Case Report

Microscopic Polyangiitis Secondary to Silica Exposure ∗

Juliana Vega Miranda,a Luis Fernando Pinto Peñaranda,b Javier Darío Márquez Hernández,b Carlos Jaime Velásquez Franco b,c,∗

a Medicina Interna, Universidad Pontificia Bolivariana, Medellín, Colombia
b Grupo de Reumatología, Hospital Pablo Tobón Uribe, Medellín, Colombia
c Facultad de Medicina, Escuela de Ciencias de la Salud, Universidad Pontificia Bolivariana, Medellín, Colombia

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ABSTRACT

There is sufficient evidence of the capacity of silica to induce autoimmunity in patients with some type of genetic susceptibility. There are several autoimmune diseases related to this exposure (rheumatoid arthritis, Sjögren’s syndrome, sarcoidosis, and systemic sclerosis). Nodular silicosis (clinical expression of this exposure in lungs) generates apoptosis, inflammation, loss of tolerance and a respiratory burst. There is evidence that relates silica with induction of antineutrophil cytoplasmic antibodies, but, until it is better explained, the reports of systemic vasculitis secondary to silica exposure are inconclusive. We describe the case of a patient with a history of occupational exposure to silica who developed microscopic polyangiitis.

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Poliangitis microscópica secundaria a exposición a sílice

RESUMEN

Existe suficiente evidencia de la capacidad de la sílice de inducir autoinmunidad en pacientes con algún tipo de susceptibilidad genética. Existen varias enfermedades autoinmunes relacionadas con esta exposición (artritis reumatoide, síndrome de Sjögren, sarcoidosis, esclerosis sistémica). La silicosis nodular (expresión clínica pulmonar de esta exposición) genera fenómenos de apoptosis, inflamación, pérdida de la tolerancia y explosión respiratoria. También se ha descrito la inducción de anticuerpos antititoplasma del neutrófilo con este mineral, pero hay reportes no concluyentes de vasculitis sistémicas secundarias a la exposición a la sílice. Se describe el caso de un paciente con antecedente de exposición ocupacional a sílice que desarrolla una poliangitis microscópica.

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Introduction

Silica exposure has been linked to autoimmune diseases (rheumatoid arthritis, systemic sclerosis, Sjögren’s syndrome, and sarcoidosis). An association has been described between this mineral and the presence of ANCA but, to the best of our knowledge, there are no conclusive reports of induction of systemic vasculitis.

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∗ Corresponding author.
E-mail addresses: carjaivel@hotmail.com, carjaivel@gmail.com (C.J. Velásquez Franco).

Clinical Observation

The patient was a 34-year-old male who, for the past 6 years, worked as a marble installer. He had other relevant medical history. He came to the clinic after 7 days of progressive edema of the lower limbs, which progressed to anasarca, fatigue, weakness and oliguria. Review of systems: Negative. His blood pressure was 160/100 mmHg and he was pale; the rest of the examination provided no relevant data. The patient developed nephritic and nephrotic syndromes. The results of paraclinical studies on admission are seen in Table 1.

The hepatic function tests, antinuclear antibodies, antibodies against core extractable antigens, anti-DNA, complement levels, antistreptolysin, hepatitis B and C, ELISA for human immunodeficiency virus and syphilis serology were negative. ANCA were
Table 1
Initial Paraclinical Data.

<table>
<thead>
<tr>
<th>Test</th>
<th>Result</th>
<th>Normal value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Creatinine (mg%)</td>
<td>13</td>
<td>0.8–1.2</td>
</tr>
<tr>
<td>Urea nitrogen (mg%)</td>
<td>82</td>
<td>0.2–10</td>
</tr>
<tr>
<td>Urinalysis</td>
<td>30 erythrocytes/AP; 500 mg protein, erythrocyte and leukocyte cylinders (5–10/AP)</td>
<td></td>
</tr>
<tr>
<td>Protein in 24-h urine (mg)</td>
<td>5400</td>
<td>Less than 120</td>
</tr>
<tr>
<td>Hemoglobin (g/L)</td>
<td>8.9</td>
<td>12–16</td>
</tr>
</tbody>
</table>

Fig. 1. Renal biopsy evidence of glomerulonephritis with segmental necrotizing extracapillary proliferation (long arrow) with marked interstitial fibrosis and tubular atrophy (short arrow).

The patient was hemodynamically unstable, with high ventilatory requirements, and died.

Discussion

The initial organ affected with silica exposure was the lung (silicosis). Necrosis and apoptosis of alveolar macrophages, production of proinflammatory cytokines (tumor necrosis factor, interleukin-1), increased cytotoxic T cell survival, decreased regulatory cells and enhanced reactive oxygen species are phenomena generated. In general, the patient had few symptoms; dyspnea should prompt a complication or another associated entity.

This has also been linked with the production of ANCA; they may be directed against various antigens (proteinase 3, myeloperoxidase, lactoferrin and bactericidal permeability increasing protein). To the best of our knowledge, reported cases of vasculitis secondary to silica are controversial.

Bartůnková et al. analyzed 86 individuals exposed to silica for at least 5 years; presence of ANCA was more frequent in those exposed (17.1%) than in controls (3.6%). The odds ratio (OR) was 5.04 (95% CI, 1.2–21.2).5

The case being reported is similar to those described in the literature: Tervaert et al. explored the relationship between silica and the development of systemic vasculitis, finding an OR for renal failure after rapidly progressive glomerulonephritis (95% CI, 1.37–4.60) and 6.5 for pulmonary vasculitis (95% CI, 1.4–11.6).9

Hogan et al., in a case–control study, found an increased risk of small-vessel vasculitis associated with ANCA after high exposure to silica (5 years) (OR 1.9, 95%, 1–3.5, P=0.05).10 The pathophysiology is similar to that of silicosis, but the target cell is a neutrophil.

Conclusion

This case illustrates the probable association between severe microscopic polyangiitis and prolonged exposure to silica. Given the limited number of reported cases of this association, the prevalence of silicosis among those exposed and the frequency of development of autoimmune diseases remain unknown.

Ethical Responsibilities

Protection of people and animals. The authors declare that no experiments have been performed on humans or animals.

Data confidentiality. The authors declare that they have followed the protocols of their workplace regarding the publication of data from patients and that all patients included in the study have received sufficient information and have given their written informed consent to participate in the study.

Right to privacy and informed consent. The authors state that no patient data appears in this article.

Conflict of Interest

The authors declare no conflicts of interest.

References