

BioSerenity USA, Inc.

Attention: Application Examiner

Re: Covered List, located at <https://www.fcc.gov/supplychain/coveredlist> and Report & Order 22-84 located at <https://docs.fcc.gov/public/attachments/DOC-389524A1.pdf>

Applicant: [**BioSerenity USA, Inc.**]

FCC ID: [2A7VI-ACCV3]

To whom it may concern,

Pursuant to FCC Report & Order 22-84 section IV. Interim Freeze Order, paragraphs 264, 265, and 266, as an authorized representative of [**BioSerenity USA, Inc.**], I attest that the equipment for which we are seeking equipment authorization through certification is not prohibited from receiving an equipment authorization; and [**BioSerenity USA, Inc.**] is not an entity identified on the Covered List or an affiliate or subsidiary of an entity identified on the Covered List, established pursuant to 47 CFR §1.50002.

I HEREBY CERTIFY THAT I am authorized to make the representations above on behalf of the company and agree to immediately notify Intertek if there is any change in the status identified above.

 12/19/2022

[*Madubuike OKAFOR*]

[*Director of QMS & Regulatory Affairs for US Medical Devices*]

[*BioSerenity USA, Inc.*]