Refractory cutaneous polyarteritis nodosa with threat of loss of feet: successful treatment with rituximab

Julie E. Stephan1 and David A. Minna2*

1MPAS, PA-C UT Southwestern Medical Center Dallas, Texas, USA. 2MD, FACR UT Southwestern Medical Center Dallas, Texas, USA. *Correspondence: david.minna@utsouthwestern.edu

Abstract
Cutaneous Polyarteritis Nodosa (CPAN) is a rare, single organ vasculitis of unknown etiology which presents with livedo reticularis and tender nodules involving the lower extremity in 97% of cases. Diagnosis is established by skin biopsy demonstrating segmental leukocytoclastic vasculitis and fibrinoid necrosis with workup excluding any systemic involvement (Figure 1). The presence of IgM antiphosphatidylserine-prothrombin complexes has led to the hypothesis that CPAN is a localized arthrus reaction. CPAN is characterized as a benign, chronic, relapsing vasculitis to be treated conservatively. As a single organ vasculitis, CPAN can affect arteries of any size. Due to CPAN's predilection for the lower extremities, involvement of the limb at or below the ankle's major arteries may result in loss of digits and potentially feet. The case presented involves CPAN in which the main arteries to the feet are occluded. The patient had been refractory to therapy with findings of early digital necrosis. Following adequate skin biopsy to establish a diagnosis, we initiated targeted therapy with rituximab, azathioprine, and high dose steroids with reversal of the ischemic process. It is the authors' opinion that CPAN should be treated more aggressively from initial diagnosis to prevent morbidity and potentially abort a chronic relapsing course.

Keywords: Cutaneous polyarteritis nodosa, rituximab therapy, refractory cutaneous polyarteritis nodosa, CPAN, cutaneous PAN

Introduction
The 2012 Chapel Hill Consensus Conference Nomenclature of Vasculitides defined Systemic Polyarteritis Nodosa (PAN) as a non-ANCA associated necrotizing arteritis of medium or small arteries. This delimitation of PAN makes it a rare disease (PAN: 31/106, GPA: 160/106, MPA: 94/106) [2] the single organ presentation of CPAN is rarer still. The true incidence is unknown [3] it has been estimated to be <3% of all cutaneous vasculitides [4]. CPAN is characterized as a "benign, chronic, relapsing vasculitis" [5] requiring conservative management with NSAIDs, low dose steroids, and possibly metotrexate; as opposed to, the malignant course of systemic PAN which necessitates aggressive therapy with high dose steroids and immunosuppressive agents [1]. However, CPAN has been reported to cause distal extremity necrosis resulting in amputation [6,7]. When CPAN involves the distal lower extremity's intermediate size arteries, there is an increased risk of loss of digits and limb and aggressive therapy must be considered.

CPAN is a rare diagnosis which may be missed in favor of other more prevalent vasculitides. There are limited initial laboratory findings, while the clinical findings of tender nodules, livedo reticularis, and palpable purpura are associated with multiple diagnoses increases the probability of a missed CPAN diagnosis. This may be compounded by insufficient biopsy material [3], especially with foot and ankle involvement.

Presented here is a case of CPAN in which bilateral vascular occlusion at the level of the ankles threaten loss of digits and potentially feet which reversed with combination therapy of high dose steroids, rituximab, and azathioprine.

Case presentation
The patient is a 38-year-old white female who presented to the outpatient rheumatology service with a ten month history of progressive bilateral foot pain. The right mid foot and forefoot were first involved, followed by the left ankle and hindfoot. Initially, no findings of swelling, temperature variation, skin...
doi: 10.7243/2055-7000-4-1

discoloration or rash had been present. She had no history of cutaneous, pharyngeal, respiratory infections, new medications, or illicit drug use. At four weeks from onset of symptoms, she developed signs of ischemia, acral cyanosis, palpable purpuric lesions of the right forefoot and left hind foot.

The patient returned for re-evaluation with a community rheumatologist two months after onset of symptoms with a workup including negative ANA, DNA, ENA, ANCA panel, hepatitis B and C serologies, and cryoglobulins. The patient's hypercoagulability profile was negative except for a positive phosphatidylserine IgM. From this data, a presumptive diagnosis of vasculitis had been made which prompted therapy of prednisone 20 mg and methotrexate 12.5 mg weekly. The patient had been maintained on this regimen for six months with progression of other symptoms. Subsequently, a skin biopsy from the right lateral foot demonstrated focal changes of leukocytoclastic vasculitis. The shallowness of the biopsy without biopsy of an artery limited diagnostic interpretation.

Admission to the hospital was later necessary due to further worsening of her symptoms. Upon admission, her vital signs were normal. Her Skin exam revealed the following: “dark and purple” feet with non-blanching, tender, purpuric lesions; right toes were cyanotic and tender to palpation; distal pulses were palpable but not rated. The remainder of her physical exam was within normal limits. Her labs were significant for a leukocytosis of 14.6, an anemia with hemoglobin of 10.7, elevated platelet count of 599, sedimentation rate of 30 (0-20), phosphatidylserine IgM of 50 (0-24), and an ANA of 1:160. The remainder of her serologies was unrevealing including DNA, ENA, complements, ANCA panel, hepatitis Band C serologies, cryoglobulins, and her hypercoaguable profile. Of note, the patient also demonstrated a normal trans-esophageal echocardiogram and lower extremity doppler study for deep vein thrombosis. The patient underwent CT angiograms of her abdomen, pelvis, and bilateral lower extremities which revealed no abnormalities to the level of the ankles at which point the right and left anterior tibial and dorsalis pedis arteries abruptly taper or cut off. In addition, the left foot demonstrated the left posterior tibial artery to be the sole vascular supply to the left foot, the left anterior tibial artery occluded at the ankle, and the peroneal artery occluded in the distal calf. Similarly, the right foot shows no in-line flow to the foot, the posterior tibial artery occludes at the level of the calcaneous, the anterior tibial artery occluded at the ankle, and the peroneal artery supplies collaterals to the foot (Figures 2-5) Finally, the digital arteries are irregular, bilaterally, consistent with the diagnosis of vasculitis. In view of her diagnosis of vasculitis, she was discharged on 60 mg prednisone daily and continued on methotrexate 12.5 mg weekly.

One week after her discharge, the patient presented to our clinic. At this time, patient presented with livedo changes of bilateral feet, tender palpable purpuric lesions on the right medial forefoot and left medial hind foot, acral cyanosis of the right 1st and 4th toes with early necrosis of the digital pulps (Figures 6-8). Her dorsalis pedis pulses were non palpable bilaterally; posterior tibial pulses were palpable 2/4 on left foot only. Her neurologic exam was within normal limits. Her labs again showed leukocytosis, thrombocytosis, anemia, elevated c reactive protein, and normal urinalysis and creatinine. A skin biopsy was performed from the digital pulp of the right 1st toe demonstrating an arteriole, inflamed by neutrophils, histiocytes, and lymphocytes with fibrin deposition in the muscular wall, and occlusion of lumen (Figure 1). This pattern is consistent with PAN.

Taken together, the objective evidence led to the establishment of a CPAN diagnosis. Accordingly, therapy was initiated with 60 mg prednisone daily, azathioprine 100 mg daily, and rituximab 1000 mg day 0 and day 15. After eight weeks from initiation of therapy, her diffuse foot pain had resolved and skin color improved, but patient had developed a painful 1 cm ulcer on tip of her right 1st toe (Figures 9 and 10).
Figure 3. Lt Foot: Lt Post Tibial Artery is sole supply of Lt foot. Lt ant-tibial artery occludes at ankle. Peroneal artery occludes in distal calf. Irregularity of all digital arteries consistent with dx vasculitis.

Figure 4. Rt Foot: There is no in-line flow to the Rt foot. Post tibial artery occludes at level of calcaneus. The ant tibial artery occludes at the ankle. Peroneal artery supplies collaterals to foot. Digital arteries are irregular consistent with dx of vasculitis.

Figure 5. Rt Foot: There is no in-line flow to the Rt foot. Post tibial artery occludes at level of calcaneus. The ant tibial artery occludes at the ankle. Peroneal artery supplies collaterals to foot. Digital arteries are irregular consistent with dx of vasculitis.

Discussion
Cutaneous Polyarteritis Nodosa, a rare single organ vasculitis, whose pathology is a segmental, necrotizing, leukocytoclastic vasculitis, may affect muscular arteries of any size. The initial manifestation is livedo reticularis which is also associated with antiphospholipid syndrome. Findings of CPAN do not fulfill criteria for antiphospholipid syndrome; however, antiphospholipid antibodies have been hypothesized to be involved in its pathogenesis. In one study, 81% of patients with CPAN
were reported to have IgM antiphosphatidylserine-prothrombin complexes with all controls negative. The hypothesis is prothrombin bound to apoptotic endothelial cells induces anti-PS/PT antibody production. These immunoglobulin complexes then activate the classical complement pathway causing CPAN. Direct immunofluorescence shows IgM deposition in vessel walls in 60% of specimens and C3 in 40% of specimens with no deposits at the dermal-epidermal junction. It is postulated that a localized arthrus reaction is occurring in the vessel wall [3,8]. As a result, the chronic, relapsing course of CPAN is indicative of persistent antibody production and varying concentration of antigen. Therefore, the reduction of the inciting clone of B cells would reduce immune complex formation to inhibit the inflammatory process and restrain the chronic, relapsing course. In other words, the elimination of the B cell clone would prevent recurrence of CPAN.

The non-systemic pattern of involvement in CPAN is portrayed as a benign process warranting conservative management with nonsteroidals and/or colchicine, unless the disease proves to be refractory to conservative management [3,5]. Refractory disease is characterized by one or more of following: persistence of cutaneous lesions, severe pain in involved region, ulcerations, necrosis, fever, myalgias, arthralgias, dysesthesias, or neuropathy [3]. The reported prevalence of the aforementioned in CPAN cases is as follows: skin ulcers develop in 50%, arthralgias 69%, myalgias 31%, neuropathy 22%, and fever 25-30% [3]. At the time of diagnosis, refractory CPAN should be the expected disease course rather than the exception. It would be prudent and in the patient’s longterm interest to institute an aggressive course of therapy unless contraindicated.

Symptom amelioration in 78% patients with CPAN has been reported with warfarin therapy [9]. Anticoagulation is premised on the presence of phosphatidylserine prothrombin complexes and antibodies with findings of thrombosis in cases of CPAN. Corticosteroid therapy in doses from 30mg to 1mg/kg of prednisone is recommended in refractory cases; steroids control acute exacerbations of CPAN. Remissions have been reported, but upon tapering the steroid dose exacerbations of CPAN are generally reported [3]. Steroid sparing therapies reported for CPAN include hydroxychloroquine, dapsone, azathioprine, methotrexate, IVlg, and cyclophosphamide [10-13]. Although immunosuppressives have been reported to be effective in CPAN, there are no controlled prospective trials to evaluate their efficacy. Use of immunosuppressives has been reserved for the most serious and unresponsive cases of CPAN [3]. Rituximab has been reported to be effective in a single case of CPAN with digital gangrene that was refractory to therapy with steroids and cyclophosphamide; no recurrence was reported at six months [14].

Our patient had occlusions of all lower extremity vessels save for the left posterior tibialis artery. She had been treated with steroids and methotrexate for six months with progression of her symptoms. Use of cyclophosphamide for this case was avoided since the use of cyclophosphamide in a 38 year old female may induce a premature menopause (>50% incidence) unless pre-medicated with leuprolide [15]. Considering the immediate threat of loss of digits, significant potential for loss of foot, and probable irreversibly compromised arterial supply to feet which future recurrences of CPAN would
exacerbate, we decided to infuse rituximab 1000 mg day 0 and day 15 with concomitant use of azathioprine 100mg and 60 mg prednisone daily. Rituximab has been shown to be effective in other types of vasculitides for induction and maintenance therapy [16-18].

Eight weeks from her initial rituximab infusion, the patient had resolution of her diffuse bilateral foot pain, improvement of skin color, and improved nail bed perfusion. The cyanosis of her right 1st and 4th toes had resolved, but the right 1st toe despite improved perfusion developed a 1 cm full thickness ulcer of the digital pulp at the base of digit. The patient was able to weight bear and walk without limitations. Her steroids have been tapered to 20 mg daily and azathioprine continued. It is planned to repeat the rituximab dose of 1 gram at day 0 and 15 at six months. Following her second cycle of rituximab, therapy will be based from clinical observation. Since the presence of IgM phosphatidylserine-prothrombin (PST/PT) complexes have been reported in 81% of patients with CPAN [8]; PST/PT complexes have been used as a diagnostic marker for CPAN, but have neither been studied in regards to disease activity, nor success of therapy. Assessing the vascular status in this patient is problematic. To identify the pathology a catheter angiogram was required, so repetition of this study is not considered prudent. Doppler studies could be helpful if re-cannulization of one or more arteries occurs. A three phase nuclear scan may allow quantification of vascular perfusion.

**Conclusion**

The development of severe acral cyanosis, attendant ischemia, and possible threat of necrosis generally triggers a differential diagnosis of 1) peripheral vascular disease, 2) embolic phenomena, 3) hypercoaguable state, thrombotic vasculopathy, 4) vasospastic disease/Reynaud's and 5) vasculitis. The first four can be evaluated with physical exam, noninvasive imaging, and laboratory testing. Localized vasculitis requires tissue biopsy which must be placed in context with the patient's history. When the involved region is distal, such as the foot or hand, obtaining a suitable biopsy can be difficult and non-diagnostic or misleading if too superficial. Since CPAN is hypothesized to be caused by local deposition of antigen antibody complexes of IgM antiphosphatidylserine and prothrombin in the arterial wall causing a necrotizing, segmental leukocytoclastic vasculitis, a targeted therapy with rituximab could be chosen with a high level of confidence of benefit. While selection of appropriate therapy is always important, it was critical in this case with the severity of vascular compromise and the threat of loss of digit or limb by ischemic necrosis. Consideration was also given to the long term adverse effects of cyclophosphamide, as well as, the increased complexity and frequency of therapy. The use of rituximab affords the chance of reducing or eliminating the inciting B cell line with the intention of altering the typical chronic, relapsing course of CPAN. In the patient presented, recurrence could cause loss of digit or limb. It is the authors' opinion that CPAN should be treated more aggressively from initial diagnosis to prevent morbidity and potentially abort a chronic, relapsing course.

**List of abbreviations**

ANA: Antinuclear antibodies
ENA: Extractable nuclear antigens; includes Anti-smith antibody, Ro and La antibodies, Jo1 antibodies, Sc170 antibodies, and ribonuclear protein antibodies
ANCA: Anti-neutrophilic cytoplasmic antibodies; includes c and p ANCA, myeloperoxidase, and protein 3

**Competing interests**

The authors declare that they have no competing interests.

**Authors' contributions**

<table>
<thead>
<tr>
<th>Authors' contributions</th>
<th>JES</th>
<th>DAM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Research concept and design</td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Collection and/or assembly of data</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Data analysis and interpretation</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Writing the article</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Critical revision of the article</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Final approval of article</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Statistical analysis</td>
<td>--</td>
<td>--</td>
</tr>
</tbody>
</table>

**Publication history**

Editor: Antonio G Tristano, Medical Center Carpetana, Spain.
Received: 09-Jan-2017 Final Revised: 20-Feb-2017
Accepted: 01-Mar-2017 Published: 11-Mar-2017

**References**

8. Kawakami T, Yamazaki M, Mizoguchi M and Soma Y. High titer of anti-


Citation:
http://dx.doi.org/10.7243/2055-7000-4-1