					FAX: <u>1-888-267-3277</u> For Medicare Advantage Part B: Please Use Medicare Request Form	
Please indicate: Start of treatment: Start date/ / /						
Continuation of therapy: Date of last treatment / /						
Precertification Requested By:		Phone:		Fax:		
A. PATIENT INFORMATION						
First Name:	Last Name:			DOB:		
Address:		City:		State:	ZIP:	
Home Phone: Work Phone:		Cell Phone:		Email:		
Patient Current Weight: lbs or kgs Patier						
B. INSURANCE INFORMATION			5			
	coverage?	🗌 Yes 🗌 No				
	-	-	Carrier Name:			
	Insured:					
Medicare: Yes No If yes, provide ID #:	Medic	aid: 🗌 Yes 🛛	No If yes, pro	ovide ID #:		
C. PRESCRIBER INFORMATION						
First Name:	Last Name:		(Check one,): 🗌 M.D. 🗌 D.	.O. 🗌 N.P. 🗌 P.A.	
Address:		City:		State:	ZIP:	
Phone: Fax:	St Lic #:	NPI #:	DEA #:		UPIN:	
Provider Email:	Office Contact Name:		1	Phone:	•	
Specialty (Check one): Oncologist Hematologis	st 🔲 Other:					
D. DISPENSING PROVIDER/ADMINISTRATION INFO						
Place of Administration: Self-administered Physician's Office Bioscript Phone: Coram Phone: Other: Agency Name: Phone: Outpatient Facility: Facility Name: Phone: Outpatient Infusion Center: Center Name: Addministration code(s) (CPT): Address: PRODUCT INFORMATION		Dispensing Provider/Pharmacy: Patient Selected choice Physician's Office Retail Pharmacy Specialty Pharmacy Other: Name:		macy		
Rolvedon (eflapegrastim-xnst) Dose: Directions for Use:						
F. DIAGNOSIS INFORMATION - Please indicate primary ICD code and specify any other where applicable.						
Primary Indication:						
G. CLINICAL INFORMATION - Required clinical information must be completed in its entirety for all precertification requests.						
For All Requests (clinical documentation required): Yes No Is the patient completing an existing chemotherapy regimen that requires current use of the requested medication to remain unchanged? If yes, indicate start date of chemotherapy regimen: / Yes No Has the patient had a contraindication, intolerance, or ineffective response to Neulasta on Neulasta Onpro (pegfilgrastim)? Yes No Hairy cell leukemia Yes No Will the requested medication be used for treatment of neutropenic fever following chemotherapy? Hematopoietic acute radiation syndrome Yes No Will the requested medication be used for the treatment of radiation-induced myelosuppression following a radiological/nuclear incident?						

Active don® (eflapegrastim-xnst)

Continued on next page

Aetna Precertification Notification Phone: 1-866-752-7021 (TTY: 711)

Rolvedon[®] (eflapegrastim-xnst) Medication Precertification Request

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♥aetna

 Aetna Precertification Notification

 Phone:
 <u>1-866-752-7021</u> (TTY: <u>711)</u>

 FAX:
 <u>1-888-267-3277</u>

(All fields must be completed and legible for precertification review.)

For Medicare Advantage Part B: Please Use Medicare Request Form

(All lields	must be completed and legible for precent	ilication review.)	Please Use Medicare Request Form		
Patient First Name	Patient Last Name	Patient Phone	Patient DOB		
G CLINICAL INFORMATION (continued)	Required clinical information must be comp	latad in its antiraty for all prosorti	ication requests		
Yes No Will the requested medical For which of the following indications is Primary prophylaxis of febrile neutrope	th myelosuppressive anti-cancer therapy ation be used in combination with any other ing chemotherapy at the same time as they ation be administered with a weekly chemo is the requested medication being presc enain a patient with a solid tumor or non- meceived, is currently receiving, or will be re- incidence of febrile neutropenia? eceived, is currently receiving, or will be re- lt in a 10-19% incidence of febrile neutrope- eceived, is currently receiving, or will be re- lt in less than 10% of febrile neutropenia? Does the patient have at least two patient- Please select the patient's risk factors belo Active infections, open wounds, or rece Bone marrow involvement by tumor pro Previous chemotherapy or radiation the Poor performance status Pore performance status Previous episodes of FN Other serious co-morbidities, including cardiovascular disease; please explain: Please select the patient's risk factors belo Cartive infections, open wounds, or rece Bone marrow involvement by tumor pro Previous chemotherapy or radiation the Poor performance status Por performance status Por performance status Previous episodes of FN Other; please explain: Neidered to be at high risk for febrile neutro Please select the patient's risk factors belo Active infections, open wounds, or rece Age greater than or equal to 65 years Bone marrow involvement by tumor pro Previous chemotherapy or radiation the Poor performance status Poor performance status Poor performance status Poor performance status Previous episodes of FN Other serious co-morbidities, including cardiovascular disease; please explain: Previous episodes of FN Other serious co-morbidities, including cardiovascular disease; please explain: Persistent neutropenia Other bone marrow compromise, como please explain: poenia in a patient with a solid tumor or nor experienced a neutropenic complication or	y colony stimulating factor product receive radiation therapy? therapy regimen without breaks? ribed? yeloid malignancy ceiving myelosuppressive anti-ca ceiving myelosuppressive anti-ca elated risk factors? w (select all that apply): nt surgery ducing cytopenias rapy renal dysfunction, liver dysfunction myenia because of bone marrow of w (select all that apply): nt surgery ducing cytopenias rapy renal dysfunction, liver dysfunction myenia because of bone marrow of w (select all that apply): nt surgery ducing cytopenias rapy renal dysfunction, liver dysfunction renal dysfunction, liver dysfunction rapy renal dysfunction, liver dysfunction rapy	ts within any chemotherapy cycle? Incer therapy that is expected to result neer therapy that is neer therapy that is on, HIV infection, compromise or comorbidity?		
Yes No For the planned chemotherapy cycle, will the patient receive the same dose and schedule of chemotherapy as the previous cycle (for which primary prophylaxis was not received)?					
Other (please explain):					
□ Stem cell transplantation-related indications					
Other - Please explain:					
H. ACKNOWLEDGEMENT					
Request Completed By (Signature Requin	red):		Date: //		
Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent insurance act, which is a crime and subjects such person to criminal and civil penalties.					

The plan may request additional information or clarification, if needed, to evaluate requests.