

Important Information and User's Manual amendments (for USA)

Changed the FDA statement and changed labels

Statements of compliance on the manufacturer's ID label on the bottom of the projector

Original	Complies with 21 CFR 1040.10 and 1040.11 except for conformance as a Risk Group 2 LIP as defined in IEC 62471-5:Ed. 1.0. For more information see <u>Laser Notice No. 57</u> , dated May 8, 2019.
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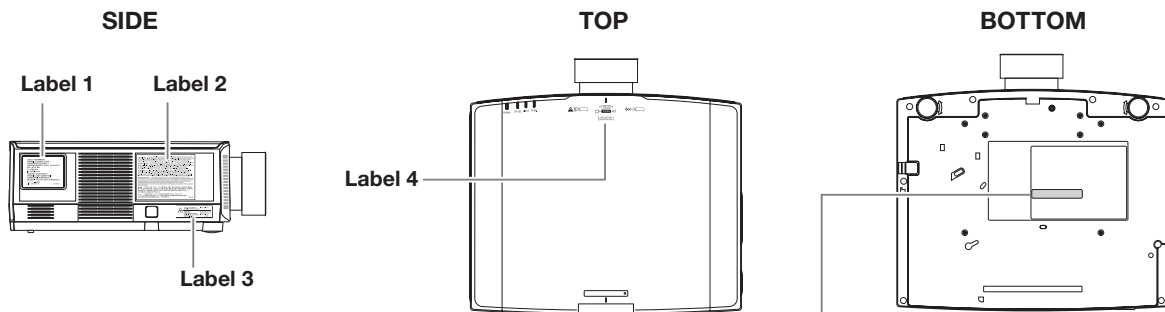


Correct	Complies with FDA performance standards for laser products except for deviations pursuant to <u>Laser Notice No. 50</u> , dated June 24, 2007.
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Labels

! WARNING

- The following labels are stuck on the projector.

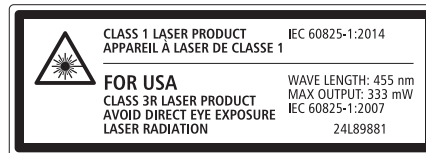


Changed the Statements of compliance

Label 1 Laser explanatory label



Label 3 * Laser explanatory label for USA



Label 2 Lens replacement caution label



Label 4 * Laser aperture label



* The Label 3 and 4 are added for Laser Notice No. 50.

