Crusted ("Norwegian") scabies in a specialist HIV unit: successful use of ivermectin and failure to prevent nosocomial transmission

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A nosocomial outbreak of scabies in a specialist inpatient HIV unit resulted from a patient admitted with crusted scabies. Treatment of his infestation with topical scabicides alone failed and he remained infectious for several weeks. His infestation was then eradicated with combined topical treatment and oral ivermectin. In total, 14 (88%) out of 19 ward staff became symptomatic, and 4 (21%) had evidence of scabies on potassium hydroxide examination of skin scrapings. The ward infection control policy was changed to distinguish patients with crusted scabies from those with ordinary scabies. A second patient with crusted scabies was treated with combined oral and topical therapy early in his admission and nursed with more stringent isolation procedures. No nosocomial transmission occurred and his infestation responded readily to treatment. Patients with crusted scabies require strict barrier nursing if nosocomial transmission is to be avoided. Ivermectin combined with topical scabicides may be a more efficacious treatment than topical scabicides alone in such patients.

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Introduction
Scabies is a frequent problem among patients infected with human immunodeficiency virus (HIV). Patients unable to mount an adequate immune response to the infestation develop atypical rashes, often involving large areas of the body, and are considerably more infectious than patients with a typical rash. The most extreme form is crusted, so-called Norwegian scabies, which was first reported in association with HIV in 1986. Treatment failure is frequently encountered when topical scabicides are used to treat crusted scabies, owing to poor penetration of the crusted areas. Ivermectin is an oral alternative to topical treatment that has been successfully used to treat patients with both ordinary and crusted scabies.

Six reports of nosocomial transmission of scabies have been published in which the index cases were HIV-infected patients with crusted scabies. In most cases misdiagnosis of the rash predisposed to nosocomial transmission. This report describes an outbreak on an HIV ward that occurred despite immediate diagnosis, and discusses appropriate preventative measures and the response of the index case and a second patient to combined topical treatment and ivermectin.

Case 1: Index Case
The index case was a 45 year old white homosexual male who was admitted for management of worsening HIV encephalopathy and a rash. He had a previous history of astasia with associated eczema. Scabies had been diagnosed 5 months previously and treated with malathion with worsening of the eczema. The CD4+ lymphocyte count was 0-11 (NR = 0-35-2-2) × 10^3/l. On admission, the patient had widespread popular erythema with excoriations over most of the body with areas of thick crust particularly over the scalp and beard area. Scabies was suspected by the admitting nurses, and the patient was isolated; microscopic examination of skin scrapings treated with 10% potassium hydroxide subsequently confirmed the diagnosis.

He was treated on days 1 and 3 of admission with topical malathion (Derbac M). Following this the erythema and itch increased and live scabies mites were found still to be present. Because of the lack of effect of topical treatments and the distress produced by treatment, further topical treatment was delayed for 12 days until most of the crusted areas of skin had been removed with dilute chlorhexidine gluconate solution. Topical applications of carbaryl (Derbac C) on five occasions over the following 14 days failed to clear the infestation. Oral ivermectin (200 mcg/kg) was then given on a named patient basis, and two courses of topical malathion were applied; subsequent skin scrapings were negative and the general appearances of the skin improved. In total, the patient's admission lasted 51 days, and he was infectious for at least the first 29 days.

Secondary cases
A total of 19 nursing and medical staff worked on the ward during the index patient's admission, of whom 16 (14 nurses, 2 doctors) had direct physical contact with him. Of the staff with direct patient contact, 14 (88%) developed itching within the first 14 days of his
admission. None of the other patients on the ward, and none of the three staff who did not have contact with the index patient became symptomatic. In 12 staff itching lesions were limited to the forearms and/or hands. Of the 14 affected staff, 9 were seen by the hospital dermatologists; 2/9 (22%) had definite scabies with identification of mites, 2 (22%) had skin scrapings suggestive of scabies with typical egg shells but no mites seen, and 5 (56%) had normal skin scrapings. Several members of staff had empirically treated themselves with scabicides before seeing the dermatologist.

**Infection Control Measures**

On admission the index patient was isolated in a single room. Staff attending him wore gloves and plastic aprons to minimise skin to skin contact. When secondary transmission had been confirmed, all symptomatic staff and their partners treated themselves irrespective of parasitological confirmation. Further spread did not occur. The failure of infection control led us to review our policy for dealing with crusted scabies in hospitalised patients.

**Changes to our infection control guidelines resulting from the outbreak**

We now distinguish between crusted and ordinary scabies in our infection control guidelines. Staff and visitors attending patients with crusted scabies now wear gowns rather than aprons, in addition to gloves, until skin scrapings have been shown to be negative. Skin scrapings from the patient are repeated when the patient becomes asymptomatic and again 2 and 4 weeks after treatment. The infectious period is minimised by using both ivermectin and topical scabicide as initial treatment. Staff members who inadvertently have skin to skin contact with a patient with crusted scabies are advised to treat themselves prophylactically with a topical scabicide. Staff members who develop symptoms of rash or itching are seen urgently by the hospital dermatologists.

The patient’s room is regularly damp dusted to decrease environmental mites. The patient’s linen is treated as infectious.

**Case 2**

A 38 year old white homosexual man with a CD4+ lymphocyte count of 0-01 (NR

0-35–2-2) × 10⁹/l was admitted with a pruritic, papular rash on his trunk and limbs, with crusting on the hands, feet and buttocks (fig). Skin scrapings from the hands showed numerous scabies mites. The patient was isolated and treated with oral ivermectin (120 mcg/kg) together with a topical application of malathion. His itching resolved within 24 hours and his rash within 10 days of treatment. The patient was attended by a total of 2 doctors and 9 nurses. No nosocomial transmission of scabies was attributed to this admission.

**Discussion**

Crusted scabies is caused by the same mite, *Sarcoptes scabei*, that causes typical scabies in otherwise healthy individuals, and the difference in the two forms results from the host response.¹ The highly contagious nature of crusted scabies reflects the high number of mites required to produce the diffuse, erythematous, crusted rash. Until the HIV epidemic crusted scabies was most commonly reported in elderly patients, many of whom were institutionalised and debilitated for a variety of reasons. HIV infection is an increasingly frequent risk factor.²

Nosocomial transmission from HIV seropositive patients with crusted scabies has been reported on a larger scale than the outbreak described here, with transmission to other patients, relatives, and ancillary staff such as radiographers, laundry staff and chaplain in addition to nurses and doctors.³ Crusted scabies is considerably more contagious than ordinary scabies and brief skin to skin contact with a patient who has crusted scabies may be sufficient for transmission to occur. In addition, patients with crusted scabies may remain infectious for long periods because of difficulty eradicating the infestation from heavily crusted areas of skin, even when repeated applications of scabicides are used together with keratolytic agents. The use of oral ivermectin for the treatment of human ectoparasite infestations is a relatively recent development.¹ Ivermectin is an effective treatment of ordinary scabies without the addition of topical agents;³ however, treatment failure occurred in 1 of 2 patients with crusted scabies and AIDS treated with ivermectin alone.⁴ Our patients tolerated ivermectin well, and had a rapid response to ivermectin combined with topical malathion with no recurrence in either case. A rapid response to ivermectin and topical salicylic acid in 2 HIV negative patients with crusted scabies has been recently reported.⁵ At present ivermectin can only be obtained in the UK on a named patient basis.

The extent of our outbreak was probably

*Hands of patient 2 showing crusted scabies.*
limited by immediate diagnosis and isolation of the patient, but transmission still occurred to staff who had direct contact with him, despite use of aprons and gloves. Forearm skin contact while lifting the patient may have occurred. Of the nine symptomatic staff who were dermatologically assessed, four (44%) had microscopic evidence of scabies infection. Itching in staff with negative skin scrapings may have been psychological in origin, but in some may have represented allergic reactions to immature mites following prior sensitisation to mite antigens.

The occupational health service must be prepared to investigate outbreaks without delay; this requires rapid access to specialist dermatology services for microscopic confirmation of the diagnosis in early cases. In the case of an established outbreak, treatment of all staff and patients who have had contact with the index case may be warranted, although there are currently no guidelines concerning the treatment of asymptomatic staff, or symptomatic staff without a definite diagnosis. Our experience has been that without such measures concerns remain among ward staff about continued transmission.

The treatment of crusted scabies with topical scabicides often requires multiple applications and early use of ivermectin orally should be considered. Nosocomial transmission of scabies from AIDS patients is a risk, especially on HIV units. The risk of transmission is greatly increased in the case of crusted scabies and guidelines to prevent and respond to nosocomial transmission should be in place.