Disturbances in human arm movement trajectory due to mild cerebellar dysfunction

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Abstract

The temporal structure of arm movements was studied in nine cerebellar patients with mild impairment of the upper limbs and in six age-matched control subjects. The experimental paradigm consisted of visually guided, step tracking movements about the elbow. Movements ranged from 10° to 70° in amplitude and were made under different instructions (fast, fast/accurate, accurate). As in normal subjects, cerebellar patients were able to scale peak velocity with movement amplitude. This relationship was highly linear under all instruction conditions. Similar relationships existed between movement duration and amplitude. In contrast to normal subjects who produced movements with nearly symmetric velocity profiles, movements made by cerebellar patients were characterised by short acceleration and long deceleration durations. The degree of asymmetry was directly related to movement duration but was unaffected by movement peak velocity. Acceleration durations did not increase beyond 300 ms even in movements lasting up to 1s. These findings demonstrate that, despite little or no obvious impairment of the limb during routine examination, the temporal structure of voluntary movements in cerebellar patients is clearly disturbed. This supports the view that the production of an optimal movement trajectory is under cerebellar influence.

In his elegant treatise concerning motor deficits following gunshot wounds to the cerebellum, Holmes1 described a specific set of movement abnormalities which still provides the framework for clinical diagnosis of cerebellar dysfunction. In addition to hypotonia, tremor and ataxia, cerebellar patients often exhibit abnormalities in the rate, range, accuracy and force of goal-directed voluntary movements as well as irregularities in the performance of alternating movements. Since Holmes' pioneering work, cerebellar involvement in movement planning and execution has been the focus of considerable study. However, the precise role that the cerebellum plays in the control of voluntary movement is still unclear.

The widely accepted notion of movements being either “fast” or “slow” led Kornhuber2 to suggest separate, central mechanisms responsible for each movement type. On the basis of disrupted eye movements in patients with cerebellar atrophy, Kornhuber proposed that the cerebellum was necessary in “the translation of the spatial concept of the movement . . . into time”. While the cerebellum was considered to represent a clock mechanism involved in the timing of muscle activity producing a rapid movement, the basal ganglia were thought to act as a ramp generator for slow movements.

That the cerebellum may be preferentially involved in the generation of fast movements has been supported by both primate and clinical studies. It has been shown, for example, that cooling of the cerebellar nuclei in monkeys results in movements which exhibit more than one velocity peak during the dynamic phase, a phenomenon referred to by Brooks as “discontinuous”.4 Such movements are typically characterised by delayed onset times, hypermetria and the presence of terminal oscillations.5-7 Similar disturbances in movement range and speed are seen in cerebellar patients performing fast tracking movements about the thumb5 and elbow6 and during rapid isometric movements of the thumb and index finger.6

It has been shown, however, that cerebellar lesions also give rise to specific deficits during the production of slow, continuous tracking movements. Errors in movement rate and amplitude results in a marked reduction in tracking accuracy and movements resemble a series of intermittent responses performed at inappropriately high velocities.11-13

Part of the difficulty in determining the precise role for the cerebellum during voluntary movement is the generalised assumption that movements can be classified as either “fast” or “slow”. While such a classification is appealing in its simplicity, it is often not possible to determine the boundary between movement types on the basis of a single movement parameter. Kinematic analyses have shown that, despite large differences in speed, human arm movements share common organisational principles. Thus, for example, maximum velocity increases linearly with movement amplitude.16-18

More recently, the dynamic or temporal structure of movements has been the focus of increased attention since time-symmetric velocity profiles appear to be characteristic of many movement types which differ in amplitude, direction and speed and made...
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The observation that the time course during the dynamic phase of movements is preserved under different conditions has added further support to the theory that movement trajectory is the controlled variable underlying movement planning and execution. In addition to commonality of specific kinematic features, many movements share similar electromyographic activation patterns. A triphasic (agonist-antagonist-agonist) burst pattern, once thought to be characteristic of only the most rapid movements, has been described for movements of a wide range of speeds and amplitudes. Furthermore, it has been recently shown that modulation of movement-related phasic EMG activity is directly related to the dynamic characteristics of the intended movement.

Since it has been suggested that the cerebellum plays a crucial role in movement timing, it would seem reasonable to hypothesise that generation of a common, temporal profile across different movement conditions may be under cerebellar control. Experiments were therefore designed to examine the dynamic features of single-joint, step tracking movements in patients exhibiting cerebellar symptoms. To minimise problems in determining, for example, movement end point, only patients with mild impairment of the upper extremities were selected. Unlike previous studies involving cerebellar patients, the present paradigm permitted an analysis of graded changes in movement speed. Thus, by altering amplitude and/or instruction, a broad range of velocities could be produced.

The results show that, for movements about the elbow, patients with mild cerebellar symptoms appear unable to produce time symmetric movements across a range of movement amplitudes and speeds. Thus, disturbances in the dynamic phase of voluntary movements may be present despite little or no impairment of the limb during routine clinical testing.

Methods

Subjects
Nine patients diagnosed with cerebellar disease were studied (eight male, one female, aged 37 to 63 years, mean age: 52 years). Following a complete clinical assessment, patients were selected on the basis of age, location of lesion, visual acuity and degree of upper limb impairment. Patients over the age of 65 years were excluded to avoid the possibility of age-related changes in motor performance. Patients suffering from demyelinating disease (for example, multiple sclerosis) or exhibiting major brainstem involvement were also excluded. All patients were ambulatory and presented only mild impairment of the upper limbs.

The clinical data are summarised in table 1. Four patients (WD, RB, KS, EH) suffered from chronic cerebellar disease of between two and five years duration. Using Harding’s classification, these patients were diagnosed as suffering from ideopathic late onset cerebellar ataxia with patients RB and KS of the Marie–Foix–Alajouanine type; EH, group C; and WD, probable group C. Four patients (BR, HF, KB, VR) presented acute cerebellar symptoms due to vascular injury. BR suffered from occlusion of both vertebral arteries with complex collateralisation of the right cerebellar hemisphere. HF suffered from an occlusion of the right posterior cerebellar artery. In patient KB an embolism of the right superior cerebellar artery was diagnosed and in VR, a large, ischaemia induced cyst located in the left cerebellar hemisphere. EZ had a solid metastasis (hypernephroma) located in the right cerebellar hemisphere. In cases where symptoms were bilateral, the dominant arm was tested. All patients gave informed consent for the procedures involved. Control studies were performed on six subjects with no known history of motor dysfunction (four male, two female, 30 to 60 years, mean age: 45 yrs).

Experimental paradigm

The experimental setup used in these studies was similar to that used in previous studies of step tracking movements about the elbow. Subjects were seated comfortably and grasped a manipulandum handle. The arm was abduced 90° and supported along the entire length of the forearm. The manipulandum was pivoted beneath the elbow and moved freely in the horizontal plane.

An oscilloscope placed approximately 1 metre in front of the subject was used to display target and handle position. The target appeared as two vertical lines, 8 mm apart. The target switched at a regular interval (every 5s) between two fixed positions equidistant about an elbow angle of 90 deg.

Table 1  Clinical data for cerebellar patients

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Sex</th>
<th>Length of Illness</th>
<th>Diagnosis</th>
<th>Tremor Arm</th>
<th>Head/Trunk</th>
<th>Ataxia Arm</th>
<th>Stance/Gait</th>
<th>Dyssarthis</th>
<th>Oculomotor</th>
<th>Upper Limb</th>
<th>Lower Limb</th>
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<td>+</td>
<td>+</td>
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<tr>
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<td>F</td>
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<td>&quot;</td>
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<td>2</td>
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<td>&quot;</td>
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<td>2</td>
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<tr>
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<td>47</td>
<td>M</td>
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<td>&quot;</td>
<td>2</td>
<td>3</td>
<td>2</td>
<td>3</td>
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<td>+</td>
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<td>0</td>
<td>++</td>
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</tr>
</tbody>
</table>

*see text for detailed description of diagnosis

*rating of clinical symptoms: 0 = normal, 1-5 = mild to severe disturbance.
(180° deg equivalent to full extension). The targets were not mechanically detectable and were not bounded by mechanical stops. Position of the manipulandum handle was displayed as a thin vertical line.

Subjects were instructed to place the handle cursor within the target zone according to the given instruction. Instructions were: 1) “move accurately” with an emphasis on placing the handle cursor exactly in the middle of the target zone, 2) “move fast but accurately” with an emphasis on avoiding any overshoot of the target zone and 3) “move as fast as possible” where speed was stressed.

During each experimental session, the subject was asked to make a series of step-tracking movements about the elbow. Each trial consisted of 30 flexion-extension movements. Target amplitude varied from 10° to 70° but was kept constant during any given trial. Accurate movements were performed first, followed by fast/accurate movements and finally movements made as fast as possible. Within each instruction block (for example, “accurate”), target amplitude varied randomly from trial to trial. The task was not a reaction time task in that the subject was not required to minimize movement onset time relative to movement of the target. Each subject was allowed 2–3 minutes of practice before data sampling. Recording sessions never exceeded one hour.

Data recording and analysis
Angular position and velocity of the manipulandum handle were recorded respectively from a potentiometer and tachometer mounted beneath the pivot point of the handle. Data were digitised on-line with an effective sampling rate of 250 Hz. Each movement was analysed individually using automatic computer programmes. Timing points were determined from the differentiated velocity signal using an acceleration threshold of 120°/s². End of movement was arbitrarily defined as that point where the subject was within 3° of target centre. Using this definition, 5–10% of movements were considered not of the correct amplitude and were discarded. Computer selected timing points were later confirmed by visual inspection of plotted records. Statistical evaluation of group differences was determined using a two-tailed Student’s t test where appropriate.

Results
Interaction between peak velocity, movement duration and amplitude
All patients were able to perform reproducible, step-tracking movements after a short period of practice. Typical records of arm velocity associated with 70° movements made as fast as possible are shown in fig 1 for a normal (A) and two cerebellar patients (B, C). Movements made by normal subjects were highly stereotyped with bell-shaped, symmetrical velocity profiles. The records in B (cerebellar 1, patient WD) were representative of movements made by most of the patients under investigation: a smooth acceleratory phase followed by a slower, more prolonged period of deceleration. In C (cerebellar 2, patient KB) show velocity records obtained from the most severely affected patient. In this patient, individual movements were often characterised by either a secondary, low amplitude velocity peak or a period of relatively constant velocity. All patients were able to smoothly terminate each movement within the target zone and the degree of terminal overshoot in movements made as fast as possible was no greater in the patient than in the normal group. Flexion and extension movements were qualitatively similar and thus only data associated with flexion movements are presented.

Patients were able to modulate movement velocity in response to changes in either target amplitude or instruction. This is shown in fig 2. Peak velocity increased with increasing movement amplitude regardless of instruction. For a given movement amplitude, peak velocity depended on the patient’s strategy or the instruction given (“accurate” versus “fast”). Movement duration also changed with both amplitude and instruction. Despite large
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Figure 3 Amplitude dependent scaling of peak velocity. Individual linear regression lines are plotted for each patient under each instruction condition. Pooled data from normal subjects are represented by the dashed regression lines. Slope, intercept and correlation coefficient (r) values for each patient are given in table 2.

Figure 4 Changes in movement duration with amplitude. Individual linear regression lines are plotted for each patient under each instruction condition. Pooled data from normal subjects are represented by the dashed regression lines. Slope, intercept and correlation coefficient (r) values for each patient are given in table 2.

differences in amplitude, duration and peak velocity across the range of movements examined, most patients produced movements with smooth, unimodal velocity profiles. As with normal subjects, terminal oscillations were only seen during movements made as fast as possible.

In all patients peak velocity increased linearly with movement amplitude. Best fit regression lines for each patient are shown in fig 3. Values of regression parameters given in table 2 indicate that the relationship was highly linear across subjects. Linear scaling of peak velocity with amplitude was preserved regardless of instruction. However, the gain (slope) of the relationship was instruction dependent, being higher for fast than for accurate movements. For moderate speed movements (fast/accurate), maximum velocity did not differ greatly from mean normal values. Movements made as rapidly as possible were, in general, slower in the cerebellar group, particularly at larger movement amplitudes. Accurate movements greater than 20° amplitude were performed at slightly higher speeds than that observed in the normal group.

As stated earlier, movement duration increased with amplitude. While this relationship was linear for each instruction in all patients with two exceptions (EZ—fast/accurate; KS accurate), slope and intercept values were highly variable across patients (fig 4, table 2). The time required for movement completion generally reflected the maximum speed attained during the movement (fig 3). Thus, compared to normal values, movement durations were slightly longer for fast movements and, on average, slightly shorter for accurate movements. For moderate speed (fast/accurate) movements, however, movement duration tended to be longer in the cerebellar group despite peak velocities being within normal limits (fig 2).

To summarise, a highly linear relationship

### Table 2 Summary of linear regression analysis

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<thead>
<tr>
<th>Case</th>
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<th>slope</th>
<th>r</th>
<th>Peak Velocity—Amplitude</th>
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<td>r</td>
<td>Accurate intercept</td>
<td>slope</td>
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<table>
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<th>slope</th>
<th>r</th>
<th>Movement Duration—Amplitude</th>
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<td>r</td>
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<td>slope</td>
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</table>

*Data unavailable
*mean regression values for normal subjects
*movements too slow for on-line sampling time
between peak velocity and movement amplitude was preserved in patients with mild cerebellar symptoms with the slope of the relationship instruction dependent. A similar relationship between movement duration and amplitude occurred but was more variable across subjects.

**Acceleration duration/deceleration duration relationship**

Despite relatively normal scaling of peak velocity with amplitude, movements made by cerebellar patients were asymmetric in their time course (fig 1 B, C; fig 2 accurate and fast/accurate). This is illustrated in fig 5 where data have been plotted for all patients, amplitudes and instructions. Each data point is the mean value for an individual patient performing movements of a particular amplitude and instruction. For relatively short duration (<500 ms) movements, acceleration and deceleration durations were approximately equal and close to the mean normal curve (dashed line). However, as total movement duration increased in the patients, deceleration duration became disproportionately longer compared to acceleration duration. In normal subjects, acceleration duration was graded up to approximately 500 ms while, in contrast, none of the patients exhibited movements with acceleration durations greater than 300 ms. As a result, the relationship between acceleration duration and deceleration duration was well described by a logarithmic function which gave a slightly better goodness of fit ($r = 0.84$) than did a linear function ($r = 0.78$).

The degree of movement asymmetry, determined by the ratio of acceleration duration to deceleration duration (symmetry ratio = SR), appeared to be both amplitude and instruction dependent. Figure 6 shows that medium to large amplitude movements (30–70°) were consistently characterised by skewed velocity profiles and that slower (accurate) movements were more asymmetric than movements made as fast as possible. In contrast, the time course of 10° movements was not significantly different from normal values regardless of instruction. This was also true for all movements made as fast as possible independent of movement amplitude. Movements made by normal subjects were nearly symmetric in their time course at all amplitudes and under all instructions (SR range: 0.8–0.9; mean SR = 0.85).

To determine if the asymmetry in movement profiles was best correlated with total movement duration as suggested in fig 5 and 6 or with peak velocity (since, for example, longer duration movements were associated with lower speeds), movement duration and peak velocity were plotted separately as a function of SR. This is shown in fig 7. As can be seen in A, the degree of skewness was clearly dependent on movement duration. Only at movement durations of less than 500 ms did SR values approach the normal range (fig 6). As movement duration increased, SR decreased to values in which deceleration duration was twice as long as acceleration duration (SR < 0.5). The relationship between movement duration and SR was highly correlated and best described by an exponential function ($r = 0.82$).

Surprisingly, the degree of movement asymmetry was not affected by the maximum speed of the movement (fig 7B). Movements with low SR values (<0.5) and thus highly asymmetric were associated with a broad range of maximum velocities as was also the case for relatively symmetric movements (SR ≥ 0.8). Therefore, in movements in which both speed and duration increase, the resulting velocity skewness appears to be a result of the increase in total movement duration rather than the associated increase in peak velocity.

**Changes in movement variability**

In addition to being temporally asymmetric, movements made by cerebellar patients were, in general, more variable than normal subjects (fig 8). Although mean values were consistently greater in the cerebellar group regardless of
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Discussion

During the planning of either simple, single-joint or more complex, multi-joint movements, the central nervous system is concerned with two problems: 1) where to move and 2) how to move. In the first case, changes in the relative levels of tonic muscle activity acting about a given joint are sufficient to move the limb to a new position as long as the demand for speed does not exceed limitations imposed by viscoelastic forces.2,26 Movements produced in such a manner are generally too slow to meet task requirements, however, and thus how one moves becomes a central factor in determining the appropriate motor output. The results of this study show that, while patients with mild cerebellar dysfunction of the upper extremities can move successfully to externally determined target positions, the time course of the movement is disturbed. Moreover, these changes in movement trajectory are not simply due to alterations in or between specific movement parameters such as maximum speed or amplitude but appear to result from inadequate scaling of acceleration duration.

It is well known that, in normal subjects, the relationship between maximum speed and amplitude is highly linear.14-16 The slope of the relation is dependent on instructional set, being greater for movements made as fast as possible than for movements made accurately.15 The results presented here show that cerebellar patients were also capable of scaling peak velocity over a seven-fold increase in amplitude and that linearity between these two parameters was independent of speed and/or accuracy demands. Amplitude dependent increases in movement speed have also been described for slow, continuous tracking movements during cooling of the dentate

![Graph showing effect of movement duration and peak velocity on symmetry ratio.](image)

**Figure 7** Effect of movement duration and peak velocity on symmetry ratio. Each graph shows data obtained from all patients. Each averaged (n = 12) data point corresponds to movements made by an individual patient at a given amplitude and instruction. The relationship between movement duration and symmetry ratio was best described by an exponential function (r = 0.82). No significant effect of peak velocity on symmetry ratio could be determined.

Instruction or amplitude, significant differences occurred primarily in the fast/accurate condition. The degree of movement variability thus appeared to be task specific in that the demand for both speed and accuracy resulted in more variability than when only speed or accuracy was emphasised. No clear effect of movement amplitude on variability could be discerned. Variability in peak velocity, however, was consistently greater in small movements (10°), particularly for movements made as fast as possible and those made fast and accurately.

![Diagrams showing variability of peak velocity, movement duration and symmetry ratio.](image)

**Figure 8** Variability of peak velocity, movement duration and symmetry ratio. Coefficient of variation values for normals (open bars) and cerebellar patients (solid bars) are plotted for each kinematic variable as a function of amplitude and instruction. Asterisks indicate level of significant difference between groups as determined by a two-tailed Student's t test analysis (* = p < 0.05; ** = p < 0.01; *** = p < 0.005).
nucleus in primates and for rapid wrist movements in Parkinson patients. Taken together, these findings suggest that the relationship between peak velocity and amplitude is relatively immune to disruption of either the cerebellum or basal ganglia.

Not surprisingly, movements made as fast as possible were slower in the cerebellar group compared to normals. As the demand for accuracy increased, differences in peak velocity were minimal. It must be noted, however, that the ± 3° error allowed for final position may not have been stringent enough to force a significant slowing of movement velocity in the cerebellar group.

Miall et al.\(^7\) have recently suggested that the cerebellum may act to finely tune movements by limiting peak velocity. In this study, accurate movements were associated with slightly higher peak velocities compared to normal values. These movements, however, were still performed smoothly and with the required degree of accuracy and thus it is most likely that any increase in speed simply reflects differences in the subjective interpretation of the instruction. Development of more “ballistic” and less accurate movements which has been reported for continuous, pursuit tracking following dentate inactivation in primates and in cerebellar patients performing similar tracking tasks may be explained both by differences in the severity of the lesion and the type of tracking task employed. In the first case, we chose to examine patients with mild disturbances of the upper limbs. It is therefore possible that any inability to properly adjust maximum speeds during step tracking movements is not observed in the early stages of cerebellar dysfunction.

Secondly, there is growing evidence to suggest that afferent information plays a greater role in the generation of slow, pursuit tracking movements than, for example, more rapid, step tracking movements where only start and end positions are determined. Schieber and Thach have hypothesised that the cerebellum may act to regulate muscle spindle sensitivity via independent control of the fusimotor system. Thus, by presetting spindle sensitivity, the cerebellum could optimise afferent feedback which would be of particular value during slow tracking requiring constant monitoring of performance errors.

Although the relationships between peak velocity, movement duration and amplitude were preserved in cerebellar patients, the moment to moment time course was found to be disturbed. Specifically, this alteration in movement dynamics took the form of skewed velocity profiles. In normal subjects, movements exhibit temporally symmetric velocity profiles where the time spent in accelerating and decelerating the limb are approximately equal. Symmetrical profiles have been described for elbow and speech movements as well as movements of the vocal folds. More complex movements such as those involved in reaching are also symmetrical when made in either the horizontal or vertical planes and under different load conditions.

Together, these observations have lent support to the hypothesis that movement trajectory is centrally determined and may reflect a basic organising principle underlying movement generation. Mathematical modelling has shown that time symmetric profiles can result from minimising the rate of change of acceleration to produce movements in the most energy efficient manner. Thus, movements of different amplitudes, for example, might be produced by a relatively simple scaling of a base trajectory profile. In our study, movements made by cerebellar patients, regardless of pathology, were consistently characterised by short acceleration and long deceleration phases. This asymmetry in movement profile was independent of peak velocity but covaried with movement duration. Thus, large amplitude movements made slowly and accurately exhibited a marked asymmetry while rapid, small amplitude movements were, in fact, slightly more symmetric than normals.

Why cerebellar patients perform step tracking movements with temporally asymmetric profiles is not clear. One partial explanation may lie in the observation that, despite total movement durations of over 1s, the duration of the acceleratory phase never exceeded 300ms. Recent experiments in normal subjects have clearly demonstrated that the temporal structure of movements depends upon the precise timing of phasic drive to opposing muscle groups. For example, shifts in velocity profiles from short to long acceleration durations while maintaining total movement duration constant are accomplished by an increase in the duration of the initial agonist burst. Additional studies have shown that the duration of this burst is directly related to the duration of the acceleratory phase, independent of changes in mean acceleration. Thus, in our study, it is likely that the duration of the initial agonist burst was not continuously graded as movement duration increased. This would, in turn, result in movements with short acceleration durations and skewed velocity profiles.

The finding that cerebellar patients appeared unable to appropriately scale acceleration duration contrasts with other studies in which cerebellar dysfunction led to a prolongation of acceleration duration. Flament and Hore found, for example, that inactivation of the dentate nucleus in monkeys resulted in prolonged accelerations which were associated with an increase in initial agonist burst duration as well as a delay in onset of antagonist activity. Increased initial agonist burst duration has also been described for cerebellar patients during fast elbow flexion movements but accompanying alterations in movement kinematics were not reported. Although appropriate modulation of initial agonist burst duration may have been affected in the present study, it can be inferred from the patients’ ability to increase peak velocity with amplitude that control of burst magnitude was not disturbed, or at least not to the same extent as with duration.

It cannot be assumed, a priori, that altera-
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