MAKENA® (hydroxyprogesterone caproate injection)

LENGTH OF AUTHORIZATION: UP TO 5 MONTHS

CLINICAL NOTES:
Makena is a progestin indicated to reduce the risk of preterm birth in women with a singleton pregnancy (a pregnancy with only one baby) who have a history of singleton spontaneous preterm birth. Makena is not intended for use in women with multiple gestations or other risk factors for preterm birth.

Florida Medicaid covers prescribed drugs that combine two or more drugs when all of the following are met:
- At least one of the drugs in the compound drug is covered by Florida Medicaid.
- The compounded drug is not otherwise commercially available.
- The drug is compounded as prescribed for the recipient to treat his or her specific condition. (Prescribed Drug Services Coverage Policy, 2017)

REVIEW CRITERIA:
- Must be ≥ 16 years of age.
- Must be currently pregnant with only one baby (a singleton pregnancy) at 16 to 37 weeks gestation confirmed by supporting documentation or diagnosis code(s).
- Documentation of having a history of singleton spontaneous preterm birth must be submitted.
- Makena is not intended for use in women with multiple gestations or other risk factors for preterm birth.

DOSING AND ADMINISTRATION:
- Administer once weekly.
- Begin treatment between 16 weeks, 0 days and 20 weeks, 6 days of gestation.
- Continue administration once weekly until week 37 (through 36 weeks, 6 days) of gestation or delivery, whichever occurs first.

Dosage Forms:
- Available as Makena auto-injector 275mg administered subcutaneously once weekly.
- Available as Makena (single or multi-dose vials) 250mg administered intramuscularly once weekly.