



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration
10903 New Hampshire Avenue
Document Control Center - WO66-G609
Silver Spring, MD 20993-0002

Qubyx Software Technologies Inc.
Marc Leppla
Director, CTO
501 Silverside Road, Suite 105
Wilmington, Delaware 19809

May 26, 2017

Re: K171229

Trade/Device Name: DELL UP3017 with QUBYX PerfectLum bundle
Regulation Number: 21 CFR 892.2050
Regulation Name: Picture Archiving And Communications System
Regulatory Class: Class II
Product Code: PGY
Dated: April 20, 2017
Received: April 27, 2017

Dear Marc Leppla:

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 Part 801); 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 regulation (21 CFR Part 801), 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,

A handwritten signature in black ink, appearing to read "Robert Ochs", is positioned over a large, light blue, semi-transparent "FDA" logo.

Robert Ochs, Ph.D.
Director
Division of Radiological Health
Office of In Vitro Diagnostics
and Radiological Health
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES
Food and Drug Administration

Indications for Use

Form Approved: OMB No. 0910-0120
Expiration Date: January 31, 2017
See PRA Statement below.

510(k) Number (if known)

K171229

Device Name

Dell UP3017 with QUBYX PerfectLum bundle

Indications for Use (Describe)

The DELL UP3017 with QUBYX PerfectLum is intended to be used for displaying and viewing medical images, for review and analysis by trained medical practitioners.

The DELL UP3017 can be used only in conjunction with QUBYX PerfectLum.

The device can not be used in primary image diagnosis in mammography.

The device can not be used for a life-support system.

The device does not contact with the patient.

The device is intended for prescription use.

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.

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510(K) Summary

as required by 807.92

1. Company Identification

Qubyx Software Technologies Inc.
501 Silverside Road, Suite 105
Wilmington, DE
19809
USA
Tel: +1 518 618 1012

2. Official Correspondent

Mr. Marc Leppla

President and CTO (Chief Technical Officer)
leppla@qubyx.com

3. Date of Submission

Feb 20, 2017

4. Device Trade name

DELL UP3017 with QUBYX PerfectLum bundle

5. Common/Usual Name

Image display system, Color LCD Monitor, image monitor/display

6. Classification Number

Medical displays classified in Class II per 21 CFR 892.2050

7. Predicate device 1

Name: DELL U3014

Manufacturer: Qubyx Software Technologies Inc.

510(k) number: K131601

8. Device description

The DELL UP3017 with QUBYX PerfectLum is a 30" color display for medical viewing. It provides 2560 x 1600 resolution with an adjustable Look Up Table and a 10 bit panel. It is combined with QUBYX PerfectLum and PerfectLum remote management, a user-friendly DICOM calibration and AAPM TG18 verification software suite. The software allows setting the display function to DICOM, displaying test pattern and performing acceptance and constancy tests.

9. Intended use

The DELL UP3017 with QUBYX PerfectLum is intended to be used for displaying and viewing of digital images, for review and analysis by trained medical practitioners.

The DELL UP3017 must only be used in conjunction with QUBYX PerfectLum.

The device must not be used in primary image diagnosis in mammography.

The device can not be used for a life-support system and does not contact with the patient.

10. Comparison table – predicate device 1

	DELL UP3017 with PerfectLum	Predicate device DELL U3014 510(k) number: K131601
Panel Type	IPS	IPS
Panel size	30" viewable	30" viewable
Native Resolution	2560 x 1600	2560 x 1600
Aspect Ratio	16:10	16:10
Pixel Pitch	0.25 mm	0.25 mm
Brightness (typical)	350 cd/m2	350 cd/m2
Contrast Ratio (typical)	1:1000	1:1000
Viewing Angle (typical)	178° Vert., 176° Hor.	178° Vert., 178° Hor.
Displayable Colors	1.07 billion colors	1.07 billion colors
Backlight	White LED	RGB LED
DICOM calibration and AAPM verification software	bundled	bundled
Indications for Use	The DELL UP3017 with QUBYX PerfectLum is intended to be used for displaying and viewing medical images, for review and analysis by trained medical practitioners. The DELL UP3017 can be used only in conjunction with QUBYX PerfectLum. The device can not be used in primary image diagnosis in mammography. The device can not be used for a life-support system. The device does not contact with the patient.	The DELL U3014 with QUBYX PerfectLum is intended to be used for displaying and viewing medical images, for review and analysis by trained medical practitioners. The DELL U3014 can be used only in conjunction with QUBYX PerfectLum. The device can not be used in primary image diagnosis in mammography. The device can not be used for a life-support system. The device does not contact with the patient.

DELL UP3017 is the direct successor of the DELL U3014 model in the product line of UltraSharp displays, so the DELL UP3017 with PerfectLum has exactly the same technical characteristics that predicate device DELL U3014 with PerfectLum, except backlight type. Backlight type of the U3014 model is RGB LED, while backlight type of the UP3017 model is White LED. The next generation used a white light LED to save on power. The comparison table shows that the subject device (DELL UP3017 with PerfectLum) has the same intended use as the predicate device.

Although the devices have different backlight types, these differences do not make the subject device less safe and reliable, so the subject device fits for displaying and viewing of digital images for diagnosis by trained physicians as the predicate device does.

Both devices are compliant with DICOM Part 14 GSDF and AAPM TG18 standards, which is tested and verified by University of Arizona. To verify DICOM and AAPM compliance for the subject device, AAPM acceptance test and DICOM conformance test were also performed by QUBYX.

Details of testing:

To verify DICOM conformance, a DICOM conformance test was performed, using QUBYX PerfectLum software and an X-Rite i1 Display Pro measurement device.

The test procedure was generated by the software in accordance with the requirements of the DICOM standard. It consisted of measurement steps, where the meter measured display's characteristics and the software recorded them. Then the software analyzed the results in comparison with target values, defined by DICOM standard, and generated the report, stating that the display is DICOM-conformant.

The display device has successfully passed DICOM conformance test, so it is compliant with DICOM Part 14 GSDF standard. So is the predicate device, so the two devices are substantially equivalent in this regard.

To verify AAPM TG18 conformance, an acceptance test was performed, using QUBYX PerfectLum software and an X-Rite i1 Display Pro measurement device.

The test procedure was generated by the software in accordance with the requirements of the AAPM TG18 standard and consisted of measurement and visual parts.

During the measurement steps, the meter measured display's characteristics and the software recorded them. During the visual steps, the user analyzed test patterns, generated by the software in accordance with AAPM standard. The software recorded the user's answers. Then the software analyzed the results in comparison with target values, defined by AAPM standard, and generated the report, stating that the display passes AAPM TG18 acceptance test.

The display device has successfully passed AAPM TG18 acceptance test, so it is compliant with AAPM TG18 standard and can be used as a primary category display for interpretation of medical images. The same is true for the predicate device, so the two devices are substantially equivalent in this regard.

Both devices have the same indications for use.

The new and predicate devices are substantially equivalent in the areas of technical characteristics, general function, application and intended use.