

**Guangdong Launca Medical Device Technology Co., Ltd.**  
Room 901-908 and 914-916, Building 5, No.1 Yanfa Road, Songshan Lake Park,  
Dongguan, Guangdong, China

Federal Communications Commission  
Authorization and Evaluation Division  
Equipment Authorization Branch  
7435 Oakland Mills Road  
Columbia, MD 21046

2023-7-5

To whom it may concern:

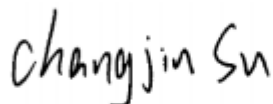
We, **Guangdong Launca Medical Device Technology Co., Ltd.** , hereby certify that the equipment for which authorization is sought (FCC ID: 2BBRU-DL300W) is not "covered" equipment prohibited from receiving an equipment authorization pursuant to § 2.903 of the FCC rules.

We also certify that, as of the date of the filing of the application, we are not identified on the Covered List (<https://www.fcc.gov/supplychain/coveredlist>) as an entity producing "covered" equipment.

By signing this form, we confirm the above and that we are aware of the application requirements listed under § 2.911(b) and (d)(5) and (6).

The information provided in this letter is based on the referenced rule parts of Title 47 of the CFR as well as KDB 986446 D01.

Sincerely,



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Signature

Changjin Su

Manager

Guangdong Launca Medical Device Technology Co., Ltd.