The Fabulous Fibroblast

By Warren Hammer, MS, DC, DABCO

One of the most exciting things about manual methods such as Graston, active release, fascial manipulation, friction massage and others is that mechanical loading stimulates the proliferation of fibroblasts. Fibroblasts are the most common cells of connective tissue. These cells, among many other functions, synthesize the extracellular matrix and collagen, which represents the structural framework for our tissues. Lest we forget, the extracellular matrix (ECM) represents everything outside of the cells. Besides providing a structural support to our cells, within the ECM are gels of polysaccharides and fibrous proteins that fill the interstitial spaces and provide a compression buffer against all loads.

The composition of the extracellular matrix determines the physical properties of connective tissues – whether they become tendons, ligaments, etc. So, the collagen, glycosaminoglycans, glycoproteins, reticular and elastic fibers all found in the ECM are made by fibroblasts. Collagen is considered the most important ECM component since fibrillar collagen takes most of the tensile loading. Can you imagine that we help create these changes by frictioning tissues?1-2

Fibroblasts also play an essential role in wound healing. After the initial injury to connective tissue and blood vessels, growth factors cause fibroblasts to enter the wound, increase in number and start synthesizing...
new collagen, creating new granulation tissue and assisting in remodeling. The extracellular matrix of granulation tissue is created and modified by fibroblasts. At first, the fibroblasts produce type III collagen, a weaker form of the structural protein; and later produce the stronger, long-stranded type I collagen that appears in the scar tissue.

A scar is collagen deposited by fibroblasts during repair. Collagens I to III represent 80-90 percent of the collagen in the body,\(^5\) with the majority being type I. Fascia is a type I form of collagen. Graston Technique and fascial manipulation both work on the premise that small amounts of trauma to an area initiate an inflammatory process, triggering a healing cascade by enhancing the proliferative invasion of blood, nutrients, and fibroblasts to the region, which results in collagen deposition and maturation.\(^6\)

One of the main arguments for the use of mechanical load on conditions such as tendinosis is that "fibroblasts in tendinosis are extremely active metabolically and there is a great capacity for the production of collagen. A fibroblast-driven process would be expected to integrate old and new collagen in order to contribute to the final stability of the matrix."\(^7\) Fibroblasts can also convert into myofibroblasts necessary in wound strengthening by extracellular collagen fiber deposition, and wound contraction for closing wounds since myofibroblasts express -smooth-muscle actin. Upon resolution of the injury, these activated fibroblasts (myofibroblasts) undergo apoptosis (programmed cell death).

Tendons that undergo high rates of stretching may be more susceptible to inflammation and eventual degeneration due to the stretching of fibroblasts. Cyclic stretching of fibroblasts, and especially increasing the frequency of the stretching increases the production of pro-inflammatory cyclooxygenase enzyme (COX-1, COX-2) and prostaglandin-E2. COX-1 and COX-2 convert arachidonic acid into prostaglandins. So, overstimulation of the fibroblasts may be responsible for repetitive-motion problems. Stretching also may cause an alignment of the tendon fibroblasts.\(^8\)

Recent studies have shown how eccentric exercises may be more beneficial than concentric exercises regarding the rehabilitation of muscles and tendons.\(^9\) There is reason to believe that the effect of the load pattern of eccentric exercise creates greater stimulation of fibroblasts, increasing collagen synthesis and thereby stimulating the healing of the injured tissue.\(^10\)

Finally, Langevin\(^11\) poses the theory that connective tissue, especially the fibroblasts are part of a whole body cell to cell communication signaling network. She states that fibroblasts exhibit active cytoskeletal responses within minutes of tissue lengthening. "Analogous cell-to-cell signaling involving calcium and/or
ATP may exist within connective tissue and may be accompanied by active tissue contraction or relaxation. One can envisage the whole-body web of connective tissue involved in a dynamic, body-wide pattern of cellular activity fluctuating over seconds to minutes reflecting all externally and internally generated mechanical forces acting upon the body.

Author’s note: The International Myofascial Pain Conference will be held Sept. 3-4; the 2nd Annual Myofascial Pain Conference takes place Oct. 1-2. For more information, visit [www.fascialconference.com](http://www.fascialconference.com).

References


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