

Weekly/Monthly  
**Buvidal**<sup>®</sup>

**BUPRENORPHINE**

PROLONGED-RELEASE  
SOLUTION FOR INJECTION

# Pharmacy administration of Buvidal

## ADMINISTRATION RECORD FORM

For Pharmacist use only - not to be given to the patient.

For further information visit [www.buvidal.co.uk](http://www.buvidal.co.uk), where you will find a number of resources, including a Buvidal administration video and patient testimonials.

# Patient Details

Name: \_\_\_\_\_

Date of birth: \_\_\_\_\_

Name of keyworker: \_\_\_\_\_

Name of prescriber: \_\_\_\_\_

Date: \_\_\_\_\_

1. Did patient have any side effects after their last Buvidal injection?  
For example: at the site of injection etc. Yes  No

If yes, please state here and report as detailed in prescribing information:

2. Did patient have any withdrawal symptoms or cravings? Yes  No

If yes, please state here:

3. Has patient used any drugs on top of their Buvidal treatment? Yes  No

If yes, please state here:

Is there a current and fully correct Patient Specific Direction (PSD) in place? Yes  No

Dose confirmed with patient and cross-checked with prescription/PSD  mg

Opportunity provided to discuss any current medication, drug or alcohol use Yes  No

Opportunity provided to discuss possible side effects Yes  No

Are there any newly diagnosed medical conditions or prescribed medications since last review / administration appointment? Yes  No

If yes, please state here:

Has patient taken any drugs that haven't been prescribed in the last 24 hours? Yes  No

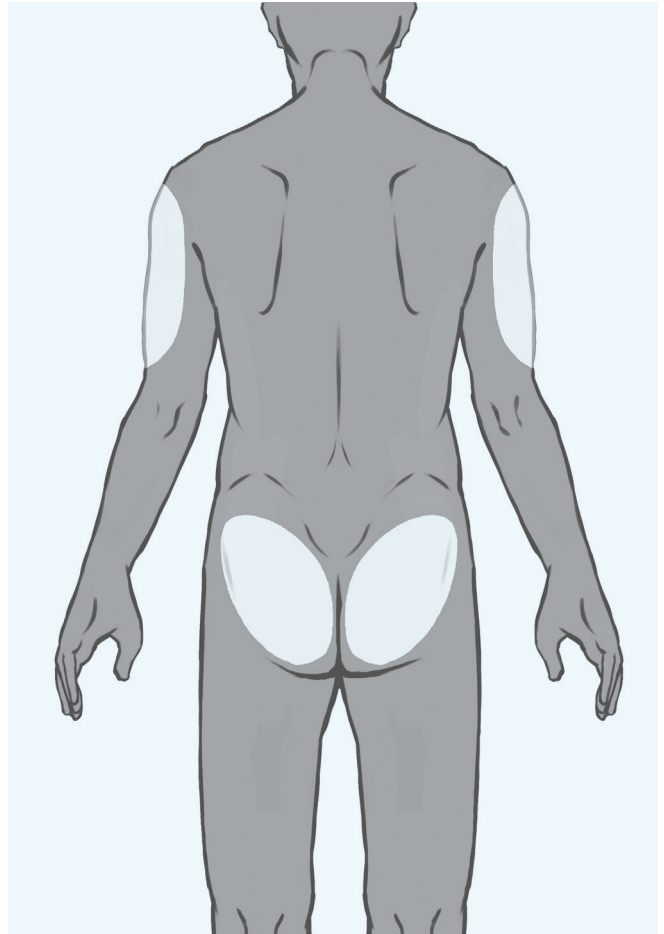
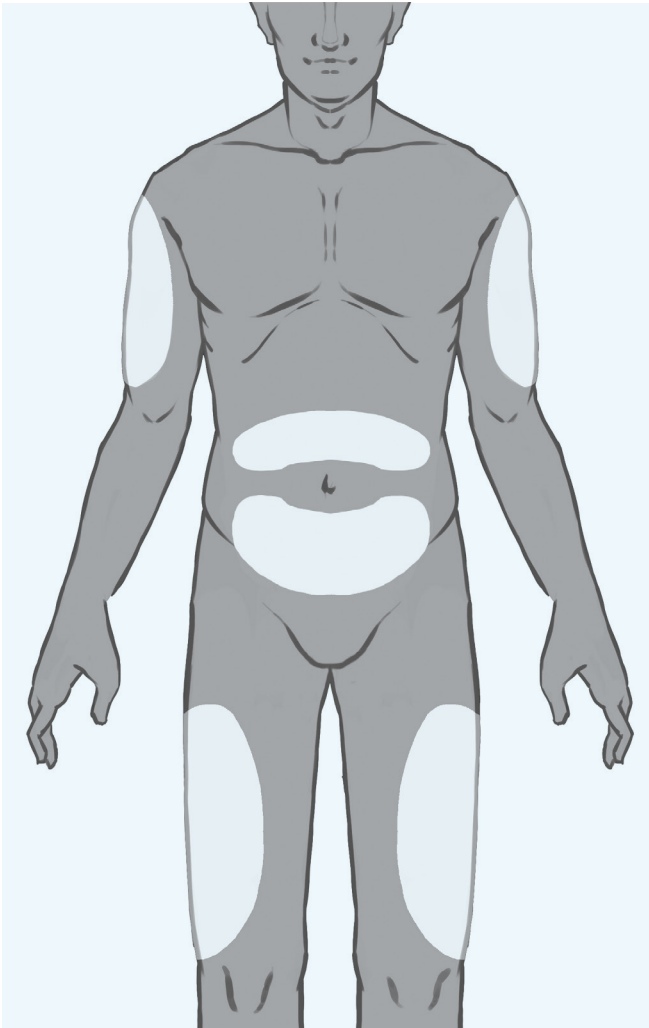
If yes, please state here:

Batch Number: \_\_\_\_\_ Expiry Date: \_\_\_\_\_

Date of next appointment: \_\_\_\_\_

Appointment card given to patient Yes  No

# Patient Details



Indicate where Buvidal was given by placing an 'X' on the body image

Patient consent (signature): \_\_\_\_\_

Pharmacist name: \_\_\_\_\_

Pharmacist GPhC number and signature: \_\_\_\_\_

1. On a scale of 1-10, with 1 being not satisfied and 10 being extremely satisfied, how satisfied was the patient with the Buvidal administration service they received today?

1 <input type="checkbox"/>	2 <input type="checkbox"/>	3 <input type="checkbox"/>	4 <input type="checkbox"/>	5 <input type="checkbox"/>	6 <input type="checkbox"/>	7 <input type="checkbox"/>	8 <input type="checkbox"/>	9 <input type="checkbox"/>	10 <input type="checkbox"/>
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2. Is the patient happy with the service they received today? Yes  No

3. What could have improved the patient's experience today?

# Prescribing Information for Buvidal® (buprenorphine prolonged-release solution for injection)

Please refer to the Summary of Product Characteristics (SmPC) before prescribing

**Active ingredient:** Buprenorphine. Prolonged-release solution for injection in pre-filled syringes. Weekly injection (8 mg, 16 mg, 24 mg, 32 mg) or monthly injection (64 mg, 96 mg, 128 mg, 160 mg).

**Indication:** Treatment of opioid dependence within a framework of medical, social and psychological treatment. Treatment is intended for use in adults and adolescents aged 16 years or over.

**Dosage:** To avoid precipitated withdrawal, initiate when objective and clear signs of mild to moderate withdrawal are evident, considering the duration of action of the opioid, time since last dose and degree of opioid dependence. Do not start until  $\geq 6$  hours after last heroin or short-acting opioid. Reduce methadone to  $\leq 30$  mg/day and start Buvidal®  $\geq 24$  hours after the last methadone dose. Buvidal® may trigger withdrawal symptoms in methadone-dependent patients. **Initiation in patients not already receiving buprenorphine:** Patients not previously exposed to buprenorphine, administer 4 mg sublingual buprenorphine and observe for an hour to confirm tolerability. Recommended starting dose of Buvidal® is 16 mg, with one or two additional 8 mg doses at least 1 day apart (target dose of 24 mg or 32 mg during the first week). The dose for the second week is the total dose administered during the first week. May transfer to monthly Buvidal® after four weeks and once stabilised. **Switching from sublingual buprenorphine:** Switch directly to weekly or monthly Buvidal®, starting on the day after the last sublingual buprenorphine dose. See SmPC for dose recommendations. **Maintenance:** Weekly or monthly as needed. One supplemental Buvidal® 8 mg dose may be administered between regular weekly or monthly doses (except 160 mg). The maximum dose is 32 mg weekly, with an additional 8 mg dose, or 160 mg monthly. Weekly doses may be administered up to 2 days before or after the weekly time point, and monthly doses may be administered up to 1 week before or after the monthly time point. If a dose is missed, administer the next dose as soon as practical. **Termination:** Consider prolonged-release characteristics and any withdrawal symptoms. If switching to sublingual buprenorphine, do so one week after the last weekly dose or one month after the last monthly dose of Buvidal®. **Elderly:** No dosing recommendations over 65 years. Consider renal and hepatic function.

**Administration:** Administration of Buvidal® is restricted to healthcare professionals only. For subcutaneous administration only. Inject slowly and completely into sufficient subcutaneous tissue of the buttock, thigh, abdomen, or upper arm area. Do not re-inject the same injection site for at least 8 weeks (each area can have multiple injection sites).

**Contraindications:** Hypersensitivity to buprenorphine or excipients. Severe respiratory insufficiency. Severe hepatic impairment. Acute alcoholism or *delirium tremens*.

**Special warnings and precautions for use:** Must not be administered intravenously, intramuscularly or intradermally. Monitor for any attempts to remove the depot. Some precautions associated with buprenorphine class. **Prolonged-release properties** of the product should be considered during treatment. Patients with concomitant medicines and/or comorbidities should be monitored for signs and symptoms of toxicity, overdose or withdrawal. **Respiratory depression:** Deaths reported with buprenorphine. Care in respiratory insufficiency. **CNS depression:** Buprenorphine may cause drowsiness. **Dependence:** Chronic administration of buprenorphine can produce opioid dependence. **Serotonin syndrome:** Concomitant serotonergic agents (e.g. monoamine oxidase inhibitors, selective serotonin re-uptake inhibitors, serotonin and noradrenaline re-uptake inhibitors or tricyclic antidepressants) may result in serotonin syndrome, a potentially life-threatening condition - if clinically warranted, observe carefully, particularly during initiation and dose increases and consider reducing or discontinuing therapy if serotonin syndrome is suspected. **Hepatitis, hepatic events and hepatic**

**impairment:** Recording of baseline liver function tests and viral hepatitis status recommended. Hepatic injury reported with buprenorphine. Caution with buprenorphine in moderate hepatic impairment – monitor for signs and symptoms of opioid withdrawal, toxicity and overdose. Monitor hepatic function regularly. **Drug withdrawal syndrome (GB):** Before starting any opioids, discuss withdrawal strategy with the patient. Dose tapering over weeks or months may be required. Risk of neonatal withdrawal syndrome following use in pregnancy. **Precipitation of opioid withdrawal syndrome:** Buprenorphine products have precipitated withdrawal symptoms in opioid-dependent patients when administered before the agonist effects from recent opioid use or misuse have subsided. **Renal impairment:** Caution in severe renal impairment. **QT-prolongation:** Caution with other medicines that prolong the QT interval and in patients with a history of long QT syndrome or other risk factors for QT prolongation. **Acute pain management:** A combination of opioids with high mu-opioid receptor affinity, non-opioid analgesics and regional anaesthesia might be necessary. Monitor and titrate, considering potential risk of overdose and/or death. **Sleep-related breathing disorders:** Opioids can cause sleep-related breathing disorders. **Opioid class effects:** See SmPC for details. **Interactions:** See SmPC for buprenorphine interactions. **Pregnancy and lactation:** Caution – see SmPC for details. **Driving and operating machines:** Minor to moderate influence, including drowsiness, dizziness or impaired thinking – likely to be pronounced by alcohol or CNS depressants. See SmPC for details of what individual patients should be told by the prescriber.

**Undesirable effects:** **Very common:** insomnia, headache, nausea, hyperhidrosis, drug withdrawal syndrome, pain. **Common:** infection, influenza, pharyngitis, rhinitis, lymphadenopathy, hypersensitivity, decreased appetite, anxiety, agitation, depression, hostility, nervousness, abnormal thinking, paranoia, medical dependence, somnolence, dizziness, migraine, paraesthesia, syncope, tremor, hypertonia, speech disorders, lacrimal disorder, mydriasis, miosis, palpitations, vasodilation, hypotension, cough, dyspnoea, yawning, asthma, bronchitis, constipation, vomiting, abdominal pain, flatulence, dyspepsia, dry mouth, diarrhoea, gastrointestinal disorder, rash, pruritus, urticaria, arthralgia, back pain, myalgia, muscle spasms, neck pain, bone pain, dysmenorrhoea, injection site reactions (pain, pruritus, erythema, swelling, reaction, induration, mass), peripheral oedema, asthenia, malaise, pyrexia, chills, neonatal withdrawal syndrome, chest pain, abnormal liver function tests. **Other:** urinary retention, injection site reactions (abscess, ulceration and necrosis). See SmPC for further details.

**Overdose:** Apply general supportive measures, closely monitoring and treating respiratory and cardiac status. Consider long duration of action of buprenorphine and prolonged release from the depot.

**Package quantities and UK net price:** 1 pre-filled syringe per pack. Weekly injection (8 mg (0.16 ml), 16 mg (0.32 ml), 24 mg (0.48 ml), 32 mg (0.64 ml)): £55.93. Monthly injection (64 mg (0.18 ml), 96 mg (0.27 ml), 128 mg (0.36 ml), 160 mg (0.45 ml)): £239.70. **Marketing authorisation numbers:** GB: PLGB 42800/0001, PLGB 42800/0003-9. **ROI and NI:** EU/1/18/1336/001-7, EU/1/18/1336/009. **Legal category:** POM. **Marketing authorisation holder:** Camurus AB, Ideon Science Park, SE-223 70 Lund, Sweden. Email: [Camurus.uk@camurus.com](mailto:Camurus.uk@camurus.com) Additional information available on request.

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Adverse events should be reported. Reporting forms and information can be found at [www.mhra.gov.uk/yellowcard](http://www.mhra.gov.uk/yellowcard) (or search for MHRA Yellow Card in the Google Play or Apple App Store) for the UK and <http://www.hpra.ie/homepage/about-us/report-an-issue> for Ireland. Adverse events should also be reported to Camurus AB via email: [safety@camurus.com](mailto:safety@camurus.com)