

# FEDERAL COMMUNICATIONS COMMISSION

## DECLARATION OF CONFORMITY (DoC)

Equipment: Electronic stethoscope  
Trademark(s) and Model(s): iMED+ / DS101, Omni-Steth  
Manufacturer: IMEDIPLUS INC.  
FCC ID in case other parts of this equipment are subject to certification:  
2AM7NDS101

**This device complies with Part 15 of the FCC Rules.**

**Operation is subject to the following two conditions:**

- (1) this device may not cause harmful interference, and**
- (2) this device must accept any interference received, including interference that may cause undesired operation.**

The following test reports are subject to this declaration:

Test report number: Issue date:  
**MW/2018/10049 Feb. 21, 2018**

The following manufacturer/importer/entity is responsible for this declaration:

Company name: IMEDIPLUS INC.  
Name/Title (legal representative): CHEN, CHIH-HAO/Special Assistant  
Address: 2F, 12 Sheng Yi Rd. Sec. 2, Chupei City, Hsinchu  
County 302, Taiwan (R.O.C).  
Phone: +886-3-6587700 ext.602  
Fax: +886-3-6589535  
E-mail: elio.chen@imediplus.com  
Date: 2018/03/05  
Signature:

