Effect of Osaterone Acetate Administration on Prostatic Regression Rate, Peripheral Blood Hormone Levels and Semen Quality in Dogs with Benign Prostatic Hypertrophy

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(Received 15 March 2000/Accepted 25 December 2000)

ABSTRACT. The effects of osaterone acetate (OSA), which is an anti-androgen agent being developed as a therapeutic drug for benign prostatic hypertrophy (BPH) in dogs, on the degree of prostatic regression and semen qualities were investigated. Prostatic regression was compared between dogs with and without orchidectomy. Five male beagles aged 5–9 years were used in the experiment. OSA was orally administered at doses of 0.2 mg/kg and 0.5 mg/kg for one week. The prostatic regression rate one week after the end of administration was 62.6% on average. In the orchidectomized group, the mean regression rate one week after orchidectomy was 60.1%. However, the prostate became enlarged 6 months after administration, compared to the size prior to administration. The above findings suggested that OSA is clinically applicable as a therapeutic drug for BPH in dogs, and inhibits prostatic hypertrophy during the early phase.

KEY WORDS: canine BPH, osaterone acetate, prostatic volume.
tion, orchidectomy was performed on 4 dogs in which the PV had returned to the same size as that prior to administration. Orchidectomy was performed under general anesthesia. As a pretreatment, 0.1 mg/kg of atropine sulfate was subcutaneously administered followed by intravenous administration of 16–20 mg/kg thiopental sodium (Ravonal®) 20 min later, and anesthesia was maintained with isoflurane.

**Measurement of PV:** PV was measured prior to initiation of OSA administration and four days, one and two weeks, and one month after the administration, then every one month thereafter until 6–7 months. In the orchidectomized group, the PV was measured by the same schedule until two months after orchidectomy. The CT was performed under the same general anesthesia used for the orchidectomy.

**Measurements of peripheral blood LH and testosterone (T) levels:** Blood samples for measurements of peripheral blood LH and T levels after OSA administration were collected prior to (three days before and on the administration day) and every week after initiation of administration. Blood was sampled three times per day (10:00, 12:00, 14:00). LH and T levels were measured three times per day, and the mean value was regarded as the value of that day. In blood sampling, heparin sodium, an anticoagulant, was used. Three milliters of blood were collected at each sampling from the cephalic vein, and immediately centrifuged at 600 G for 15 min using a cold centrifuge (KR-20000T, KUBOTA Co., Japan). The plasma was stored at –30°C until hormone measurement. Plasma LH and T levels were measured by the RIA method we previously reported [5].

**Semen quality:** Semen quality was examined on the same days as blood sampling. Semen was collected in three fractions by the previously reported [8] procedure. In the semen quality, semen volume, sperm motility, sperm count, sperm viability, sperm abnormality and semen pH were examined.

**Statistical analysis:** The results of this experiment were analyzed by Student’s t-test. A p value less than 0.05 was regarded as significant.

**PV:** Regarding the volume prior to administration as 100%, the changes in PV in the OSA treatment groups are shown individually in the 0.2 mg/kg group, and presented as the means ± SE in the 0.5 mg/kg and orchidectomy groups in Fig. 1.

In the 2 animals in the OSA 0.2 mg/kg treatment group, the PV regressed to 77.7% and 63.1% one week after administration, respectively. The maximal regression rate was observed at two weeks. As in the 0.2 mg/kg group, the maximal regression rate was attained two weeks after administration in the 0.5 mg/kg group. After the maximal regression rate was attained, PV gradually increased in both treatment groups, and the prostate had recovered to almost the original size 6 months after administration.

PV rapidly decreased after orchidectomy, and regressed to 34.8% on average after two months. PV was clearly smaller in the castrated group than in the 0.5 mg/kg OSA-treated group, however, no difference was observed in the percent decrease between PV one week after the end of OSA administration and that one week after castration.

**Peripheral blood LH and T levels:** Because there were no differences in either the peripheral blood LH or T level after OSA administration between the two treatment groups, the levels are presented as the means ± SE of 3 animals in the 0.5 mg/kg group in Fig. 2.

In the OSA 0.5 mg/kg group, the LH level was above 5 ng/ml on many days 1–2 months after administration. However, the level was maintained at a level of 5 ng/ml or lower during the period 3–5 months after the end of administration.

The T level prior to OSA administration was 2.3–3.8 ng/ml (3.1 ± 0.3 ng/ml). The peripheral blood T level during the three months after OSA administration was maintained at 2 ng/ml or lower, showing a clearly decreased level com-
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pared to the level prior to administration (p<0.05, p<0.01). However, the level slightly increased thereafter, remaining at 2–3 ng/ml.

Semen quality: Because there were no differences in the semen quality after OSA administration between the two treatment groups, the semen quality is presented as the means ± SE of three animals in the 0.5 mg/kg group in Fig. 3.

The semen volume was maintained at about 10 ml for four months after administration, then increased to 15–20 ml. The sperm count varied greatly among days of semen collection, but was within a range of 2–4 × 10⁶. The sperm abnormality showed a high level of about 10% from four weeks after administration for approximately 1.5 months, after which it clearly decreased and was maintained at 5% or lower. The transiently increased abnormality was observed in the tail region.

No changes were observed in the other semen qualities such as sperm motility and viability and semen pH.

One-week oral administration of OSA at a dose of 0.2 or
0.5 mg/kg to dogs with prostatic hypertrophy showed a marked effect of 62.6% prostatic mean regression rate one week after the final administration day. In the orchidectomized group, the prostatic mean regression rate one week after orchidectomy was 60.1%, showing that prostatic regression caused by OSA administration occurred soon and rapidly thereafter. Therefore, for the mechanism of PV regression induced by OSA administration during the early phase, Takezawa et al. [13] reported that OSA directly decreased the dihydrotestosterone and androgen receptor contents in the prostate.

The period after OSA administration until the prostate was enlarged to the same size as that prior to administration was six months. These findings suggest that it is desirable to concurrently use a CMA implant with oral OSA administration to inhibit prostatic hypertrophy for a prolonged period in BPH dogs. Tsumagari et al. [14] reported that a one year or longer effect can be expected with oral OSA administration, but they did not describe the size of the prostate. It is necessary to investigate whether the clinical symptoms when the prostatic size returned to the original size are the same as those prior to administration.

For several months after OSA administration, the peripheral blood T level was maintained at a level clearly lower than that prior to administration. However, the peripheral blood LH level did not change during this period. This finding suggested that due to OSA administration, the testicular secretion of T directly inhibited the LH level, not via the pituitary hormone.

Regarding the semen quality, OSA administration transiently increased sperm abnormality. The peripheral blood T level decreased during this period. This reduction in the T level changed the secretory discharge in the epididymis, which may have increased the incidence of abnormality in the sperm tail region.

In conclusion, although oral OSA administration transiently decreased the peripheral blood T level, it did not markedly affect either the LH level or semen quality, suggesting that OSA is clinically applicable as a therapeutic drug for BPH in dogs.

REFERENCES