| | MEDICARE FORM Lupron Depot [®] (leuprolide acetate for depot suspension) Medication Precertification Request Page 1 of 3 (All fields must be completed and legible for precertification review.) reatment: Start date// tion of therapy, Date of last treatment// | | | | | For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974 For other lines of business: Please use other form. Note: Lupron Depot is non- preferred. The preferred product is Eligard. Firmagon is also a preferred product. | |
|---|---|---|--|---|--------------------------|---|--|
| Precertification Requested By | : | | Phor | ne: | Fax | : | |
| A. PATIENT INFORMATION | | | | | | | |
| First Name: | | Last Name: | • | | DOB: | I | |
| Address: | | | City: | | State: | ZIP: | |
| Home Phone: | Work Phone: | | Cell Phone: | | Email: | | |
| Patient Current Weight: Ibs | - | ent Height: inche | es or <u> </u> | Allergies: | | | |
| B. INSURANCE INFORMATION | | | | | | | |
| Aetna Member ID #: | | Does patient have oth | | | | | |
| Group #: Insured: | | If yes, provide ID#: Insured: | | Carrier Name: | | | |
| | | | | | | | |
| | | IVIO | edicald: 📋 Yes | ☐ No If yes, pro | vide ID #: | | |
| C. PRESCRIBER INFORMATIO First Name: | N | Last Name: | | (Check (|)ne)· □ M D | . 🗌 D.O. 🗌 N.P. 🗌 P. <i>I</i> | |
| Address: | | Last Name. | City: | (Onech e | State: | ZIP: | |
| Phone: Fax | x: | St Lic #: | NPI #: | DEA #: | 0.0.101 | UPIN: | |
| Provider Email: | | Office Contact Name | | | Phone: | - | |
| Agency Name: Administration code(s) (CPT): | Phone: | | _ | y Pharmacy | | | |
| Address: | 01-1- | 210 | | | | | |
| City: Phone: | | | | | | | |
| TIN: | | | NPI: | | | | |
| NPI: | | | _ | | | | |
| E. PRODUCT INFORMATION | | | | | | | |
| Request is for: Lupron Depot (I | euprolide acetate fo | r depot suspension) D |)ose: | F | requency: | | |
| F. DIAGNOSIS INFORMATION | Please indicate prim | ary ICD code and spec | ify any other whe | ere applicable. | | | |
| Primary ICD Code: | | Secondary ICD Cod | e: | Other | ICD Code: | | |
| Yes 🗌 No | ocumentation require r Lupron Depot-PED? upron Depot-PED form sex cord-stromal turn of dose is being require d drug being prescribe Is the patient undergo Will the patient receive unner Stage of puberty umors | ed for all requests): n for this request. nors, prostate cancer, re- ested: 3.75 mg 7 d for pubertal hormonal so bing gender transition? ve the requested drug con- the patient has reached: | ecurrent salivary .5 mg 11.25 suppression in an ncomitantly with g Stage I Sta | gland tumors india mg □ 22.5 mg □ adolescent patient? ender-affirming horn age II □ Stage III [| cations only:] 30 mg | - | |

For Ohio MMP:



MEDICARE FORM

Lupron Depot[®] (leuprolide acetate for depot suspension) Medication Precertification Request Page 2 of 3

(All fields must be completed and legible for precertification review.)

For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974

For other lines of business: Please use other form.

Note: Lupron Depot is nonpreferred. The preferred product is Eligard. Firmagon is also a preferred product.

| Patient First Name Patient Last Name Patient Phone Patient DOB G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. Patient salivary gland tumors | | | | | | | |
|---|---|--|--|--|--|--|--|
| | | | | | | | |
| | | | | | | | |
| Recurrent salivary gland tumors | | | | | | | |
| | | | | | | | |
| ☐ Yes ☐ No Is the tumor androgen receptor positive? | | | | | | | |
| For breast cancer, endometriosis, ovarian cancer, preservation of ovarian function, recurrent menstrual related attacks in acute porphyria or uterine | | | | | | | |
| leiomyomata (fibroids) indication only: | | | | | | | |
| Please select which Lupron Depot dose is being requested: 3.75 mg 11.25 mg | | | | | | | |
| Breast cancer | | | | | | | |
| Please indicate the patient's hormone receptor (HR) status: HR-positive HR-negative Unknown | | | | | | | |
| | | | | | | | |
| Ovarian cancer Please select: Epithelial ovarian cancer Fallopian tube cancer Primary peritoneal cancer Malignant sex cord-stromal tumor | | | | | | | |
| □ Preservation of ovarian function | | | | | | | |
| ☐ Yes ☐ No Is the patient premenopausal and undergoing chemotherapy? | | | | | | | |
| Prevention of recurrent menstrual related attacks in acute porphyria | | | | | | | |
| ☐ Yes ☐ No Is the requested drug being requested to prevent recurrent menstrual related attacks in acute porphyria? | | | | | | | |
| Yes No Is the requested drug being prescribed by, or in consultation with, a physician experienced in the management of porphyrias? | | | | | | | |
| Uterine leiomyomata (fibroids) | | | | | | | |
| Yes No Does the patient have a diagnosis of anemia (for example, Hct less than or equal to 30% and/or Hgb less than or equal to 10 g/dL)? | | | | | | | |
| \square Yes \square No Will the requested drug be used prior to surgery for uterine fibroids? | | | | | | | |
| For Continuation Requests (clinical documentation required for all requests): | | | | | | | |
| For gender dysphoria, malignant sex cord-stromal tumors, prostate cancer, recurrent salivary gland tumors continuation requests only: | | | | | | | |
| Please select which Lupron Depot dose is being requested: 🗌 3.75 mg 📋 7.5 mg 📋 11.25 mg 📋 22.5 mg 🔲 30 mg 🔲 45 mg | | | | | | | |
| Gender dysphoria | | | | | | | |
| Yes No Is the requested drug being prescribed for pubertal hormonal suppression in an adolescent patient? | | | | | | | |
| \longrightarrow Yes \square No Is the patient undergoing gender transition? | | | | | | | |
| Yes No Will the patient receive the requested drug concomitantly with gender-affirming hormones? | | | | | | | |
| └──────────────────────────────────── | n | | | | | | |
| ☐ Malignant sex cord-stromal tumors | | | | | | | |
| Yes No Has the patient experienced an unacceptable toxicity or disease progression while receiving the requested drug? | | | | | | | |
| □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ □ | | | | | | | |
| ☐ Yes ☐ No Has the patient had prior therapy with Lupron Depot within the last 365 days? | | | | | | | |
| 🗌 Yes 🔲 No Has the patient experienced clinical benefit while receiving the requested drug (e.g., serum testosterone less than 50ng/dl)? | | | | | | | |
| Yes No Has the patient experienced an unacceptable toxicity while receiving the requested drug? | | | | | | | |
| Recurrent salivary gland tumors | | | | | | | |
| Yes No Has the patient experienced an unacceptable toxicity or disease progression while receiving the requested drug? | | | | | | | |

Continued on next page



MEDICARE FORM

Lupron Depot[®] (leuprolide acetate for depot suspension) Medication Precertification Request Page 3 of 3

(All fields must be completed and legible for precertification review.)

For Ohio MMP: FAX: 1-855-734-9389 PHONE: 1-855-364-0974

For other lines of business: Please use other form.

Note: Lupron Depot is nonpreferred. The preferred product is Eligard. Firmagon is also a preferred product.

| Patient First Name | Patient Last Name | Patient Phone | Patient DOB | | | | | |
|--|--|--------------------------------------|-----------------------------|--|--|--|--|--|
| | | | | | | | | |
| G. CLINICAL INFORMATION (continued) – Required clinical information must be completed in its entirety for all precertification requests. | | | | | | | | |
| For breast cancer, endometriosis, ovarian cancer, preservation of ovarian function, recurrent menstrual related attacks in acute porphyria or uterine | | | | | | | | |
| fibroids continuation requests only: | | | | | | | | |
| Please select Lupron Depot dose for the following indications: 🗌 3.75 mg 🔲 11.25 mg | | | | | | | | |
| Breast cancer | | | | | | | | |
| Please indicate the patient's hormone receptor (HR) status: 🗌 HR-positive 🔲 HR-negative 🔲 Unknown | | | | | | | | |
| Yes No Has the patient experienced clinical benefit while receiving the requested drug? | | | | | | | | |
| ☐ Yes ☐ No Has the patient experienced an unacceptable toxicity while receiving the requested drug? | | | | | | | | |
| Endometriosis | | | | | | | | |
| Yes No Has the patient received previous therapy with the requested medication or Lupaneta Pack? | | | | | | | | |
| └──────────────────────────────────── | | | | | | | | |
| Yes No Is the patient's bone mineral density within normal limits? | | | | | | | | |
| How long has the patient received previous therapy with the requested drug and Lupaneta Pack? months | | | | | | | | |
| Ovarian cancer | | | | | | | | |
| Please select: 🗌 Epithelial ovarian cancer 🔲 Fallopian tube cancer 🔲 Primary peritoneal cancer 🔲 Malignant sex cord-stromal tumor | | | | | | | | |
| Yes INo Has the patient experienced clinical benefit while receiving the requested drug? | | | | | | | | |
| Yes No Has the patient experienced an unacceptable toxicity while receiving the requested drug? | | | | | | | | |
| Preservation of ovarian function | | | | | | | | |
| _ Yes □ No Is the patient premenopaus | | | | | | | | |
| Prevention of recurrent menstrual related | | | | | | | | |
| Yes No Is the requested medication being requested to prevent recurrent menstrual related attacks in acute porphyria? | | | | | | | | |
| 🗌 Yes 🔲 No Is the requested medication being prescribed by, or in consultation with, a physician experienced in the management of porphyrias? | | | | | | | | |
| Uterine leiomyomata (fibroids) | | | | | | | | |
| ☐ Yes ☐ No Has the patient received previous therapy with the requested drug or Lupaneta Pack? | | | | | | | | |
| → Yes No Does the patient have a diagnosis of anemia (for example, Hct less than or equal to 30% and/or Hgb less than or equal to 10g/dL)? | | | | | | | | |
| How long has the patient received previous therapy with the requested drug and Lupaneta Pack? months | | | | | | | | |
| └──────────────────────────────────── | patient have a diagnosis of anemia (for ex 10g/dL)? | ample, Hct less than or equal to | 30% and/or Hgb less than or | | | | | |
| └───> □ Yes [| No Will the requested drug be used pr | ior to surgery for uterine fibroids? | | | | | | |
| H. ACKNOWLEDGEMENT | | | | | | | | |
| Request Completed By (Signature Require | ed): | | Date: / / | | | | | |
| Any person who knowingly files a request for authorization of coverage of a medical procedure or service with the intent to injure, defraud or deceive | | | | | | | | |
| any insurance company by providing materially false information or conceals material information for the purpose of misleading, commits a fraudulent | | | | | | | | |

insurance act, which is a crime and subjects such person to criminal and civil penalties.

The plan may request additional information or clarification, if needed, to evaluate requests.