A rigorous model of reflex function indicates that position and force feedback are flexibly tuned to position and force tasks

Winfred Mugge · David A. Abbink · Alfred C. Schouten · Julius P. A. Dewald · Frans C. T. van der Helm

Received: 24 March 2009 / Accepted: 5 August 2009 / Published online: 28 August 2009 © The Author(s) 2009. This article is published with open access at Springerlink.com

Abstract This study aims to quantify the separate contributions of muscle force feedback, muscle spindle activity and co-contraction to the performance of voluntary tasks (“reduce the influence of perturbations on maintained force or position”). Most human motion control studies either isolate only one contributor, or assume that relevant reflexive feedback pathways during voluntary disturbance rejection tasks originate mainly from the muscle spindle. Human ankle-control experiments were performed, using three task instructions and three perturbation characteristics to evoke a wide range of responses to force perturbations. During position tasks, subjects \((n = 10)\) resisted the perturbations, becoming more stiff than when being relaxed (i.e., the relax task). During force tasks, subjects were instructed to minimize force changes and actively gave way to imposed forces, thus becoming more compliant than during relax tasks. Subsequently, linear physiological models were fitted to the experimental data. Inhibitory, as well as excitatory force feedback, was needed to account for the full range of measured experimental behaviors. In conclusion, force feedback plays an important role in the studied motion control tasks (excitatory during position tasks and inhibitory during force tasks), implying that spindle-mediated feedback is not the only significant adaptive system that contributes to the maintenance of posture or force.

Keywords Human motion control · Neuromusculoskeletal modeling · Afferent feedback · Golgi tendon organs · Muscle spindles · Muscle force feedback

Introduction

The presently accepted view is that Ia afferents act as excitatory input and Ib afferents as inhibitory input to the alpha motoneuron (Donelan and Pearson 2004). Is this the only possible role of Ia and Ib reflex pathways, or does this viewpoint result from the experimental conditions during which these pathways were studied (passive or statically loaded)? In experimental studies on walking
patterns in decerebrate cats, evidence for a reversal from 
Ib inhibition to Ib excitation was found (Pearson and 
Collins 1993). Also from a theoretical study (Prochazka 
et al. 1997), it was concluded that force feedback is 
generally inhibitory (during stance), but that locomotion 
requires excitatory force feedback, augmenting the muscle 
stiffness. More recently, indirect evidence of adaptation in 
force feedback was found during human locomotion 
(Geyer et al. 2003; Grey et al. 2007), as well as experi-
mental evidence suggesting that in humans Ib inhibition is 
only present in non-loaded situations such as during sit-
ting and being in supine positions (Faist et al. 2006). 
However, in the latter case, possible contributions of Ia 
reflex pathways to the measured activity could not be 
conclusively ruled out.

The aim of this study was to simultaneously quantify 
contributions of both Ia and Ib pathway activity to the 
overall motion control behavior, not only during passive 
conditions, but also during tasks that require the preser-
vation of posture or force. When attempting to understand 
the control of human motion during various functional 
voluntary tasks, one needs to investigate how the neuro-
muscular system acts as a whole and how the individual 
components of the neuromuscular system interact and 
contribute to its overall dynamic behavior. The field of 
control engineering provides tools to investigate dynamic 
systems, such as the neuromuscular system, and to suc-
cessfully estimate the contribution of the individual sys-

tem components (e.g., Kearney et al. 1997; Kiemel et al. 
2006; Van der Helm et al. 2002; Van der Kooij et al. 2005).

Human motion control behavior in control theoretic 
terms

The essential components of the human motion control 
system can be compared to those of a robot: a linkage 
system (skeleton), actuators (muscles), a sensor system 
(proprioceptors) and a controller (the central nervous sys-
tem or CNS), which is connected to the actuators and 
sensors (via the nerves). Several control strategies can be 
recognized, for example feed-forward (voluntary, planned 
and anticipated movements) and feedback (reflexive 
movements). Unpredictable disturbances, which occur 
during many activities of daily living, cannot be antici-
pated. This is why understanding mechanisms underly-
ing sensory feedback is crucial (Shadmehr and Krakauer 
2008).

Apart from relatively slow feedback that requires cor-
tical processes (e.g., visual feedback), two principal stra-
egies that allow humans to change the resistance of a limb 
to unpredictable perturbations exist:

1. instantaneous resistance, arising from passive struc-
tures and from (co-)contracted muscles (increased 
contraction enlarges the muscle visco-elasticity);

2. reflexive feedback from sensory organs in the muscle 
(proprioceptors) and from tactile sensors in the skin.

Both mechanisms are effective during posture mainte-
nance, each with its advantages and disadvantages. 
Reflexive feedback only provokes muscle activity in 
response to a perturbation, making it energy-efficient. On 
the other hand, the inherent neural transport latency limits 
its effectiveness at higher frequencies. The effect of co-
contraction is instantaneous and, therefore, also effective at 
higher frequencies at the expense of greater energy 
consumption.

The role of proprioceptive reflexes in motion control has 
been the subject of both physiological and experimental 
studies. Physiological studies have described the two 
known proprioceptors in detail: muscle spindles (MS) and 
Golgi tendon organs (GTO). MS are sensors parallel to the 
muscle fibers, which provide feedback to the CNS through 
Ia afferents (providing muscle stretch and velocity infor-
mation) and through II afferents (mainly providing muscle 
stretch information). GTO are located in the muscle tendon 
and provide feedback through Ib afferents (providing 
muscle force information).

Many experiments have been performed that aim to 
isoalte the reflexive contribution by studying electromy-
ográfical (EMG) and mechanical responses to pertur-
bations. Literature shows that these responses depend on 
task instruction (Hammond 1956; Crago et al. 1976; 
Rothwell et al. 1980; Jaeger et al. 1982; Doemges and 
Rack 1992a, b; Abink 2006; Pruszynski et al. 2008), the 
level of muscle contraction (Jaeger et al. 1982), the dis-
placement amplitude (Cathers et al. 1999; Stein and 
Kearney 1995), the bandwidth of the perturbation signal 
(Van der Helm et al. 2002) and the mechanical load 
which the subject interacts with (De Vlugt et al. 2002). 
Comparisons and results between the many studies are 
difficult to interpret due to the abundance of experimental 
conditions: the measured joint, the type of perturbation 
(force or position, transient or continuous), and, often not 
explicitly stated, the task instruction. The reflexive con-
tribution is usually attributed to MS-mediated feedback. 
Variations in the reflexive feedback contribution are 
ascribed to presynaptic Ia inhibition and gamma 
activation, according to the ‘fusimotor set’ hypothesis 
(Prochazka et al. 1985). Both are established physiologi-
cal mechanisms for the muscle spindles. Physiological 
literature has shown mechanisms that may realize varia-
tions in GTO activity: “Ib fibers bear adequate synaptic 
equipment to receive substantial presynaptic inhibition” 
(Rudomin and Schmidt 1999).
Evoking a wide range of motion control behavior

This study aims to evoke a broad range of control strategies of the subjects and as such provoke reflex modulation. For example, in dynamic experiments a position task can be expected to involve mainly position feedback, while a force task is likely to benefit from force feedback (Mugge et al. 2009). In this study, subjects were required to perform three tasks in face of continuous torque perturbations, according to the following task instructions:

- **Force task (FT)**, i.e., maintain force, minimize force deviations.
- **Relax task (RT)**, i.e., passive behavior, minimize muscle activity.
- **Position task (PT)**, i.e., maintain position, minimize position deviations.

The dynamic control behavior as a whole can be quantified by the admittance: a measure of the displacement magnitude due to a force or torque, as a function of frequency. A small admittance (simply put: high stiffness) results in relatively small displacements in response to an external torque perturbation. This behavior is effective during PTs, such as posture maintenance or perturbation suppression. Daily life examples include steering a car on a bumpy road or holding an umbrella in a storm. It is hypothesized that during PT, the admittance is decreased as a result of co-contraction combined with excitatory MS feedback (Van der Helm et al. 2002). No information is yet available about the role of GTO feedback during PTs. RTs are hypothesized to result in negligible co-contraction and little MS and GTO feedback, effectively evoking a large admittance. FTs require a constant force to be exerted on an object (e.g., maintaining the pressure of a pencil on paper on the bus or holding a full cup of coffee during turbulence in a plane). During FT, force deviations are minimized, which is best accomplished by being compliant. It is hypothesized that FT performance is enhanced by active reflexes, helping to give way to the force perturbations, thereby reducing the overall stiffness of the system. Therefore, behavior with a large admittance is expected and even more compliant than during RT. Excitatory MS feedback and co-contraction increase stiffness (and damping) and are therefore counterproductive during FTs, while inhibitory GTO feedback decreases stiffness, improving FT performance. Therefore, it is hypothesized that FT entails inhibitory GTO feedback and little or no MS feedback, thus resulting in assistive muscle contractions, reducing the contact force in the absence of co-contraction. The hypotheses are summarized in Table 1.

![Table 1](image)

To test these hypotheses, first the admittances of ten subjects performing the three tasks in face of continuous random torque perturbations were experimentally obtained.
weight of the foot. This allowed the subject to relax the lower leg muscles in the desired configuration.

Before the experiment, the subjects received specific instructions on how to perform the three tasks: the force task (FT), relax task (RT) and the position task (PT). To prevent drift from the reference, visual feedback was presented on a 17-inch monitor mounted in front of the subject during FT and PT. During PT, the gas pedal angle was plotted against a horizontal reference line indicating the target angle ($10^\circ$ pedal depression), whereas during FT the torque exerted on the gas pedal was plotted against a horizontal reference line indicating the target torque. The target torque was equal to the weight of the foot at the desired position (approximately 28 N at foot contact, again $10^\circ$ pedal depression), so that no muscle force was required. During RT, the monitor was turned off.

For each of the three tasks, three torque perturbation signals were designed yielding nine conditions (each a combination of a perturbation bandwidth and a task instruction). Each condition was presented to the subject four times for averaging purposes; yielding 12 trials per task (and 36 trials in total). The order in which subjects had to perform the three tasks was randomized for each subject, the order of the trials was randomized as well, but grouped to corresponding task instructions to improve overall task perception. Every task instruction was accompanied by a 5-min break to prevent fatigue and 10 min of training.

**Signal recording**

The torque on the gas pedal $T_c(t)$ was measured by a force sensor (Interface, 1210BXH-300, positioned at a moment arm ($d_1$) of 76.2 mm). The gas pedal angle $\theta_{\text{pedal}}$ was measured by an angle encoder. The signals were recorded via analog-to-digital conversion at 250-Hz sample frequency and digitally stored. Furthermore, differential surface electrodes (Delsys) recorded EMG of four muscles: ankle plantarflexors (pushing down), gastrocnemius medialis (GM), gastrocnemius lateralis (GL) and soleus (SOL); and dorsiflexor (pulling up), tibialis anterior (TA). The
EMG signals were pre-amplified, low-pass filtered at 450 Hz and high-pass filtered at 20 Hz, before being digitized at 1,250 Hz by a separate system. For each trial, the recorded EMG was rectified, normalized to the EMG during maximum voluntary contraction (MVC) and finally averaged over time. Dorsiflexor EMG consisted of EMG of TA only, while plantarflexor EMG was the average of EMG of GM, GL and SOL muscles. The EMG during MVC was obtained before and after the experiment and averaged. The EMG normalization allowed for inter-subject comparison.

The first 3 s and the last second of all measured signals were excluded from analysis to reduce the influence of possible transient effects (due to the onset of the perturbation or an imminent stop).

**Perturbation signals**

The three perturbation signals were designed in the frequency domain to have rectangular spectra containing dominant power from 0.1 to 0.2 1.0 and 2.0 Hz. Power was applied to two adjacent frequency points, to enable frequency averaging to reduce estimator variance (Jenkins et al. 2008) and is shown in Fig. 2. The model linearization systems with no linear relation to one for a linear system without noise (linearization or measurement noise).

Model parameterization

To obtain physiologically relevant parameters, a neuromusculoskeletal model was fitted to the experimental data. The model describes the admittance with 13 physiologically interpretable parameters: 5 that represent condition-dependent parameters and 8 that represent condition-independent parameters. The condition-dependent parameters describe the strength of reflexive feedback from muscle spindles (position feedback gain $k_p$, velocity feedback gain $k_v$), Golgi tendon organs (force feedback gain $k_f$) and muscle visco-elasticity (muscle stiffness $k_a$, muscle viscosity $b_a$). The condition-independent parameters all described processes that were assumed not to change with task instruction or perturbation bandwidth, but should be able to vary between subjects: i.e., ankle inertia ($I_a$), tendon stiffness ($k_t$), reflexive time delays (for MS $\tau_{ms}$, for GTO $\tau_{gto}$), second-order activation dynamics (relative damping $d_c$, eigen-frequency $f_c$) and contact dynamics (damping $b_c$ and stiffness $k_c$). The full model has been described previously (Schouten et al. 2008) and is shown in Fig. 2. The model linearization and derivation are described in Appendices 1 and 2. The outcome of a parameter fit can strongly depend on the choice of model structure and parameter boundaries. As a result, an incorrect choice could affect the validity of the conclusions of this study. A substantial part of “Discussion” will address the ability of the model to describe the measured behavior, as well as to what extent the conclusions depend on the chosen model assumptions. Particularly, the effect of long-latency reflexes instead of short-latency reflexes and the omission of GTO feedback are assessed.

The complete neuromusculoskeletal model describes the admittance according to:

$$H_{model} = \frac{\theta_{pedal}}{T_c} = \left[ H_c^{-1} + H_i + H_{ms}H_{act}H_{filter}} \right]$$

where $H_c$ describing the contact dynamics, $H_i$ the inertial dynamics, $H_{ms}$ the muscle visco-elasticity, $H_{act}$ the muscle

The procedure estimates the admittance without any predefined model structure, but does assume linearity, which can be checked by the coherence according to:

$$\Gamma^2_{DD}(f) = \frac{|\hat{S}_{DD}(f)|^2}{\hat{S}_{DD}(f) \cdot \hat{S}_{DD}(f)}$$

The squared coherence function $\Gamma^2_{DD}$ ranges from zero for systems with no linear relation to one for a linear system without noise (linearization or measurement noise).

Non-parametric system identification

The admittance was estimated in the frequency domain, using a closed-loop frequency domain identification procedure (Van der Helm et al. 2002), according to:

$$H_{TD}(f) = \frac{\hat{S}_{DD}(f)}{\hat{S}_{DT}}$$

in which $\hat{S}_{DD}(f)$ denotes the cross-spectral density of external torque perturbation $D(f)$ and gas pedal angle $\theta_{pedal}(f)$, and $\hat{S}_{DT}$ denotes the cross-spectral density of external torque perturbation $D(f)$ and gas pedal torque $T_c(f)$.
spindle activity and $H_{\text{act}}$ the muscle activation dynamics. The equation was condensed using the following equations:

$$H_{\text{filter}} = \frac{\text{H}_{\text{tendon}}}{\text{H}_{\text{tendon}} + \text{H}_{\text{ve}}}$$

$$H_{\text{filter}} = \frac{1}{1 - \text{H}_{\text{filter, tendon}} \left( -\text{H}_{\text{GTO}} + \left( \frac{\text{H}_{\text{ve}}}{\text{T}_{\text{musk)}}} \right) \text{H}_{\text{act}} \right)}$$

where $H_{\text{tendon}}$ describes the tendon stiffness, and $H_{\text{GTO}}$ the Golgi tendon organ activity.

**Parameter fit procedure**

The parameter fit was done in the frequency domain, to the admittances estimated from noise-free signals generated using auto-regressive models (Van den Hof et al. 1995). The 13 parameters were identified by fitting the model on the frequency response function (FRF) of an ARMAX approximation of the measured signals, a technique known to improve the fit (De Vlugt et al. 2006).

The fit procedure was guided by parameter boundaries (see Table 2), which served to prevent unrealistic parameter values and bad convergence. As a fit criterion, the total of the squared logarithmic difference in admittance was minimized (Pintelon et al. 1994) over all conditions and frequencies, fitting all parameters for each subject in one optimization, according to:

$$E = \sum_{\text{cond}(k)} \sum_{\text{freq}(m)} \log \left[ \frac{H_{\text{ARMAX}}}{H_{\text{Model}}} \right]^2$$

with $\text{freq}(I)$ representing the lowest frequency, $\text{freq}(m)$ the highest, $\text{cond}(I)$ the first condition (task x bandwidth), $\text{cond}(k)$ the last condition, $H_{\text{ARMAX}}$ the FRF of the ARMAX approximation to the measurement data and $H_{\text{model}}$ the FRF of the model. The error $E$ was summed over all frequencies and all experimental conditions. Five parameters were estimated for each of the nine experimental conditions and 8 parameters were kept constant over all conditions, resulting in a total of 53 parameters per subject.

**Parameter fit validation**

As a measure of the validity of the ARMAX approximation as well as of the model fit, the variance accounted for (VAF) was calculated for both steps (Eq. 5). The VAF relates how much of the variance of a signal ($u_1$) is captured by a predicted signal ($u_2$). A VAF of 100% means that $u_2$ perfectly describes $u_1$.

The VAF for the position was calculated from measured position ($u_1$) and the model-predicted position ($u_2$), yielded by time domain simulation of the model with the fitted parameters, according to:

$$\text{VAF} = \left[ 1 - \frac{\sum_{q(1)} \left( u_1(t) - u_2(t) \right)^2}{\sum_{q(1)} (u_1(t))^2} \right] \cdot 100\%$$

with $q(1)$ representing the first sample and $q(n)$ the last sample of the time signal. In this study, the first sample was chosen after 1.6 s to exclude transient effects of filters (ARMAX) and integrators (parametric model). The same procedure was followed to calculate the VAF for force.

**Statistical analysis**

The effect of task and perturbation bandwidth on the admittance was tested for significance using a repeated measures ANOVA ($p < 0.05$) on the magnitude of the admittance over all subjects, for each frequency point. Effects on the five condition-dependent parameters were also tested for significance using a repeated-measures ANOVA over all subjects, for each parameter.

**Results**

**Time domain analysis**

Figure 3 shows a time segment of a few seconds, to illustrate the difference between FTs and PTs. First of all, it shows the task-dependent difference in magnitude of the torque perturbations and contact torques at the same
angular deviations: an indication of a difference in admittance. Second, during FTs, the resultant torque $T_c$ is almost zero, while during PTs $T_c$ counteracts the perturbation more. Also, PTs entail much EMG activity.

An intriguing difference between FTs and PTs, is that during FTs the EMG activity is in phase with the torque, while during PTs it is in anti-phase. In other words, during FTs subjects actively used muscle activation strategies that give way to the perturbations, while during PTs subjects counteracted the perturbations.

Figure 4 shows the mean EMG level for all conditions. During PTs significantly ($p < 0.01$) more EMG activity was found compared to FTs and RTs, indicating increased muscle co-contraction. FTs entail a slight, but significant ($p < 0.01$), increase in EMG with respect to RTs. Within the tasks no significant effect of bandwidth on the mean EMG level was found.

Non-parametric frequency domain identification

As hypothesized, the admittance was substantially influenced by task instruction. Figure 5 presents the admittance during four repetitions of one subject performing the three task instructions. The largest admittance occurs during FT (i.e., minimized force deviations), a smaller, but still large, admittance during RT (i.e., minimized muscle activity), and a small admittance during PT (i.e., minimized position deviations). These task-dependent changes in admittance were consistent over all subjects: a significantly ($p < 0.01$) higher admittance was found for FTs with respect to RTs up to 3.0 Hz, and a significantly ($p < 0.01$) lower admittance was found for PTs with respect to RTs up to 1.0 Hz. The (squared) coherence was high (not shown, $\approx 0.9$ for all frequencies) indicating that the input–output behavior can be considered linear with low levels of noise.

In addition, significant effects of perturbation bandwidth were found. As can be seen from Fig. 6, perturbations with dominant power up to a lower frequency result in improved task performance: an increased admittance during FTs and a decreased admittance during PTs.

Parametric frequency domain identification

The eight condition-independent parameters (limb inertia, tendon stiffness, reflexive time delays, muscle activation dynamics and contact visco-elasticity) are shown in
Table 3 (mean ± SD over subjects). Figure 7 shows the five condition-dependent parameters (position and velocity feedback originating from MS, force feedback originating from GTO, and muscle visco-elasticity). The effect of task instruction on the condition-dependent parameters is in accordance with the hypotheses (inhibitory GTO during FT, excitatory MS during PT), except for two remarkable results. In addition to the inhibitory GTO feedback during FT, inhibitory MS feedback was found, and in addition to the excitatory MS feedback during PT, excitatory GTO feedback was found.

Adaptations in force feedback from GTO

The effect of task instruction on the force feedback gain $k_f$ was substantial. The gain was even shown to change sign (i.e., from inhibitory to excitatory). Significantly ($p < 0.05$) more inhibiting GTO feedback ($k_f = 1.2$) was found during FTs at bandwidths 0.7 and 1.2 Hz with respect to all other conditions, including the FT with 2.0 Hz perturbation bandwidth ($k_f = 0.2$), indicating a bandwidth effect during FTs.

Significantly ($p < 0.01$) more excitatory GTO feedback ($k_f = -1.0$) was found during PTs with respect to RTs ($k_f = 0.1$) and FTs ($k_f = 1.2$). A significant ($p < 0.05$) bandwidth effect on the GTO gain was found during PTs as well.

Adaptations in position and velocity feedback from MS

The results for the MS position and velocity feedback gains ($k_p$ and $k_v$) show more variability, and were generally less interpretable. The gains did not change significantly with bandwidth, and a task effect was only found for $k_v$: during PTs an inhibitive gain ($k_v = -17 \text{ Nm rad}^{-1}$) was estimated for 0.7 Hz and 1.2 Hz perturbation bandwidth, significantly ($p < 0.05$) lower than for other conditions.

Adaptations in muscle visco-elasticity

Even though muscle viscosity and elasticity are physiologically related, they were modeled independently. Still, the parameters jointly changed with task instruction: during PTs, substantially increased stiffness ($k_a = 550 \text{ Nm rad}^{-1}$)
and damping ($b_a = 10$ Nm s/rad) were found compared to FTs and RTs ($k_a = 180$ Nm/rad, $p < 0.01$ and $b_a = 7$ Nm s/rad, $p < 0.05$). Bandwidth did not significantly influence the muscle visco-elasticity. These results correspond with the EMG level results of the time domain analysis, and show that the parameter estimation retrieved the physiological relation between muscle viscosity and elasticity.

**Validation**

The fit procedure yielded a parameterized model that accurately captures the dynamics: Fig. 8 shows the magnitude and phase of the admittances during FT, RT and PT as estimated with the parameter fit (solid line), with the ARMAX-approximated signals (dashed line) and with the measured signals (dotted line).

The VAFs between the model-predicted signals (contact torque and angular position) and the ARMAX approximations are presented in Table 4, together with the VAFs between the ARMAX approximations and the measured signals. All VAFs were generally very good for RTs and PTs, but lower during FTs.

**Discussion**

The effect of task effect and perturbation bandwidth on admittance

Consistent with previous studies, task instruction had a substantial effect on joint admittance (Hammond 1956; Crago et al. 1976; Rothwell et al. 1980; Jaeger et al. 1982; Doemges and Rack 1992a, b; Abbink 2006; Pruszynski et al. 2008). As hypothesized, for all subjects the largest admittance was found during FTs, a smaller admittance during RTs, and the smallest admittance was found during PTs. During FTs, slightly more EMG activity was found compared to RTs (see Fig. 4), indicating that subjects used their muscles to actively give way. During PTs, much more EMG activity for all relevant muscles was found compared to RTs.

Similar results were found for FT, RT and PT in single-joint experiments in the same (Abbink 2007) and other joints (wrist) and in multi-joint systems such as controlling a steering column and side-stick in an airplane (Lasschuit et al. 2008), in which also an increased admittance for the FT was found with respect to the RT.

The perturbation bandwidth significantly influenced the admittance for both FTs and PTs: the lower the perturbation bandwidth, the better the subjects performed their task. This bandwidth effect was previously reported for PTs during arm posture control (Van der Helm et al. 2002), where it was explained by a trade-off between performance and stability. Theoretically, a large amount of position feedback activity would be optimal during a PT. However, due to the neural conduction and synaptic delays, position (and velocity) feedback causes oscillations at its eigen-frequency, which is counterproductive to task performance. Therefore, when a perturbation contains substantial power near this eigen-frequency, the feedback gains are suppressed. The small bandwidth perturbations in this study evoke low-frequency admittance changes that are substantially larger than reported in previous literature (Colebatch and McCloskey 1987; Hore et al. 1990). It is presently unclear why these differences exist. Possibly, the present method elicited more reflexive activity due to the provoked functional behavior (task instruction with motivating visual feedback) and the perturbation properties that caused reflexive activity to be exceptionally effective (Mugge et al. 2007).

Unexpected result: excitatory GTO feedback

The model results section shows that the model that included inhibitory as well as excitatory MS and GTO yields accurate fits for all conditions. This may be surprising, given the prevailing neurophysiologic view of
GTO being essentially inhibitory and MS excitatory. Could a model with only MS yield the same fits? The solid line in the left panel of Fig. 9 shows the results of a typical parameter fit for PT, using a model comprising all parameters except the GTO force feedback gain $k_f$. The reduced model could not accurately capture the dynamics of the admittance during FT and PT, resulting in bad fits, substantially reduced VAFs and unrealistic parameter values. Could a model with only inhibitory GTO yield accurate fits? The implementation of strictly inhibitory GTO muscle force feedback did result in accurate fits for FT; however, the behavior during PT can only be accurately described with a model that incorporates excitatory muscle force feedback.

Neurophysiologic background for inhibitory and excitatory reflexive feedback

Is there a physiological basis for the assumption that MS activity and GTO activity can be both excitatory and inhibitory? The overall effect of Ia afferent information on the efferent output of the alpha motor neuron originates from a direct excitatory connection (monosynaptic, but affected by presynaptic inhibition) and from indirect connections through excitatory and inhibitory interneurons in the spinal cord (Jankowska and McCrea 1983). The pathways with inhibitory interneurons can result in an overall inhibitory effect of Ia on the alpha motor efferent output (Stienen et al. 2007). Shared interneuronal circuits resulting in inhibitory and excitatory effects of Ib afferent input to the alpha motor neurons have been described (Jankowska and McCrea 1983) in the cat.

Experimental findings from literature report excitatory GTO feedback during walking (Faist et al. 2006; Grey et al. 2007). Apparently, the same neurophysiologic mechanisms responsible for excitatory Ib feedback during walking apply also for the non-walking PT in this study.
More specifically, results from our parameter fit suggest an important behavioral role for GTO activity in face of low-frequent perturbations: inhibitory to increase the admittance during FTs and excitatory to decrease the admittance during PTs. The slightly excitatory MS position feedback during PTs did not differ significantly from the other tasks, and only the inhibitory velocity feedback was significant. This inhibitory feedback was not expected, since on its own it increases the admittance, degrading PT performance. A separate model sensitivity analysis at the ankle using the NMCLab Graphical User Interface (Schouten et al. 2008) yielded evidence that slightly inhibitory velocity feedback has a stabilizing effect near the GTO’s eigen-frequency, allowing more excitatory GTO feedback: an interesting notion that demands additional research. Moreover, it also revealed that \( k_p \) and \( k_v \) had less impact on the fit accuracy than \( k_f \) (which may explain the large variability in the estimated muscle spindle feedback gains).

The effect of task and perturbation bandwidth on estimated parameters

The parameter identification revealed how reflexive feedback and muscle co-contraction interact to yield the observed admittances. As hypothesized, FTs evoked inhibitory force feedback gains. The corresponding small time delays (25–30 ms) indicate that the force feedback originates from reflexive GTO activity, rather than from slower tactile feedback. The admittance increase of FT with respect to RT was not accompanied by significant changes in intrinsic visco-elasticity or MS feedback, suggesting that the main contributor to the increased admittance was the GTO feedback. As was hypothesized for the RT, little muscle visco-elasticity, MS feedback or GTO feedback were present, indicating that during these perturbations an RT constitutes a good reference condition rather than being well suited for studying reflexes. The small reflexive feedback that was present during RTs was slightly inhibitory for the GTO and slightly excitatory for the MS, which corresponds to the role for the two proprioceptors commonly found in experimental studies that do not consider the impact of functional task instruction.

With regard to the effect of perturbation bandwidth, the parameter identification confirms previous findings (Van der Helm et al. 2002) that at lower perturbation bandwidths, more MS velocity feedback is present than at larger bandwidths. In the present study, the same bandwidth dependency was found for force feedback during FTs: a larger GTO gain was estimated at lower bandwidths. Note that the changes in admittance cannot be explained by changes in muscle visco-elasticity, since no significant changes were found with bandwidth (which was supported by the fact that the mean EMG did not vary with perturbation bandwidth).

Limitation 1: assumption of linearity

The presented method can only be used to describe the linear response to the perturbations. For the current experimental conditions, the linear response generally dominates the observed behavior, as can be concluded from the high coherences and the high VAFs of the linear model fit. However, during FTs the VAFs were lower though the coherences were high. Apparently, the linear part of the behavior could be accurately measured and the model could accurately describe that behavior. However, the lower VAFs indicated that some non-linear behavior was present that could not be described by the model. Close inspection of measured torque and position signals showed that power was present on frequencies where the perturbation did not contain power. Some part of the subject’s response did not have a linear relation with the perturbation, which is likely due to intermittent control actions. Most subjects indicated to have more difficulty performing FTs than the other tasks, in particular with the FT with 2.0 Hz perturbation bandwidth. Note that during FTs, the force can be controlled at different operating points, which allows some drift in the position (reducing the position VAF, but hardly influencing the frequency domain fit).

Limitation 2: parameter interplay

The human body has redundant mechanisms at its disposal to realize a wide range of neuromuscular dynamics. Consequently, a model that aims to describe the separate contributions of these mechanisms is over-parameterized by definition. The parameter fit procedure utilizes the specific characteristics of each mechanism in the frequency domain. However, due to substantial interplay between these modeled mechanisms, multiple combinations of parameters can yield approximately the same admittance. The parameter fit procedure converged to a unique solution by fitting the model to a diverse set of conditions simultaneously, while assuming certain condition-independent parameters (e.g., inertia, reflexive time delays), which effectively reduced the number of parameters to be fitted.

Confidence in the validity of the parameters was obtained from the good fits in both frequency (Fig. 8) and time domain (Table 4), by good parameter convergence from different starting values and by the fact that the measured mean EMG resembled the estimated visco-elasticity. Moreover, parameter values that have been previously determined (e.g., Kirsch and Kearney 1997) through other means (e.g., inertia, tendon and muscle stiffness, and
reflexive time delays) converged to similar values as found in the current study.

Limitation 3: spinal versus cortical feedback

Another concern in the choice of the model structure is the possible presence of other feedback pathways: visual and tactile feedback might also be a contributing factor to the low-frequent changes in admittance. However, motion control in response to only visual feedback is possible for up to 6 rad/s (~1.0 Hz) at best (McRuer and Jex 1967); the visual control loop is characterized by a relatively large time delay (above 200 ms). Although the time delay for the tactile loop is smaller (approximately 120 ms), it is still substantially larger than the delays estimated in this study for GTO and MS. Visual and tactile feedback is therefore too slow to explain the significant increase in admittance up to 3.0 Hz during FTs. Still, there is a potential contribution at low frequencies, and this interplay may lead to reflex gains different from those found in the current study. To investigate this, a brief model study was done where the neuromusculoskeletal model was supplemented with visual and tactile feedback, modeled as feedback gains and large time delays. Parameter estimation of that model resulted in visual and tactile feedback gains explaining some of the low-frequent adaptations. As a result, the reflexive feedback gains were estimated to be somewhat lower, but still the same qualitative trends were observed in the reflexive parameters (GTO sign change from FT to PT and bandwidth effect). Moreover, without reflexive feedback, the model could not accurately describe the observed admittances. In other words, while the quantitative magnitude of the estimated reflexive gains in this study may not be accurate, the qualitative adaptations in GTO and MS feedback are needed to explain the observed behavior.

Limitation 4: short-latency versus long-latency reflexive feedback

In addition to cortical feedback pathways ("Limitation 3"), an influence of slow spinal feedback can be expected (Grey et al. 2001; Petersen et al. 1998). After all, medium- and long-latency reflexes have been shown to be task dependent (Crago et al. 1976; Rothwell et al. 1980). Implementation of every single physiologically known pathway into the model would certainly result in a more detailed, but also a greatly over-parameterized model (see "Limitation 2"). To investigate whether the main reflexive contribution to the measured behavior was from short or long-latency reflexes, the model parameterization was repeated with a neuromusculoskeletal model where the short-latency reflexes were replaced by long-latency reflexes (time delay fit boundaries of 80–120 ms). The solid line in the right panel of Fig. 9 shows the parameter fit with the long-latency reflexes for a typical subject. The parameterization still fitted high feedback gains to yield the large low-frequent admittance changes. However, due to their larger time delays, a trough in the modeled admittance appeared around 1 Hz, which did not correspond to the measured admittance. Note that although it can be concluded from this result that short-latency reflexes are required to accurately fit the data, this does not rule out contributions by medium- and long-latency reflexes.

The fact that short-latency reflexes are required to accurately fit the data in this study could be because it focused on single-joint behavior. It has been suggested in literature that the most important function of the short-latency reflexes is to compensate for muscle yielding in stretched muscles (Nichols and Houk 1976; Crago et al. 1976; Allum et al. 1982; Allum and Mauritz 1984; Sinkjær et al. 1996). Long-latency reflexes are primarily involved in multi-joint movements (Gielen et al. 1988; Soechting and Lacquaniti 1988; Perreault et al. 2006; Kurtzer et al. 2008), where coordinated muscle activation patterns (such as synergies) are controlled by supraspinal brain regions.

Measuring GTO and MS feedback simultaneously

The common approach to examine reflexive behavior, based on applying transient perturbations (such as the deep tendon reflex), cannot distinguish between GTO and MS feedback. One way to distinguish contributions of GTO and MS activity is to isolate reflexive activity in response to only muscle stretch, or only muscle force (Doornik and Sinkjær 2008). However, that approach does not apply to motion control tasks where both GTO and MS activity may be present. MS, GTO and muscle visco-elasticity each contribute to the muscle force and mutually affect each other. The current study used closed-loop modeling as a way to distinguish between GTO and MS activity, and as such is the first to identify how GTO and MS feedback act together simultaneously to realize task performance.

Conclusions

The experiment shows that the dynamic human response to perturbations (i.e., the admittance) is strongly influenced by task instruction and the perturbation bandwidth. Parameter identification of a physiological model provides insight into how feedback from Golgi tendon organs, muscle spindles and muscle co-contraction simultaneously contribute to the observed control behavior.

Relax tasks evoked a large admittance, realized by little co-contraction or reflexive activity. A continuous FT results in an even larger admittance, mainly caused by
inhibitory Golgi tendon organ feedback, accompanied by little co-contraction. A continuous PT results in a much smaller admittance, mainly caused by high levels of co-contraction and excitatory feedback from Golgi tendon organs, accompanied by excitatory position feedback and inhibitory velocity feedback from muscle spindles. Moreover, during continuous force and position tasks with large bandwidth perturbations, the reflexive feedback is suppressed, most likely to maintain stability.

Although many questions remain about the functionality of Golgi tendon organs, the results indicate that the strength of Golgi tendon organ feedback is adaptable and that it plays an important role during continuous tasks where force or position needs to be controlled.

Acknowledgments We thank E. de Vlugt for support. This research was funded by the Dutch Government Grant BSIK03016.

Open Access This article is distributed under the terms of the Creative Commons Attribution Noncommercial License which permits any noncommercial use, distribution, and reproduction in any medium, provided the original author(s) and source are credited.

Appendix 1: model derivation

The measured behavior in the main article can be described by a linear motion control model based on a linearized Hill-type muscle model (Van der Helm et al. 2002), of which the linearization is described in Appendix 2.

To assess the influence of the model parameters on the frequency domain and time domain characteristics of the admittance, a graphical user interface was developed previously (Schouten et al. 2008). For clarification, in this study, the equations from which the model is derived are presented below. Note that the model is represented in the frequency domain where $s$ denotes the Laplace operator (see Fig. 2).

The torque that accelerates the total inertia is the sum of the gas pedal torque $T_c(s)$ and the opposing torque $T_{\text{muscle}}(s)$ generated by the muscles, according to:

$$T_{\text{muscle}}(s) = H_{\text{act}}(s)A(s) - H_{\text{ve}}(s)[\theta_{\text{ankle}}(s) - \theta_{\text{tendon}}(s)]$$

where $H_{\text{act}}$ describes the activation dynamics (the process of active muscle force buildup following a neural activation signal $A$). $H_{\text{ve}}$ describes the intrinsic muscle visco-elasticity according to:

$$H_{\text{ve}}(s) = k_v + b_v s$$

with $k_v$ the total intrinsic stiffness and $b_v$ the total intrinsic damping of (already contracted) muscles. Visco-elasticity can be actively increased by muscle co-contraction, so the two gains are allowed to vary between the conditions.

The tendon angle depends on the tendon stiffness $k_{\text{tendon}}$ and the muscle torque $T_{\text{muscle}}$ according to:

$$\theta_{\text{tendon}}(s) = H_{\text{tendon}}(s)T_{\text{muscle}}(s) = k_{\text{tendon}} T_{\text{muscle}}(s)$$

$H_{\text{act}}$ is approximated by a second-order system:

$$H_{\text{act}}(s) = \frac{2f_a k_v}{s^2 + 2d_a k_v + f_a^2}$$

with eigen-frequency $f_a$ and relative damping $d_a$ and with $\omega_a = 2\pi f_a$.

The neural activation signal $A(s)$ is modeled as:

$$A(s) = U(s) - H_{\text{gto}}(s)T_{\text{muscle}}(s) - H_{\text{ms}}(s)[\theta_{\text{ankle}}(s) - \theta_{\text{tendon}}(s)]$$

where $U(s)$ represents deviations from the mean supraspinal command. Note that $U(s)$ does not contain the mean descending command that results in co-contraction (which is modeled by intrinsic visco-elasticity $k_a$ and $b_a$). Effectively, $U(s)$ can be assumed to be zero because of the unpredictable nature of the perturbations. $A(s)$ is therefore only the result of feedback from muscle spindles ($H_{\text{ms}}$) and Golgi tendon organs ($H_{\text{gto}}$).

$H_{\text{ms}}$ models the activity of muscle spindles as a lumped position and velocity feedback with a time delay, according to:

$$H_{\text{ms}}(s) = (k_p + k_v s) e^{-\tau_{\text{ms}}}$$

The parameters $k_p$ and $k_v$ represent the gains of, respectively, the monosynaptic stretch length and stretch velocity feedback. A single short-latency time delay $\tau_{\text{ms}}$ is used to model the elapsed time due to the neural signal traveling from the spindles to the spinal cord and back to the muscle (De Vlugt et al. 2002). Muscle spindles are often reported to have an excitatory effect on the alpha
A positive gain $k_f$ denotes an inhibitory effect, increasing the admittance. Since literature has indicated that GTOs can deliver excitatory feedback as well, a negative gain is also allowed in this model.

The pedal rotations not only cause joint rotations, but also displacement of skin and underlying tissues. This effect is modeled as contact dynamics:

$$T_c(s) = H_c(s)\left[\theta_{\text{pedal}}(s) - \theta_{\text{ankle}}(s)\right]$$

with $H_c(s)$ modeled as:

$$H_c(s) = b_c s + k_c$$

where $b_c$ and $k_c$ respectively denote the viscous and elastic part of the contact dynamics. Since in the experiment the foot is strapped to the pedal, these values are expected to be constant over all experimental conditions.

Equations 6–10 are combined into one equation, as can be found in the main text as Equation 3.

**Appendix 2: model linearization**

The model is based on previous research on the linearization of a multiplicative Hill-type model (Stroeve 1998), in which the muscle force is described by the multiplication of the activation signal $a$, the force–length relationship $f_l(L_m)$, the force–velocity relationship $f_v(V_m)$ and the maximal force $F_{max}$:

$$F_s = a \cdot f_l(L_m) \cdot f_v(V_m) \cdot F_{max} \quad (11)$$

The first-order linearized activation term is a partial derivative of $F_s$ to $a$, and is approximated by:

$$\frac{\partial F_s}{\partial a} = f_l(L_m) \cdot f_v(V_m) \cdot F_{max} \quad (12)$$

The variations of muscle force $\delta F_s$ caused by changing muscle activation $\delta a$ around an operating point can now be described as:

$$\delta F_s = \left[f_l(L_m) \cdot f_v(V_m) \cdot F_{max}\right] \cdot \delta a + \left[a \cdot f_v(V_m) \cdot F_{max} \cdot \frac{\partial f_l(L_m)}{\partial L_m}\right] \cdot \delta L_m + \left[a \cdot f_l(L_m) \cdot F_{max} \cdot \frac{\partial f_v(V_m)}{\partial V_m}\right] \cdot \delta V_m, \quad (13)$$

which can be rewritten in the nomenclature of the current study as:

$$T_{\text{muscle}} = \left[f_l(L_m) \cdot f_v(V_m) \cdot F_{max}\right] \cdot A + [k_a] \theta_{\text{muscle}} + [b_a] \theta_{\text{muscle}} \quad (14)$$

with $T_{\text{muscle}}$ now denoting deviations in muscle force ($\delta F_s$), $A$ deviations in muscle activation ($\delta a$) due to reflexive activity, $\theta_{\text{muscle}}$ deviations in muscle stretch ($\delta L_m$), and $\theta_{\text{muscle}}$ the deviations in muscle stretch velocity ($\delta V_m$). Note that the intrinsic stiffness and damping due to already contracted muscles is described by $k_a$ and $b_a$, respectively. The term $[f_l(L_m) \cdot f_v(V_m) \cdot F_{max}]$ is for this model implicitly incorporated in the loop gain of each of the reflexive feedback loops ($k_m, k_v, k_f$).

**Appendix 3**

See Tables 2, 3, and 4.

**Table 2 Lower and upper parameter boundaries**

| Parameters                          | Lower boundary | Upper boundary |
|-------------------------------------|----------------|----------------|
| Ankle inertia, $I_a$ (kg m$^2$)     | 0.01           | 0.6            |
| Tendon stiffness, $k_{\text{tendon}}$ (Nm/rad) | 100           | 10,000         |
| Muscle spindle time delay, $\tau_{\text{ms}}$ (s) | 0.015         | 0.05           |
| Golgi tendon organ time delay, $\tau_{\text{gto}}$ (s) | 0.015         | 0.04           |
| Relative damping activation dynamics, $d_a$ (s) | 0.5           | 1.5            |
| Eigen-frequency activation dynamics, $f_a$ (Hz) | 2             | 10             |
| Contact dynamics viscosity, $b_c$ (Nms/rad) | 1             | 100            |
| Contact dynamics stiffness, $k_c$ (Nm/rad) | 100           | 2,000          |
| Muscle viscosity, $b_a$ (Nms/rad) | 0.01           | 20             |
| Muscle stiffness, $k_a$ (Nm/rad) | 100            | 1,500          |
| Muscle stretch velocity feedback gain, $k_{\text{v}}$ (Nms/rad) | -40           | 40             |
| Muscle stretch length feedback gain, $k_{\text{sl}}$ (Nm/rad) | -1,000        | 1,000          |
| Muscle force feedback gain, $k_f$ (-) | -10           | 10             |

**Table 3 Condition-independent parameters**

| Parameters                          | Mean ± SD (n = 10) |
|-------------------------------------|--------------------|
| Ankle inertia, $I_a$ (kg m$^2$)     | 0.078 ± 0.04       |
| Tendon stiffness, $k_{\text{tendon}}$ (Nm/rad) | 5.074 ± 2.369   |
| Muscle spindle time delay, $\tau_{\text{ms}}$ (s) | 0.038 ± 0.009   |
| Golgi tendon organ time delay, $\tau_{\text{gto}}$ (s) | 0.025 ± 0.009   |
| Relative damping activation dynamics, $d_a$ (-) | 0.99 ± 0.32   |
| Eigen-frequency activation dynamics, $f_a$ (Hz) | 2.68 ± 0.78   |
| Contact dynamics viscosity, $b_c$ (Nms/rad) | 24.8 ± 13.6   |
| Contact dynamics stiffness, $k_c$ (Nm/rad) | 826 ± 410   |
Table 4 Validation of the ARMAX approximations and parameter fits

| Task       | Bandwidth (Hz) | ARMAX                      | Parameter fit                      |
|------------|----------------|----------------------------|------------------------------------|
|            |                | VAF $T_c$ (SD) (%)         | VAF $\theta_{\text{pedal}}$ (SD) (%) | VAF $T_c$ (SD) (%)         | VAF $\theta_{\text{pedal}}$ (SD) (%) |
| Force      | 0.7            | 84.3 (±5.6)                | 65.2 (±10.9)                       | 39.0 (±9.6)                | 85.1 (±7.9)                       |
|            | 1.2            | 88.6 (±4.8)                | 50.1 (±14.7)                       | 69.7 (±2.9)                | 81.5 (±13.1)                      |
|            | 2.0            | 91.4 (±5.3)                | 35.6 (±12.6)                       | 85.7 (±2.0)                | 73.6 (±14.6)                      |
| Relax      | 0.7            | 99.8 (±0.1)                | 99.1 (±0.3)                        | 93.3 (±1.0)                | 94.5 (±1.9)                       |
|            | 1.2            | 99.7 (±0.1)                | 98.8 (±0.2)                        | 94.7 (±0.9)                | 96.4 (±1.5)                       |
|            | 2.0            | 99.7 (±0.1)                | 98.4 (±0.3)                        | 94.9 (±1.1)                | 95.4 (±1.7)                       |
| Position   | 0.7            | 99.8 (±0.1)                | 78.4 (±7.9)                        | 94.9 (±0.4)                | 78.8 (±22.3)                      |
|            | 1.2            | 99.8 (±0.1)                | 81.9 (±6.2)                        | 97.0 (±0.2)                | 84.1 (±15.7)                      |
|            | 2.0            | 99.8 (±0.2)                | 88.5 (±5.6)                        | 97.8 (±0.2)                | 91.2 (±8.4)                       |

Mean and standard deviation of the variance accounted for (VAF) of the ARMAX-approximated signals to the experimental data (left) and model-predicted to the ARMAX-approximated signals (right); calculated for both gas pedal torque ($T_c$) and gas pedal depression angle ($\theta_{\text{pedal}}$) for all subjects.

References

Abbink DA (2006) Neuromuscular analysis of haptic gas pedal feedback during car following. Dissertation, Delft University of Technology, Delft University Press

Abbink DA (2007) Task instruction: the largest influence on human operator control dynamics. In: Proceedings of world haptics, Tsukuba, Japan, 22–24 March 2007, pp 206–211

Allum JH, Mauritz KH (1984) Compensation for intrinsic muscle stiffness by short-latency reflexes in human triceps surae muscles. J Neurophys 52:797–818

Allum JH, Mauritz KH, Vögele H (1982) The mechanical effectiveness of the short-latency reflexes in the human triceps surae muscles revealed by ischaemia and vibration. Exp Brain Res 48:153–156

Cathers I, O’Dwyer N, Neilson P (1999) Dependence of stretch reflexes on amplitude and bandwidth of stretch in human wrist muscle. Exp Brain Res 129:278–287

Colebatch JG, McCloskey DI (1987) Maintenance of constant arm position or force: reflex and volitional components in man. J Physiol 386:247–261

Crago PE, Houk JC, Hasan Z (1976) Regulatory actions of human stretch reflex. J Neurophysiol 39(5):925–935

De Vlugt E, Van der Helm FCT, Schouten AC (2002) Adaptation of reflexive feedback during arm posture to different environments. Biol Cybern 87:10–26

De Vlugt E, Schouten AC, Van der Helm FCT (2006) Quantification of intrinsic and reflexive properties during multijoint arm posture. J Neurosci Methods 155:328–349

Doemges F, Rack PMH (1992a) Changes in the stretch reflex of the human first dorsal interosseus muscle during different tasks. J Physiol (Lond) 447:563–573

Doemges F, Rack PMH (1992b) Task-dependent changes in the response of human wrist joints to mechanical disturbance. J Physiol (Lond) 447:575–585

Donelan JM, Pearson KG (2004) Contribution of force feedback to ankle extensor activity in decerebrate walking cats. J Neurophysiol 92:2093–2104

Doornik J, Sinkjær T (2008) Robotic platform for human gait analysis. IEEE Trans Biomed Eng 54(9):1696–1702

Faist M, Hoefer C, Hodapp M, Dietz V, Berger W, Duyens J (2006) In humans Ib facilitation depends on locomotion while suppression of Ib inhibition requires loading. Brain Res 1076:87–92

Geyer H, Seyfarth A, Blickhan R (2003) Positive force feedback in bouncing gaits? Proc Bioc Sci 270(1529):2173–2183

Gielen CCAM, Ramaekers L, Van Zuylen EJ (1988) Long-latency stretch reflexes as co-ordinated functional responses in man. J Physiol 407:273–292

Grey MJ, Ladouceur M, Andersen JB, Nielsen JB, Sinkjær T (2001) Group II muscle afferents probably contribute to the medium latency soleus stretch reflex during walking in humans. J Physiol 534(3):925–933

Grey MJ, Nielsen JB, Mazzaro N, Sinkjær T (2007) Positive force feedback in human walking. J Physiol 581:99–105

Hammond PH (1956) The influence of prior instruction to the subject on an apparently involuntary neuro-muscular response. J Physiol 132(1):17–18

Hore J, McCloskey DJ, Taylor JL (1990) Task-dependent changes in gain of the reflex response to imperceptible perturbations of joint position in man. J Physiol 429:309–321

Jaeger R, Gottlieb G, Agarwal G (1982) Myoelectric responses at flexors and extensors of the human wrist to step torque perturbations. J Neurophysiol 48:388–402

Jankowska E, McCrea DA (1983) Shared reflex pathways from Ib tendon organ afferents and Ia muscle spindle afferents in the cat. J Physiol 338:99–111

Jenkins GM, Watts DG (1968) Spectral analysis and its applications. Holden-Day, San Francisco

Kearney RE, Stein RB, Paranmeswaran L (1997) Identification of intrinsic and reflex contributions to human ankle stiffness dynamics. IEEE Trans Biomed Eng 44(6):493–504

Kiemele T, Oie K, Jeka JJ (2006) The slow dynamics of postural sway are in the feedback loop. J Neurophysiol 95:1410–1418

Kirsch R, Kearney R (1997) Identification of time-varying stiffness dynamics of the human ankle joint during an imposed movement. Exp Brain Res 114:71–85

Kurtzer IL, Pruszynski JA, Scott SH (2008) Long-latency reflexes of the human arm reflect an internal model of limb dynamics. Curr Biol 18(6):449–453

Lasschuit L, Lam TM, Mulder M, van Paassen MM, Abbink DA (2008) Measuring and modeling neuromuscular system dynamics for haptic interface design. In: Proceedings of the AAA modeling and simulation technologies conference and exhibit 18–21 August 2008, Honolulu, Hawaii

McRuer DT, Jex HR (1967) A review of quasi-linear pilot models. IEEE Trans Human Factors Electron 8(3):231–249
Mugge W, Abbink DA, Van der Helm FCT (2007) Reduced power method: how to evoke low-bandwidth behavior while estimating the full-bandwidth dynamics. In: Proceedings of the IEEE 10th international conference on rehabilitation robotics, 13–15 June 2007, pp 575–581

Mugge W, Schuurmans J, Schouten AC, Van der Helm FCT (2009) Sensory weighting of force and position feedback in human motor control tasks. J Neurophysiol 39(1):119–142

Nichols TR, Houk JC (1976) Improvement in linearity and regulation of stiffness that results from actions of stretch reflex. J Neurophysiol 39(1):119–142

Pearson KG, Collins DF (1993) Reversal of the influence of group Ia afferents from plantaris on activity in medial gastrocnemius muscle during locomotor activity. J Neurophysiol 70:1009–1017

Perreault EI, Chen K, Lewis GN (2006) Regulation of multijoint stretch reflexes during interactions with stiff and compliant environments. In: 28th Annual international conference of the IEEE, Aug 30–Sept 3 2006. Engineering in Medicine and Biology Society, pp 300–302

Petersen N, Christensen LOD, Morita H, Sinkjær T, Nielsen J (1998) Evidence that a transcortical pathway contributes to stretch reflexes in the tibialis anterior muscle in man. J Physiol 512(1):267–276

Pintelon R, Schoukens J (2001) System Identification: a frequency domain approach. IEEE Press, New York

Pintelon R, Guillaume P, Rolain Y, Schoukens J, van Hamme H (1994) Parametric identification of transfer functions in the frequency domain: a survey. IEEE Trans Autom Contr 39:2243–2260

Prochazka A, Hulliger M, Zanger P, Appenteng K (1985) ‘Fusimotor set’: new evidence for alpha-independent control of gamma-motoneurones during movement in the awake cat. Brain Res 339(1):136–140

Prochazka A, Gillard D, Bennett DJ (1997) Implications of positive feedback in the control of movement. American Physiological Society 0022-3077/97

Pruszynski JA, Kurtzer I, Scott SH (2008) Rapid motor responses are appropriately tuned to the metrics of a visuospatial task. J Neurophysiol 100:224–238

Rothwell JC, Traub MM, Marsden CD (1980) Influence of voluntary intent on the human long-latency stretch reflex. Nature 286(5772):496–498

Rudomin P, Schmidt RF (1999) Presynaptic inhibition in the vertebrate spinal cord revisited. Exp Brain Res 129:1–37

Schouten AC, Mugge W, Van der Helm FCT (2008) NMClab, a model to assess the contribution of muscle visco-elasticity and afferent feedback to joint dynamics. J Biomech 41(8):1659–1667

Shadmehr R, Krakauer JW (2008) A computational neuroanatomy for motor control. Exp Brain Res 185:359–381

Sinkjær T, Andersen JB, Larsen B (1996) Soleus stretch reflex modulation during gait in humans. J Neurophysiol 76(2):1112–1120

Soechting JF, Lacquaniti F (1988) Quantitative evaluation of the electromyographic responses to multidirectional load perturbations of the human arm. J Neurophysiol 59(4):1296–1313

Stein RB, Kearney RE (1995) Nonlinear behavior of muscle reflexes at the human ankle joint. J Neurophysiol 73(1):65–72

Stienen AH, Schouten AC, Schuurmans J, van der Helm FCT (2007) Analysis of reflex modulation with a biologically realistic neural network. J Comput Neurosci 23(3):333–348

Stroeve S (1998) Neuromuscular control model of the arm including feedback and feedforward components. Acta Psychol 100(1–2):117–131

Van den Hof PMJ, Schrama RJF, Bosgra OH, De Callafon RA (1995) Identification of normalized coprime plant factors from closed-loop experimental data. Eur J Control 1:62–74

Van der Helm FCT, Schouten AC, De Vlugt E, Brouwn GG (2002) Identification of intrinsic and reflexive components of human arm dynamics during postural control. J Neurosci 119:1–14

Van der Kooij H, van Asseldonk E, van der Helm FCT (2005) Comparison of different methods to identify and quantify balance control. J Neurosci Methods 145(1–2):175–203