Hydroxyprogesterone caproate injection (Makena®, generics)

While Makena and generic hydroxyprogesterone are commercially manufactured drugs, hydroxyprogesterone caproate may be compounded for injection through one of the pharmacies listed in Section 5.

Place of Service
Home Infusion Administration
Office Administration
Outpatient Facility Infusion

Administration
Specialty Pharmacy

Use HCPC: Makena: J1726 per 10mg

Hydroxyprogesterone caproate, (generic) not otherwise specified: J1729 per 10mg

Use code **\$9560** for 17-P services provided by **Alere** Women's and Children's Health only (formerly Matria Healthcare)

Self-Administration

May be covered under pharmacy benefit (Makena)

Condition(s) listed in policy (see criteria for details)

- Prevention of preterm labor in women who are at high risk (Makena)
- Advanced adenocarcinoma of the uterine corpus (Stage III or IV) (generic)
- Amenorrhea (primary and secondary) (generic)
- Abnormal uterine bleeding due to hormonal imbalance in the absence of organic pathology (e.g. submucous fibroids or uterine cancer) (generic)
- <u>Test for endogenous estrogen production (generic)</u>
- Production of secretory endometrium and desquamation (generic)

AHFS therapeutic class: Progestins

Mechanism of action: Hydroxyprogesterone caproate, also known as 17-P, is an esterified derivative of the naturally occurring 17α -hydroxyprogesterone which has substantial progestational activity and prolonged duration of action.

(1) Special Instructions and Pertinent Information

Makena:

- If member has a Prescription Benefit, please refer cases to Pharmacy Services for prior authorization.
- If covered under the Medical Benefit, please submit clinical information for prior authorization review via fax. Please include medical rationale why medication cannot be home self-administered.

Generic hydroxyprogesterone caproate injection:

 Covered under the Medical Benefit, please submit clinical information for prior authorization review via fax.

Makena and hydroxyprogesterone caproate may be obtained and billed by a specialty

Hydroxyprogesterone caproate (Makena®, generic)

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pharmacy.

** CRITERIA FOR HOSPITAL OUTPATIENT FACILITY ADMINISTRATION**

AAAAI Guidelines 2011, MCG™ Care Guidelines, 19th edition, 2015, Makena PI, 4/16

Members with the following benefits: **PPO, Direct Contract HMO, and when applicable, ASO/Shared Advantage**, may be required to have their medication administered at a preferred site of service, including the home, a physician's office, or an independent infusion center not associated with a hospital.

For members that cannot receive infusions in the preferred home or ambulatory setting AND meet one of the following criteria points, drug administration may be performed at a hospital outpatient facility infusion center.

ADMINISTRATION OF HYDROXYPROGESTERONE CAPROATE INJECTIONS IN THE HOSPITAL OUTPATIENT FACILITY SITE OF CARE REQUIRES ONE OF THE FOLLOWING: (Supporting Documentation must be submitted to support the need for additional clinical monitoring)

1. Patient is receiving their first dose or is being re-initiated after at least 6 months off therapy. Subsequent doses will require medical necessity for continued use in the hospital outpatient facility site of care.

O

Additional clinical monitoring is required during administration as evidenced by one of the following:

- 2. Patient has experienced <u>a previous severe adverse event</u> on the medication based on documentation submitted.
- 3. Patient <u>continues to experience moderate to severe adverse events</u> on the medication based on documentation submitted, despite receiving premedication such as acetaminophen, steroids, diphenhydramine, fluids, etc.
- 4. Patient is clinically unstable based on documentation submitted.
- 5. Patient is physically or cognitively unstable based on documentation submitted.

(2) Prior Authorization/Medical Review is required for the following condition(s)
All requests for hydroxyprogesterone caproate injection NOT LISTED in section 3 must be sent for clinical review and receive authorization prior to drug administration or claim payment.

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<u>Prevention of preterm labor in women who are at high risk (i.e. prior history of preterm labor) -</u> *Makena*

- Must be a singleton pregnancy, AND
- Treatment with hydroxyprogesterone caproate is being initiated within 16 weeks, 0 days and 24 weeks, 6 days of gestation, AND
- Patient has a history of prior preterm singleton delivery before 37 weeks gestation.

Covered Doses

Makena vial: Up to 250 mg IM once weekly

Makena auto-injector: Up to 275 mg SC once weekly

Coverage Period

Cover for up to 36 weeks, 6 days gestation

ICD-10: O20.0, O60.00, O60.02, O60.03

Abnormal uterine bleeding due to hormonal imbalance in the absence of organic pathology (e.g. submucous fibroids or uterine cancer) – generic hydroxyprogesterone caproate injection

Patient has had an inadequate response, intolerance or contraindication to an oral progestin

Covered Doses

375mg IM x1

For cyclic therapy with estradiol valerate injection: 250mg IM on day 15 of 28-day cycles for up to 4 cycles

Coverage Period

One time dose as monotherapy

For cyclic therapy with estradiol valerate injection: up to four 28-day cycles

ICD-10: N93.8

Advanced adenocarcinoma of the uterine corpus (Stage III or IV) – generic hydroxyprogesterone caproate injection

 Patient has had an inadequate response, intolerance or contraindication to medroxyprogesterone (NCCN category 2A-recommended treatment)

Covered Doses

Up to 7g per week

Coverage Period

Indefinite (if patient had clinical benefit)

ICD-10: C54.1-C54.3, C54.9

Amenorrhea (primary and secondary) - generic hydroxyprogesterone caproate injection

Patient has had an inadequate response, intolerance or contraindication to an oral progestin

Covered Doses

Hydroxyprogesterone caproate (Makena®, generic)

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375mg IM x1

For cyclic therapy with estradiol valerate injection: 250mg IM on day 15 of 28-day cycles for up to 4 cycles

Coverage Period

One time dose as monotherapy

For cyclic therapy with estradiol valerate injection: up to four 28-day cycles

ICD-10: N91.0, N91.1, N91.2

(3) The following condition(s) <u>DO NOT</u> require Prior Authorization/Preservice

All requests for hydroxyprogesterone caproate injection NOT LISTED in section 3 must be sent for clinical review and receive authorization prior to drug administration or claim payment.

<u>Production of secretory endometrium and desquamation – generic hydroxyprogesterone caproate injection</u>

Covered dose:

375mg x1

For cyclic therapy with estradiol valerate injection: 250mg on day 15 of 28-day cycles for up to six months

ICD-10: N85.8

Test for endogenous estrogen production – generic hydroxyprogesterone caproate injection

Covered dose:

250mg x 2 IM injections; separated by 4 weeks

ICD-10 includes but is not limited to: E22.8

(4) This Medication is NOT medically necessary for the following condition(s)

<u>Blue Shield's research indicates there is inadequate clinical evidence to support off-label use of this</u> drug for the following conditions (Health and Safety Code 1367.21):

- Prevention of preterm labor in women pregnant with multiple gestations
- Prevention of preterm labor in pregnant women with a short cervical length and no prior history of preterm birth

Coverage for a Non-FDA approved indication, requires that criteria outlined in Health and Safety Code § 1367.21, including objective evidence of efficacy and safety are met for the proposed indication.

<u>Please refer to the Provider Manual and User Guide for more information.</u>

(5) Additional Information

How supplied: Makena vial: 250mg (single dose) 1250mg (multi-dose)

Makena auto-injector:

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275 mg/1.1 mL (single-use)

Hydroxyprogesterone caproate (generic): 1250mg (multi-dose vial)

The chemical name for Makena and hydroxyprogesterone caproate is pregn-4-ene-3,20-dione, 17[(1-oxohexyl) oxy], and is also known as 17 alpha-hydroxyprogesterone caproate or 17P.

Hydroxyprogesterone is commercially available but may also be prepared by specialty pharmacies. Below is a list of some specialty pharmacies which may be supplying this product:

- Alere Women's and Children's Health (formerly Matria Healthcare)
- Wedgewood Pharmacy
- Vitality
- Freedom Pharmacy

(6) References

- AHFS®. Available by subscription at http://www.lexi.com
- American College of Obstetricians and Gynecologists. Prediction and Prevention of Preterm Birth. Practice Bulletin Number 130, October 2012.
- DrugDex®. Available by subscription at http://www.micromedexsolutions.com/home/dispatch
- Hydroxyprogesterone caproate injection prescribing information. McGuff Pharmaceuticals, Inc. August 2015.
- Makena® prescribing information. AMAG pharmaceuticals. 2017.
- Meis PJ, et al. Prevention of recurrent preterm delivery by 17 alpha hydroxyprogesterone caproate. N Engl J Med 2003; 348: 2379-2385.
- Northern A, et al. Follow-up of children exposed in utero to 17-hydroxyprogesterone caproate compared with placebo. American College of Obstetricians and Gynecologists. Vol 110, No 4, October 2007.
- Rouse DJ, et al. A trial of 17 alpha-hydroxyprogesterone caproate to prevent prematurity in twins. N Eng J Med 2007; 357: 454-461.
- Use of Progesterone to Reduce Preterm Birth. The American College of Obstetricians and Gynecologists. Volume 112. no. 4 October 2008.

(7) Policy Update

Date of last revision: 3O2018 Date of next review: 3Q2019

Changes from previous policy version:

No clinical change to policy following annual routine review

BSC Drug Coverage Criteria to Determine Medical Necessity Reviewed by P&T Committee

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