Relugolix: New Oral Gonadotropin-Releasing Hormone (GnRH) Antagonist

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ABSTRACT

With the recent development of Gonadotropin-Releasing Hormone (GnRH) analogs, physicians have manipulated gonadotropin secretion and, consequently, to treat sex steroid-dependent hormonal disorders such as central precocious puberty in children, and benign prostatic hypertrophy and prostate cancer in men, and endometriosis, leiomyoma and breast cancer in women. GnRH agonists can cause transient deterioration of clinical symptoms associated with an initial surge of gonadotropins and gonadal hormones. However, the peptide antagonists must be administered by frequent subcutaneous injection or sustained release formulation due to orally fragile bioavailability. A new oral nonpeptide GnRH antagonist, relugolix, promptly lowers pituitary gonadotropin by GnRH receptor competition lowering estrogen level in women and testosterone in men within a few days without the symptom flare-up observed with GnRH agonist. Sex steroid levels to baseline and menses resumed more quickly after the oral GnRH antagonist withdrawal. The oral GnRH antagonist may play an ideal role in therapeutic management of the estrogen or androgen-dependent disorders.

Introduction

The decapeptide hormone, Gonadotropin-Releasing Hormone (GnRH), induces gonadotropins (luteinizing hormone and follicle-stimulating hormone) secretion from the anterior pituitary in an orderly way which is important for the control of gonadal function and normal ovarian cyclicity. The action of GnRH is mediated by its binding to high affinity membrane receptors belonging to the seven-transmembrane receptor family. It is noted that sustained occupancy of the receptor with GnRH itself or a GnRH agonist caused desensitization. The GnRH agonist-induced down-regulation has been applied to manipulate gonadotropin secretion and, consequently, to treat sex steroid (estrogen or androgen)-dependent hormonal disorders such as central precocious puberty in children, and benign prostatic hypertrophy and prostate cancer in men, and endometriosis, leiomyoma and breast cancer in women [1-4].

However, GnRH agonists can cause transient worse clinical symptoms associated with an initial surge of gonadotropins and gonadal hormones [1-4]. GnRH antagonists, in contrast to GnRH agonists, achieve an immediate reduction of these hormones and thus fast onset of therapeutic effects without undesirable adverse events induced by the flare-up. To date, several peptide GnRH antagonists have been developed, including cetrorelix [5,6] and ganirelix for in vitro fertilization [7,8], and degarelix for advanced prostate cancer [9,10]. These peptide antagonists, however, had to be administered by frequent subcutaneous injection or sustained release formulation due to orally fragile bioavailability.

Many nonpeptide GnRH antagonists have been developed by several research groups. Relugolix (Relumina®), an oral nonpeptide GnRH receptor antagonist, promptly suppresses pituitary gonadotropin by GnRH receptor competition, lowering sex steroids levels within a few days, without the symptom flare-up associated with GnRH agonist [11-13]. More recently, Osuga et al. [14] demonstrated the noninferiority of relugolix compared with current GnRH analogs in reducing heavy menstrual bleeding associated with uterine leiomyoma. This novel oral GnRH antagonist showed a rapid effect on the symptoms and reduced the leiomyoma and uterine volumes. Depot preparation of GnRH agonists also reduce leiomyoma size but have a delayed onset and must be given by monthly injection. Once daily oral intake achieved a similar proportion of heavy menstrual bleeding responders at 6-12 weeks.
MacLean et al. [15] investigated relugolix effect on serum testosterone level in healthy volunteers as a possible prostate cancer treatment. The oral GnRH antagonist produced sustained reduction in testosterone level to below castrate level. Testosterone flare-up did not occur. The GnRH antagonist are an ideal option for testosterone suppression in benign prostatic hypertrophy and prostate cancer. The attractive advantages of orally available GnRH antagonists over peptide antagonists include the rapid recovery of gonadal function after drug withdrawal, since peptide antagonists are usually administered by subcutaneous injection in sustained release preparation and are unable to be discontinued during the release period even when undesirable adverse events occur.

Discontinuation after 28-day oral administration resulted in recovery of female reproductive function from a completely suppressed state in approximately 5 days, followed by continuous estrous phase for several days. The oral GnRH antagonist may play an ideal role in therapeutic management of the estrogen or androgen-dependent disorders; e.g. endometriosis, uterine myomas, hirsutism, dysfunctional uterine bleeding, premenstrual syndrome, precocious puberty, assisted reproduction, and some hormone-dependent tumors.

**Disclosure Statement**

The authors declare no conflict of interest.

**References**