Comparative Efficacy of the Combination of Topical Betamethasone Dipropionate and Calcipotriene with Betamethasone Dipropionate and Calcipotriene Alone in the Treatment of Localized Vitiligo.

Alam MN1, Wahab MA2, Husain MA3, Khondker L4, Ahmad GKA5, Chakraborty A6

Abstract

Vitiligo is an acquired skin disorder characterized by well-defined white patches that are often symmetrically distributed. The study was conducted to compare the efficacy of the combination of topical betamethasone dipropionate and calcipotriene with betamethasone dipropionate and calcipotriene alone in the treatment of localized vitiligo.

A clinical trial was carried out with the patients of vitiligo from January 2012 to August 2012. In group A, 20 patients were applied betamethasone dipropionate cream 0.05% and topical calcipotriene ointment (0.005%), in group B, 20 patients were applied betamethasone dipropionate cream 0.05% only; In group C, 20 patients were applied calcipotriene ointment 0.005% alone.

From baseline percentage reduction of total lesion in 1st follow up in group A, group B and group C were 20%, 15% and 10% respectively. At 3rd follow up in group A, group B and group C it was 50%, 37% and 30% and at 5th follow up, it was 80%, 75% and 65% respectively. ANOVA test was done and found significant difference of reduction of total lesion of vitiligo among the groups (p value < 0.05). At the completion of the study, each patient was separately graded the treated sides on a 6-point ordinal scale based on a global estimate of the change in vitiligo and found that, the very much improvement were 70%, 55% and 45%, much improvement were 20%, 30% and 35% and improvement were 10%, 15% and 20% in group A, group B and group C respectively. ANOVA test was found significant difference of success rate of patients of localized vitiligo among the groups (p value 0.005).

Both the drugs, calcipotriene and betamethasone dipropionate when used individually, were found to be equally effective in the treatment of vitiligo, but the combination of the two was found to be superior in efficacy.

Key words: Betamethasone dipropionate, calcipotriene, vitiligo.

Introduction

Vitiligo vulgaris is a pigmentary disorder with no universally efficacious therapeutic options. It is an acquired skin disorder that result from damage to and destruction of melanocytes.1 The cause is unknown but may involve genetic factors, autoimmunity, toxic metabolites and/or a higher vulnerability of melanocytes.2-4 It is affecting 0.5–2% of the population worldwide.5,6 Vitiligo can be cosmetically disfiguring and it is a stigmatizing condition, leading to serious psychologic problems in daily life.7,8 Two of the major theories of the pathogenesis of vitiligo are the autoimmune theory and the autocyctotoxicity theory.9 The autoimmune theory speculates that patients with vitiligo form autoantibodies against melanocytes. The existence of antimelanocyte surface antigen antibodies has been demonstrated, and the severity of vitiligo has proven to be related to the amount of antibodies present.10,11 Patients have numerous treatment options available, but none is universally effective. Even among patients who respond to treatment there is a high potential for relapse. Treatment of vitiligo is a challenge.12 The most widely prescribed therapies are PUVA and topical corticosteroids. PUVA is not recommended for children because of concern over its long term side effects, and prolonged use of topical steroids has the potential to cause cutaneous atrophy, including telangiectasia and perioral dermatitis.13

Calcipotriene is a synthetic analog of vitamin D3 (calcitriol: 1, 25(OH)D3) that has been shown to have immunomodulating and immunosuppressive actions.14 Receptors for 1, 25(OH)D3 the active form of vitamin D, have been demonstrated on keratinocytes, melanocytes and fibroblasts, and on immunologically active cells. Melanocytes and keratinocytes within vitiligenous lesions have shown defective calcium uptake. Based on this observation, recently investigators demonstrated that calcipotriene can be effective for vitiligo, both as monotherapy and in combination in Bangladeshi population.15,16

1. Corresponding Author: Dr. Mohd. Nurul Alam
   MBBS, FCPS, DDV, MCPS.
   Assistant Professor, Department of Dermatology & Venereology
   Ibn Sina Medical College & Hospital, Kallyanpur, Dhaka
2. Professor Lt. Col (Retd.) Dr. Md. Abdul Wahab
   MBBS, FCPS, DDV, MCPS, FRCP.
   Professor, Department of Dermatology & Venereology
   Bangabandhu Sheikh Mujib Medical University, Dhaka
3. Dr. Major (Retd.) Md Anwar Husain
   MBBS, DDV, MCPS.
   Associate Professor, Department of Dermatology & Venereology
   Ibn Sina Medical College & Hospital, Kallyanpur, Dhaka
4. Dr. Lubna Khondker
   MBBS, FCPS, DDV, MCPS, MPH.
   Assistant Professor, Department of Dermatology & Venereology
   Bangabandhu Sheikh Mujib Medical University, Dhaka
5. Dr. Gulam Kazem Ali Ahmad
   MBBS, FCPS, DDV.
   Specialist in Dermatology and Venereology
   Bangabandhu Sheikh Mujib Medical University, Dhaka
6. Dr. Anjana Chakraborty
   MBBS, DDV
   Specialist in Dermatology and Venereology
   Bangabandhu Sheikh Mujib Medical University, Dhaka
Materials and Methods

It was a clinical trial, from January 2012 to August 2012. Patients of vitiligo attending outpatient department of Dermatology and Venereology, Bangabandhu Sheikh Mujib Medical University (BSMMU), Dhaka were the study population and non-probability sampling method was followed in this study. Inclusion criteria were all localized vitiligo patients, age more than 5 years of both sexes. The exclusion criteria were generalized vitiligo patients, previous skin malignancy, treatment for vitiligo with corticosteroid agents, vitamin D analogues, or tacrolimus within the last 3 months, pregnancy and lactation, renal or hepatic disease, lupus erythematosus, severely ill patients and patients or attendants unwilling to take part in the study.

Procedures of collecting data

A total of 60 patients suffering from localized vitiligo were primarily selected from outdoor department of Dermatology & Venereology department, BSMMU. Complete history, general physical, dermatological and Wood’s light examinations were done for all enrolled patients. For women of reproductive age, reproductive history were carefully mentioned. History and physical findings were recorded in a structured questionnaire. Finally those patients, who were matched the inclusion and exclusion criteria according to history, physical examination and freely gave their informed consent, were selected for study. Clinical assessment at baseline and follow up visit, monthly for 5 months were taken and photographs of all lesions at baseline and after 5 months were taken for subsequent assessment and further comparison. The patients were randomized into three treatment groups. In group A, 20 patients were applied betamethasone dipropionate cream 0.05% in the morning and topical calcipotriene ointment (0.005%) in the evening, group B, 20 patients with betamethasone dipropionate cream 0.05% twice daily; In group C, 20 patients were applied calcipotriene ointment 0.005% similarly; and each individual lesion were treated daily for five months. Generally the efficacy of repigmentation therapy & safety were recorded from those used in previous studies, we performed global assessments, which were done halfway through and at the end of the study.

For each body region, the VASI was determined by the product of the area of vitiligo in hand units (which were set at 1% per unit) and the extent of depigmentation within each hand unit–measured patch (possible values of 0, 10%, 25%, 50%, 75%, 90%, or 100%). Any new depigmented patches that developed during the study were also estimated using the hand unit method and were included in the VASI calculation. Standardized assessments for estimating the degree of pigmentation to derive the Vitiligo Area Scoring Index. At 100% depigmentation, no pigment is present; at 90%, specks of pigment are present; at 75%, the depigmented area exceeds the pigmented area; at 50%, the depigmented and pigmented areas are equal; at 25%, the pigmented area exceeds the depigmented area; and at 10%, only specks of depigmentation are present. For the second measurement method, total body photographs were taken at baseline and at each monthly follow-up visit as an aid to the global clinical scoring. These 35-mm slides were used by investigators for global assessments, which were done halfway through and at the end of the study.

To compare the VASI system with ordinal scales analogous to those used in previous studies, we performed global assessments. At the completion of the study, each patient separately graded the treated sides on a 6-point ordinal scale based on a global estimate of the change in vitiligo as follows: complete improvement (100%), very much improved (76-99%), much improved (51%-75%), improved (26%-50%), minimal change (1%-25%), no change. Data analysis was performed by Statistical Package for Social Science (SPSS), and level of significance was measured by using appropriate statistical test and level of significance (p value) was set at 0.05 and confidence interval at 95%.

Results

Mean age of group A, group B and group C patients were $21.50 \pm 3.32$, $21.55 \pm 4.12$ and $22.25 \pm 4.67$ respectively. There was no significant difference of age between the groups.
Table I: Distribution of age of the patients by groups.

<table>
<thead>
<tr>
<th>Age (in years)</th>
<th>Groups</th>
<th>Groups</th>
<th>Groups</th>
<th>*p value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Group A (n=20)</td>
<td>Group B (n=20)</td>
<td>Group C (n=20)</td>
<td></td>
</tr>
<tr>
<td>13-22</td>
<td>7 (35%)</td>
<td>8 (40%)</td>
<td>7 (35%)</td>
<td>0.810</td>
</tr>
<tr>
<td>23-32</td>
<td>9 (45%)</td>
<td>6 (30%)</td>
<td>6 (30%)</td>
<td></td>
</tr>
<tr>
<td>&gt;32</td>
<td>4 (20%)</td>
<td>6 (30%)</td>
<td>7 (35%)</td>
<td></td>
</tr>
<tr>
<td>Mean ± SD</td>
<td>21.50±3.32</td>
<td>21.55±4.12</td>
<td>22.25±4.67</td>
<td></td>
</tr>
</tbody>
</table>

In distribution of the patients by main site of lesion in different groups, extremities involvement were 60%, 70% and 45%, face involvement were 30%, 20% and 35%, and trunk involvement were 10%, 10% and 20% in group A, group B and group C respectively. ANOVA test was done to measure significance and found no significant difference of main site of lesion among the groups (p>0.05).

In distribution of the patients by duration of lesion in different groups, less than 1 year duration were 45%, 70% and 35% and more than 1 year duration were 55%, 30% and 65% in group A, group B and group C respectively. ANOVA test was done to measure significance and found no significant difference of main site of lesion among the groups (p>0.05).

In the base line the score of vitiligo in group A, group B and group C were 26, 25 and 23 respectively. At 1st follow up it was 20, 22 and 19, at 3rd follow up it was 10, 15 and 12, and at 5th follow up it was 3, 8 and 6 in group A, group B and group C respectively. ANOVA test was done to measure significance and found significant difference of score of vitiligo among the groups (p<0.05).

Discussion

At the completion of the study, each patient was separately graded the treated sides on a 6-point ordinal scale based on a global estimate of the change in vitiligo and found that, the very much improvement were 70%, 55% and 45%, much improvement were 20%, 30% and 35% and improvement were 10%, 15% and 20% in group A, group B and group C respectively. ANOVA test was found significant difference of success rate of patients of localized vitiligo among the groups (p value 0.005).
and 35%, improvement were 10%, 15% and 20% in group A, group B and group C respectively (p value 0.005). These findings were consistent with other studies like Kumaran et al and Parsad et al. A trial was conducted by Kumaran et al to evaluate the effect of topical calcipotriol ointment (0.005%) and betamethasone dipropionate (0.05%) cream, given alone or in combination, in treatment of localized vitiligo. Group I patients were treated with betamethasone dipropionate (0.05%) cream twice daily. Group II patients were treated with calcipotriol ointment (0.005%) twice daily, and group III with betamethasone dipropionate (0.05%) in the morning and calcipotriol (0.005%) in the evening. Marked (50% to 75%) repigmentation was observed in 2 (13.3%), 1 (6.7%) and 4 (26.7%) patients in groups I, II and III, respectively and moderate (25-50%) repigmentation was observed in 7 (46.7%), 5 (33.3%) and 7 (46.7%) patients in groups I, II and III, respectively. They concluded that combined therapy appeared to give a significantly faster onset of repigmentation along with better stability of the achieved pigmentation. Parsad et al enrolled twenty-one patients age 5 to 17 years with vitiligo in this study and the children were advised to apply calcipotriol 50 microg/g in the evening and expose themselves to sunlight the next day for 10 to 15 minutes. Initial repigmentation occurred in the majority of children after 6 to 12 weeks of treatment. Marked to complete repigmentation was seen in 10 of 18 patients and four patients showed moderate improvement.

Our study observed that both the drugs, calcipotriol and betamethasone dipropionate when used individually as monotherapy, were found to be equally effective in the treatment of vitiligo, but the combination of the two was found to be superior in efficacy. These findings of our study were not consistent with study findings of Chiavérimi et al. Chiavérini et al carried out a study to evaluate the efficiency of topical calcipotriol monotherapy in vitiligo and concluded that topical calcipotriol in monotherapy is not an effective treatment of vitiligo. Our study findings were also consistent with findings of Parsricha et al. In my study, both the drugs, calcipotriol and betamethasone dipropionate when used individually, were found to be effective in the treatment of vitiligo. Parsricha et al tried a new approach using mini-pulse therapy with betamethasone. Forty patients having extensive and/or fast-spreading vitiligo were given 5 mg betamethasone/dexamethasone as a single oral dose after breakfast on 2 consecutive days per week. Within 1-3 months, progression of the disease was arrested in 89% of the 36 patients having active disease, while 2 patients needed an increase in the dose to 7.5 mg per day to achieve complete arrest of lesions. Within 2-4 months, 80% of the patients started having spontaneous repigmentation of the existing lesions which progressed with continued treatment. The extent of repigmentation varied in different patients and even in different lesions in the same patient. It was less than 10% in 14 (35%) patients and almost complete (> 90%) in three patients. Oral mini-pulse therapy with betamethasone seems to be an effective treatment modality to arrest the progression of vitiligo. It also induces spontaneous repigmentation.

Both the drugs, calcipotriene and betamethasone dipropionate when used individually, were found to be equally effective in the treatment of vitiligo, but the combination of the two was found to be superior in efficacy.

References

14. Lisa BT, Nanette BS. Calcipotriene and Corticosteroid


