



Medical Equipment and PPE Regulatory Reference Guide

The COVID-19 pandemic has had an unprecedented impact on the world. As healthcare providers work on the front line to combat the pandemic, the supply of ventilators, face masks and other personal protective equipment (PPE) are growing scarce. As manufacturers, many of you are in the process of supporting these ongoing needs. To assist you in the fight, we've put together this guide to share the proper protocols for U.S. regulatory submission, as provided by the FDA, CDC and ASTM, to help ensure your products get to market as quickly as possible, and into the hands of our healthcare providers.

Free Resources to support COVID-19	Relevant ASTM Testing Standards & Specifications	Recent Final Medical Device Guidance Documents	FDA Emergency Us Authorizations
Guidance for Medical	Enforcement Policy for Gowns,	FAQs of Shortages of Surgical	Questions about Personal
Gloves and Gowns	Gloves, etc.	Masks and Gowns:	Protective Equipment (PPE)
FDA Industry Guidance for Face Masks & Respirators	Enforcement Policy For Face	N95 Respirators and Surgical	MOU using 3D Printing
	Masks and Respirators	Masks (Face Masks)	between NIH, FDA, and VHA
FDA Industry Guidance for	Enforcement Policy for	Ventilator Supply Mitigation	FDA Ventilator and EUA
Ventilators	Ventilators	Strategies	Guidance
U.S Customs and Border Protection	CSMS #42124872 Filing PPE & Medical Devices	Reminder Format the correspondence in accordance with the relevant FDA guidance document to expedite response.	

For additional FDA contact information, please call: 1-888-INFO-FDA
For questions regarding the appropriate product coding, email the <u>FDA</u>.
If requesting emergency use authorizations for diagnostic devices, email the <u>FDA</u>.
If requesting emergency use authorizations for non-diagnostic devices, email the <u>FDA</u>.

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