

## Rubraca

## **Prior Authorization Request**

CVS Caremark administers the prescription benefit plan for the patient identified. This patient's benefit plan requires prior authorization for certain medications in order for the drug to be covered. To make an appropriate determination, providing the most accurate diagnosis for the use of the prescribed medication is necessary. Please respond below and fax this form to CVS Caremark toll-free at 1-866-249-6155. If you have questions regarding the prior authorization, please contact CVS Caremark at 1-866-814-5506. For inquiries or questions related to the patient's eligibility, drug copay or medication delivery; please contact the Specialty Customer Care Team: CaremarkConnect® 1-800-237-2767.

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Patient's Name:Patient's ID:		Patient's Date of Birth:				
					Ph	ysician's Name:
Specialty:Physician Office Telephone:		NPI#:				
Re	quest Initiated For:	_				
1.	What is the diagnosis?  ☐ Epithelial ovarian cancer ☐ Fallopian tube cancer ☐ Uterine leiomyosarcoma ☐ Other	☐ Primary peritoneal cancer ☐ Prostate cancer ☐ Pancreatic Adenocarcinoma				
2.	What is the ICD-10 code?					
3.	switched to a preferred product? If Yes, pleas					
4.	Is this request for continuation of therapy with	the requested product?				
5.	Is the patient currently receiving the requested product through samples or a manufacturer's patient assistance program? If unknown, answer Yes $\square$ Yes $\square$ No If Yes, skip to #9					
6.	Has the patient experienced disease progression or an unacceptable toxicity while receiving the requested drug/regimen? $\square$ Yes $\square$ No <i>No further questions</i> .					
7.	intolerable adverse event to treatment with Lyn	er, does the patient have a documented inadequate response or aparza? <i>ACTION REQUIRED: If Yes, attach supporting</i> No \(\sum \text{N/A}\) N/A diagnosis is NOT prostate or pancreatic cancer				
8.	Does the patient have a documented inadequate the preferred products (Lynparza and Zejula)?	e response or intolerable adverse event to treatment with both of $\square$ Yes $\square$ No				
9.	Is the patient currently receiving treatment with	h the requested drug?				
	☐ Yes ☐ No No further questions.	cceptable toxicity while on the current regimen?				
11.	Will the requested drug be used as a single age					
	Send completed form to: Case Review Unit,	CVS Caremark Prior Authorization Fax: 1-866-249-6155				

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## Complete the following section based on the patient's diagnosis, if applicable. Section A: Epithelial Ovarian Cancer, Primary Peritoneal Cancer, Fallopian Tube Cancer 12. Does the patient have germline or somatic BRCA-mutated disease? ACTION REQUIRED: If Yes, attach chart note(s) or test results confirming BRCA mutation status. □ Yes □ No □ Unknown 13. Will the requested drug be used as maintenance therapy? ☐ Yes ☐ No 14. What is the clinical setting in which the requested drug will be used? ☐ Recurrent disease ☐ Advanced (stage II-IV) disease, *skip to #16* ☐ Other: 15. Is the patient in a complete or partial response to platinum based chemotherapy? $\square$ Yes $\square$ No No further questions. 16. Is the patient in a complete or partial response to primary therapy? ☐ Yes ☐ No Section B: Prostate Cancer 17. What clinical setting will the requested drug be used? ☐ Metastatic disease ☐ Other 18. Is the disease castration-resistant? ☐ Yes ☐ No 19. Does the tumor have a deleterious BRCA mutation (germline, somatic, or both)? ACTION REQUIRED: If Yes, attach chart note(s) or test results confirming BRCA mutation status. ☐ Yes ☐ No ☐ Unknown 20. Has the patient been treated with androgen receptor-directed therapy? ☐ Yes ☐ No 21. Has the patient been treated with a taxane-based chemotherapy? If Yes, skip to #23 \(\simeg\) Yes \(\simeg\) No 22. Is the patient unfit for chemotherapy? $\square$ Yes $\square$ No 23. Will the patient receive concurrent therapy with a gonadotropin-releasing hormone (GnRH) analog? ☐ Yes ☐ No 24. Has the patient had a bilateral orchiectomy? \(\begin{align\*} \Pi \text{ Yes} \\ \Bigsim \text{ No} \end{align\*} Section C: Uterine Leiomyosarcoma 25. Does the patient have BRCA2-altered uterine leiomyosarcoma? ACTION REQUIRED: If Yes, attach chart note(s) or test results confirming BRCA2 mutation status. $\square$ Yes $\square$ No $\square$ Unknown 26. What is the place in therapy in which the requested drug will be used? ☐ First-line treatment ☐ Subsequent treatment 27. What is the clinical setting in which the requested drug will be used? ■ Advanced disease ☐ Recurrent disease ☐ Metastatic disease ☐ Inoperable disease ☐ Other Section D: Pancreatic Adenocarcinoma 28. What is the clinical setting in which the requested drug will be used? ☐ Metastatic disease

29. Does the tumor have a BRCA mutation (germline or somatic) or a PALB2-mutation?

☐ Other

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	ACTION RE	onfirming BRCA or PA	LB2 mutation			
		A mutation (germling B2 mutation	ne or somatic)			
30.	Has the patient weeks? □ Y		platinum-based o	chemotherapy (e.g	g., cisplatin, carboplatin)	for at least 16
31.	. Has the disease progressed during treatment with platinum-based chemotherapy (e.g., cisplatin, carboplatin) $\square$ Yes $\square$ No					
					nentation supporting k or the benefit plan :	
X_ Pre	escriber or A	uthorized Signat	ure		Date (mm/dd/	

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