DUPIXENT

 Complete the entire form and submit pages 1-2 to DUPIXENT MyWay[®] via fax at 1-844-387-9370 or Document Drop at www.patientsupportnow.org (code: 8443879370)
 For assistance, call 1-844-DUPIXEN(T) (1-844-387-4936) Option 1, Monday–Friday, 8 AM–9 PM ET

Patient name (first, MI, last)			DOB	Gender 🗆 F		
Address		•				
State ZIP	ZIP Preferred language (if not English)					
Mobile phone ()	□ Voicemail	Alternate phone ()	Preferred # Voicema		
Best time to call □8–10 AM □10 AM−12 PM □12−2 PM □2−4 PM □4−6 F	рм □6-9рм					
Email				ent in Section 8 and expressly or on behalf of the Program.		
Patient Authorizations						
I have read and agree to the Patient Authorization to Use and Disclose Health Information inclue	I have read and agree to the Patient Certifications included in Section 8.					
Patient Sign		Patient Sign				
(1 of 2) Patient signature/Legal representative if patient is <18 years (Puerto Rico <21 years	s) Date		representative if patient	is <18 years (Puerto Rico <21 years) Date		
Printed name if signed by legal representative if patient is <18 years		Representative relationship to patient if patient is <18 years				
Continue 2. Household Income Duration differentiation in the DUDU						
Section 2. Household Income Required if enrolling in the DUPIX		•	household incor	mo ⁰		
How many people live in your household? Please refer to Section 8, Patient Certifications, for additional information about the Patient Assistance F	Program.	What is your total annual household income? (Includes salary/wages, Social Security income, unemployment insurance benefits, disability income, any other inco for the household.)				
Section 3. Insurance Information Patient has no insurance. (Please fill out Se	ection 2.) \Box Attached copies	s of front and back	of primary prescription and medical ca		
rimary Rx insurance name		Primary medical insurance name				
Rx insurance phone ()	<u></u>	Insurance phone ()			
Policy ID # Group #		Policy ID #		_ Group #		
Rx BIN # Rx PCN #		Policyholder name (first/	/last)			
		Relationship to patient _				
□ I have already sent this prescription to the specialty pharma	acy.	-				
By checking the box, I acknowledge DUPIXENT MyWay will not conduct a b My preferred specialty pharmacy is				ring coverage on my patient's behalf. Fax ()		
my preieneu specially phannacy is		////		T dX ()		
Section 4. Prescriber Information						
Prescriber name	Site/facility name					
Specialty						
ddress						
City State ZIP				_ Fax ()		
Prescriber NPI #		Tax ID #				
Section 5. Diagnosis (Choose ONE) Date of diagnosis	//					
Moderate-to-severe asthma characterized by an eosinophi or with oral corticosteroid dependent asthma □ Primary c				oolyposis □ Primary diagnosis □ J33.0 Polyp of nasal cavity		
□ J45.50 Severe persistent asthma, uncomplicated						
□ J45.40 Moderate persistent asthma, uncomplicated		Other ICD-10-CM cod	de			
□ Oral corticosteroid dependent □ Eosinophilic phenotype						
		ICD-10-CM=International Classifica	ation of Diseases, Tenth Re	evision, Clinical Modification.		
Prescriber to fill out required prescription information on p	bage 2			Ν.		
				DUPIXENT		
se see accompanying full Prescribing Information or visit DU	PIXENThcp.co	<u>m</u> .		(dupilumab)Injection		
2 Sanofi and Regeneron Pharmaceuticals, Inc. All Rights Reserved. 06/2022	DUP.22.03.0025		1	100mg · 200mg · 300mg		





> Complete the entire form and submit pages 1-2 to DUPIXENT MyWay® via fax at 1-844-387-9370 or Document Drop at www.patientsupportnow.org (code: 8443879370)

FOR ENT SPECIALISTS/PULMONOLOGISTS > For assistance, call 1-844-DUPIXEN(T) (1-844-387-4936) Option 1, Monday-Friday, 8 AM-9 PM ET DOB **Prescriber Phone #**

Prescriber Address

Patient Name

Prescriber Name

NPI#

Prescriber State License #

(Required for prescribers in Puerto Rico only)

Section 6a. DUPIXENT[®] (dupilumab) Prescription Information

Section 6b. DUPIXENT[®] (dupilumab) Quick Start Program Prescription Information (For COMMERCIALLY INSURED patients)

For COMMERCIALLY INSURED patients, Quick Start may be able to provide DUPIXENT at no cost if there is a coverage delay. Fill out sections 6a and 6b completely to determine patient eligibility.

Device type (Choose ONE): □ Pre-filled syringe (100/200/300 mg) OR □ Pre-filled pen (200/300 mg) (for use in adolescents ≥12 years) Known drug allergies Quantity sufficient up to 84–day supply Refills				Device type (Choose ONE): □ Pre-filled syringe (100/200/300 mg) OR □ Pre-filled pen (200/300 mg) (for use in adolescents ≥12 years) Known drug allergies Quantity sufficient up to 28-day supply Refills			
Moderate-to-se corticosteroid	vere asth dependen	ma characterized by an eosinophilic phenotype or with oral t asthma	Moderate-to corticostero	-severe asti id depende	hma characterized by an eosinophilic phenotype or with oral nt asthma		
Patients aged ≥12 years				Initial loading dose: 400 mg SIG: 2 (200 mg/1.14 mL) injections SQ on Day 1 Subsequent (maintenance) dose: 200 mg SIG: 1 (200 mg/1.14 mL) injection every 2 weeks, starting on Day 15 Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1			
		Patients age ≥12 years		Subsequent (maintenance) dos: 300 mg SIG. 2 (300 mg/2 mL) injections SQ on Day 1 Subsequent (maintenance) dos: 300 mg SIG. 1 (300 mg/2 mL) injection SQ 2 weeks, starting on Day 15			
212 years	Dosage for patients with oral corticosteroid-dependent asthma or with co-morbid moderate-to- severe atopic dermatitis or adults with co-morbid chronic rhinosinusitis with nasal polyposis ☐ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 ☐ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks, starting on Day 15			severe at	Dosage for patients with oral corticosteroid-dependent asthma or with co-morbid moc severe atopic dermatitis or adults with co-morbid chronic rhinosinusitis with nasal pol Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ ever 2 weeks, starting on Day 15		
Patients aged 6-11 years: Weight:	Weight 15 kg to <30 kg	□ Initial and subsequent doses: 100 mg SIG: 1 (100 mg/0.67 mL) injection SQ every 2 weeks CR □ Initial and subsequent doses: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 4 weeks	Patients age 6-11 years: Weight:	Weight 15 kg to <30 kg	□Initial and subsequent doses: 100 mg SIG: 1 (100 mg/0.67 mL) ir SQ every 2 weeks ■ Initial and subsequent doses: 300 mg SIG: 1 (300 mg/2 mL) inject every 4 weeks		
kg (1 kg=2.2 lb)	Weight ≥30 kg	□ Initial and subsequent doses: 200 mg SIG: 1 (200 mg/1.14 mL) injection SQ every 2 weeks	kg (1 kg=2.2 lb) Weight ≥30 kg	Initial and subsequent doses: 200 mg SIG: 1 (200 mg/1.14 mL) inject every 2 weeks		
Patients aged 6-11 years with asthma	Weight 15 kg to <30 kg	□ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 □ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 4 weeks, starting on Day 29	Patients aged 6-11 years with asthma	Weight 15 kg to <30 kg	☐ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Da ☐ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injections every 4 weeks, starting on Day 29		
and co-morbid moderate-to- severe atopic dermatitis:	Weight 30 kg to <60 kg	□ Initial loading dose: 400 mg SIG: 2 (200 mg/1.14 mL) injections SQ on Day 1 □ Subsequent (maintenance) dose: 200 mg SIG: 1 (200 mg/1.14 mL) injection SQ every 2 weeks, starting on Day 15	and co-morbic moderate-to- severe atopic dermatitis:	Weight 30 kg to <60 kg	□ Initial loading dose: 400 mg SIG: 2 (200 mg/1.14 mL) injections SQ or □ Subsequent (maintenance) dose: 200 mg SIG: 1 (200 mg/1.14 mL) in SQ every 2 weeks, starting on Day 15		
Weight: kg (1 kg=2.2 lb)	Weight ≥60 kg	□ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 □ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks, starting on Day 15	Weight: kg (1 kg=2.2 lb	Weight ≥60 kg	□ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Da □ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) inject every 2 weeks, starting on Day 15		
Chronic rhinosi	nusitis witl	nasal polyposis	Chronic rhine	osinusitis wi	ith nasal polyposis		
Patients aged ≥18 years	□ Initial and subsequent dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks			Initial and subsequent dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 v			

Quantity sufficient up to 28-day supply Refills					
Moderate-to-severe asthma characterized by an eosinophilic phenotype or with oral corticosteroid dependent asthma					
Patients aged ≥12 years	Initial loading dose: 400 mg SIG: 2 (200 mg/1.14 mL) injections SQ on Day 1 Subsequent (maintenance) dose: 200 mg SIG: 1 (200 mg/1.14 mL) injection SQ every 2 weeks, starting on Day 15				
	Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks, starting on Day 15				
	Dosage for patients with oral corticosteroid-dependent asthma or with co-morbid moderate-to- severe atopic dermatitis or adults with co-morbid chronic rhinosinusitis with nasal polyposis Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks, starting on Day 15				
Patients aged 6-11 years: Weight:	Weight 15 kg to <30 kg	□ Initial and subsequent doses: 100 mg SIG: 1 (100 mg/0.67 mL) injection SQ every 2 weeks OR □ Initial and subsequent doses: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 4 weeks			
kg (1 kg=2.2 lb)	Weight ≥30 kg	□ Initial and subsequent doses: 200 mg SIG: 1 (200 mg/1.14 mL) injection SQ every 2 weeks			
Patients aged 6-11 years with asthma	Weight 15 kg to <30 kg	□ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 □ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 4 weeks, starting on Day 29			
and co-morbid moderate-to- severe atopic dermatitis:	Weight 30 kg to <60 kg	□ Initial loading dose: 400 mg SIG: 2 (200 mg/1.14 mL) injections SQ on Day 1 □ Subsequent (maintenance) dose: 200 mg SIG: 1 (200 mg/1.14 mL) injection SQ every 2 weeks, starting on Day 15			
Weight: kg (1 kg=2.2 lb)	Weight ≥60 kg	☐ Initial loading dose: 600 mg SIG: 2 (300 mg/2 mL) injections SQ on Day 1 ☐ Subsequent (maintenance) dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks, starting on Day 15			
Chronic rhinosinusitis with nasal polyposis					
Patients aged ≥18 years	☐ Initial and subsequent dose: 300 mg SIG: 1 (300 mg/2 mL) injection SQ every 2 weeks				

Sign Prescriber signature (No stamps) Dispense as written

Prescriber signature (No stamps) Substitution permitted

2

Collaborating MD name

(Nurse practitioner/physician assistant) NPI

Prescriber Certification: My signature certifies that the person named on this form is my patient; the information provided on this application, to the best of my knowledge, is complete and accurate; that therapy with DUPIXENT is medically necessary; and that I have prescribed DUPIXENT to the patient not bits from from the patient is form as my patient; the information provided to Regeneron Pharmaceuticals, Inc., Sanofi US, and their affiliates and agents (the "Alliance") is for the use of *DUPIXENT* is medically necessary; and that I have prescribed DUPIXENT to the facilitate the filling of my patient's prescription; to assess, if applicable, my patient's information provided to Regeneron Pharmaceuticals, Inc., Sanofi US, and their affiliates and agents (the "Alliance") is for the use of *DUPIXENT MyWay* solely to verify my patient's information in accordance with applicable is atterned and the approved indication. I understand that my patient's information provided to Regeneron Pharmaceuticals, Inc., Sanofi US, and their affiliates health information provided to the patient as sole and other support programs; and to otherwise administer *DUPIXENT MyWay* for the patient. I certify that I have obtained my patient's agreement that they would like to receive the Services and Other support programs; and to otherwise administer *DUPIXENT MyWay* to conduct a benefits investigation for my patient and to accurate its integration in accordance with applicable. Lauthorize *DUPIXENT MyWay* to conduct a benefits investigation for my patient and to accurate its integrate the individually identifiable health insurance otherwise indicated. I understand that any free product distributed through the *DUPIXENT MyWay* to conduct a benefits investigation for my patient and to accurate its insurance otherwise indicated. I understand that any free product distributed for sale. I consent to *DUPIXENT MyWay* to conduct a benefits information provide addinal information about DUPIXENT to the patient as adecarate ithat the product dist

If you are a New York prescriber, please use an original New York State prescription form. The prescriber is to comply with his/her state-specific prescription requirements, such as e-prescribing, state-specific prescription form, fax language, etc. Non-compliance with state-specific requirements could result in outreach to the prescriber.



Date

Please see accompanying full Prescribing Information or visit DUPIXENThcp.com.

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06/2022 DUP.22.03.0025

Date

Prescription Information

Section 7. Authorization to Use and Disclose Health Information

DUPIXENT

Enrollment Form

FOR ENT SPECIALISTS/PULMONOLOGISTS

Patient: Please read the following carefully, then date and sign where indicated in Section 1 on page 1

I authorize my healthcare providers and staff (together, "Healthcare Providers"), my health insurer, health plan or programs that provide me healthcare benefits (together, "Health Insurers"), and any specialty pharmacies ("Specialty Pharmacies") that dispense my medication to disclose to Regeneron Pharmaceuticals, Inc., Sanofi US, and their affiliates and agents (together, the "Alliance") health information about me, including information related to my medical condition and treatment, health insurance coverage and claims, and prescription (including fill/refill information) related to my prescription for DUPIXENT[®] (dupilumab) therapy ("My Information"). I understand the disclosure to the Alliance will be for the purposes of enrolling me in, and providing certain services through the "*DUPIXENT MyWay* Program," including:

- to determine if I am eligible to participate in DUPIXENT MyWay coverage assistance programs, patient assistance programs, or other support programs
- to investigate my health insurance coverage for DUPIXENT injection
- to obtain prior authorization for coverage
- to assist with appeals of denied claims for coverage
- for the operation and administration of the DUPIXENT MyWay Program
- to refer me to, or to determine my eligibility for, other programs, or alternative sources of funding or coverage that may be available to provide assistance to me with the costs of my medication
 - I understand that the Alliance may de-identify My Information and use it in performing research, education, business analytics, marketing studies, or for other commercial purposes, including linkage with other de-identified information the Alliance receives from other sources. I understand that members of the Alliance may share My Information, including identifiable health information, among themselves in order to de-identify it for these purposes and as needed to perform the Services or to communicate with me by mail, telephone, or email, or, if I indicate my agreement and consent in Section 1 on page 1, by text. I understand and agree that the Alliance may use My Information for these purposes and may share My Information with my Healthcare Providers, Health Insurers and Specialty Pharmacies.
 - I understand and agree that my Healthcare Providers, Health Insurers, and Specialty Pharmacies may receive remuneration from the Alliance in exchange for disclosing My Information to the Alliance and/or for providing me with support services in connection with the DUPIXENT MyWay Program.

Once My Information has been disclosed to the Alliance, I understand that federal privacy laws may no longer protect it from further disclosure. However, I also understand the Alliance has agreed to protect My Information by using and disclosing it only for the purposes allowed by me in this Authorization or as otherwise required by law.

I understand that I do not have to sign this Authorization. A decision by me not to sign this Authorization will not affect my ability to obtain medical treatment, payment for treatment, insurance coverage, access to health benefits or Alliance medications from covered entities such as Health Care Providers, Health Insurers, and Specialty Pharmacies. However, if I do not sign this Authorization, I understand that I will not be able to participate in the *DUPIXENT MyWay* Program.

I understand that this Authorization expires 18 months from the date support is last provided under the Program, or until my local state law requires expiration, subject to applicable law, unless and until I withdraw (take back) this Authorization before then, or as otherwise required by law. Further, I understand that I may withdraw this Authorization at any time by mailing or faxing a written request to *DUPIXENT MyWay* at 1800 Innovation Point, Fort Mill, SC 29715; Fax: 1-844-387-9370. Withdrawal of this Authorization will end my participation in the *DUPIXENT MyWay* Program and will not affect any disclosure of My Information based on this Authorization made before my request is received and processed by my Healthcare Providers, Health Insurers, and Specialty Pharmacies.

I understand that I may request a copy of this Authorization.



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Section 8. Patient Certifications

Patient: Please read the following carefully, then date and sign where indicated in Section 1 on page 1

I am enrolling in the *DUPIXENT MyWay* Program (the "Program") and authorize Regeneron Pharmaceuticals, Inc., Sanofi US, and their affiliates and agents (together the "Alliance") to provide me services under the Program, as described in the Program Enrollment Form and as may be added in the future. Such services include medication and adherence communications and support, medication dispensing support, coverage and financial assistance support, disease and medication education, injection training, and other support services (the "Services").

If enrolling in the DUPIXENT MyWay Copay Card Program, I understand that Copay Card information will be sent to my designated specialty pharmacy along with my prescription, and any assistance with my applicable cost-sharing or copayment for DUPIXENT[®] (dupilumab) injection will be made in accordance with the Program terms and conditions.

If I am completing Section 2, I certify that the information I have set forth in Section 2, including my household income, is true and accurate to the best of my knowledge. I also agree that the "Alliance" may verify my eligibility for the *DUPIXENT MyWay* Patient Assistance Program, and I understand that such verification may include contacting me or my healthcare provider for additional information and/or reviewing additional financial, insurance, and/or medical information. I authorize the Alliance 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, the Alliance 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 the Alliance 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 *DUPIXENT MyWay* 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 *DUPIXENT MyWay* Patient Assistance Program is conditioned upon timely verification of income. In addition, I agree to notify *DUPIXENT MyWay* if my insurance situation changes.

I authorize the Alliance to contact me by mail, telephone, or email, or, if I indicate my agreement and consent on page 1, by text,^a 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 (together, the "Communications"). I understand that I may be contacted by the Alliance in the event that I report an adverse event. I understand that I do not have to enroll in the Program or receive the Communications, and that I can still receive DUPIXENT injection, as prescribed by my Healthcare Provider. I may opt out of receiving Communications, individual support services offered by the Program, including the *DUPIXENT MyWay* Copay Card, or opt out of the Program entirely at any time by notifying a Program representative by telephone at 1-844-387-4936 or by sending a letter to *DUPIXENT MyWay*, 1800 Innovation Point, Fort Mill, SC 29715. I also understand that the Services may be revised, changed, or terminated at any time.

I understand that my health information, contact information, and other information I, my healthcare provider, and others share with Regeneron Pharmaceuticals, Inc., Sanofi US, and their affiliates and agents (together the "Alliance") is collected to provide me with the assistance I request and for other business purposes of the Alliance, as described in their privacy policy, which is available at regeneron.com/privacy-policy. Depending on where I live, I may have certain rights with respect to my privacy information, including the request to access or delete my personal information. I am aware that Regeneron may not be required to fulfill my requests in certain circumstances. I understand that to exercise these rights, I may contact the Privacy Office by emailing dataprotection@regeneron.com or by calling 844-835-4137. I may reference Sanofi's Global Privacy Policy at sanofi.com/our-responsibility/sanofi-global-privacy-policy for further information regarding these rights with respect to Sanofi US.

Text Messaging Consent:

^aI acknowledge that by checking the Text Messaging Consent box on page 1, I expressly consent to receive text messages from or on behalf of the Program at the mobile telephone number(s) that I provide.

I confirm that I am the subscriber for the mobile telephone number(s) provided, and I agree to notify the Alliance promptly if any of my number(s) change in the future. I understand that my wireless service provider's message and data rates may apply. I understand that I can opt out of future text messages at any time by texting SMSSTOP to 39771 and 69929 from my mobile phone, and that I can get help for text messages by texting SMSHELP to 39771 and 69929. I also understand that additional text messaging terms and conditions may be provided to me in the future as part of an opt-in confirmation text message. Message and data rates may apply.

I understand that my consent is not required as a condition of purchasing any goods or services from Regeneron Pharmaceuticals, Inc., Sanofi US, or their affiliates.

You may keep a copy of this form for your records.



Please see accompanying full Prescribing Information or visit <u>DUPIXENThcp.com</u>.



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