# sanofi

### MAINE RESIDENTS ONLY APPLICATION



# MAINE INSULIN SAFETY NET PROGRAM FOR SANOFI INSULINS

The Act to Create the Insulin Safety Net Program ("the Act"), effective as of March 1, 2022, mandates that insulin manufacturers create an Insulin Safety Net Program ("the Program") to provide Maine (ME) residents who meet all eligibility criteria with their insulin prescription at no cost. Sanofi is administering the ME Insulin Safety Net Program through Sanofi Patient Connection<sup>®</sup>, a patient assistance program that helps patients get access to their medications.

### What are the eligibility criteria mandated by the Act?

In order to be eligible for this Program, ME requires you to meet the following requirements:

- You must be a resident of the State of Maine (ME) with a valid ME identification card that indicates ME residency in the form of an ME identification card, driver's license, or permit. If the individual is under the age of 18, the individual's parent or legal guardian shall provide proof of residency.
- You must not be enrolled in MaineCare.
- You must not be enrolled in prescription drug coverage through an individual or group health plan that limits the total amount of costsharing that an enrollee is required to pay for a 30-day supply of insulin, including copayments, deductibles, or coinsurance of \$75 or less, regardless of the type or amount of insulin needed.
- You must have an annual household income of ≤400% of the current Federal Poverty Level.
- You must not be eligible to receive healthcare coverage through a federally funded program or receive prescription drug benefits through the United States Department of Veterans Affairs.
- If you are enrolled in Medicare Part D, in addition to the criteria above, you must also spend at least \$1,000 on prescription drugs in the current calendar year.

## How do I apply?

To apply for Patient Assistance Connection, all information must be complete and include the following:

#### **Patient Information:**

• Complete all relevant information on page 2, and sign and date the REQUIRED patient authorizations for HIPAA consent and income verification on page 2.

#### Healthcare Provider:

- Ask your healthcare provider (HCP) to complete page 3 and sign and date it.
- Ask your HCP to mail, fax, or submit through the Provider Portal your completed application (only pages 2 and 3 are needed).

Provide proof of ME residency (copy of ME ID card, driver's license, or permit). If you are under 18, your parent or legal guardian needs to provide residency proof. Missing information may delay processing of your application. Your completed application should be submitted by your HCP via mail or fax, including your completed application and residency proof to:



US Mail Sanofi Patient Connection PO Box 222138 Charlotte, NC 28222-2138



#### What happens next?

Once we receive your application and proof of ME residency:

- 1. We will notify you within 5-7 business days if we require additional information to process your application.
- 2. Once we have all of the required information, we will review it to see if you meet all of ME's eligibility criteria to qualify for the Program within 3 business days.

If you are eligible:

- 1. You and your HCP will receive a letter within 10 business days notifying you of enrollment.
- 2. You will be enrolled for 12 months. If you are a Medicare Part D patient, you will be enrolled through the end of the calendar year. Your eligibility is renewable upon a redetermination of eligibility.
- 3. Your medication will be sent directly to your home in approximately 5-7 business days from when you are approved.

If you do not qualify for the Program, we will send you and your HCP a letter within 10 business days with the reason for denial.



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# **Section 1. Patient Information**

Patient first name					MI		Last name
SSN							DOB
Address							City
State					Zip		Preferred language (if not English)
Phone number (		)					
Email							
Household size	1	2	3	4	5	Other:	Annual household income

I permit Sanofi Patient Connection<sup>®</sup> to speak with the following person and/or organization about the information on this application and the status of my application request.

Patient representative/organization name Relationsh	ip to patient Phone
Patient Authorizations Sanofi Patient Connection does <u>not</u> charge any fees for this service; application pr a third party completing this application on your behalf are not required by nor remi	ocessing, medication, and shipping are all offered at no cost. Any fees charged to you by tted to Sanofi.
I have read and agree to the HIPAA Consent included in <b>Section 6</b> on page 4.	I have read and agree to the Patient Certifications regarding receiving communications from Sanofi Patient Connection included in <b>Section 8</b> on page 5.
PATIENT SIGN (REQUIRED)	PATIENT SIGN (OPTIONAL)
(1 of 3) Patient signature/Legal representative if patient is <18 years Date I have read and agree to the Income Verification included in Section 7 on page 4. PATIENT SIGN (REQUIRED)	(3 of 3) Patient signature/Legal representative if patient is <18 years Date You must check boxes in <b>Section 8</b> and return page 5 with signature to opt in.
(2 of 3) Patient signature/Legal representative if patient is <18 years Date	

Section 2. Insurance Information (Please provide a copy of the patient's insurance card[s], both front and back, if available)

Insurance? Yes No	If yes, is it Medicare Part D?			
Primary insurance name		Secondary insurance name		
Insurance phone ( )		Insurance phone ( )		
Policy ID #	Group #	Policy ID #	Group #	
Policyholder name (first/last)		Policyholder name (first/last)		
Relationship to patient	DOB	Relationship to patient	DOB	





D	DEC	CDI	RED	TO	FILL	
	<b>NLO</b>	UNI	DLN			

# Please fill out and return this form (prescriber and patient signature required for all applications)

# Section 3. Treatment and Prescribing Information (See Section 5 for supported products)

Patient name					DOB		
Concurrent med	dications						
Product #1			Vials	Product #2			/ials
ICD-10 Code			Pens N/A	ICD-10 Code			Pens I∕A
Frequency				Frequency			
Maximum dose	per day	Q	ty	Maximum dose per d	ау	Qty	
If required by a	oplicable state-specif	ic law, please attac	ch copie	s of prescriptions on offici	al state prescription	forms.	
Section / Pr	escriber Informa	ation					
Prescriber name				State where license	ed		
License #	NF	ว่า #		Tax ID #	DEA #	ŧ	
Site/facility nam	e			Office contact name	 Э		
Type Clinic	Physician office	Outpatient hosp	oital Ir	npatient hospital	Phone	e()	
Facility address	*				Fax (	)	
City				State	Zip Co	ode	
Collaborating P	hysician Name						
*Sanofi product mu	ist be shipped to the sign	ing prescriber's office o	or hospital	address authorized by the pres	criber and not to a third <sub>i</sub>	party.	
am authorized under patient's personal ide information provided i Program and to other I have not received, n office or hospital addi	state law to prescribe and di ntification, medical, and insu is for the sole use of the Pro wise administer the Sanofi F or will I receive, any benefit ress. My signature certifies ti	spense the requested me rance information to Sanc gram to verify my patient's Patient Connection Progra from Sanofi or their agent hat any prescription produ	dication. I of ofi US and/o s insurance m and relat s or repres- icts receive	my knowledge. I certify that the Sar certify that I have obtained from my or Sanofi Cares North America and a coverage, to assess, if applicable, ted services. I understand that I am entatives for prescribing a Sanofi pi d from this Program will be used for sought from any payer, patient, or o	patient all required written a their agents and representa patient's eligibility for partic under no obligation to pres oduct. The facility address r the above-named patient o	authorization for the re atives. I understand th cipation in the Patient scribe any Sanofi prod noted above in Sectio only and will not be re	elease of my nat any Assistance luct and that on 4 is my sold nor
(REQUIRED)	iber signature (REQI	JIRED – no stamps	5)	Printed na	me		Date
Section 5. Pro	oducts Available	e with Patient	Conne	ection			

- Admelog® (insulin lispro injection) 100 Units/mL
- Apidra® (insulin glulisine injection) 100 Units/mL
- Insulin Glargine U-300 Injection (1.5 mL or 3.0 mL pens)\*

- Lantus® (insulin glargine injection) 100 Units/mL
- Soliqua® 100/33 (insulin glargine & lixisenatide) injection 100 Units/mL and 33 mcg/mL
- Toujeo® (insulin glargine) Injection 300 Units/mL(1.5 mL or 3.0 mL pens)\*

Full US prescribing information for all Sanofi Patient Connection supported products can be accessed at <u>www.sanofipatientconnection.com/medications-available</u>. \*Regular SoloStar® is packaged as 3 pens per pack, 450 units/pen; dials up to 80 units per single injection. Max SoloStar® is packaged as 2 pens per pack, 900 units/pen; dials up to 160 units per single injection; Max pen dials in 2-unit increments.

## **Additional Information**

- · Sanofi Patient Connection ships most medications in a 90-day supply.
- A representative from Sanofi may contact you for follow-up on any adverse event you may report regarding a Sanofi product.

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# Section 6. Authorization to Use and Disclose Health Information (REQUIRED) *Patient: Please read the following carefully, then date and sign where indicated in Section 1 on page 2.*

HIPAA Consent: I authorize my healthcare providers and staff; my health insurer, health plan, or programs that provide me health benefits (together, "Health Insurers") to disclose to, Sanofi US, its affiliated companies (ie, Sanofi Pasteur U.S. and Genzyme, a Sanofi Company), Sanofi Cares North America, and authorized third party agents involved in administration of this Program, (collectively "Program Sponsor"), health information about me, including information related to my medical condition, treatment, health insurance coverage, claims, prescriptions, and referral to and enrollment in this Program for purposes of determining my participation in, and administering, the Program, which may include contacting me as well as my doctor/healthcare provider, office/hospital staff, insurer (public/private) or others. I understand a representative from Sanofi may contact me for follow-up on any adverse event I may report regarding a Sanofi product. I authorize and consent to release of identifiable information about me including medical, financial and insurance records and information as required for participation in the Program. I understand that identifiable information about me will be kept confidential and will not be further used or disclosed except to administer the Program, or as required by law. I understand that information I authorize to be disclosed may be re-disclosed and is no longer protected by Federal privacy regulations. I agree that this authorization is voluntary and that I may refuse to sign this authorization. Refusal to sign will not affect my ability to obtain treatment but I will not be able to participate in this Program. Unless revoked, this authorization shall remain in effect throughout my participation in the Program, including subsequent reapplication as required. I may withdraw this authorization at any time by written notification to my doctor/healthcare provider; however, withdrawal of authorization will terminate my participation in this Program and will not affect information already disclosed under this Authorization. I understand that it is my responsibility to follow up with my prescriber or the Program to make sure that my re-orders, as appropriate, are requested in a timely manner by my healthcare provider so I do not run out of medication. I understand that Sanofi US and Sanofi Cares North America reserve the right at any time and without notice to modify or change eligibility criteria or discontinue this Program. I understand that I may withdraw (take back) this Authorization at any time by calling 1.888.847.4877.

# Section 7. Income Verification (REQUIRED) *Patient:* Please read the following carefully, then date and sign where indicated in Section 1 on page 2.

**Income Verification:** I authorize Sanofi Patient Connection (SPC) under the Fair Credit Reporting Act to use my demographic information to access reports on my individual credit history from consumer reporting agencies. I understand that, upon request, SPC will tell me whether an individual consumer report was requested and the name and address of the agency that furnished it. I further understand and authorize SPC to use any consumer reports about me and information collected from me, along with other information they obtain from public and other sources, to estimate my income in conjunction with the Patient Assistance Program eligibility determination process, if applicable. I further understand that no free product may be submitted for reimbursement to any payer, including Medicare and Medicaid; and no free product may be sold, traded, or distributed for sale. If approved for the SPC Patient Assistance Program, I will not seek to have the value of any medication provided to me under this program counted toward my true-out-of-pocket (TrOOP) cost for prescription drugs for my Medicare Part D Plan. Continuation in the SPC Patient Assistance Program is conditioned upon timely verification of income. In addition, I agree to notify SPC if my insurance situation changes.



PATIENT CHECK (OPTIONAL)

> PATIENT CHECK

(OPTIONAL)

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# Section 8. Patient Certifications (OPTIONAL) *Patient: Please read the following carefully, then date and sign where indicated in Section 1 on page 2.*

By checking this box, I authorize the Program to collect and use my personal information to contact me by mail, telephone, or email, with information about the Program, disease state, and products, promotions, services, and research studies, and to ask my opinion about such information and topics, including market research and disease-related surveys. I further authorize the Program to de-identify my health information and use it in performing research, including linkage with other de-identified information the Program receives from other sources, education, business analytics, marketing studies, or for other commercial purposes. I agree that the Program may use my health information for these purposes.

□ By checking this box, I authorize the Program to share my personal information with its contracted vendors to contact me by mail, telephone, or email, with information about the Program, disease state, and products, promotions, services, and research studies, and to ask my opinion about such information and topics, including market research and disease-related surveys. I further authorize the Program to share identifiable health information with one another in order to de-identify it for these purposes and as needed to perform the Services or to send the communications listed above (the "Communications"). I agree that the Program may share my health information with my doctors, specialty pharmacies, and insurers for these purposes.

I understand that I may be contacted by the Program in the event that I report an adverse event associated with a Sanofi product. I understand that I do not have to opt in to receive the Communications, and that I can still receive patient assistance through the Program, as prescribed by my physician. I may opt out of receiving the Communications offered by the Program, at any time by notifying a Program representative by telephone at 1-800-633-1610 or by email via <a href="https://cscontactus.sanofi.us/customerInformation.aspx">https://cscontactus.sanofi.us/customerInformation.aspx</a>. US residents may opt out or "unsubscribe" from future communications from Sanofi via the following website. For further information regarding privacy rights and how Sanofi uses personal information, please reference our Privacy Policy at <a href="https://www.sanofi.com/en/sanofi-us-privacy-policies">https://www.sanofi.com/en/sanofi-us-privacy-policies</a>. I also understand that the Services may be revised, changed, or terminated at any time.