DISCHARGE DATE:

PATIENT ENROLLMENT

touchpoints Support Services

Prescription only valid if faxed FAX COMPLETED FORM TO: 1-877-329-8484

TP ID# (TOUCHPOINTS USE ONLY):

TOUCHPOINTS PHONE: 1-800-848-4876

PRESCRI	BFR IN	IFOR	MATI	ON
T IL SCILL				

Staff Contact E-mail

(First)

Name

Address

City

Date of Birth

Home Phone #

Email Address

Alcohol Dependence

ICD-9

303.

303.

303.

303.

303.

Other:

PATIENT INFORMATION

Best Day to Call 🗆 M 🛛 T

Best Time to Call D Morning

ICD-10

F10.

F10._

F10.

F10.

Check if patient has concurrent medications

F10.

Prescriber Name*				
State License #	DEA #			
Prescriber Phone #	NPI #			
Facility Name	Fax #			
Address				
City	State	Zip Code		
Staff Contact Name				
Staff Contact Phone #				

(Last)

Gender

State

🗆 TH

PATIENT DIAGNOSIS—Please complete the diagnosis code(s) you would like to use

ICD-10

F11.

F11.

F11.

F11.

F11.

Patient's concurrent medication:

Opioid Dependence

Afternoon

 $\Box W$

INSTRUCT PATIENT TO LIST ALTERNATE DESIGNEE OR CONTACTS ON PAGE 2.

by filling in the additional digits. (A list of codes can be found on page 3)

ICD-9

304.

304.

304.

304.

304.

Other:

Mobile Phone #

Evening

🗆 F

□ Male □ Female

Zip Code

Patient has tried and failed the

Please list any known allergies to

medications or other substances:

following medication(s):

PLEASE COMPLETE ALL FIELDS TO AVOID PROCESSING DELAYS

ADMITTANCE DATE:

INJECTION PROVIDER/SPECIALTY PHARMACY INFORMATION

Will your office/facility be injecting VIVITROL?

Yes, ALL doses

 $\hfill\square$ No, please locate an Injection Provider or refer to Provider below

Provider Name

Provider Address

Provider Phone #

Preferred specialty pharmacy (if applicable)

Special shipping instructions/restrictions

PATIENT INSURANCE INFORMATION

Payment Method 🗆 Insured 🗆 Paying out-of-pocket

ATTACH A COPY OF BOTH SIDES OF THE PATIENT'S INSURANCE CARD(S).

IF NOT AVAILABLE, COMPLETE SECTION BELOW.

Carrier Phone #

Group ID #

PRIMARY INSURANCE / MEDICAL INSURANCE

Insurance Type 🗆 HMO 🔅 PPO 🔅 Medicaid 🔅 Medicare

Carrier Name Policyholder Name

Relationship to Patient

Policyholder Employer Name

Policy #

PHARMACY BENEFIT PLAN (PBM) - Required for Co-Pav card activation: see page 3

PBM Name	
Policyholder Name	
Relationship to Patient	PBM Phone #
Policyholder Employer Name	
Policy #	Rx Grp:
Rx BIN #	Rx PCN:

PRESCRIPTION INFORMATION Patient Name Date VIVITROL 380 mg x 1 unit Inject 380 mg IM every 4 weeks or every 1 month Provider State License

Refill ______ times (Complete refills to minimize interruption in monthly VIVITROL therapy)

By signing below, I verify that the information provided in this Touchpoints enrollment form is complete and accurate to the best of my knowledge. I understand that Alkermes reserves the right at any time and for any reason, without notice, to modify this Touchpoints enrollment form or to modify or discontinue any services or assistance provided through Touchpoints. Finally, I authorize Alkermes, United BioSource Corporation, Armada Health Care, LLC, and OPUS Health as my designated agents to use and disclose my patient's health information as necessary to verify the accuracy of any information provided, to provide reimbursement services through Touchpoints, to forward the above prescription, by fax or other mode of delivery, to a pharmacy for fulfillment, and (as applicable) to assess my patient's leigibility for co-pay assistance.

PROVIDER ATTEST	ATION		* Prescriber signatur	re must be the same a	s the prescriber name above
Prescriber's Signature		(If applicable) Prescriber's Signature (No Stamps Allowed)	Dispense as Written Substitution Permitted	Date of Signature	

PLEASE SEE <u>IMPORTANT SAFETY INFORMATION</u> ON PAGE 4. PLEASE SEE <u>PRESCRIBING INFORMATION</u> AND <u>MEDICATION GUIDE</u>, OR VISIT <u>VIVITROL.COM</u>. PLEASE REVIEW MEDICATION GUIDE WITH PATIENTS.

PATIENT ENROLLMENT

PATIENT REPRESENTATIVE

By signing below, I authorize my Designee(s), listed below, to receive administrative information related to my treatment, such as appointment reminders, and to make decisions on my behalf—for which I will remain liable—regarding delivery of VIVITROL [®] (naltrexone for extended-release injectable suspension). Alkermes is not liable for any decision(s) made by the Designee(s) or actions taken in reliance on such Designee(s) decisions.				
Please list any Designees authorized to receive administrative information related to my treatment:				
Designee Name (1)	Relationship	Phone #		
Designee Name (2)	Relationship	Phone #		
Patient's Signature	Dat	te of Signature		

PATIENT AUTHORIZATION FOR USE/DISCLOSURE OF HEALTH INFORMATION

By signing below, **I authorize: 1.** my prescribing physician, **2.** the healthcare provider designated to administer VIVITROL to me ("Administering HCP"), **3.** one or more network specialty pharmacies[†] (collectively"Providers") to use and disclose to Alkermes, United BioSource Corporation, Intouch Solutions, OPUS Health, my Designee(s) listed above (collectively"Recipients") my medical and other information set forth on the first page of this form, including information about my treatment with VIVITROL (taken together, "Information") **for the specific purposes** of allowing Recipients to facilitate: **1.** ordering, delivering, and administering VIVITROL, **2.** conducting reimbursement verification and obtaining payment from my Health Plan(s), **3.** providing me with educational and therapy support services by mail, text-messaging, e-mail, and/or telephone, **4.** referring me to, or determining my eligibility for, other programs, foundations or alternative sources of funding or coverage to help me with the costs of VIVITROL and **5.** providing me with support services, which may include sending me product information materials and treatment reminders. **Information May Be Further Disclosed:** I understand that Information disclosed pursuant to this authorization could be re-disclosed by a Recipient and may no longer be protected by federal privacy law (HIPAA). **For California Residents:** California law prohibits the person receiving your health information from making further disclosure of it, unless another authorization for such additional disclosure is obtained from you or unless such disclosure is specifically required or permitted by law.

I understand that signing this authorization is voluntary and if I do not sign this authorization it will not affect my ability to obtain treatment from my Providers or obtain insurance or insurance benefits. I understand, however, that if I do not sign this authorization, I will not be eligible to receive the educational, support or other services described above. I understand I have the right to receive a copy of this authorization after I sign. I understand that the disclosure of my Information by my Providers may result in remuneration to one or more network specialty pharmacies.

I may withdraw this authorization at any time by mailing or faxing a written request to Touchpoints Reimbursement Support, 1670 Century Center Parkway, Memphis, TN 38134. Withdrawal of this authorization will end further uses and disclosures of my Information by Providers when they receive notice of my withdrawal except to the extent those uses and disclosures have been made in reliance upon this authorization and as permitted by applicable law. This authorization expires on the sooner to occur of (1) five years from the date indicated below or (2) the maximum period permitted by applicable state law, unless I withdraw it earlier.

Patient's Signature

Parent/Guardian/Legal Representative's Signature[‡]

Date of Signature

Authority/Relationship to Patient

† NETWORK SPECIALTY PHARMACIES

Acaria; Accredo; Aetna Specialty Pharmacy; AllCare Specialty Pharmacy; Amber; Armada Healthcare, LLC and its members; Bioscrip Specialty Pharmacy; Blount Specialty Pharmacy; Briova; Chartwell; Cigna Specialty Pharmacy; Community Pharmacy; CVS Caremark; Diplomat; Fairview; Humana Specialty Pharmacy; ICORE; Intermountain Homecare; KanCare Specialty Pharmacy; Medicine Shoppe; OptumRx; Orsini Healthcare; Perform Specialty; Pharmacy Advantage; Presbyterian; Prime; Providence Specialty Pharmacy; Transition Patient Services; Walgreens Specialty Pharmacy

For information about joining this network contact 1-800-VIVITROL (1-800-848-4876).

+ If patient is a minor without capacity to act alone under state law, signature of patient and parent/guardian/legal representative is required.

PLEASE SEE <u>IMPORTANT SAFETY INFORMATION</u> ON PAGE 4. PLEASE SEE <u>PRESCRIBING INFORMATION</u> AND <u>MEDICATION GUIDE</u>, OR VISIT <u>VIVITROL.COM</u>. PLEASE REVIEW MEDICATION GUIDE WITH PATIENTS.

PATIENT ENROLLMENT

CO-PAY INFORMATION FOR ELIGIBLE PATIENTS §

- □ (Check if "yes") I would like to receive co-payment assistance from Alkermes.
- 🗆 l certify that l am at least 18 years old, l am being treated for opioid dependence after detox or alcohol dependence.

Please confirm that you understand the eligibility requirements for the VIVITROL Co-pay Savings Program:

Co-payment assistance for VIVITROL is not valid for prescriptions that are purchased (in full or in part) by any federal, state or government funded program. Such programs includes, but are not limited to:

- Medicare, including Medicare Part D or Medicare Advantage plans
- Medicaid, including Medicaid Managed Care and Alternative Benefit Plans ("ABPs") under the Affordable Care Act
- Medigap
- Veterans Administration ("VA")
 Department of Defense ("DoD")
- •TRICARE®
- State funded programs such as medical or pharmaceutical assistance programs

If you use benefits from state, federal or government funded programs, such as those listed above, to help pay for your prescription for VIVITROL, you will no longer be eligible to participate in this program. **Do you agree to** follow these requirements and that you will NOT purchase your VIVITROL prescription with benefits from state, federal or government funded programs?

□ Yes □ No Patient's Signature

Date of Signature

INJECTION PROVIDER/SPECIALTY PHARMACY SELECTION INFORMATION (AS APPLICABLE)

If you have requested injection services for your patient, Touchpoints will provide a selection of several injectors based on geographic proximity to your patient's address listed on the enrollment form (from closest to farthest from such address).

These injection providers are listed on the VIVITROL Provider Locator¹¹ at www.VIVITROL.com.

These options will be provided to you for your patient. We will also contact the selected injection services provider to help coordinate injection services.

Touchpoints may coordinate and route patient prescriptions for VIVITROL among the specialty pharmacies participating in the Touchpoints Program ("network specialty pharmacies"). Prescriptions for product sent to Touchpoints are forwarded to network specialty pharmacies for fulfillment based on insurance plan requirements, specialty pharmacy capabilities, healthcare provider selection, patient preference, and pharmacy performance in filling VIVITROL prescriptions. Inclusion as a network specialty pharmacy is voluntary and free of charge for specialty pharmacies. For information about joining this network contact 1-800-VIVITROL (1-800-848-4876).

Alcohol Dependence: (ICD-9) 303.00 Acute alcoholic intoxication, unspecified drunkenness 303.01 Acute alcoholic intoxication, continuous drunkenness 303.90 Other and unspecified alcohol dependence, unspecified alcohol dependence, continuous drunkenness 303.91 Other and unspecified alcohol dependence, continuous drunkenness 303.92 Other and unspecified alcohol dependence, episodic drunkenness 303.93 Other and unspecified alcohol dependence, in remission (ICD-10) F10.2 Alcohol dependence - F10.20 Uncomplicated	 - F10.220 Uncomplicated - F10.221 Delirium - F10.229 Unspecified F10.230 Uncomplicated - F10.230 Uncomplicated - F10.231 Delirium - F10.232 With perceptual disturbance - F10.239 Unspecified F10.24 With alcohol-induced mood disorder F10.25 Alcohol dependence with alcohol-induced psychotic disorder - F10.250 With delusions - F10.250 Unspecified F10.250 Unspecified F10.250 With alcohol-induced psychotic disorder - F10.250 With alcohol-induced psychotic disorder - F10.250 Unspecified F10.26 With alcohol-induced persisting amnestic disorder F10.27 With alcohol-induced persisting dementia F10.28 Alcohol dependence with other alcohol-induced disorders - F10.280 Alcohol dependence with other alcohol-induced disorders 	 - F10.282 Alcohol dependence with alcohol-induced sleep disorder - F10.288 Alcohol dependence with other alcohol-induced disorder F10.29 With unspecified alcohol-induced disorder Opioid Dependence: (ICD-9) 304.00 Opioid type dependence, unspecified abuse 304.01 Opioid type dependence, continuous abuse 304.02 Opioid type dependence, episodic abuse 304.03 Opioid type dependence, in remission 304.70 Combinations of opioid type drug with any other drug dependence, continuous abuse 304.71 Combinations of opioid type drug with any other drug dependence, continuous abuse 304.72 Combinations of opioid type drug with any other drug dependence, episodic abuse 304.73 Combinations of opioid type drug with any other drug dependence, episodic abuse 304.73 Combinations of opioid type drug with any other drug dependence, episodic abuse 304.73 Combinations of opioid type drug with any other drug dependence, episodic abuse 	 F11.21 In remission F11.22 Opioid dependence with intoxication F11.220 Uncomplicated F11.221 Delirium F11.222 With perceptual disturbance F11.229 Unspecified F11.23 With withdrawal F11.24 With opioid-induced mood disorder F11.25 Opioid dependence with opioid-induced psychotic disorder F11.250 With delusions F11.251 With hallucinations F11.28 Opioid dependence with other opioid-induced disorder F11.28 Opioid dependence with other opioid-induced sexual dysfunction F11.28 Opioid dependence with other opioid-induced sexual dysfunction F11.282 Opioid dependence with other opioid-induced sleep disorder F11.283 Opioid dependence with other opioid-induced sleep disorder

§ Eligibility for Sponsored Co-pay Assistance: Offer valid only for prescriptions for FDA-approved indications. Patients must be at least 18 years old. If patients are purchasing their VIVITROL prescriptions with benefits from Medicare, including Medicare Part D or Medicare Advantage plans; Medicaid, including Medicaid Managed Care or Alternative Benefit Plans ("ABPs") under the Affordable Care Act; Medigap; Veterans Administration ("VA"); Department of Defense ("DoD"); TriCare®; or any similar state funded programs such as medical or pharmaceutical assistance programs, they are not eligible for this offer. Void where prohibited by law, taxed or restricted. Alkermes, Inc. reserves the right to rescind, revoke or amend these offers without notice.

II Enrollment in the Locator is voluntary and free of charge and, along with the provider-specific information in the Provider Locator, is based on healthcare provider responses. Inclusion in the Locator does not imply a referral, recommendation or endorsement by Alkermes. Alkermes has not independently verified the qualifications of any healthcare provider included in the Locator. We recommend that you research the credentials, qualifications, and experience of each provider before confirming an appointment. Alkermes shall not be liable to you or to anyone for any decision made or action taken in reliance on this information.

PLEASE SEE <u>IMPORTANT SAFETY INFORMATION</u> ON PAGE 4. PLEASE SEE <u>PRESCRIBING INFORMATION</u> AND <u>MEDICATION GUIDE</u>, OR VISIT <u>VIVITROL.COM</u>. PLEASE REVIEW MEDICATION GUIDE WITH PATIENTS.

IMPORTANT SAFETY INFORMATION FOR VIVITROL® (naltrexone for extended-release injectable suspension)

INDICATIONS

VIVITROL is indicated for:

- Treatment of alcohol dependence in patients who are able to abstain from alcohol in an outpatient setting. Patients should not be actively drinking at the time of
 initial VIVITROL administration.
- · Prevention of relapse to opioid dependence, following opioid detoxification.
- VIVITROL should be part of a comprehensive management program that includes psychosocial support.

CONTRAINDICATIONS

VIVITROL is contraindicated in patients:

• Receiving opioid analgesics

- · With current physiologic opioid dependence
- In acute opioid withdrawal
- Who have failed the naloxone challenge test or have a positive urine screen for opioids
- · Who have exhibited hypersensitivity to naltrexone, polylactide-co-glycolide (PLG), carboxymethylcellulose, or any other components of the diluent

WARNINGS/PRECAUTIONS

Vulnerability to Opioid Overdose: Because VIVITROL blocks the effects of exogenous opioids for approximately 28 days after administration, patients are likely to have a reduced tolerance to opioids after opioid detoxification. As the blockade dissipates, use of previously tolerated doses of opioids could result in potentially life-threatening opioid intoxication (respiratory compromise or arrest, circulatory collapse, etc). Cases of opioid overdose with fatal outcomes have been reported in patients who used opioids at the end of a dosing interval, after missing a scheduled dose, or after discontinuing treatment. Patients and caregivers should be told of this increased sensitivity to opioids and the risk of overdose.

Any attempt by a patient to overcome the VIVITROL blockade by taking opioids may lead to fatal overdose. Patients should be told of the serious consequences of trying to overcome the opioid blockade.

Injection Site Reactions: VIVITROL injections may be followed by pain, tenderness, induration, swelling, erythema, bruising, or pruritus; however, in some cases injection site reactions may be very severe. Injection site reactions not improving may require prompt medical attention, including, in some cases, surgical intervention. Inadvertent subcutaneous/adipose layer injection of VIVITROL may increase the likelihood of severe injection site reactions. Select proper needle size for patient body habitus, and use only the needles provided in the carton. Patients should be informed that any concerning injection site reactions should be brought to the attention of their healthcare provider.

Precipitation of Opioid Withdrawal: Withdrawal precipitated by administration of VIVITROL may be severe. Some cases of withdrawal symptoms have been severe enough to require hospitalization and management in the ICU. To prevent precipitated withdrawal, patients, including those being treated for alcohol dependence:

• Should be opioid-free (including tramadol) for a minimum of 7–10 days before starting VIVITROL.

Patients transitioning from buprenorphine or methadone may be vulnerable to precipitated withdrawal for as long as two weeks.

Patients should be made aware of the risk associated with precipitated withdrawal and be encouraged to give an accurate account of last opioid use.

Hepatotoxicity: Cases of hepatitis and clinically significant liver dysfunction have been observed in association with VIVITROL. Warn patients of the risk of hepatic injury; advise them to seek help if experiencing symptoms of acute hepatitis. Discontinue use of VIVITROL in patients who exhibit acute hepatitis symptoms.

Depression and Suicidality: Alcohol- and opioid-dependent patients taking VIVITROL should be monitored for depression or suicidal thoughts. Alert families and caregivers to monitor and report the emergence of symptoms of depression or suicidality.

When Reversal of VIVITROL Blockade Is Required for Pain Management: For VIVITROL patients in emergency situations, suggestions for pain management include regional analgesia or use of non-opioid analgesics. If opioid therapy is required to reverse the VIVITROL blockade, patients should be closely monitored by trained personnel in a setting staffed and equipped for CPR.

Eosinophilic Pneumonia: Cases of eosinophilic pneumonia requiring hospitalization have been reported. Warn patients of the risk of eosinophilic pneumonia and to seek medical attention if they develop symptoms of pneumonia.

Hypersensitivity Reactions: Patients should be warned of the risk of hypersensitivity reactions, including anaphylaxis.

Intramuscular Injections: As with any IM injection, VIVITROL should be administered with caution to patients with thrombocytopenia or any coagulation disorder.

Alcohol Withdrawal: Use of VIVITROL does not eliminate nor diminish alcohol withdrawal symptoms.

ADVERSE REACTIONS

Serious adverse reactions that may be associated with VIVITROL therapy in clinical use include severe injection site reactions, eosinophilic pneumonia, serious allergic reactions, unintended precipitation of opioid withdrawal, accidental opioid overdose, and depression and suicidality. The adverse events seen most frequently in association with VIVITROL therapy for alcohol dependence include nausea, vomiting, injection site reactions (including induration, pruritus, nodules, and swelling), muscle cramps, dizziness or syncope, somnolence or sedation, anorexia, decreased appetite or other appetite disorders. The adverse events seen most frequently in association with VIVITROL in opioid-dependent patients also include hepatic enzyme abnormalities, injection site pain, nasopharyngitis, insomnia, and toothache.

Alkermes

PLEASE SEE <u>PRESCRIBING INFORMATION</u> AND <u>MEDICATION GUIDE</u>, OR VISIT <u>VIVITROL.COM</u>. PLEASE REVIEW MEDICATION GUIDE WITH PATIENTS.

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