

Texas Standard Prior Authorization Request Form for Prescription Drug Benefits

NOFR002 | 0615 Texas Department of Insurance

Please read all instructions below before completing this form.

Please send this request to the issuer from whom you are seeking authorization. **Do not send this form** to the Texas Department of Insurance, the Texas Health and Human Services Commission, or the patient's or subscriber's employer.

Beginning September 1, 2015, health benefit plan issuers must accept the Texas Standardized Prior Authorization Request Form for Prescription Drug Benefits if the plan requires prior authorization of a prescription drug or device.

In addition to commercial issuers, the following public issuers must accept the form: Medicaid, the Medicaid managed care program, the Children's Health Insurance Program (CHIP), and plans covering employees of the state of Texas, most school districts, and The University of Texas and Texas A&M Systems.

Intended Use: Use this form to request authorization **by fax or mail** when an issuer requires prior authorization of a prescription drug, a prescription device, formulary exceptions, quantity limit overrides, or step-therapy requirement exceptions. An Issuer may also provide an **electronic version of this form** on its website that you can complete and submit electronically, through the issuer's portal, to request prior authorization of a prescription drug benefit.

Do not use this form to: 1) request an appeal; 2) confirm eligibility; 3) verify coverage; 4) request a guarantee of payment; 5) ask whether a prescription drug or device requires prior authorization; or 6) request prior authorization of a health care service.

Additional Information and Instructions:

Section I - Submission:

Enter the name and contact information for the issuer or the issuer's agent that manages or administers the issuer's prescription drug benefits, as applicable. An issuer or agent may have already prepopulated its contact information on the copy of this form posted on its website.

Section VI – Prescription Compound Drug Information:

List the quantities of ingredients in units of measure (mg, ml, etc.).

Section VIII - Patient Clinical Information:

Enter ICD Version 9 or 10, as applicable.

Section IX — Justification:

In the space provided or on a separate page:

- Provide pertinent clinical information to justify requests for initial or ongoing therapy, or increases in current dosage, strength, or frequency.
- Explain any comorbid conditions and contraindications for formulary drugs.
- Provide details regarding titration regimen or oncology staging, if applicable.
- Provide pertinent information about any step-therapy exception, if applicable.

Attach supporting clinical documentation (medical records, progress notes, lab reports, etc.), if needed.

Note: Some issuers may require more information or additional forms to process your request. If you think more information or an additional form may be needed, please check the issuer's website before faxing or mailing your request.

TEXAS STANDARDIZED PRIOR AUTHORIZATION REQUEST FORM FOR PRESCRIPTION DRUG BENEFITS

SECTION I — S	SUBMISSIO	N										
Submitted to:				Phone:			Fax:			Date:		
SECTION II —	REVIEW											
time frar	ne may seri	Review Requeste iously jeopardize Prescriber's Des	the life or	_			-	•		_		
			signee.									
SECTION III —	PATIENT 1	INFORMATION		D.I.			500					
Name:				Phone:						/lale Other	Uı	emale nknown
Address:				City:						State:	ZIP Co	de:
Issuer Name (i	f different f	from Section I):	Membe	Member or Medicaid ID #:				Group #:				
BIN # (if available):			PCN (if	PCN (if available):				Rx ID # (if available):				
SECTION IV —	- Prescrib	ER INFORMATIO	ON									
Name:				NPI#:				Specialty:				
Address:				City:						State: ZIP Code:		de:
Phone: Fax:				Office Contact Name:					Contact Phone:			
		TION DRUG INFO ug, identify all			Section VI,	below.)						
Requested Dru	ıg Name:											
Strength: Route of Administration:				Quantity: Days' Supply:			upply:	Expected Therapy Duration:				
	-	edge this medica		nrovii	mate date 1	herany ir	nitiated:					1
For Provider A			incrupy (up	рголп	nate date (пстару п	milatea					/
			NDC#:				D	ose Per Ad	ministr	ation:		
		TION COMPOU										
Compound Dru	ug Name:											
Ingredient		NDC#	Qı	uantity	Ingredient				ND	C#	Quantity	

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SECTION VII – Requested Dev		Expec	Expected Duration of Use:			HCPCS Code (If applicable):			
	— PATIENT CLINICAL INFORM	IATION							
Patient's diagn	osis related to this request:						on:	ICD Code:	
	ollowing information to the both has taken for this diagnosis:	est of your kn	owledge)						
Drug Name		Strength	Frequency		orted and Sto eximate Dur		escribe Response, Reasc for Failure, or Allergy		
Drug Allergies:					Height (if a	applicable):	Wei	ght (if applicable)	
	ratory values and dates (attac		w):						
Date		Test				Value			
SECTION IX —	JUSTIFICATION (SEE INSTRUC	TION PAGE SE	CTION IX)						
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