Effect of Cernitin Pollen-Extract on Experimental Nonbacterial Prostatitis in Rats

Toshiyuki Kamijo,1 Shigeru Sato,2 and Tadaichi Kitamura1
1Department of Urology, Faculty of Medicine, University of Tokyo, Japan
2Ohme Research Laboratories, Tobishi Pharmaceutical Co., Ltd., Tokyo, Japan

BACKGROUND. The treatment for chronic nonbacterial prostatitis (NBP) has not been established. Cernitin pollen-extract (CN-009) is reported to have therapeutic effects for NBP. The effects and mechanisms of CN-009 were investigated.

METHODS. Ten-month-old rats were used with administration of estradiol after castration, which were similar to human NBP histologically. Since CN-009 consists of T-60 and GBX, these drugs were administered, respectively. The prostate was evaluated histopathologically including glandular damage (epithelial score), stromal ratio and immunohistochemical assays for epithelial function (PAP), stromal evaluation (Vimentin), cell proliferation (PCNA) and apoptosis (deoxyuridine triphosphate biotin nick end-labeling (TUNEL)).

RESULTS. Controls revealed severe acinar gland atrophy and stromal proliferation. CN-009 showed diminished these damages. Epithelial score was better (P<0.01) and PAP positive materials were more abundant in CN-009 and GBX than in Controls. The stromal ratio was lower in CN-009 (P<0.01) and T-60 (P<0.05). There was no difference for PCNA positive cells in the epithelium and stroma, and TUNEL positive cells in epithelium. While, the number of TUNEL positive cells in the stroma of CN-009 and T-60 increase (P<0.01).

CONCLUSIONS. These findings suggest than CN-009 protects acinar epithelial cells mainly by GBX and also inhibits stromal proliferation in association with enhanced apoptosis mainly by T-60. Prostate 49: 122-131, 2001. © 2001 Wiley-Liss, Inc.

KEYWORDS: cernitin pollen-extract; apoptosis; chronic prostatitis; sex-hormone induced prostatitis

INTRODUCTION

The chronic prostatitis syndromes have been recognized; chronic bacterial prostatitis (CBP), chronic nonbacterial prostatitis (NBP) and prostatodynia. NBP is the most frequent disorder of 64% in these three diseases [1]. The etiology of NBP is unknown. A number of organisms or other factors have been reported to be the possible causes for NBP. They are Trichomonas vaginalis, Chlamydia trachomatis, genital mycoplasmas, staphylococci, coryneforms, genital viruses [2], biofilms [3], stagnation of prostatic secretion, autoimmune disease, allergy, disorder of sex hormone and psychological effects [4,5]. For the treatment of CBP or NBP, antibiotics of new-quinolone or tetracycline have been administered. However, many cases resist these treatments [6]. CN-009 is a pollen extract, which contains 20:1 ratio of powdered aqueous and organic extract. It is essentially a microbial digest of a standardized mixture of eight plant species grown at the Scania area in southern Sweden. The active ingredients consist of water-soluble (T-60) and fat-soluble (GBX) fractions [7,8]. It was reported that CN-009 showed urine discharge action [9,10], anti-prostatic hypertrophic action [7] and anti-inflammatory effects to the prostate [11] in a preliminary study. Since Ask-Upmark [12] reported CN-009 showed an efficacy to prostatitis, it has been used for the treatment of chronic prostatitis with...
high therapeutic effects. However, the mechanisms for these effects remain unknown.

To assess the mechanisms of the anti-prostatitis effect by CN-009, the present study was performed using a nonbacterial prostatitis rat model [13,14] induced by 17β-estradiol administration and castration.

**MATERIALS AND METHODS**

**Sex Hormone-Induced Nonbacterial Prostatitis Model**

Ten month-old Wistar aged male rats were purchased from Japan Slc Co. (Tokyo, Japan). The rats were housed in a climatised environment with a 12-hr light/ dark cycle, 40-70% humidity. Food and water were supplied ad libitum. The rats were castrated under ether anesthesia, and then 17β-estradiol (Sigma, MI) 0.25 mg/ 2 ml/ kg diluted by sesame oil, as an inducer for prostatitis, was subcutaneously injected into the back of rats for 30 days from 1 day after castration [13,14].

**Experimental Structure and Schedule**

CN-009 was suspended for 630 or 1,260 mg/ 5 ml with 1% HCO-60 (Japan Surfactant, Tokyo, Japan). T-60 and GBX were similarly prepared for 1,200 and 60 mg/ 5 ml, respectively. Testosterone (TS) (Wako Chemicals, Tokyo, Japan), as a positive control, was diluted for 2.5 mg/ 2 ml with corn oil (Yuro Chemical, Tokyo, Japan).

The experimental structure is shown in Table I and the experimental schedule is illustrated in Figure 1. The rats were divided into seven groups consisting of Sham-operation (Sham-ope), Control, CN-009 630, CN-009 1260, T-60, GBX and TS with five or six animals in each group.

In the Sham-ope group, the rats were treated with only Sham-castration and without any drugs. In the Control group, the rats were injected subcutaneously with 17 β-estradiol for 30 days from the day following castration and administered orally with only 1% HCO-60 5 ml/ kg for 14 days from Day 17. In the CN-009 630, CN-009 1260, T-60 and GBX groups, similar protocols were performed with oral administration of CN-009 630, CN-009 1260, T-60 1200 and GBX 60 mg/ kg, respectively. Also in the TS group, the rats were injected subcutaneously with 17 β-estradiol for 30 days from the next day of castration. After 14 days, TS 2.5 mg/ kg was injected subcutaneously for 14 days. All studies were conducted in accordance with institutional guidelines of animal care and in accordance with Japanese Government Animal Protection and Management Law.

**Prostate Weights and Histopathological Evaluation**

The rats were sacrificed on the day following the final administration. The prostate was extirpated and weighed. Relative prostatic weight was calculated from body weight and absolute weight.

After fixation in 10% neutral buffered formalin, each prostate was cut into coronal blocks. The tissue samples were dehydrated and embedded in paraffin. Sections (3-4 μm thickness) were stained with Hematoxyline-Eosin (HE), Periodic acid Shiff (PAS) and Masson’s tri-chrome. The specimens were evaluated histopathologically.

**Immunohistochemistry**

Immunohistochemistry studies were performed with anti-prostatic acid phosphatase (PAP), and Viementin. PAP staining was performed for the evaluation of glandular epithelial function. In
PAP stained specimens, anti-PAP polyclonal antibody (Chemicon International, New York, NY) was diluted by PBS including 0.1% BSA of a 1:100 ratio, and incubated for 2 hr at 37°C. Biotinylated anti-rabbit IgG and the avidin-biotin peroxidase complex (ABC) method was performed. Unitect rabbit immunohistochemistry detection systems (Oncogene Science, New York, NY) were reacted by those methods. Vimentin staining was performed for the evaluation of stromal proliferation. An ImmunoCruz staining system (Santa Cruz BioTech., Santa Cruz, CA) for Vimentin staining was used according to the manufacturer's instructions.

Cell Proliferation and Apoptosis

Cell proliferation and apoptosis were investigated with proliferating cell nuclear antigen (PCNA) and terminal deoxynucleotidyl tranferase mediated deoxyuridine triphosphate biotin nick end-labeling (TUNEL), respectively. PCNA staining was performed with PCNA staining kit (ZYMED Laboratories, South San Francisco, CA). TUNEL method was performed with ApoTag Peroxidase In Situ Apoptosis Detection kit (Intervene, New York, NY). In PCNA and TUNEL specimen, 5,000 cells were counted under a microscope in glandular epithelial cells and stromal cells, respectively.

Acinar Epithelial Score and Stromal Area Ratio

To evaluate glandular damage, acinar epithelial cells were classified and scored, as follows: columnar (2 points), cuboidal (1 point), squamous-like (0 point) shape. Three different pathologists without any information judged the score. Using this scoring evaluation, 20 acinar glands of each specimen were investigated. To assess stromal proliferation, all areas of the specimen and the glandular area were measured using a digitizer (Graph Tech, Tokyo, Japan) with photomicrographs. Using these findings, the stromal ratio was calculated.

Statistical Analysis

All experiments were repeated at least twice. Each value was demonstrated as the mean±SD. Dunnett's test if in equal variance, or non-parametric Dunnett's test if in unequal variance between treatment groups and Control group was performed after 1-way ANOVA followed by Bartlett variance analysis test. Mann-Whitney U test was performed between the Sham-ope and Control groups.

RESULTS

Body and Prostate Weights (Fig. 2)

There was no significant difference in body weight among the CN-009 630, CN-009 1260, T-60, GBX and TS groups compared with the Control group. Absolute and relative prostate weights were significantly (P<0.01) decreased in the Control group compared with the Sham-ope group (Fig. 2) in the CN-009 630, CN-009 1260, T-60 and GBX 60 groups, there was no difference compared with the Control group. In the TS group, absolute and relative prostate weights were very close to the Sham-ope group and were significantly different (P<0.01) from other groups.

Histopathology and Immunohistochemistry (PAP and Vimentin Staining)

In the Sham-ope group, the prostate was larger than in other groups. Acinar glands were roundly shaped. The acinar lumen was filled with eosinophilic materials. Acinar epithelial cells were cylindrical with a normally situated nucleus and the supranuclear spaces of these cells
Effect of Cernitin Pollen-Extract on Experimental Nonbacterial Prostatitis in Rats

contained secretory materials, which were strongly stained with PAP antibody. A few fibrous tissues were found in the stroma (Figs. 3A and 4A). Vimentin positive cells were few, and the Vimentin positive area was small (data not shown).

In the control group, the prostate was atrophic. Acinar glands were irregularly shaped. The acinar lumen was poor with pale stained eosinophilic materials and filled with inflammatory cell infiltrations mainly characterized by neutrophils. Acinar epithelial cells were flattened similar to a squamous cell. A few secretory materials in the epithelial cells were poorly reacted with PAP antibody. The stroma showed severe proliferation with many lymphocyte and monocyte infiltrations and marked fibrosis with fibroblasts (Figs. 3B and 4B). The stroma was stained very strongly with Vimentin. The Vimentin positive area was significantly increased (data not shown). In the CN-009 630 group, the findings were basically identical with the Control group (data not shown).

In the CN-009 1260 group, acinar glands were more roundly shaped than in the Control group. Acinar epithelial cells were cuboidal, and the supranuclear spaces contained secretory materials stained with anti-PAP that were much more abundant than in the Control group. Inflammatory cell infiltrations into the acinar lumen were diminished. The stroma showed mild proliferation with a few lymphocytes, monocytes and mild fibrosis with fibroblasts (Figs. 3C and 4C). The Vimentin positive area was much less than that of the Control group (data not shown).

In the T-60 group, acinar epithelial cells were more roundly shaped than in the Control group. Although inflammatory cell infiltrations into the lumen were found, stromal cell infiltrations (Fig. 3D), the Vimentin positive cells were also less than that of the Control group (data not shown).

In the GBX group, acinar epithelial cells were more cuboidal than in the Control group. Epithelial cells contained secretory materials stained with anti-PAP, which was basically identical with the CN-009 1260 group. Diminished cell infiltration into the lumen was found (Fig. 3E). However, the stroma showed a proliferative condition with many lymphocyte and monocyte infiltrations and marked fibrosis with many fibroblasts. The stroma was stained strongly with Vimentin, and the positive area was markedly increased (data not shown).

In the TS group, acinar glands were roundly shaped. The acinar lumen was filled with eosinophilic materials with a few cell infiltrations. Acinar epithelial cells were cylindrical and the supranuclear spaces contained many secretory materials with reactive anti-PAP. However, the stroma was stained strongly with Vimentin and showed mild proliferation with fibroblasts (data not shown).
Cell Proliferation and Apoptosis (PCNA and TUNEL Positive Cell Counts (Fig. 5))

No significant differences among the groups were observed in the PCNA positive cell counts in epithelial cells (Fig. 6) or in stromal cells (Fig. 7). In the Sham-ope group, a few TUNEL positive cells were found (Fig. 5A). The findings of the Control group were basically identical with the Sham-ope group (Fig. 5B). In the CN-009 1260 group, TUNEL positive cells in the stroma were more abundant than in the Sham-ope and Control groups (Fig. 5C). In TUNEL positive cell counts, no significant differences were observed in acinar epithelial cells (Fig. 8). However, in the stroma, TUNEL positive cells were significantly (P<0.05) increased in the CN-009 1260 group or T60 group compared with the Control group (Fig. 9).

Acinar Epithelial Score (Fig. 10)

In the Control group, acinar epithelial score was significantly lower (P<0.01) than that of the Sham-ope group. In comparison with the Control group (Fig. 10), the acinar epithelial score was significantly higher (P<0.01) in the CN-009 1260, GBX, and TS groups.

Stromal Area Ratio (Fig. 11)

In the Control group, the stromal area ratio was significantly higher (P<0.01) than that of Sham-ope group in comparison with the Control group.
In comparison with the Control group (Fig. 11) the stromal area ratio of the CN-009 1260 was significantly (P<0.01) lower. The T-60 group was also significantly (P<0.05) lower than the Control group. However, there was no difference between other groups.

**Discussion**

Although chronic prostatitis is a common disease, it is very difficult to treat effectively.

Typical clinical findings are decreased potential, perineal or scrotal pain, urethral discharge and lower urinary tract irritative symptoms. The prostate gland is irregularly indurated and the numbers of leukocytes in expressed prostatic secretion are increased [15]. Pathological findings of this disease are chronic inflammation characterized by aggregates of lymphocytes in the stroma and
acute inflammation characterized by the presence of neutrophic polymorphonuclear leukocytes in the lumen of acinar glands [15-17]. Pathological definition of chronic prostatitis is different from clinical definition for the urologists. Clinical definition has been the combination of a clinical symptom and inflammatory cells in expressed prostatic secretion. The pathological inflammation of the prostate was reported to be not frequent in patients with symptoms of chronic prostatitis/chronic pelvic pain syndrome [16].

In experimental animals, Lewis, Wistar and Copenhagen rats have a high incidence of spontaneous nonbacterial prostatitis [14]. Administration of exogenous 17β-estradiol can induce 100% of the incidence on prostatitis in old Wistar rats [18] and castration also has a similar effect [13, 18]. Naslund et al. [13] reported that histopathological findings were very similar between spontaneous nonbacterial prostatitis and estradiol-induced prostatitis in rats [13]. These histopathological findings in rat spontaneous nonbacterial age-dependent prostatitis demonstrated several similarities to pathologically defined chronic prostatitis in human [19, 20]. These findings suggested that this rat model would be a useful model for the study of the treatment of human chronic prostatitis. Therefore, we decided to investigate the effects and mechanisms of CN-009 using a nonbacterial prostatitis rat model [13, 14] induced by 17β-estradiol injection and castration.

No differences in the prostate weight were found in CN-009 630, CN-009 1260, T-60, and GBX
groups compared with the Control group. Since
the weight of the prostate is mostly determined
by the amount of residual secretory fluid, these
findings may indicate that CN-009 cannot
prevent the reduction of secretory prostatic fluid.

In the CN-009 1260 group, we observed roundly
shaped acinar glands, cuboidal acinar epithelial
cells containing secretory materials with positive
PAP staining and diminished cell infiltrations into
the lumen compared with the Control group. The
acinar epithelial score was significantly
increased. CN-009 could protect acinar epithelial
function and cell shape against nonbacterial
inflammation. GBX had a similar effect to CN-
009 in the acinar glands. T-60 was not effective
in the acinar epithelial function of this rat model.
Therefore, GBX may play a central role for the
protection of epithelial damage in the NBP. The
effect of GBX was similar to that of TS. However,
GBX does not contain androgenic activity, because GBX has no effect on normal
and hypertrophic prostate (unpublished data,
1968). Accordingly, this protective effect of GBX
is discriminated from TS effect. In an in vitro
study, GBX was reported to inhibit the
cyclooxygenases and 5-lipoxygenases in the
biosynthesis of the prostaglandins and
leucotrienes [21]. Since prostaglandins and
leucotriens enhance inflammatory cell
infiltrations, GBX may protect against
inflammation into the acinar lumen by inhibition
of these enzymes. Furthermore, CN-009
showed an inhibition on the heat-induced
hemolysis, which is correlated to lysosomal
membrane stability [11]. CN-009 appears to
stabilize a lysosomal membrane, recover cell function and protect against degeneration of the acinar epithelium.

In addition, T-60 was shown to inhibit the growth of an immortal prostate cancer cell line in vitro [22]. However, their mechanisms are unknown. In the present study, the ratio of stromal area was significantly decreased in the CN-009 1260 and T-60 groups. Stromal TUNEL positive cell counts were increased in these groups. Therefore, CN-009 and T-60 may inhibit stromal cell proliferations by enhanced apoptosis. Although the exact mechanism of this process is unclear, several speculations are possible such as the direct effect by the apoptosis of fibroblast, and the indirect effect by the apoptosis of lymphocytes throught the the inhibition of several cytokines, such as several interleukins.

Further laboratory studies are necessary to elucidate the exact mechanisms of this compound.

Since no toxicological effects have been shown even in long-term administration, CN-009 is thought to be a safe drug [6, 23]. Here we reported the effects and mechanisms of CN-009 on rat experimental nonbacterial prostatitis model. CN-009 will also be a safe and effective against human nonbacterial chronic prostatitis.

In conclusion, CN-009 can work as a potent anti-inflammatory agent against chronic prostatitis. The present findings suggest that GBX, a fat soluble fraction of CN-009, protects the function and shape of acinar glandular epithelium and T-60, a water soluble fraction of CN-009, inhibits stromal cell proliferations in association with enhanced apoptosis.

---

Image 1: Effects of CN-009 on acinar epithelial score of the prostate. Each column represents the mean ± SD. **Significantly different from the Sham-op group at P < 0.01. #Significantly different from the Control group at P < 0.01.

Image 2: Effects of CN-009 on stromal ratio in the prostate. Each column represents the mean ± SD. **Significantly different from the Sham-op Group at P < 0.01. *Significantly different from the Control group at P < 0.01.
Acknowledgements

We thank M. Komukai, M. Ishii, F. Kimura and E. Higaki for their technical assistance.

References