



Test Verification of Conformity

Verification Number: 201225119GZU -VOC002

On the basis of the referenced test report(s), sample(s) tested of the below product have been found to comply with the standards harmonized with the directives listed on this verification at the time the tests were carried out. Other standards and Directives may be relevant to the product. This verification is part of the full test report(s) and should be read in conjunction with it <them>.

Once compliance with all product relevant  mark directives are verified, including any relevant e.g. risk assessment and production control, the manufacturer may indicate compliance by signing a Declaration of Conformity themselves and applying the mark to products identical to the tested sample(s).

Applicant Name & Address:	Qierling (Beijing) Health Technology Co., Ltd. No 101-42,101-43(Dongsheng district),9th Floor, No 1 Building, No 8th, Heiquan Road, Haidian District, Beijing City.
Product Description:	Air Purifiering Disinfector
Models/Type References:	DS-S800, DS-X1000W
Ratings & Principle Characteristics:	220V-240V, 50Hz, 90W for model DS-S800 220V-240V, 50Hz, 120W for model DS-X1000W
Brand Name(s):	
Standard(s)/Directive(s):	Refer to Appendix
Verification Issuing Office Name & Address:	Intertek Testing Services Shenzhen Ltd. Guangzhou Branch Room 02, & 101/E201/E301/E401/E501/E601/E701/E801 of Room 01 1-8/F., No. 7-2. Caipin Road, Science City, GETDD, Guangzhou, Guangdong, China
Date of Tests:	06 January 2021-07 February 2021
Test Report Number(s):	201225119GZU-001 201225119GZU-002 201225119GZU-003 201225119GZU-004 201225119GZU-005 201225119GZU-006

Additional information in Appendix.



Signature

Name: Helen Ma

Position: Team Leader

Date: 26 February 2021

APPENDIX: Test Verification of Conformity

This is an Appendix to Test Verification of Conformity Number: 201225119GZU -VOC002

Standard(s)/Directive(s):

ETSI EN 300 328 V2.2.2 (2019-07)
ETSI EN 300 330 V2.1.1 (2017-02)
ETSI EN 301 489-17 V3.2.4(2020-09)
ETSI EN 301 489-1 V2.2.3(2019-11)
ETSI EN 301 489-1 V2.2.0 (2017-03)
ETSI EN 301 489-3 V2.1.1 (2019-03)
EN 55014-1:2017+A11:2020
EN IEC 61000-3-2:2019
EN 61000-3-3:2013+A1:2019
EN 55014-2:2015
EN IEC 62311:2020

Radio Equipment Directive (2014/53/EU) – Article 3.1(a)(health), 3.1(b) & article 3.2

Helen Ma

Signature

Name: Helen Ma

Position: Team Leader

Date: 26 February 2021

This Verification is for the exclusive use of Intertek's client and is provided pursuant to the agreement between Intertek and its Client. Intertek's responsibility and liability are limited to the terms and conditions of the agreement. Intertek assumes no liability to any party, other than to the Client in accordance with the agreement, for any loss, expense or damage occasioned by the use of this Verification. Only the Client is authorized to permit copying or distribution of this Verification. Any use of the Intertek name or one of its marks for the sale or advertisement of the tested material, product or service must first be approved in writing by Intertek. The observations and test/inspection results referenced in this Verification are relevant only to the sample tested/inspected. This Verification by itself does not imply that the material, product, or service is or has ever been under an Intertek certification program.