



UnitedHealthcare Pharmacy
Clinical Pharmacy Programs

Program Number	2020 P 2059-7
Program	Prior Authorization/Medical Necessity Oxford – Buprenorphine/Naloxone Products
Medication	Buprenorphine/naloxone products: Bunavail*and Suboxone * (Brand Only)
P&T Approval Date	7/2015, 10/2016, 3/2017, 9/2017, 9/2018, 7/2019, 7/2020
Effective Date	10/1/2020; Oxford: 10/1/2020

1. Background

Suboxone* and Bunavail* are Schedule III narcotic medications available under the Drug Abuse Treatment Act (DATA) of 2000 for the treatment of opioid dependence. Only qualified doctors with the necessary DEA (Drug Enforcement Agency) identification number can prescribe or dispense buprenorphine products for opioid addiction therapy.

Bunavail* and Suboxone* contain an opiate antagonist to guard against misuse. Intravenously administered naloxone will block the effect of opiates and cause withdrawal symptoms.

This program requires a member to meet treatment criteria prior to the coverage of buprenorphine/naloxone combination products. It also requires the member to try the preferred combination product buprenorphine/naloxone (generic Suboxone) or Zubsolv prior to receiving coverage for Bunavail*, Suboxone *.

2. Coverage Criteria^a:

A. Initial Authorization

1. **Suboxone*** (Brand Only) **or Bunavail*** will be approved based on **all** of the following criteria:

a. The prescriber is qualified under DATA 2000 to prescribe buprenorphine products as indicated by a unique DEA identification number

-AND-

b. The patient is being treated for opioid dependence

-AND-

c. The medication is not being used solely for pain management

-AND-

d. Both of the following:

i. **One** of the following:

- (a) Submission of medical records (e.g. chart notes) documenting an inadequate response to a minimum 30-day trial of Zubsolv. (30-day trial must be completed prior to Prior Authorization/Medical Necessity request.)
- (b) The member has experienced adverse effects or has a contraindication to Zubsolv. (List manifestation of the adverse reaction or reason for contraindication and the prescriber's expectation for an alternative experience with Bunavail or Brand Suboxone)

-AND-

ii. **One** of the following:

- (a) Submission of medical records (e.g. chart notes) documenting an inadequate response to a minimum 30-day trial of buprenorphine/naloxone (generic Suboxone). (30-day trial must be completed prior to Prior Authorization/Medical Necessity request.)
- (b) The member has experienced adverse effects or has a contraindication to generic buprenorphine/naloxone (generic Suboxone). (List manifestation of the adverse reaction or reason for contraindication and the prescriber's expectation for an alternative experience with Bunavail or Brand Suboxone)

Authorization will be issued for 12 months.

B. Reauthorization:

1. **Suboxone*** (Brand Only) **or Bunavail*** will be approved based on the following criterion:

a. Documentation of positive clinical response

Authorization will be issued for 12 months.

^a State mandates may apply. Any federal regulatory requirements and the member specific benefit plan coverage may also impact coverage criteria. Other policies and utilization management programs may apply.

3. Additional Clinical Rules:

- Supply limits may be in place.
- Bunavail, Suboxone (Brand Only) are typically excluded from coverage. Tried/Failed criteria may be in place. Please refer to plan specifics to determine exclusion status.
- Notwithstanding Coverage Criteria, UnitedHealthcare may approve initial and re-authorization based solely on previous claim/medication history, diagnosis codes (ICD-10) and/or claim logic. Use of automated approval and re-approval processes varies by program and/or therapeutic class.

4. References:

1. Suboxone [package insert]. Richmond, VA: Indivior Inc.; October
2. Center for Substance Abuse Treatment. Clinical Guidelines for the Use of Buprenorphine in the Treatment of Opioid Addiction. Treatment Improvement Protocol (TIP) Series 40. DHHS Publication No. (SMA) 04-3939. Rockville, MD: Substance Abuse and Mental Health Services Administration, 2004.
3. Bunavail [package insert]. Raleigh, NC: BioDelivery Sciences International, Inc.; October 2019.

Program	Prior Authorization/Medical Necessity Oxford – Buprenorphine/Naloxone Products
Change Control	
Date	Change
7/2015	New Program
10/2016	Annual update. Decreased reauthorization period from 24 months to 12 months. Updated references.
3/2017	Administrative update. Removed requirement for medical record submission to verify opioid dependence diagnosis. Updated references.
9/2017	Changed reference from intolerance to adverse reaction to Zubsolv. Removed DEA waiver requirement.
9/2018	Annual review. Removed reference to brand Suboxone tablets (brand no longer available). Updated references.
7/2019	Removed generic Suboxone and buprenorphine/naloxone tablets from medications covered by criteria. Added criteria for DATA2000 prescriber.
7/2020	Annual review. Updated references. Clarified timing of 30 day trial.