You are being given this Fact Sheet because your healthcare provider believes it is necessary to treat you with a device, called an infusion pump, with the controlled infusion of medications, nutrition, called total parenteral nutrition or TPN, and/or other fluids. This Fact Sheet contains information to help you understand the benefits and risks of using infusion pumps and infusion pump accessories (such as the tubing and catheters that allow the pump to deliver medication) for the controlled infusion of medications, TPN, and/or other fluids. This Fact Sheet is specific to infusion pumps and infusion pump accessories that were authorized by FDA under an emergency use authorization (EUA) for these devices available at https://www.fda.gov/media/138057/download.

After reading this Fact Sheet, if you have questions or would like to discuss the information provided, please talk to your doctor, nurse, or other healthcare provider.

What is COVID-19?
COVID-19 is a disease caused by the SARS-CoV-2 virus. The virus, which can cause mild to severe respiratory illness, was first identified in Wuhan, China, and has now spread globally, including the United States. The current information available to characterize the spectrum of clinical illness associated with COVID-19 suggests that symptoms include cough, shortness of breath or difficulty breathing, fever, chills, muscle pain, headache, sore throat, or new loss of taste or smell.

What do I need to know about infusion pumps and infusion pump accessories?
A healthcare provider may choose to treat you with an infusion pump and infusion pump accessories if you require controlled infusion of medications, TPN, and/or other fluids. Controlled infusion means that the pump can be programmed to give you continuous (constant), intermittent (occasional), and bolus (a dose given all at once) infusions. Infusion pumps are an integral part of patient care during the COVID-19 pandemic because patients undergoing mechanical ventilation, among other medical interventions, require controlled infusion using at least one infusion pump.

Certain infusion pumps and infusion pump accessories have been authorized under an EUA for emergency use by healthcare providers to treat conditions caused by COVID-19 with the controlled infusion of medications, TPN, and/or other fluids due to shortages.

What are the known and potential benefits and risks of infusion pumps and infusion pump accessories?
Potential benefits of infusion pumps and infusion pump accessories include:

- Controlled flow of medications, TPN, and/or other fluids into a patient.

For infusion pumps with remote monitoring or remote manual control features or administration sets and other infusion pump accessories with increased length, maintaining a safe physical distance between the clinician and patient affected by COVID-19.

• Where can I go for updates and more information? The most up-to-date information on COVID-19 is available at the CDC General webpage: https://www.cdc.gov/COVID19. In addition, please also contact your healthcare provider with any questions/concerns.
• Have a problem with device performance? Report adverse events to MedWatch by submitting the online FDA Form 3500 (https://www.accessdata.fda.gov/scripts/medwatch/index.cfm?action=reporting.home) or by calling 1-800-FDA-1088.

FACT SHEET FOR PATIENTS
Emergency Use of Infusion Pumps and Infusion Pump Accessories During the COVID-19 Pandemic
May 13, 2020

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You should discuss any questions or concerns with your health care provider. You have the option at any time to refuse or stop treatment with this device. If you choose to decline or stop treatment with this device, you should discuss any alternative treatment options with your healthcare provider.

Potential risks of infusion pumps and infusion pump accessories include:

- Over or under delivery of therapy (especially medications).
- Other infusion delivery error, including free flow, and line occlusion.
- Air emboli (when air gets into the blood stream).
- Pump programming error from remote manual controller malfunction.
- Delayed infusion resulting from faster battery depletion due to remote manual control functionality.
- Malfunction of infusion pump alarms and/or patient monitoring features.
- User error when healthcare providers may not be familiar with new pumps or new pump features.

What is an EUA?
The United States FDA has authorized emergency use of infusion pumps and infusion pump accessories to treat conditions caused by COVID-19 with the controlled infusion of medications, TPN, and/or other fluids available under an emergency access mechanism called an EUA. The EUA is supported by the Secretary of Health and Human Services (HHS) declaration that circumstances exist to justify the emergency use of these devices for COVID-19, and the totality of available scientific evidence indicates that it is reasonable to believe that infusion pumps and infusion pump accessories that meet certain criteria may be effective to treat conditions caused by COVID-19 with the controlled infusion of medications, TPN, and/or other fluids.

The EUA for infusion pumps and infusion pump accessories for COVID-19 is current for the duration of the COVID-19 declaration justifying emergency use of these devices (unless terminated or revoked (after which the products may no longer be used).

These infusion pumps and infusion pump accessories have not undergone the same type of review as an FDA-approved or cleared device. FDA may issue an EUA when certain criteria are met, which includes that there are no adequate, approved, and available alternatives. In addition, the FDA decision is based on the totality of scientific evidence available showing that it is reasonable to believe that infusion pumps and infusion pump accessories meet certain criteria. It is effective to treat conditions caused by COVID-19 with the controlled infusion of medications, TPN, and/or other fluids.

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