HEALTH OPTIONS

DMMA Approved: 08/2024

Request for Prior Authorization for Lantidra (donislecel-juju)

website Form – <u>www.highmarkhealthoptions.com</u>
Submit request via: Fax - 1-855-476-4158

All requests for Lantidra (donislecel-juju)require a Prior Authorization and will be screened for medical necessity and appropriateness using the criteria listed below.

Lantidra (donislecel-juju) Prior Authorization Criteria:

Coverage may be provided with a diagnosis of **Type I Diabetes** and the following criteria is met:

- Documentation the member has had Type I diabetes for more than 5 years accompanied by all the following despite intensive insulin management efforts (including insulin, devices, and education):
 - o Unable to reach target HbA1C due to repeated episodes of severe hypoglycemia
 - O At least one episode of severe hypoglycemia in the past 3 years defined as an event with symptoms compatible with hypoglycemia in which the member required the assistance of another person, and which was associated with either a blood glucose level < 50 mg/dL (2.8 mmol/L) or prompt recovery after oral carbohydrate, intravenous glucose, or glucagon administration
 - O Prescriber attestation that the member reports reduced awareness of hypoglycemia as defined by the absence of adequate autonomic symptoms at capillary glucose levels of < 54 mg/dL (3 mmol/L) AND the member is unable to prevent repeated severe hypoglycemia events using intensive diabetes management (including insulin, devices, and education)
- Prescriber by or in consultation with an endocrinologist
- The member must not have any of the following:
 - o Co-existing cardiac disease defined by one of the following:
 - Recent (within past 6 months) myocardial infarction
 - Angiographic evidence of non-correctable coronary artery disease
 - Evidence of ischemia on functional cardiac exam (with a stress echo test recommended for members with a history of ischemic disease).
 - Heart failure > New York Heart Association (NYHA) II
 - History of stroke within the past 6 months
 - Active alcohol or substance abuse-includes cigarette smoking (must be abstinent for six months). Active alcohol abuse should be considered using the current National Institute on Alcohol Abuse and Alcoholism (NIAAA) definitions.
 - Psychiatric disorder making the member not a suitable candidate for transplantation, e.g., schizophrenia, bipolar disorder, or major depression that is unstable or uncontrolled on current medication.
 - o History of non-adherence to prescribed regimens
 - o Active infection including hepatitis C, hepatitis B, HIV
 - TB (by history or currently infected as evidenced by a positive QuantiFERON® -TB Gold test or under treatment for suspected TB)
 - Any history of malignancies except squamous or basal skin cancer. Any member found to have squamous or basal cancer is required to have it removed prior to transplant.

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- Body Mass Index (BMI) > 27 kg/m2.
- \circ C-peptide response to glucagon stimulation (1 mg IV) (any C-peptide ≥ 0.3 ng/mL)
- O Creatinine clearance < 80 mL/min/1.73 m2 by 24-hour urine collection. If corrected creatinine clearance is < 80 and serum creatinine is < 1.2 mg/dl, then a nuclear renal scan is required to determine glomerular filtration rate.
- Serum creatinine consistently > 1.5 mg/dL
- o Macroalbuminuria (urinary albumin excretion rate > 300 mg/24h)
- O Baseline Hb < 12 gm/dL in women or < 13 gm/dL in men
- O Baseline liver function tests (LFT) outside of normal range (An initial LFT test panel with any values > 1.5 times normal upper limits will exclude a member without a retest. A re-test for any values between normal and 1.5 times normal should be made, and if the values remain elevated above normal limits, the member will be excluded.)
- Untreated proliferative retinopathy
- Positive pregnancy test, intent for future pregnancy, or male members' intent to procreate, unwilling to follow effective contraceptive measures, or presently breastfeeding
- o Insulin requirement > 0.7 IU/kg/day
- HbA1c > 12%
- Hyperlipidemia (fasting LDL cholesterol > 130 mg/dL, treated or untreated; and/or fasting triglycerides > 200 mg/dL)
- Under treatment for a medical condition requiring chronic use of steroids other than a previous organ transplant
- Use of coumadin or other antiplatelet or anticoagulant therapy, or member with PT INR > 1.5. Low dose aspirin is allowed after transplantation.
- History of Factor V deficiency
- Addison's disease
- Allergy to radiographic contrast material
- o Symptomatic cholecystolithiasis
- o Acute or chronic pancreatitis
- Symptomatic peptic ulcer disease
- o Severe unremitting diarrhea, vomiting, or other gastrointestinal disorders that could interfere with the ability to absorb oral medications
- o Received live attenuated vaccine(s) within 2 months
- The requested dose and frequency is in accordance with FDA-approved labeling, nationally recognized compendia, and/or evidence-based practice guidelines
- Must be age-appropriate according to FDA-approved labeling, nationally recognized compendia, or evidence-based practice guidelines
- Must be used in conjunction with concomitant immunosuppression
- **Initial Duration of Approval:** 12 months
- Reauthorization criteria
 - O Documentation the member has not achieved independence from exogenous insulin within one year of infusion OR within one year after losing independence from exogenous insulin after a previous infusion
- Reauthorization Duration of Approval: 12 months (maximum 3 infusions per lifetime)



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LANTIDRA (DONISLECEL-JUJU) PRIOR AUTHORIZATION FORM

as applicable to Highmark Health Options Pharmacy Services. FAX: (855) 476-4158 If needed, you may call to speak to a Pharmacy Services Representative. PHONE: (844) 325-6251 Mon – Fri 8:00 am to 7:00 pm PROVIDER INFORMATION Requesting Provider: NPI: Provider Specialty: Office Address: Office Phone: Office Fax: MEMBER INFORMATION Member Name: DOB: Member ID: Member weight: Height: REQUESTED DRUG INFORMATION Medication: Strength:
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Directions: Quantity: Refills:
Is the member currently receiving requested medication? Yes No Date Medication Initiated:
Is this medication being used for a chronic or long-term condition for which the medication may be necessary for the life of the
patient? Yes No
Billing Information
This medication will be billed: at a pharmacy OR medically, JCODE:
Place of Service: Hospital Provider's office Member's home Other
Place of Service Information
Name: NPI:
Address: Phone:
MEDICAL HISTORY (Complete for ALL requests)
Diagnosis: ICD Code:
How long has the member been diagnosed with Type I diabetes?
Does the member have any of the following (please check all that apply):
the inability to reach target HbA1C due to repeated episodes of severe hypoglycemia
History of at least one episode of severe hypoglycemia in the past 3 years defined as an event with symptoms compatible with
hypoglycemia in which the member required the assistance of another person, and which was associated with either a blood glucose
level < 50 mg/dL (2.8 mmol/L) or prompt recovery after oral carbohydrate, intravenous glucose, or glucagon administration
reduced awareness of hypoglycemia as defined by the absence of adequate autonomic symptoms at capillary glucose levels of <
54 mg/dL (3 mmol/L) AND the member is unable to prevent repeated severe hypoglycemia events using intensive diabetes
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LANTIDRA (DONISLECEL-JUJU) PRIOR AUTHORIZATION FORM (CONTINUED) – PAGE 2 OF 2

Please complete and fax all requested information below including any progress notes, laboratory test results, or chart documentation as applicable to Highmark Health Options Pharmacy Services. FAX: (855) 476-4158

If needed, you may call to speak to a Pharmacy Services Representative. PHONE: (844) 325-6251 Mon – Fri 8:00am to 7:00pm MEMBER INFORMATION Member Name: DOB: Height: Member ID: Member weight: **MEDICAL HISTORY (Complete for ALL requests)** Body Mass Index (BMI) > 27 kg/m2. \square C-peptide response to glucagon stimulation (1 mg IV) (any C-peptide \geq 0.3 ng/mL) ☐ Creatinine clearance < 80 mL/min/1.73 m2 by 24-hour urine collection. If corrected creatinine clearance is < 80 and serum creatinine is < 1.2 mg/dl, then a nuclear renal scan is required to determine glomerular filtration rate. Serum creatinine consistently > 1.5 mg/dL Macroalbuminuria (urinary albumin excretion rate > 300 mg/24h) \square Baseline Hb < 12 gm/dL in women or < 13 gm/dL in men Baseline liver function tests (LFT) outside of normal range (An initial LFT test panel with any values > 1.5 times normal upper limits will exclude a member without a re-test. A re-test for any values between normal and 1.5 times normal should be made, and if the values remain elevated above normal limits, the member will be excluded.) Untreated proliferative retinopathy Positive pregnancy test, intent for future pregnancy, or male members' intent to procreate, unwilling to follow effective contraceptive measures, or presently breast-feeding Insulin requirement > 0.7 IU/kg/day \square HbA1c > 12% Hyperlipidemia (fasting LDL cholesterol > 130 mg/dL, treated or untreated; and/or fasting triglycerides > 200 mg/dL) Under treatment for a medical condition requiring chronic use of steroids other than a previous organ transplant Use of coumadin or other antiplatelet or anticoagulant therapy, or member with PT INR > 1.5. Low dose aspirin is allowed after transplantation. History of Factor V deficiency Addison's disease Allergy to radiographic contrast material Symptomatic cholecystolithiasis Acute or chronic pancreatitis Symptomatic peptic ulcer disease Severe unremitting diarrhea, vomiting, or other gastrointestinal disorders that could interfere with the ability to absorb oral Received live attenuated vaccine(s) within 2 months Will the therapy be used conjunction with concomitant immunosuppression? \(\subseteq \text{Yes} \) No REAUTHORIZATION Has the member achieved independence from exogenous insulin? Yes No Date of last infusion: Date member lost independence from exogenous insulin: How many infusions has the member had? SUPPORTING INFORMATION or CLINICAL RATIONALE **Prescribing Provider Signature** Date