains of *Streptococcus pneumoniae*, *Streptococcus pyogenes*, and *Streptococcus sanguinis* (formerly *S. sanguis*).

In microbial detection limit testing, a total of 360 paired sets inoculated with 0.5 mL, 5.0 mL blood at target inoculum levels of 0 to 1 and 1 to 10 CFU per vial were evaluated. This study was designed to assess the capability of the BD BACTEC blood culture media tested to detect one CFU, when present. Of the 360 paired sets tested, 196 grew and detected in both devices, 42 detected in glass vials only, 57 detected in plastic vials only, and 65 did not detect in either. There were a total of 107 pair sets that were not detected in plastic vials, in which 36 showed organism growth on the inoculum plate: *Neisseria meningitidis* (5 CFU), *Haemophilus parainfluenzae* (4 CFU), *Staphylococcus epidermidis* (2 CFU), 1 CFU each for *Candida albicans*, *Candida glabrata*, *Escherichia coli*, *Staphylococcus aureus*, and *Streptococcus sanguinis*. The remaining 71 pair sets showed no organism growth (0 CFU) on the inoculum plate: *Cryptococcus neoformans*, *Enterococcus faecalis*, *Haemophilus influenzae*, *Haemophilus parainfluenzae*, *Micrococcus luteus*, *Neisseria gonorrhoeae*, *Neisseria meningitidis*, *Staphylococcus epidermidis*, and *Streptococcus pneumoniae*.

In additional microbial detection limit testing, a total of 180 paired sets inoculated with 3 mL blood at target inoculum levels of 0 to 1 and 1 to 10 CFU per vial were evaluated. This study was designed to assess the capability of the BD BACTEC blood culture media tested to detect one CFU, when present. Of the 180 paired sets tested, 104 grew and detected in both devices, 23 detected in glass vials only, 19 detected in plastic vials only, and 34 did not detect in either. There were a total of 57 paired sets that were not detected in plastic vials, in which 23 showed organism growth on the inoculum plate: 1 CFU each for *Candida albicans*, *Cryptococcus neoformans*, *Haemophilus influenzae*, *Neisseria gonorrhoeae*, *Pseudomonas aeruginosa*, *Staphylococcus aureus*, *Streptococcus pneumoniae*, and *Streptococcus sanguinis*. The remaining 34 paired sets showed no organism growth (0 CFU) on the inoculum plate: *Candida glabrata*, *Enterococcus faecalis*, *Escherichia coli*, *Haemophilus parainfluenzae*, *Micrococcus luteus*, *Neisseria meningitidis*, *Staphylococcus epidermidis*, and *Streptococcus pneumoniae*.

AVAILABILITY

Cat. No. Description

442020 BD BACTEC™ Peds Plus™/F Medium, Case of 50 vials

REFERENCES

1. Wallis, C. et al. 1980. Rapid isolation of bacteria from septicemic patients by use of an antimicrobial agent removal device. *J. Clin. Microbiol.* 11:462–464.
2. Applebaum, P.C. et al. 1983. Enhanced detection of bacteremia with a new BACTEC resin blood culture medium. *J. Clin. Microbiol.* 7:48–51.
3. Pohlman, J.K. et al. 1995. Controlled clinical comparison of Isolator and BACTEC 9240 Aerobic/F resin bottle for detection of bloodstream infections. *J. Clin. Microbiol.* 33:2525–2529.
4. Clinical and Laboratory Standards Institute. 2005. Approved Guideline M29-A3. Protection of laboratory workers from occupationally acquired infections, 3rd ed. CLSI, Wayne, Pa.
5. Garner, J.S. 1996. Hospital Infection Control Practices Advisory Committee, U.S. Department of Health and Human Services, Centers for Disease Control and Prevention. Guideline for isolation precautions in hospitals. *Infect. Control Hospital Epidemiol.* 17: 53–80.
6. U.S. Department of Health and Human Services. 2007. Biosafety in microbiological and biomedical laboratories, HHS Publication (CDC), 5th ed. U.S. Government Printing Office, Washington, D.C.
7. Directive 2000/54/EC of the European Parliament and of the Council of 18 September 2000 on the protection of workers from risks related to exposure to biological agents at work (seventh individual directive within the meaning of Article 16(1) of Directive 89/391/ EEC). Official Journal L262, 17/10/2000, p. 0021–0045.
8. Flayhart, D. et al. 2007. Comparison of BACTEC Plus blood culture media to BacT/Alert FA blood culture media for detection of bacterial pathogens in samples containing therapeutic levels of antibiotics. *J. Clin. Microbiol.* 45:816–821.
9. Howden, R.J. 1976. Use of anaerobic culture for the improved isolation of *Streptococcus pneumoniae*. *J. Clin. Pathol.* 29:50–53.

Technical Information: In the United States contact BD Technical Service and Support at 1.800.638.8663 or bd.com.

Change History

Revision	Date	Change Summary
(05)	2019-09	<p>Converted printed instructions for use to electronic format and added access information to obtain the document from BD.com/e-labeling.</p> <p>In Warnings and Precautions section, added recommendation to perform molecular testings on positive blood cultures according to standard-of-care practices and manufacturer's instructions for use.</p>

US Customers only: For symbol glossary, refer to bd.com/symbols-glossary



Manufacturer / Производител / Výrobce / Fabrikant / Hersteller / Κατασκευαστής / Fabricante / Tootja / Fabricant / Proizvodač / Gyártó / Fabbricante / Аткарушы / 제조업체 / Gamintojas / Ražotājs / Tilvirker / Producēt / Producent / Производитель / Výrobca / Proizvodač / Tillverkare / Üretici / Виробник / 生产厂商



Use by / Использовайте до / Spotrebujte do / Brug før / Verwendbar bis / Xhríetjé éwç / Usar antes de / Kasutada enne / Date de péremption / 사용 기한 / Upotrijebiti do / Fehlasználhatóság dátuma / Usare entro / Дейн пайдалануѓа / Naudokite iki / Izletot līdz / Houdbaar tot / Brukes for / Stosować do / Prazo de validade / A se utiliza pánha la / Использовать до / Použíte do / Upotrebiti do / Använd före / Son kullanım tarihi / Використати доділе / 使用截止日期
 YYYY-MM-DD / YYYY-MM (MM = end of month)
 ГГГГ-ММ-ДД / ГГГГ-ММ (ММ = края на месецада)
 RRRR-MM-DD / RRRR-MM (MM = konec měsíce)
 AAAA-MM-DD / AAAA-MM (MM = slutning af måned)
 JJJJ-MM-TT / JJJJ-MM (MM = Monatsende)
 EEEE-MM-HH / EEEE-MM (MM = τέλος του μήνα)
 AAAA-MM-DD / AAAA-MM (MM = fin del mes)
 AAAA-KK-PP / AAAA-KK (KK = kuu lopp)
 AAAA-MM-JJ / AAAA-MM (MM = fin du mois)
 GGGG-MM-DD / GGGG-MM (MM = kraj mjeseca)
 ÉÉÉÉ-HH-NN / ÉÉÉÉ-HH (HH = hónap utolsó napja)
 AAAA-MM-GG / AAAA-MM (MM = fine mese)
 ЖЮЮЮК-АА-КК / ЖЮЮЮК-АА / (AA = айданы соңы)
 YYYY-MM-DD/YYYY-MM (MM = 월말)
 MMMM-MM-DD / MMMM-MM (MM = ménésio pabaiga)
 GGGG-MM-DD / GGGG-MM (MM = mēnēša beigas)
 JJJJ-MM-DD / JJJJ-MM (MM = einde maand)
 AAAA-MM-DD / AAAA-MM (MM = slutten av måneden)
 RRRR-MM-DD / RRRR-MM (MM = koniec miesiąca)
 AAAA-MM-DD / AAAA-MM (MM = fin do měsíce)
 AAAA-LL-ZZ / AAAA-LL (LL = sfârșitul lunii)
 ГГГГ-ММ-ДД / ГГГГ-ММ (ММ = конец месяца)
 RRRR-MM-DD / RRRR-MM (MM = koniec mesiaca)
 GGGG-MM-DD / GGGG-MM (MM = kraj meseca)
 AAAA-MM-DD / AAAA-MM (MM = slutet av månaden)
 YYYY-AA-GG / YYYY-AA (AA = ayin sonu)
 PPPP-MM-ДД / PPPP-MM (MM = кінець місяця)
 YYYY-MM-DD / YYYY-MM (MM = 月末)



Catalog number / Каталожен номер / Katalogové číslo / Katalognummer / Αριθμός καταλόγου / Número de catálogo / Katalooginumber / Numéro catalogue / Kataloški broj / Katalógu szám / Numero di catalogo / Каталог номірі / 카탈로그 번호 / Katalogo / numeris / Kataloga numurs / Catalogus nummer / Numer katalogowy / Număr de catalog / Номер по каталогу / Katalógové číslo / Kataloški broj / Katalog numerasi / Номер за каталогом / 目录号



Authorized Representative in the European Community / Оторизиран представител в Европейската общност / Autorizovaný zástupce pro Evropském společenství / Autoriseret repræsentant i De Europæiske Fællesskaber / Autorisierte Vertreter in der Europäischen Gemeinschaft / Εξουσιοδοτημένος αντιπρόσωπος στην Ευρωπαϊκή Κοινότητα / Representante autorizado en la Comunidad Europea / Volitatud esindaja Euroopa Nõukogus / Reprézentant autorisé pour la Communauté européenne / Autorizuaruun predstavnik u Europskui uniji / Meghatalmazott képviselő az Európai Közösségen / Rappresentante autorizzato nella Comunità Europea / Европа кауымдастырындығы үекінетті екін / 유럽 공동체의 위원 대표 / Igaliotasis atstovas Europos Bendrijoje / Plinvaroitis pārstāvis Eiropas Kopienā / Bevoegde vertegenwoordiger in de Europese Gemeenschap / Autoriseret representant i EU / Autoryzowane przedstawicielstwo we Wspólnocie Europejskiej / Representante autorizado na Comunidade Europeia / Reprézentantul autorizat pentru Comunitatea Europeană / Уполномоченный представитель в Европейском сообществе / Autorizovaný zástupce v Evropskom spoločenstve / Autorizované predstavništvo v Evropskej unii / Auktoriseraad representant i Europeiska gemenskapen / Avrupa Topluluğu Yetkili Temsilcisi / Упновоаженний представник у країнах ЄС / 欧洲共同体授权代表



In Vitro Diagnostic Medical Device / Медицински уред за диагностика ин vitro / Lékařské zařízení určené pro diagnostiku in vitro / In vitro diagnostisk medicinsk anordning / Medizinisches In-vitro-Diagnostikum / In vitro биохимический импринт схема / Dispositivo médico para diagnóstico in vitro / In vitro diagnostika meditsinskiyaparatur / Dispositif médical de diagnostic in vitro / Medicinska pomagala za In Vitro Dijagnostiku / In vitro diagnozszkai orvosi eszköz / Dispositivo medicale per diagnostica in vitro / Жасанды жағдайда хүргізетін медициналық диагностика аспабы / In Vitro Diagnostic 의료 기기 / In vitro diagnostikos prietais / Medicinas ierīces, ko lietū in vitro diagnostikā / Medisch hulpmiddel voor in-vitro diagnostiek / In vitro diagnostisk medical utstyr / Urządzenie medyczne do diagnostyki in vitro / Dispositivo médico para diagnóstico in vitro / Dispositiv medical pentru diagnostic in vitro / Медицинский прибор для диагностики in vitro / Medicínska pomôcka na diagnostiku in vitro / Medicinski uredaj za in vitro diagnostiku / Medicinteknisk produkt för in vitro-diagnostik / In Vitro Diagnostik Tibbi Cihaz / Медицинский пристрой для диагностики in vitro / 体外诊断医疗设备



Temperature limitation / Температурни ограничения / Teplotní omezení / Temperaturbegrenzung / Temperaturbegrenzung / Περιορισμοί θερμοκρασίας / Limitación de temperatura / Temperatura / Temperaturai piirang / Limites de température / Dozvoljena temperatura / Hörmésekleti határ / Limiti di temperatura / Температурны шектегү / 온도 제한 / Laikymo temperatūra / Temperatūras ierobežojumi / Temperatuurlimiet / Temperaturbegrenzung / Ograniczenie temperatury / Limites de temperatura / Limite de temperatūrā / Ограничение температуры / Ohraničenie teploty / Ograničenje temperature / Temperaturgräns / Sicaklık sınırlaması / Обмеження температури / 温度限制



Batch Code (Lot) / Код на партидата / Kód (číslo) šarže / Batch-kode (lot) / Batch-Code (Charge) / Κωδικός παρτίδας (παρτίδα) / Código de lote (lote) / Partii kood / Numéro de lot / Lot (kod) / Tétel száma (Lot) / Codice batch (lotto) / Топтама коды / 배치 코드(로트) / Partijos numeris (LOT) / Partijas kods (laidiens) / Lot nummer / Batch-kode (parti) / Kod partii (seria) / Código do lote / Cod de serie (Lot) / Код партии (лот) / Kód série (šarža) / Kod serije / Partinummer (Lot) / Parti Kodu (Lot) / Код партии / 批号 (亚批)



Contains sufficient for <n> tests / Съдържанието е достатъчно за <n> теста / Dostatečné množství pro <n> testů / Indeholder tilstrækkeligt til <n> tests / Ausreichend für <n> Tests / Περιέχει епокрк пообогатяло за <n> εξετάσεις / Contenido suficiente para <n> pruebas / Kullaldane <n> testimendi jaoks / Contenu suffisant pour <n> tests / Sadržaj za <n> testova / <n> tesztet elegendő / Contenuto sufficiente per <n> test / <n> тесттери үшін жеткілікті / <n> 테스트가 충분히 포함됨 / Pakankamas kiekis atlikti <n> testų / Satur pietiekami <n> párbaudém / Inhou volodoende voor "n" testen / Innholder tilstrekkelig til <n> tester / Zawiera ilość wystarczającą do <n> testów / Conteúdo suficiente para <n> testes / Continut suficient pentru <n> teste / Достаточно для <n> тестов(a) / Obsah vystačí na <n> testov / Sadržaj dovoljan za <n> testova / Innehåller tillräckligt för <n> analyser / <n> test için yeterli malzemeler / Вистачить для аналізу: <n> / 足够进行 <n> 次检测



Consult Instructions for Use / Направете справка в инструкциите за употреба / Prostidujte pokyny k použití / Se brugsanvisningen / Gebrauchsanweisung beachten / Συμβουλεύτε τις οδηγίες χρήσης / Consultar las instrucciones de uso / Luggedi kasutusjuhendit / Consulter la notice d'emploi / Koristi upute za upotrebu / Olvassa el a használati utasítást / Consultare le istruzioni per l'uso / Пайдалану нұсқаулығымен танысын алыңыз / 사용 지침 참조 / Skaitykite naudojimo instrukcijas / Skafit lietosanás pamáčibú / Raadpleeg de gebruiksaanwijzing / Se i bruksanvisningen / Zobacz instrukcję użytkowania / Consultar as instruções de utilização / Consultați instrucțiunile de utilizare / См. руководство по эксплуатации / Poznij Pokyny na používanie / Pogledajte uputstvo za upotrebu / Se bruksanvisningen / Kullanım Talimatları'na başvurun / Див. інструкції з використання / 请参阅使用说明



Do not reuse / Не използвайте отново / Nepoužívejte opakovane / Ikke til genbrug / Nicht wiederverwenden / Μην επαναχρησιοποιείτε / No reutilizar / Mitte kasulada korduvat / Ne pas réutiliser / Не користи поново / Egyszer használatos / Non riutilizzare / Пайдаланбандыз / 재사용 금지 / Tik vienkartiniam naudojimui / Nelietot atkārtoti / Niet opnieuw gebruiken / Kun til engangsbruk / Nie stosować powtórnie / Não reutilize / Nu refolositi / Не использовать повторно / Nepoužívajte opakovane / Ne upotrebljavajte ponovo / Fár ej áteranvändas / Tekrar kullanmayın / Не використовувати повторно / 请勿重复使用



Serial number / Серийн номер / Sériové číslo / Seriennummer / Seriennummer / Σεριακός αριθμός / Nº de serie / Serianumber / Numéro de série / Serijski broj / Sorozatszám / Numero di serie / Топтамалық немірі / 일련 번호 / Serijos numeris / Sérías numurs / Serie nummer / Numer seryjny / Número de série / Număr de serie / Серийный номер / Серійна марка / Номер серії / 序列号



For IVD Performance evaluation only / Само за оценка качеството на работа на IVD / Pouze pro vyhodnocení výkonu IVD / Kun til evaluering af IVD ydelse / Nur für IVD-Leistungsbewertungszwecke / Môvo yhľaduľou počas výskumu IVD / Sóly para la evaluación del rendimiento en diagnóstico in vitro / Ainult IVD seadme hindamiseks / Réservez à l'évaluation des performances IVD / Samo u znanstvene svrhe za In Vitro Dijagnostiku / Kizárolag in vitro diagnosztikához / Solo per valutazione delle prestazioni IVD / Жасанды жағдайда «пробирка ишінде», диагностикада тек жұмысты бағапау ушін / IVD 성능 평가에 대해서만 사용 / Tik IVD prietais veikimo charakteristikoms tikrinti / Vientij IVD darbības novērtēšanai / Uitsluitend voor doelstreffendheidsonderzoek / Kun for evaluering af IVD-ydelse / Tylko do oceny wydajności IVD / Uso exclusivo para avaliação de IVD / Umai para evaluararea performanței IVD / Только для оценки качества диагностики in vitro / Určené iba na diagnostiku in vitro / Samo za procenu učinka u in vitro diagnostici / Endast för utvärdering av diagnostisk användning in vitro / Yalnızca IVD Performans değerlendirme için / Тільки для оцінювання якості діагностики in vitro / 仅限 IVD 性能评估

For US: "For Investigational Use Only"



Lower limit of temperature / Долен лимит на температурата / Dolní hranice teploty / Nedre temperaturgrænse / Temperaturuntergrenze / Катырға орьо ѡнеруулары / Límite inferior de temperatura / Alumine temperaturipirip / Limite inférieure de température / Najniżza dozwolona temperatura / Alsó hőmérsékleti határ / Limite inferiori di temperatura / Температуралың теменгі рүсгән шері / 하한 온도 / Žemiausiai laikymo temperatūra / Temperatūras zemakā robeža / Laagste temperatuurlimiet / Nedre temperaturgrense / Dolna granica temperatury / Limite mínima de temperatura / Limită minimă de temperatură / Нижний предел температуры / Spodná hranica teploty / Donja granična temperature / Nedre temperaturgräns / Sicaklıktı alır sınırı / Минимальна температура / 温度下限



Control / Контролно / Kontrola / Kontrol / Kontrolle / Μόρτυρας / Kontroll / Contrôle / Controllo / Controllo / Бақылау / Контроль / Kontroll / Kontrol / Kontrole / Kontrol / Kontrol / Kontroll / Контроль / 对照



Positive control / Позитивният контрол / Pozitivní kontrola / Positiv kontroll / Positive Kontrolle / Θετικός μάρτυρας / Control positivo / Positivne kontroll / Contrôle positif / Pozitívna kontrola / Pozitív kontroll / Controllo positivo / Он бақылау / 양성 컨트롤 / Teigiamma kontrolé / Pozitív kontrole / Positiveve kontrole / Kontrola dodatnia / Controlo positivo / Control pozitiv / Поможительный контроль / Pozitif kontrol / Позитивный контроль / 阳性对照试剂



Negative control / Отрицателен контрол / Negativní kontrola / Negativ kontrol / Negative Kontrolle / Αρρυτικός μάρτυρας / Control negativo / Negativne kontroll / Contrôle négatif / Negativna kontrola / Negativ kontroll / Controlo negativo / Негативный контрол / Negativ kontroll / Negativ kontrole / Negatiieve controle / Kontrola ujemna / Controlo negativo / Control negativ / Отрицательный контроль / Negatif kontrol / Негативный контроль / 阴性对照试剂



Method of sterilization: ethylene oxide / Метод на стерилизация: этиленов оксид / Způsob sterilizace: etylenoxid / Steriliseringsmetode: ethylenoxid / Sterilisationsmethode: Ethylenoxid / Μέθοδος αποστείρωσης: αιθαλεοξείδιο / Método de esterilización: óxido de etileno / Steriliseerimismetood: etüleenoksidi / Méthode de stérilisation : oxyde d'éthylène / Metoda sterilizacije: etilen oksid / Sterilizálás módszere: etilén-oxid / Metodo di sterilizzazione: ossido di etilene / Стерилизация әдісі – этилен тотығы / 소독 방법: 에틸렌옥사이드 / Sterilizavimo būdas: etileno oksidas / Sterilizēšanas metode: etilēnoksīds / Gesteriliseerd met behulp van ethylenoxide / Steriliseringsmetode: etylenoksid / Metoda sterilizacije: tlenek etilu / Método de esterilização: óxido de etileno / Metodā da sterilizare: oxid de etilēnā / Метод стерилизации: этиленоксид / Metoda sterilizacije: etilén-oxid / Metoda sterilizacije: etilen oksid / Sterilizasyon yöntemi: etilen oksit / Метод стерилизацији: этиленоксидом / 灭菌方法: 环氧乙烷



Method of sterilization: irradiation / Метод на стерилизация: иридиация / Způsob sterilizace: bestrálení / Steriliseringsmetode: bestrålning / Sterilisationsmethode: Bestrahlung / Μέθοδος αποστείρωσης: ακτινοβολία / Método de esterilización: irradiación / Steriliseerimismetood: kiirgus / Méthode de stérilisation : irradiation / Metoda sterilizacije: zračenje / Sterilizálás módszere: besugárzás / Metodo di sterilizzazione: irradiazione / Стерилизация әдісі – иридиация / Метод да sterilizare: napromienianie / Método de esterilização: irradiação / Metodā da sterilizare: iadiere / Метод стерилизации: облучение / Metoda sterilizacije: ozračenje / Metoda sterilizacije: ozračavanje / Steriliseringsmetod: strålnin / Sterilizasyon yöntemi: irradiasyon / Метод стерилизацији: опрометненням / 灭菌方法: 辐射



Biological Risks / Биологични рискове / Biologická rizika / Biologisk fare / Biogegefährung / Виолюкіо кілдунор / Riesgos biológicos / Biologgiled riskid / Risques biologiques / Biolojik rizik / Biologialag veszélyes / Rischio biologico / Биологиялық тәуекелдер / 생물학적 위험 / Biologinis pavojus / Biologiskie riski / Biologisch risiko / Biologisk risiko / Zagrożenie biologiczne / Perigo biológico / Riscuri biologice / Биологическая опасность / Biologické riziko / Biološki rizici / Biologisk risk / Biyolojik Riskler / Биологична небезпека / 生物学风险



Caution, consult accompanying documents / Внимание, направете справка в придружащите документи / Pozor! Prostidujte si přiloženou dokumentaci! / Forsiktig, se ledsgagende dokumenter / Achtung, Begleiddokumente beachten / Προσοχή, συμβουλεύτε τα συνοδευτικά έγγραφα / Precaucción, consultar la documentación adjunta / Ettēvaatust! Luggedi kaasnevad dokumentatsiooni / Attention, consulter les documents joints / Upozorenje, koristi prateču dokumentaciju / Figuele! Olvassa el a mellékelt tájékoztatót / Attenzione: consultare la documentazione allegata / Абайланың, тиісті құжаттармен танысының / 주의, 동봉된 설명서 참조 / Démésio / Zürükte pridedamus dokumentus / Plesardziba, skafit pavaddokumentus / Voorzichtig, raadpleeg bijgevoegde documenten / Forsiktig, se vedlagt dokumentasjon / Należy zapoznać się z dołączonymi dokumentami / Cuidado, consulte a documentação fornecida / Atenție, consultați documentele însoțitoare / Внимание: см. прилагаемую документацию / Výstraha, pozri sprievodné dokumenty / Pažnja! Pogledajte priložena dokumenta / Obs! Se medföljande dokumentation / Dikkat, birlikte verilen belgelere başvurun / Увага: див. супутну документацію / 小心：请参阅附带文档。



Upper limit of temperature / Горен лимит на температурата / Horní hranice teploty / Øvre temperaturgrænse / Temperaturobergrenze / Ану́тэро орьо ѡнеруулары / Límite superior de temperatura / Ülémire temperaturipirip / Limite supérieure de température / Gornja dozvoljena temperatura / Felső hőmérsékleti határ / Limite superiore di temperatura / Температуралың түркіштегі жогары рүсгән шері / 상한 온도 / Aukščiausiai laikymo temperatūra / Augščiųjā temperatūras robeža / Hoogste temperatuurlimiet / Øvre temperaturgrense / Górnja granica temperatury / Limite máximo de temperatura / Limită maximă de temperatură / Верхний предел температуры / Horná hranica teploty / Gornja granična temperature / Øvre temperaturgräns / Sicaklıktı üst sınırı / Максимальна температура / 温度上限



Keep dry / Пазете сухо / Skladujte v suchém prostředi / Opbevares tørt / Trocklagern / Фулдьте то стөгүү / Mantener seco / Hoida kuivas / Conserver au sec / Držati na suhom / Száraz helyen tartandó / Tenere all'asciutto / Құрғак күйінде үста / 건조 상태 유지 / Laikyite sausai / Uzglabāt sausu / Droog houden / Holdes tørt / Przechowywać w stanie suchym / Manter seco / A se feri de umezelalā / Не допускать попадания влаги / Uchovávajte v suchu / Držite na suvom mestu / Förvaras torrt / Kuru bir şekilde muhafaza edin / Берегти від вологи / 请保持干燥



Collection time / Время на събиране / Čas odberu / Opsamlingstidspunkt / Entnahmehrzeit / Ørta суллюгүс / Hora de recogida / Kogumisaeg / Heure de prélevement / Sati prikupljanja / Mintavétel időpontja / Ora di raccolta / Жинау үақыты / 수집 시간 / Paémimo laikas / Savákšanas laiks / Verzameltijd / Tid prøvetaking / Godzina pobrania / Hora de coleitta / Ora colectării / Время сбора / Doba odberu / Vremea prikupljanja / Uppsamlingstid / Toplama zamanı / Час забора / 采集时间



Peel / Обелегте / Otevřete zde / Ábn / Abziehen / Аткоклъйтте / Desprender / Koorida / Décoller / Otvorit skini / Húzza le / Staccare / Үстінгі қабатын алып таста / 売き开け / Pliešti čia / Atlīmēt / Schillen / Trekk av / Oderwač / Destacar / Se dezlipește / Otklepnit / Odtrhnite / Olijuštiti / Dra isăr / Ayırma / Відклепні / 撕下



Perforation / Перфорация / Perforace / Perforering / Διάτρηση / Perforación / Perforatsioon / Perforacija / Perforálás / Perforazione / Tecik tecsy / 절취 선 / Perforacija / Perforācija / Perforatie / Perforacja / Perfuração / Perforare / Перфорация / Perforácia / Perforasyon / 穿孔





Do not use if package damaged / Не използвайте, ако опаковката е повредена / Neroužívejte, je-li obal poškozený / Má ikke anvendes hvis emballagen er beskadiget / Inhal beschädigter Packungsnicht verwenden / Μη χρησιμοποιείτε εάν η συσκευασία έχει υποστεί ζημιά. / No usar si el paquete está dañado / Mitte kasutada, kui pakend on kahjustatud / Ne pas l'utiliser si l'emballage est endommagé / Ne koristiti ako je oštećeno pakiranje / Ne használja, ha a csomagolás sérült / Non usare se la confezione è danneggiata / Erep paket bûzylgan болса, пакетаның / Пакет, як онын түркімі жағынан күлгін болса, пакетаның / Ne uporabljajte, če je pakovanje poškodenje / Nie używac, jeśli opakowanie jest uszkodzone / Não usar se a embalagem estiver danificada / A nu se folosi dacă pachetul este deteriorat / Не использовать при повреждении упаковки / Neroužívajte, ak je obal poškodený / Ne koristite ako je pakovanje oštećeno / Använd ej om förpackningen är skadad / Ambalaj hasar görmüse kullanmayın / Не використовувати за пошкодженої упаковки / 如果包装破损, 请勿使用



Keep away from heat / Газете от топлина / Nevystavujte přílišnému teplu / Má ikke udsættes for varme / Vor Wärme schützen / Краткото то макрия атпó тη θερμότητα / Mantener alejado de fuentes de calor / Hoida eemal valgusest / Protéger de la chaleur / Držati dalje od izvora topline / Óvja a melegtől / Tenerе lontano dal calore / Салын жерде сакта / 열을 피해야 함 / Laikyt atokiau nuo šilumos šaltinių / Sargāt no karstuma / Beschermen tegen warmte / Má ikke utsettes for varme / Przechowywać z dala od źródła ciepła / Manter ao abrigo do calor / A se feri de căldură / Не нарревать / Uchovávajte mimo zdroju tepla / Držite dalje od toplote / Får ej utsättas för värme / Isidan uzak tutun / Берегти від дії тепла / 请远离热源



Cut / Срежете / Odstrňhnete / Klip / Schneiden / Коят / Cortar / Lõigata / Découper / Reži / Vágja ki / Tagliare / Kecici / 잘라내기 / Kirpti / Nogrezt / Knippen / Kutt / Odciąć / Cortar / Decupať / Отрезать / Odstrihnite / Iseći / Klipp / Kesme / Rozřízati / 剪下



Collection date / Дата на събиране / Datum odběru / Opsamlingsdato / Entnahmedatum / Ημερομηνία συλλογής / Fecha de recogida / Kogumiskuupäev / Date de prélèvement / Dani prikupljanja / Mintavétel dátuma / Data di raccolta / Жиганган табекуну / 수집 날짜 / Paémimo data / Savákhšanas datums / Dato pravetakning / Data pobrania / Data de colheita / Data colectării / Data сбора / Dátum odberu / Datum prikupljanja / Uppsamlingsdatum / Toplama tarihi / Дата забора / 采集日期



µL/test / µL/rect / µL/Test / µL/εξέταση / µL/prueba / µL/teszt / µL/테스트 / µL/анализ / µL/检测



Keep away from light / Газете от светлина / Nevystavujte světu / Má ikke udsættes for lys / Vor Licht schützen / Краткото то макрия атпó тη φωτ / Mantener alejado de la luz / Hoida eemal valgusest / Conserver à l'abri de la lumière / Držati dalje od svjetla / Fény nem érheti / Tenere al riparo dalla luce / Қарашыланған жерде ұста / 빛을 피해야 함 / Laikyt atokiau nuo šilumos šaltinių / Sargāt no gaismas / Niet blootstellen aan zonlicht / Má ikke utsettes for lys / Przechowywać z dala od źródła światła / Manter ao abrigo da luz / Feriți de lumină / Хранить в темноте / Uchovávajte mimo dosahu svetla / Držite dalje od svetlosti / Får ej utsättas för ljus / Işiktan uzak tutun / Берегти від дії світла / 请远离光线



Hydrogen gas generated / Образуваен в водород газ / Možnost úniku phynného vodíku / Frembringer hydrogengas / Wasserstoffgas erzeugt / Δημιουργία αερίου υδρογόνου / Producción de gas de hidrógeno / Vesinikaasi tekkitäytä / Produit de l'hydrogène gazeux / Sadṛži hydrogen vodik / Hidrogén gáz fejleszt / Produzione di gas idrogeno / Газетек сутері пайда болды / 수소 가스 생성됨 / İşskiria vandenilio dujas / Rodas üdegradis / Waterstofgas gegeneréerd / Hydroengass generert / Powoduje powstawanie wodoru / Produção do gás de hidrogénio / Generare gaz de hidrogen / Выделение водорода / Vyrobené použítm vodíka / Osloboda se vodoník / Genererad vätgas / Açıga çıkan hidrojen gazi / Реакция в видленням водню / 会产生氢气



Patient ID number / ИД номер на пациент / ID pacienta / Patientens ID-nummer / Patienten-ID / Αριθμός αναγνώρισης ασθενούς / Número de ID del paciente / Patsiendi ID / No d'identification du patient / Identifikacijski broj pacijenta / Beleg azonosító száma / Numero ID paciente / Пациенттн идентификациялық немірі / 환자 ID 번호 / Paciente identifikavimo numeris / Pacienta ID numurs / Identificatienummer van de patiënt / Pasientens ID-nummer / Numer ID pacienta / Número da ID do doente / Număr ID pacient / Идентификационный номер пациента / Identifikačné číslo pacienta / ID broj pacijenta / Patientnummer / Hasta kimlik numarası / Идентификатор пацієнта / 患者标识号



Fragile, Handle with Care / Чупливо, Работете с необходимото внимание. / Krehké. Při manipulaci postupujte opatrne. / Forsigtig, kan gå i stykker. / Zerbrechlich, vorsichtig handhaben. / Εύθραυστο. Χειρίστε το με προσοχή. / Frágil. Manipular con cuidado. / Óm, káisítse eltevállalk. / Fragile. Manipuler avec précaution. / Lomljivo, rukujte pažljivo. / Törékeny! Óvatosan kezelendő. / Fragile, maneggiare con cura. / Сынъш, абылай пайдаланыңыз. / 조심 깨지기 쉬운 처리 / Trapu, elkítés atsargiai. / Trauslis; rikoties uzmanīgi / Breekaar, voorzichtig behandelen. / Ømtålig, håndter forsiktig. / Krucha zawartość, przenosić ostrożnie. / Frágil, Manuseio com Cuidado. / Fragil, manipulați cu atenție. / Хрупкое! Обращаться с осторожностью. / Krehké, vyzádjuje sa opatrná manipulácia. / Lomljivo - rukujte pažljivo. / Bräckligt. Hantera försiktigt. / Kolay Kirılır, Dikkatli Taşınır. / Тендентна, звертатися з обережністю / 易碎, 小心轻放

Rx Only

This only applies to US: "Caution: Federal Law restricts this device to sale by or on the order of a licensed practitioner." / S'applique uniquement aux États-Unis: "Caution: Federal Law restricts this device to sale by or on the order of a licensed practitioner." / Vale solo per gli Stati Uniti: "Caution: Federal Law restricts this device to sale by or on the order of a licensed practitioner." / Gilt nur für die USA: "Caution: Federal Law restricts this device to sale by or on the order of a licensed practitioner." / Solo se aplica a los EE.UU.: "Caution: Federal Law restricts this device to sale by or on the order of a licensed practitioner."



bd.com/e-labeling

KEY-CODE: 500008334

Europe, CH, GB, NO:		+800 135 79 135	
International:		+31 20 794 7071	
AR	+800 135 79 135	LT	8800 30728
AU	+800 135 79 135	MT	+31 20 796 5693
BR	0800 591 1055	NZ	+800 135 79 135
CA	+1 855 805 8539	RO	0800 895 084
CO	+800 135 79 135	RU	+800 135 79 135
EE	0800 0100567	SG	800 101 3366
GR	00800 161 22015 7799	SK	0800 606 287
HR	0800 804 804	TR	00800 142 064 866
IL	+800 135 79 135	US	+1 855 236 0910
IS	800 8996	UY	+800 135 79 135
LI	+31 20 796 5692	VN	122 80297



Becton, Dickinson and Company
7 Loveton Circle
Sparks, MD 21152 USA

EC REP

Benex Limited
Pottery Road, Dun Laoghaire
Co. Dublin, Ireland

ATCC® is a trademark of American Type Culture Collection.

BD, the BD Logo, BACTEC, FOS, Luer-Lok, Peds Plus, Safety-Lok, and Vacutainer® are trademarks of Becton, Dickinson and Company or its affiliates. © 2019 BD. All rights reserved.