ABSTRACT

Background: Anogenital warts are growths caused by human papilloma virus infection, which causes significant psychosocial morbidity. Intraliesional immunotherapy is a promising treatment that is able to stimulate a delayed-type hypersensitivity reaction to different antigens as well as wart tissue. This treatment is suggested to increase CD4 T helper 1 lymphocyte activity to destroy human papilloma virus.

Objective: To measure the level of serum soluble CD4 (sCD4) in patients with anogenital after intralesional tuberculin purified protein derivative (PPD) injection.

Patients and Methods: This study was carried out as a pre–post interventional study on 49 patients with anogenital warts presented to the dermatology clinic and the andrology clinic, Suez Canal University Hospital. Each patient was injected 10U (0.2ml) of PPD intralesionally in the largest or mother wart with 2-week interval till complete cure or six injections, whichever was closer. Serum sCD4 was measured before and after injection using sandwich ELISA technique.

Results: Complete clinical wart recovery was obtained in 26 (53%) patients and mostly was after the sixth session (60.4%) of injection. Mean level of serum sCD4 was significantly higher after than before PPD injection (18.47±10.4 vs. 44.48±20.67ng/ml, respectively). Serum sCD4 levels were significantly correlated to clinical response rates, as the higher the level of serum sCD4, the more the wart recovery.

Conclusion: Intraliesional PPD injection is a promising effective modality in the treatment of anogenital warts. CD4 cells may have a role in anogenital wart eradication by PPD injection and significantly related to clinical response rates.

Key Words: Anogenital warts, immunotherapy, purified protein derivative, soluble CD4.

INTRODUCTION

Viral warts are growths or tumors affecting skin and mucous membranes caused by infection with human papilloma virus (HPV)\(^1\). Anogenital wart infection is considered one of the commonest sexually transmitted infections that occur in the genital, perigenital, and perianal regions in sexually active males and females and cause marked stress, anxiety, and interpersonal difficulties\(^2\).

Regulation of an effective immune reaction against different invading pathogens in human body is mainly dependent on CD4+T cells through release of distinctive cytokines after their activation and differentiation on exposure to those pathogens and also through its ability to stimulate different cell types as innate immune cells, cytotoxic T cells, B-lymphocytes and non immune cells\(^3\).

Therapeutic options in anogenital wart management depend on many factors like size, number, distribution and morphology of lesions, as well as treatment adverse effects, cost, availability and convenience, physician experience, and patient needs and preference. The general management of warts includes one of two methods: the first is the ordinary destructive method and the second is immunotherapy\(^4\).

Immunotherapy is a type of treatment that depends on using antigens that stimulate or suppress immune system to help in the treatment of malignancies, infections, immunological disorders, and others\(^5\). Immunotherapy in anogenital warts uses the capacity of the immune system to recognize certain viral, fungal, and bacterial antigens to initiate a delayed hypersensitivity reaction, which expands the ability of the immune system to perceive and remove the HPV\(^6\).

Cell-mediated immune response had been reported as
a suggested immune change found in patients with genital wart responding to treatment[7]. The CD4+ helper T cell is stimulated through antigen-presenting MHCII molecules, which leads to adaptive humoral immunity effective against warts[8].

Purified protein derivative (PPD) has a role in the treatment of viral warts and being capable of eliciting a delayed-type hypersensitivity at site of injection into patients previously sensitized with tuberculosis bacilli, as previously sensitized CD4+ T lymphocytes, attracted to the skin test site, proliferate and produce cytokines[9].

Intralesional PPD produces a powerful proinflammatory impulses and appeals antigen-presenting cells, which identify HPV fragments in the affected tissue causing a robust adaptive immune reaction that eradicates the wart infection. PPD has been found to be associated with the production of Thelper 1 cytokines such as interleukin (IL)-4, IL-5, IL-8, IL-12, interferon (IFN)-γ, and tumor necrosis factor α, which stimulate cytotoxic and NK cells that produce a destructive immune reaction against HPV[10].

Rahim et al.[11] assessed CD4 and CD8 levels in serum of patients infected with genital warts and reported that level of CD4 is lower in patients with genital warts than in controls in contrast to level of CD8, which was found to be higher in patients than in control group; statistical relationships were significant.

The current study was designed to find out the role of CD4 in the effect of PPD in treatment of anogenital warts through measurement of the level of serum soluble CD4 (sCD4) before and after intralesional injection of tuberculin PPD in anogenital warts.

PATIENTS AND METHODS

This study was a pre–post interventional study carried out on patients with anogenital warts recruited from the andrology clinic and the dermatology clinic, Suez Canal University Hospital, Faculty of Medicine, Suez Canal University, during the period from February 2019 to March 2020. A total of 49 patients who accepted to participate in this study according to planned eligibility criteria were randomly chosen. Patients included in the study were male patients with diagnosis of external anal or genital warts and aged more than 18 years. Patients excluded from the study were patients with active tuberculosis, chronic or debilitating diseases, and patients receiving immunosuppressive drugs, systemic corticosteroids, or other therapeutic modalities for anogenital warts, especially immunotherapy during the time of the study or in the last 3 months.

This study was approved by Ethical Committee Board of the Faculty of Medicine, Suez Canal University. Ethical Committee approval number was 3581, dated 9/9/2018. Clear informative medical consent containing aim of the study, rationale, objectives, methodology, expected benefits, and complications of the study was signed by all patients.

All patients were subjected to the following: thorough history taking was obtained and included personal history, present history (duration of infection, sites affected, illegal or extramarital relationships, presence of similar lesions in partners or current affection with other sexually transmitted infections, and previous therapeutic modalities for warts), past medical, and drug history to exclude conditions that interfere with tuberculin PPD injection.

Local examination included careful genital inspection to define number, site, and size of anogenital warts and detection of any genital dermatomes present. Total body skin examination was also performed to exclude presence of other dermatological conditions or any distant warts.

Tuberculin purified protein derivative injection

The vaccine used was PPD of a human strain of mycobacterium tuberculosis. It was manufactured by Vacsera Company (Cairo, Egypt). Each vial contained 2-ml solution, and concentration was 5TU (tuberculin unit) per 0.1ml solution.

Tuberculin test was done first by intradermal injection of 5TU (0.1ml solution) of tuberculin PPD in the left forearm. Presence of a well-defined indurated bleb at the site of injection measured about 10mm in diameter within 48–72h after injection meant that patient was sensitized[12].

After tuberculin test, sensitized patients were injected with tuberculin PPD intralesionally; 10U (0.2ml solution) were used per dose in the largest warts every 2 weeks till complete clinical cure or maximum of six injections, whichever was closer[13].

The clinical assessment was performed by the researchers before the study begins and every injection session. Assessment was done for warts in the injected sites and warts in distant anatomical sites.

Assessment of the result of injection was done using the following scale[14]: complete response: complete disappearance of all warts and return of normal skin markings; partial response: regression in warts’ size by 25–99% and no response: regression in size of warts by 0–25%.

The most common adverse effects were pain, edema, fever, mild erythema, and postinflammatory hypopigmentation. These adverse effects happened in a minority of patients and were treated symptomatically by analgesics and anti-inflammatory medications, and in all patients, these adverse effects did not lead to discontinuation of the study.

Assessment of wart recurrence was done by regular follow-up visits every month for 3 months in patients who experienced complete cure of warts to detect any recurrence.
either in same sites of injection or in neighboring sites.

**Assessment of serum soluble CD4**

The assay was done for all patients before PPD injection and after complete cure of warts or six sessions of PPD injection whichever was closer, according to the instructions provided by the manufacturer using sCD4 ELISA Kit (Product Id ABIN512865; antibodies-online GmbH, Aachen, Germany). The used sCD4 kit had human reactivity, colorimetric detection using Sandwich ELISA technique. Detection range was 5–100ng/ml, and the minimum detection limit was 5ng/ml.

Partial responders and nonresponders were examined carefully by the researchers and reassessed. Patients were shifted to other management plan such as cryotherapy, ablative CO2 laser or podophyllin, other immunotherapy materials, or combination of different modalities till complete recovery.

**Statistical analysis**

Data were analyzed using IBM SPSS software package, version 20.0. (IBM Corp., Armonk, New York, USA). Qualitative data were described using frequency and percentage. Quantitative data were described using range (minimum and maximum), mean, SD, and median. Significance of the obtained results was judged at the 5% level.

**RESULTS**

This study was carried out on 49 patients with external anogenital warts. The mean age of the studied group was 31.59±8.34 years, and most of them were less than 30 years old (55.1%). Mean duration of warts was 1.0–16.0 months, and 65.3% of patients had duration less than 8 months. Mean number of warts was 16.76±11.91 and ranged from 5 to 34 warts. Warts in distant anatomical sites were found in 18.4% of patients. Most of the patients completed their sessions and the mean number of sessions was 5.47±0.53, and the least number of sessions was three sessions. The most common sites of warts were shaft of penis (55.2%) and pubic area (44.9%), and the least affected area was perianal area (6.1%). Only 20.4% of patients experienced adverse effects to PPD injection; these adverse effects included local pain, edema, mild erythema, and fever and postinflammatory hypopigmentation (Table 1).

| Parameters                        | n (%)         |
|-----------------------------------|---------------|
| Age (years)                       |               |
| ≤30                               | 27 (55.1)     |
| >30                               | 22 (44.9)     |
| Minimum–maximum                   | 21.0–55.0     |
| Mean±SD                           | 31.59±8.34    |
| Duration of warts (months)        |               |
| ≤8                                | 32 (65.3)     |
| >8                                | 17 (34.7)     |
| Minimum–maximum                   | 1.0–16.0      |
| Mean±SD                           | 5.73±4.07     |
| Number of warts                   |               |
| Minimum–maximum                   | 5.0–34.0      |
| Mean±SD                           | 16.76±11.91   |
| Number of sessions                |               |
| Minimum–maximum                   | 3.0–6.0       |
| Mean±SD                           | 5.47±1.9      |
| Site of anogenital warts          |               |
| Shaft of penis                    | 27 (55.1)     |
| Pubic area                        | 22 (44.9)     |
| Scrotum                           | 10 (20.4)     |
| Frenulum of penis                 | 9 (18.4)      |
| Perianal area                     | 3 (6.1)       |
| Warts in distant anatomical sites |               |
| Present                           | 9 (18.4)      |
| Absent                            | 40 (81.6)     |
Clinical response to intralesional tuberculin PPD injection revealed that 26 (53%) patients completely recovered from warts, 16 (32.7) patients partially recovered, and seven (14.3%) patients were nonresponders. Complete clinical response occurred in most of the patients at the sixth session of injection (60.4%). Warts in distant anatomical sites were found in six of the 26 completely recovered patients; five (83.3%) of them totally recovered from these distant warts. Intralesional tuberculin PPD injection in warts is a safe method with minimal tolerable adverse effects. In the present study, we found that 79.6% of patients had no adverse reactions to tuberculin, and adverse effects were found only in 20.4%, in the form of pain (100%), mild erythema (80%), transient postinflammatory hypopigmentation (30%), fever (20%), and transient edema at the site of injection (20%). Recurrence occurred in four (15.4%) of the 26 completely recovered patients during the 3-month period after completion of sessions (Table 2).

**Table 2:** Frequency distribution of the studied patients according to clinical response to purified protein derivative injection, adverse effects and recurrence rate

| Clinical response to PPD                  | N=49 [n(%)] |
|------------------------------------------|-------------|
| No response (0–25%)                      | 7 (14.3)    |
| Partial response (25–99%)                | 16 (32.7)   |
| Complete response (100%)                 | 26 (53)     |
| Session of complete response (N=26)      |             |
| Third session                            | 1 (3.8)     |
| Fourth session                           | 3 (11.5)    |
| Fifth session                            | 6 (23.1)    |
| Sixth session                            | 16 (61.6)   |
| Response of distant warts in completely recovered patients (N=6) |             |
| Complete response                        | 5 (83.3)    |
| Partial or no response                   | 1 (16.7)    |
| Occurrence of Adverse reactions to PPD injection (N=49) |             |
| No adverse reaction                      | 39 (79.6)   |
| Adverse reaction                         | 10 (20.4)   |
| Adverse reactions to PPD injection (N=10) |             |
| Pain                                     | 10 (100)    |
| Mild erythema                            | 8 (80)      |
| Hypopigmentation                         | 3 (30)      |
| Fever                                    | 2 (20)      |
| Transient edema                          | 2 (20)      |
| Recurrence after completeresponse (N=26) |             |
| No                                       | 22 (84.6)   |
| Yes                                      | 4 (15.4)    |

PPD, purified protein derivative.

Level of serum sCD4 was significantly increased after intralesional tuberculin PPD injection as mean serum sCD4 was 18.47±10.4 versus 44.48±20.67 ng/ml before and after injection, respectively ($P<0.001$) (Table 3).

Correlation between different demographic parameters and clinical response of intralesional tuberculin PPD injection revealed that age was significantly related to response; the younger the age, the better the response. The number of warts was also significantly related to response, as the smaller the number, the better the response. Duration of warts was also significantly related to response, as the shorter the duration, the better the response ($P<0.001$). Site of warts was not a determinant factor in clinical response as there was no significant statistical relationship between site and clinical response (Table 4).
response revealed that before PPD injection level of sCD4 was not significantly related to clinical response, whereas after injection, serum sCD4 levels were significantly correlated \( (P<0.001) \) to response, as the more the clinical response rate, the higher the level of serum sCD4 (Table 5).

**Table 3:** Comparison of serum levels of serum soluble CD4 before and after injection with intralesional tuberculin purified protein derivative injection \( (N=49) \)

| sCD4 (ng/ml) | Before injection \( (N=49) \) | After injection \( (N=49) \) | \( t \) | \( P \) |
|-------------|-------------------|------------------|------|------|
| Minimum–maximum | 7.20–55.42 | 10.60–82.51 | | |
| Mean±SD | 18.47±10.4 | 39.48±20.67 | 5.386* | <0.001* |
| Median | 19.0 | 29.0 | |
| Change | ↑19.01±35.81 | | | |

\( * \) paired t test.

sCD4, serum soluble CD4.

\( P \): \( P \) value for comparing between sCD4 levels before and after injection.

*Statistically significant at \( P \) value less than or equal to 0.05.

**Table 4:** Comparison of age, number, duration and site of anogenital warts, and serum levels of serum soluble CD4 clinical response after intralesional tuberculin purified protein derivative injection \( (N=49) \)

| Variables | Clinical response | \( H \) | \( P \) |
|-----------|-----------------|------|------|
| Number of warts | No response \( (N=7) \) | Partial response \( (N=16) \) | Complete response \( (N=26) \) | Test of significance | \( p \) |
| Mean±SD | 26.71±1.98 | 19.56±6.87 | 13.69±3.66 | 34.744* | 0.021* |
| Duration of warts (months) | Mean±SD | 12.57±1.99 | 11.31±1.08 | 4.23±1.63 | 37.266* | <0.001* |
| Site of warts [\( n (% \)] | \( \chi^2 \) | \( \chi^2 \) test | | |
| Shaft of penis | 4 | 57.1 | 7 | 43.8 | 16 | 61.5 | 1.336 | 0.524 |
| Pubic area | 4 | 57.1 | 9 | 56.3 | 9 | 34.6 | 2.397 | 0.295 |
| Scrotum | 0 | 0.0 | 6 | 37.5 | 4 | 15.4 | 4.332 | 0.078 |
| Frenulum | 0 | 0.0 | 3 | 18.8 | 6 | 23.1 | 1.572 | 0.520 |
| Perianal | 0 | 0.0 | 2 | 12.5 | 1 | 3.8 | 1.504 | 0.715 |
| Serum soluble CD4 (ng/ml) | After PPD injection | F | | |
| Minimum–maximum | Mean±SD | 7.20–40.45 | 16.45–54.20 | 15.89–55.42 | 2.871 | 0.067 |
| Mean±SD | 18.86±11.82 | 18.34±9.22 | 16.11±11.88 | | |
| After PPD injection | Minimum–maximum | Mean±SD | 10.60–48.50 | 15.60–80.64 | 21.00–82.51 | 32.758* | <0.001* |
| Mean±SD | 27.73±8.82 | 32.84±11.46 | 49.23±18.37 | |

\( \chi^2 \), \( \chi^2 \) test; \( F \), analysis of variance test; \( H \), Kruskal–Wallis test; \( MC \), Monte Carlo; PPD, purified protein derivative; sCD4, serum soluble CD4.

\( P \): \( P \) value for association between clinical response and age, number, duration, site of warts and serum soluble CD4.

*Statistically significant at \( P \) value less than or equal to 0.05.
**Table 5:** Comparison of serum levels of serum soluble CD4 and clinical response after intralesional tuberculin purified protein derivative injection (N=49)

| sCD4 (ng/ml) | Clinical response | Test of | P     |
|--------------|-------------------|---------|-------|
|              | No response (N=7) | Partial response (N=16) | Complete response (N=26) | significance |       |
| Before PPD injection |                     |                     |                     | F     | P     |
| Minimum–maximum | 7.20–40.45         | 16.45–54.20         | 15.89–55.42         |       |       |
| Mean±SD       | 18.86±11.82        | 18.34±9.22          | 16.11±11.88         | 2.871 | 0.067 |
| After PPD injection |                   |                     |                     | F     | P     |
| Minimum–maximum | 10.60–48.50        | 15.60–80.64         | 21.00–82.51         | 32.758* | <0.001* |
| Mean±SD       | 27.73±8.82         | 32.84±11.46         | 49.23±18.37         |       |       |

F, analysis of variance test; PPD, purified protein derivative; sCD4: serum soluble CD4.
P: P value for association between clinical response and CD4.
*Statistically significant at P value less than or equal to 0.05.

**DISCUSSION**

Anogenital warts are a very common sexually transmitted condition in both males and females that cause marked quality of life deterioration especially sexual life and couple interpersonal communications. AGW are caused by HPV infection, which has more than 100 subtypes; the most common strains are 6 and 11 subtypes[15]. Immunotherapy is a promising new line of management of numerous and difficult-to-treat anogenital warts as it can cause complete wart removal with minimal invasion, injury, pain, scarring, and relatively short duration in some cases as well as they improve the host immune reactivity response against HPV, which leads to high recovery and little recurrence rates[13]. Immunotherapy is considered an economical therapeutic modality for extensive neglected wart infections and subsequently can be of extraordinary incentive in developing countries[16].

In the same line, Shaheen et al.[17] reported that PPD immunotherapy injection showed 60% clearance of target warts and distant warts with significant increase in circulating IL4.Nimbalkar et al.[13] was in agreement with our study, as they found that of 45 patients, 28 (62.2%) showed complete clearance, eight (17.8%) patients showed partial clearance, and nine (20%) patients showed no improvement. Higher results were reported by Elela et al.[18], who found that complete clearance was seen in 94.1% of the cases when intralesional PPD was administered in the wart tissue versus success rates of 96% with the intradermal injection.

On the contrary, Kus et al.[19] used intralesional tuberculin PPD injection in 18 patients with recalcitrant warts and found that complete cure was found in 5/18 (29%) patients, partial response was found in 10/18 (59%) patients, and no response was found in two (12%) patients. This cure rate is lower than the cure rate in our study and previously mentioned studies, and this can be explained by a small sample size, lower number of sessions, and wider interval of injections (3 vs. 2 weeks) in our study.

Cellular immunity has the main responsibility in fighting wart infection. Spontaneously recovering warts show exceptional epidermal and dermal release of CD4+ activated memory lymphocytes in comparison with nonrecovered warts. HPV proteins antibodies have been significantly found in the serum of HPV-infected patients, but the role of these antibodies is uncertain as they do not correlate with the wart regression[20]. Additionally, Critchlow et al.[21] found that defective T-cell-mediated
immune response can be the cause of prolonged and resistant infection with anogenital warts.

In 2017, Singh et al.[7] studied the functional characterization of HPV6 and HPV11 antigen-specific CD4+ and CD8+ T-cell responses from serum of patients with anogenital warts and reported that frequency of both CD4+ IFN-γ+ and CD8+ IFN-γ+ cells in patients with anogenital wart was significantly lower than healthy controls. In this study, they recommended that their findings may be helpful in designing future immunomodulation methods that can be used as adjunct immunotherapy to simulate host immune response in patients of anogenital warts. This might help in prevention of recurrence of warts after treatment[7].

In the current work, we measured the level of sCD4 in serum of patient with anogenital warts before and after intralesional tuberculin PPD injection. The level of serum sCD4 was significantly increased after than before injection as mean serum sCD4 level was 18.47±10.4 versus 44.48±20.67, respectively (P<0.001). Before tuberculin PPD injection, level of sCD4 was not significantly correlated to clinical cure rates, whereas after injection, serum sCD4 levels were significantly related (P<0.001) to clinical recovery rates, as the more clinical response rate, the higher the level of serum sCD4.

Rahim et al.[11] studied the relationship between serum levels of both sCD4 and sCD8 and genital wart activity, and they reported that level of sCD4 was lower in patients versus controls (5.47±0.12 vs. 10.98±0.19 ng/ml, respectively; P=0.001), which was opposite to the level of serum sCD8, which was higher in patients than control group (38.56±1.90 vs. 16.86±0.23 ng/ml, respectively).

Findings in our study were in concordance indirectly with results reported by Rahim et al.[11] and Singh et al.[7], with some difference in study design and management. In the current study, we were the first to measure the level of CD4 cells in patients with anogenital warts in relation to immunotherapy modality. We assumed that CD4 cells had an effective role in mechanism of action of intralesional tuberculin PPD as a treatment method for anogenital warts. This was proved by the higher levels of serum sCD4 before injection than after injection and significant statistical correlation between sCD4 level and rates of clinical response. Another clue for this hypothesis is regression of distant or neighboring warts in other sites rather than injection sites, which denotes that tuberculin PPD intralesional injection causes significant systemic immune reaction may be dependent on CD4 cells and causes wart recovery.

Limitations of the study were the small sample size, making the results difficult to be generalized; sensitivity of the vaccine as well as ease to be spoiled by heat and light, with continuous need to strict cold chain maintenance; bad patient compliance as intolerance to adverse effects and boring from repeated injections; shame and stigma of genital wart infections; difficulties in assessment of partners; and data regarding effect of immunotherapy with PPD and level of sCD4 being limited to be compared with our results.

CONCLUSION

Immunotherapy with intralesional tuberculin PPD is an effective, dependent economic modality of treatment of genital warts with good cosmetic results and lower recurrence rates. Serum sCD4 levels were higher in patients with anogenital warts treated with intralesional tuberculin PPD injection than pretreatment levels and were significantly correlated to wart clinical response rates.

CONFLICT OF INTEREST

1. Pretet JL, Charlo JM, Mougin C. Virological and carcinogenic aspects of HPV. Bull Acad Natl Med 2007; 191:611–623.
2. Lopaschuk CC. New approach to managing genital warts. Can Fam Physician 2013; 59:731–736.
3. Mahnke YD, Beddall MH, Roederer M. OMIP-017: human CD4+ helper T-cell subsets including follicular helper cells. Cytometry A 2013; 83:439–440.
4. Vender R, Bourcier M, Bhatia N, and Lynde C. Therapeutic options for external genital warts. J Cutan Med Surg 2013; 17 (Suppl 2): S61–S67.
5. Guo L, Zhang H, Chen B. Nivolumab as programmed death-1 (PD-1) inhibitor for targeted immunotherapy in tumor. J Cancer 2017; 8:410.
6. Chauhan P S, Mahajan V K, Mehta K S, Rawat R, and Sharma V. The efficacy and safety of intralesional immunotherapy with measles, mumps, rubella virus vaccine for the treatment of common warts in adults. Indian Dermatol Online J 2019; 10:19.
7. Singh M, Thakral D, Kar H K, Rishi N, Sharma P K, and Mitra D K. Distinct clinico-immunological profile of patients infected with human papilloma virus genotypes 6 and 11. Virusdisease 2017. 28:200–204.
8. Crotty S. T follicular helper cell differentiation, function, and roles in disease. Immunity 2014; 41:529–542.
9. Macdonald G. Harrison’s internal medicine, -by AS Fauci, DL Kasper, DL Longo, E. Braunwald, SL Hauser, JL Jameson and J. Loscalzo. Intern Med J 2008; 38:932–932.
10. Mohamed EM, El Taieb MA, Abd El-sabour GA. Intralesional vitamin D3 versus purified protein derivatives in the treatment of multiple cutaneous warts: comparative study. Egypt J Hosp Med 2019; 76:3589–3594.

11. Rahim S, Ibraheam IA, Radi NK. Evaluation of CD4 and CD8 in patients infected with genital wart caused by human papilloma virus in Babylon province/Iraq. J Glob Pharma Technol 2017; 9:372–375.

12. Nayak S, Achariya B. Mantoux test and its interpretation. Indian Dermatol Online J 2012; 3:2.

13. Nimbalkar, A., Pande, S., Sharma, R. and Borkar, M., Tuberculin purified protein derivative immunotherapy in the treatment of viral warts. Indian J Drugs Dermatol 2016; 2:19.

14. Jaisinghani, A.K., Dey, V.K., Suresh, M.S. and Saxena, A., Bacillus Calmette–Guerin immunotherapy for recurrent multiple warts: an open-label uncontrolled study. Indian J Dermatol 2019; 64:164.

15. Lynch MD, Cliffe J, Morris-Jones R. Management of cutaneous viral warts. BMJ 2014; 348:g3339.

16. Rajashekar TS, Amulya R., Sathish S, Kumar S. Comparative study of intralesional BCG and PPD in the treatment of multiple cutaneous warts. Indian J Clin Exp Dermatol 2018; 2018:1–6.

17. Shaheen, M.A., Salem, S.A.M., Fouad, D.A. and El-Fatah, A.A.A. Intralesional tuberculin (PPD) versus measles, mumps, rubella (MMR) vaccine in treatment of multiple warts: a comparative clinical and immunological study. Dermatol Ther 2015; 28:194–200.

18. Elela I, Elshahid AR, Mosbeh AS. Intradermal vs intralesional purified protein derivatives in treatment of warts. Golf J Dermatol Venereol 2011; 18:21–26.

19. Kus, S., Ergun, T., Gun, D. and Akin, O., Intralesional tuberculin for treatment of refractory warts. J Eur Acad Dermatol 2005; 19:515–516.

20. Majewski S, Jablonska S. Immunology of HPV infection and HPV-associated tumors. Int J Dermatol 1998; 37:81–95.

21. Critchlow, C.W., Hawes, S.E., Kuypers, J.M., Goldbaum, G.M., Holmes, K.K., Surawicz, C.M. and Kiviat, N.B., Effect of HIV infection on the natural history of anal human papillomavirus infection. AIDS 1998; 12:1177–1184.