



Medtronic

Manage your diabetes
with up to **96% fewer injections***1

MiniMed™ 630G
insulin pump system**



* Compared to multiple daily injections.

** Optional continuous glucose monitoring (CGM)/sensor use.



Components of insulin pump therapy**

1. MiniMed™ 630G insulin pump*

A small, durable and waterproof†† device that delivers insulin. Contains two compartments for the reservoir and a AA battery.

2. Extended infusion set and reservoir†

An infusion set delivers insulin from the reservoir in the pump to your body through a small, thin tube in your set.

3. Continuous glucose monitoring (CGM)

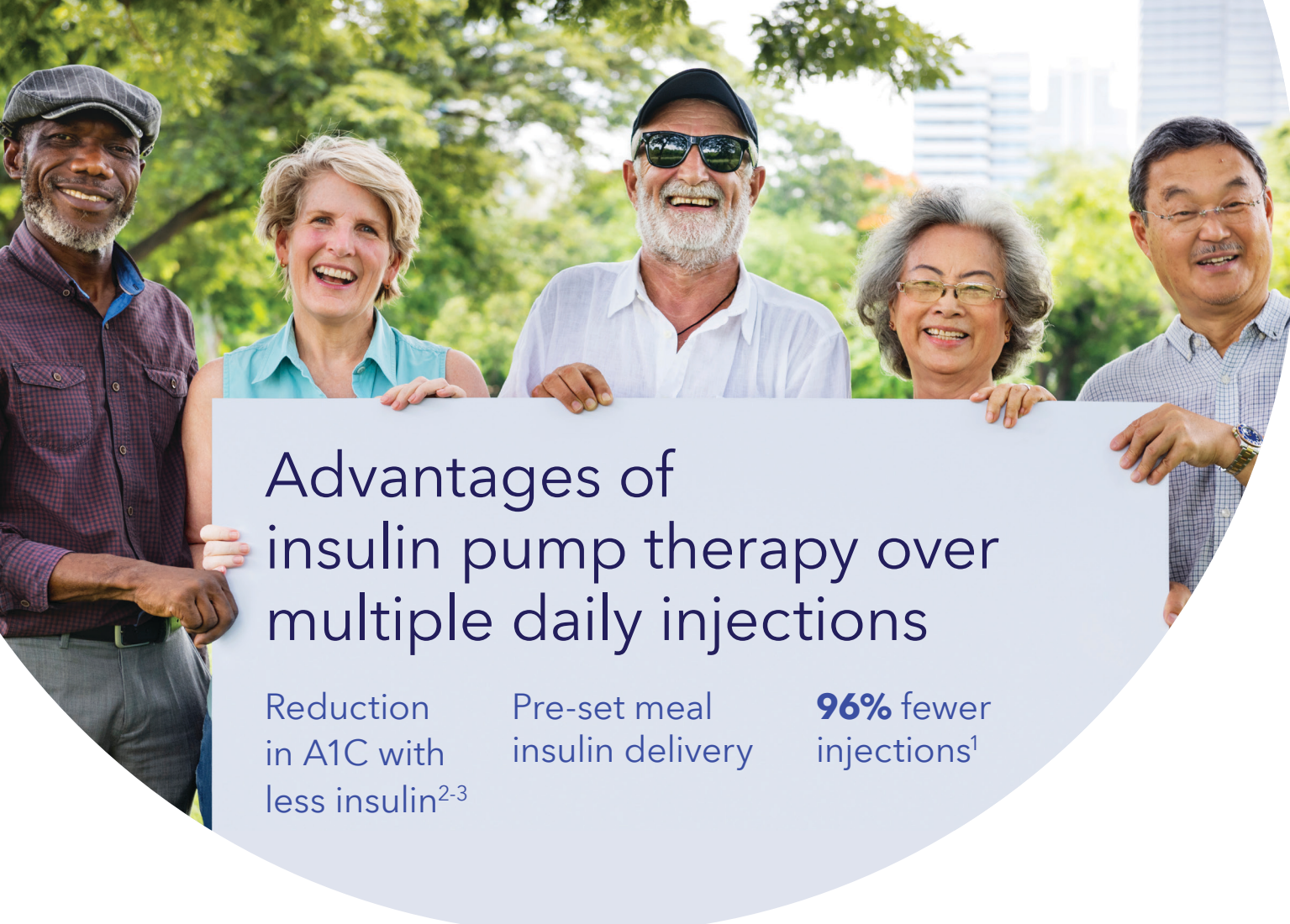
Optional CGM lets you see your glucose readings every five minutes in real-time.

* Optional CGM/sensor use.

** The products shown are for illustration purposes only. Not actual size.

† Extended infusion set and reservoir are sold separately

†† At the time of manufacture and when the reservoir and tubing are properly inserted, your pump is waterproof. It is protected against the effects of being underwater to a depth of up to 12 feet (3.6 meters) for up to 24 hours. This is classified as IPX8 rating. See user guide for more details. The sensor and transmitter are water-resistant at 8 feet (2.4 meters) for up to 30 minutes. CGM readings may not be transmitted from the CGM to the pump while in water.



Advantages of insulin pump therapy over multiple daily injections

Reduction in A1C with less insulin²⁻³

Pre-set meal insulin delivery

96% fewer injections¹

Long-term complication reduction



Kidney damage reduced up to 54%²



Nerve damage reduced up to 60%²



Cardiovascular damage reduced up to 41%²



Eye damage reduced up to 63%⁴

99%
of U.S. health plans have demonstrated access⁵



MiniMed™ insulin pump therapy Provides better management of type 2 diabetes



Programmed mealtime dosing

Preset meal bolus options are available for breakfast, lunch, snack, and dinner so you may not have to count carbs.



Less insulin

May reduce total daily insulin* by up to 19%³ and no longer requires long-acting insulin for a bonus cost savings.



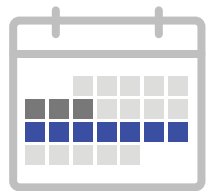
Predictive alerts

With CGM, the pump lets you know up to 30 minutes ahead of a high or low.

* Individual results may vary.

Medtronic Extended infusion set

The first and only infusion set with up to 7 days of wear



- Other infusion sets
- Medtronic Extended infusion set



x 4

site changes a month*

vs.



x 120

injections per month*



With a pump, you change your infusion set site rather than give yourself an injection



* A month is defined as 4 weeks

Diabetes takes no breaks. Neither do we.

24-hour technical support

Our support team is available around the clock to answer all your product, service, and software questions.

Training and education

Our experienced training and education team can help you get the most from your new therapy.

Global assistance

Our global service gives you peace-of-mind by ensuring you have the support you need when you need it, wherever you go.

Medtronic Assurance

Our commitment provides you with options during challenging times; from receiving supplies at a lower cost to flexible payment plans.

Call **888-847-6019** to learn more about the MiniMed™ 630G insulin pump.

1. Medtronic Data on file. Extended Wear reference (Half as often) REF 10395. Jan. 2021.
2. Reznik Y, Cohen O, Aronson R, et al.; OpT2mise Study Group, Insulin pump treatment compared with multiple daily injections for the treatment of type 2 diabetes (OpT2mise): a randomized open label controlled trial. *Lancet*. 2014; 384(9950): 1265-1272.
3. Aronson R, Reznik Y, Conget I, et al. Sustained efficacy of insulin pump therapy, compared with multiple daily injections, in type 2 diabetes: 12-month data from the OpT2mise randomized trial. *Diabetes Obesity and Meta*. 2016; 18:500-507.
4. The Diabetes Control and Complications Trial Research Group. The effect of intensive treatment of diabetes on the development and progression of longer-term complications in insulin-dependent diabetes mellitus. *N Engl J Med*. 1993;324:977-986.
5. Data on File.

Important Safety Information: MiniMed™ 630G system with SmartGuard™ technology

Indicated for the continuous delivery of insulin, at set and variable rates, for the management of diabetes mellitus. MiniMed™ 630G system is approved for ages 14 years or older with Guardian™ Sensor 3 and MiniMed™ 630G system is approved for ages 16 years or older with Enlite™ sensor. Both systems require a prescription. Insulin infusion pumps and associated components of insulin infusion systems are limited to sale by or on the order of a physician and should only be used under the direction of a healthcare professional familiar with the risks of insulin pump therapy. Pump therapy is not recommended for people who are unwilling or unable to perform a minimum of four blood glucose tests per day. Pump therapy is not recommended for people who are unwilling or unable to maintain contact with their healthcare professional. Pump therapy is not recommended for people whose vision or hearing does not allow recognition of pump signals and alarms. Insulin pumps use rapid-acting insulin. If your insulin delivery is interrupted for any reason, you must be prepared to replace the missed insulin immediately. Replace the infusion set every 48-72 hours, or more frequently per your healthcare professional's instructions. Insertion of a glucose sensor may cause bleeding or irritation at the insertion site. Consult a physician immediately if you experience significant pain or if you suspect that the site is infected. The information provided by CGM systems is intended to supplement, not replace, blood glucose information obtained using a blood glucose meter. A confirmatory fingerstick using a CONTOUR®NEXTLINK2.4 meter is required prior to making adjustments to diabetes therapy. Always check the pump display when using a CONTOUR®NEXTLINK2.4 meter, to ensure the glucose result shown agrees with the glucose results shown on the meter. Do not calibrate your CGM device or calculate a bolus using a result taken from an Alternative Site (palm) or a result from a control solution test. If a control solution test is out of range, please note that the result may be transmitted to your pump when in the "Always" send mode. It is not recommended to calibrate your CGM device when sensor or blood glucose values are changing rapidly, e.g., following a meal or physical exercise. The MiniMed™ 630G system is not intended to be used directly for preventing or treating hypoglycemia but to suspend insulin delivery when the user is unable to respond to the Suspend on low alarm and take measures to prevent or treat hypoglycemia themselves. Therapy to prevent or treat hypoglycemia should be administered according to the recommendations of the user's healthcare provider.

WARNING: The SmartGuard™ Suspend on low feature will cause the pump to temporarily suspend insulin delivery for two hours when the sensor glucose reaches a set threshold. Under some conditions of use the pump can suspend again, resulting in very limited insulin delivery. Prolonged suspension can increase the risk of serious hyperglycemia, ketosis, and ketoacidosis.

Before using the SmartGuard™ feature, it is important to read the SmartGuard™ feature information in the User Guide and discuss proper use of the feature with your healthcare provider. See www.medtronicdiabetes.com/importantssafetyinformation and the appropriate user guides for additional important details.

The Extended Infusion Set is indicated for up to 7 days of wear for the subcutaneous infusion of insulin from an infusion pump. It is NOT indicated for intravenous (IV) infusion or the infusion of blood or blood products. Inaccurate medication delivery, infection and/or site irritation may result from improper insertion and maintenance of the infusion site. Before insertion, clean the insertion site with isopropyl alcohol. Remove the needle guard before inserting the infusion set. If using this infusion set for the first time, do the first set-up in the presence of your healthcare professional. Do not leave air in the infusion set. Prime completely. Check frequently to make sure the soft cannula remains firmly in place as you may not feel pain if it pulls out. The soft cannula must always be completely inserted to receive the full amount of medication. If the infusion site becomes inflamed, replace the set, and use a new site until the first site has healed. Replace the infusion set if the tape becomes loose, or if the soft cannula becomes fully or partially dislodged from the skin. Regularly replace the infusion set as indicated in the instructions for use, or per the insulin labeling, whichever duration is shorter. See www.medtronicdiabetes.com/importantssafetyinformation and the appropriate user guides for additional important details.

Reservoir (Extended): The Extended Reservoir requires a prescription and is indicated for the subcutaneous infusion of insulin from compatible Medtronic insulin pumps and infusion sets. Refer to your Medtronic insulin pump user guide for a list of compatible insulins and infusion sets. This reservoir is contraindicated for the infusion of blood or blood products. Do not use with two or three day infusion sets. The reservoir may only be used for up to seven days with use of the Medtronic Extended infusion set. Use of the reservoir with two or three day infusion sets may lead to hyperglycemia or diabetic ketoacidosis. Do not use the reservoir for more than seven days. Using the reservoir for more than seven days may result in the delivery of too little insulin, which may cause hyperglycemia. Reservoir and transfer guard are sterile, nonpyrogenic, and for single use. For more details, see <https://bit.ly/importantssafety>.

Medtronic Diabetes Continuous Glucose Monitoring (CGM) Systems: The information provided by MiniMed CGM systems is intended to supplement, not replace, blood glucose information obtained using a home glucose meter. A confirmatory finger stick is required prior to treatment. Insertion of a glucose sensor may cause bleeding or irritation at the insertion site. Consult a physician immediately if you experience significant pain or if you suspect that the site is infected. Please visit www.medtronicdiabetes.com/about/safety.html for additional details.

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