A Case of Pseudochromohidrosis Cured with Oral Erythromycin and Topical Clindamycin

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Received Date: 14-07-2021; Accepted Date: 16-08-2021; Published Date: 24-08-2021

Abstract

Apocrine chromohidrosis and eccrine pseudochromohidrosis are both rare skin presentations. In case of former patient presents with coloured sweats in apocrine distributions usually axilla, breast etc., while later presents with coloured sweat in eccrine distribution.

Such cases need to be reported after other possibilities with similar presentations have been ruled, and need special mention in literature as mostly they are misdiagnosed as factitious.

Hereby, I am also reporting one such case of pseudochromohidrosis which needed attention as it was causing anxiety to both patient and family members and its proper diagnosis and treatment brought a great relief to them.

Keywords

Chromohidrosis; Pseudochromohidrosis; Rare Apocrine; Eccrine; Coloured Sweat

Introduction

Chromhidrosis and pseudochromhidrosis are rarely reported skin conditions. Chromhidrosis is the production of colored sweat from sweat glands containing lipofuscin pigments and mostly
apocrine in origin and pseudochromhidrosis is production of colorless sweat, which becomes
colored when it reaches the skin and comes in contact with extrinsic agents such as
chromogenic bacterial products, fungi, dyes, paint, and colored foods [1-3]. Rarely,
chromohidrosis of eccrine origin may be due to excessive amount of colouring agents or dyes
consumed in diet and later secreted in sweat.

Through this case report, the author wants to draw the attention of readers towards an
extraordinary case of pseudochromohidrosis, as such cases often lead to social isolation
because of fear of humiliation among children.

A nine year old girl, resident of Varanasi, presented to the skin outpatient department of
Heritage Institute of Medical Sciences (HIMS), Bhadvar, Varanasi with complaints of sudden
onset reddish discoloration of her face starting from under eyelids downwards on the face and
neck with progressively advanced area of involvement in each episode. The patient’s skin and
mucosa were normal on examination. Patient had no discoloration of axillary sweat. Patient’s
condition had been previously diagnosed as irritant contact dermatitis and systemic lupus
erthematosus and she had received oral and topical steroid and hydroxychloroquin
respectively without any success in treatment. Further, history revealed by her mother
emphasized that patient developed red discoloration when she played or sweated for some
time.

History of any deliberate application of coloured items on face was negative. Patient’s mother
was requested for a video call when such lesions appeared, a video recording of same to be
done and to wipe lesions with alcohol swab, in the same video.

Patient’s photograph has also been taken with red discolouration on face (Fig. 1) (For details,
check: https://athenaeumpub.com/wp-content/uploads/JDR-02-026.mp4)

Patient’s family members never suffered such illness in past. Investigations including
antinuclear antibody, coagulation profile and urine examination was normal. Microbiological
examination of red colour material showed no growth.

The condition was finally diagnosed as pseudochromhidrosis. Treatment advised was
erthromycin tablet 500 mg twice daily for 5 days and topical clindamycin gel twice daily on
whole face for 1 month.

Patient did not develop any discoloration in follow up visits till 2 months after the last day of
treatment.
Discussion

This patient had pseudochromhidrosis can be concluded from the fact that she developed such discoloration only on face and patient had no history of any dietary intake of any coloured food or any drug or dye or any topical application, and face being a habitat for corneybacterium and other such bacteria which can cause sweat discoloration.

Further, negative ANA report and absence of photosensitivity and systemic symptoms ruled out systemic lupus erythematosus.

In a similar report Koley and Mandal didn’t find any growth of microorganism in culture or microscopic examination [4].

Thami and Kanwar have also reported such case of red pseudochromhidrosis on face in which no growth of microorganism was seen [1].

Ohanna Eva Constance Burggraaff and colleagues in 2019 reported a case of pseudochromhidrosis and they found no growth of organisms in microbiological examination nor did they find lipofuscin granules in histopathology of lesional skin biopsy [5].

A similar case of blue pseudochromhidrosis was reported by Christoffer Aam Ingvaldsen and his colleagues in 2020 but in this case growth of Bacillus cereus was seen in microbiological examination which is a pigment producing bacillus [6].
The absence of growth of any organism in various studies and ours can be explained by the fact that many commensal organisms are also present on face and may interfere with the growth of concerned organisms.

But, we should always advise microbiological examinations in such extraordinary cases to establish correct diagnosis and report them.

Moreover, pseudochromhidrosis is a diagnosis of exclusion as it responds to antibiotics oral and topical unlike chromhidrosis in which treating physician focuses on reducing sweat production.

**Conclusion**

Though, in our case it was quite difficult to differentiate between chromhidrosis and pseudochromhidrosis, but response of later to macrolide antibiotics as erythromycin and azithromycin, indicated that, there are high chances of cornebacterium, being a culprit in such cases.

The purpose of reporting this case is to draw the attention of dermatologists to such cases which mimics factitious diseases and many a times is a cause of emotional trauma due to social stigma of going in public.

**References**