

NC Pharmacy Prior Approval Request for Opioid Dependence Therapy Agents

Beneficiary Information

1. Beneficiary Last Name: _____ 2. First Name: _____
3. Beneficiary ID #: _____ 4. Beneficiary Date of Birth: _____ 5. Beneficiary Gender: _____

Prescriber Information

6. Prescribing Provider NPI #: _____ Provider Fax #: _____
7. Requester Contact Information - Name: _____ Phone #: _____ Ext. _____

Drug Information

8. Drug Name: _____ 9. Strength: _____ 10. Quantity Per 30 Days: _____
11. Length of Therapy (in days): up to 30 Days 60 Days 90 Days 120 Days 180 Days 270 Days 365 Days

Clinical Information

For Coverage of Buprenorphine/Naloxone SL Films, and Zubsolv:

1. Has the beneficiary Failed one preferred drug? **Yes** **No** Please List: _____
1a. Allergic Reaction 1b. Drug-to-drug interaction. Please describe reaction: _____
2. Previous episode of an unacceptable side effect or therapeutic failure. Please provide clinical information: _____
3. Clinical contraindication, co-morbidity, or unique patient circumstance as a contraindication to preferred drug(s).
Please provide clinical information: _____
4. Age specific indications. Please give patient age and explain: _____
5. Unique clinical indication supported by FDA approval or peer reviewed literature. Please explain and provide a general reference: _____
6. Unacceptable clinical risk associated with therapeutic change. Please explain: _____

For Coverage of Buprenorphine Sublingual Tablets:

7. Does the Beneficiary have a diagnosis of Opioid Dependence? **Yes** **No**
8. Is the beneficiary unable to use Suboxone Film? **Yes** **No** If Yes, please specify one or more of the following conditions)
 Beneficiary is pregnant: Please Provide Estimated Due Date: _____ **Max Length of Therapy is 270 Days**
 Beneficiary is breast feeding **Max Length of Therapy is 60 Days (can be renewed)**
 Beneficiary has an allergy to naloxone (rashes, hives, pruritis, bronchospasm, angioneurotic edema and anaphylactic shock) **Max Length of Therapy is 365 Days**
 Other condition Please List: _____
9. Has the prescriber reviewed the controlled substances reporting system database prior to writing the prescription to ensure that concomitant opioid use is not occurring? **Yes** **No**
10. Is the maximum daily dose less than or equal to 32 mg/day? **Yes** **No**

Signature of Prescriber: _____ Date: _____

(Prescriber Signature Mandatory)

I certify that the information provided is accurate and complete to the best of my knowledge, and I understand that any falsification, omission, or concealment of material fact may subject me to civil or criminal liability.