

Initiating benefits investigation is easy



For prescribers

- Complete the required Prescriber Information and Clinical Information sections on pages 1-3
- Complete the required Treatment Location Information section on page 2 (DARZALEX®, PROCRIT®, and YONDELIS® only)
- Complete the optional Prescription Information and the Preferred Pharmacy Information sections on page 3 (ERLEADA® and ZYTIGA® only)
- If prior authorization assistance is NOT needed, check the appropriate box in the Prior Authorization section on page 1 to opt out



Fax the completed and signed Benefits Investigation Form to Janssen CarePath at 855-998-4422



For your patients/caregivers

- Complete or have your patient complete the Patient Information and Insurance Information sections on page 4
- As requested by your patient, complete or have your patient complete the Janssen CarePath Savings Program section on page 5 to determine eligibility
- If you do not have a signed Business Associate Agreement (BAA) on file with Janssen CarePath, have your patient read, sign, and date the Patient Authorization on page 6
 - Give your patient a copy of the signed Patient Authorization form and keep the original for your records

Here's what happens next



For prescribers

Janssen CarePath will:

- Medical Benefit: Confirm receipt of requests within 2 hours and verify benefits within 1 to 2 business days
- Pharmacy Benefit: Verify benefits within 4-6 business hours
- Provide you with a verification of benefits and call your patient to review the benefits



For your patients/caregivers

Janssen CarePath will:

- Call your patient to review the benefits and provide you with a verification of benefits
- Inform your patient about cost support options and offer your patient care coordination support services with the infusion provider or specialty pharmacy
- Enroll your eligible patient with commercial or private health insurance in the Janssen CarePath Savings Program, if requested by your patient



PROCRIT®
EPOETIN ALFA



Erleada®
(apalutamide) 60 mg tablets

Zytiga®
abiraterone acetate

Medical Benefit

Pharmacy Benefit



Need help?

Call **877-CarePath** (877-227-3728)
Monday–Friday, 8:00 AM–8:00 PM ET

Please see full Prescribing Information for [DARZALEX®](#), [YONDELIS®](#), [ERLEADA®](#), and [ZYTIGA®](#).

Please see full Prescribing Information, including Boxed Warnings and Medication Guide for [PROCRIT®](#). Provide the Medication Guide to your patients and encourage discussion.

YONDELIS® (trabectedin) is under license from Pharma Mar, S.A.

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1. Prescriber Information—to be completed by Physician (Required)

Prescriber Name (First, Last) _____ Specialty _____

Practice Name _____ Office Contact _____

Address _____

City _____ State _____ ZIP _____

Email _____ Phone _____ Fax _____

Medicaid/Medicare Provider # _____ Tax ID # _____

State License # _____ UPIN/NPI # _____

2. Prior Authorization—to be completed by Physician (Optional)

Automatically provided with benefits investigation. You may opt out by checking the box below.

Prior Authorization Form Assistance and Status Monitoring: Janssen CarePath assists your office in providing the requirements of the patient’s health plan related to prior authorization for treatment with the medication specified on this form. Assistance includes obtaining the health-plan-specific prior authorization form and providing it based upon the patient-specific information provided on this form. The partially completed prior authorization form, if received from the health plan, will be provided to your office for possible completion and submission in the office’s sole discretion. Janssen CarePath also actively monitors the status of prior authorization submission to the patient’s plan and provides status updates to your office with respect to this patient’s prior authorization for treatment with the medication specified on this form.

I do **NOT** wish to receive Prior Authorization Form Assistance or Status Monitoring

By providing your information and information about your patient on the Benefits Investigation Form, you are requesting the services described on this form. The information you provide will only be used by Johnson & Johnson Health Care Systems Inc., our affiliates, and our service providers involved in delivering these services. You may withdraw your request for these services by calling 877-CarePath (877-227-3728). Our [Privacy Policy](#) governs the use of the information you provide. By providing the information and submitting this form, you indicate you read, understand, and agree to these terms.

Patient insurance benefits investigation and other Janssen CarePath program offerings are provided by third-party service providers for Janssen CarePath, under contract with Johnson & Johnson Health Care Systems Inc. on behalf of Janssen Pharmaceuticals, Inc., Janssen Biotech, Inc., and Janssen Products, LP (Janssen). Janssen CarePath is not available to patients participating in the Patient Assistance Program offered by Johnson & Johnson Patient Assistance Foundation. The availability of information and assistance may vary based on the Janssen medication, geography and other program differences. Janssen CarePath assists healthcare providers (HCPs) in the determination of whether treatment could be covered by the applicable third-party payer based on coverage guidelines provided by the payer, and patient information provided by the HCP under appropriate authorization following the provider’s exclusive determination of medical necessity. This information and assistance are made available as a convenience to patients, and there is no requirement that patients or HCPs use any Janssen product in exchange for this information or assistance. Janssen assumes no responsibility for and does not guarantee the quality, scope, or availability of the information and assistance provided. The third-party service providers, not Janssen, are responsible for the information and assistance provided under this program. Each HCP and patient is responsible for verifying or confirming any information provided. All claims and other submissions to payers should be in compliance with all applicable requirements.



3. Clinical Information for Benefits Investigation: IV Only—to be completed by Physician (Required)

Medication

DARZALEX® (daratumumab) PROCRT® (epoetin alfa) YONDELIS® (trabectedin)

Treatment Information

Primary Diagnosis Code _____ Primary Diagnosis Indication _____

Approximate Date of Patient's Diagnosis (mm/dd/yyyy) _____

Secondary Diagnosis (Optional) _____

Dosage Form and Strength _____ No. of Vials _____

Administration _____

Patient Weight _____ lbs _____ kg

Has the patient started therapy with the medication specified above? Yes No

If yes, what date did the patient start therapy? (mm/dd/yyyy) _____

Additional information regarding treatment (if applicable to benefits verification)

DARZALEX® only:

Monotherapy
 Combination Therapy

If Combination, list medications:

Prior Medications/Treatments:

PROCRT® only:

Initial HCT _____ %

Initial Hb _____ g/dL

For cancer patients, is the patient on chemotherapy? Yes No

Is the patient a nephrology patient? Yes No

If nephrology patient, what is the patient's:

Serum creatinine _____ mg/dL

Creatinine clearance _____ mL/min

Is the patient taking PROCRT® preoperatively? Yes No

If yes, surgery type _____

YONDELIS® only:

Patient Height _____ ft _____ in

Patient BSA _____

Has the patient taken a prior chemotherapy? Yes No

If yes, what prior chemotherapy has the patient taken?

Anthracycline
 Ifosfamide
 Other _____

Please investigate benefits for YONDELIS® infused through an ambulatory pump through a central venous catheter

4. Treatment Location Information: IV Only—to be completed by Physician (Required)

Treatment Location Type

Prescribing MD's Office Non-prescribing MD's Office Home Infusion/Infusion Provider Company
 Hospital Outpatient Hospital Inpatient Other _____

Provider Information

If prescribing MD's office, the fields below do not need to be completed if information is the same as the Prescriber Information section.

Provider Name (First, Last) _____ Physician Specialty _____

Practice Name _____

Address _____

City _____ State _____ ZIP _____

Site Phone _____ Site Fax _____

Insurance Provider # _____ Tax ID # _____

Please see full Prescribing Information for [DARZALEX®](#) and [YONDELIS®](#).

Please see full Prescribing Information, including Boxed Warnings and Medication Guide for [PROCRT®](#). Provide the Medication Guide to your patients and encourage discussion.



5. Clinical Information for Benefits Investigation: Orals Only—to be completed by Physician (Required)

Medication

ERLEADA® (apalutamide) 60 mg Tablet Dosing: 240 mg PO once daily with or without food Quantity: _____

ZYTIGA® (abiraterone acetate) 250 mg Tablet Dosing: _____ mg PO _____ daily on an empty stomach Quantity: _____

500 mg Film-Coated Tablet Dosing: _____ mg PO _____ daily on an empty stomach Quantity: _____

Treatment Information

Primary Diagnosis Code: C61 Primary Diagnosis Indication: Malignant neoplasm of prostate

Approximate Date of Patient's Diagnosis (mm/dd/yyyy) _____

6. Prescription Information: to be completed by Physician (Optional)

If requesting benefits investigation only, do not complete this section. The prescription is only valid if received by fax. If not faxed, prescription must be submitted on state-specific blank, if applicable for your state.

Patient Name (First, MI, Last) _____ Date of Birth _____

Rx ERLEADA® 60 mg Tablet

Directions: Take 240 mg PO once daily with or without food Quantity _____ Refills # _____

Rx ZYTIGA® 250 mg Tablet 500 mg Film-Coated Tablet

Directions: Take _____ mg PO _____ daily on an empty stomach Quantity _____ Refills # _____

Initial Dosing: For patients with baseline moderate hepatic impairment (Child-Pugh Class B), reduce the ZYTIGA® starting dose to 250 mg once daily (see Dose Medication Guidelines for more information). Do not use ZYTIGA® in women who are or may become pregnant and patients with baseline severe hepatic impairment (Child-Pugh Class C). Refer to the ZYTIGA® full PRESCRIBING INFORMATION, including the following sections: INDICATIONS AND USAGE, CONTRAINDICATIONS, DOSAGE AND ADMINISTRATION, WARNINGS AND PRECAUTIONS, ADVERSE REACTIONS, DRUG INTERACTIONS, and USE IN SPECIFIC POPULATIONS prior to initiating treatment.

Rx Prednisone 5 mg tablet

Directions: Take _____ Quantity _____ Refills # _____

Prednisone is required to be taken with ZYTIGA®; however, it is optional to include on this Benefits Investigation Form. You may provide a prescription direct to the patient to be filled at a pharmacy that can fill the script. NOTE: Janssen CarePath will not investigate benefits for prednisone. Please refer to full Prescribing Information for complete information prior to initiating treatment.

Prescriber Name (if different from page 1) _____

Address _____

City _____ State _____ ZIP _____

Phone _____ Fax _____

PRESCRIBER SIGNATURE (NO STAMPS) REQUIRED. I certify that therapy with the Janssen medication indicated above is medically necessary for this patient. I will be supervising the patient's treatment accordingly, and I have reviewed the current Prescribing Information for the Janssen medication indicated above. I authorize Janssen CarePath to act on my behalf for the limited purposes of transmitting this prescription to the appropriate pharmacy designated by me, the patient, or the patient's plan.

Prescriber Signature >> (Dispense as written) _____ Date _____

Prescriber Signature >> (Substitutions allowed) _____ Date _____

Supervising Physician Signature >> (If applicable) _____ Date _____

Supervising Physician Name _____

7. Preferred Pharmacy: Orals Only—to be completed by Physician (Optional)

As the treating physician, I have discussed preference for a Specialty Pharmacy (SP) with this patient. This patient prefers use of the SP indicated below. I authorize Janssen Biotech, Inc., and its representatives to fax this prescription to: **1.** The SP designated below, provided it is approved by this patient's plan. **2.** If the SP designated is not a plan-approved SP, then to an SP approved by this patient's plan. **3.** If there is no preferred SP indicated, then to any SP approved by this patient's plan.

Preferred Specialty Pharmacy _____ Self-Dispensing Pharmacy

Please see full Prescribing Information for **ERLEADA®** and **ZYTIGA®**.



8. Patient Information (Required)

Name (First, MI, Last) _____ Language English Spanish

Male Female Date of Birth (mm/dd/yyyy) _____

Address _____

City _____ State _____ ZIP _____

Primary Email _____ Secondary Email (Optional) _____

Primary Phone _____ Secondary Phone (Optional) _____ Best Time to Contact _____

Caregiver/Contact _____
(A caregiver/contact is someone who can be contacted in place of the patient)

Home/Cell Phone _____ Work Phone _____ Best Time to Contact _____

I authorize Janssen CarePath to leave a message, including the name of the Janssen medication indicated on this form, if I am unavailable when they call.

If I cannot be reached, I authorize Janssen CarePath to contact my caregiver.

I prefer and authorize Janssen CarePath to contact my caregiver in place of me.

9. Insurance Information (Required)

Please provide insurance information for all health insurance coverage you may have.

Please see attached front and back copy of insurance card.

Primary Medical Insurance: required for DARZALEX® (daratumumab), YONDELIS® (trabectedin), PROCRIT® (epoetin alfa)

Primary Insurance Carrier _____ Phone _____

Cardholder Name (First, MI, Last) _____ Relationship to Cardholder _____

Cardholder Employer _____ Policy # _____ Group # _____

Secondary Medical Insurance (Optional)

Secondary Insurance Carrier _____ Phone _____

Cardholder Name (First, MI, Last) _____ Relationship to Cardholder _____

Cardholder Employer _____ Policy # _____ Group # _____

Prescription Drug Insurance: required for ERLEADA® (apalutamide), ZYTIGA® (abiraterone acetate)

Prescription Drug Insurer _____ Card BIN # _____ Phone _____

Cardholder Name (First, MI, Last) _____ Relationship to Cardholder _____

Cardholder Employer _____ Policy # _____ Group # _____

Please investigate out-of-network benefits.

Please see full Prescribing Information for [DARZALEX®](#), [YONDELIS®](#), [ERLEADA®](#), and [ZYTIGA®](#).

Please see full Prescribing Information, including Boxed Warnings and Medication Guide for [PROCRIT®](#). Provide the Medication Guide to your patients and encourage discussion.



10. Janssen CarePath Savings Program (Optional)

Eligible patients using commercial insurance can save on out-of-pocket Janssen medication costs. See program requirements at [JanssenCarePath.com](https://www.JanssenCarePath.com).

I would like Janssen CarePath to check my eligibility for and enroll me into the Janssen CarePath Savings Program if the results of this benefits investigation determine I have commercial or private health insurance.

Rebate Type for DARZALEX® (daratumumab) or YONDELIS® (trabectedin)

Please select how you would like to receive your rebate, if the consent above is checked and benefits investigation is for DARZALEX® or YONDELIS®

Load Funds onto Card Mail Check to Patient Mail Check to Provider* (please select one option below)
 Prescriber Office Treatment Location

*By selecting this option, I understand that I am requesting that Janssen CarePath Savings Program rebate check(s) will be sent on my behalf to the designated provider for payment of my out-of-pocket Janssen medication costs. I also understand that I may, at any time, call Janssen CarePath and elect for the rebate check(s) to be sent directly to me.

Eligibility Questions

1. Do you currently have commercial or private health insurance that you will use for your Janssen medication, including commercial insurance provided through an employer or former employer, provided to you as a federal or state employee, and insurance you pay for yourself, as well as plans available through state and federal healthcare exchanges?

- Yes**, I have commercial or private health insurance that I will use for my Janssen medication
 No, I do not have commercial or private health insurance that I will use for my Janssen medication

2. Do you confirm that you will NOT seek reimbursement from any state or federal government-subsidized healthcare program to cover a portion of the Janssen medication costs such as Medicare Parts A, B, C (also known as Medicare Advantage Plan), D, and Medicare Supplement, Medicaid, TRICARE, Department of Defense, or Veterans Administration?

- Yes**, I confirm that I will NOT seek reimbursement from any state or federal government-subsidized program for my Janssen medication
 No, I may seek reimbursement from a state or federal government-subsidized healthcare program for my Janssen medication

3. Do you confirm that you will not submit out-of-pocket costs paid by this program as a claim for payment to any third-party payer, pharmaceutical patient assistance foundation, or account such as a Flexible Spending Account (FSA), a Health Savings Account (HSA), or a Health Reimbursement Account (HRA)?

- Yes**, I confirm that I will NOT submit out-of-pocket costs paid by this program as a claim for payment to any third-party payer, pharmaceutical patient assistance foundation, or account
 No, I may submit out-of-pocket costs paid by this program as a claim for payment to a third-party payer, pharmaceutical patient assistance foundation, or account

Please see full Prescribing Information for [DARZALEX®](#) and [YONDELIS®](#).



- Patients should read the Patient Authorization and sign electronically or download, print, and sign.
 - Completed form may be uploaded to Patient Account or Provider Portal, faxed to Janssen CarePath at 855-998-4422, or mailed to address below.
- Patients can access a copy of completed form in their Janssen CarePath Account – My Profile.

My signature on this Patient Authorization Form confirms that I authorize each of my physicians, pharmacists, including any specialty pharmacy that receives my prescription for a Janssen medication and other healthcare providers (together, "Healthcare Providers") and each of my health insurers (together, "Insurers") to disclose my protected health information, including but not limited to information related to my medical condition and treatment, my health insurance coverage, my name, address, telephone number, insurance plan and/or group numbers (together, "Protected Health Information") to Johnson & Johnson Health Care Systems Inc., its affiliated companies, agents and representatives (together, "Janssen"), including providers of alternate sources of funding for prescription drug costs, and other approved service providers authorized to manage, administer, and/or support Janssen CarePath programs, Janssen CarePath Account for Patients, and Provider Portal for their Healthcare Providers for the purposes described below.

Specifically, I authorize Janssen to receive, use, and disclose my Protected Health Information in order to (i) enroll me in, determine my eligibility for, and contact me about, Janssen medication support programs; (ii) provide me with educational materials, information, and services related to my Janssen medication; (iii) verify, investigate, assist with, and coordinate my coverage for my Janssen medication with my Insurers; (iv) coordinate prescription fulfillment; (v) assist with analyses related to the quality, efficacy, and safety of my Janssen medication, and patient access to and adherence to my Janssen medication; (vi) to share and provide access to, information generated by Janssen CarePath that may be useful for my care, and; (vii) to improve, develop, and evaluate Janssen CarePath, its offerings, and materials. I also understand that pharmacies that ship my medication may be paid to share this information with Janssen CarePath to help provide the offerings requested for me. Furthermore, I understand that my Protected Health Information will not be used or disclosed by Janssen for any other purpose without my prior authorization unless permitted by law or unless information that specifically identifies me is removed. I understand that Janssen will make every effort to keep my information private. Further, I understand that if my information is accidentally shared, federal privacy laws do not require that the person/party receiving it not disclose the information further and that such information provided to a third party may no longer be protected by federal privacy laws.

I understand that I am not required to sign this Patient Authorization Form. My choice about whether to sign will not change the way my Healthcare Providers or Insurers treat me. If I refuse to sign the Patient Authorization Form, or revoke my authorization later, I understand that this means I will not be able to participate or receive assistance from Janssen CarePath.

This authorization will last until I am no longer participating in Janssen CarePath, or accessing my Janssen CarePath Account. I understand that I may cancel or revoke this Authorization at any time by mailing a letter requesting such cancellation to Janssen CarePath, 2250 Perimeter Park Drive, Suite 300, Morrisville, NC 27560 or by informing my Healthcare Providers and Insurers in writing that I do not want them to share any information with Janssen. I further understand that cancellation or revocation will not affect Janssen's ability to use and disclose Protected Health Information that it has received prior to its receipt of my cancellation and revocation of participation in the program. My authorization will also end if Janssen CarePath support programs or the Janssen CarePath Account is discontinued. Furthermore, I understand that I have the right to see or copy the Protected Health Information my Healthcare Providers or Insurers have given to Janssen.

Patient name: _____ Date of birth (mm/dd/yyyy): _____

Patient address: _____

City: _____ State: _____ ZIP Code: _____

Patient sign here: _____ Date: _____

If the patient cannot sign, patient's legally authorized representative must sign below:

By: _____ Date: _____

(Signature of person legally authorized to sign for patient)

Describe relationship to patient and authority to make medical decisions for patient:

Janssen CarePath
2250 Perimeter Park Drive, Suite 300
Morrisville, NC 27560
Fax 855-998-4422

