Frequency of chronic venous insufficiency in patients of Schamberg’s disease

Yousuf Abd Mallick, Sidra Akhlaq Hussain*

Department of Dermatology, The Indus Hospital, Main Campus-Korangi, Karachi
* Department of Dermatology, Abbasi Shaheed Hospital, Karachi

Abstract

Objective To determine the frequency of chronic venous insufficiency in patients of Schamberg’s disease.

Methodology The study was cross-sectional & conducted in Dermatology outpatient department at Saifee Hospital Trust, Karachi, Pakistan. The duration of study was eleven months i.e. from April 2017 to March 2018. Clinically diagnosed, 60 cases of Schamberg’s disease were included via non probability consecutive sampling technique. Age of patients was 18 years and above. Clinical examination for varicose veins and Doppler ultrasonography of the lower extremities were performed and findings were recorded.

Results In our study, the mean age was 44.43±13.83 years. Out of 60 patients, 27 (45%) were male and 33 (55%) were female. Only 13 (21.67%) patients had a positive family history of Schamberg’s disease. In our study, 18 (30%) patients had clinically detected varicose veins of lower limbs while variable degrees of venous insufficiency were detected in 28 (46.67%) patients on Doppler ultrasound of the lower extremities.

Conclusion The number of patients with chronic venous insufficiency in Schamberg’s disease is significantly higher and it might be one of the main predisposing factors for this disease. However, further case-control studies are required to calculate its prevalence and to confirm its aetiopathogenic association.

Key words Schamberg’s disease, chronic venous insufficiency, Doppler ultrasound, frequency, varicose veins.

Introduction

Schamberg’s disease (SD) is also known as Purpura Pigmentosa Progressiva.1 World-wide Schamberg’s disease is the most common type of pigmented purpuric dermatoses (PPD).2 PPD represents a rare group of diseases which clinically manifest as various forms of purpura and pigmentation on the skin. Despite its rarity, PPD have been reported in all races and ethnicities across the globe. Domenico Majocchi (1849–1929), an Italian dermatologist, described the first ever PPD in 1896 as “purpura annularis telangiectaticum” which later on was given the name- Majocchi’s disease.3 After that many other types of PPD were reported and described in the literature. The current list now includes Schamberg’s disease, Majocchi’s disease, lichen aureus, pigmented purpuric lichenoid dermatitis of Gougerot and Blum, itching purpura of Lowenthal and eczematid-like purpura of Doucas and Kapetanakis.4,5 Unilateral variant of SD, linear pigmented purpura and
granulomatous variant of pigmented purpura are amongst the very rare types.\(^4\)

All these disorders appear similar on histopathology and share common features which include narrowing of the lumen and endothelial swelling of superficial cutaneous blood vessels, extravasation of erythrocytes, perivascular T lymphocyte infiltration and marked haemosiderin deposition in dermis and dermal macrophages.\(^5\)

The hallmark of Schamberg’s disease is progressive, asymptomatic and non-palpable pigmentary changes.\(^6\) Clinically, SD manifests as orange-red flat macules and patches of variable sizes with diagnostic “cayenne pepper” spots on the borders. Old lesions transform to yellow-brown colour. These lesions are oval or irregular in outline and pinpoint petechiae usually found inside the patches.\(^5\) SD has a very chronic course, sometimes it is of life-long duration. The disease continues to appear in successive crops over many years mostly at same places. The site of predilection is lower extremities but it can occur on trunk, thighs, buttocks; even arms and forearms can be affected.\(^1,5\) SD is completely asymptomatic, mostly occurs in adult population but it can also affect children\(^7\) and males are more often affected than females.\(^6\) \((\text{Figure 1 & 2})\)

There is no definitive treatment, neither cure is available for SD. Topical and oral corticosteroids, phototherapy, PUVA, photodynamic therapy,\(^8\) intense pulsed light (IPL),\(^9\) griseofulvin, pentoxifylline,\(^7\) cyclosporine, oral rutoside with ascorbic acid,\(^4\) colchicine\(^7,10\) and fractional non-ablative 1540nm erbium:YAG laser\(^11\) have been used with variable success. Sometimes, the pigmentation is resistant to all known treatments and besides psychological support, nothing can be offered by physicians and dermatologists.

\(46\)
was designed to determine the frequency of chronic venous insufficiency (CVI) in patients of Schamberg’s disease.

**Patients and Methods**

This study was conducted in OPD of Dermatology department, Saifee Hospital Trust, Karachi. The data collection was done for 11 months from April 2017 to March 2018. The cases were selected via non probability consecutive sampling technique. We took a sample of 60 adult subjects meeting the selection criteria for this study after written informed consent.

Clinically diagnosed, untreated cases of Schamberg’s disease for at least 6 months, of either sex group, age 18 years and above, with or without family history of SD were selected for the study. Patients with lower limb injuries, congenital and skeletal abnormalities, vascular malformations, malignancies, pregnant & lactating mothers and patients under treatment for SD were excluded.

SD is diagnosed clinically and then detailed clinical examination for varicose veins was performed in the OPD in lying down and standing positions. Trendelenburg and Tourniquet tests were performed. The Doppler ultrasound (DUS) of venous systems of both lower limbs was performed in ultrasound department by using 7.5 MHz Doppler ultrasound probe. The room temperature was maintained between 18°C and 22°C throughout the procedure. The standard protocol was followed. This consists of three steps:

1. DUS was first performed in orthostatic position.
2. Then DUS examination of the superficial veins of the lower limbs was performed in standing position. The patient was standing upright on a stool and the limb under examination was rotated outward.
3. Then DUS was performed in supine position. The patient was lying on a padded bed.

All steps were followed in all patients. All findings were recorded on a specified form by the researcher. Permission from the institutional ethical review committee was taken prior to the conduction of the study.

The data feeding and analysis was performed on computer package SPSS (statistical package of social sciences) version 21.0. Clinical characteristics were summarized in terms of frequencies and percentages for qualitative variables (gender, family history, clinically varicose veins and Doppler findings). Mean±S.D for quantitative variable (age) was done. Stratification was done with regard to gender, family history of SD and clinically varicose condition to see the effect of outcome and post-stratification Fisher Exact test applied. *p* <0.05 was considered as significant.

**Results**

Out of 60 patients, 27 (45%) were male and 33 (55%) were female. The mean age was 44.43±13.83 years. Only 13 (21.67%) patients had a positive family history of Schamberg’s disease. In our study, 18 (30%) patients had clinically detectable varicose veins of both lower limbs. Doppler ultrasound (DUS) of the lower extremities showed variable degrees of venous insufficiency in 28 (46.67%) patients while 32 (53.33%) patients showed normal DUS findings. *p*-value was <0.001.

Out of these 28 patients; 23 (82.14%) showed insufficient superficial venous system and 5 (17.85%) patients showed insufficiency of both deep and superficial venous systems while none
of the patients showed isolated insufficient deep venous system. All those 18 (30%) patients who have clinically detected varicose veins of lower limbs showed signs of CVI on DUS. So, we divided patients into 3 groups based on DUS findings; first with insufficient superficial venous system (superficial group), second with insufficient both deep and superficial venous systems (both group) and third with normal DUS findings group.

Stratification for gender with respect to DUS findings showed that 10 males (43.5%) and 13 females (56.5%) had insufficient superficial venous system; 3 males (60%) and 2 females (40%) had insufficient both deep and superficial venous systems; and 14 males (43.8%) and 18 females (56.3%) showed unaffected or normal venous systems. $p$-value was 0.854.

Stratification for family history with respect to DUS findings showed that 7 (30.4%) patients had positive while 16 (69.6%) patients had negative family history of SD in the superficial group. 1 (20%) patient had positive while 4 (80%) patients had negative family history of SD in both groups. 5 (15.6%) patients had positive while 27 (84.4%) patients had negative family history of SD in the normal DUS findings group. $p$-value was 0.365.

Stratification for clinically varicose condition with respect to DUS findings showed that 14 (60.9%) patients had and 9 (39.1%) patients did not have varicose veins on examination in the superficial group. 4 (80%) patients had and 1 (20%) patient did not have varicose veins on examination in both group. In the normal DUS findings group, all 32 (100%) patients had normal examination and none of the patients showed clinically detectable varicose veins. $p$-value was <0.001.

**Discussion**

In 1901, Schamberg’s Disease (Purpura Pigmentosa Progressiva) was first described\(^\text{17}\) as a disorder which causes discoloration of the skin, predominantly of the lower extremities but it can also involve buttocks, trunk and even upper extremities. The exact cause of SD still remains unknown.\(^\text{17}\) Multiple cofactors are reported in the literature which are thought to be associated with SD. The most important cofactor is venous hypertension.\(^\text{20}\) Other important ones are orthostatic pressure, recurrent lower limb infections, regular strenuous exercises, capillary fragility, ingestion of some chemicals and various drugs.\(^\text{16,20}\) The pathological manifestations of chronic venous insufficiency (CVI) develop as a consequence of venous hypertension.\(^\text{21}\) First of all tissue edema develops, followed by red cell extravasation and perivascular fibrin deposition leading to impaired arterial flow along with some other local immune disturbances. There is also concurrent damage to the lymphatic system which leads to impaired clearance of cellular metabolites.\(^\text{21}\) Hence hemosiderin deposition becomes a predominant feature of CVI.\(^\text{11}\)

The earliest clinical manifestation of venous hypertension is lower limb edema,\(^\text{21}\) followed by venous stasis, venous eczema, trophic skin lesions, pigmentation, lower limb pain and variable degrees of varicosities.\(^\text{19}\) Other serious and mostly irreversible cutaneous complications which develop subsequently in patients of CVI are atrophie blanche, lipodermatosclerosis, venous ulcers, livedoid vasculopathy, PPD and Pseudo-Kaposi’s sarcoma (Acroangiodermatitis).\(^\text{21}\) Most of the venous ulcers are caused by incompetence of the superficial venous system of the lower limb.\(^\text{22}\)

There is a list of systemic disorders which are found to be associated or reported with SD. It
includes diabetes mellitus, thyroid disorders, hereditary spherocytosis, rheumatoid arthritis, systemic lupus erythematosus (SLE), osteomyelitis, hepatic diseases & porphyria, hematological disorders and various malignancies. But their exact relation and contribution towards disease development is yet to be identified. Furthermore, psychological issues such as anxiety and depression are also common in patients suffering from CVI and its complications. The recurrent nature and resistant to treatment course of the disease upset the mental well-being of sufferers and make them vulnerable to various psychological diseases.

There is generally a male predominance in Schamberg’s disease but our study showed a subtle female dominance i.e. 55%. Similar results were also reported by Kim et al. and Cho et al. from Korea. We reported a positive family history of SD in 21.67% which no other study has documented yet. It indicates that some genetic predisposition to this disease also exists. Our study reported varicose dilatation in 30% of patients. Gönül et al. reported this in 20% of patients. We reported venous insufficiency in 46.67% of patients. Gönül et al. also reported this in 75% of patients.

To the best of our knowledge, this is the first study in Pakistan which reports chronic venous insufficiency by Doppler ultrasound in patients of SD. Detection of varicose veins clinically, and early detection of CVI on DUS necessitates management of venous hypertension and referral to the vascular surgeon. Early referral, multidisciplinary team care and prompt treatment may save patients of SD not only from serious and irreversible cutaneous complications but also from psychological diseases like anxiety and depression.

Conclusion

Schamberg’s disease is reported as a consequence of venous disease for decades but little is known about its aetiology and pathogenesis. We have reported a high frequency (46.67%) of chronic venous insufficiency among patients of Schamberg’s disease. It might be the main predisposing factor for this disorder. Further multi-centre, large scale studies are required to build its aetiopathogenic role in development of Schamberg’s disease.

References