

## FDA REQUIREMENTS FOR

## **High Demand COVID-19 Devices**

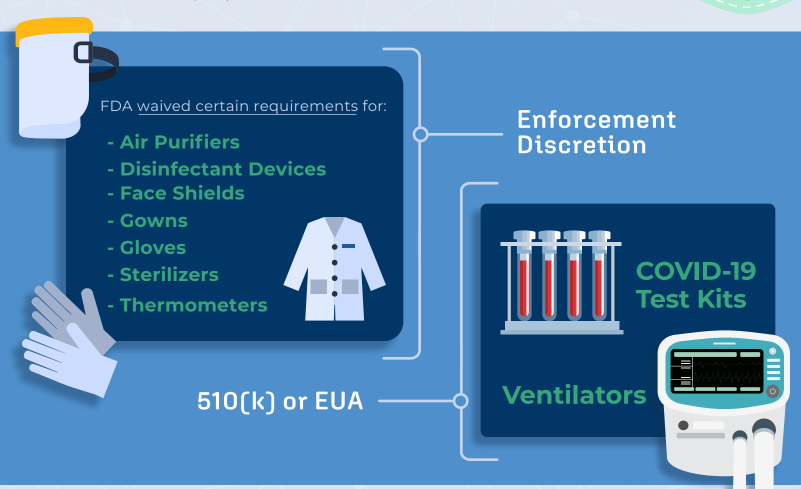
Registrar Corp can answer any questions you may have about product codes and FDA requirements specific to your medical devices.

## **Masks**

The following is not an all inclusive list of FDA product codes. Click the product codes below for the FDA definition of each device.



- \* FDA has temporarily waived registration and 510(k) requirements for face masks and CDC-recognized respirators
- \*\* N95 respirators must have NIOSH-approval, CDC-recognition, or Emergency Use Authorization (EUA)



Per FDA's Enforcement Policy, manufacturers may request EUA to market certain products without a 510(k).

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