June 14, 2021

bioMérieux, Inc
Esther Hernandez
Regulatory Affairs Specialist
595 Anglum Rd.
Hazelwood, Missouri 63042

Re: K211136
Trade/Device Name: VITEK 2 AST-Gram Negative Imipenem/Relebactam (<0.25/4 - >16/4 µg/mL)
Regulation Number: 21 CFR 866.1645
Regulation Name: Fully Automated Short-Term Incubation Cycle Antimicrobial Susceptibility System
Regulatory Class: Class II
Product Code: LON, LTW, LTT
Dated: April 15, 2021
Received: April 16, 2021

Dear Esther Hernandez:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmin/pmn.cfm identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part...
801 and Part 809); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.


For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance) and CDRH Learn (https://www.fda.gov/training-and-continuing-education/cdrh-learn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

Ribhi Shawar -S

Ribhi Shawar, Ph.D. (ABMM)
Chief
General Bacteriology and Antimicrobial Susceptibility Branch
Division of Microbiology Devices
OHT7: Office of In Vitro Diagnostics and Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure
Indications for Use

VITEK® 2 AST-Gram Negative Imipenem/Relebactam is designed for antimicrobial susceptibility testing of Gram negative bacilli and is intended for use with the VITEK® 2 and VITEK® 2 Compact Systems as a laboratory aid in the determination of in vitro susceptibility to antimicrobial agents. VITEK® 2 AST-Gram Negative Imipenem/Relebactam is a quantitative test. Imipenem/Relebactam has been shown to be active against most strains of the microorganisms listed below, according to the FDA label for this antimicrobial.

Active in vitro and in clinical infections:
- Acinetobacter calcoaceticus-baumannii complex
- Klebsiella (Enterobacter) aerogenes
- Enterobacter cloacae
- Escherichia coli
- Klebsiella pneumoniae
- Pseudomonas aeruginosa
- Citrobacter freundii
- Klebsiella oxytoca

In vitro data are available, but clinical significance is unknown:
- Citrobacter koseri
- Enterobacter asburiae

The VITEK® 2 Gram-Negative Susceptibility Card is intended for use with the VITEK® 2 Systems in clinical laboratories as an in vitro test to determine the susceptibility of clinically significant aerobic Gram-negative bacilli to antimicrobial agents when used as instructed.

Type of Use (Select one or both, as applicable)

☐ Prescription Use (Part 21 CFR 801 Subpart D)  ☐ Over-The-Counter Use (21 CFR 801 Subpart C)

CONTINUE ON A SEPARATE PAGE IF NEEDED.
510(k) SUMMARY

VITEK® 2 AST-GN Imipenem/Relebactam

A. 510(k) Submission Information:

Submitter’s Name: bioMérieux, Inc.
Address: 595 Anglum Road
          Hazelwood, MO  63042
Contact Person: Esther Hernandez
               Regulatory Affairs Specialist
Phone Number: 314-731-8841
Fax Number: 314-731-8689
Date of Preparation: April 08, 2021

B. Device Name:

Formal/Trade Name: VITEK® 2 AST- Gram Negative
                   Imipenem/Relebactam (≤ 0.25/4 – ≥ 16/4 μg/mL)
Classification Name: 21 CFR 866.1645
                     Fully Automated Short-Term Incubation Cycle
                     Antimicrobial Susceptibility System
                     Product Code LON
Common Name: VITEK® 2 AST-GN Imipenem/Relebactam

C. Predicate Device:

VITEK® 2 AST-GN Imipenem/Relebactam
(K193572)

D. Device Description:

The principle of the VITEK® 2 AST cards is based on the microdilution minimum inhibitory
concentration (MIC) technique reported by MacLowry and Marsh (1) and Gerlach (2). The
VITEK® 2 AST card is essentially a miniaturized, abbreviated and automated version of the
doubling dilution technique (3).
Each VITEK® 2 AST card contains 64 wells. A control well which only contains microbiological culture media is resident on all cards. The remaining wells contain premeasured portions of a specific antibiotic combined with culture media. The bacterial or yeast isolate to be tested is diluted to a standardized concentration with 0.45 – 0.5% saline before being used to rehydrate the antimicrobial medium within the card. The VITEK® 2 System automatically fills, seals and places the card into the incubator/reader. The VITEK® 2 Compact has a manual filling, sealing and loading operation. The VITEK® 2 Systems monitor the growth of each well in the card over a defined period of time. At the completion of the incubation cycle, a report is generated that contains the MIC value along with the interpretive category result for each antibiotic contained on the card.

VITEK® 2 AST-GN Imipenem/Relebactam has the following concentrations in the card: 0.25/4, 1/4, 4/4 and 16/4 µg/mL (equivalent standard method concentration by efficacy in µg/mL).

E. Substantial Equivalence Information

The similarities and differences of the VITEK 2 AST-GN Imipenem/Relebactam when compared to the predicate device, VITEK 2 AST-GN Imipenem/Relebactam (K193572), are described in the following table. The only difference between both devices is the addition of an Indications for Use, Acinetobacter calcoaceticus-baumannii complex.

<table>
<thead>
<tr>
<th>Item</th>
<th>Device: VITEK® 2 AST-GN Imipenem/Relebactam</th>
<th>Predicate: VITEK® 2 AST-GN Imipenem/Relebactam (K193572)</th>
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<tbody>
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<td><strong>Similarities</strong></td>
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### Test Methodology

- Automated quantitative antimicrobial susceptibility test for use with the VITEK® 2 and VITEK® 2 Compact Systems to determine the in vitro susceptibility of microorganisms

### Antimicrobial Agent

- Imipenem/Relebactam

### Inoculum

- Saline suspension of organism

### Test Card

- Gram-negative (AST-GN) Susceptibility Card

### Analysis Algorithms

- Growth Pattern Analysis

### Instrument

- VITEK® 2 and VITEK® 2 Compact
VITEK® 2 AST-GN Imipenem/Relebactam
Traditional 510(k) Submission

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**F. Intended Use:**

VITEK® 2 AST-Gram Negative Imipenem/Relebactam is designed for antimicrobial susceptibility testing of Gram Negative bacilli and is intended for use with the VITEK® 2 and VITEK® 2 Compact Systems as a laboratory aid in the determination of *in vitro* susceptibility to antimicrobial agents. VITEK® 2 AST-Gram Negative Imipenem/Relebactam is a quantitative test. Imipenem/Relebactam has been shown to be active against most strains of the microorganisms listed below, according to the FDA label for this antimicrobial.

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F. Performance Overview and Conclusion:

VITEK® 2 AST-GN Imipenem/Relebactam demonstrated substantially equivalent performance when compared with the CLSI broth microdilution reference method, as defined in the FDA Class II Special Controls Guidance Document: Antimicrobial Susceptibility Test (AST) Systems; Guidance for Industry and FDA (Issued August 28, 2009).

The Premarket Notification (Traditional 510[k]) presents data in support of VITEK® 2 AST-GN Imipenem/Relebactam. An external evaluation was conducted with fresh and stock clinical isolates, as well as a set of challenge strains. The external evaluations were designed to confirm the acceptability of VITEK® 2 AST-GN Imipenem/Relebactam by comparing its performance with the CLSI broth microdilution reference method incubated at 16-20 hours. The data is representative of performance on both the VITEK® 2 and VITEK® 2 Compact instrument platforms.

The VITEK 2 AST-GN Imipenem/Relebactam demonstrated acceptable performance of 98.2% Essential Agreement and 98.2% Categorical Agreement for the Acinetobacter calcoaceticus-baumannii complex compared with the reference method.

References:

