Abarelix (Plenaxis™)

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**Drug name:** Abarelix is marketed under the trade name Plenaxis™ (Praecis Pharmaceuticals Inc., Waltham, MA).

**Classification:** Gonadotropin-releasing hormone (GNRH) antagonist.

**Action:** GNRH, also known as leutinizing hormone-releasing hormone (LHRH), binds to the anterior pituitary gonadotropic cells and stimulates the secretion of leutinizing hormone (LH) and follicle stimulating hormone (FSH). Once LH and FSH enter the circulatory system, they act on the testes, causing them to produce testosterone and dihydrotestosterone (DHT), both of which are considered androgens. LH and FSH are needed for the prostate gland to function normally and also have a role in the sustenance of malignant prostate tissue. Abarelix inhibits GNRH from binding to gonadotropic cells, thus preventing the secretion of LH and FSH. The cascade of hormone secretion is interrupted at a very early stage, preventing the production of significant levels of the male hormones testosterone and DHT, which are known to lead to the proliferation of androgen-dependent and androgen-sensitive prostate cancer cells. Unlike currently available LHRH agonists, abarelix does not induce an initial surge of testosterone or other known androgens.

**Indications:** Abarelix currently is indicated for the palliative treatment of men with advanced symptomatic prostate cancer in whom LHRH agonist therapy is not appropriate or who refuse surgical castration and meet at least one of the following criteria.

- Risk of neurologic complications resulting from metastatic disease
- Obstruction of the ureters or outlet of the urinary bladder because of localized encroachment or metastatic disease
- High levels of bone pain from skeletal metastases despite ongoing use of narcotic analgesia

As with the administration of any hormonal agent for the treatment of prostate cancer, abarelix is not intended to be curative. It can, however, provide patients with symptomatic relief from the complications listed previously and palliation.

**Metabolism:** Studies have indicated that abarelix is metabolized in the liver without evidence of cytochrome p 450 involvement.

**Excretion:** No detectable metabolites were found in the urine. Approximately 13% of unmetabolized abarelix was found in urine after a 15 mcg/kg intramuscular (IM) injection. Renal clearance of abarelix was 14.4 L per day (10 ml per minute) following administration of 100 mg of the drug.

**Absorption:** After IM administration of 100 mg of abarelix, absorption peaked with a concentration level of 43.4 ng/ml observed approximately three days following injection.

**Effect on blood counts:** No evidence suggests that abarelix has a significant effect on bone marrow production.

**Adverse reactions and events:** The most significant adverse event associated with the administration of abarelix is an immediate onset systemic allergic reaction most often occurring within eight minutes of administration. The exact nature of the reported allergic reactions remains unclear. Phase IV clinical studies are under way to make this determination. The incidence rate of this type of severe allergic reaction was reported in 3.7% of patients with advanced symptomatic prostate cancer and 1.1% of all patients with prostate cancer. These immediate onset reactions have been reported to occur following any administration of abarelix, including the initial dose. The cumulative risk of such a reaction increases with the duration of treatment. Other adverse reactions to abarelix that have occurred with increased frequency include hot flashes (79%), sleep disturbance (44%), pain (31%), breast enlargement (30%), breast or nipple tenderness (20%), back pain (17%), constipation (15%), peripheral edema (15%), dizziness (12%), headache (12%), upper respiratory infection (URI) (12%), and diarrhea (11%). Approximately 20% of patients who received abarelix in clinical trials experienced prolonged QT interval changes on electrocardiograms. Whether this was a direct result of drug administration, androgen deprivation therapy, or other unknown variables has not been determined.

**Route and dosage:** The recommended dosage of abarelix is 100 mg administered as an IM injection into the buttock on days 1, 15, 29, and every four weeks thereafter. Because of the possibility of a localized skin reaction, injection sites in the right and left buttocks should be alternated with each subsequent injection.

**Dilution and reconstitution:** Abarelix is supplied as a single-dose, preservative-free, sterile powder that, when reconstituted with

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