Case Report: Pansclerotic Morphea-Clinical Features, Differential Diagnoses and Modern Treatment Concepts

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Pansclerotic morphea (PSM) is a rare skin disease characterized by progressive stiffening of the skin with or without the typical superficial skin changes usually seen in morphea (localized scleroderma). Standard therapy, consisting of a combination of systemic glucocorticoids and methotrexate or mycophenolate mofetil, does rarely stop disease progression, which may lead to severe cutaneous sclerosis and secondary contractures. Little is known about the efficacy of newer biologicals such as abatacept, a fusion protein antibody against CTLA-4, or tocilizumab, a fully humanized IL-6R antibody, in the treatment of this pathology. We present the case of an 8 years old girl with an unusual, progressive stiffening of the skin, which was eventually diagnosed as pansclerotic morphea. A treatment with systemic glucocorticoids and methotrexate combined with tocilizumab led to a good clinical response within 2 months after initiation. In this paper, we discuss differential diagnoses to be considered and this new promising treatment option based on a case review of the literature.

Keywords: pansclerotic morphea, stiff skin, scleroderma, tocilizumab, IL-6, case report

INTRODUCTION

Diseases associated with stiff skin are extremely rare in children (1). The differential diagnoses include a generalized presentation of localized scleroderma (especially the deep variant of pansclerotic morphea (PSM)], systemic sclerosis, scleroderma-like disorders such as sclerema and eosinophilic fasciitis, or stiff skin disease. Differentiating these disorders is not always straightforward.

In this case report we describe an 8 years old child, presenting with a slowly progressing skin tightness after an initial infiltrated plaque at the anterior neck. Laboratory values, lung function tests, capillaroscopy, cardiac and ophthalmologic investigations were largely unremarkable. Skin MRI showed oedema of muscle fasciae. The final diagnosis was pansclerotic morphea, based on course and histopathology. We highlight the diagnostic dilemma and show the excellent response
to a treatment combining systemic glucocorticoids, methotrexate and tocilizumab.

CASE REPORT

An 8-year-old girl in good general health, presented with an irregular infiltrated plaque on the anterior neck of 3 months duration (Figure 1A). The family reported a streptococcal throat infection treated with antibiotics 1 week prior to onset. Antistreptolysin antibodies were high at that time. A skin biopsy, showed mucin accumulation between thickened collagen fibers suggestive of scleredema Buschke.

However, over the following months, the child developed a slowly progressive stiffness and reduced mobility of the back, shoulders and hips, subsequently extending to the wrists, fingers and knee with restriction of the flexibility. Raynaud phenomenon was absent.

After seven months, she presented at our consultation with a generalized wooden hard, infiltrated skin, most prevalent at the posterior thighs and lower back, only sparing her face, feet, hands and fingers [modified Rodnan Skin Score (mRSS) = 29/51]. The original neck lesion had disappeared (Figures 1B,C). Skin was adherent to the underlying tissues on palpation, with some nodular infiltrations and impossible to fold. Her general health remained well. She was able to continue school activities, tennis and artistic gymnastics although she felt more and more restrained. There were no episodes of fever or other systemic complaints. Rheumatologic investigation showed restricted movements of the wrists and knees, without clear signs of arthritis, therefore thought to be secondary to the tight skin.

At that time a deep skin biopsy of the thigh showed an enlarged dermis with a sparse perivascular lymphohistiocytic infiltrate, without eosinophils, and coarse hypertrophic collagen fibers. They invaded part of the hypodermis, underneath the sweat glands, surrounding fat globuli, causing compression on skin adnexes but did not invade deeper layers. A Masson stain confirmed the presence of coarse collagen but only minimal mucin was present (Figures 2A–C).

Blood-work was unremarkable with only at the time of the second biopsy minimally elevated eosinophils, slight hypergammaglobulinemia, but negative auto-antibodies (rheumatoid factor, ANA esp. anti-dsDNA, Scl70, RNP, SS-A, SS-B, SmD-1, Jo-1, anti-β2GP1, anti-cardiolipin, lupus anticoagulant) and normal kidney function, liver and muscle enzymes. On whole body MRI widespread oedema of fascia and subcutis was present (Figure 2D).

Lung function tests, capillaroscopy, cardiac and ophthalmologic investigations were within normal range. This together with absence of acral sclerosis and Raynaud
phenomenon makes a diagnosis of systemic sclerosis highly improbable.

Due to this severe clinical status, suspecting PSM, 7 months after the first symptoms, an aggressive combined treatment with glucocorticosteroid pulses (methylprednisolone 30 mg/kg/day for 3 days) followed by prednisone 2 mg/kg/day orally on a tapering regimen over 6 months, methotrexate 16.5 mg/m²/week and tocilizumab 10 mg/kg intravenously followed by weekly subcutaneous injections of 4.5 mg/kg/week were started. Our patient was encouraged to continue her sports activities, instead of adding physiotherapy, to stimulate her mobility skills.

Already after 1 month, a slight improvement was noted, which became more obvious after 4 months (Figure 1D). The skin felt less tight and hardened (mRSS = 18/51). With the exception of hyperpigmentation of a skin region where a band aid was placed, there was no alteration in skin coloration. Gymnastic skills improved drastically and this was confirmed by clinical examination showing less restriction in wrists and knees mobility as well as by disappearance of oedema on MRI (Figure 2E). Further improvement occurred the following months when she remained under methotrexate and tocilizumab treatment alone (mRSS = 6/51). The hyperpigmented patch disappeared slowly. Skin returned to normal, 1 year after start of treatment (mRSS = 0/51). Further lowering of therapy will be considered after 1 year, if no relapse occurs.

**DISCUSSION**

Stiffening of the skin associated with systemic symptoms, Raynaud phenomenon, characteristic lesions on capillaroscopy, and the presence of auto-antibodies esp. ANA, are diagnostic for systemic sclerosis, which is an extremely rare disease in childhood.

When systemic complaints and Raynaud phenomenon are absent, pansclerotic morphea (PSM), scleroderma-like disorders especially scleredema and eosinophilic fasciitis or stiff skin disease have to be considered (Table 1) (1). At onset, as our case illustrates, patients may be difficult to categorize, as there is a lack of clear diagnostic criteria, of disease markers and pathophysiology is not well-known, as isn’t the relationship between those entities. Illustrating this, there are many case reports showing clinically and/or histologically overlapping features of different entities (2–6).

The clinic and histopathology of the first lesion in the neck (although not in the typical posterior location), preceded by a streptococcal infection, were suggestive for scleredema (1) in this particular case. In children this disorder is usually caused by an infection, runs a very acute course and can lead to skin stiffness, which tends to involve preferentially the back neck and shoulder girdle. On histology, the disease is characterized by mucin accumulation between coarse collagen fibers. Spontaneous resolution is the rule within months. However, in our case, the development of a severe involvement of the pelvic girdle,
protracted course and the findings of the second biopsy proved this first diagnosis wrong.

Stiff skin disease usually is most marked at the pelvic girdle as in our patient. However it is a monoclonal disease, caused by an heterozygous mutation in the fibrillin-1 gene (FBN1), with onset always before the age of 6 years in the diffuse form. Moreover, this disease tends to remain stable or is only minimally progressive over the years (7, 8).

Eosinophilic fasciitis has a subacute onset and tends to affect extremities, trunk and neck but spares hands and feet (1, 2). Histopathology shows inflammation and fibrosis mainly at the fasciae and in the lower subcutis (6). However, in our case, although fascia was affected on MRI, on histology deep subcutis and fascia were spared. Moreover, skin and blood eosinophilia were absent in most analyses. Response to corticosteroids, is usually excellent, however, relapse after stopping is the rule. Therefore, the most likely diagnosis in this patient was PSM. This disease, a rare (<1%) subtype of localized scleroderma (9), is characterized by near total body surface involvement, with circumferential lesions, sparing fingers and toes that usually extend in subcutaneous tissue, and may affect fascia, muscle and bone. It has a more insidious onset than eosinophilic fasciitis and post-infectious scleredema, symptoms progressively appearing over months (10). Atrophic hyperpigmented lesions and joint contractures (therefore called disabling PSM) are usually present but may also develop later in the course of disease. On histopathology fibrosis and inflammatory infiltrates always involve lower dermis and upper subcutis, sometimes deeper tissues may also be affected (6). In about one third of

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**TABLE 1** | Typical clinical characteristics of PSM, eosinophilic fasciitis, scleredema and stiff skin syndrome.

|                         | Pansclerotic morphea (PSM) | Eosinophilic fasciitis | Scleredema | Stiff skin syndrome |
|-------------------------|-----------------------------|------------------------|------------|---------------------|
| Age at onset            | Infants to teenagers rare in adults | Infants to adults | Child (post-infectious), adults | Toddler to infant, M: 1,6 a; max: 7a (sporadic or familial) |
| Location at onset       | All body areas possible | Extremities | Back of the neck | Pelvic and shoulder girdles |
| Final distribution      | Diffuse, circumferential, centrifugal extension; sparing hands and feet, progressive | Diffuse; sparing face and trunk | Face, head and neck, trunk and shoulders; sparing extremities spontaneous resolution within months (exc. 2 years) | Diffuse |
| Skin (characteristic changes) | Hard to touch, hypo- or hyper-pigmentation, atrophic sclerotic plaques, ulcers | Painful induration of the skin | Induration of the skin, Pigmentation, Peau d'orange | Rock-hard skin bound to underlying Tissues, Hypertrichosis, Hyperpigmentation |
| Articulations           | Restricted joint mobility, contractures | Restricted joint mobility, contractures | Restricted joint mobility | Restricted joint mobility, contractures |
| Systemics symptoms      | Asthenia | Secondary: compartment syndrome, dysphagia, dyspnea | Exceptional | Exceptional (cardiac) |
| Capillaroscopy          | Normal | Normal | Normal | Not reported |
| Biology                 | ANA in 30% of cases; eosinophilia and hypergammaglobulinemia: inconstant | ANA negative; eosinophilia and hypergammaglobulinemia (may be absent) | Epidemis: normal | Epidemis: normal |
| Histology               | Epidermis: thinned | Epidermis: normal | Epidemis: normal | Epidemis: normal |
|                         | Dermis/hypodermis (may extend into fascia); inflammation and thickened collagen bundles | Dermis: normal | Dermis: thickened x3-4, full of collagen fibers Containing clear spaces full of mucin ++ | Dermis: increased mucin and fibroblast, horizontal collagen fibers |
|                         | Perivascular inflammation; lymphocytes and plasma cells; Eosinophils may be present | Hypodermis and fascia: edema, inflammation and thickened collagen bundles | Hypodermis and fascia: Edema, Inflammation: lymphocytes, eosinophils, histiocytes and plasma cells | Hypodermis: adipocytes, entrapment |
|                         | IPM | Inconstant oedema of hypodermis and fascia | Oedema of hypodermis and fascia | Fascia: thickened and hyalinized |

*Prayer sign: when the patient is unable to oppose the palmar surfaces of both hands with extended wrists. Groove sign: depression along the course of the superficial veins. M, median.
patients, auto-antibodies are present and sometimes eosinophilia and hypergammaglobulinemia can be seen.

Much progress has been made in clarifying the role of profibrotic cytokines such as TGFβ, IL-4 and IL-6 [for a review see (11)] in scleroderma pathogenesis. More specifically, IL-6 is a pro-inflammatory cytokine produced by B-cells, T-cells, monocytes and fibroblasts. It is required for differentiation of Th17-cells. IL-6 also regulates fibroblasts activity, stimulates collagen production and inhibits the synthesis of collagenases. In systemic sclerosis it has been demonstrated that IL-6 blockade reverses TGFβ activation (12). Via these actions, IL-6 has a role in the fibrotic pathways (11, 13). High serum levels of IL-6 have been described in patients with localized scleroderma (13–17).

According to literature, treatment of PSM is disappointing (18). Disease progression, despite classical treatments (systemic glucocorticoids, methotrexate or mofetil mycophenolate) is

| TABLE 2 | Pediatric case reports of PSM treated with tocilizumab. |
|---|---|---|---|---|---|---|---|
| References | Age at onset (years) | Previous treatments | Disease duration at onset of TCZ | Dose of tocilizumab | Concomitant treatment | Result |
| Martini et al. (19) | 4 | Prednisolone (pulse and oral) MTX 15 mg/m²/w MMF 700 mg/m²/d Imatinib 200 mg/d | +/+144 m | IV 8 mg/kg/8 w | Still on Imatinib? None? | Rapid reduction of inflammation follow-up (FU) 18 m |
| Patient 2 | 4 | Prednisolone (pulse and oral) MTX 10–15 mg/m²/w MMF 700 mg/m²/d | +/+ 68 m | IV 8 mg/kg/8 w | Prednisone + MTX + MMF | Inactivation of lesions allowing reduction and stop of TCZ, FU of 24 m after stop still inactive |
| Foeldvari et al. (20) | 7 | MTX 15 mg/m²/week MMF 1,200 mg/day | 81 m | IV 8 mg/kg every 2 weeks | Prednisone + Tacrolimus | No new lesions; decrease in erythema and skin thickness after 31 m |
| Patient 2 | 2 | MTX 15 mg/m²/week MMF 40 mg/kg/day Etanercept 0.8 mg/kg/week | 95m | IV 8 mg/kg every 4 weeks | MMF | No new lesions, decrease of erythema, skin thickness stable after 23 m |
| Patient 3 | 7 | MTX 24 mg/m²/w | 9 m | SC 6.5 mg/kg every 3 weeks | MTX | No new lesions, decrease of skin thickness after 9 m |
| Zang et al. (21) | 6 | Prednisolone pulses MTX MMF | +/+42 m | IV 300 mg/4 w | MTX | Improvement within months |
| Soh et al. (18) | 4,5 | Naproxen, prednisolone (pulses and oral) MTX (15 mg/m²/w) MMF 300 mg BID IVIG Rituximab | +/+ 96 m | dose not specified | Prednisone + MTX + Ig IV Hydroxychloroquine and naproxen | After 3 months no effect |
| This case report | 9 | none | 7 m | IV 10 mg/kg followed by SC 4.5 mg/kg/w | Prednisolone pulse and oral MTX 16,5 mg/m²/w | Improvement of skin infiltration started after 1 m and was marked after 4 m of treatment, skin returned to normal after 12 m: mRSS from 29 to 0/51 |

*mLoSSI modified Localized Scleroderma Skin Severity Index.
**PGA-A Physician Global Assessment of Disease Activity score. mRSS, modified Rodnan Skin Score.
often reported. New treatments targeting pathways stimulating collagen production such as IL-6 (tocilizumab), or broader T-cell activation (abatacept) have been evaluated. Until now, four reports, on tocilizumab treatment of seven longlasting pediatric PSM cases, not responding to standard treatments, have been published. There was a partial response (especially on activity scores), in 6 out of 7 patients (18–21), within months. Details about these and our patients are given in Table 2. Positive experiences with abatacept are especially reported in patients with systemic sclerosis and different types of morphea. However, there are no patients with PSM treated at this moment as far as we know (22, 23).

In our case the combination of systemic glucocorticoids, methotrexate and tocilizumab led to a rapid, complete and sustained healing. The specific role of tocilizumab is difficult to evaluate, because all three medications were started simultaneously. However, considering the disappointing results of standard treatments, it seems likely that it played a major role in reversing the fibrosis in our patient. Furthermore, the response in this case suggests that early treatment with tocilizumab is even more effective, leading to complete healing.

In conclusion, with our case, we discuss the differential diagnoses of stiff skin in children and describe some potential difficulties in making an early diagnosis of PSM. This case also suggests that drugs targeting IL-6 might be a good option to consider early in the treatment of PSM, when it might be most beneficial, in this disease where standard treatments usually are disappointing.

**DATA AVAILABILITY STATEMENT**

The original contributions generated for this study are included in the article/supplementary material, further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

Written informed consent was obtained from the minor(s)’ legal guardian/next of kin for the publication of any potentially identifiable images or data included in this article.

**AUTHOR CONTRIBUTIONS**

AS-B was the first physician to see the patient. SV, M-AM, RN, CS, MH, and FD followed the patient clinically. SB and DH reviewed the histopathology. SV and M-AM wrote the first draft of the article. All authors contributed to and agreed the final draft.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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