BAYER HEALTHCARE
JENNIFER GREGORY
REGULATORY AFFAIRS SPECIALIST
430 SOUTH BEIGER ST.
MISHAWAKA IN  46544

Re: K150942
Trade/Device Name: Contour Next USB Blood Glucose Monitoring System
Regulation Number: 21 CFR 862.1345
Regulation Name: Glucose test system
Regulatory Class: II
Product Code: NBW, LFR
Dated: October 29, 2015
Received: October 30, 2015

Dear Jennifer Gregory:

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. 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 Parts 801 and 809); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.
If you desire specific advice for your device on our labeling regulations (21 CFR Parts 801 and 809), please contact the Division of Industry and Consumer Education at its toll-free number (800) 638 2041 or (301) 796-7100 or at its Internet address http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm. Also, please note the regulation entitled, “Misbranding by reference to premarket notification” (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH’s Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Industry and Consumer Education at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours,

Katherine Serrano -S

For:  Courtney H. Lias, Ph.D.
    Director
    Division of Chemistry and Toxicology Devices
    Office of In Vitro Diagnostics
    and Radiological Health
    Center for Devices and Radiological Health

Enclosure
Indications for Use

The Contour Next USB blood glucose monitoring system is an over the counter (OTC) device utilized by persons with diabetes in home settings for the quantitative measurement of glucose in whole blood, and is for single-patient use only and should not be shared.

The system consists of a Contour Next USB blood glucose meter, Contour Next test strips and Contour Next control solutions.

The Contour Next USB blood glucose monitoring system is indicated for use with fresh capillary whole blood samples drawn from the fingertip and palm only. Alternative site testing should be done only during steady state times (when glucose is not changing rapidly).

The Contour Next USB blood glucose monitoring system is not intended for the diagnosis of or screening for diabetes mellitus and is not intended for use on neonates.

The Contour Next Test Strips are for use with the Contour Next USB blood glucose monitoring system for the quantitative measurement of glucose in whole blood.

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.

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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The burden time for this collection of information is estimated to average 79 hours per response, including the time to review instructions, search existing data sources, gather and maintain the data needed and complete and review the collection of information. Send comments regarding this burden estimate or any other aspect of this information collection, including suggestions for reducing this burden, to:

Department of Health and Human Services
Food and Drug Administration
Office of Chief Information Officer
Paperwork Reduction Act (PRA) Staff
PRASstaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."
510(k) Summary

Date prepared: November 23, 2015

According to the requirements of 21 CFR 807.92, the following information is being submitted in sufficient detail to provide an understanding of the basis for a determination of substantial equivalence.

1) Submitter
Jennifer Gregory
Regulatory Affairs Specialist
Bayer Healthcare LLC
430 South Beiger Street
Mishawaka, IN 46544
Telephone: (574) 256-3447
Fax: (574) 256-3519

2) Device name:
Trade name: Bayer Contour® Next USB Blood Glucose Monitoring System
Common name: Blood Glucose Meter
Classification name: Blood Glucose Test System, Over-the-Counter, 75 NBW, LFR (21 CFR § 862.1345)

3) Predicate device:
Contour Next USB Blood Glucose Meter (Reference: Contour Next USB Blood Glucose Monitoring System (K121087))

4) Device description:
The Contour Next USB Blood Glucose Meter consists of a small handheld blood glucose meter that utilizes dry reagent test strips for the measurement of glucose in capillary whole blood by persons with diabetes. Liquid control solution is used to check the performance of the system. The meter, together with the test strips and control solutions, is referred to as the Contour Next USB Blood Glucose Monitoring System.

The chemical principle utilized for both the predicate and modified devices is based on measurement of electrical current caused by the reaction of glucose in the blood with chemicals on the reagent strip. The blood sample is drawn into the tip of the reagent strip through capillary action. Glucose in the sample reacts with FAD glucose dehydrogenase (FAD-GDH) enzyme on the reagent strip. The electrons generated by this reaction are shuttled to an
5) Intended Use: The Contour Next USB blood glucose monitoring system is an over the counter (OTC) device utilized by persons with diabetes in home settings for the quantitative measurement of glucose in whole blood, and is for single-patient use only and should not be shared.

The system consists of a Contour Next USB blood glucose meter, Contour Next test strips and Contour Next control solutions.

The Contour Next USB blood glucose monitoring system is indicated for use with fresh capillary whole blood samples drawn from the fingertip and palm only. Alternative site testing should be done only during steady state times (when glucose is not changing rapidly).

The Contour Next USB blood glucose monitoring system is not intended for the diagnosis of or screening for diabetes mellitus and is not intended for use on neonates.

The Contour Next Test Strips are for use with the Contour Next USB blood glucose monitoring system for the quantitative measurement of glucose in whole blood.

Data demonstrating substantial equivalence

The Contour Next USB Blood Glucose Meter consists of a small handheld blood glucose meter that is substantially equivalent to the predicate device, the Contour Next USB Blood Glucose Meter (K121087). The modified and predicate devices use the same glucose calculation algorithm. Both devices also use dry reagent test strips for the measurement of glucose in capillary whole blood by persons with diabetes and liquid controls to check the performance of the system. The same Contour Next test strips and Contour Next control solutions are used by both the modified and predicate devices.

A detailed comparison of the characteristics featured between the modified and predicate devices is provided in the tables below:
### Summary of the Technological Characteristics of the Modified Device Compared to Predicate

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Predicate Contour Next USB (K121087)</th>
<th>Contour Next USB (Modified Device)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test Strip</td>
<td>Contour Next Test Strips</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Control Solution</td>
<td>Contour Next Control Solution (Level 1 and 2)</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Detection Method</td>
<td>Amperometric</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Measuring Range</td>
<td>20-600 mg/dL</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Sample Volume</td>
<td>0.6 µL</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Countdown time displayed</td>
<td>5 Seconds</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Illuminated Strip Port</td>
<td>Yes</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Operational Buttons</td>
<td>4</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Battery Type</td>
<td>Rechargeable (3.4-4.2V)</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Operating Temperature Range</td>
<td>41°-113° F</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Operating Humidity Range</td>
<td>10-93% RH</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Hematocrit Range</td>
<td>15%-65%</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Meter life</td>
<td>5 Years</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Validated Product Used for Cleaning and Disinfection</td>
<td>Clorox Germicidal wipes</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Meal Markers</td>
<td>Yes</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Characteristic</td>
<td>Predicate Contour Next USB (K121087)</td>
<td>Contour Next USB (Modified Device)</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>--------------------------------------</td>
<td>-----------------------------------</td>
</tr>
<tr>
<td>Calibration/Coding</td>
<td>Autocoding (no coding for users)</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>User Interface</td>
<td>Alphanumeric, Iconic, Native Language</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Display (technology)</td>
<td>Graphical (OLED)</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Display Visibility</td>
<td>Day and night</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Communication Port</td>
<td>USB Interface</td>
<td></td>
</tr>
<tr>
<td>Communication Link to Computer</td>
<td>Direct USB connection or optional USB cable</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>Test Results in Memory</td>
<td>2000 Results</td>
<td>Same as Predicate</td>
</tr>
<tr>
<td>User Contact Materials/Surface Finish</td>
<td>Polycarbonate/ABS with AS plastics with mirror or matte finish</td>
<td>Same as Predicate</td>
</tr>
</tbody>
</table>
## DIFFERENCES from Predicate

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Predicate Contour Next USB (K121087)</th>
<th>Contour Next USB (Modified Device)</th>
<th>Risk Assessment Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Improved detection of test strips that may have been exposed to a chemical that can degrade the mediator.</td>
<td>No</td>
<td>Yes</td>
<td>The error check improves the ability of the modified meter to detect exposed test strips and provide an error message instead of a high biased result.</td>
</tr>
<tr>
<td>Improved detection of un-mixed control solution.</td>
<td>No</td>
<td>Yes</td>
<td>The error check improves the ability of the modified meter to detect un-mixed control test solutions and provide an error message instead of a high biased result.</td>
</tr>
<tr>
<td>Improved detection of sample 'perturbation' during a test.</td>
<td>No</td>
<td>Yes</td>
<td>The error check improves the ability of the modified meter to detect a sample that is disturbed during the countdown period and provide an error message instead of a biased result.</td>
</tr>
</tbody>
</table>
Summary of Performance testing

Bench testing was conducted to ensure that the error checks acted as intended and gave error messages for:

• test strips that had been exposed to a reducing agent
• un-mixed control test solution
• samples that had been disturbed during the test countdown

Software verification testing was conducted to ensure that no good results were classified as errors as a result of the modifications.

Equivalency testing was conducted to ensure the performance of the modified and predicate meters was not statistically different.

An internal user study was conducted to ensure the modifications did not impact the blood glucose result accuracy of the Contour Next USB system.

Conclusions from Performance Evaluations

Based on the outcome of the performance testing conducted, the modified Contour Next USB Blood Glucose Meter is substantially equivalent to the predicate Contour Next USB Blood Glucose Meter (K121087).