A recent contribution to this journal, Camellino and colleagues [1] reported that asymptomatic inflammation of the interspinous bursas is a common occurrence in patients with polymyalgia rheumatica. Thus, they confirm the findings of three other studies which found florid inflammation (which was felt to be symptomatic) at these sites between the spinous processes of adjacent vertebrae. Why would this be so when bursitis at more accessible locations is not recognized as a part of this syndrome? One possible explanation might be found in the proximity of each interspinous bursa to its adjacent ligamentum flavum [2]. If the fundamental process is an auto-immune response to elastic tissue, then this most elastic of all anatomical structures would be an appropriate target and a 'sympathetic effusion' might be expected in adjacent tissues such as the interspinous bursas.

A similar situation could explain the characteristic synovitis of large joints in polymyalgia. There, scanning techniques demonstrate an unequivocal inflammatory cause for the joint pain and stiffness, but the effusions are small, synovial fluid findings are unimpressive, and the synovitis is not destructive. This, too, could represent an immune response to aging elastic tissue of the joint capsule with the synovitis being secondary.

Possible elastic antigenicity is not a new idea in giant cell arteritis [3]. Localization of inflammatory cells around the internal elastic laminae of temporal artery biopsies was prominent from the first recognition of this entity and remains so today [4]. Apparent elastic fragments within giant cells, immunoglobulin deposits adjacent to the internal elastic lamina, proliferative responses of lymphocytes to arterial tissue in vitro, and a striking response of mononuclear cells to elastin peptides all lend support to this idea, although none of these findings have been replicated convincingly [5].

Polymyalgia rheumatica can be excruciating, giant cell arteritis can be lethal, and both conditions respond dramatically to appropriate therapy. Yet their diagnosis can be difficult, and their common link has not been found. Confirmation of elastic antigenicity would provide much-needed understanding of the basic pathophysiology and could thereby facilitate both diagnosis and management. Does not someone want to take another look at the possibility that these findings and concepts may be held together with an elastic band?

Competing interests
The author declares that he has no competing interests.

Author's information
I am now an Emeritus Professor of Medicine at the University of Washington where I have been since beginning my fellowship in 1966. From my original mentor, Andy Healey, I derived a career-long interest in polymyalgia and giant cell arteritis. This included early contributions on familial aggregation of these conditions, angiographic studies of involved temporal arteries, and immunoglobulin deposition in arterial biopsies. In addition, I have had a long-held, but unpublished, suspicion that the synovitis in polymyalgia is secondary to an immune response in the underlying joint capsule. The recent reports of bursitis in interspinous bursas, which abut upon the ligamentae flava, provide an opportunity to share this hypothesis with young immunologists who might be in a position to confirm it. More importantly, it provides Andy’s many rheumatologist friends with the sad news of his recent death.

Acknowledgements
Most of what I know about polymyalgia and giant cell arteritis was learned from the late LA Healey.

Published online: 28 April 2015

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