Effects of depomedroxyprogesterone acetate on the development and maintenance of *Candida albicans* in the vagina of oophorectomized Wistar rats (*Rattus norvegicus*)

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The objective of this study was to determine the effect of medroxyprogesterone acetate (DMPA) on the development and maintenance of *Candida albicans* in the vagina of oophorectomized Wistar rats. The animals were divided into negative control groups (NCG), which received injections of sterile saline; positive control groups (PCG), which were given injections of estradiol valerate; and progesterone groups (PG), which were given injections of Depo-Provera®. After one week of hormonal induction, vaginal infection by *C. albicans* was induced in all the groups and detected by vaginal yeast culture and Papanicolaou smear. In addition, scanning and transmission electron microscopy images were obtained to confirm the vaginal infection by yeast in PG. A difference in progesterone levels in PG was observed between the basal level and after hormonal induction (P<0.0001). In this group, 100% of the rats acquired vaginal infection in the first week, but did not maintain it until the third week. The pharmaceutical brand of DMPA was effective for inducing the metestrus or diestrus phase of the estrous cycle in rats, similar to the use of pure progesterone. In contrast to estrogen treatment, progesterone alone could not support an experimental vaginal infection by *C. albicans* for any significant period of time.


O objetivo do presente estudo foi determinar os efeitos do acetato de depomedroxyprogesterona (ADMP) no desenvolvimento e manutenção de *Candida albicans* na vagina de ratas Wistar ooforectomizadas. Os animais foram divididos em grupos controle negativos (GCN), que receberam injeções de salina estéril; grupos controle positivos (GCP), que receberam injeções de valerato de estradiol; e grupos progesterona (GP), nos quais foram feitas injeções de Depo-Provera®. Após uma semana da aplicação hormonal, foi induzida a infecção vaginal por *C. albicans* em todos os grupos, detectada por cultura para leveduras vaginais e esfregaço de Papanicolaou. Foram feitas ainda imagens por microscopia eletrônica de varredura e transmissão para confirmar a infecção pela levadura no GP. Foram observados diferentes níveis de progesterona em GP, entre os valores basais e após a indução hormonal (P<0.0001). Neste grupo, 100,0% das ratas contraíram a infecção vaginal na primeira semana, mas não a mantiveram até a terceira semana. A forma farmacêutica de ADMP foi efetiva em induzir as fases de metaestro e diestro do ciclo estral das ratas, da mesma forma que usando progesterona pura. Em contraste com o que ocorre no tratamento com estrógeno, a progesterona não pôde manter a infecção vaginal experimental por *C. albicans* por um período significativo de tempo.

INTRODUCTION

Vulvovaginal candidiasis (VVC) is a pathology caused by the abnormal growth of yeast-like fungi in the mucosa of the female genital tract. It is an infection of both the vulva and the vagina caused by commensal yeasts, especially Candida albicans that inhabits the vaginal mucosa (Sobel, 1993; Sobel, 2007). This infection is characterized by itching, burning, dyspareunia, and a grumous cream-like vaginal discharge. The symptoms worsen during the pre-menstrual period, when vaginal acidity increases (Sobel, 1993).

VVC is sometimes difficult to treat, and represents a worldwide health problem of indisputable importance (Corsello et al., 2003; Patel et al., 2004). The diagnosis and treatment of VVC, together with lost productivity, result in an estimated cost of US$ 1 billion per year in the USA, where this infection is the second most common cause of vaginal infections (close to 25%), after bacterial vaginosis. Recently, the incidence of VVC has increased markedly worldwide. It is estimated that 75% of adult women have at least one episode of VVC during their lives, 40-50% of these will experience recurrences, and 5% will reach the recurrent character (RVVC), which is defined as the occurrence of three to four episodes of VVC in a period of 12 months in the absence of a recognized predisposing factor (Corsello et al., 2003).

The attachment of C. albicans to mucosal surfaces has been shown to be an important step in the infectious process, particularly in the oral cavity and vaginal mucosa (Irie et al., 2006). The pathogenesis of this infection involves the initial adherence of the yeasts to the vaginal mucosa, followed by asymptomatic colonization, from which the yeasts may reach the status of infectious agent (symptomatic vaginitis). This occurs when the colonization site of the host becomes favorable for the development of yeasts, normally as a consequence of some predisposing factor, such as previous colonization by yeast, the diminished immunological response observed in immunosuppressive diseases, diabetes mellitus, pregnancy, or chronic use of corticoids. The use of antibiotics, estrogen therapy, minor traumas such as the sexual act, the habit of wearing tight or synthetic clothing, and diet also seem to contribute (Corsello et al., 2003; Patel et al., 2004; Sobel, 2007). In the absence of these factors, clinical observations show that VVC occurs predominantly during the luteal phase of the menstrual cycle, when the levels of estrogen and principally progesterone are high. In contrast, pre-menstrual girls and post-menopausal women who are not receiving hormone replacement therapy, rarely exhibit this infection (Kalo-Klein, Witkin, 1989). However, the mechanisms by which these hormones, principally progesterone, act in VVC are not fully known (Fidel et al., 2000; Miller et al., 2000).

The behavior of the epithelia and of the cervical and vaginal stroma in response to sexual steroids is similar to that of humans, i.e., changes take place in the cellular pattern of the vaginal epithelium, depending on the phase of the estrous cycle (Mandal, Zuckerman, 1951). Therefore, experimental models of VVC in female rats have been extremely useful in the identification of factors concerning the influences of hormones on the infection, the virulence of the yeasts, and the susceptibility to and the treatment of the infection (Kalo-Klein, Segal, 1988; Kalo-Klein, Witkin, 1989; Junqueira et al., 2005). In these models, the key to obtaining a persistent infection is the state of pseudo-estrus, which is often induced through the expensive subcutaneous administration of estradiol benzoate (Clemons et al., 2004; Junqueira et al., 2005). Models using progesterone are scarce.

The purpose of the present study was to determine the effects of medroxyprogesterone acetate (Depo-Provera®) (DMPA) on the development and maintenance of Candida albicans in the vagina of oophorectomized rats.

MATERIALS AND METHODS

Animals

Oophorectomized Wistar rats (Rattus norvegicus) weighing 200 to 300 g and 70 days old, supplied by the Central Animal House of the State University of Maringá-PR (UEM), were employed. The animals were kept at a temperature of 22 ºC, under a light-dark cycle of 12 hours, and received a standard ration of chow and water ad libitum. This research was approved by the Committee on Ethical Conduct in the Use of Animals of UEM (Protocol No. 013/2006, statement No. 050/2006). The rats were used in the experiments after a period of adaptation of about seven days at the animal house.

Groups of rats per experiment

Each experimental day consisted of two control groups (CG) and three test groups (PG-progesterone groups), with each group containing five rats (total =75 rats). The study was carried out using these groups on three different days. Two different control groups were used: a negative control group (NCG), given subcutaneous injections of sterile saline (0.9% NaCl); and a positive control group (PCG), given subcutaneous injections of estradiol valerate (Sigma, Switzerland) dissolved in sesame seed
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oil at 0.2 mg/wk/rat, both administered as fractions three times/wk for four weeks. The infection was induced in all groups as described below.

**Induction of metestrus and diestrus in the progesterone groups (PG)**

The model of infection was carried out as described for mice (Fidel *et al.*, 2000), but instead using rats and a pharmaceutical brand of depomedroxyprogesterone acetate. Briefly, subcutaneous injections of Depo-Provera® (Aventis) were given at a concentration of 100 mg/wk/rat, administered as fractions three times/wk for four weeks, to obtain the state of metestrus and diestrus. This schedule was used because in the estrous cycle, progesterone secretion increases during metestrus and diestrus, decreasing thereafter (Mandal, Zuckerman, 1951; Fidel *et al.*, 2000). The hormonal activity was checked through the analysis of the cells of the vaginal fluid. Vaginal material was collected every three days after hormonal induction for four weeks. The vaginal fluid was obtained with the aid of a plastic pipette, after administration of 20 µL sterile saline in the vagina of the rats. A drop of this material was placed between a glass slide and coverslip and observed fresh by ordinary optical microscopy under 10X and 40X objectives. Three types of cells were identified: round and nucleated cells are epithelial cells; irregular cells without a nucleus are the cornified cells; and the small round cells are the leukocytes. The proportion among these types was used to determine the estrous cycle phases. A metestrus smear consists of the same proportion of leukocytes, cornified, and nucleated epithelial cells whereas a diestrus smear primarily consists of a predominance of leukocytes (Mandal, Zuckerman, 1951).

Hormonal activity was also checked by determining the blood level of progesterone in PG and NCG, and estrogen in PGC. Three days after hormonal induction, and weekly for four weeks, soon after the collection of the vaginal fluid, blood was collected for the determination of hormones, after 12-h fasting. The rats’ tails were cut, and the blood was collected in tubes containing an anti-clotting agent (fluoride EDTA). The tails were immediately cauterized to accelerate healing. The rats were then returned to their cages, and the ad libitum supply of water and chow was reestablished. Blood hormones were determined through solid-phase competitive chemiluminescent enzymatic immunoassay. The procedure was totally automated (IMMULITE 2000 – DPC MEDLAB Equipment) and results expressed as pg/mL (NCCLS, 1998).

**Experimental infection**

A human vaginal isolate of *C. albicans* identified at the Medical Mycology section of the Laboratory of Teaching and Research in Clinical Analyses (LEPAC) of UEM was used. This isolate had been stored since identification in 10% glycerinated water at -20 °C. The yeast isolate was identified through classical (Kurtzmann, Fell, 1998) and molecular methods (Sugita *et al.*, 2002). For the preparation of the suspension, the yeast was reactivated through seeding onto SDAC- Sabouraud Dextrose Agar (Sigma Chemical, St. Louis, USA) supplemented with 50 mg/mL of chloramphenicol (Sigma Chemical, St. Louis, USA) and incubated at 25 °C/48-72 h. An inoculation of the yeast at a concentration of 5x10⁸ CFU/mL (colony-forming units), counted in a Neubauer chamber, was used.

To induce the infection, 80 µL of the yeast suspension was introduced intra-vaginally one week after the beginning of hormonal induction. The animals were followed weekly to monitor the infection, from the first week after infection until week three (PG, NCP, PCG). Samples from the vaginal fluid, collected as described previously, were placed in Petri dishes containing SDAC, and incubated at 25 °C/48-72 h. After fungal growth, the CFU was counted. The infection was considered positive if at least one yeast colony grew. The same procedure was followed in the rats from the control groups. In addition to the culture, the vaginal samples were prepared as smears on glass slides, and stained for Papanicolaou cytology (Bibbo, 1997).

**Scanning (SEM) and transmission (TEM) electron microscopy**

After one week of infection, one rat each from the PG and the PCG, was sacrificed with an overdose of anesthetics (Ketamine and Xylazine) and the vaginas removed. For SEM, after the washes these were fixed in 2.5% glutaraldehyde solution dissolved in 0.1 M cacodylate buffer (Sigma Chemical, ST. Louis, USA) and dehydrated in an ascending series of alcohol. The critical point was obtained using a Balzers CPD-010 (Balzers Instruments, Balzers, Liechtenstein) with carbonic gas. Metallization in gold was done on a Balzers SCD-030 (Balzers Instruments, Balzers, Liechtenstein). Documentation was carried out on a JEOL-JSM 6360 LV scanning electron microscope (Jeol Ltda, Tokyo, Japan) at the Center of Electron Microscopy – Federal University of Paraná.

For TEM, after fixation in Karnowiski (2% paraformaldehyde, 2.5% glutaraldehyde in 0.1 M cacodylate buffer, pH 7.2 at 4 °C) the vaginal specimens were washed in 0.1 M cacodylate buffer pH 7.2 at 4 °C. Subsequently, the
material was post-fixed in 2% osmium tetroxide in 0.1 M cacodylate buffer pH 7.2 for one hour. The contrasting in the blocks was done in 2% uranyl acetate for two hours. The material was dehydrated in an ascending series of alcohols and then placed in acetone. The impregnation and inclusion occurred in Epon-812 resin. The sections were produced on a Sorval Porter Blum MT-2 ultra-microtome using glass and diamond blades. The ultra-thin sections were contrasted in aqueous solution of 2% uranyl acetate and lead nitrate/acetate. The observation of the material was made under a JEOL 1200EX II transmission electron microscope at the Center of Electron Microscopy of the Federal University of Paraná.

Statistical analysis

The results were analyzed using a t test and analysis of variance for comparisons between the hormone concentration at the basal level, and after the injections of the hormone, and for the percentage of the rats infected for one experimental week. A p value of less than 0.05 was considered significant. The tests were carried out with Graph Pad Prism® version 4.0 software (Graph Pad Software Inc.).

RESULTS AND DISCUSSION

In PG, 90.0% of the rats had smears of the diestrus type, and 10.0% metestrus whereas in PCG, 100% were in pseudoestrus. In NCG, the smears showed that the rats were in various phases of the estrous cycle. Figure 1 shows an unstained native vaginal smear from PG rats, visualized under a light microscope.

Therefore, the pharmaceutical brand of DMPA proved effective for inducing the metestrus or diestrus phase of the estrous cycle in rats, similarly to other studies using pure progesterone (Kinshiman, Collard, 1986; Fidel et al., 1993). This experimental model is both economically and operationally accessible, and may help to clarify important aspects in the pathogenesis of infection.

In the NCG, the blood progesterone (as mean values) ranged from 9.0 to 11.5 pg/mL (P> 0.05) (Figure 2). These rats, although infected with the yeast, did not develop vaginal infections.

In the PCG, the basal levels of estrogen were between 44.9 and 55.9 pg/mL; and after hormonal administration they ranged from 1384.0 to 1579.0 pg/mL (P< 0.05) (Figure 3 (A). Figure 3 (B) shows that 100% of these rats acquired a vaginal infection in the first week, and retained it for the three weeks of the experiment.

FIGURE 1 - Photomicrographs of unstained native vaginal smear from rats, observed under a light microscope with 25, 10, and 40X objective lenses, respectively, after administration of depomedroxyprogesterone acetate 100.0 mg/wk/rat. The proportion of the three types of cells was used for determination of estrous cycle phases. The round and nucleated cells are epithelial cells (E); irregular cells without a nucleus are the cornified cells (CC); and the small round cells are the leukocytes (L). A metestrus smear consists of the same proportion of leukocytes, cornified, and nucleated epithelial cells (A); and a diestrus smear primarily consists of a predominance of leukocytes (B).

FIGURE 2 - Blood progesterone concentration (pg/ml) determinations (as mean values) of the negative control group (NCG) at basal level, and seven days after sterile saline injection. *P > 0.05 for the progesterone concentration compared at basal level and after sterile saline injection.

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In the PG, the basal levels of progesterone were between 7.8 and 16.9 pg/mL; after hormonal administration, values ranged from 188.0 to 211.0 pg/mL (P< 0.05) (Figure 4A). In this group, 100% of the rats acquired a vaginal infection in the first week, and 0% retained it after the third week (P< 0.05) (Figure 4B). The CFU/mL of *C. albicans* in PG rats ranged from 275 to 5000 in the first week, and from 19 to 66 in the second week. No growth was observed in the third week (P< 0.05) (Figure 4C).

In contrast to estrogen (Figure 3B), progesterone treatment alone could not support an experimental vaginal infection for any significant period of time (Figure 4B, C). Other investigators have reported similar results, although these used pure progesterone rather than a pharmaceutical brand for human treatment (Kinshiman, Collard, 1986; Fidel *et al*., 1993). This phenomenon also occurs in women taking progesterone contraceptives such as Depo-Provera. Miller *et al.* (2000) reported a decreased proportion of positive *Candida* vaginal cultures (from 32% to 8%) after only 3 months of using DMPA as a contraceptive. Similarly in this study, there were no symptomatic *Candida* infections among women who had used DMPA for longer than 1 year. Fidel *et al.* (2000) also suggested that progesterone treatment in women and rodents alone could not support a vaginal infection by *Candida*, because of a lack of, or reduced, influence of endogenous estrogen. This theory was studied by Cundy *et al.* (1998), who reported that serum estradiol
The Papanicolaou staining of the rat vaginal smears showed the presence of yeasts and pseudohyphae only in the PCG and PGs, as illustrated in Figure 5C. It was also possible to observe the phases of the estrous cycle (Figure 5). This staining may be used for infection and hormonal control.

In addition to ordinary optical microscopy of the vaginal fluid, the scanning and transmission electron microscopy images confirmed the vaginal infection by *Candida albicans* in PG. However, this methodology cannot be used in many research laboratories because it is difficult to use in experimental models of VVC in female rodents, where cultures and Papanicolaou smears are more feasible. In Figure 6, SEM showing the surface of the vaginal epithelium of female rats in diestrus, infected with *Candida albicans*. In (A) and (B), a few blastoconidial yeasts can be seen, adhered as colonies. The yeasts adhered to the epithelium in an external location relative to the epithelial surface. In Figure 7 (A) and (B), the integrity of the epithelium can be seen by TEM, as well as several leukocytes in the lumen of the vaginal canal. In (c), TEM showing several non-adherent *C. albicans* yeasts in the lumen of the vaginal canal.

Kinsman, Collard (1986) suggested that host factors in the rat vaginal smears showed the presence of yeasts and pseudohyphae only in the PCG and PGs, as illustrated in Figure 5C. It was also possible to observe the phases of the estrous cycle (Figure 5). This staining may be used for infection and hormonal control.

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(E2) levels are markedly reduced in human DMPA users.

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Concerning the hypothesis that the presence of a cornified epithelium predisposes to VVC, our results also demonstrated the presence of abundant cornified cells in PCG, as previously described by other authors (Smith *et al.*, 1975; Kinsman, Collard, 1986). However, because
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some authors hold that adherence is lesser in cornified epithelial cells and greater in intermediate epithelial cells (IC) of the vaginal epithelium Segal et al., 1984; Kalo-Klein, Segal, 1988; Irie et al., 2006), further studies to assess these results should be undertaken. In addition, some clinical observations have established that VVC occurs more frequently during the luteal phase of the menstrual cycle, when progesterone is secreted in a greater quantity than estrogen (Kalo-Klein, Witkin, 1989; Miller et al., 2000). Recent research efforts have revealed that yeast cells can respond to human steroids (Smith et al., 1975; Banerjee et al., 2008). Banerjee et al., (2008) listed a total of 99 genes of C. albicans that are regulated by progesterone, and their correlation with drug resistance, virulence, morphogenesis, and general stress response. This explained, in part, the mechanisms by which progesterone acts in VVC pathogenesis.

FIGURE 7 - Yeasts and vaginal epithelium of rats observed by transmission electron microscopy (TEM), after establishment of diestrus and induced vulvovaginal candidiasis. In (A) and (B), the integrity of the epithelium can be seen, as well as several leukocytes in the lumen of the vaginal canal. In (C), TEM showing several Candida albicans yeasts in the lumen of the vaginal canal (bar = 0.5 μm).

CONCLUSIONS

Our results showed that the pharmaceutical brand of DMPA was effective for inducing the metestrus or diestrus phase of the estrous cycle in rats, similarly to other studies using pure progesterone. In contrast to estrogen, progesterone treatment alone could not support an experimental vaginal infection for any significant period of time. The Papanicolaou staining of the rat vaginal smears showed the presence of yeasts, and also revealed the phases of the estrous cycle. Further in vivo studies are necessary to assess the specific role of progesterone and other hormonal methods of contraception, and yeast colonization and infection. The experimental model used in the present study can be applied in this research.

REFERENCES


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