The Highly Effective Use of Topical Zinc Pyrithione in the Treatment of Psoriasis: A Case Report
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Introduction
Psoriasis affects approximately 20% of the population. The clinical course is dynamic, capricious, and replete with flares and remissions.

Currently there are many good anti-psoriatic treatment choices including topical agents (steroids, tars, anthrallin, and calcipotriene), phototherapeutics (UVB, PUVA), and systemic treatments (methotrexate, cyclosporine, and etretinate). Often a combination of treatment modalities is required for clinical success. Unfortunately, phototherapeutic modalities can be labor intensive and inconvenient. Systemic treatments have significant side effect considerations. Consequently, any effective and safe topical agent is a welcomed addition to our therapeutic armamentarium. We present here a report of a case using a novel topical preparation of zinc pyrithione for the treatment of psoriasis. Topical zinc pyrithione appears to be a safe and effective treatment for psoriasis.

We have recently become aware of a new anti-psoriatic treatment containing zinc pyrithione (which is also recognized as the active ingredient in a major anti-dandruff shampoo). The effectiveness of zinc pyrithione to treat seborrheic dermatitis and psoriasis has been well documented. Since some believe that seborrheic dermatitis and psoriasis may lie on different ends of the same spectrum, it seemed reasonable to try a topical preparation of zinc pyrithione to treat psoriasis. The mechanism of action of zinc pyrithione on psoriasis and seborrheic dermatitis has been reported to be anti-proliferative via DNA interactions, anti-yeast, antiseptic, and keratinolytic mechanisms.

We report here the clinical results of a psoriatic patient treated with a topical spray containing zinc pyrithione.
Methods

A 39 year old man with a 14 year history of moderate plaque psoriasis was instructed to spray the topical zinc pyrithione preparation, twice per day, to the left elbow. The right elbow was to be used as a control and received no treatment of any kind during the trial. The treatment preparation contains zinc pyrithione (0.25%) in a vehicle containing isopropyl myristate.

Results

Figure 1 depicts the baseline condition of the patient's elbows before trial initiation. Figure 2 depicts the patient's elbows 3 weeks after applying the zinc pyrithione preparation to the left elbow only. At the end of the 3 week treatment period the left (treated) elbow was essentially clear. The right (control) elbow remained virtually unchanged, if not slightly worse. The patient also reported a complete disappearance of pruritus approximately 3 days into treatment, on the zinc pyrithione preparation treated plaque only, that was sustained throughout the treatment period.

Discussion

The result in this case is impressive. There was marked reduction in scale, erythema, and pruritus. The patient reported no significant side effects during and after the treatment period. At a 6 week follow up the patient remained lesion free on the treatment plaque, with no additional applications. Perhaps a maintenance treatment plan (e.g. BID on weekends only), or the immediate re-treatment of any new/ recurring plaque may reduce the frequency and/ or severity of future flares.

Although the exact mechanism of action of the "activated" zinc pyrithione preparation is unknown it may be speculated that the possible anti-proliferative mechanism of action of zinc pyrithione may involve the regulation of DNA transcription factors containing "zinc finger" binding domains. It is also well recognized that many enzymes require the binding of metal ions for activation. Perhaps some of these (zinc requiring) enzymes or transcription factors play a key role in the regulation of cellular proliferation. It is also well known that a deficiency of zinc produces a disease state (acrodermatitis enteropathica) that includes psoriasiform lesions. Perhaps the vehicle or the activated form of
zinc pyrithione permits a physiologic level of zinc to be reached in the target cells, either epidermal and/or lymphocytic. The combination of zinc pyrithione with the other vehicle including isopropyl myristate may also have significant additional influence on clinical effectiveness.

In this preliminary report, an aerosol preparation of zinc pyrithione (0.25%) in a vehicle containing isopropyl myristate appears to be a safe and effective treatment for psoriasis. Each ml of the liquid preparation will cover an area the size of a palm, approximately 6-8 times.

Because of the promising results in this case report and a plethora of anecdotal reports, we are initiating a 60 patient, double-blind, vehicle-controlled clinical study evaluating the use of topical zinc pyrithione for the treatment of psoriasis. If additional controlled research confirms these encouraging results, topical zinc pyrithione may represent one of the major advances in the treatment of psoriasis.

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**References**


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