Full length Research Paper

Investigation of *Demodex* spp. in perinea areas and faces of young men visiting the urology policlinic

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It has been reported that types of *Demodex* can be transmitted to other people through close contact and they can play a pathogenic role in rosacea, acne vulgaris, perioral dermatitis, seborrhoeic dermatitis, micropapillary-pruritic dermatitis, and blepharitis. Types of *Demodex* are reported to be located in various places of human body. The purpose of the present study was to detect the presence of *Demodex* spp. in the samples taken from the faces and perinea areas of the young men using standardized surface skin biopsy (SSSB). Accordingly, samples were obtained using SSSB method from the faces and perinea areas of 200 men aged 19 - 34, which were then covered with entellan and sent to the parasitological laboratory. *Demodex* spp. was found in 42.0% of the samples taken from perinea area and in 83.0% of the samples taken from subjects' faces, while *Enterobius vermicularis* was detected in two cases. One of the samples taken from was *Demodex brevis*, while the others were *Demodex folliculorum*. It was concluded that, given that species of *Demodex* can be found in perinea area under conditions of collective accommodation, specimens should be taken from patients consulting with pruritus or allergic reactions in order to diagnose the parasite, and treatment protocol should be planned according to the results of the analysis.

Key words: *Demodex* spp., facies, perinea and cheek.

INTRODUCTION

It has been reported that among the *Demodex* spp. species *Demodex folliculorum* and *Demodex brevis* usually settle in the pilosebase units in the hair folicules on human face. It is also stated that *D. folliculorum* survive alone or in groups in follicular spaces, whereas *D. brevis* live alone in the depts of sebaseosis glands and the thin and long structures of the mites are suitable for these places (Erbagci and Ozgoztasi, 1998; Wesolowska et al., 2005).

It has been reported that the immunologic reactions developed against the parasite as a result of the proliferation of *D. folliculorum* due to immunological defects play a role in emergence of skin lesions (Dong and Duncan, 2006). It has also been reported that

species cies of *Demodex* are found in various places of human body including nasolabial region, base of eyelashes, chin, forehead, outer ear canal, nipple, back, penis and hips (Baima and Sticterling, 2002; Nutting, 1976). The first case of *Demodex* was detected by (Saygi et al., 1984) in perennial area using cellophane tape method.

Methods used for diagnostic purposes include cellophane tape, skin scraping, punch biopsy and standardized surface skin biopsy (SSSB). Among these diagnostic methods SSSB is an effective one in terms of detecting the mite intensity of the parasite per cm², since it is able to collect completely the follicular content together with the surface part of the stratum corneum where the species of *Demodex* inhabit (Erbagci and Ozgoztasi, 1998; Forton et al., 2005). It was aimed in this study to detect the presence of *Demodex* spp. In the samples taken from the faces and perinea areas of the young men with SSSB.

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MATERIALS AND METHODS

Prior to the study, health staff working in urology policlinic were given an applied in-service training by a parasitologist about *Demodex* species and their parasitism, the purpose of the study and planned procedures, how to obtain specimens from women's perinea area using SSSB method. The specialized doctors who would obtain the specimens practiced on volunteering subjects. After the surface part of the stratum corneum together with the hair follicles on the specimens was observed, the evaluation began.

The research group included 200 male patients aged 19 - 34 who visited the urology policlinics. Among the patients, those who were detected to have pruritus and blushes in the perinea area based on their histories were informed comprehensively about the present research by the specialized doctor, and specimens taken from the perinea area of the volunteering subjects using SSSB method were sent to the parasitological laboratory after being covered with entellan. Specimens were examined in parasitological laboratories through light microscope at X100 and X400 magnification. Even if one *Demodex* spp. was observed, it was considered as positive. Parasite intensity of 5 and more per cm² of samples taken from face was considered as positive. Patients with *Demodex* spp. positive were first informed about the results and forwarded to the relevant policlinic.

Statistical analyses were done using Student $\it t$ test for independent samples, Yates' corrected chi -square test, and Fisher's exact test. The data were presented in terms of mean values, standard deviation, number, and percentage. The statistical significance was considered as p < 0.05 and analyses were done using SPSS 13.0 software.

RESULTS

Specimens were taken from faces and perinea areas of 200 men aged 19 - 34 using SSSB method, covered with entellan, and sent to the parasitological laboratory. The examination revealed the presence of *Demodex* spp. among 42.0% of the specimen from perinea (Figure 1) and among 83.0 % of the specimens from face (Figure 3 - 4), and *Enterobius vermicularis* in 2 of the specimens from perinea (Figure 2). One of the specimens from face was *D. brevis*, while the others were *D. folliculorum*. The relationship between the presence of *Demodex* on face and pruritus complaints is shown in Table 1.

Fisher's Exact Test was used in Table 1, p = 0.54. No significant relationship was found between the presence of *Demodex* on face and pruritus complaint (p = 0.54). The relationship between pruritus in perinea area and the presence of *Demodex* is given in Table 2. Yates' corrected chi-square was used in Table 2, which revealed no significant relationship (p = 0.69). The percentages on the co-existence of *Demodex* on face and perinea area are shown in Table 3. Yates' corrected chi-square was used in Table 3, which revealed no significant relationship (p = 0.54). The statistics about age, barrack size (number of men in a barrack) and presence of *Demodex* are shown in Table 4. Independent samples test was applied, and no significant differences in age and barrack size were found between presence and absence of *Demodex*.

DISCUSSION

With regard to the pathological and clinical symptoms caused by different types of *Demodex*, some researchers consider the inhabitance of *Demodex spp* in pilosebase follicles as harmless, while others have reported that D. folliculorums can play an etiopathogenic role in rosacea, acne vulgaris, blepharitis, perioral dermatitis, pustular folliculitis, papular -pustular lesions on hairy skin, and pustular lesions in acquired immune deficiency syndrome (Dong and Duncan, 2006; Mathieu and Wilson, 2000; Wesolowska et al., 2005) . In various researches about the epidemiology of types of *Demodex*. (Akdeniz et al., 2002) found that the density of D. folliculorum among diabetic patients was significantly higher than control group. Similarly, in a research on patients with chronic kidney failure, Ozçelik et al. (2007) found the parasite in 12.76% of the 47 patients. In a different study Ding and Huang (2005) examined the outer ear canal secretion of 613 healthy high school students and found Demodex in 11.58% of them. Also Karaman et al. (2008) reported the presence of Demodex spp. in 15.3% of the hairy skin biopsies diagnosed keratinized cist, trikelemmal cist, nevus, dermatitis, car-cinoma and inflammation. Forton et al. (1993) reported that in specimens obtained from 49 patients with rosacea using SSSB they found the density of D. folliculorum about 10.8/cm², which was significantly higher than the control group. Moreover, the researchers stated that SSSB is an effective method measure D. folliculorum density easily. Again in a research to evaluate the D. folliculorum pathogenesis in rosacea Abd-El et al. (1997) found the mite density as 28.6/cm² in randomly chosen 16 patients with papulopustular rosacea problem and 6.9/cm² in control group. Similarly, Roihu et al. (1998) reported the presence of *Demodex* in 51% of 80 patients with rosacea, 28% of the 40 patients with egsema and 31% of the 40 patients with lupus erythematosus.

Sener et al. (2009) aimed to detect the positivity of Demodex spp. inbiopsy specimens of skin diagnosed as nevus. In the study the specimens obtained from 110 patients diagnosed with nevus 43 (39.1%) out of 110 specimens were detected to have Demodex spp. Demodex colonization augmented in nevi can be explained by the possible affinity of the parasite to the melanin pigment. It was reported that the prevalence of Demodex species increases as the patients grow older. Aycan et al. (2007) reported Demodex prevelance in 20% of the 20 age group and in 53.5% of the 21 age group. In their study to evaluate the relation between presence of Demodex spp. and age (Baysal et al. 1997) found 8.3% positivity among 11 - 15 age group and 12.7% positivity among 16 - 20 age group. It has been reported that Demodex types do not appear among children and increase as of adolescence and peak in older ages. This finding can be interpreted as a result of lack of considerable differences between the ages of the participating



Figure 1. Adult Demodex spp. 100X.



Figure 2. E. vemicularis egg 100X.

men. As far as the relevant literature was scanned, no researches about the epidemiology of *Demodex* have been reported on the face and perinea area of the young men. Yet, Ugra et al. (2009) examined the specimens obtained from the perianal area of 100 men and found no *Demodex* presence. As a result of the present study, we

found *Demodex* spp. in 42.0% of the specimens from perinea and in 83.0% of the specimens from face, and *E. vermicularis* in two of the specimens taken from 0.2% *Taenia* spp. eggs, and 0.2% *Phthirus pubis*. Saygi et al. (1984) reported *Demodex spp.* in perennial area material using cellophane tape method. These findings are similar



Figures 3 and 4. Adult *Demodex* spp. in specimens from face 100X.

Table 1. The relationship between the presence of *Demodex* on face and pruritus complaint.

| | Demodex on face | | | | | | |
|------------------|-----------------|------|--------|------|--------|-------|--|
| | Yes | | No | | Total | | |
| | Number | (%) | Number | (%) | Number | (%) | |
| Pruritus on face | | | | | | | |
| Yes | 29 | 16.3 | 149 | 83.7 | 178 | 100.0 | |
| No | 5 | 22.7 | 17 | 77.3 | 22 | 100.0 | |
| Total | 34 | 17.0 | 166 | 83.0 | 200 | 100.0 | |

Table 2. The relationship between the presence of *Demodex* and pruritus in perinea.

| | Demodex in perinea | | | | | | |
|---------------------|--------------------|------|--------|------|--------|-------|--|
| | Yes | | No | | Total | | |
| | Number | (%) | Number | (%) | Number | (%) | |
| Pruritus in perinea | | | | | | | |
| Yes | 101 | 59.2 | 70 | 40.9 | 171 | 100.0 | |
| No | 16 | 55.2 | 13 | 44.8 | 29 | 100.0 | |
| Total | 117 | 58.5 | 83 | 41.5 | 200 | 100.0 | |

Table 3. Presence of *Demodex* on face and perinea.

| | Pruritus in perinea | | | | | | |
|-----------------|---------------------|------|--------|------|--------|-------|--|
| | No |) | Yes | | Total | | |
| | Number | (%) | Number | (%) | Number | (%) | |
| Demodex on face | | | | | | | |
| No | 22 | 64.7 | 12 | 35.3 | 34 | 100.0 | |
| Yes | 95 | 57.2 | 71 | 42.8 | 166 | 100.0 | |
| Total | 117 | 58.5 | 83 | 41.5 | 200 | 100.0 | |

Table 4. Distributional statistics on age group, barrack size and presence of *Demodex*.

| | | Age | Barrack | | |
|--------------------|-----|----------------|---------|-----------------|------|
| | n | Mean ± SD | Р | Mean ± SD | Р |
| Demodex on face | | | | | |
| No | 34 | 21.3 ± 2.5 | 0.15 | 50.4 ± 43.9 | 0.63 |
| Yes | 166 | 21.9 ± 2.6 | | 46.4 ± 43.7 | |
| Demodex in perinea | | | | | |
| No | 117 | 21.8 ± 2.3 | 0.56 | 49.0 ± 49.3 | 0.44 |
| Yes | 83 | 22.0 ± 2.8 | | 44.5 ± 34.2 | |

to the results obtained in the present study.

It was concluded that, given that species of *Demodex* can be found in perinea area under conditions of collective accommodation, specimens should be taken from patients consulting with pruritus or allergic reactions in order to diagnose the parasite, and treatment protocol should be planned according to the results of the analysis.

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