Zebra Medical Vision, Ltd.                                      November 26, 2019
% Flair Bar
VP Operations
Shefayim Commercial Center
PO Box 25
Shefayim, 6099000
ISRAEL

Re: K192320
  Trade/Device Name: HealthCXR
  Regulation Number: 21 CFR 892.2080
  Regulation Name: Radiological computer-assisted prioritization software
  Regulatory Class: Class II
  Product Code: QFM
  Dated: November 3, 2019
  Received: November 6, 2019

Dear Flair Bar:

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/cfpmn/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
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,

Thalia T Mills, Ph.D.
Director
Division of Radiological Health
OHT7: Office of In Vitro Diagnostics and Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure
510(k) Number (if known)
K192320

Device Name
HealthCXR

Indications for Use (Describe)
The Zebra HealthCXR device is a software workflow tool designed to aid the clinical assessment of adult Chest X-Ray cases with features suggestive of pleural effusion in the medical care environment. HealthCXR analyzes cases using an artificial intelligence algorithm to identify suspected findings. It makes case-level output available to a PACS/workstation for worklist prioritization or triage. HealthCXR is not intended to direct attention to specific portions or anomalies of an image. Its results are not intended to be used on a stand-alone basis for clinical decision-making nor is it intended to rule out pleural effusion or otherwise preclude clinical assessment of X-Ray cases.

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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Office of Chief Information Officer
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5. 510 (k) Summary

510(K) Summary - HealthCXR
Zebra Medical Vision Ltd.

510(k) Number – K192320

I. Applicant’s Name: Flair Bar, VP Operations
   Zebra Medical Vision Ltd.
   Shefayim Commercial Center
   PO Box 25
   Shefayim, 6099000
   ISRAEL
   Telephone: +972-9-8827795
   Fax: +972-9-8827795

   Date Prepared: November 22, 2019

II. Device

   Trade Name: HealthCXR

   Classification Name: QFM - Radiological Computer-Assisted Prioritization Software

   Regulation Number: 892.2080

   Classification: Class II, Radiology

III. Predicate Device:
The HealthCXR device is substantially equivalent to the following device:

<table>
<thead>
<tr>
<th>Proprietary Name</th>
<th>HealthPNX</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premarket Notification</td>
<td>K190362</td>
</tr>
<tr>
<td>Classification Name</td>
<td>Radiological Computer-Assisted Prioritization Software</td>
</tr>
<tr>
<td>Regulation Number</td>
<td>21 CFR 892.2080</td>
</tr>
<tr>
<td>Product Code</td>
<td>QFM</td>
</tr>
<tr>
<td>Regulatory Class</td>
<td>II</td>
</tr>
</tbody>
</table>

IV. Device Description
The HealthCXR solution is a software product that automatically identifies suspected findings on chest x-rays (e.g. pleural effusion) and notifies PACS/workstation of the presence of this critical finding in the scan. This notification allows for worklist prioritization of the identified scan and assists clinicians in viewing the prioritized scan before others. The device aim is to aid in prioritization and triage of radiological medical images only.

The software is automatic and is capable of analyzing PA or AP chest x-rays. If a suspected finding is found in a scan, the alert is automatically sent to the PACS/workstation used by the radiologist or to a standalone desktop application in parallel with the ongoing standard of care. The PACS/workstation prioritizes and displays the study through its worklist interface. The standalone desktop application, Zebra Worklist, includes a compressed preview image meant for informational purposed only and is not intended for diagnostic use.

The HealthCXR device works in parallel to and in conjunction with the standard care of workflow. After a chest x-ray has been performed, a copy of the study is automatically retrieved and processed by the HealthCXR device. The device performs the analysis of the study and returns a notification about the relevant pathology to the PACS/workstation which prioritizes it through the worklist interface or alternatively, the Zebra Worklist will notify the user through the standalone desktop application. The clinician is then able to review the study earlier than in standard of care workflow.

The software does not recommend treatment or provide a diagnosis. It is meant as a tool to assist in improved workload prioritization of critical cases. The final diagnosis is provided by a radiologist after reviewing the scan itself.

The following modules compose the HealthCXR software for Pleural Effusion:

**Data input and validation:** Following retrieval of a study, the validation feature assessed the input data (i.e. age, modality, view) to ensure compatibility for processing by the algorithm.

**Pleural Effusion algorithm:** Once a study has been validated, the algorithm analyzes the frontal chest x-ray for detection of suspected finding suggestive of pleural effusion.

**IMA Integration feature:** The study analysis and the results of a successful study analysis is provided to IMA, to then be sent to the PACS/workstation for prioritization.

**Error codes feature:** In the case of a study failure during data validation or the analysis by the algorithm, an error is provided to the system.

**V. Intended Use/Indication for Use:**

The Zebra HealthCXR device is a software workflow tool designed to aid the clinical assessment of adult Chest X-Ray cases with features suggestive of pleural effusion in the medical care
environment. HealthCXR analyzes cases using an artificial intelligence algorithm to identify suspected findings. It makes case-level output available to a PACS/workstation for worklist prioritization or triage. HealthCXR is not intended to direct attention to specific portions or anomalies of an image. Its results are not intended to be used on a stand-alone basis for clinical decision-making nor is it intended to rule out pleural effusion or otherwise preclude clinical assessment of X-Ray cases.

VI. Technological Characteristics Compared to Predicate Device:

The technological characteristics, e.g., overall design, mechanism of action, mode of operation, performance characteristics, etc., and the intended use of the HealthCXR device are substantially equivalent to the predicate device cited above.

A comparison of the technological characteristics with the predicate is summarized below.

<table>
<thead>
<tr>
<th>Technological Characteristics</th>
<th>Proposed Device HealthCXR</th>
<th>Predicate Device HealthPNX (K190362)</th>
<th>Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indication for Use/Intended Use</td>
<td>The Zebra HealthCXR device is a software workflow tool designed to aid the clinical assessment of adult Chest X-Ray cases with features suggestive of pleural effusion in the medical care environment. HealthCXR analyzes cases using an artificial intelligence algorithm to identify suspected findings. It makes case-level output available to a PACS/workstation for worklist prioritization or triage. HealthCXR is not intended to direct attention to specific portions or anomalies of an image. Its results are not intended to be used on a stand-alone basis for clinical decision-making nor is it intended to rule out Pleural Effusion or otherwise preclude clinical assessment of X-Ray cases.</td>
<td>The Zebra Pneumothorax device is a software workflow tool designed to aid the clinical assessment of adult Chest X-Ray cases with features suggestive of Pneumothorax in the medical care environment. HealthPNX analyzes cases using an artificial intelligence algorithm to identify suspected findings. It makes case-level output available to a PACS/workstation for worklist prioritization or triage. HealthPNX is not intended to direct attention to specific portions or anomalies of the image. Its results are not intended to be used on a stand-alone basis for clinical decision-making nor is it intended to rule out Pneumothorax or otherwise preclude clinical assessment of X-Ray cases.</td>
<td>Similar expect for lesion type</td>
</tr>
<tr>
<td>Notification-only, parallel workflow tool</td>
<td>Yes</td>
<td>Yes</td>
<td>Same</td>
</tr>
<tr>
<td>User</td>
<td>Radiologist</td>
<td>Radiologist</td>
<td>Same</td>
</tr>
<tr>
<td>Radiological images format</td>
<td>DICOM</td>
<td>DICOM</td>
<td>Same</td>
</tr>
<tr>
<td>---------------------------</td>
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</tr>
<tr>
<td>Identify patients with prespecified clinical condition</td>
<td>Yes</td>
<td>Yes</td>
<td>Same</td>
</tr>
<tr>
<td>Clinical condition</td>
<td>Pleural Effusion</td>
<td>Pneumothorax</td>
<td>Different but as per the product classification definition, both identify “time sensitive imaging.”</td>
</tr>
<tr>
<td>Alert to finding</td>
<td>Yes; notification flagged for review on hospital worklist or Zebra list</td>
<td>Yes; notification flagged for review on hospital worklist.</td>
<td>Similar, HealthCXR can be directly integrated for notification on the hospital worklist or on the Zebra list application. Both notifications operate in parallel with the standard of care.</td>
</tr>
<tr>
<td>Independent of standard of care workflow</td>
<td>Yes; No cases are removed from worklist</td>
<td>Yes; No cases are removed from worklist</td>
<td>Same</td>
</tr>
<tr>
<td>Modality</td>
<td>X-Ray</td>
<td>X-Ray</td>
<td>Same</td>
</tr>
<tr>
<td>Body part</td>
<td>Chest</td>
<td>Chest</td>
<td>Same</td>
</tr>
<tr>
<td>Artificial Intelligence algorithm</td>
<td>Yes</td>
<td>Yes</td>
<td>Same</td>
</tr>
<tr>
<td>Limited to analysis of imaging data</td>
<td>Yes</td>
<td>Yes</td>
<td>Same</td>
</tr>
<tr>
<td>Aids prompt identification of cases with indicated findings</td>
<td>Yes</td>
<td>Yes</td>
<td>Same</td>
</tr>
<tr>
<td>Preview Image</td>
<td>Presentation of a preview of the study for initial assessment, not meant for diagnostic purposes. The device operated in parallel with the standard of care, which remains the default option for all cases.</td>
<td>Presentation of notification for initial assessment not meant for diagnostic purposes. The device operates in parallel with the standard of care, which remains the default option for all cases.</td>
<td>Similar, HealthCXR provides an additional thumbnail view of the original exam as preview only on the Zebra list or Radiology Assistant, not for diagnostic use.</td>
</tr>
<tr>
<td>Multiple operating points</td>
<td>Yes; 2 optional operating points</td>
<td>No; single operating point</td>
<td>Different, but both operating points are substantially equivalent to the performance predicate device and comply with DEN 170073 Special control 1(iii).</td>
</tr>
</tbody>
</table>
VII. Performance Data:

Safety and performance of HealthCXR has been evaluated and verified in accordance with software specifications and applicable performance standards through Software Development and Validation & Verification Process to ensure performance according to specifications, User Requirements and Federal Regulations and Guidance documents, “Guidance for the Content of Premarket Submissions for Software Contained in Medical Devices”.

The performance of the HealthCXR device has been validated in a performance study for triage of time sensitive chest X-Ray cases. The data included a retrospective cohort of 554 anonymized Chest X-ray cases from the USA and Israel, including 276 cases positive for Pleural Effusion and 278 cases negative for Pleural Effusion, as well as confounding imaging factors. The validation data set was truthed (ground truth) by three US Board-Certified Radiologists (truthers). The stand-alone detection accuracy was measured on this cohort respective to the ground truth.

The HealthCXR device detection accuracy met the accuracy performance goals for AUC, sensitivity, and specificity for two defined operating points. Overall, the HealthCXR was able to demonstrate an area under the curve (AUC) of 0.9885 (95% CI: [0.9815, 0.9956]), which is both comparable to the predicate device, and exceeds the required technical method under the QFM product code. The device establishes effective triage based on an AUC >95%. The sensitivity and specificity of the HealthCXR was reported for two operating points, the first for equal sensitivity and specificity and the second operating point for a higher specificity. At both operating points, the HealthCXR performance met the performance goal and was found to be substantially equivalent to the predicate device, HealthPNX (K190362). The first operating point showed a sensitivity of 96.74% (95% CI: [92.79; 96.48]) and a specificity of 93.17% (95% CI: [89.57; 95.58]). The second, “high-specificity” operating point, reported a sensitivity of 93.84% (95% CI: [90.36;96.12%]) and a specificity of 97.12% (95% CI: [94.43 ;98.53]).

In addition, we assessed the performance time of the HealthCXR that reflects the time it takes for the device to analyze the study and send a notification to the worklist. The HealthCXR device mean processing time was 27.76 seconds for the “Equal Sensitivity and Specificity” operating point, and 20.18 seconds for the “High Specificity” operating point, a timing performance that is substantially equivalent to the predicate. This establishes that the HealthCXR meets its intended use statement.

VIII. Conclusion
The subject HealthCXR device and the HealthPNX predicate device are both software-only devices intended to aid in triage of radiological images, independent of standard of care workflow. The labeling of both devices are limited to the categorization of exams and are not to be used in-lieu of full patient evaluation or relied upon to make or confirm diagnosis.

Both devices operate in parallel to the standard of care workflow in the sense that they do not change the original image, do not provide any marking, and do not remove cases from the standard of care. The minor differences between the subject device and the predicate raise no new issues of safety or effectiveness. In addition, performance testing demonstrates that the HealthCXR performs as intended. The HealthCXR device is therefore substantially equivalent to the HealthPNX predicate.