An Actin-Myosin Machine

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The Problem: A long standing goal in engineering is to exploit the unique designs of biological systems to guide the development of autonomous biomimetic machines that exhibit agility, strength and speed in a variety of natural environments. Most critical to this effort is the development of an actuator that behaves like muscle. Although important research has been conducted to advance actuator function, engineering science has not yet produced an effective artificial muscle. Polymer gels provide stresses and displacements comparable to muscle, but their contractile velocity is far too slow for many applications[1]. The goal of our work is to investigate the feasibility of using animal-derived muscle as an actuator for robotic applications in the millimeter to centimeter size scale.

Motivation: Muscle is employed almost exclusively by animals for actuation, from sub-millimeter organisms to blue whales. Using muscle to power movement has certain advantages. Muscle generates force quietly, allowing predators to move within close proximity of prey. Muscle is also adaptive and responds to varying work loads by modulating its structure to meet specific tasks demands. Still further, muscle is efficient. Working aerobically, muscle can generate up to 4000kJ of work from just 1 Kg of glucose. And for its size, muscle can generate a large isometric force, enabling the extremities of organisms to be lightweight but strong.

Approach: In our work we are specifically engineering muscle tissue for machine actuation. Genetic, chemical, electromechanical and temperature interventions are being employed to enhance muscle robustness and contractile function in vitro. Two types of muscle tissue are examined: native and cultured tissues from genetically-modified mice (Figure 1) and native whole muscle from non-mammalian sources such as marine invertebrates. Once engineered, the contractility and robustness of these tissues are characterized and comparisons are made to current artificial muscle technologies.

Difficulty: Perhaps researchers in the past did not consider muscle a viable actuator because of tissue robustness problems. It is true that native whole muscle, once extracted from an animal, only retains contractility for several days using standard organ culture techniques[2, 3]. However, recent advances in biology suggest that enhancing tissue robustness in vitro may now be an achievable goal. Preliminary experiments by Rosenthal and collaborators at Massachusetts General Hospital suggest that IGF-I genetic interventions (Figure 1) may improve muscle contractility and robustness. IGF-I genetic manipulations have been shown to enhance the tetanic force of native wildtype muscle by as much as 25 percent[4]. There is also preliminary evidence that these interventions promote cell differentiation and robustness in cultured tissues grown from transgenic mice.

Impact: A muscle actuator may offer certain advantages over contemporary motor technologies. As with real muscle, an engineered muscle could offer a transduction efficiency far superior to that of electric motors, or shape memory alloys, powered by battery or fuel cell. Although high efficiencies can be achieved from gasoline engines, stealth quietness cannot; gas powered autonomous robots are noisy and produce environmentally unfriendly bi-products. In distinction, engineered muscle would generate force quietly with biodegradable bi-products. Finally, there remains the possibility of manufacturing engineered muscle at low cost. Starting from the cells of an animal, it may be possible to grow hundreds of muscle actuators.

Figure 1: The forelimb of a wildtype mouse (left) and a genetically-modified mouse (right) are shown. The pronounced muscular hypertrophy in the transgenic animal is the result locally-acting IGF-I.
**Future Work:** Once we have successfully engineered a muscle actuator, we will then build a robotic fish in which a compliant tail is powered by muscle. Once built, the fish will be our first actin-myosin machine, a muscle-driven robot that feeds on glucose.

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**References:**


