



Research Article

Sleep Quality in Psoriasis Patients and its Relations with Possible Affecting Factors

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Abstract

Objectives: Psoriasis (PS) is a chronic, immune-mediated inflammatory skin disease. It can lead significant effects on health-related quality of life (HRQoL) and other outcomes. In the current study, investigation of sleep quality (SQ), and its possible relations with factors which can affect it were aimed.

Methods: A total of 74 subjects from both sexes were enrolled in this study, between January and July, 2017. Patients were evaluated with their demographics, body mass index (BMI), Psoriasis Area Severity Index (PASI), Pittsburgh Sleep Quality Index (PSQI), Psoriasis Quality of Life Index (PQLI), Self Perception Scale (SPS). Additionally, accompanying chronic diseases, disease duration and severity of pruritus were recorded. Obtained PSQI values were compared with the mentioned parameters in terms of significance of their relations with it. SPSS, version 24, 2016 was used to analyse the data, and significance was evaluated with p values of <0.05, 0.01, and 0.001, and rho (r) values of <0.2, =0.2-0.4, =0.4-0.6, =0.6-0.8 and >0.8.

Results: Thirty-seven female and 37 male were studied. The mean age of total of study population was 47.21±13.91. Mean BMI and mean duration were 30.09±4.68 kg/m², and 10.58±9.1 months. Mean values of PASI, PSQI, SPS, and PQLI of the study group were 19.79±16.99, 9.14±5.09, 142.12±23.83, and 21.94±16.31, respectively. Approximately thirty-one percent of them had at least one chronic disease. Alcohol and smoking rates were 17.56%, 50%. PASI was positive/strongly correlated with PQLI, and negative/weakly correlated with SPS. No correlation was detected between PSQI values, and, age, gender, BMI, and SPS values. PSQI was moderately correlated with PQLI, diabetes mellitus (DM), and pruritus severity, whereas it was weak correlated with PASI, hypertension (HT), thyroid diseases and disease duration. PASI and DM showed predictive effect on SQ.

Conclusion: SQ is affected from certain factors such as QoL, disease severity, disease duration, pruritus severity, accompanying disorders such as HT, DM and thyroid diseases, in which disease severity and DM have predictive effects on SQ in PS patients. Controls of disease activation and prevention of progression in DM may provide to keep of SQ in PS.

Keywords: Causality; demographic factors; diabetes mellitus; psoriasis; sleep; quality of life.

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Psoriasis (PS) is a chronic, immune-mediated inflammatory skin disease of genetic basis, which characterized with sharp limited, erythemasquamous plaques. Its prevalence is between 1.5% and 5%. The disease can easily be diagnosed with chronic and frequently recurrent bright-red colored plaques which are coated with silvery and brittle scales, on the extensor surfaces of extremities, knees, elbows, hip, and scalp.^[1] The disease has a great impact on quality of life (QoL) due to the fact that both having a chronic and repetitive course and being a visual and cosmetic problem, that can almost be comparable to the impacts caused by systemic diseases. Indeed, especially in recent years, it has begun to be defined as a disease spectrum or a multisystemic disease rather than being a skin-confined disease.^[2] These comorbidities have been described as cardiovascular diseases, psychological/psychiatric disorders, inflammatory disorders, sexual dysfunctions, fatty liver disease, alcoholism, smoking, metabolic syndrome as a whole or its individual components such as hypertension (HT), dyslipidemia, high blood glucose, or atherosclerosis.^[3] On the other hand, sleeping is both a stationary state and active renovation phase of an organism in which whole body is recharged and prepared for life during this process.^[3, 4] It is well known that sleep disorders result in some changes in immune functions. Additionally, it has been proposed that these dysfunctions in sleep patterns can lead to a chronic inflammation in the metabolism, an increasing in activation of the diseases, and so a decreasing in Health-related Quality of Life (HRQoL). It is thought that the mechanism of these effects can be dependent on underlying dysfunctions in immune regulatory system. The importance of improvement of sleep dysfunctions should not be ignored in the management of chronic inflammatory diseases.^[4] Some recent reports regarding PS and sleep disturbances suggest that another comorbidity for PS is obstructive sleep apnea/hypopnea (OSAHS) syndrome, and there is a higher prevalence in relation to the general population. Although it is thought that the relationship stemmed from the increased prevalence of obese PS patients, strongly probable effects of inflammatory mediators should not be neglected.^[5] Similarly, Shutty et al.^[6] stated positive relation between insomnia and PS. Moreover, it is suggested that, pruritus, pain and emotional factors might have predictor effects on sleep disturbances in PS patients, and sleep disorders might be risk factors for the development and activation of PS.^[7] However, the exact relationship between the PS and sleep disorders is still inconsistent, and available data on detailed questioning of sleep quality (SQ) and related conditions are very limited and unsatisfying.^[3, 5-8] Thus, in this study investigating of overall SQ, sleep subcomponents, and determining of correlations of SQ and factors that can affect it, in PS patients were aimed.

Methods

A total of 74 subjects (37 male and 37 female) were enrolled in this study, who referred to our dermatology clinic between January and July, 2017. After the Local Ethics Committee approval, and the required written informed consent was obtained, volunteer patients were included in the study. PS diagnosed with clinical and/or histopathological examination findings. Inclusion criteria were as follows: volunteering to work, ≥ 18 years of age, clinical and/or histopathological diagnosis of PS, having the ability to understand the questions. Subjects with a cognitive impairment, psychiatric and other dermatological disorders, and ones who underwent any systemic therapy for their PS except for topical therapies were excluded from the study. Subjects' age, gender, smoking and drinking habits, any coexistent chronic diseases (diabetes mellitus (DM), hypertension (HT), heart disease, chronic kidney disease, chronic liver disease, thyroid disease) and disease duration were questioned. After the heights and weights of the subjects were measured, Body Mass Index (BMI) was calculated as weight (kilograms) divided by height (square meters) for an estimate of obesity. The evaluation of body weight was made as < 25 = normal and ≥ 25 = overweight, whereas determining of obesity was made as follows: non obese = ≤ 29.9 kg/m², and obese = ≥ 30 kg/m².^[9] PS severity was evaluated using Psoriasis Area Severity Index (PASI) scoring which is the golden standard for determine in disease severity in PS.^[10] PASI is an evaluation of the average redness, thickness and desquamation of the lesions (grades 0-4), weighted by the area of involvement. Minimum score is 0 (no disease) and maximum score is 72 (maximal disease), whereas values of 10 and above are considered moderate/severe disease. SQ was evaluated with the Pittsburgh Sleep Quality Index (PSQI).^[11] It is a self-administered questionnaire that subjectively measures SQ over the previous month. The questionnaire includes 19 questions that must be answered by patients, and five questions that must be rated by their bed or room partner (if exists). The questions form the component scores, each of which has a range of 0-3 points. A score of 0 indicates no difficulty in sleep, while a score of 3 indicates severe difficulty. The seven component scores which subjective SQ, sleep latency, sleep duration, habitual sleep efficiency, sleep disturbances, use of sleeping medication and daytime dysfunction finally give one total score (0-21 points). A total score of ≥ 5 indicates bad SQ. This system was tested and validated by Agargun et al.,^[12] in the Turkish population. Disease related QoL of the subjects was evaluated with the Psoriasis Quality of Life Index (PQLI) which is determined by a likert type survey. This PS-specific scale was first developed in our coun-

try by Aydemir et al., in 2003, and was integrated to QoL studies associated with PS, after its validity and reliability were confirmed. The scale is comprised of 17 questions organized in 3 categories which include disease findings, social and sexual live properties of patients, and use of disease-specific medication. It is entirely composed of yes-no questions and all "yes" answers should further be rated in 4 different severity categories. There is no any cut-off value for this index. It provides a categorical evaluation in QoL of PS patients. As PQLI score increases, disease related QoL is proportionally and adversely impacted. Pruritus was assessed in 5 categories according to answers given in question five of PQLI scale (no, yes/but it does not affect at all, yes/it affects a little, yes/it is quite effective, yes/very effective).^[13] Perception of body image was determined by Self Perception Scale (SPS) which is developed by Secord and Jourard in 1953. This scale is consist of a total of forty questions in five categories which regards different body images. The Turkish validity and reliability were performed by Hovardaoğlu et al.^[14] in 1992. It is composed of 40 items which each is related to an organ or a part of the body. Each item includes five-grade answer options (I do not like it at all, I do not like it, I'm undecided about it, I like it, I like it very much) that can takes values from 1 to 5. The cutt of point of the scale is 135, and total score can take values from 40 to 200. A value under 135 indicates low self-perception. The study was conducted in accordance with the "World Medical Association Declaration of Helsinki, ethical principles for medical researches involving human subjects, 2013".

Statistical Analysis

Obtained data was evaluated with the *Statistical Package for Social Sciences* (SPSS Statistics for Mac, version 24, New York, USA, 2016). Descriptive statistics were used as *mean, standart deviation (SD), median, and frequency*. The normality assumption of the variables was checked with *Kolmogorov-Simironov test*. Because our all variables were distributed asymmetric, *Mann-Whitney U non-parametric analysis* was used for comparing different parameters. *Pearson correlation test* was used for determining in relation of qualitative and quantitative variables which were distributed normal, whereas *Sperarman's rho test* was used for ones which were distributed asymmetric. A p value of <0.05 was considered as significant in binary comparisons, whereas p values of <0.05, 0.01 and 0.001 were considered as significant in relations in which they were interpreted as weak, moderate, and strong relation, respectively. Correlation results were interpreted according to rho (r) values, as follows: r<0.2 is very weak/no correlation, r=0.2-0.4 is weak correlation, r=0.4-0.6 is moderate correlation, r=0.6-0.8 is strong correlation, and, r>0.8 is very strong correlation. Fi-

nally, predictivity of determinant factors which show positive correlation on SQ was detected with multiple linear regression analysis method.

Results

A total of 74 PS subject were enrolled in the study (37 female/37 male). General characteristics of study population are seen in Table 1. Mean age of the subjects was 47.21±13.91 (min. 21/max. 73). Mean BMI of subjects was 30.09±4.68 kg (min. 21.9/max. 42.9), and majority of them (n=36, 48.6%) was obese. Approximately half of them (n=31, 41.89%) had at least one chronic disease, which were 25 hypertension (33.78%), 20 diabetes mellitus (27.02%), 5 thyroid diseases (6.75%), 13 alcohol user (17.56%) and 37 smoker (50%), respectively. Subjects' SQ was evaluated based on the total PSQI scores. Mean PSQI of the subjects was 9.14±5.09 (min.1 /max. 20), and most subjects (n=58, 78.3%) had bed SQ. Subjects' mean PASI value was found as 19.79±16.99 (min.1.5/ max. 70). A little more of the subjects had moderate/severe disease severity (n=46, 62.16%). Mean score for PLQI was 21.94±16.31 (min. 0/max.51), and there was very significant difference between PASI groups according to PLQI values (p<0.001). Mean SPS was 142.12±23.83 (min. 63/max.199). The difference in SPS values according to PASI groups was significant (p<0.05). Mean disease duration was 10.58±9.10 years (min.1/max.45 years). The linear relationships with PASI and PLQI scores, and, PASI and

Table 1. General characteristics of totally of study population

Variables	n (%)	Mean (±SD)	Min	Max
Gender (Female/Male)	37(50)/37(50)			
Age		(47.21±13.91)	21	73
BMI *		30.09±4.68	21.90	42.90
Disease duration (year/s)		10.58±9.10	1	45
PASI**		19.79±16.99	1.5	70
SPS***		142.12±23.83	63	199
PLQI ****		21.94±16.31	0	51
PSQI*****		9.14±5.09	1	20
PSQI components				
Sleep quality		1.83±0.82	0	3
Sleep latency		1.59±1.10	0	4
Sleep duration		1.39±1.20	0	3
Sleep efficiency		0.85±1.10	0	3
Sleep disturbances		1.41±0.57	0	3
Use of sleeping drug		0.68±0.79	0	3
Daytime dysfunction		1.55±0.99	0	3

BMI*: Body Mass Index; PLQI****: Psoriasis Life Quality Index; PASI**:
Psoriasis Area Severity Index; PSQI*****: Pittsburgh Sleep Quality Index;
SPS***: Self perception scale.

SPS scores are seen in Table 2. There was positive and very strong correlation between PASI and PLQI values ($r=0.823$), whereas negative and weak correlation was obtained between PASI and SPS scores ($r=-0.287$). Comparison of the PSQI values according to cut-off values of SPS, PASI and BMI (overweight/normal) is seen in Table 3. Moderate/severe PASI groups had worse SQ than the other group, whereas there was no difference according to SPS and BMI groups. Additionally, SQ was not different in obese and non-obese persons. Correlations of PSQI values and demographic and disease related other factors affecting on SQ are seen in Table 4. No any linear relationships were detected between PSQI values, and, age, gender, BMI, and SPS values, whereas they were positively correlated with disease duration, PASI, PLQI, DM, HT and thyroid diseases. Strengths of these correlations were weak for disease duration (0.329), PASI ($r=0.363$), HT ($r=0.256$) and thyroid diseases ($r=0.248$), whereas moderate correlations were obtained for PLQI ($r=0.433$) and DM ($r=0.409$). There was significant relationship with PSQI values and pruritus, PSQI was moderately correlated ($r=0.424$) with severity of pruritus. Factors showing both positive correlation with PSQI values and have significant predictivity on SQ are seen in Table 5. According to this stepwise analysis, only PASI and DM had meaningful predictive effects on PSQI values, while other parameters did not show any predictivity.

Table 2. Comparison of the PSQI values of study population according to cut-off values of SPS, PASI and BMI.

n=74	Median (±SD)	U	p	Effect size
SPS*				
<135, (n=23)	8.91±3.89 (9)	576.50	0.907	0.00
≥135, (n=51)	9.25±5.57 (8)			
PASI**				
<10 Weak, (n=28)	6.10±3.30 (6)	289.50	<0.0001 [†]	0.46
≥10 Moderate-Severe, (n=46)	11.00±5.12 (12)			
BMI***				
<25, (n=10)	8.20±4.46 (9)	281.50	0.542	0.07
≥25, (n=64)	9.29±5.19 (9)			

Mann-Whitney U test; [†]p<0.0: SPS*: Self perception scale; PASI**: Psoriasis Area Severity Index; BMI***: Body Mass Index.

Table 3. Correlations of disease severity with PLQI and SPS values

n=74	PLQI*	SPS**
PASI***	0.823▲▲	-0.287▲

Spearman's rho Korelasyon; ▲▲p<0.01, ▲p<0.05; PLQI*: Psoriasis Life Quality Index; SPS**: Self perception scale; PASI***: Psoriasis Area Severity Index.

Discussion

PS is one of the 5 diseases which the World Health Organization (WHO) reported as having a great burden on patients' QoL.^[15] It has physical, social and psychological negative effects on patients' health.^[6, 7] On the other hand, sleep is an essential requirement for daily functioning and health. Jensen et al.^[16] stated in their study that PS patients had poor overall SQ and insomnia in the rates of 53.9% and 25% compared to 21.9% and 10.5 of controls, respectively. They indicated that these differences were statistically significant, and itch was strongly associated with all sleep-related outcomes in PS patients. Biçici et al.^[17] also reported that overall SQ was worse in PS than in their controls. Additionally, patients with worse SQ had higher rates of depression and anxiety scores, and pruritus had significant effect on worsening in SQ. Melikoğlu stated that 60% of PS patients had poor PSQI values, and their means were significantly different from healthy controls.^[18] Gowda et al.^[7] indicated that PS patients had important sleep disturbance

Table 4. Correlations of PSQI values and demographic factors, disease duration and some other factors affecting on sleep quality

Variables	PSQI* values
Age	0.212
Gender	0.200
BMI**	-0.078
sDisease Duration	0.329▲▲
PASI***	0.363▲▲
SPS****	-0.081
PLQI*****	0.433▲▲
Hypertension	0.256▲
Diabetes mellitus	0.409▲▲
Thyroid diseases	0.248▲
Alcohol consumption	0.183
Smoking	0.045
sPruritus severity	0.424▲▲

sSpearman's rho Korelasyon; Pearson Korelasyon; ▲▲p<0.01, ▲p<0.05; r<0.2= very weak or no correlation; r=0.2-0.4 weak correlation; r=0.4-0.6 moderate correlation; r=0.6-0.8 strong correlation, r>0.8 very strong correlation; PSQI*: Pittsburgh Sleep Quality Index; BMI**: Body Mass Index; PASI***: Psoriasis Area Severity Index; SPS****: Self perception scale; PLQI*****: Psoriasis Life Quality Index.

Table 5. Factors showing both positive correlations with PSQI values and have significant predictivity on SQ

Determinant factors	B	S.E	β
Constant factor	14.702	2.190	
Diabetes mellitus	4.347	1.153	0.382
PASI	0.099	0.030	0.332

Multiple linear regression analysis (R²=0.17 in first step and 0.28 in second step in stepwise model); PASI*: Psoriasis Area Severity Index.

compared to those of healthy controls, whereas Shutty et al.^[6] reported that the of bed SQ in PS was 4.3 times more higher than healthy controls. Similarly, in Wong et al.^[19] and Taçiet al.'s^[20] studies, sleep disorders were found significantly higher in PS patients compared to the healthy population, and they stated that this situation was related with patients' fatigue, anxiety and poor QoL. However, Stinco et al.^[21] did not find any difference in SQ between PS patients and healthy controls.

In accordance with most of aforementioned literature, we found overall poor SQ (PSQI ≥ 5) in our subjects in the rate of 78.3%, and there was not any difference according to age or genders. We detected significant difference between the subjects having low (<10) and high (≥ 10) PASI values, and subjects having higher PASI values had worse SQ. In Stinco et al.^[21] and Duffin et al.^[22] studies they did not find any correlation with PASI and PSQI values. However, Strober et al.^[23] found a correlative but non predictor relationship between PASI and PSQI. Similar to the last study, our PASI scores showed meaningful and positive correlation with PSQI values. Although strength of this correlation was weak, PASI was found as a predictive factor for SQ. Mean duration of our subjects was 10.58 ± 9.10 years. Similar to PASI, there was positive but weak correlation between the PSQI and disease duration. It did not meet our expectations, because both sleep disorders have been associated with chronic inflammatory processes,^[22] and higher PASI values and long disease duration are usually related with higher disease activity and increased chronic inflammation in PS, respectively.^[10, 22] However, these inconsistencies related with correlations of PASI and disease duration might be resulted from a great distances in minimum and maximum values of PASI (min.1.5/ max. 70) and of durations (min.1/ max.45 years), compared to the relatively small number of subjects. Because, these variables were distributed asymmetric and taking account of outliers in an asymmetric group analysis may change interpretation of results depending on the coefficient preferred.^[24]

On the other hand, PS leads to a lot of consequences, such as cardio/occlusive vascular diseases, anxiety/depression, inflammatory bowel diseases, uveitis, arthritis, erectile dysfunction, fatty liver, alcoholism, smoking, lymphomas, osteoporosis, Parkinson's disease, obesity, HT, dyslipidemia, DM, atherosclerosis or metabolic syndrome.^[3] Similar to the literature, 48.6% of our subjects was obese. Although high BMI and obesity have been identified as risk factors for poor SQ, the relationship is still conflicting.^[4] Indeed, our PSQI values did not show any difference according to overweight and normal BMI-groups, and also obese and non-obese groups. Thirty-one of our subjects had at least one chronic disease (HT, DM, thyroid diseases). These findings

were complied with the usual literature. It is known that increase in blood pressure leads to changins in SQ, sleep quantity, and sleep fragmentation,^[25] and DM has been described as a risk factor for poor SQ.^[4] On the other hand, although some reports suggest that thyroid disorders can decrease in SQ as lead to OSAHS, some others suggest that there is no direct correlation between the two condition.^[26]

Considering the correlations with PSQI values, we found positive but weak correlations between the PSQI scores, and, HT and thyroid diseases, whereas DM showed positive and moderate correlation. Additionally, DM had a predictivity on SQ. Approximately seventeen of our subjects was alcohol user, but quantity of consumption in most of them was no more than two standart beer, or 2 glasses of wine, and only at the weekends. Although alcohol is frequently used for become tranquilized and as self medication for insomnia, it leads to disruptions in sleep architecture and reduction in the total night REM, especially when consumption is chronic and excessive (>4 standard drinks/day).^[27] However, we did not find any correlation between the two condions which might be related to the low amount of consumption of our subjects.

Fifty percent of our subjects was smoker. It is well known that there is strong relation between smoking cessation and poor SQ. On the other hand, some authors reported that positive association between smoking and poor SQ which is related with stimulation and inhibition of different neurotransmitters in sleep-regulating ventrolateral preoptic region, whereas others suggest that no or negative relationship.^[28] We did not find any correlation between SQ and smoking.

Another factor related with sleep disorders in PS is pruritus. Although the exact relationship is not known between SQ and pruritus, it has been suggested that the pruritus in PS patients might be related with both mechanical stimulation of pruritus and PS-related psychologic factors such as depression.^[17] Gowda et al.,^[7] Biçici et al.^[17] and Amatya et al.^[29] found that SQ negatively affected from pruritus in PS. We also detected significant positive relationship between the two condition, and there was moderately correlation between the severity of pruritus and poor SQ.

SQ is one of the most important factors affecting QoL, and humans need an awarage of 8 hours of nocturnal sleep to have more energy and performance, better cognition, improved memory, alertness, and heathier immunity. In the evaluation of overall QoL in PS patients, usually HRQoL-scales or dermatologic QoL-scale (DQoL) have been used. However, it is well known that PS is not only life-restrictive condition, but also causes important psychological consequences because it leads to significant visual impairment,

and this relation can not be neglected.^[13] Thus, we preferred to use PQOL scale to detect PS-specific QoL, instead of the other scales.

Indeed, PQOL is rather similar to a psychological scale than being solely skin scale, which measures especially PS-related physic, social, mental and psychological restrictions in different components of everyday life, such as hair dressing, sleeping with partners, sexual relation, go to restaurant, shake hands, wear a decollete dress. PS patients feel shame and usually hesitate at these activities.^[13, 30] It is suggested that barriers in QoL in PS patients are developed due to essentially mental and psychological restrictions because poor skin appearance. Body image (BI) is a multidimensional concept, including thoughts, emotions, feelings and behaviors related to physical appearance, and attitude to body. It began to change when a patient suffers from any kind of dermatosis, and finally leads to decreased in QoL, accompanying anxiety, depression, social phobias and adaptation disorders.^[30]

Our moderate/severe PS group also had worse PQOL scores than been in the low-PASI group, and the difference is significant, and our findings were consistent with literature. We also found positive and moderate correlation between the PLQI and PSQI values, and PLQI showed predictive effect on SQ.

The importance of outer aspect and attitude to one's body are the key factors of human mentality. Skin appearance is associated with adaptive mental functioning and well-being, thus plays an important role in the BI, self-assessment, self-perception (SP), self-esteem, and establishing satisfactory relations with others. Because PS lesions are usually visible, they result in negative mental consequences, and subsequently distortion in the BI. Studies usually suggest a strong relationship between PS lesions and negative SP due to deteriorated health, unfavorable appearance and lack of satisfaction with appearance of body, sexual inhibition, and restricted physical performance, however irrelevant results have also been taken.^[30]

Our mean SPS score was 142.12 ± 23.83 . PSQI did not show significant difference according to SPS groups, and no correlation was detected in the two condition. Interestingly, although 75.67% of our subjects had moderate and severe disease, SP skills of them were evaluated as inadequate in only 31.08% of them. Thus, we thought that only severity of skin lesions is not enough to decrease in SP of PS patients, and obtained irrelevant results between SQ and SP might be affected from this reason.

Our study showed that SQ was moderately correlated with factors of QoL, pruritus, and DM, whereas it was no or weakly correlated with others, in PS patients. But only

PASI values and DM had predictive effect on SQ. However, broad-based and controlled further studies are needed to criticize our results.

Conclusion

SQ may be affected from certain factors such as QoL, disease severity, disease duration, accompanying disorders such as HT, DM, thyroid diseases, and pruritus in PS patients, in which QoL, pruritus, and DM may be more correlated with it. Disease severity and DM have predictive effects on SQ. Controls of disease activation and prevention of progression in DM may facilitate to keep SQ in PS patients.

Disclosures

Ethics Committee Approval: The study was approved by the Local Ethics Committee.

Peer-review: Externally peer-reviewed.

Conflict of Interest: None declared.

References

- Gülekon A. Psoriasis ve benzeri dermatozlar. In: Tüzün Y, Güner MA, Serdaroğlu S, Oğuz O, Aksungur VL, editors. 3rd ed. Dermatoloji. Nobel Medical Bookstore: Istanbul; 2008. pp. 745-64.
- Langley RG, Krueger GG, Griffiths CE: Psoriasis: epidemiology, clinical features, and quality of life. *Ann Rheum Dis* 2005;64:18-23.
- Machado-Pinto J, Diniz Mdos S, Bavoso NC. Psoriasis: new comorbidities. *An Bras Dermatol*. 2016;91:8-14.
- Berhanu H, Mossie A, Tadesse S, Geleta D. Prevalence and associated factors of sleep quality among adults in Jimna Town, Southwest Ethiopia: A Community –based cross-sectional study. *Sleep Disord* 2018;22:8342328.
- Ranjbaran Z, Keefer L, Stepanski E, Farhadi A, Keshavarzian A. The prevalence of sleep abnormalities to chronic inflammatory conditions. *Inflamm Res* 2007;56:51-7.
- Shutty BG, West C, Huang KE, et al: Sleep disturbances in psoriasis. *Dermatol Online J* 2013;19:1.
- Gowda S, Goldblum OM, McCall WV, Feldman SR. Factors affecting sleep quality in patients with psoriasis. *J Am Acad Dermatol* 2010;63:114-23.
- Buslau M, Benotmane K. Cardiovascular complications of psoriasis: does obstructive sleep apnea play a role *Acta Derm Venereol*. 1999;79:234.
- Appropriate body-mass index for Asian populations and its implications for policy and intervention strategies". WHO Expert Consultation. *The Lancet*. 2004;363: 157-63.
- Feldman SR, Krueger GG. Psoriasis assessment tools in clinical trials. *Ann Rheum Dis*. 2005;64:65-8.
- Buysse DJ, Reynolds CF, Monk TH, Berman SR, Kupfer DJ. The Pittsburgh Sleep Quality Index: a new instrument for psychiatric prac-

- tice and research. *Psychiatry Res.* 1989;28:193-213.
12. Agargun MY, Kara H, Anlar O. The validity and reliability of the Pittsburgh Sleep Quality Index. *Turk Psikiyatri Derg* 1996;7:107-15.
 13. Aydemir Ö, Kamer Gündüz, Danacı AE, Errmertcan AT. Psoriasisli Hastalarda Yaşam Kalite Ölçeği Geliştirilmesi. *Turkderm.* 2003;37:189-195.
 14. Hovardaoğlu, S. Vücut Algısı Ölçeği. *Psikiyatri, Psikoloji, Psikofarmakoloji Dergisi (3P). Testler Özel Eki.* 1992;1:26-27.
 15. Wohlrab J, Fiedler G, Gerdes S, Nast A, Philipp S, Radtke MA, et al. Recommendations for detection of individual risk for comorbidities in patients with psoriasis. *Arch Dermatol Res.* 2013;305:91-98.
 16. Jensen P, Zachariae C, Skov L, Zachariae R. Sleep disturbance in psoriasis - a case-controlled study. *Br J Dermatol* 2018 Apr 28. doi: 10.1111/bjd.16702. [Epub ahead of print]
 17. Biçici F, Hayta SB, Akyol M, Özçelik S, Çınar Z. Evaluation of sleep quality in patients with psoriasis. *Turkderm* 2015;49: 208-12.
 18. Melikoglu M. Sleep Quality and its Association with Disease Severity in Psoriasis. *Eurasian J Med* 2017;49:124-7
 19. Wong ITY, Chandran V, Li S, Gladman DD. Sleep Disturbance in Psoriatic Disease: Prevalence and Associated Factors. *J Rheumatol* 2017;44:1369-74.
 20. Taçi D, Galimberti R, Amaya-Guerra M, et al. Improvement in aspects of sleep with etanercept and optional adjunctive topical therapy in patients with moderate-to-severe psoriasis: results from the PRISTINE trial. *J Eur Acad Dermatol Venereol* 2014;28:900-6.
 21. Stinco G, Trevisan G, Piccirillo F, Di Meao N, Nan K, Deroma L, et al: Psoriasis vulgaris does not adversely influence the quality of sleep. *G Ital Dermatol Venerol* 2013;148:655-9.
 22. Duffin KC, Wong B, Horn EJ, Krueger GG: Psoriatic arthritis is a strong predictor of sleep interference in patients with psoriasis. *J Am Acad Dermatol* 2009;60:604-8.
 23. Mukaka MM. A guide to appropriate use of Correlation coefficient in medical research. *Malawi Med J* 2012; 24:69–71.
 24. Seravalle G, Mancia G, Grassi G. Sympathetic Nervous System, Sleep, and Hypertension. *Curr Hypertens Rep* 2018;20:74.
 25. Bielicki P, Przybyłowski T, Kumor M, Barnaś M, Wiercioch M, Chazan R. Thyroid Hormone Levels and TSH Activity in Patients with Obstructive Sleep Apnea Syndrome. *Adv Exp Med Biol* 2016;878:67-71.
 26. Wilkinson AN, Afshar M, Ali O, Bhatti W, Hasday JD, Netzer G, Verceles AC. Effects of binge alcohol consumption on sleep and inflammation in healthy volunteers. *J Int Med Res.* 2018 Jan 1:300060518782020. doi: 10.1177/0300060518782020. [Epub ahead of print]
 27. Boakye D, Wyse CA, Morales-Celis CA, Biello SM, Bailey MES, Dare S, et al. Tobacco exposure and sleep disturbance in 498 208 UK Biobank participants. *J Public Health (Oxf)* 2017 Aug 30:1-10. doi:10.1093/pubmed/fox102. [Epub ahead of print]
 28. Amatya B, Wennersten G, Nordlind K. Patients perspective of pruritus in chronic plaque psoriasis: a questionnaire-based study. *J Eur Acad Dermatol Venereol* 2008;22:822-6.
 29. Rosińska M, Rzepa T, Szramka-Pawlak B, Żaba R. Body image and depressive symptoms in person suffering from psoriasis. *Psychiatr Pol* 2017;51:1145-52.