

Disentangling acute motor deficits and adaptive responses evoked by the loss of cerebellar output

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Using a unique cerebellar disruption approach in non-human primates, this study provides **valuable** new insight into how cerebellar inputs to the motor cortex contribute to reaching. The findings **convincingly** demonstrate that reaching movements following cerebellar disruption slow down because of both an acute deficit in producing muscle activity as well as a progressive decline in compensating for limb dynamics. This work will be of interest to neuroscientists and clinicians interested in cerebellar function and pathology.

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Abstract

Patients with cerebellar damage experience various motor impairments, but the specific sequence of primary and compensatory processes that contribute to these deficits remains unclear. To clarify this, we reversibly blocked cerebellar outflow in monkeys engaged in planar reaching tasks. This intervention led to a spatially selective reduction in hand velocity, primarily due to decreased muscle torque, especially in movements requiring high inter-joint torque coupling. When examining repeated reaches to the same target, we found that the reduced velocity resulted from both an immediate deficit and a gradually developing compensatory slowing to reduce passive inter-joint interactions. However, the slowed hand velocity did not account for the fragmented and variable movement trajectories observed during the cerebellar block. Our findings indicate that cerebellar impairment results in motor deficits due to both inadequate muscle torque and an altered motor control strategy for managing impaired limb dynamics. Additionally, impaired motor control elevates noise, which cannot be entirely mitigated through compensatory strategies.

Introduction

The ability to coordinate well-timed movements greatly relies on the cerebellum as evident from studies of individuals with cerebellar ataxia (Beppu, Suda et al. 1984). The cerebellum regulates the timing of movements by predicting the mechanical interactions between adjacent joints (Ivry and Keele 1989, Manto, Bower et al. 2011), a computation that requires an internal model of feedforward control (Shidara, Kawano et al. 1993, Shadmehr and Mussa-Ivaldi 1994, Wolpert, Ghahramani et al. 1995, Wolpert, Miall et al. 1998, Kawato 1999, Sainburg, Ghez et al. 1999, Popa, Hewitt et al. 2013). However, motor impairments exhibited by individuals with cerebellar deficits extend much beyond poor motor timing. For example, during the acute phase of cerebellar injury, there is a loss of muscle tone and weakness of voluntary movements (Holmes 1917, Goldstein 1927, Holmes 1939). Cerebellar patients also exhibit movement-related sensory deficits such as an impaired visual perception of stimulus movement and perception of the limb position sense (i.e., proprioception or state estimation) during active movements (Therrien and Bastian 2015, Weeks, Therrien et al. 2017).

Clinical studies of cerebellar patients have yielded conflicting insights into the pathophysiology caused by a loss of cerebellar function. Some researchers suggest that motor impairments stem primarily from an inability to compensate for inter-joint interactions (Bastian, Martin et al. 1996, Bastian, Zackowski et al. 2000), while others argue that insufficient muscle torque generation is the central issue (Topka, Konczak et al. 1998, Boose, Dichgans et al. 1999). These discrepancies likely arise from the variability among individuals with chronic cerebellar deficits, where primary and compensatory effects are intertwined and difficult to separate. Moreover, individuals with cerebellar lesions often have differences in lesion etiology and associated damage beyond the cerebellum itself. As a result, the specific sequence of events that leads to motor impairments following the loss of cerebellar signals remains unclear.

To investigate this issue, we trained monkeys to perform a planar reaching task with arm support provided by an exoskeleton. We then reversibly disrupted cerebellar communication with other neural structures using high-frequency stimulation (HFS) of the superior cerebellar peduncle, assessing the impact of this perturbation on subsequent movements. Although our approach primarily affects cerebellar output to the motor cortex, it also disrupts fibers carrying input signals (e.g., spinocerebellar) and pathways to various subcortical targets (e.g., cerebello-rubrospinal). Thus, our manipulation broadly interferes with cerebellar communication. Previously, we have shown that HFS can effectively and reversibly interfere with the normal outflow of cerebellar signals (Nashef, Cohen et al. 2019) since this perturbation accurately replicated the behavioral deficits identified in cerebellar patients (Bastian 1997) and the changes in motor cortical activity that appear after dentate cooling in monkeys (Meyer-Lohmann, Conrad et al. 1975). In the present study, we found that the loss of cerebellar outflow led to reduced hand velocity during reaching movements, reflecting two parallel underlying mechanisms. First, there was a reduction in muscle torques, even for movement directions where inter-joint interactions were low. Second, the reduction in hand velocity exhibited a spatial tuning and was greater in directions that generated large inter-joint interactions, thereby reflecting an inability to compensate for limb dynamics. The time course of these two mechanisms was identified by analyzing the sequences of movements made to the same target during the cerebellar block. During such movements, hand velocity was low initially, reflecting a primary muscle torque deficit, and declined further in successive trials, revealing an adaptive motor strategy to minimize inter-joint interactions. Finally, we found that the decrease in velocity during cerebellar block could not fully explain the noisy and decomposed movements observed under these conditions, which may be due to an impaired state estimate of the limb that leads to the faulty control of inter-joint interactions. Taken together, we show that cerebellar output is essential for generating sufficient muscle torques to allow for fast and stereotypical movements while overcoming the

complex inter-joint interactions inherent in such actions. The loss of this output triggers a sequence of early primary effects followed by compensatory responses aimed at mitigating these impairments. Together these processes result in significantly slower movements.

Results

Slower reaching movements in cerebellar block are due to weakness and impaired limb dynamics

We trained four adult female monkeys (*Macaca fascicularis*) to sit in a primate chair with their upper limb supported by an exoskeleton (KINARM system) and perform planar center-out movements (**Figure 1a**). The spatial location of the targets enabled us to dissociate between shoulder-dominant (lateral targets) vs. shoulder-elbow movements (diagonal and straight targets respectively). Each trial began with the monkey holding a cursor within a central target. After 500–800 ms, one of eight peripheral targets appeared (cue). Following a 450–850 ms delay, the central target disappeared ('go' signal), prompting the monkey to reach the peripheral target within 500–900 ms and briefly hold the cursor there (100–150 ms) to receive a reward. Between successive trials, monkeys repositioned the cursor back to the central target. The experiment was conducted across several sessions (i.e., days). Each session was divided into 3–4 sub-sessions. Each sub-session comprised of a block of control trials (~80) followed by a block of trials (~50) with high-frequency stimulation (130 Hz) of the superior cerebellar peduncle.

Cerebellar block significantly reduced the ability of the monkeys to perform the task as measured by the success rate across all the recorded sessions (mean success rate in control = 86.5%, CI [85.8, 87.2] vs. cerebellar block = 75.3%, CI [73.9, 76.7], $p < 0.001$; Supplementary table S1). Since the monkeys had to meet strict timing criteria to perform the movements successfully during the task, we began by examining the hand kinematics. **Figure 1b** shows profiles of hand velocity in sample trials collected during the control (blue) and cerebellar block (vermillion) conditions. This example shows that movements exhibited a significant reduction in the peak hand velocity during cerebellar block. However, when compared across the targets, we found that the reduction in hand velocity was strongly modulated by the target direction (1-way ANOVA of the change in peak hand velocity, the effect of cerebellar block: $p = 0.01$, the effect of targets: $p < 0.001$; Supplementary table S2). *Post hoc* comparisons further revealed that the effect of cerebellar block on outward reaching movements (targets 1–4) i.e., movements involving shoulder flexion was significantly higher than on inward reaching movements (targets 5–8) i.e., movements involving shoulder extension (**Figure 1c**; mean decrease of 9.7%, CI [8.2, 11.2], $p < 0.001$ for targets 1–4 compared to mean decrease of 2.5%, CI [0.9, 4.0], $p = 0.21$ for targets 5–8) even though there was no difference in the peak hand velocity between the movements to the two target groups during control ($p = 0.88$, Supplementary tables S3a and b).

Previous studies have demonstrated that Purkinje neurons of the cerebellar cortex exhibit weaker modulation during task-irrelevant movements compared to task-related movements (Pi, Fakharian et al. 2024, Hage, Fakharian et al. 2025). Therefore, we examined whether the impact of the cerebellar block was limited specifically to target-directed, reward-associated movements or if it generalized to all movements performed during the task. To this end, we compared peak hand velocities between target-directed (within-trial, reward-relevant) movements and subsequent return-to-center (out-of-trial) movements. Our analysis revealed that cerebellar block resulted in a significantly larger reduction in peak velocity for target-directed movements compared to return-to-center movements (7.5%, CI [6.5, 9.0] vs. 1.7%, CI [-1.8, 3.0]; $p < 0.001$; Supplementary Figure S1). These findings suggest that the cerebellum plays a preferential role in optimizing reward-associated, within-trial movements rather than equally influencing all movements.

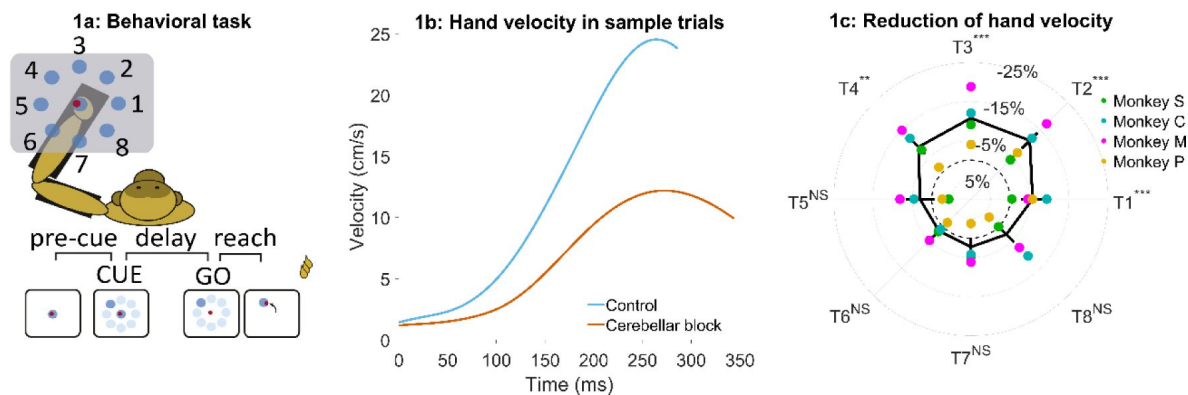


Figure 1

Effect of cerebellar block on peak hand velocity.

(a) Schematic of the monkey performing the delayed reaching task in the KINARM exoskeleton to eight peripheral targets on the horizontal plane. The locations of the eight targets are numbered. (b) Movement-onset aligned hand velocity profiles during a sample control and cerebellar block trial performed by monkey C to target 2. Entry of the control cursor into the peripheral target marked the end of the movement for the trial. Since the monkeys did not have to stop their movements fully for the trial to end (see Methods for a detailed explanation of this), the traces appear cutoff at the beginning of the deceleration/stopping phase of the movement. (c) Target-wise effect of the cerebellar block on peak hand velocity. For each session, the target-wise reduction in the median peak hand velocity during the cerebellar block trials was computed relative to that of control trials. The depicted values are the means $\pm$ 95% confidence intervals across all sessions pooled from all four monkeys. The means of individual monkeys are overlaid. The dashed circle indicates no change. Statistical significance is denoted as follows: $p \geq 0.05$ NS, $p < 0.05$ *, $p < 0.01$ **, $p < 0.001$ ***. [T1-8: Targets 1-8]

In the task used in our study, hand movements were made by controlling the muscle torque at the shoulder and elbow joints (Scott 2004 [↗](#)). However, simultaneous movements of both joints also generated coupling torques due to passive interactions between limb inertial properties and joint velocities (Bernstein 1967 [↗](#), Hollerbach and Flash 1982 [↗](#)). To identify the target-specific muscle and coupling torques, we employed an inverse dynamics model of the upper limb (Fagg, Ojakangas et al. 2009 [↗](#)) which enabled us to distinguish between the different contributions to the net torque experienced by each joint. According to this model, the net torque at each joint is the sum of both the active muscle torque and the passive coupling torque acting at that joint (see Methods section for details).

Using the inverse dynamics model, we found that the target-specific reduction in hand velocities was produced by changes in muscle commands (e.g., **Figures 2a, b** [↗](#)). Specifically, we observed that the muscle torque impulses, calculated by integrating the torque profiles over the early phase of movements when acceleration was positive, were reduced in a target-specific manner during the cerebellar block (**Figure 2c** [↗](#); Supplementary Table S4-5). This reduction was particularly pronounced at the shoulder joint. Furthermore, as in the case of hand velocities, *post hoc* comparisons revealed that the shoulder joint experienced a significantly greater reduction in muscle torque for targets 1-4 vs. 5-8 (**Figure 2d** [↗](#); mean decrease of 10.6%, CI [9.0, 11.2], $p < 0.001$ for targets 1-4 compared to mean decrease of 3.3%, CI [-1.3, 5.2] $p = 0.21$ for targets 5-8). In contrast, the slight reduction in elbow torques due to the cerebellar block did not reach significance for any of the targets (**Figure 2e** [↗](#)). Collectively, these findings indicate that during the cerebellar block, there is a reduction in shoulder muscle torque specifically for targets involved in outward-reaching movements, which leads to the target-specific reduction in hand velocity.

Several studies have shown that individuals with cerebellar deficits exhibit an inability to correctly compensate for inter-joint interactions (Bastian, Martin et al. 1996 [↗](#), Bastian, Zackowski et al. 2000 [↗](#)). Therefore, we asked whether the reduction in muscle torques and hand velocities due to the cerebellar block could be associated with the coupling torques generated during movements in various directions. **Figures 3a and b** [↗](#) show the coupling torque profiles at the shoulder and elbow joints respectively during sample trials. We observed that the coupling torque impulses varied in a target-dependent manner during both the control and cerebellar block conditions (**Figure 3c** [↗](#)). Furthermore, the reduction in peak hand velocities during cerebellar block was correlated to the net coupling torque impulse (calculated as the sum of absolute coupling torque impulses at both joints) during control, particularly for the outward reaching movements (**Figure 3d** [↗](#); targets 1-4: partial correlation coefficient to account for the variability across monkeys, $\rho = 0.32$, CI [0.23, 0.41], $p < 0.001$). This indicated that the effect of the cerebellar block on hand velocity during the outward reaching movements was strongest for targets with high coupling torques. Nevertheless, despite the significant correlation between the reduced hand velocity and coupling torques, cerebellar block reduced the hand velocity even in outward reaches where the overall coupling torque was low i.e., movements to target 1, mean reduction in hand velocity = 5.8%, CI [2.8, 8.7], $p < 0.001$ (movements to all outward targets involved both shoulder and elbow joints, but the contribution of each joint varied in a target dependent manner. Specifically, movements to targets 1 exhibited negligible net torque at the elbow, meaning the elbow remained largely stationary during these movements as shown in Supplementary Figure S2. Consequently, the shoulder experienced correspondingly low coupling torques as shown in **Figure 3c** [↗](#)). This finding indicated a reduction in the motor command which occurred in addition and beyond the effect attributable to the impaired control of coupling torques alone.

Overall, the cerebellar block produced a significant impact on the kinematics and dynamics of outward reaching movements. Specifically, the cerebellar block caused slower hand velocities due to a systematic, target- and joint-specific reduction in muscle torques for these movements, which strongly correlated with the coupling torques. Additionally, a reduction in muscle torques was also

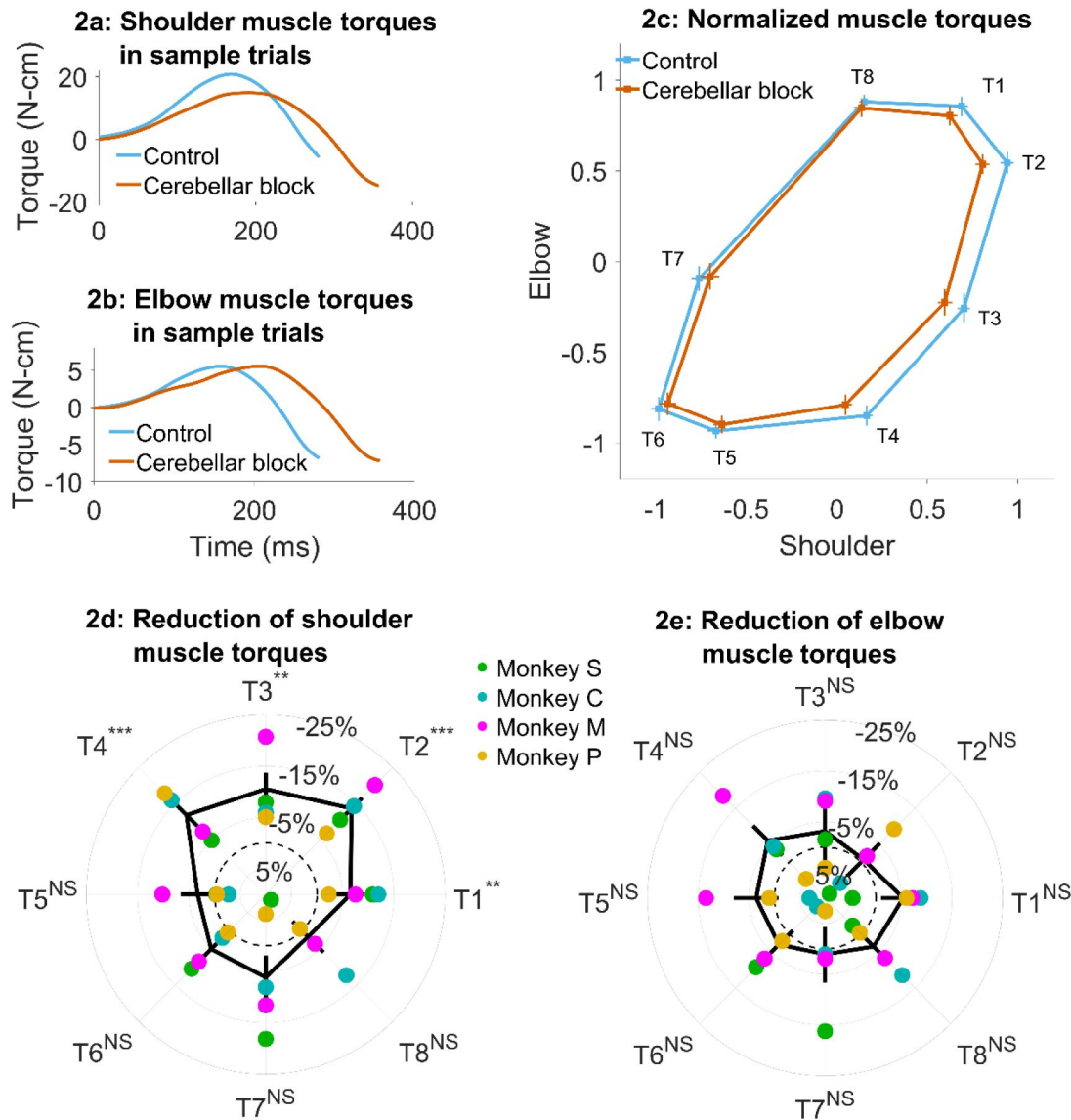


Figure 2

Effect of cerebellar block on muscle torques.

Movement-onset aligned shoulder (a) and elbow (b) muscle torque profiles during a sample control and cerebellar block trial performed by monkey C to target 2. Entry of the control cursor into the peripheral target marked the end of the movement for the trial. Since the monkeys did not have to stop their movements fully for the trial to end (see Methods for a detailed explanation of this), the traces appear cutoff at the beginning of the deceleration/stopping phase of the movement. (c) Normalized muscle torque impulse at the shoulder vs. elbow per target during control and cerebellar block. For each session, we computed the target-wise median muscle torque impulse during the acceleration phase of the movement across the control and cerebellar block trials. They were then normalized by the maximum absolute torque impulse across all the targets. The depicted values are the means across all the sessions pooled from the data of all four monkeys. The 95% confidence interval of the means is indicated by the horizontal and vertical bars for the shoulder and elbow joints respectively. Target-wise effect of cerebellar block on shoulder (d) and elbow (e) muscle torque impulse. For each session, the target-wise reduction in the median torque impulse during the cerebellar block trials was computed relative to the maximum absolute value of the target-wise medians observed during the control trials. The depicted values are the means $\pm$ 95% confidence intervals across all sessions pooled from all four monkeys. The means of individual monkeys are overlaid. The dashed circle indicates no change. Statistical significance is denoted as follows: $p \geq 0.05$ NS, $p < 0.05$ *, $p < 0.01$ **, $p < 0.001$ ***. [T1-8: Targets 1-8]

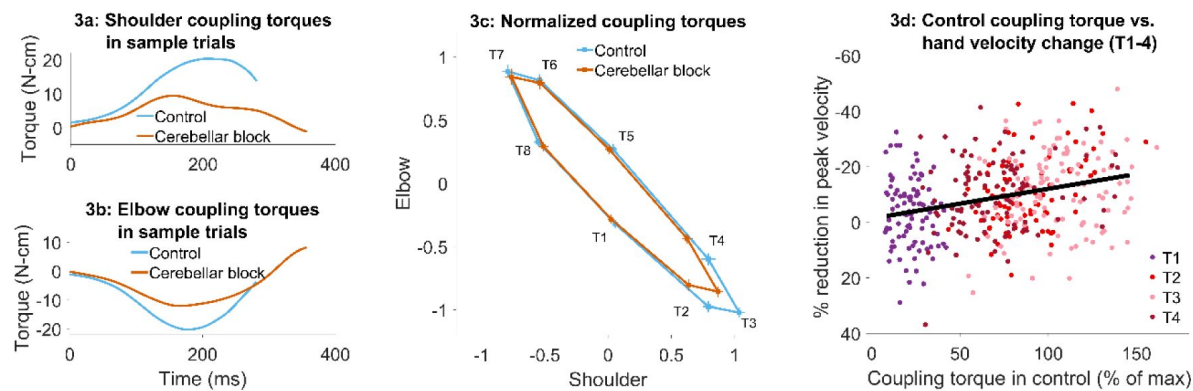


Figure 3

Effect of cerebellar block on coupling torques.

Movement-onset aligned shoulder (a) and elbow (b) coupling torque profiles during a sample control and cerebellar block trial performed by monkey C to target 2. Entry of the control cursor into the peripheral target marked the end of the trial. (c) Normalized coupling torque impulse at the shoulder vs. elbow per target during control and cerebellar block. For each session, we computed the target-wise median coupling torque impulse during the acceleration phase of the movement across the control and cerebellar block trials. They were then normalized by the maximum absolute torque impulse across all the targets. The depicted values are the means across all the sessions pooled from the data of all four monkeys. The 95% confidence interval of the means is indicated by the horizontal and vertical bars for the shoulder and elbow joints respectively. (d) The effect of the cerebellar block on peak hand velocity (y-axis) is plotted against the total coupling torque experienced across both joints during control (x-axis) for the outward reaches (T1-4). The effect of the cerebellar block on peak hand velocity was measured by the target-wise change in the median peak hand velocity during the cerebellar block trials relative to that of control trials per session (negative percentage values indicate a stronger reduction due to the cerebellar block). This measure was then correlated to the target-wise median coupling torque during the control trials of the corresponding sessions. The median coupling torque was taken as the sum of the absolute coupling torque impulse at both the shoulder and elbow joints per trial. These values were then normalized by the median across all trials of all sessions per monkey (to yield percentage units like the y-axis). The depicted data contains all the sessions pooled from all four monkeys with individual points representing each of the four targets (color-coded) per session. The black line overlayed on the scatter indicates the least-squares linear regression fit to the data points. [T1-8: Targets 1-8]

observed during the outward reaching movements with low coupling torques, indicating an additional component of muscle torque insufficiency that further exacerbated motor impairment following cerebellar block.

Hand velocity exhibits an adaptive trend during repeated movements to the same target during cerebellar block

The advantage of using a reversible cerebellar block in an animal model is the ability to track the temporal profile of motor impairments induced by this condition, allowing us to distinguish between primary and compensatory effects. To this end, we assessed changes in hand kinematics produced by the cerebellar block across successive trials. Specifically, we examined the evolution of peak hand velocities for outward reaches with low (target 1) vs. high coupling torques (data pooled for movements to targets 2-4) during blocks of control and cerebellar block trials. **Figure 4a** [↗](#) illustrates this procedure. Since the targets were presented in a pseudorandom order during each block of trials, we collected for each block the tested target, while keeping its order of presentation to generate a temporally accurate sequence of trials, all directed towards the same single target.

We found that during the control condition, peak hand velocity remained stable across successive trials. In contrast, cerebellar block affected hand velocity in a trial- and target-dependent manner (**Figure 4b, c** [↗](#)). For movements to target 1, hand velocity was lower than control but unaffected by the temporal sequence of target presentation (trial type $\times$ trial sequence interaction effect: $p = 0.69$, Supplementary Table S6). However, for movements to targets 2-4, hand velocity was lower than control in the early trials and declined further with successive trials (trial type $\times$ trial sequence interaction effect: $p = 0.007$, Supplementary Table S7).

To further quantify the effect of trial sequence on movements to targets 2-4, we performed *post hoc* comparisons (**Figure 4d** [↗](#)). These comparisons revealed that during the cerebellar block, hand velocity was 4.3% (CI [1.0, 7.3], $p < 0.001$) lower than control during the first 1-2 trials. With successive trials, the hand velocity declined further before reaching an asymptote during the late trials 11-20 at 8.2% (CI [5.7, 10.7], $p < 0.001$) lower than the control.

Taken together these results reveal that the hand velocity of outward-reaching movements with high coupling torques was already low initially during cerebellar block, reflecting a primary deficit. Over and above this, there was a further progressive decline of the hand velocity during successive movements, most likely due to an adaptive response to the inability to compensate for coupling torques. Furthermore, this adaptive response was not observed in the case of movements where the coupling torques were low, and therefore inter-joint interactions did not play a significant role.

Slow movements during the cerebellar block cannot fully account for the impaired motor timing

Individuals with cerebellar deficits often exhibit impaired motor timing, leading to movements that are poorly coordinated. Normally, smooth execution of multi-joint movements relies on precise temporal synchronization across joints. However, cerebellar lesions disrupt this temporal coordination, resulting in asynchronous, sequential joint activation resulting in movement decomposition as described earlier (Bastian, Martin et al. 1996 [↗](#)). Moreover, the disrupted temporal control of joint activation introduces variability from trial to trial, causing increased spatial variability in the resulting movement trajectories (Day, Thompson et al. 1998 [↗](#)). Thus, movement decomposition and trajectory variability can both be viewed as direct consequences of impaired temporal coordination due to cerebellar dysfunction. These properties of movement are also velocity-dependent (Bastian, Martin et al. 1996 [↗](#), Osu, Morishige et al. 2015 [↗](#)). Therefore, it is conceivable that the combination of primary deficit and adaptive reduction in velocity to reduce

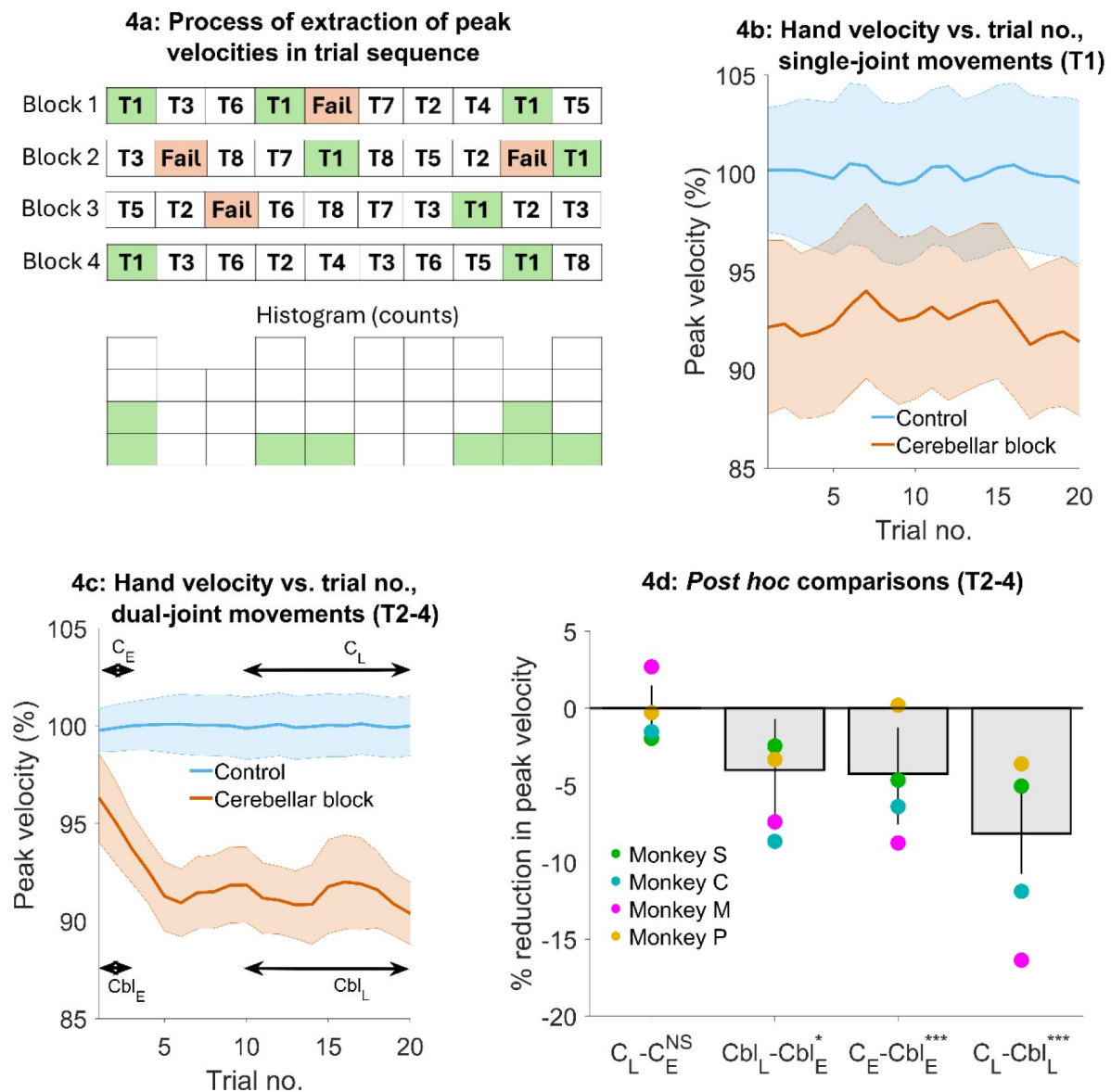


Figure 4

Effect of the cerebellar block on hand velocity across successive trials.

(a) Schematic of the process of extraction of peak hand velocity for movements to a particular target from blocks of control/cerebellar block trials while preserving the order in which the target was presented. For each block, the peak velocity for movements to a particular target (in this example target 1) was extracted while retaining the trial numbers in which it was presented in the block. This enabled us to examine the evolution of the peak velocity across the sequence of presentation of the target during control vs. cerebellar block. (b) Mean $\pm$ 95% confidence intervals of the peak hand velocity for movements to target 1 across all the trial blocks pooled from all four monkeys. For each monkey, the peak hand velocities were normalized to the median peak velocity of the early trials 1-2 from all the control blocks. (c) Same as in (b) but for targets 2-4. (d) Post hoc comparisons between early trials 1-2 and late trials 11-20 during control and cerebellar block. Statistical significance is denoted as follows: $p \geq 0.05$ NS, $p < 0.05$ *, $p < 0.01$ **, $p < 0.001$ ***. [E: Early, L: Late, C: Control, Cbl: Cerebellar block; T1-8: Targets 1-8]

the effects of coupling torque observed in our results is the main driver of abnormal motor timing. To evaluate this, we examined movement decomposition and trajectory variability during the outward reaching movements (which exhibited a significant reduction in hand velocity during cerebellar block) and measured the impact of hand velocity on these measures.

To quantify movement decomposition, we normalized the movement times of individual trials to range from 0 to 1 and identified the bins where either the shoulder or the elbow joint paused (speed < 20°/s) while the other joint continued moving (Bastian, Martin et al. 1996). In control trials, decomposition was primarily observed during the early phase of movement, where joint velocities are low making this measure more sensitive (**Figure 5a** and Supplementary Figure S3a). However, during cerebellar block trials, decomposition occurred throughout the movement (**Figure 5b** and Supplementary Figure S3b) similar to individuals with cerebellar deficits (Bastian, Martin et al. 1996). To quantify this effect, we calculated a decomposition index for each trial, representing the proportion of movement time during which the movement was decomposed, as defined above. Overall, the cerebellar block increased movement decomposition by 53.5% (CI [42.2, 66.0], $p < 0.001$). Next, we asked whether movement decomposition was mainly due to lower hand velocities. We therefore selected a subset of control trials that matched the cerebellar block trials in their peak velocities. Movement decomposition in these control trials was significantly lower than velocity-matched cerebellar block trials ($p < 0.001$; **Figure 5c**; see Supplementary Figure S3c for target-wise effects), even though these control trials themselves had 11.0% (CI [5.2, 17.0], $p = 0.03$) higher decomposition than all control trials together. This indicates that even at matched velocities, movements during cerebellar block were more decomposed relative to the control.

Next, we assessed the effect of cerebellar block and hand velocity on trajectory variability. To this end, we first aligned the hand trajectories to have the same starting position and then rotated them so that their endpoint was located on the positive Y-axis (**Figure 5d**; for visualization in joint position space see Supplementary Figures S4a and b). Subsequently, we defined the error in each trajectory as the maximal perpendicular distance from the Y-axis. The standard deviation of the errors across all trials was used to quantify the trial-to-trial trajectory variability. When we compared the subset of velocity matched control and cerebellar block trials, we found that cerebellar block exhibited 34.6% (CI [26.2, 43.2], $p < 0.001$) higher trajectory variability (**Figure 5e**; see Supplementary Figure S4c for target-wise effects). Normally, slower movements are also less variable due to the speed-accuracy tradeoff (Plamondon and Alimi 1997). Indeed, the trajectory variability in this subset of slower control trials was 5.5% (CI [0.9, 9.9], $p = 0.02$) lower than that of all control trials. In other words, despite slower movements, cerebellar block increased trajectory variability.

Finally, we quantified how cerebellar block affected movement decomposition across successive trials similar to our analysis of hand velocities in **Figure 4**. The cerebellar block induced increase in movement decomposition was unaffected by the trial-sequence (trial type $\times$ trial sequence interaction effect: $p = 0.43$, Supplementary Table S8 and Supplementary Figure S5). This further demonstrates that movement decomposition remained high during cerebellar block despite slower movement speeds.

Discussion

Individuals with cerebellar deficits exhibit altered motor behavior related mostly to poor timing and coordination of voluntary movements. Understanding the impairments that follow cerebellar lesions is often complicated by the fact that the studies of this question mostly rely on a heterogeneous population due to various levels and types of cerebellar pathologies. Here we addressed this question by using high-frequency stimulation to reversibly block cerebellar output to the motor cortex in behaving monkeys wearing an exoskeleton (Nashef, Cohen et al. 2019).

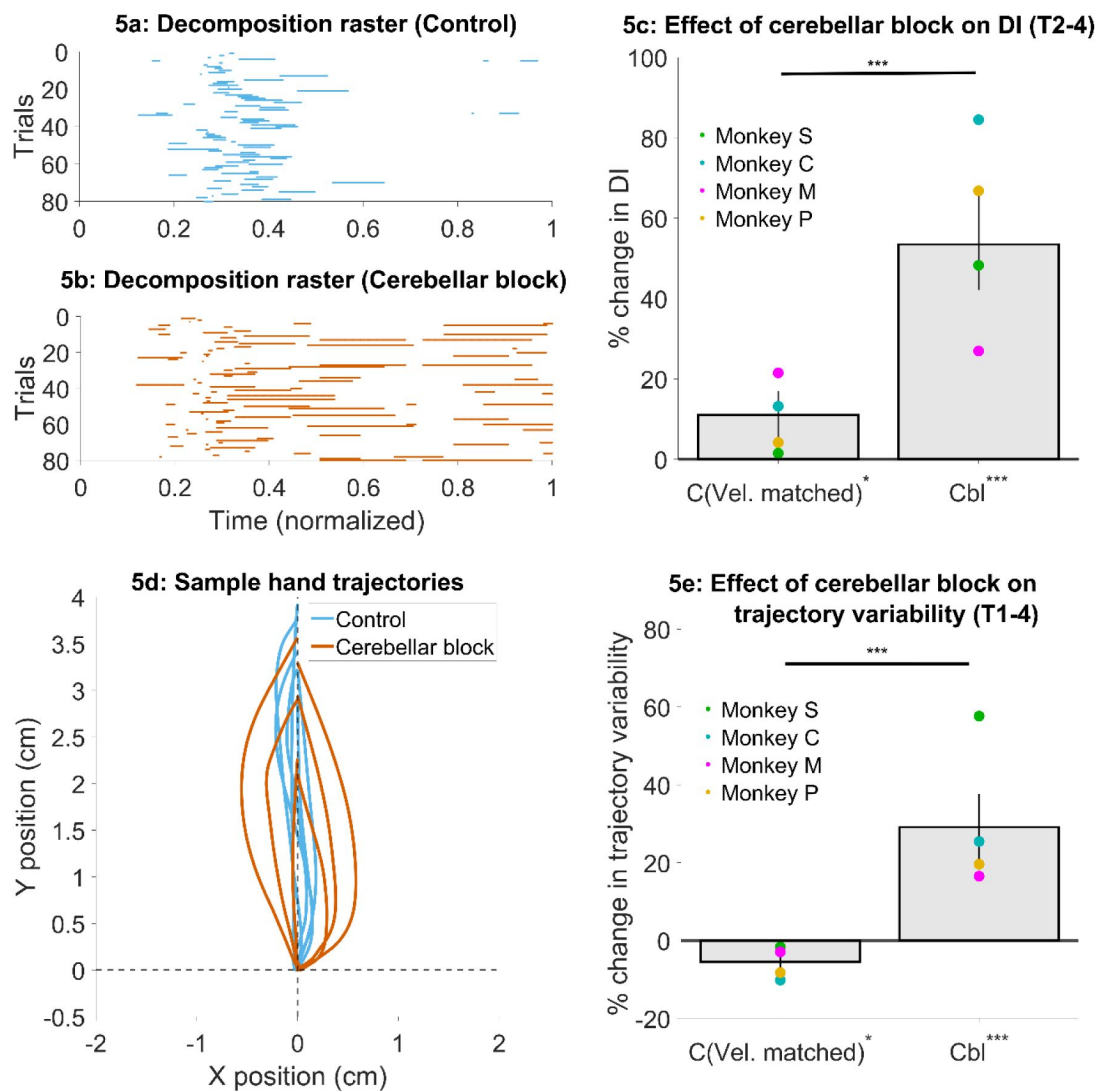


Figure 5a-c

Effect of the cerebellar block on the decomposition of movements.

(a) Binary decomposition raster of sample trials for movements made to target 2 by monkey C during control. The duration of each trial was normalized to 0-1 (bin width = 0.001) and then decomposition was computed for each bin based on whether either (but not both) of the shoulder or the elbow joint velocity was less than 20°/s. (b) Same as in (a) but for movements to target 2 by monkey C during cerebellar block. (c) Change in decomposition index (i.e., the proportion of the movement time during which the movement was decomposed, as defined above) for movements to targets 2-4 during velocity-matched control and cerebellar block trials relative to all control trials. The change in median decomposition was computed for each session. The depicted values are the mean $\pm$ 95% confidence intervals across all sessions pooled from all four monkeys. The individual means of each monkey are overlaid. **Figure 5d-e: Effect of the cerebellar block on inter-trial trajectory variability.** (d) Sample hand trajectories during control vs. cerebellar block trials for movements to target 2 by monkey C. The starting point of the trajectories was shifted to the origin and then they were rotated about the origin so that their endpoints lie on the positive Y-axis. (e) Change in inter-trial trajectory variability for movements to targets 2-4 during velocity-matched control and cerebellar block trials relative to all control trials. The trajectory variability was measured as the standard deviation of the maximum perpendicular distance of the trajectories from the Y-axis after transforming them as in (d). The change in trajectory variability for the cerebellar block trials was computed relative to the control trials for each session. The depicted values are the mean $\pm$ 95% confidence intervals across all sessions pooled from all four monkeys. The individual means of each monkey are overlaid. Statistical significance is denoted as follows: $p \geq 0.05$ NS, $p < 0.05$ *, $p < 0.01$ **, $p < 0.001$ ***. [C: Control, Cbl: Cerebellar block; T1-8: Targets 1-8; DI: Decomposition index]

and comparing the post-block motor behavior to control conditions. Our results indicate that cerebellar block leads to decreased hand velocity, particularly during the outward reaching movements via two mechanisms: a reduction in muscle torques suggesting motor weakness, and spatially tuned decreases in velocity in targets where large inter-joint interactions occur, indicating a failure to compensate for limb dynamics. Analyzing the time course of these impairments revealed partially overlapping processes: an initial reduction in hand velocity due to the inability to generate sufficient muscle torques, followed by a further progressive decline in hand velocities during subsequent movements during the cerebellar block. This secondary decline was observed only in the case of movements with high inter-joint interactions, suggesting an adaptive strategy to minimize these interactions. Finally, we observed increased motor noise in terms of movement decomposition and trial-to-trial trajectory variability which were independent of the slower movements observed during the cerebellar block.

Target-dependent reduction in hand velocity and coupling torques during reaching movements during the cerebellar block

Cerebellar block led to a target-specific reduction in hand velocity, even though the arm was fully supported by the exoskeleton and thus the limb weight did not play any role. This observation replicates findings made in previous studies that have documented the slowing of movements in individuals with cerebellar lesions (Wild and Dichgans 1993 [↗](#), Topka, Konczak et al. 1998 [↗](#), Konczak, Schoch et al. 2005 [↗](#)) as well as in monkeys that underwent dentate cooling (Tsujimoto, Gemba et al. 1993 [↗](#)). It was argued that the reduction in hand velocity may be a compensatory strategy to counteract the impaired ability to manage inter-joint coupling torques in the absence of cerebellar input to the motor cortex (Wild and Dichgans 1993 [↗](#), Bastian and Thach 1995 [↗](#), Bastian, Martin et al. 1996 [↗](#), Beer, Dewald et al. 2000 [↗](#)). Since coupling torques in multi-joint movements are generated in a velocity-dependent manner (Hollerbach and Flash 1982 [↗](#), Virji-Babul and Cooke 1995 [↗](#)), slowing down the movement not only simplifies its kinetics (by reducing the impact of coupling torques) but also allows more time for slower visual feedback to aid in guiding the movement (Jeannerod 1988 [↗](#)). Our findings, which demonstrate a correlation between the extent of hand velocity reduction during cerebellar block and the magnitude of coupling torque under control conditions, further corroborate this hypothesis and confirm our approach for reversible inactivation of cerebellar output as a model for ataxia. Furthermore, our model enabled us to study this effect rigorously and systematically while dissociating it from the other primary effects of cerebellar block as discussed below.

Muscle torque deficit is an acute response to the cerebellar block

Poor compensation for coupling torque is a fundamental deficit in cerebellar ataxia (Bastian, Zackowski et al. 2000 [↗](#)). However, early studies have reported the occurrence of muscle weakness (asthenia) and hypotonia acutely following cerebellar injury in humans (Haines, Manto et al. 2007 [↗](#)) or experimental lesions in animals (Luciani 1893 [↗](#), Bremer, Roger et al. 1935 [↗](#), Fulton and Dow 1937 [↗](#), Granit, Holmgren et al. 1955 [↗](#)). Both Luciani (Luciani 1893 [↗](#)) and Holmes (Holmes 1917 [↗](#)) described a triad of cerebellar deficits to be atonia (low muscle tone), asthenia (weakness of voluntary movements), and astasia (oscillation of the head and trunk). Importantly, muscle weakness is typically characteristic of the acute phase of injury, especially in the upper extremity, and tends to normalize gradually over weeks to months, depending on the severity of the injury (Konczak, Pierscianek et al. 2010 [↗](#)). It is therefore conceivable that studies in humans with cerebellar deficits, which often included subjects at the chronic stage of the lesion, may have underestimated the contribution of muscle weakness which by that time was already less pronounced.

We explored the acute outcome of the cerebellar block, a phase in which agonist muscle weakness may play a more prominent role. Importantly, we showed that deficits in the self-generated muscle torque appear immediately after the cerebellar block, resulting in lower hand velocity already in the first few trials of movements. This kind of fast response is less likely to be a strategic response

aimed at mitigating the loss of cerebellar signals. This finding is consistent with the initial weakness followed by a gradual recovery of function observed in acute lesions of other neural structures involved in movement control, including the primary (Pressman and Rosen 2015 [↗](#)) and premotor cortices (Freund 1985 [↗](#), Freund and Hummelsheim 1985 [↗](#)). It also aligns with the understanding that cerebellar output provides a strong excitatory drive to the motor cortex (Shinoda, Yamazaki et al. 1982 [↗](#), Hore and Flament 1988 [↗](#), Nashef, Cohen et al. 2018 [↗](#)) and several subcortical descending motor systems (Ruigrok 2013 [↗](#)), beyond its well-known role in motor timing and coordination. Following the initial reduction in hand velocity observed in our study, there was a further gradual decline across subsequent trials specifically for the movements with high coupling torques, a behavior profile that is more consistent with strategic effort, most likely in response to the inability to compensate for limb dynamics.

The muscle torque, representing the sum of all muscle forces acting at a joint, can also be reduced by co-contraction of agonist and antagonist muscles rather than by agonist weakness alone. However, in cerebellar ataxia, co-contraction has been proposed as a strategy to stabilize stationary joints during decomposed multi-joint movements (Bastian, Martin et al. 1996 [↗](#)). In our experiments, this strategy would likely emerge gradually following cerebellar block similar to the adaptive slowing of movements aimed at reducing inter-joint interactions. We therefore argue that the observed acute deficiency in muscle torque was mainly driven by agonist weakness. Our argument is further supported by previous studies which attributed reduced agonist muscle activity as a cause for slowing of voluntary movements in individuals with cerebellar lesions (Hallett, Berardelli et al. 1991 [↗](#), Wild, Klockgether et al. 1996 [↗](#)).

In summary, the slower movements in the absence of cerebellar signals are likely due to two main factors: (i) a primary inability to generate sufficient muscle torques due to the lack of synchronized recruitment of the cortical and sub-cortical descending motor systems via the cerebellar output, and (ii) a secondary adaptive strategy aimed at compensating for the impaired prediction of limb dynamics due to the loss of signals from the cerebellum.

Insufficient muscle torques vs. poor inter-joint coordination

Unlike most previous clinical studies that examined motor deficits in individuals with chronic cerebellar injuries, our study utilized a reversible model of cerebellar block. This approach enabled us to track the temporal evolution of motor deficits from the onset of the block within the same study participant. Although several studies have explored the nature of motor deficits following cerebellar injuries, there remains considerable debate regarding the fundamental mechanisms driving these deficits. Some studies have suggested that the inability to compensate for inter-joint interactions is the primary driver of the observed motor abnormalities in cerebellar injury (Bastian, Martin et al. 1996 [↗](#), Bastian, Zackowski et al. 2000 [↗](#)). In contrast, other studies have emphasized the inability to generate sufficient muscle torques as the core deficit in cerebellar ataxia while claiming that inter-joint coordination is unaffected following cerebellar lesions (Topka, Konczak et al. 1998 [↗](#), Boose, Dichgans et al. 1999 [↗](#)). Given these conflicting findings, our study aimed to systematically explore the entire spectrum of motor deficits using a model of reversible cerebellar block. First, we found that the deficits in muscle torque are more pronounced for the reaching movements as compared to the retrieval movements. We identified both an acute insufficiency in muscle torque generation and the inability to compensate for coupling torques as potential mechanisms underlying the observed slowness during the reaching movements. Moreover, we delineated the time course of these deficits, providing insights into their evolution from the onset of cerebellar block.

The asymmetric effect of cerebellar block on outward vs. inward reaching movements

We observed a notable distinction in the impact of cerebellar blockage on outward vs. inward reaching movements. At a collective level, this variance could be attributed to the disparity in magnitude of coupling torque impulse, with movements towards targets 2-4 exhibiting 18.7% (CI [13.1, 24.1], $p = 0.008$) higher coupling torque impulse compared to those towards targets 6-8. However, this explanation does not hold consistently when examined at the individual target level. For instance, while the coupling torque was higher for movements towards target 7, the effects of cerebellar blockage were less pronounced compared to those observed for target 1, where the coupling torques are considerably lower. There are two possible explanations for this discrepancy. One possibility is that the effect of cerebellar block exhibits a bias in the direction of shoulder flexion (i.e., for movements to targets 1-4). Several early studies in cats (Hare, Magoun et al. 1936 [\[1\]](#), Chambers 1947 [\[2\]](#)), non-human primates (Magoun, Hare et al. 1935 [\[3\]](#)), and humans (Nashold and Slaughter 1969 [\[4\]](#)) have all reported a bias towards the activation of ipsilateral flexors of the arm following stimulation of the deep cerebellar nuclei and the superior cerebellar peduncle. Furthermore, lesioning of these structures led to the reduction of ipsilateral flexor tone and power (Schneider and Crosby 1963 [\[5\]](#), Nashold and Slaughter 1969 [\[4\]](#)). The cerebellar output influences the excitability of spinal interneurons either directly (Bantli and Bloedel 1975 [\[6\]](#), Asanuma, Thach et al. 1980 [\[7\]](#), Sathyamurthy, Barik et al. 2020 [\[8\]](#)) or via indirect relays in the subcortical descending motor systems like the rubrospinal and the reticulospinal pathways (Ruigrok 2013 [\[9\]](#)). Since the spinal interneurons influence the activity of flexor muscles twice as often as extensor muscles (Perlmutter, Maier et al. 1998 [\[10\]](#)), this might explain why the cerebellar block had a stronger effect on movements involving shoulder flexion (i.e., outward reaching movements) relative to shoulder extension (i.e., inward reaching movements) in our study. Alternatively, the initial position of the limb in our experimental setup (i.e., on the central target) might have been favorable for movements in the direction of shoulder extension, thereby making them easier to perform.

Asynchronous joint movements and motor variability occur despite slower movements in cerebellar block

Abnormally high motor variability (Day, Thompson et al. 1998 [\[11\]](#), Timmann, Watts et al. 1999 [\[12\]](#), Schlerf, Xu et al. 2012 [\[13\]](#)) and decomposition of multi-joint movements (Bastian, Martin et al. 1996 [\[14\]](#)) are core deficits in cerebellar patients. We showed that the cerebellar block decomposed the dual-joint movements into temporally isolated single-joint actions and led to an increased trial-to-trial spatial noise. These two measures can be affected by the reduction in movement velocities. Our results showed that even though movement decomposition increased with a reduction in velocity in control trials, they were still significantly lower than that of velocity-matched cerebellar block trials. Second, the trial-to-trial spatial noise in hand trajectories increased during cerebellar block despite the slower movement speeds. The cerebellum is known to participate in the planning of movements by providing an internal model of the limb which is necessary for predicting the sensory outcomes of motor commands (Wolpert, Miall et al. 1998 [\[15\]](#)). For this purpose, the cerebellum aids in estimating the state of the limb by integrating the sensory information of its last known state with predictions of its response to the latest motor command (Miall, Christensen et al. 2007 [\[16\]](#), Miall and King 2008 [\[17\]](#)). Therefore, in the absence of cerebellar signals, there are programming errors in the motor command required to launch the limb accurately toward a target (Day, Thompson et al. 1998 [\[11\]](#)). These errors may in turn translate into the noisy joint activation patterns and variable hand trajectories observed in our results. Moreover, the persistence of these deficits independently of the movement speeds observed during cerebellar block indicates that they cannot be compensated for, unlike the impaired prediction of inter-joint interactions which are strategically dealt with by reducing the movement velocity.

In summary, our study systematically investigated the temporal progression and distinct mechanisms of motor deficits in a controlled, reversible animal model of cerebellar block. These include an acute onset muscle torque deficit, impaired compensation of limb dynamics, and increased motor noise. We aim to further explore the neural correlates of these deficits in future studies.

Materials and Methods

Experimental subjects

This study was performed on four adult female monkeys (*Macaca fascicularis*, weight 4.5-8 kg). The care and surgical procedures of the subjects will be in accordance with the Hebrew University Guidelines for the Use and Care of Laboratory Animals in Research, supervised by the Institutional Committee for Animal Care and Use.

Behavioral task

Each monkey was trained to sit in a primate chair and perform planar, center-out movements. During the task, the left upper arm was abducted 90° and rested on cushioned troughs secured to links of a two-joint exoskeleton (KINARM, BKIN Technologies Ltd.). The sequence of events in each trial of this task was as follows. First, the monkey located a cursor (projected on a horizontal screen immediately above its arm) within a central target (2-3 cm in diameter). After 500-800 ms, a randomly selected peripheral target (out of 8 evenly distributed targets) was displayed. The monkey had to wait 450-850 ms until the central target disappeared (i.e., the ‘go’ signal) and then reached the cued peripheral target (2-3 cm in diameter) located 4-5 cm away. For monkeys C and M, we inserted a 200 ms grace period before the ‘go’ signal to encourage predictive timing. The onset of movement within this time before the ‘go’ signal did not abort the trial. We limited the time to reach the peripheral target after the ‘go’ signal to 500 ms for monkeys M and C and 800-900 ms for monkeys S and P. Following successful reach, the monkeys were rewarded with a drop of applesauce. It is important to note that the monkeys were required to hold at the target briefly—100 ms for Monkeys S and P, and 150 ms for Monkeys C and M— before receiving the reward. However, given the size of the targets and the velocity of movements, it often happened that the monkeys didn’t have to stop their movements fully to obtain the reward. Importantly, we relaxed the task’s requirements (by increasing the target size and reducing the temporal constraints) to enable the monkeys to perform a sufficient number of successful trials under both the control and the cerebellar block conditions. This was necessary as we found that strict criteria for these parameters yielded a very low success rate in the cerebellar block condition. Given these practical constraints, we note that this task design is likely suboptimal for studying endpoint accuracy which is an important aspect of cerebellar control (Low, Thanawalla et al. 2018 [DOI](#), Becker and Person 2019 [DOI](#), Calame, Becker et al. 2023 [DOI](#)).

Insertion of the stimulating electrode into the superior cerebellar peduncle

After the training was completed, a square recording chamber (24x24 mm) was attached to the monkey’s skull above the upper limb-related area of the motor cortex in a surgical procedure under general anesthesia. To insert a chronic stimulating electrode into the ipsilateral superior cerebellar peduncle (SCP), a round chamber (diameter = 19 mm) was also implanted above the estimated insertion point (as per stereotactic measurements) during the surgery for the cortical chamber implantation. Subsequently, a post-surgery MRI was used to plan the trajectory for the electrode insertion into the SCP. After a recovery and re-training period, a bi-polar concentric electrode (Microprobes for Life Science Inc.) was inserted through the round chamber along the

planned trajectory. During the process of insertion, the evoked multi-unit activity in the primary motor cortex was simultaneously monitored to identify the precise location where the stimulating electrode should be secured (Nashef, Cohen et al. 2018 [DOI](#), Nashef, Rapp et al. 2018 [DOI](#)).

High-frequency stimulation to block the cerebellar outflow to the motor cortex

The superior cerebellar peduncle was blocked using high-frequency stimulation (HFS). This stimulation protocol consisted of a train of biphasic rectangular-pulse stimuli (where each phase was 200 μ s) applied at 130 Hz. Each train was delivered at a fixed intensity of 150-250 μ A. In our earlier study, we demonstrated that by using this manipulation we can successfully replicate the impaired motor timing and coordination characteristic of cerebellar ataxia (Nashef, Cohen et al. 2019 [DOI](#)). Furthermore, we also demonstrated that at the motor cortical level, this manipulation leads to a loss of response transients at movement onset and decoupling of task-related activity, similar to findings observed in studies where the dentate nucleus was cooled (Meyer-Lohmann, Conrad et al. 1975 [DOI](#)). Deep brain stimulation (DBS) is a perturbation similar in properties to the HFS we used, which is routinely used in clinical practice for similar purposes of interfering with neural circuitries that underwent pathological changes (Perlmutter and Mink 2006 [DOI](#)) although in both applications there is no accurate measure for the extent of the block (i.e., the fraction of fibers which become ineffective in these conditions) other than the behavioral consequences. It is possible that HFS probably imposes only a partial block of the CTC system.

It is important to note that the superior cerebellar peduncle is known to carry input and output fibers to and from several neural structures (Naidich, Stein et al. 2013 [DOI](#)). Therefore, we cannot rule out the fact that the effect of HFS may be mediated in part through pathways other than the cerebello-thalamo-cortical pathway (as mentioned in the Discussion section). However, in primates, the cerebellar-thalamo-cortical pathway greatly expanded (at the expense of the cerebello-rubro-spinal tract) in mediating cerebellar control of voluntary movements (Horne and Butler 1995 [DOI](#)). The cerebello-subcortical pathways diminished in importance throughout evolution (Padel, Angaut et al. 1981 [DOI](#), Nathan and Smith 1982 [DOI](#), ten Donkelaar 1988 [DOI](#)). Previously we found that the ascending spinocerebellar axons which enter the cerebellum through the superior cerebellar peduncle (SCP) are weakly task-related and the descending system is quite small (Cohen et al. 2017). Nevertheless, we acknowledge that HFS disrupts cerebellar communication to the rest of the nervous system broadly, rather than solely the cerebello-thalamo-cortical pathway.

Data collection protocol

Each monkey performed the experimental task for several days (sessions). A total of 46, 29, 36, and 54 sessions of data were collected from Monkeys S, C, M, and P, respectively. Each session was divided into three or four sub-sessions. Each sub-session included blocks of control trials (~80 trials) and trials with the superior cerebellar peduncle blocked using high-frequency stimulation (~50 trials) as described above. During the task performance, the following behavioral data was obtained at 1000 Hz using the KINARM's motor encoders: (i) Hand position in the x-y plane of movement, (ii) Elbow and shoulder joint angular positions, velocities, and accelerations. The lengths of the upper arm and forearm of each monkey were measured as the distances from the head of the humerus to the lateral epicondyle and from the lateral epicondyle to the palm, respectively. The inertial properties of the upper limb were estimated using the monkey's body weight (Cheng and Scott 2000 [DOI](#)).

Processing of the behavioral data

All data analysis was performed in MATLAB R2021b (MathWorks Inc.). The raw kinematic data was low-pass filtered by using a second order Butterworth filter (cutoff = 10Hz) and then epoched into trials. For this purpose, the start and end events were taken as the time of presentation of the

cue (i.e., the peripheral target) and the time of entry of the control cursor into the peripheral target, respectively. This cutoff meant that a major portion of the extracted segment of each trial was the acceleration phase of the movement. The movement onset was defined as the time when the radial hand velocity exceeded 5% of its peak. Subsequently, all the trials were zero-centered around the calculated movement onset. We then applied a two-link inverse dynamics model of the upper arm to the kinematic data to compute the coupling and muscle torques acting on the elbow and shoulder joints (Fagg, Ojakangas et al. 2009 [DOI](#)). The inverse dynamics equations of motion for the combined system consisting of the monkey's arm and the associated moving components of the KINARM were used to compute the joint torques:

$$T_{me} = T_{ne} - T_{ce} \quad (i)$$

$$T_{ms} = T_{ns} - T_{cs} \quad (ii)$$

where T_{ne} and T_{ns} are the net torques (proportional to the joint's acceleration), T_{ce} and T_{cs} are passive coupling torques, and T_{me} , T_{ms} are the active muscle torques acting at the elbow and shoulder joints respectively. The expanded formulas for computing each of the above terms are provided by equations 1a-2e in the methods section C (Fagg, Ojakangas et al. 2009 [DOI](#)). It must be noted that the 'muscle' torques computed here include the effects of the actively generated muscle forces as well as the viscoelastic effects that result from the musculoskeletal system and the KINARM (Smith and Zernicke 1987 [DOI](#)). Data from some of the same animals was presented in prior work where we showed that during reaching, cerebellar block resulted in reduced reach velocity, increased trajectory curvature, and shoulder-elbow dyscoordination (Nashef, Cohen et al. 2019 [DOI](#)). In this study, we extend these findings by using inverse-dynamics modeling to explore the relationship between altered kinematics and joint torques. Furthermore, the analyses exploring progressive velocity changes across trials under the cerebellar block and the relationship of motor noise to movement velocity are newly reported in this study.

To strictly focus on feedforward control, we could have examined the measured variables in the first 50-100 ms of the movement which has been shown to be unaffected by feedback responses (Todorov and Jordan 2002 [DOI](#), Pruszynski, Kurtzer et al. 2008 [DOI](#), Pruszynski and Scott 2012 [DOI](#), Crevecoeur and Scott 2013 [DOI](#)). However, in our task, the amplitude of movements made by our monkeys was small, and therefore the response measures we used were too small in the first 50-100 ms for a robust estimation. Also, fixing a time window led to an unfair comparison between control and cerebellar block trials, in which velocity was significantly reduced and therefore movement time was longer. Therefore, we used the peak velocity, torque-impulse at the peak velocity, and maximum deviation of the hand trajectory as response measures.

Statistical analysis

We used linear mixed effects analysis to evaluate the effects of predictor variables of interest on the computed response variables. This allowed us to account for the variability across the monkeys which was modeled as a random effect. Each data sample for this model was obtained by calculating the median of all the trials for each unique combination of the response variable(s) per session. We computed the change in the measured outcome variable due to the cerebellar block per session using the following formula:

$$\frac{\text{Median response during cerebellar block} - \text{median response during control}}{\text{Median response during control}}$$

For post-hoc comparisons, we used the 'coefTest' function from MATLAB's 'fitlme' to perform hypothesis testing on the fixed effects. This involved specifying the contrast matrices to evaluate specific hypotheses regarding the differences in the response variables across the levels of the predictor variables. The 'coefTest' function allowed us to obtain p-values for these comparisons. The computed p-values were then corrected using the Benjamini-Hochberg procedure to control

the false discovery rate for multiple post-hoc comparisons, wherever applicable (Haynes 2013 [↗](#)). A significance level of 0.05 was used for all comparisons in this study. For estimation of the group means and their confidence intervals, we pooled the per session data from all monkeys after ensuring that effects are consistent across the monkeys. For further clarity, we have also overlaid the means of the measured variables from individual monkeys on all our plots of summary statistics.

Data availability statement

Data and code used in this manuscript are available publicly as a GitHub repository (Sinha, Israely et al. 2024 [↗](#)).

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Additional files

Supplementary information [↗](#)

References

- Asanuma C., Thach W. T., Jones E. G. (1980) **Nucleus interpositus projection to spinal interneurons in monkeys** *Brain Research* **191**:245–248 [Google Scholar](#)
- Bantli H., Bloedel J. R. (1975) **The action of the dentate nucleus on the excitability of spinal motoneurons via pathways which do not involve the primary sensorimotor cortex** *Brain Research* **88**:86–90 [Google Scholar](#)
- Bastian A. J. (1997) **Mechanisms of Ataxia** *Physical Therapy* **77**:672–675 [Google Scholar](#)
- Bastian A. J., Martin T. A., Keating J. G., Thach W. T. (1996) **Cerebellar ataxia: abnormal control of interaction torques across multiple joints** *Journal of Neurophysiology* **76**:492–509 [Google Scholar](#)
- Bastian A. J., Thach W. T. (1995) **Cerebellar outflow lesions: A comparison of movement deficits resulting from lesions at the levels of the cerebellum and thalamus** *Annals of Neurology* **38**:881–892 [Google Scholar](#)
- Bastian A. J., Zackowski K. M., Thach W. T. (2000) **Cerebellar Ataxia: Torque Deficiency or Torque Mismatch Between Joints?** *Journal of Neurophysiology* **83**:3019–3030 [Google Scholar](#)
- Becker M. I., Person A. L. (2019) **Cerebellar Control of Reach Kinematics for Endpoint Precision** *Neuron* **103**:335–348 [Google Scholar](#)
- Beer R. F., Dewald J. P. A., Rymer W. Z. (2000) **Deficits in the coordination of multijoint arm movements in patients with hemiparesis: evidence for disturbed control of limb dynamics** *Experimental Brain Research* **131**:305–319 [Google Scholar](#)
- Beppu H., Suda M., Tanaka R. (1984) **Analysis of cerebellar motor disorders by visually guided elbow tracking movement** *Brain* **107**:787–809 [Google Scholar](#)
- Bernshtein N. A. (1967) **The co-ordination and regulation of movements** Oxford: Pergamon Press Oxford [Google Scholar](#)
- Boose A., Dichgans J., Topka H. (1999) **Deficits in phasic muscle force generation explain insufficient compensation for interaction torque in cerebellar patients** *Neuroscience Letters* **261**:53–56 [Google Scholar](#)
- Bremer F., Roger G.-H., Binet L. (1935) **Le cervelet** *Traité de physiologie* :39–134 [Google Scholar](#)
- Calame D. J., Becker M. I., Person A. L. (2023) **Cerebellar associative learning underlies skilled reach adaptation** *Nat Neurosci* **26**:1068–1079 [Google Scholar](#)
- Chambers W. W. (1947) **Electrical stimulation of the interior of the cerebellum in the cat** *Am. J. Anat* **80**:55–93 [Google Scholar](#)
- Cheng E. J., Scott S. H. (2000) **Morphometry of Macaca mulatta forelimb. I. Shoulder and elbow muscles and segment inertial parameters** *Journal of Morphology* **245**:206–224 [Google Scholar](#)

- Crevecoeur F., Scott S. H. (2013) **Priors Engaged in Long-Latency Responses to Mechanical Perturbations Suggest a Rapid Update in State Estimation** *PLOS Computational Biology* **9**:e1003177 [Google Scholar](#)
- Day B. L., Thompson P. D., Harding A. E., Marsden C. D. (1998) **Influence of vision on upper limb reaching movements in patients with cerebellar ataxia** *Brain* **121**:357–372 [Google Scholar](#)
- Fagg A., Ojakangas G., Miller L., Hatsopoulos N. (2009) **Kinetic Trajectory Decoding Using Motor Cortical Ensembles** *IEEE transactions on neural systems and rehabilitation engineering : a publication of the IEEE Engineering in Medicine and Biology Society* **17**:487–496 [Google Scholar](#)
- Freund H.-J (1985) **Clinical aspects of premotor function** *Behavioural Brain Research* **18**:187–191 [Google Scholar](#)
- Freund H. J., Hummelsheim H. (1985) **Lesions of premotor cortex in man** *Brain : a journal of neurology* **108**:697–733 [Google Scholar](#)
- Fulton J. F., Dow R. S. (1937) **The Cerebellum: A Summary of Functional Localization** * *The Yale Journal of Biology and Medicine* **10**:89–119 [Google Scholar](#)
- Goldstein K (1927) **Das Kleinhirn** *Handbuch der normalen und pathologischen Physiologie* **10**:222–222 [Google Scholar](#)
- Granit R., Holmgren B., Merton P. A. (1955) **The two routes for excitation of muscle and their subservience to the cerebellum** *The Journal of Physiology* **130**:213–224 [Google Scholar](#)
- Hage P., Fakharian M. A., Shoup A. M., Pi J. S., Sedaghat-Nejad E., Orozco S. P., Jang I. K., Looi V., Elseweifi H. Y., Mohammadrezaei N., Vasserman A. N., Arginteanu T., Shadmehr R. (2025) **Control of tongue movements by the Purkinje cells of the cerebellum** *bioRxiv* [Google Scholar](#)
- Haines D. E., Manto M.-U., Glickstein M. (2007) **Clinical symptoms of cerebellar disease and their interpretation** *The Cerebellum* **6**:141–156 [Google Scholar](#)
- Hallett M., Berardelli A., Matheson J., Rothwell J., Marsden C. D. (1991) **Physiological analysis of simple rapid movements in patients with cerebellar deficits** *J Neurol Neurosurg Psychiatry* **54**:124–133 [Google Scholar](#)
- Hare W. K., Magoun H. W., Ranson S. W. (1936) **Electrical stimulation of the interior of the cerebellum in the decerebrate cat** *American Journal of Physiology-Legacy Content* **117**:261–266 [Google Scholar](#)
- Haynes W (2013) **Benjamini-Hochberg Method** In: Dubitzky W., Wolkenhauer O., Cho K.-H., Yokota H., editors. *Encyclopedia of Systems Biology* New York, NY: Springer New York pp. 78–78 [Google Scholar](#)
- Hollerbach J. M., Flash T. (1982) **Dynamic interactions between limb segments during planar arm movement** *Biological Cybernetics* **44**:67–77 [Google Scholar](#)
- Holmes G (1917) **The symptoms of acute cerebellar injuries due to gunshot injuries** *Brain: A Journal of Neurology* **40**:461–535 [Google Scholar](#)
- Holmes G (1939) **The cerebellum of man** *Brain* **62**:1–30 [Google Scholar](#)

Hore J., Flament D. (1988) **Changes in motor cortex neural discharge associated with the development of cerebellar limb ataxia** *Journal of Neurophysiology* **60**:1285–1302 [Google Scholar](#)

Horne M. K., Butler E. G. (1995) **The role of the cerebello-thalamo-cortical pathway in skilled movement** *Progress in Neurobiology* **46**:199–213 [Google Scholar](#)

Ivry R. B., Keele S. W. (1989) **Timing Functions of The Cerebellum** *Journal of Cognitive Neuroscience* **1**:136–152 [Google Scholar](#)

Jeannerod M (1988) **The neural and behavioural organization of goal-directed movements** New York, NY, US: Clarendon Press/Oxford University Press [Google Scholar](#)

Kawato M (1999) **Internal models for motor control and trajectory planning** *Current Opinion in Neurobiology* **9**:718–727 [Google Scholar](#)

Konczak J., Pierscianek D., Hirsiger S., Bultmann U., Schoch B., Gizewski E. R., Timmann D., Maschke M., Frings M. (2010) **Recovery of Upper Limb Function After Cerebellar Stroke** *Stroke* **41**:2191–2200 [Google Scholar](#)

Konczak J., Schoch B., Dimitrova A., Gizewski E., Timmann D. (2005) **Functional recovery of children and adolescents after cerebellar tumour resection** *Brain* **128**:1428–1441 [Google Scholar](#)

Low A. Y. T., Thanawalla A. R., Yip A. K. K., Kim J., Wong K. L. L., Tantra M., Augustine G. J., Chen A. I. (2018) **Precision of Discrete and Rhythmic Forelimb Movements Requires a Distinct Neuronal Subpopulation in the Interposed Anterior Nucleus** *Cell Reports* **22**:2322–2333 [Google Scholar](#)

Luciani L (1893) **Il Cerevelletto. Nuovi Studi di Fisiologia Normale e Patologica** *Philosophical Review* **2**:4 [Google Scholar](#)

Magoun H. W., Hare W. K., Ranson S. W. (1935) **Electrical stimulation of the interior of the cerebellum in the monkey** *American Journal of Physiology-Legacy Content* **112**:329–339 [Google Scholar](#)

Manto M., Bower J., Conforto A., Delgado-García J., Guarda S., Gerwig M., Habas C., Hagura N., Ivry R., Mariën P., Molinari M., Naito E., Nowak D., Taib N., Pélisson D., Tesche C., Tilikete C., Timmann D. (2011) **Consensus Paper: Roles of the Cerebellum in Motor Control-The Diversity of Ideas on Cerebellar Involvement in Movement** *Cerebellum* **11**:457–487 [Google Scholar](#)

Meyer-Lohmann J., Conrad B., Matsunami K., Brooks V. B. (1975) **Effects of dentate cooling on precentral unit activity following torque pulse injections into elbow movements** *Brain Research* **94**:237–251 [Google Scholar](#)

Miall R. C., Christensen L. O. D., Cain O., Stanley J. (2007) **Disruption of State Estimation in the Human Lateral Cerebellum** *PLOS Biology* **5**:e316 [Google Scholar](#)

Miall R. C., King D. (2008) **State Estimation in the Cerebellum** *The Cerebellum* **7**:572–576 [Google Scholar](#)

Naidich T. P., Stein E. G., Lento P. A., Kleinman G. M., Fowkes M. E., Carpenter D. M. (2013) **CHAPTER 16 - Cerebellum** In: Naidich T. P., Castillo M., Cha S., Smirniotopoulos J. G., Saunders

W.B., editors. *Imaging of the Brain* pp. 328–347 [Google Scholar](#)

Nashef A., Cohen O., Harel R., Israel Z., Prut Y. (2019) **Reversible Block of Cerebellar Outflow Reveals Cortical Circuitry for Motor Coordination** *Cell Reports* **27**:2608–2619 [Google Scholar](#)

Nashef A., Cohen O., Israel Z., Harel R., Prut Y. (2018) **Cerebellar Shaping of Motor Cortical Firing Is Correlated with Timing of Motor Actions** *Cell Reports* **23**:1275–1285 [Google Scholar](#)

Nashef A., Rapp H., Nawrot M. P., Prut Y. (2018) **Area-specific processing of cerebellar-thalamo-cortical information in primates** *Biological Cybernetics* **112**:141–152 [Google Scholar](#)

Nashold B. S., Slaughter D. G. (1969) **Effects of Stimulating or Destroying the Deep Cerebellar Regions in Man** *Journal of Neurosurgery* **31**:172–186 [Google Scholar](#)

Nathan P. W., Smith M. C. (1982) **The rubrospinal and central tegmental tracts in man** *Brain : a journal of neurology* **105**:223–269 [Google Scholar](#)

Osu R., Morishige K.-i., Nakanishi J., Miyamoto H., Kawato M. (2015) **Practice reduces task relevant variance modulation and forms nominal trajectory** *Scientific Reports* **5**:17659 [Google Scholar](#)

Padel Y., Angaut P., Massion J., Sedan R. (1981) **Comparative study of the posterior red nucleus in baboons and gibbons** *Journal of Comparative Neurology* **202**:421–438 [Google Scholar](#)

Perlmutter J. S., Mink J. W. (2006) **Deep brain stimulation** *Annual Review of Neuroscience* **29**:229–257 [Google Scholar](#)

Perlmutter S. I., Maier M. A., Fetz E. E. (1998) **Activity of Spinal Interneurons and Their Effects on Forearm Muscles During Voluntary Wrist Movements in the Monkey** *Journal of Neurophysiology* **80**:2475–2494 [Google Scholar](#)

Pi J. S., Fakharian M. A., Hage P., Sedaghat-Nejad E., Muller S. Z., Shadmehr R. (2024) **The olivary input to the cerebellum dissociates sensory events from movement plans** *Proc Natl Acad Sci U S A* **121**:e2318849121 [Google Scholar](#)

Plamondon R., Alimi A. M. (1997) **Speed/accuracy trade-offs in target-directed movements** *Behavioral and Brain Sciences* **20**:279–303 [Google Scholar](#)

Popa L., Hewitt A., Ebner T. (2013) **Purkinje Cell Simple Spike Discharge Encodes Error Signals Consistent with a Forward Internal Model** *Cerebellum* **12** [Google Scholar](#)

Pressman P., Rosen H. J. (2015) **Chapter 33 - Disorders of Frontal Lobe Function** In: Zigmond M. J., Rowland L. P., Coyle J. T., editors. *Neurobiology of Brain Disorders* San Diego: Academic Press pp. 542–557 [Google Scholar](#)

Pruszynski J. A., Kurtzer I., Scott S. H. (2008) **Rapid Motor Responses Are Appropriately Tuned to the Metrics of a Visuospatial Task** *Journal of Neurophysiology* **100**:224–238 [Google Scholar](#)

Pruszynski J. A., Scott S. H. (2012) **Optimal feedback control and the long-latency stretch response** *Exp Brain Res* **218**:341–359 [Google Scholar](#)

Ruigrok T. J. H. (2013) **Cerebellar Influences on Descending Spinal Motor Systems** In: Manto M., Schmahmann J. D., Rossi F., Gruol D. L., Koibuchi N., editors. *Handbook of the Cerebellum and*

Cerebellar Disorders Dordrecht: Springer Netherlands pp. 497–528 [Google Scholar](#)

Sainburg R. L., Ghez C., Kalakanis D. (1999) **Intersegmental Dynamics Are Controlled by Sequential Anticipatory, Error Correction, and Postural Mechanisms** *Journal of Neurophysiology* **81**:1045–1056 [Google Scholar](#)

Sathyamurthy A., Barik A., Dobrott C. I., Matson K. J. E., Stoica S., Pursley R., Chesler A. T., Levine A. J. (2020) **Cerebellospinal Neurons Regulate Motor Performance and Motor Learning** *Cell Reports* **31**:107595 [Google Scholar](#)

Schlerf J. E., Xu J., Klemfuss N. M., Griffiths T. L., Ivry R. B. (2012) **Individuals with cerebellar degeneration show similar adaptation deficits with large and small visuomotor errors** *Journal of Neurophysiology* **109**:1164–1173 [Google Scholar](#)

Schneider R. C., Crosby E. C. (1963) **The Interplay Between Cerebral Hemispheres and Cerebellum in Relation to Tonus and Movements** *Journal of Neurosurgery* **20**:188–198 [Google Scholar](#)

Scott S. H (2004) **Optimal feedback control and the neural basis of volitional motor control** *Nature Reviews Neuroscience* **5**:532–545 [Google Scholar](#)

Shadmehr R., Mussa-Ivaldi F. (1994) **Adaptive representation of dynamics during learning of a motor task** *The Journal of Neuroscience* **14**:3208–3224 [Google Scholar](#)

Shidara M., Kawano K., Gomi H., Kawato M. (1993) **Inverse-dynamics model eye movement control by Purkinje cells in the cerebellum** *Nature* **365**:50–52 [Google Scholar](#)

Shinoda Y., Yamazaki M., Futami T. (1982) **Convergent inputs from the dentate and the interpositus nuclei to pyramidal tract neurons in the motor cortex** *Neuroscience Letters* **34**:111–115 [Google Scholar](#)

Sinha N., Israely S., Ben Harosh O., Harel R., Dewald J. P. A., Prut Y. (2024) **Disentangling acute motor deficits and adaptive responses evoked by the loss of cerebellar output** *GitHub*

Smith J. L., Zernicke R. F. (1987) **Predictions for neural control based on limb dynamics** *Trends in Neurosciences* **10**:123–128 [Google Scholar](#)

ten Donkelaar H. J (1988) **Evolution of the red nucleus and rubrospinal tract** *Behavioural Brain Research* **28**:9–20 [Google Scholar](#)

Therrien A. S., Bastian A. J. (2015) **Cerebellar damage impairs internal predictions for sensory and motor function** *Current Opinion in Neurobiology* **33**:127–133 [Google Scholar](#)

Timmann D., Watts S., Hore J. (1999) **Failure of Cerebellar Patients to Time Finger Opening Precisely Causes Ball High-Low Inaccuracy in Overarm Throws** *Journal of Neurophysiology* **82**:103–114 [Google Scholar](#)

Todorov E., Jordan M. I. (2002) **Optimal feedback control as a theory of motor coordination** *Nature Neuroscience* **5**:1226–1235 [Google Scholar](#)

- Topka H., Konczak J., Dichgans J. (1998) **Coordination of multi-joint arm movements in cerebellar ataxia: Analysis of hand and angular kinematics** *Experimental Brain Research* **119**:483–492 [Google Scholar](#)
- Topka H., Konczak J., Schneider K., Boose A., Dichgans J. (1998) **Multijoint arm movements in cerebellar ataxia: Abnormal control of movement dynamics** *Experimental Brain Research* **119**:493–503 [Google Scholar](#)
- Tsujimoto T., Gemba H., Sasaki K. (1993) **Effect of cooling the dentate nucleus of the cerebellum on hand movement of the monkey** *Brain Research* **629**:1–9 [Google Scholar](#)
- Virji-Babul N., Cooke J. D. (1995) **Influence of joint interactional effects on the coordination of planar two-joint arm movements** *Experimental Brain Research* **103**:451–459 [Google Scholar](#)
- Weeks H. M., Therrien A. S., Bastian A. J. (2017) **Proprioceptive Localization Deficits in People With Cerebellar Damage** *The Cerebellum* **16**:427–437 [Google Scholar](#)
- Wild B., Dichgans J. (1993) **Cerebellar ataxia in ataxic hemiparesis? A kinematic and EMG analysis** *Movement Disorders* **8**:363–366 [Google Scholar](#)
- Wild B., Klockgether T., Dichgans J. (1996) **Acceleration deficit in patients with cerebellar lesions. A study of kinematic and EMG-parameters in fast wrist movements** *Brain Research* **713**:186–191 [Google Scholar](#)
- Wolpert D. M., Ghahramani Z., Jordan M. I. (1995) **An Internal Model for Sensorimotor Integration** *Science* **269**:1880–1882 [Google Scholar](#)
- Wolpert D. M., Miall R. C., Kawato M. (1998) **Internal models in the cerebellum** *Trends in Cognitive Sciences* **2**:338–347 [Google Scholar](#)

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Reviewer #1 (Public review):

Summary:

In a previous work Prut and colleagues had shown that during reaching, high frequency stimulation of the cerebellar outputs resulted in reduced reach velocity. Moreover, they showed that the stimulation produced reaches that deviated from a straight line, with the shoulder and elbow movements becoming less coordinated. In this report they extend their previous work by addition of modeling results that investigate the relationship between the kinematic changes and torques produced at the joints. The results show that the slowing is not due to reductions in interaction torques alone, as the reductions in velocity occur even for movements that are single joint. More interestingly, the experiment revealed evidence for decomposition of the reaching movement, as well as an increase in the variance of the trajectory.

Strengths:

This is a rare experiment in a non-human primate that assessed the importance of cerebellar input to the motor cortex during reaching.

Weaknesses:

None

Reviewer #2 (Public review):

This manuscript asks an interesting and important question: what part of 'cerebellar' motor dysfunction is an acute control problem vs a compensatory strategy to the acute control issue? The authors use a cerebellar 'blockade' protocol, consisting of high frequency stimuli applied to the cerebellar peduncle which is thought to interfere with outflow signals. This protocol was applied in monkeys performing center out reaching movements and has been published from this laboratory in several preceding studies. I found the take-home-message broadly convincing and clarifying - that cerebellar block reduces muscle activation acutely particularly in movements that involve multiple joints and therefore invoke interaction torques, and that movements progressively slow down to in effect 'compensate' for these acute tone deficits. The manuscript was generally well written, data were clear, convincing and novel. The key strengths are differentiating acute from sub-acute (within session but not immediate) kinematic consequences of cerebellar block.

<https://doi.org/10.7554/eLife.105152.3.sa2>

Reviewer #3 (Public review):**Summary:**

In their revised manuscript, Sinha and colleagues aim to identify distinct causes of motor impairments seen when perturbing cerebellar circuits. This goal is an important one, given the diversity of movement related phenotypes in patients with cerebellar lesion or injury, which are especially difficult to dissect given the chronic nature of the circuit damage. To address this goal, the authors use high-frequency stimulation (HFS) of the superior cerebellar peduncle in monkeys performing reaching movements. HFS provides an attractive approach for transiently disrupting cerebellar function previously published by this group. First, they find a reduction in hand velocities during reaching, which was more pronounced for outward versus inward movements. By modeling inverse dynamics, they find evidence that shoulder muscle torques are especially affected. Next, the authors examine the temporal evolution of movement phenotypes over successive blocks of HFS trials. Using this analysis, they find that in addition to the acute, specific effects on torques in early HFS trials, there was an additional progressive reduction in velocity during later trials, which they interpret as an adaptive response to the inability to effectively compensate for interaction torques during cerebellar block. Finally, the authors examine movement decomposition and trajectory, finding that even when low velocity reaches are matched to controls, HFS produces abnormally decomposed movements and higher than expected variability in trajectory.

Strengths:

Overall, this work provides important insight into how perturbation of cerebellar circuits can elicit diverse effects on movement across multiple timescales.

The HFS approach provides temporal resolution and enables analysis that would be hard to perform in the context of chronic lesions or slow pharmacological interventions. Thus, this study describes an important advance over prior methods of circuit disruption in the monkey, and their approach can be used as a framework for future studies that delve deeper into how additional aspects of sensorimotor control are disrupted (e.g., response to limb perturbations).

In addition, the authors use well-designed behavioral approaches and analysis methods to distinguish immediate from longer-term adaptive effects of HFS on behavior. Moreover, inverse dynamics modeling provides important insight into how movements with different kinematics and muscle dynamics might be differentially disrupted by cerebellar perturbation.

Remaining comments:

The argument that there are acute and adaptive effects to perturbing cerebellar circuits is compelling, but there seems to be a lost opportunity to leverage the fast and reversible nature of the perturbations to further test this idea and strengthen the interpretation. Specifically, the authors could have bolstered this argument by looking at the effects of terminating HFS - one might hypothesize that the acute impacts on joint torques would quickly return to baseline in the absence of HFS, whereas the longer-term adaptive component would persist in the form of aftereffects during the 'washout' period. As is, the reversible nature of the perturbation seems underutilized in testing the authors' ideas. While this experimental design was not implemented here, it seems like a good opportunity for future work using these approaches.

The analysis showing that there is a gradual reduction in velocity during what the authors call an adaptive phase is convincing. While it is still not entirely clear why disruption of movement during the adaptive phase is not seen for inward targets, despite the fact that many of the inward movements also exhibit large interaction torques, the authors do raise potential explanations in the Discussion.

<https://doi.org/10.7554/eLife.105152.3.sa1>

Author response:

The following is the authors' response to the previous reviews

Reviewer #1 (Public review):

Summary:

In a previous work Prut and colleagues had shown that during reaching, high frequency stimulation of the cerebellar outputs resulted in reduced reach velocity. Moreover, they showed that the stimulation produced reaches that deviated from a straight line, with the shoulder and elbow movements becoming less coordinated. In this report they extend their previous work by addition of modeling results that investigate the relationship between the kinematic changes and torques produced at the joints. The results show that the slowing is not due to reductions in interaction torques alone, as the reductions in velocity occur even for movements that are single joint. More interestingly, the experiment revealed evidence for decomposition of the reaching movement, as well as an increase in the variance of the trajectory.

Strengths:

This is a rare experiment in a non-human primate that assessed the importance of cerebellar input to the motor cortex during reaching.

Weaknesses:

None

Reviewer #1 (Recommendations for the authors):

The authors have answered my questions adequately and I have no further comments.

Reviewer #2 (Public review):

This manuscript asks an interesting and important question: what part of 'cerebellar' motor dysfunction is an acute control problem vs a compensatory strategy to the acute control issue? The authors use a cerebellar 'blockade' protocol, consisting of high frequency stimuli applied to the cerebellar peduncle which is thought to interfere with outflow signals. This protocol was applied in monkeys performing center out reaching movements and has been published from this laboratory in several preceding studies. I found the takehome-message broadly convincing and clarifying - that cerebellar block reduces muscle activation acutely particularly in movements that involve multiple joints and therefore invoke interaction torques, and that movements progressively slow down to in effect 'compensate' for these acute tone deficits. The manuscript was generally well written, data were clear, convincing and novel. The key strengths are differentiating acute from subacute (within session but not immediate) kinematic consequences of cerebellar block.

Reviewer #2 (Recommendations for the authors):

I think the manuscript is good as is. That said, it would have been nice to see more of the behavioral outcomes in Figure 5 (e.g. decomposition and trajectory variability) analyzed longitudinally like the velocity measurements in Fig. 4. This would clearly strengthen the insight into acute and compensatory components of cerebellar motor deficits.

The two behavioral measures of motor noise used in our study are movement decomposition and trajectory variability (Figure 5). Since trajectory variability is measured across trials we could not analyze this measure longitudinally as a function of trial number. However, following the reviewer's advice, we examined movement

decomposition for successive trials in control vs. cerebellar block for movements to targets 2-4 similar to the analysis of hand velocity in figure 4. We found no interaction effect between trial sequence x cerebellar block on movement decomposition. This result is consistent with our conclusion that noisy joint activation occurs independently of adaptive slowing of multi-joint movements. We have updated our main text (lines 293-299) and supplementary information (supplementary figure S5 and supplementary table S8) to include this result.

Reviewer #3 (Public review):

Summary:

In their revised manuscript, Sinha and colleagues aim to identify distinct causes of motor impairments seen when perturbing cerebellar circuits. This goal is an important one, given the diversity of movement related phenotypes in patients with cerebellar lesion or injury, which are especially difficult to dissect given the chronic nature of the circuit damage. To address this goal, the authors use high-frequency stimulation (HFS) of the superior cerebellar peduncle in monkeys performing reaching movements. HFS provides an attractive approach for transiently disrupting cerebellar function previously published by this group. First, they find a reduction in hand velocities during reaching, which was more pronounced for outward versus inward movements. By modeling inverse dynamics, they find evidence that shoulder muscle torques are especially affected. Next, the authors examine the temporal evolution of movement phenotypes over successive blocks of HFS trials. Using this analysis, they find that in addition to the acute, specific effects on torques in early HFS trials, there was an additional progressive reduction in velocity

during later trials, which they interpret as an adaptive response to the inability to effectively compensate for interaction torques during cerebellar block. Finally, the authors examine movement decomposition and trajectory, finding that even when low velocity reaches are matched to controls, HFS produces abnormally decomposed movements and higher than expected variability in trajectory.

Strengths:

Overall, this work provides important insight into how perturbation of cerebellar circuits can elicit diverse effects on movement across multiple timescales.

The HFS approach provides temporal resolution and enables analysis that would be hard to perform in the context of chronic lesions or slow pharmacological interventions. Thus, this study describes an important advance over prior methods of circuit disruption in the monkey, and their approach can be used as a framework for future studies that delve deeper into how additional aspects of sensorimotor control are disrupted (e.g., response to limb perturbations).

In addition, the authors use well-designed behavioral approaches and analysis methods to distinguish immediate from longer-term adaptive effects of HFS on behavior. Moreover, inverse dynamics modeling provides important insight into how movements with different kinematics and muscle dynamics might be differentially disrupted by cerebellar perturbation.

In this revised version of the manuscript, the authors have provided additional analyses and clarification that address several of the comments from the original submission.

Remaining comments:

The argument that there are acute and adaptive effects to perturbing cerebellar circuits is compelling, but there seems to be a lost opportunity to leverage the fast and reversible nature of the perturbations to further test this idea and strengthen the interpretation. Specifically, the authors could have bolstered this argument by looking at the effects of terminating HFS - one might hypothesize that the acute impacts on joint torques would quickly return to baseline in the absence of HFS, whereas the longer-term adaptive component would persist in the form of aftereffects during the 'washout' period. As is, the reversible nature of the perturbation seems underutilized in testing the authors' ideas. While this experimental design was not implemented here, it seems like a good opportunity for future work using these approaches.

We agree with the reviewer that examining the effect of the cerebellar block on immediate post-block washout trials in future studies will be insightful.

The analysis showing that there is a gradual reduction in velocity during what the authors call an adaptive phase is convincing. While it is still not entirely clear why disruption of movement during the adaptive phase is not seen for inward targets, despite the fact that many of the inward movements also exhibit large interaction torques, the authors do raise potential explanations in the Discussion.

The text in the Introduction and in the prior work developing the HFS approach overstates the selectivity of the perturbations. First, there is an emphasis on signals transmitted to the neocortex. As the authors state several times in the Discussion, there are many subcortical targets of the cerebellar nuclei as well, and thus it is difficult to disentangle target-specific behavioral effects using this approach. Second, the superior cerebellar peduncle contains both cerebellar outputs and inputs (e.g., spinocerebellar). Therefore, the selectivity in perturbing cerebellar output feels overstated. Readers would benefit from a more agnostic claim that HFS affects cerebellar communication with the

rest of the nervous system, which would not affect the major findings of the study. In the revised manuscript, the authors do provide additional anatomical and evolutionary context and discuss potential limitations in the selectivity of HFS in the Materials and Methods. However, I feel that at least a brief mention of these caveats in the Introduction, where it is stated, "we then reversibly blocked cerebellar output to the motor cortex", would benefit the reader.

Following the advice of the reviewer, we have now revised the introduction section of our manuscript in the following way (lines 61-67):

“...We then reversibly disrupted cerebellar communication with other neural structures using high-frequency stimulation (HFS) of the superior cerebellar peduncle, assessing the impact of this perturbation on subsequent movements. Although our approach primarily affects cerebellar output to the motor cortex, it also disrupts fibers carrying input signals (e.g., spinocerebellar) and pathways to various subcortical targets (e.g., cerebellorubrospinal). Thus, our manipulation broadly interferes with cerebellar communication...”

Reviewer #3 (Recommendations for the authors):

Typo on line 102; "subs-sessions"

We have corrected this typographical error in our revised manuscript (line 106).

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