

Please provide the information below, print your answers, attach supporting documentation, sign, date and return to our office as soon as possible to expedite this request.

Please FAX responses to: (800) 869-7791. Phone: (855) 322-4082.

Patient	Date of Birth	Molina Member ID#	
Pharmacy Name	Pharmacy NPI	Telephone Number	Fax Number
Prescriber	Prescriber NPI	Telephone Number	Fax Number
Medication and Strength			Qty/Days Supply
Directions for Use			
<p>1. Is this request for a continuation of therapy? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes, is there documentation of a positive clinical benefit? <input type="checkbox"/> Yes <input type="checkbox"/> No</p> <p>2. Indicate patient's diagnosis: <input type="checkbox"/> Moderate to severe opioid use disorder <input type="checkbox"/> Other. Specify: _____</p> <p>3. Select from the following for your patient and complete associated question(s):</p> <p><input type="checkbox"/> Patient is pregnant. Estimated delivery date (EDD): _____ Was pregnancy confirmed with a lab test by the provider? <input type="checkbox"/> Yes <input type="checkbox"/> No Is buprenorphine prescriber managing patient's pregnancy? <input type="checkbox"/> Yes <input type="checkbox"/> No Has patient been stable on buprenorphine/naloxone for at least 8 weeks? <input type="checkbox"/> Yes <input type="checkbox"/> No</p> <p><input type="checkbox"/> Patient is breastfeeding. Delivery date: _____</p> <p><i>Patients approved based on breastfeeding, will be approved for 12 months following delivery. Transition to a buprenorphine/naloxone combination product is required for ongoing treatment thereafter.</i></p> <p><input type="checkbox"/> Patient has experienced a documented serious allergic or idiosyncratic reaction to the buprenorphine/naloxone combination product. Chart notes documenting reaction are required.</p> <p><input type="checkbox"/> Patient has continued to experience severe nausea or daily headache after a 7-day trial of buprenorphine/naloxone sublingual tablet and sublingual film formulations.</p>			

Indicate formulations tried for at least 7 days (check all that apply):

- Sublingual film
- Sublingual tab

4. Best practice is to limit patients to a 7-day supply at a time for the first month of treatment.

Indicate the intended day supply per fill for your patient: 7 day 14 day 28 day
If over a 7-day supply is indicated:

- Is the reason due to transportation complications? Yes No

If no, provide reason: _____

- Has patient demonstrated evidence of stability (8 weeks of treatment) taking buprenorphine monotherapy or buprenorphine/naloxone? Yes No

If yes, how long has patient been clinically stable? _____

Prescriber Signature	Prescriber Specialty	Date
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Notice Prohibiting Redisclosure of Alcohol or Drug Treatment Information

This information has been disclosed to you from records protected by Federal confidentiality rules (42 CFR part 2). The Federal rules prohibit you from making any further disclosure of this information unless further disclosure is expressly permitted by the written consent of the person to whom it pertains or as otherwise permitted by 42 CFR part 2. A general authorization for the release of medical or other information is NOT sufficient for this purpose. The Federal rules restrict any use of the information to criminally investigate or prosecute any alcohol or drug abuse patient.