Virtual biomechanics: a new method for online reconstruction of force from EMG recordings

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EVEN SIMPLE MOVEMENTS REQUIRE the coordinated recruitment of multiple muscles. The net joint torques required to perform a movement can be computed from the observable kinematics by the method of inverse dynamics, but these torques might be achieved by many different combinations of individual muscle forces (Bernstein 1967). This is the so-called “redundancy problem,” and how the central nervous system (CNS) selects specific patterns of muscle activation to perform a given task remains one of the most critical unresolved questions in motor control. Indeed, the ability to resolve redundancy is central to the attractiveness of several influential motor control schemes, including optimal control, motor primitives, and hierarchical sensorimotor control (e.g., d’Avella et al. 2003; Haruno and Wolpert 2005; Loeb et al. 1999; Todorov 2004; Todorov and Jordan 2002). Most previous approaches to test predictions of these theories involved observing how muscle activation patterns vary according to task under natural conditions. However, because of the high degree of correlation among muscle activation and limb kinetics and kinematics, many theoretical models might predict similar muscle activation solutions in a given natural situation. A more direct test of a particular hypothesis about the nature of the movement control system would be to see how it responds to changes in the musculoskeletal plant, which could be designed to probe and disambiguate solutions predicted by different theoretical models. Making such changes physically such as by surgical intervention is not ethical with human subjects. Furthermore, it tends to involve a prolonged recovery period, during which adaptations may be occurring but cannot be studied. In this paper, we present a virtual reality method in which subjects control an animated model of their musculoskeletal system in real-time by means of their electromyographic (EMG) signals.

There is extensive literature devoted to modeling methods that allow forward simulation of joint torques or kinematics from estimates of muscle activation. The approaches described previously vary from, at one extreme, models that seek to provide realistic simulations of the physiological processes of excitation-contraction coupling and the mechanical application of muscle forces to generate torques on the skeleton (e.g., Buchanan et al. 2004; Cheng and Loeb 2008; Erdemir et al. 2007; Tsianos et al. 2012) to approaches at the other extreme that are based purely on associations between measured EMG parameters and movement outcomes without consideration of muscle physiology or musculoskeletal mechanics (e.g., Seifert and Fuglevand 2002). Neither of these extreme approaches provides a method for the online reconstruction of joint torque from EMG records that is ideal for subsequent experimental manipulation according to our requirements. On the one hand, approaches based purely on statistical associations between muscle activation and movement outcomes provide no opportunity to manipulate specific aspects of the musculoskeletal plant. On the other hand, the accuracy of force reconstructions produced by a comprehensive neuromuscular-skeletal model remains contingent on accurate settings of the numerous parameters of such models, most of which are difficult to assess on an individual basis (e.g., Sartori et al. 2012). Furthermore, an ideal musculoskeletal model that perfectly represents a subject’s biomechanics would still be insufficient to guarantee the quality of reconstructions because EMG records do not provide a perfectly accurate representation of muscle activation. For instance, EMG signals are subject to contamination by electrical activity of nearby muscles and represent only a fraction of all active motor units in the target muscle (Hug 2011; Staudenmann et al. 2010).
Because an accurate biomechanical model requires an accurate measurement of muscle activation for forward simulation of muscle force, and because the means to measure muscle activation accurately are unavailable, we designed a practical approach whereby a virtual representation of muscle biomechanics was defined that best reconstructs limb force when combined with EMG recordings. In this approach, the inaccuracy of the biomechanical representation is intended to compensate for imperfections of EMG recordings in a manner that best reconstructs force when both are combined. In separate experiments, we demonstrate that the technique works in different contexts of relatively low isometric force at the wrist joint, for which the EMG to force relationship is approximately linear. In particular, we show that the goodness of force reconstruction was similarly high for surface and fine-wire recordings, which are differentially affected by cross talk and vary in the degree to which they represent activity of the overall muscle. We also show that the technique is sensitive to biomechanical changes elicited by a modification of the forearm posture and is therefore suitable to address the important question of how the nervous system tunes motor commands to the biomechanics of the current posture (Buneo et al. 1997; Sergio and Kalaska 1997, 2003). Because it starts from the most intuitive relationship between muscle activity and force and enables virtual change in the biomechanics of any muscle, the technique offers novel opportunities to explore the nature of the adaptive controller embodied by the nervous system.

MATERIALS AND METHODS

Virtual Biomechanics

Overview. Fagg et al. (2002) proposed an optimal model that produces muscle activation patterns qualitatively similar to those observed experimentally for the biomechanics of a known muscle (Fig. 1A). Our virtual biomechanics technique consists in using a similar optimization procedure but in the opposite direction, i.e., to extract a representation of muscle biomechanics from observed muscle activations (Fig. 1B) and then combine the virtual biomechanics with real-time EMG to reconstruct force online (Fig. 2). First, we introduce the Fagg et al. (2002) model; second, we illustrate how using optimization in the reverse direction enables extraction of the virtual biomechanics from muscle activations; and third, we show how combining the virtual biomechanics with real-time EMG recordings enables accurate online force reconstruction.

The Fagg et al. (2002) model. For the biomechanics of a given muscle, Fagg et al. (2002) proposed a method to determine the overall activations of the various wrist muscles without requiring direct information such as EMG. It is based on the assumption that the CNS would minimize the summed squared activation across all muscles. In this model, the extrinsic direction of action (up/down and right/left) of each muscle is defined by a two-element pulling vector, $P_i$, and muscles contribute to the endpoint movement along their vector of action with a length proportional to their activation levels, $a_i$. The endpoint movement is described by the two-element vector $x$:

$$x = \sum_{i=1}^{5} P_i a_i$$

where $A$ is the set of five muscles. The authors then consider the minimization of the following two criteria error function (endpoint error and muscle activation):

$$E = \frac{1}{2} ||x_{targ} - x||^2 + \frac{\lambda}{2} ||a||^2$$

subject to $a_i \geq 0$ for all $i \in A$, where $x_{targ}$ is a vector representing the target location, $\lambda$ is a regularization parameter set to 0.02, $a$ is the muscle activation vector, and $||.||$ denotes the magnitude of a vector. The authors showed that minimizing this cost function produced muscle activation patterns that were qualitatively similar to those observed experimentally in EMG recordings. In particular, these patterns exhibit a cosinelike recruitment of muscles as a function of movement directions and reproduce the observed discrepancies between directions for which muscles are preferentially used and their direction of action (Fig. 1).

Extracting muscle virtual biomechanics from EMG. Assuming that we know the muscle activation patterns (e.g., recorded experimentally) but not the biomechanics, we use optimization in the direction opposite to that used in the Fagg et al. (2002) model to extract a representation of muscle biomechanics from the known muscle activations (Fig. 1B). This was achieved by determining the set of pulling vectors, $P_i$, that resulted in the best aiming performance, i.e., that minimizes endpoint errors, $E = ||x_{targ} - x||^2$, when combined with the actual muscle activation, $a$. To this end, we used a custom coordinate descent algorithm with the following steps. 1) Assign random values to the initial set of pulling vectors in the physiological range of muscle force and direction. 2) Pick a muscle at random and modify its pulling vector by changing its endpoint by a step in 4 orthogonal directions. The target errors associated with each of the 5 pulling vectors (i.e., the
original and the 4 modified for that muscle) were then calculated as
the summed squared error between targets and reconstructed reaches,
and the pulling vector that produced the lowest cost was retained.
3) One iteration of the model was said to be completed when each
muscle had been optimized once. 4) The whole model was iterated
until the overall cost converged to a low value. For the steady-state
isometric tasks presented here, an exact solution can be obtained by
separate minimization of the squared horizontal and vertical errors
using the ordinary least-squares method. For all subjects and experi-
mental sessions reported, we have checked that the coordinate descent
algorithm used successfully converged to the exact solution. The
coordinate descent method will be necessary if the virtual biome-
chanical modeling method is extended to tasks involving nonlinear
dynamic terms.

**Online force reconstruction.** Figure 2 illustrates how we used EMG
recordings during isometric force production at the wrist to extract the
virtual biomechanics as indicated previously and how it was com-
bined with real-time EMG to reconstruct force online. To generate the
time-independent patterns of muscle activation used to extract the
virtual biomechanics, rectified EMG for each muscle was first aver-
aged over the steady, holding phase of the force on target (i.e., during
a time window from 300 to 1,000 ms after movement onset, while the
task was to achieve force targets with a movement time between 150
and 250 ms, and to hold the force cursor on target for 1 s). As indicated
later in this section, participants performed 6 consecutive reaches to each
of the 16 targets, and averages over the 5 last reaches were used to
compute the time-independent muscle tuning curves (i.e., 1st reach
discarded). Once the virtual biomechanics were extracted, each pulling
vector is simply multiplied by $a_i(t)$ and the resultants summed to generate
the time course of the reconstructed force, $\hat{F}(t)$. The activation values
$a_i(t)$ are assumed to be linearly related to the rectified and filtered (1.3-Hz
low-pass) EMG signals, normalized to the largest EMG value obtained
for each in a series of maximal voluntary contractions (MVC) in different
directions (see below).

$$\hat{F}(t) = \sum_{i \in A} P_i \cdot a_i(t)$$

**Experiments**

We tested our virtual biomechanics technique for online force
reconstruction in different experimental context involving participants
to reach isometric force targets in various directions. The technique
was first evaluated with surface EMG (experiment 1, $n = 6$) and
second with fine-wire electrodes (experiment 2, $n = 6$) with the
forearm in a neutral posture (i.e., forearm midpronate as displayed Fig.
2). Then, we tested the sensitivity of the technique to changes in
biomechanics elicited by variation of the forearm orientation along the
supination/pronation axis (experiment 3, $n = 6$). In all experiments, the
real and reconstructed forces were compared during the reaching
task performed with a visual cursor that represented the real force
produced. EMG patterns observed in that context were also compared
with EMG patterns produced when the reaching task was performed
with online reconstructed force as the visual cursor.

**Subjects.** Twelve healthy, right-handed subjects (all men, age
23–38 yr) volunteered for this study. When subjects participated in
more than one experiment ($n = 4$), testing sessions were separated by
at least 3 wk. All subjects had normal or corrected-to-normal vision.
They all gave informed written consent before the experiment, which
was approved by the local ethics committee and conformed to the
Declaration of Helsinki.

**General experimental procedure.** Subjects sat 80 cm from a com-
puter display positioned at eye level. The right hand was maintained
in a custom-made manipulandum with the forearm in one of three
possible positions: in a neutral position for experiments 1 and 2
(midway between pronation and supination as displayed Fig. 2) and in
80° pronation or −80° supination for experiment 3. The elbow was
kept at 110° with the forearm parallel to the table and supported by a
custom-built device similar to that used in a previous study (de Rugy
and Carroll 2010). The wrist was fixed by an array of adjustable
supports contoured to fit the hand at the metacarpal-phalangeal joints
(12 contacts) and the wrist just proximal to the radial head (10
contacts). This allowed wrist forces to be applied without the need for
a gripping force. Wrist forces were recorded using a 6-degrees-of-
freedom force/torque transducer (45E15A-I63-A 400N60S; JR3,
Woodland, CA) coupled with the wrist manipulandum.
Real-time visual feedback of either the real wrist forces or the reconstructed wrist forces was presented on the visual display. Targets were presented at 16 radial positions around the center of the display (i.e., 22.5° apart). In the neutral position, flexion/extension corresponded to the horizontal axis (flexion left), and radial/ulnar deviation corresponded to the vertical axis (radial deviation up). In the 2 other rotated forearm postures, the visual feedback was rotated with the forearm such that the movement of the cursor matched the force direction in external space.

In all experiments, a block of 32 MVC trials was 1st conducted for each subject with the forearm in the neutral posture. This block was used to normalize the activity of each muscle during the aiming task to the maximal EMG obtained in that muscle during MVC toward any target direction. Each of the 16 target directions was presented twice in a randomized order. For each direction, subjects were asked to raise their force rapidly to the maximal extent while maintaining the force direction within a delineated range of ±8° of target direction. Maximal forces were held for ~2 s. Fifteen seconds were allowed for rest before the next target appeared in another direction.

Each experiment contained “force-driven” block(s) in which the visual cursor used to reach targets represented the real force and “EMG-driven” blocks in which the cursor represented the reconstructed force. Each force-driven block consisted of 96 trials (6 trials for each of the 16 target directions) in which a low level of force (i.e., 22.5 N, which represents ~20% MVC for the subjects tested) was required to reach targets. This level of force was identical across all subjects and chosen to reduce the possibility of fatigue. Each trial began only if the cursor was maintained <5% of the target distance from the origin continuously for 200 ms. The origin was calibrated to 0 force along both axes (wrist relaxed) before each block. A random foreperiod (1–2 s) elapsed before a single target appeared coincident with a brief tone. Participants were asked to move the cursor to the target with a movement time of between 150 and 250 ms, defined as the time between 10 and 90% of the radial distance to the target, and to hold the cursor continuously for 1 s within the target zone (a trapezoid ±8° from target direction by 10% of radial distance to target). A high-pitched tone signaled that the target had been acquired. If the target was not acquired within 2 s of target presentation, a low-pitched tone indicated the end of the trial. A 2nd tone (200 ms after the 1st) indicated whether the movement time was correct (high tone) or not (low tone), and a bar graph provided visual feedback of the movement time in relation to the prescribed time window. Both the target and cursor disappeared at target acquisition or trial end, and ±1 s elapsed before the start of the next trial. For each block, 6 consecutive trials were conducted for each of 16 randomly ordered targets. EMG-driven blocks were identical to force-driven blocks with the only exception that the real force feedback was replaced by the reconstructed force.

In experiments 1 and 2, each participant performed one force-driven block immediately followed by an EMG-driven block. In experiment 3, participants performed the same 2 blocks but both in the pronated posture and in the supinated posture (4 blocks total).

**EMG procedures.** Bipolar EMG signals were recorded from extensor carpi radialis longus (ECRL), extensor carpi radialis brevis (ECRB), flexor carpi radialis (FCR), flexor carpi ulnaris (FCU), and extensor carpi ulnaris (ECU) muscles either with self-adhesive surface electrodes (experiment 1, 12-mm diameter recording surface, 2-cm interelectrode distance) or with fine-wire intramuscular electrodes (experiments 2 and 3, 75-μm diameter, 2 mm stripped from insulation for recording sites, single wires inserted at 1.5-cm interelectrode distance, dipole axes approximately parallel to the long axis of muscles). Signals were band-pass filtered from 30 Hz to 1 kHz, amplified 200–5,000 times (P511; Grass Instrument, Astro-Med, West Warwick, RI), and sampled at 2 kHz. Electrode locations were determined according to procedures previously reported (Selvanayagam et al. 2011).

**Data reduction and analysis.** Muscle tuning curves, or the time-independent muscular activity as a function of target direction, were determined for each muscle as the mean rectified EMG during the hold phase of the task (i.e., in a time window from 300 to 1,000 ms after movement onset) averaged over 5 trials to each target (the 1st of the 6 consecutive trials to each target was discarded to prevent the uncertainty about target direction from contaminating the data).

The spatiotemporal match between the real and reconstructed forces, or the goodness of force reconstruction, was quantified by defining a multivariate $r^2$ similar to that used in d’Avella et al. (2006):

$$r^2 = 1 - \frac{\text{SSE}}{\text{SST}}$$

$$= 1 - \frac{1}{\sum_{j=1}^{M} \sum_{i=1}^{N} \left| F(t) - \hat{F}(t) \right|^2}$$

where SSE is the sum of the squared residuals, SST is the sum of the squared residual from the mean force vector ($\bar{F}$), $M$ is the set of trials in a block, and $N$ is the set of time samples per trials. Note that with this calculation, negative values of $r^2$ might occur with particularly poor force reconstruction (i.e., if $\text{SSE} > \text{SST}$).

To test specific hypothesis, the goodness of force reconstruction was computed in four instances after using alternate methods of force reconstruction. In experiment 1, force was additionally reconstructed using a set of virtual pulling vectors computed while completely ignoring either one or two muscles. When ignoring one muscle, each of the five muscles was selectively ignored (i.e., only our pulling vectors determined that best reach the targets), and the goodness of force reconstruction was averaged over the five combinations of one muscle ignored. When ignoring two muscles, all possible pairs of two muscles were considered, and the goodness of force reconstruction was averaged over all combinations. In experiment 1, force was also reconstructed using a set of virtual pulling vectors constrained by morphometric data from cadaver (Loren et al. 1996). Specifically, the directions and relative magnitudes of the pulling vectors were fixed according to data from Loren et al. (1996), and the overall magnitude only was optimized to reach best the targets when combined with muscle activities recorded for each subject. This was designed to compare our method with force reconstructed using a realistic biomechanical model of the wrist. In experiment 4, force was additionally reconstructed using the set of virtual pulling vectors extracted from muscle tuning curves obtained in the other forearm posture. This was designed to ascertain that our reconstruction method is sensitive to biomechanical changes elicited by the different postures. Differences between goodness of force reconstruction obtained for different eros were tested using independent-samples $t$-tests, and differences between goodness of force reconstruction from different reconstruction methods within the same experiment were tested using paired-samples $t$-tests.

In experiments 1 and 2, the time-independent pattern of muscle activities (i.e., muscle activity averaged per target direction) were analyzed using 3-way repeated-measures ANOVAs [2 feedback conditions (force-driven vs. EMG-driven) × 5 muscles × 16 directions of force targets]. In experiment 3, a 4-way repeated-measures ANOVA was conducted using posture (pronation vs. supination) as an additional factor. The significance level was set to $\alpha = 0.05$.

**RESULTS**

**Surface EMG: Experiment 1**

The force reconstructed with our virtual biomechanics on the EMG-driven blocks of this experiment explained 92.0% of the variance of the real force signals (i.e., $r^2 = 0.920 \pm 0.035$, mean ± SD). Figure 3 illustrates that this proportion of explained variance deteriorates when the technique is applied with one missing muscle ($t = 15.34, P < 0.0005; r^2 = 0.671$ ±
When the technique was applied with fine-wire recordings, the EMG-based force reconstruction explained 90.5% of the variance of the real force signals (i.e., $r^2 = 0.905 \pm 0.026$). This proportion of explained variance is similar ($t = 0.85, P = 0.439$) to that obtained with surface electrodes in experiment 1. Figure 5B also illustrates that as for surface recordings, the technique applied with fine-wire recording produces muscle tuning curves that match well between force-driven and EMG-driven conditions. As for experiment 1, the repeated-measures ANOVA revealed strong muscles × directions interaction ($F(60, 300) = 40.87, P < 0.0005$) but no feedback conditions × muscles × directions interaction ($F(60, 300) = 1.28, P = 0.09$), indicating strong muscular tuning that is similar for force-driven and EMG-driven conditions.

**Sensitivity to Biomechanical Changes with Forearm Posture: Experiment 3**

When we tested the sensitivity of the technique to changes in biomechanics elicited by changes in posture, we found that the EMG-based force reconstruction applied to the two different forearm postures tested explained 89.5% of the variance of the real force signals (i.e., $r^2 = 0.895 \pm 0.052$). This proportion of explained variance is very different ($t = 6.78, P < 0.0005$) from that obtained when force is reconstructed for a given posture (e.g., pronation) with the set of pulling vectors extracted from the other posture (e.g., supination; $r^2 = 0.099 \pm 0.529$). Figure 6 illustrates for a representative subject that muscle tuning curves were markedly different for the task performed in supination and in pronation but very similar between force-driven and EMG-driven conditions. The repeated-measures ANOVA revealed strong muscles × directions interaction ($F(60, 300) = 43.79, P < 0.0005$), a strong postures × muscles × directions interaction ($F(60, 300) = 11.93, P < 0.0005$), but no feedback conditions × postures × muscles × directions interaction ($F(60, 300) = 1.23, P = 0.13$). This indicates that muscle tuning curves were strongly tuned to the directions of force targets, that this tuning was different for the two different postures, but could not be distinguished between force-driven and EMG-driven conditions.

**DISCUSSION**

Our goal was to develop a technique that enables accurate online force reconstruction from imperfect EMG recordings. Instead of seeking an accurate biomechanical model, we employed an alternative, practical approach whereby a virtual representation of muscle biomechanics is defined that best reconstructs force when combined with available EMG recordings. The virtual biomechanics method was applied during two-dimensional isometric force at the wrist in a controlled musculoskeletal configuration that restricted changes in muscle length and moment arm. We demonstrated that the technique works for various experimental contexts in which we varied the recordings methods as well as the muscle biomechanics. This method assumes a linear relationship between muscle activation and rectified EMG, which is likely to be true for the relatively small range of isometric forces produced by our subjects in this task.
Robustness to Limitations from EMG Recordings

Two major issues associated with EMG recordings are cross talk and representativeness (Hug 2011; Staudenmann et al. 2010). Cross talk refers to a contamination of EMG signals by electrical activity of nearby muscles, and representativeness refers to the proportion of active motor units captured by the signal. Because electrodes are directly inserted into the target muscle, fine-wire recordings are less subject to cross talk than surface recordings that are more remote and less precisely positioned relative to muscles (Selvanayagam et al. 2012). Fine-wire recordings, however, are more selective to the part of the muscle into which the electrodes are inserted and therefore less representative of the overall activity of the target muscle. By showing that our technique works equally well with both types of recordings, we demonstrated that it is robust to both of their associated limitations. It is worth noting that both cross talk and selective sampling of muscle fibers in the vicinity of the electrodes would affect force reconstructions using forward simulations of an accurate biomechanical model. Although these limitations could be substantially reduced using high-density EMG (Staudenmann et al. 2010), our reconstruction method obviates the need for this technology, which would be expensive and demanding to incorporate the multiple muscles needed for force reconstruction in various directions as achieved here.

Sensitivity to Biomechanical Changes with Posture

As muscle length and moment arms change with musculoskeletal configuration, so changes the torque generation of muscles and therefore the muscle activation patterns capable of
generating a given joint torque in a given posture. How the nervous system adjusts motor commands to the biomechanics of the current posture is a key problem of motor control, which has been explored by simultaneously assessing changes in muscle biomechanics, in muscle activities, and in neural activities at various levels of the CNS (Buneo et al. 1997; Kakei et al. 1999, 2001; Sergio and Kalaska 1997, 2003; Yanai et al. 2008). For instance, nonhuman primate studies reported systematic changes in muscle activity selected to produce force at the wrist in different forearm orientation (Kakei et al. 1999, 2001). In a recent study in which we assessed these changes in humans, we showed in particular that FCR displayed higher activity and broader tuning during force produced with the forearm in a pronated position compared with a supinated position (de Rugy et al. 2012a). A similar pattern is visible in our data displayed Fig. 5 that has also been replicated in our condition in which the task was controlled with force reconstructed from EMG instead of real force. We therefore demonstrated that our force reconstruction technique fully captures biomechanical changes associated with the different forearm postures. This is important because it means that we can use this technique to address the question of how the nervous system tunes motor commands to the biomechanics of the current posture. In fact, we have recently addressed this question by simulating the biomechanics of a different posture to show that participants initially compensate for this perturbation using a linear scaling of their original pattern of muscle activity (i.e., the pattern that corresponds to the real posture; de Rugy et al. 2012b).

Potential Contributions from Hand and Finger Muscles

The current force reconstruction technique has ignored potential contributions from the numerous (i.e., 19) hand and finger muscles that cross the wrist and that have nonnegligible moment arms in wrist flexion/extension and radial/ulnar deviation (e.g., see Fig. 3 in Gonzalez et al. 1997 for a visual representation of these moment arms at the wrist). We believe that the contribution from these muscles to the task was not high at the relatively low level of force involved here (i.e., ~20% MVC). This is because in our device the hand was fitted at the metacarpal-phalangeal joints such that the fingers were hanging in the air with no mechanical contact to the device and subjects were instructed to prevent any forcing or gripping that could bring the fingers in contact with the device. In this context, transmission of force from finger and hand muscles to the device is still possible through cocontraction that would maintain steady finger positions. Cocontractions between opposing finger extensors and flexors would be typically paralleled by opposing wrist extension and flexion moments, thereby reducing the net contribution at the wrist joint. However, the pulling vector discrepancies with respect to the anatomically constrained model appear to involve mostly underrepresented radial torques (see Fig. 4), which are particularly large for thumb extensors and flexors (Gonzalez et al. 1997). To rule out such a contribution, it would be necessary to obtain selective EMGs from at least a representative sampling of the thumb and finger muscles uncontaminated by cross talk from the nearby and simultaneously active wrist muscles. To incorporate such contributions into the model, it would be necessary to obtain quantitative EMGs from all 19 of these muscles during both their maximal activation and a sufficiently rich set of tasks for which their activity would be differentiated. Thus it is important to recognize that the virtual biome-

Fig. 5. Example of muscle tuning curves obtained in force-driven condition and in EMG-driven condition for 2 postures (i.e., ~80° supination and +80° pronation) of 1 subject in experiment 3. The arrows indicate the wrist direction used for reference (i.e., radial deviation). EMG activities are normalized to maximal EMG, and a scale bar indicates 0.1 normalized units (n.u.).

Fig. 6. Example of muscle tuning curves obtained in force-driven condition and in EMG-driven condition for the same subject in experiment 1 (A; surface EMG) and in experiment 2 (B; fine-wire EMG). EMG activities are normalized to maximal EMG, and a scale bar indicates 0.1 normalized units (n.u.).
chanical model is not intended to generate an accurate representation of the work of specific anatomical muscles. It is intended to provide a useful experimental tool to understand sensorimotor adaptation and learning when studied within the context of a well-defined task and subject-specific parameterization of the model. The robust quality of the force reconstructions presented here suggests that this goal has been met.

Differences with Existing EMG-Based Force Reconstruction

Our method is related to a previous method whereby coefficients that relate EMG to force are determined for individual muscles from a data set that requires production of force in various directions and knowledge of muscle moment arms (An et al. 1983; Buchanan et al. 1993; Messier et al. 1971). This approach, termed “coefficient methods” (Buchanan et al. 1993), would be similar to the method described here had we optimized only the magnitude of the muscle pulling vectors, i.e., by constraining the vector directions according to biomechanical data. A drawback of this method is that it implies knowledge of muscle moment arms, which requires magnetic resonance imaging to be accurately assessed on an individual basis (Buchanan et al. 1993). Kutch et al. (2010) recently proposed an alternative method based on surface EMG to extract the direction of action of finger muscles. However, this method might require EMG that represents accurately the activity of the overall muscle, as can be the case with finger muscles, since we were not able to generate reliable direction of action using this method for wrist muscles. In the end, muscle direction of action was unnecessary as our method is free of a priori biomechanical knowledge. We believe that this enabled additional flexibility that was beneficial to the high correlation obtained between forces reconstructed online and real forces.

Another related method was recently proposed whereby a matrix factorization algorithm is applied to surface EMG to extract control signals for prostheses (Jiang et al. 2009; see also Kamavuako et al. 2012). Using forces at the wrist as control signals and a generative model of surface EMG that assumes hard-wired muscle synergies, Jiang et al. (2009) obtained a goodness of force reconstruction that was comparable with that of the present experiments in the absence of simulated muscle cross talk (90.2% variance of the force signals explained) and that degraded with the level of cross talk. In an experiment using eight pairs of recording electrodes arbitrarily placed (equally spaced) around the forearm, they obtained a variance of force signals explained that dropped to 77.5%. The method presented here uses EMG signals that can be identified with specific muscles but compensates for their inevitable shortcomings of selectivity and sampling. This resulted in low residual errors for predicted force as well as the ability to simulate specific changes in musculoskeletal function in virtual force experiments. Neither the matrix factorization algorithm nor the probabilistic method of Seifert and Fuglevand (2002) would be suitable for this application.

Adaptation to Novel Virtual Biomechanics

Over the last few decades, the literature on sensorimotor adaptation has been dominated by two main classes of perturbations: force-field, whereby a force is applied to an end-effector, and sensorimotor shifts, such as in prism adaptation or visuomotor rotation (de Rugy et al. 2009; Gandolfo et al. 1996; Ghiardi et al. 1995; Shadmehr and Mussa-Ivaldi 1994; Shadmehr and Wise 2005; Simani et al. 2007; Welch et al. 1974). The present technique offers opportunities to study adaptation to a new class of perturbation, whereby the virtual biomechanics that link muscle activity to reconstructed force can be modified at will. In particular, this technique enables selective manipulation of properties of individual muscles that is not possible within the broad alteration of sensorimotor mapping induced by force fields or sensorimotor shifts. We have already used the technique to simulate the muscle-specific biomechanics of a different posture as well as the complete loss of a muscle and large amounts of signal-dependent noise added to a muscle (de Rugy et al. 2012b). In all of these conditions, we found that participants compensated for the perturbation using a linear scaling of their original pattern of muscle activity. This has important implications because, although the pattern of muscle activity used to produce force at the wrist is reasonably well-reproduced by optimization models (Diedrichsen et al. 2010; Fagg et al. 2002; Haruno and Wolpert 2005), how the nervous system achieves this behavior remains largely unresolved. For instance, our previous results appear inconsistent with online optimization of muscle activities, as this should have elicited a reoptimization that was not observed when faced to conditions of novel biomechanics. Instead, the observed scaling of the original pattern suggests an important role of the lower sensorimotor circuitry, which might not be readily available to adaptation. Although these results hold only for the brief time scale tested so far, the method presented here could in principle be used to assess adaptation over much longer time scales.

The force reconstruction method allows modifications of the virtual biomechanics that are limited only by imagination. For instance, one might take advantage of the facts that wrist muscles switch their functional relationship depending on the direction of action and that the spinal cord circuitry is known to be intimately related to this adjustable functional relationship (Pierrot-Deseilligny and Burke 2005; Raphael et al. 2010). During wrist extension, the extensor muscles function as agonists, and the flexor muscles function as antagonists, but during radial/ulnar deviation, the extensor muscles (as well as the flexor muscles) oppose each other. Adjacent muscles could therefore be considered as “partial synergists” because they switch from agonist to antagonist based on the direction of action, and diagonal muscles that are farthest apart from each other as “true antagonists” because they always oppose each other (for example, FCR and ECU). The implication of this functional organization on sensorimotor adaptation could be tested using the virtual biomechanics to simulate different arrangements of muscles that would vary the degree of integrity of their functional relationships. Importantly, the current technique has the potential to start from the most intuitive relationship between muscle activity and force (i.e., the natural relationship) before introducing modifications. Myoelectric controllers for prosthetic limbs typically aim for intuitive mappings to reduce learning requirements (Hargrove et al. 2009; Parker et al. 2006; Zhou et al. 2007). However, Radhakrishnan et al. (2008) reported that learning a novel nonintuitive arrangement was more feasible when it involved muscles that were less functionally related. The technique presented here offers new opportunities to explore these issues.
REFERENCES


