Clinical Study

An Evaluation of Therapy with Fluconazole 150 mg Tablets Compared to Fluconazole 150 mg Tablets Plus Dermoxen Lenitiva Cream in The Time to Reduce Symptomatology in Women with Vulvovaginal Candidiasis

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Abstract. Aim of the study. Authors investigated first of all the time to onset of first relief of symptoms. Secondary measures included the time to overall relief of symptoms and the reoccurrence rate over the first 45 days after the first visit. Methods. A randomized, open-label, parallel study evaluated 47 women with moderate to severe symptoms of Vulvo Vaginal Candidosis (VVC). Patients were divided into two groups of treatment: group 1 followed a therapy with Fluconazole 150 mg tablets, while group 2 had a therapy based on Fluconazole 150 mg tablets coadjuvated by Dermoxen Lenitiva cream. Results. The time at which 50% of patients experienced first relief of symptoms was 24.6 hours for Group 1, while for Group 2 it was 12.4 hours (P < 0.05). There were significant differences between the two groups in respect to the time of first relief of symptoms and reoccurrence of infection within 45 days of treatment. Conclusions. Combined treatment with Fluconazole 150 mg tablets and by Dermoxen Lenitiva cream provides statistically significant improvement in the time of first relief of symptoms, complete relief of symptoms and relapse time in the treatment of VVC compared to fluconazole 150 mg tablets only.

Keywords: Fluconazole, Dermoxen, vulvovaginal candidiasis, vaginitis

1. Introduction

Recurrent vulvovaginal candidiasis (RVVC) is a debilitating chronic infectious condition. It is defined as four or more acute inflammatory episodes of Vulvo Vaginal Candidosis (VVC), also known as vaginal yeast infection, within a year [1, 2]. The Candida albicans spp. has been known to be the main responsible organism for RVVC, accounting for 80%–85% of cases. The other cases are due to non-albicans species, with C. glabrata being the most common. Its frequency has nearly doubled over the last ten years, and it has been shown to account for 5%–15% of RVVC [3–5]. Other non-albicans species also include C. tropicalis (<5%) and C. krusei (about 1%) [6]. It has been estimated that 75% of
all women will experience at least one episode of VVC, and approximately 40% to 50% reporting a reoccurrence during their lifetime [7]. The main symptoms of yeast infection are inflammation, itching, an abnormal vaginal discharge and painful sexual intercourse and urination [8–11]. All these symptoms often cause severe discomfort, reducing quality of life of women and their partner. Acute inflammatory episodes are usually treated with antifungal drugs of the azole class. They are effective in clearing the acute infection, but they are unable to prevent recurrences, which occur on average after a few months only. Vulvovaginal candidiasis has been associated with considerable direct and indirect economic costs [12], enhanced susceptibility to HIV infection [13], and it is being investigated for a potential relationship with preterm birth [14]. Fluconazole is the only orally available imidazole with approved labeling specific for the treatment of VVC. The recommended therapy is a single 150 mg oral tablet. Many clinical studies have compared a single fluconazole 150 mg oral tablet with a number of different antifungal vaginal topical and suppository preparations. Review of these studies suggests that the overall cure rate with fluconazole 150 mg tablets is similar to that seen with other preparations [15–23]. Moreover Dermoxen Lenitiva cream, a topical formulation based on natural actives, was found to be active against itching [24], often present in women suffering from VVC, and useful to improve intimate comfort and well-being during sexual intercourse.

The purpose of this study was to compare the time of first relief of symptoms due to VVC, evaluating the results of two study groups: group I followed a therapy with Fluconazole 150 mg tablets while group 2 had a therapy based on Fluconazole 150 mg tablets plus Dermoxen Lenitiva, a soothing intimate cream produced by Ekuberg Pharma srl (Martano, Lecce- Italy). Moreover, the time for complete relief of symptoms and reoccurrence rate were evaluated, following up patients for 45 days after the end of treatment.

2. Materials and Methods

Authors developed a pilot, open-label, randomized study, conducted according to Declaration of Helsinki and approved by an Institutional review Board. Before to start, the protocol and informed consent were reviewed, approved and signed by the patients. Authors enrolled, from July 2013 to November 2013, 47 women who had symptoms of VVC, attending the Department of Gynaecology and Obstetrics of University affiliated Hospital “Vito Fazzi” (Lecce, Italy). Confirmation of current VVC infection was made by use of KOH wet mount preparation, pelvic examination and patient’s reporting of signs and symptoms. The exclusion criteria for such study were: patients suspected of having a concurrent vaginal infection (i.e., bacterial vaginosis, trichomoniasis, herpetic lesions); women with menstruation or women who know menstrual cycle comes within two days; patients with a history of use of intravaginal or systemic antifungal medication or other intravaginal products (spermicide, douche, spray, gel, cream); women with a medical history of allergies or intolerance to any of the active or non-active ingredients of the study formulations. The symptomatology was evaluated using a scale from 0 (no symptoms) to 10 (severe symptoms) for itching, redness, burning and dryness. Details of last sexual intercourse, last menstrual period, method of contraception, recent treatment, parity, contact’s symptoms and relevant past history were all recorded. After diagnosis by microscopy, the patients were treated randomly with two types of treatment: Group I (N = 24) was treated with Fluconazole 150 mg tablets (one tablet per day, for two weeks), Group II (N = 23) with a therapy based on Fluconazole 150 mg tablets plus Dermoxen Lenitiva cream (one tablet and one application of cream per day, for two weeks). Patients were recommended to record the time of dosing. The main indication was to follow the treatment in the early afternoon for a better evaluation of first relief of symptoms. Patients were given a personal diary in which they were requested to record the date and time they first started to feel relief of symptoms, and the date and time they had complete relief of symptoms. In this diary patients had to notice any adverse events or concomitant medications. Seven days after the end of the treatment, patients were required to return to hospital for a follow-up visit, during which the investigators checked the diary. 45 days after the end of the treatment, patients were required to return another time to hospital to investigate any case of relapse. The first outcome of the study was the time of first relief of symptoms. The second outcome of the study was the evaluation of complete relief of symptoms and of any case of relapse, and their time of appearance.

2.1. Statistical analysis. Two independent reviewers collected, reported and classified data. Baseline demographics (age, number of episodes of VVC in the previous 12 months and severity score) were tabulated and compared using descriptive statistics. Time of first relief of symptoms for each patient was calculated using the relative dosing time and the time reported for first relief of symptoms by each patient. Analysis of these data was performed using Kaplan–Meier estimates and 95% Confidence Interval analysis. P values less than 0.05 were considered statistically significant.

3. Results

In this pilot study, 47 women with recognized VVC were treated with two different protocols and included in two groups: Group I (N = 24) and Group II (N = 23). A total of 77 patients (33 in group I and 44 in Group II) were eligible for inclusion during first visit; 30 women were excluded according to exclusion criteria (9 in Group I and 21 in Group II). The analysis of demographic data was similar between the two groups (Table 1).
Group I (with two different protocols and included in two groups: This study involved 47 women with recognized VVC, treated 4. Discussion

hours for Group I and 64.3 for Group II ($P < 0.05$) ($\text{symptom relief within 24.6 hours versus 12.4 of Group II}$

of Group II experienced first relief of symptoms with a higher

percentage in comparison with those of Group I at all time

points. Fifty percent of patients of Group I experienced first

symptom relief within 24.6 hours versus 12.4 of Group II

($P < 0.05$). The median time for total relief was 77.3

hours for Group I and 64.3 for Group II ($P < 0.05$).

Evaluating patients of both groups after 45 days from the

eend of the treatment, a total of 5 patients (20.83%) of Group

I experienced a reoccurrence of VVC, versus one patient

(4.34%) of Group II ($P < 0.05$).

Both forms of treatment were effective in reducing the

signs and symptoms of vaginal candidiasis. All patients in

both groups had not registered any adverse events in their

diary.

In Table 2 the percentage of patients who recorded first

symptoms relief within the first 48 hours is reported. Patients

of Group II experienced first relief of symptoms with a higher

percentage in comparison with those of Group I at all time

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($P < 0.05$). The median time for total relief was 77.3

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I experienced a reoccurrence of VVC, versus one patient

(4.34%) of Group II ($P < 0.05$).

4. Discussion

This study involved 47 women with recognized VVC, treated

with two different protocols and included in two groups: Group I ($N = 24$) and Group II ($N = 23$). Both proposed
treatments registered an improvement in vaginal symptoms.

Combined treatment with Fluconazole 150 mg tablets plus

Dermoxen Lenitiva cream (one tablet and one application of cream per day, for two weeks) showed a significant

improvement compared to the treatment with Fluconazole

150 mg tablets only, about time of first relief of symptoms

of VVC. In fact, fifty percent of patients of Group II experienced first relief of symptoms after 12.4 hours versus 24.5 hours

of Group I. It has been estimated that 24 hours is a reasonable

period of time in order to evaluate general effectiveness

of therapy for VVC. For Group I, 49 % experienced first

symptom relief after 24 hours, versus 74.4% for Group II

($P < 0.05$). This is an important result coming out from

the study, because it suggests that a combined “oral and

topical” treatment is more effective in relieving symptoms

of VVC than oral or topical treatment only. Total relief was

achieved after 77.3 hours for Group I and 64.3 for Group

II. No adverse events were registered in both groups. In

particular, in this study the oral treatment was administered
to defeat the infection, using fluconazole, because the main

objective of any therapy for the treatment of vaginal can-
didiasis is the eradication of the infecting organism; while

the topical treatment was specifically addressed to reduce

the symptomatology regarding itching, redness, burning of

external intimate area in terms of time. Another outcome of

this study was to evaluate the reoccurrence time, monitoring

patients of both groups for 45 days from the first visit: a total

of 5 patients (20.83%) of Group I experienced a reoccurrence

of VVC, versus one patient (4.34%) of Group II ($P < 0.05$). This is another important result, which needs better

investigation in order to understand if and how Dermoxen

Lenitiva cream could have a coadjuvant action against yeast
(action not presented by the product). Definitely, combined

treatment with Fluconazole 150 mg tablets plus Dermoxen

Lenitiva cream (one tablet and one application of cream

day, for two weeks) more rapidly achieves first relief of

symptoms of VVC, compared to Fluconazole 150 mg tablets

only, reducing cases of RVVC.

5. Conclusion

Combined treatment with Fluconazole and Dermoxen Leni-
tiva cream should be considered as an important first line

therapy in patients presenting the signs and symptoms

of VVC and RVVC. A trial involving more patients and

more outcomes, may guarantee a better investigation of this

Table 1: Demographic data.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Group I ($N = 24$)</th>
<th>Group II ($N = 23$)</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (mean ± SD)</td>
<td>34.4 ± 6.6</td>
<td>34.1 ± 8.1</td>
<td>0.2540</td>
</tr>
<tr>
<td>Parity (mean ± SD)</td>
<td>1.1 ± 1.0</td>
<td>1.2 ± 1.0</td>
<td>0.6430</td>
</tr>
</tbody>
</table>

Table 2: Time (hours) of first symptom relief.

<table>
<thead>
<tr>
<th>Hours after dosing</th>
<th>Group I Cumulative % ($N = 24$)</th>
<th>Group II Cumulative % ($N = 23$)</th>
<th>$P$ value (*statistically significant)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>2.9</td>
<td>11.7</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>4</td>
<td>9.1</td>
<td>22.3</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>6</td>
<td>12.5</td>
<td>27.0</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>8</td>
<td>19.2</td>
<td>34.9</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>10</td>
<td>24.0</td>
<td>45.2</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>12</td>
<td>24.0</td>
<td>48.3</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>14</td>
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<td>57.1</td>
<td>$P &lt; 0.05*$</td>
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<tr>
<td>16</td>
<td>35.1</td>
<td>57.1</td>
<td>$P &lt; 0.05*$</td>
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<td>18</td>
<td>39.4</td>
<td>64.6</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>20</td>
<td>44.4</td>
<td>64.6</td>
<td>$P &lt; 0.05*$</td>
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<tr>
<td>22</td>
<td>46.7</td>
<td>69.0</td>
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<tr>
<td>24</td>
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<td>$P &lt; 0.05*$</td>
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<td>77.7</td>
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<tr>
<td>28</td>
<td>51.1</td>
<td>80.0</td>
<td>$P &lt; 0.05*$</td>
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<tr>
<td>30</td>
<td>55.6</td>
<td>80.0</td>
<td>$P &lt; 0.05*$</td>
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<tr>
<td>32</td>
<td>61.5</td>
<td>82.9</td>
<td>$P &lt; 0.05*$</td>
</tr>
<tr>
<td>34</td>
<td>61.5</td>
<td>82.9</td>
<td>$P &lt; 0.05*$</td>
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<tr>
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<td>69.8</td>
<td>82.9</td>
<td>$P &lt; 0.05*$</td>
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<tr>
<td>38</td>
<td>76.0</td>
<td>82.9</td>
<td>$P &lt; 0.05$</td>
</tr>
<tr>
<td>40</td>
<td>78.0</td>
<td>85.6</td>
<td>$P &lt; 0.05$</td>
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<td>42</td>
<td>81.8</td>
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<tr>
<td>44</td>
<td>85.9</td>
<td>92.0</td>
<td>$P &lt; 0.05$</td>
</tr>
<tr>
<td>46</td>
<td>88.0</td>
<td>93.6</td>
<td>$P &lt; 0.05$</td>
</tr>
<tr>
<td>48</td>
<td>88.0</td>
<td>95.0</td>
<td>$P &lt; 0.05$</td>
</tr>
</tbody>
</table>
finding, taking care of extending the time of follow up to 90 days.

References


Comments

1. We changed the affiliation reference of “Andrea Tinelli” to be “6” instead of “5”. Please check.

2. We changed “p” to “𝑃”. Please check throughout.

3. We split reference “4” to “4 and 5”. Please check.

4. We split reference “9” to “10 and 11”. Please check.