Degos disease – malignant atrophic papulosis or cutaneointestinal lethal syndrome: rarity of the disease

Eduardo Pirolla
Felipe Fregni
Irene K Miura
Antonio Carlos Misiara
Fernando Almeida
Esdras Zanoni

Spaulding Rehabilitation Network Research Laboratory, Harvard Medical School, Boston, MA, USA; 2University of São Paulo, Sirio Libanes Hospital, São Paulo, Brazil; 3Sirio Libanes Hospital, São Paulo, Brazil

Background: Degos disease is a very rare syndrome with a rare type of multisystem vasculopathy of unknown cause that affects the skin, gastrointestinal tract, and central nervous system. Other organs such as the kidneys, lungs, pleura, liver, heart, and eyes, can also be involved.

Objective: To highlight the incidence of Degos disease with regard to age and sex, discuss the necessity of its accurate and early diagnosis, and demonstrate the most current techniques for its diagnosis; to discuss whether early therapeutic intervention can impact patient prognosis; and to present a literature review about this disease.

Design: With a retrospective, observational, nonrandomized trial, we described the evolution of the different forms of Degos disease and referenced the literature.

Data sources: Research on rare documented cases in the literature, including two cases of potentially lethal form of the disease involving the skin and gastrointestinal system and, possibly, the lungs, kidneys, and central nervous system. A case of the benign form of the disease involving the skin was observed by the authors.

Main outcome measures: Differences between outcomes in patients with the cutaneointestinal form and skin-only form of the disease. There was one fatal outcome. We reviewed possible new approaches to diagnosis and treatment.

Results: The study demonstrated the rapid evolution of the aggressive and malignant form of the disease. It also described newly accessible Phase I diagnostic tools being currently researched as well as new therapeutic approaches.

Limitation: The rarity of the disease, with only eleven cases throughout the literature.

Conclusion: The gastrointestinal form of Degos disease can be lethal. Its vascular etiology has finally been confirmed; however, new and more accurate early diagnostic modalities need to be developed. There are new therapeutic possibilities, but the studies of them are still in the early stages and have not yet shown the full effectiveness of these new therapies.

Keywords: multisystem vasculopathy, diagnosis, lethal, gastrointestinal, Degos disease

Introduction

Degos disease most frequently presents in white male teenagers, with the initial symptom being skin lesions on the trunk (Figure 1). Sometimes, within days, the patient presents with generalized acute abdominal pain and fever. Patients in this situation are frequently treated with surgery (exploratory laparotomy). Patients experience frequent vomiting and abdominal distension that does not improve after surgery. Weight loss at an average of 6–10 kg per month is common. A repeat laparotomy is frequently performed because of intestinal occlusion by adherences or acute peritonitis. Persistent high fevers, abdominal pain, and gasporesis are very common in these cases.
At the time of admission, fever (100.4°F–104.0°F), normal blood pressure, and sinus tachycardia are frequent. Peritonitis and septicemia should be suspected. In progressive conditions of the disease, multiple lesions with a papular aspect are observed in the thorax wall, abdominal wall, and limbs.

Blood tests reveal an elevated white blood cell count (normally of more than 30,000 cells/mm³) and low levels of hemoglobin and platelets because of infection. Blood gases show respiratory alkalosis with concomitant metabolic acidosis in most cases.

Chest and abdominal computerized tomography (CT) can reveal pleural effusion in the lower lungs fields and free liquid in the abdominal cavity with an edematous bowel aspect. Paralytic ileum can be present.

Laparotomy is always indicated and can show lesions consisting of white and yellowish or pearly-rose patches (Figure 2A) with a hyperemic rim throughout the bowel, mainly in the gastrointestinal tract. One or more points of perforation can commonly be observed in the jejunum, ileum, and colon, resulting in intestinal resections (Figure 2B).

Microscopic examination can reveal poor lymphomononuclear infiltrate and no granulomatous process, vasculitis, or neoplastic involvement.

A second-look operation is commonly performed 48 hours after the first laparotomy because of the frequent occurrence of new abdominal pain and an improvement in the inflammatory conditions. A skin biopsy is important for the diagnosis of Degos disease and will exhibit epidermal atrophy and dermal sclerosis with or without vasculitis. However, the histological findings of the intestinal and skin lesion biopsies may be inconclusive.

The diagnosis of inflammatory bowel diseases, such as Crohn’s disease, frequently requires discussion of the case by multiple doctors. Corticoids, such as methylprednisolone, are the first drug that doctors use in these patients.

Many repeat laparotomies are performed because of abscesses observed on control CTs.

A dermatological evaluation of skin lesions from multiple body locations (Figures 3 and 4) may encourage new skin microscopic examinations and may reveal an expressive papillary dermal sclerosis and thrombosis in the papillary dermis and in the vessels below the lesions.

With these histopathological findings and clinical/surgical aspects patients can be diagnosed with Degos disease. Antiplatelet therapy (acetylsalicylic acid) may be initiated and can be given in combination with steroid therapy. Pentoxifylline and heparin should also be given. Interleukin, interferon and infliximab can also be added to the treatment.

As studies have shown that vasculitis is the initial skin condition of Degos disease. The objective of those research models was to select drugs that block the deleterious autoimmune response in the blood vessels (Figure 5).
New perforations and bowel necrosis can occur, making new resections, enterectomies, and stomas necessary (Figures 6 and 7).

Microscopic examination will reveal diffuse transmural ulcerations and punctate perforations associated with an occlusive fibrosis of the medium- and small-diameter arteries and veins. Vessels are often occluded by thrombosis.

Uncontrollable septicemia and persistent gastrointestinal bleeding associated with renal failure and cerebral vascular lesions are the most common causes of death in patients with Degos disease.

The median period of survival for these patients is no longer than 6 months.

**Results**

Between June 1, 1998, and December 23, 2008, three cases of Degos disease were recorded, one case of the cutaneous or benign form and two cases of the cutaneous intestinal or malignant form. In 2013, a retrospective study of Degos disease cases was conducted, which revealed a permanent lethality of the malignant form of the disease. It also showed a favorable evolution, with greater than a 20% survival rate of the benign form of the disease.²

The retrospective study demonstrated the rapid evolution of the aggressive and malignant form of the disease, as well as newly accessible diagnostic tools, diagnostic tools currently being researched in Phase I trials, and new therapeutic approaches.

**Limitation**

The main limitation to this study is based on the rarity of the disease, with only eleven cases described throughout the world medical literature.

**Discussion and systematic review**

Degos disease, also known as Köhlmeier–Degos disease and malignant atrophic papulosis (MAP), is a rare vasculopathy of unknown cause that mainly involves the skin, gastrointestinal tract, central nervous system, and, occasionally, other organs. It was first named by Degos et al in 1942, but a case was reported in 1941 by Köhlmeier,⁴ who interpreted it as thromboangiitis obliterans of the mesenteric vessels. Although this disease usually affects young people, with an age of onset between 20 and 40 years, it may present at any age, even as early as 8 months.⁵,⁶ The disease has a male predominance (3:1), and sporadic cases of familial involvement have been recorded.⁵–⁹

The involvement of the gastrointestinal tract and other organs has been noted in approximately 60% of reported cases.¹⁰

Degos disease is a truly systemic illness, and at autopsy, lesions may be found in multiple sites, including the heart, lungs, kidneys, bladder, eyes, liver, and genital and oral mucosa.¹¹–¹⁶ However, in a subgroup of patients, the disorder may pursue a benign course involving only the skin for many years.¹⁵ Skin lesions are usually the first sign of the disease.⁵ The disease has been reported to be purely cutaneous in 37% of patients.¹⁷ Clinically, one sees pink- to rose-colored papules that may appear telangiectatic and are in different stages of development. Early lesions are slightly darker, may have an elevated border, and exhibit a central porcelain-white depression. The lesions are asymptomatic (except in a few cases in which they have appeared pruritic), benign and predominantly appear on the trunk and extremities.⁵,¹⁸

The lesions affect the bowel in approximately 47% of cases. Intestinal symptoms are variable, although
any portion of the intestinal system (from the oral cavity to the anus) may be involved, and the small bowel is predominantly affected. Lesions of the gastrointestinal tract are most often associated with skin involvement. In rare cases, a bowel injury may precede cutaneous signs, hampering the diagnosis. Some patients are asymptomatic, but others may complain of indigestion, diarrhea, constipation, and symptoms of gastroparesis or abdominal distension.

Many patients are asymptomatic, and abdominal involvement may be uncovered by systematic investigations, including endoscopy (visualizing white, yellowish, or pearly-rose patches with a hyperemic rim resembling the skin lesions but larger in size), laparoscopy (showing a comparable aspect on parietal and visceral peritoneum), and angiography (showing multiple stenoses of branches of the inferior mesenteric artery and hypovascular zones on delayed pictures). Multiple red inflammatory and white atrophic maculae can be observed on the serosa of the small and large intestines, and a section of the removed specimen can show acute inflammation and ulceration of the mucosa with surrounding intense congestion. Of greater importance, a small infarction of the
intestinal may develop and cause perforation and resultant peritonitis. To avoid fistulous problems, biopsies of the bowel wall should never be taken. Instead, biopsies of the parietal peritoneum lesions should be taken.

Neurological involvement occurs in 19% of cases. The manifestation of neurological involvement is highly variable, but hemiparesis with or without aphasia, monoplegia, tetraplegia, and cranial nerve involvement have been found. Cases of ascending polyradiculitis or myelomalacia have been reported. Headache, mental dysfunction, and weakness of the limbs are also common. Sometimes, injuries of the whole eye may be observed.

The diagnosis of Degos disease is based on the following clinicopathologic finding: many papules, 2–5 mm in diameter, with the classic porcelain-white centers characteristic of Degos disease. Some cases have exhibited different clinical morphologies that corresponded to the evolutionary stages of the papules originally described by Degos. Histologically, those patients exhibited a preeminent interface reaction with squamatization of the dermo–epidermal junction, melanin incontinence, epidermal atrophy, and a developing zone of papillary dermal sclerosis that resembled the early stages of lichen sclerosus et atrophicus, in miniature. This interface reaction was invariably confined to the central portion of the punch biopsy specimen, corresponding to the central porcelain-white area observed clinically. Additional features of fully developed papules included a prominent lymphocytic vasculitis affecting the venules, a mild periadnexal infiltrate of neutrophils and/or eosinophils, and interstitial mucin deposition. According to Harvell et al, these inflammatory cells are sometimes absent. In late-stage papules, the porcelain-white areas are more developed and the lesions are flattened. Histologically, the degree of inflammation was generally sparse, and the overall picture mirrored the classic histological description of Degos disease with a central roughly wedge-shaped zone of sclerosis surrounded by atrophic epidermis and hyperkeratotic compact stratum corneum. Skin lesions typically occur on the trunk and limbs. The palm, soles, face, and scalp are characteristically free of lesions.

The accurate etiology of Degos disease is still unclear, although viral, genetic, and immune mechanisms and an anomalous fibrinolysis have all been implicated. The pathogenesis of Degos disease is still under discussion, but primary endothelial swelling with vascular thrombosis leading to tissue infarction has been proposed. The cause of the initial vascular injury is unknown, but a mononuclear vasculitis seems to be the favored mechanism. It is possible for Degos disease to be associated with other systemic diseases. Its differential diagnosis is poor, as the cutaneous lesions are pathognomonic. Yet systemic lupus erythematosus and inflammatory bowel disease may occasionally be discussed.

Approximately 15% of Degos disease patients exhibit long-term survival, with the disease often limited to the skin and with no history of catastrophic bowel or central nervous system involvement. When there is multiple organ involvement, the disease results in death within 2–3 years. The course of the disease may be as long as 20 years, but once intestinal lesions have appeared, death usually occurs within a few months. The most common causes of death are peritonitis (61%), central nervous system complications (18%), and pleuritis and/or pericarditis (16%).
There is no proven effective treatment for patients with systemic involvement. Treatment with plasma change, corticosteroids, immunosuppressants, or antiinflammatory drugs has been ineffective in reversing the course of this disease when systemic involvement is present. Some authors have reported a good response to antiplalet therapy. A publication from Japan in January 2013 showed the expression of stromal cell-derived factor (SDF)-1/ CXCL12 in Degos disease. Secretory activity by stromal and endothelial cells of the bone marrow activates SDF-1/CXCL12 of megakaryocyte precursors and is responsible for the costimulation of platelet activation. Patients with Degos disease demonstrated a high level of secretion of SDF-1/CXCL12 inflammatory cells. This cell type was located in the perivascular, intravascular, and perineural tissue. These results support the theory that Degos disease is an endothelial disease.

**Conclusion**

The gastrointestinal form of Degos disease remains a serious and lethal disease. The vascular etiology of this disease has finally been confirmed; however, new and more accurate early diagnostic modalities need to be developed. There are new therapeutic possibilities; however, studies of new treatments are still in the early stages and the full effectiveness of those treatments has not been shown.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**

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