# MINNESOTA UNIFORM FORM FOR PRESCRIPTION DRUG PRIOR AUTHORIZATION (PA) REQUESTS AND FORMULARY EXCEPTIONS

#### INSTRUCTIONS

Important: Please read all instructions and information before completing the form.

Please do NOT send this form to a patient's employer or to the Minnesota Department of Health (MDH) or to the Minnesota Administrative Uniformity Committee (AUC).

Note: This version of the form (C-2.0) is current as of October 2015, and supersedes previous versions of Minnesota Department of Health forms for PA requests and formulary exceptions.

This form will not change frequently. The form version number and most recent revision date are displayed in the lower right corner.

## Overview:

The following form is made available by the Minnesota Department of Health (MDH) pursuant to statute, to facilitate exchanges of information between prescribers and patients' insurance carriers, HMOs, Pharmacy Benefits Managers (PBMs), or other payers\* of prescription drug claims.

# Intended use and requirements:

The form is intended primarily for use by prescribers, or those designated and authorized to act on behalf of prescribers, to:

#### 1. Request an exception to a prescription drug formulary.

- Requests for formulary exceptions are requests to make nonformulary prescription drugs available to a patient as a formulary drug.
  - Minnesota Statutes, section 62J.497, Subd. 4 requires that all health care providers must submit requests for
    formulary exceptions using the uniform form, and that all payers must accept this form from health care providers.
    No later than January 1, 2011, the uniform formulary exception form must be accessible and submitted by health
    care providers, and accepted and processed by group purchasers, through secure electronic transmissions. Note: A
    previous restriction in law that facsimile was not considered "secure electronic transmission" was removed in 2010.

## 2. Request a prior authorization (PA) for a prescription drug.

- Prescription drug prior authorization requests are requests for pre-approval from a payer for specified medications or quantities of medications.
  - Minnesota Statutes, section 62J.497, subd. 5 requires that by January 1, 2016, drug PA requests must be accessible and submitted by health care providers, and accepted by payers, electronically using the NCPDP SCRIPT Standard version 2013101.

## **Additional Instructions:**

- Prescribers, or their designees, use parts A-F as applicable. Payers making the form available on their websites may prepopulate section A. Payers use section G when responding to requests.
- Payers may request additional information or clarification needed to process formulary exceptions and PA requests.
- Payers may supply additional instructions or other relevant or legally required information with their response.
- Complete section F when submitting prescription drug PA requests to the Minnesota Department of Human Services.

<sup>\*</sup> Note: The term "payers" is used to avoid possible confusion. The electronic submission and acceptance requirements of Minnesota Statutes § 62J.497, subd. 4 and 5, apply to "group purchasers". The term "group purchaser" is defined in Minnesota Statutes § 62J.03, subd. 6 and can be considered more commonly as "payer".



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S	see additional instructions	s and overview, Instr	uctions page.				
Pleas	eing used for:						
Formulary Exception	on Prior Authoriza	ntion (PA) Request	Unsure/Unknown				
A   Destination This form	n is haing submitted t	• (Pavers making this form	n available on their websites may pre-populate section A.)				
1	_						
2 411		<del></del>					
·	Secure Fav		Other:				
	Secure Fax:		Other:				
the patient's prescription benefit card ID number separate prescription benefit ID number), provide	v, please note: If the patient has pre (the "cardholder ID"). If the patient' the patient's health plan ID numbe	s prescription benefits are int r.	parate or "carved out" from the health plan benefits, provide tegrated with the health plan coverage (if there is no  Gender:				
		City, State, Zip:	<del></del>				
Health Plan or Prescription Plan:			Patient Health Plan ID Number:				
<u></u>			(OR PRESCRIPTION PLAN ID IF DIFFERENT THAN HEALTH PLAN ID)				
C   Prescriber Information	tion						
Prescriber Name (LAST, FIRST, MI):		NPI:	Specialty:				
Prescriber Business Address:		City, State, Zip:					
Health Plan or Prescription Plan:			Patient Health Plan ID Number:				
Prescriber Phone:		Prescriber Secure Fax:					
Prescriber Point of Contact (POC) Name:		POC Phone:	POC Secure Fax:				
(IF DIFFERENT THA	N PRESCRIBER)	V	NT THAN PRESCRIBER)				
Clinic/Location/Facility Name:			Clinic/Location/Facility Contact Name:				
Clinic/Location/Facility Phone:			Secure Clinic/Location/Facility Fax:				
Clinic/Location/Facility Address:		City, State, Zip:					
"X" DEA number (buprenorphine prescriber status nun	nber, always preceded by "x," issued per	r the Drug Addiction Treatment A	Act of 2000 (Data 2000)):				
D   Prescription Drug I	nformation (Med	ication information	on)				
	use the medication, e.g, daily, four t		ms, e.g., 30mg, 15mg/ml, etc. Medication "dosing schedule" s, as needed, etc. If request is for a Minnesota Department of				
Drug Being Requested:		Strength:					
(REQUESTED DRUG NAME)		(E.G., 30 MG, 15	6 MG/ML, ETC)				
		Date Therapy Initiated:					
Duration of Therapy Expected:		Authorization Start Date					
Clinical Drug Trial Request?  (NOTE: THE MINNESOTA DEPT. OF HUMAN SERVICES I Rationale for DAW?	DOES NOT COVER CLINICAL DRUG TRIALS)	Is Dispense as Written (I	DAW) Specified?				
s patient currently being treated with the drug requested?		Date Started:					



# E | Patient Clinical Information Diagnosis Related to Medication Reguest:

Diagnosis Related to Medication	request:							
Drug Allergies:				Height:		Weight:		
(IF RELEVANT TO THIS REQUEST)				(IF RELEVANT TO THIS REQUEST)		(IF RELEVANT TO THIS REQUEST)		
PREVIOUS THERAPIES TRIED / FAI "dosing schedule" is used to repo						30 mg, 15 mg/ml, etc. Medication		
Drug Name	Strength	Dosing Schedule	Date Prescribed	Date Stopped	pped Describe Adverse Reaction or Efficacy Failure			
RATIONALE FOR REQUEST (and a	lso include any additi	onal pertinent clinical informa	tion/comments regardi	ng rationale:				
F   <b>Pharmacy</b>	Informat	tion						
Pharmacy Name:	narmacy Name:			NPI: Pharmacy Phone:				
Pharmacy Address:				City State 7in:				
NDC Number for Prescription Drug Being Requested:				Pharmacy Fax:				
G   Request D	atarmin:	ation (may be	completed b					
			-	Data of Davidson				
Date Request Received by Payer:				Date of Decision:				
Payer Responder/Contact Name:			Paye					
Payer Respondent/Contact Email:			Requ	Request Approved/Denied:				
Pharmacy Authorization/Referen		ICADI E TO DAVED						
(IF APPLICABLE TO PAYER)  Comments Regarding Decision: (INCLUDE EFFECTIVE AND END DATES OF DECISION IF APPLICABLE)								
Comments negariting becision.	INCLODE EL LECTIVE AINE	PEND DATES OF DECISION II APPEL	C IDEL)					
Additional Information or Instruc				ed at a				
Note: Group purchasers may sup and processes; other notification				with their response.	Examples of additional inf	ormation might include: Appeals rights		
	.,		IL					
CONFIDENTIALITY MOTICS TO			t de foi es	. 16				
						e hereby notified that any disclosure, ease immediately notify the sender to		
arrange for its return. Thank you			,,,	,		• •		

