Date: <Insert Date>

Payer Name: <Insert Payer Name>

Payer Address: <Insert Plan Address> Payer Fax Number: <Insert Plan Fax Number>

Attn: <Appeals Department>

To Whom It May Concern:

I am writing on behalf of my patient <Insert Patient Name> to provide additional information supporting medical necessity for the treatment with WELLBUTRIN XL® (bupropion hydrochloride) extended-release tablets. Within this letter, I am providing my patient's medical history, diagnosis, a description of their previous drug treatment, and a summary of their proposed treatment plan. I have also provided my clinically based rationale supporting the medical necessity of WELLBUTRIN XL for my patient.

Patient Information:

Patient's Name		Date of Birth		
Patient's Address				
City	State	Zip Code		
Member ID #	Policy or Group #			
☐ I need approval for a drug that requires a p	rior authorization prior to t	reatment		
Medication:				
□ WELLBUTRIN XL (bupropion hydrochloride) extended-release tablets: 150 mg once daily				
□ WELLBUTRIN XL (bupropion hydrochloride) extended-release tablets: 300 mg once daily				
Date Started:	Expected Length of	Therapy:		
Diagnosis – Please list all diagnoses being codes.	g treated with WELLBUT	RIN XL and corresponding ICD-10		
□ F33 Major depressive disorder, recurrent (includes recurrent episodes of seasonal affective disorder and recurrent episodes of seasonal depressive disorder)				
□ F33.0 Major depressive disorder, recurrent, mild				
□ F33.1 Major depressive disorder, recurrent, moderate				
□ F33.2 Major depressive disorder, recurrent, severe without psychotic features				
□ F33.3 Major depressive disorder, recurrent, severe with psychotic features				
□ F33.4 Major depressive disorder, recurrent, in remission				
□ F33.8 Other recurrent depressive disorders				
□ F33.9 Major depressive disorder, recurrent, unspecified				

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	Dates of Drug Trials	Results of previous drug trials
	1	1
	2	2
	3	3
	4	4
CLINICAL RATIONALE FOR MEI	DICAL NECESSITY	
☐ Alternate drug(s) contraindica	ated or previously tried, bu	ıt with adverse outcome
☐Therapeutic Failure	• •	
□Adverse Events		
□Sexual Dysfunction		
□Anxiety		
□Suicidal Ideation		
□Other		
		significant adverse clinical outcome with
nedication change		
☐ WELLBUTRIN XL 150 mg	g once daily; duration	
☐ WELLBUTRIN XL 300 mg	g once daily; duration	
blets) is medically necessary for	my patient. Please find atta	XL (bupropion hydrochloride) extended-release ched the additional documents that support my approval, please contact me at <insert phone<="" td=""></insert>
ncerely		
Insert Healthcare Provider Name	;>	
Insert Signature>		
		ant state therapy legislation, notes and product

FOR THE PRESCRIBERS BACKGROUND INFORMATION:

INDICATION

WELLBUTRIN XL® (bupropion hydrochloride) extended-release tablets is indicated for the treatment of major depressive disorder (MDD), and for the prevention of seasonal major depressive episodes in patients with a diagnosis of seasonal affective disorder (SAD). Periodically reevaluate long-term usefulness for the individual patient.

IMPORTANT SAFETY INFORMATION

WARNING: SUICIDAL THOUGHTS AND BEHAVIORS

SUICIDALITY AND ANTIDEPRESSANT DRUGS:

Antidepressants increased the risk of suicidal thoughts and behavior in children, adolescents, and young adults in short-term trials. These trials did not show an increase in the risk of suicidal thoughts and behavior with antidepressant use in subjects aged 65 and older.

In patients of all ages who are started on antidepressant therapy, monitor closely for worsening, and for emergence of suicidal thoughts and behaviors. Advise families and caregivers of the need for close observation and communication with the prescriber.

Contraindications

WELLBUTRIN XL is contraindicated in:

- patients with a seizure disorder
- patients with a current or prior diagnosis of bulimia or anorexia nervosa, due to a higher incidence of seizures
- patients undergoing abrupt discontinuation of alcohol, benzodiazepines, barbiturates, or antiepileptic drugs
- patients taking other bupropion products, including Zyban
- patients taking a monoamine oxidase inhibitor (MAOI) or within 14 days of discontinuing MAOI treatment due
 to an increased risk of hypertensive reactions. Starting WELLBUTRIN XL in a patient treated with reversible
 MAOIs such as linezolid or intravenous methylene blue is contraindicated.
- patients with hypersensitivity to bupropion or other ingredients of WELLBUTRIN XL

Warnings and Precautions

- WELLBUTRIN XL is not approved for smoking cessation treatment; however, bupropion HCl sustained-release is approved for this use. Postmarketing reports of serious or clinically significant neuropsychiatric adverse events with smoking cessation treatment have included changes in mood (including depression and mania), psychosis, hallucinations, paranoia, delusions, homicidal ideation, aggression, hostility, agitation, anxiety, and panic, as well as suicidal ideation, suicide attempt, and completed suicide. Observe patients attempting to quit smoking with WELLBUTRIN XL for the occurrence of such symptoms and instruct them to discontinue WELLBUTRIN XL and contact a healthcare provider if they experience such adverse events.
- Bupropion is associated with a dose-related risk of seizures. The dose should not exceed 300 mg once daily.
 Increase the dose gradually. Discontinue WELLBUTRIN XL and do not restart treatment if the patient experiences a seizure. Use with extreme caution in patients with a history of seizure or cranial trauma, or in patients treated with other medications that lower the seizure threshold.
- Treatment with WELLBUTRIN XL can result in elevated blood pressure and hypertension. Assess blood pressure before initiating treatment with WELLBUTRIN XL and monitor periodically during treatment.

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- Antidepressant treatment can precipitate a manic, mixed, or hypomanic manic episode. Prior to initiating WELLBUTRIN XL, screen patients for a history of bipolar disorder and the presence of risk factors for bipolar disorder (e.g., family history of bipolar disorder, suicide, or depression). WELLBUTRIN XL is not approved for the treatment of bipolar depression.
- Depressed patients treated with bupropion have had a variety of neuropsychiatric signs and symptoms, including delusions, hallucinations, psychosis, concentration disturbance, paranoia, and confusion. Some of these patients had a diagnosis of bipolar disorder. In some cases, these symptoms abated upon dose reduction and/or withdrawal of treatment. Discontinue WELLBUTRIN XL if these reactions occur.
- The pupillary dilation that occurs following use of many antidepressant drugs including WELLBUTRIN XL may trigger an angle closure attack (Angle-Closure Glaucoma) in a patient with anatomically narrow angles who does not have a patent iridectomy.
- Anaphylactoid/anaphylactic reactions have occurred during clinical trials with bupropion, as well as rare, postmarketing reports of erythema multiforme, Stevens-Johnson syndrome, and anaphylactic shock associated with bupropion.

Adverse Reactions

• The most common adverse reactions that occurred in at least 5% of patients treated with bupropion HCl sustained-release (300 mg and 400 mg per day) and at a rate at least twice the placebo rate were: anorexia, dry mouth, nausea, insomnia, dizziness, pharyngitis, abdominal pain, agitation, anxiety, tremor, palpitation, sweating, tinnitus, myalgia, urinary frequency, and rash.

Drug Interactions

- An increased dose of bupropion may be necessary if co-administered with CYP2B6 inducers based on
 clinical exposure but should not exceed the maximum recommended dose. Bupropion inhibits CYP2D6 and
 can increase concentrations of: antidepressants, antipsychotics, beta-blockers, and Type 1C
 antiarrhythmics. Consider dose reduction when using with bupropion. Dose bupropion with caution when
 used with drugs that lower seizure threshold. CNS toxicity can occur when bupropion is used concomitantly
 with dopaminergic drugs.
- WELLBUTRIN XL can cause false-positive urine test results for amphetamines.

Use in Specific Populations

- Pregnancy: Use only if benefit outweighs potential risk to the fetus. Healthcare providers are encouraged to register patients in the Pregnancy Exposure Registry by calling 1-844-405-6185 or visiting https://womensmentalhealth.org/research/pregnancyregistry/.
- In patients with moderate to severe hepatic impairment (Child-Pugh score: 7 to 15), the maximum dose is 150 mg every other day. In patients with mild hepatic impairment (Child-Pugh score: 5 to 6) or renal impairment (glomerular filtration rate <90 mL/min), consider reducing the dose and/or frequency of dosing.
- Advise patients to read the FDA-approved patient labeling (Medication Guide).

To report SUSPECTED ADVERSE REACTIONS, contact Bausch Health at 1-800-321-4576 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

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