

Iowa Department of Human Services

FAX Completed Form To 1 (800) 574-2515

Provider Help Desk 1 (877) 776-1567

Request for Prior Authorization Lesinurad (Zurampic)

(PLEASE PRINT - ACCURACY IS IMPORTANT)

IA Medicaid Member ID #	Patient name	DOB			
Patient address					
Provider NPI	Prescriber name	Phone			
Prescriber address		Fax			
Pharmacy name	Address	Phone			
•					
Prescriber must complete all information above. It must be legible, correct, and complete or form will be returned.					
Pharmacy NPI	Pharmacy fax NDC				

Prior authorization is required for lesinurad (Zurampic). Requests for doses above the FDA approved dose will not be considered. Requests will be considered for patients when the following criteria are met:

- 1) Patient is 18 years of age or older; and
- 2) Patient has a diagnosis of hyperuricemia associated with gout; and
- 3) Patient has not achieved target serum uric acid levels or patient remains symptomatic with a maximally tolerated dose of a xanthine oxidase inhibitor (allopurinol or febuxostat) for at least 3 months; and
- 4) Patient has documentation of a previous trial and therapy failure with probenecid in combination with a xanthine oxidase inhibitor; and
- 5) Patient has an estimated creatinine clearance (eCrCl) > 45 mL/min; and
- 6) Documentation is provided lesinurad will be used in combination with a xanthine oxidase inhibitor.
 - a. If taking allopurinol, dose should be ≥ 300 mg per day (or ≥ 200 mg per day in patients with an eCrCl < 60 mL/min); and
- Patient does not have a contraindication to therapy including any of the following:
 - a. Severe renal impairment (eCrCl <30 mL/min)
- d. On dialysis

b. End stage renal disease

e. Tumor lysis syndrome

c. Kidney transplant recipient

f. Lesch-Nyhan syndrome

If criteria for coverage are met, initial requests will be given for 6 months. Continuation of therapy will be considered when the following criteria are met:

- 1) Patient continues to take medication in combination with a xanthine oxidase inhibitor.
 - a. If taking allopurinol, dose should be ≥ 300 mg per day (or ≥ 200 mg per day in patients with an eCrCl < 60 mL/min); and
- 2) Patient has an eCrCl > 45 mL/min; and
- Patient does not have a contraindication to therapy including any of the following:
 - a. Severe renal impairment (eCrCl <30 mL/min)
- d. On dialysis

b. End stage renal disease

e. Tumor lysis syndrome

c. Kidney transplant recipient

- f. Lesch-Nyhan syndrome
- 4) Documentation of a positive clinical response to lesinurad.

The required trials may be overridden when documented evidence is provided that the use of the agent(s) would be medically contraindicated.

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Non-Preferred				
Zurampic				
Strength	Dosage Instructions	Quantity	Day's Supply	
Diagnosis:				
Initial Requests:				
Target Serum Uric Acid L	.evel:			
Current Serum Uric Acid	Level (attach lab results):			
	tomatic while on a maximally tolerated		oxidase inhibitor for at	
Document trial of a xanth	nine oxidase inhibitor:			
Drug name & dose:	Tr	ial dates:		
Reason for failure:				
Document trial of a probe	enecid in combination with a xanthin	ne oxidase inhibito	or:	
Drug name & dose:	rug name & dose: Trial dates:			
Reason for failure:				
Estimated Creatinine Clea	arance (eCrCl): Da	ate calculated:		
	ombination with a xanthine oxidase inh I) mg per day in patients with an eCrCl		purinol, dose should be	
☐ Yes, provide drug name	e and dose:			
Does patient have a contr	raindication to therapy including an	y of the following:	:	
Severe renal impairment (e	eCrCl < 30 mL/min):	0		
End stage renal disease:	☐ Yes ☐ No	0		
Kidney transplant recipient	: Yes N	0		
On dialysis:	☐ Yes ☐ N	0		
Tumor lysis syndrome:	☐ Yes ☐ N	0		
Lesch-Nyhan syndrome:	☐ Yes ☐ N	0		

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Renewal Requests:

Is lesinurad being used in combination with a x 300 mg per day (or \geq 200 mg per day in patient				irinol, dose should be ≥
Yes, provide drug name and dose:	☐ No			
Estimated Creatinine Clearance (eCrCl):			culated:	
Does patient have a contraindication to ther	apy includ	ling any of th	ne following:	
Severe renal impairment (eCrCl < 30 mL/min):	☐ Yes	☐ No		
End stage renal disease:	☐ Yes	☐ No		
Kidney transplant recipient:	☐ Yes	☐ No		
On dialysis:	☐ Yes	☐ No		
Tumor lysis syndrome:	☐ Yes	☐ No		
Lesch-Nyhan syndrome:	☐ Yes	☐ No		
Provide documentation of positive clinical response	onse to lesi	nurad therapy	/:	
Attach lab results and other documentation	as necess	sary.		
Prescriber signature (Must match prescriber listed above.)			Date of submission	

IMPORTANT NOTE: In evaluating requests for prior authorization the consultant will consider the treatment from the standpoint of medical necessity only. If approval of this request is granted, this does not indicate that the member continues to be eligible for Medicaid. It is the responsibility of the provider who initiates the request for prior authorization to establish by inspection of the member's Medicaid eligibility card and, if necessary by contact with the county Department of Human Services, that the member continues to be eligible for Medicaid.

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