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Stronger proprioceptive BOLD-responses in the somatosensory cortices reflect worse sensorimotor function in adolescents with and without cerebral palsy

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\textbf{A B S T R A C T}

Cerebral palsy (CP) is a motor disorder where the motor defects are partly due to impaired proprioception. We studied cortical proprioceptive responses and sensorimotor performance in adolescents with CP and their typically-developed (TD) peers. Passive joint movements were used to stimulate proprioceptors during functional magnetic resonance imaging (fMRI) session to quantify the proprioceptive responses whose associations to behavioral sensorimotor performance were also examined.

Twenty-three TD (15 females, age: mean ± standard deviation 13.8 ± 2.3 years) and 18 CP (12 females, age: mean ± standard deviation, 13.8 ± 2.3 years; 12 hemiplegic, 6 diplegic) participants were included in this study. Participants’ index fingers and ankles were separately stimulated at 3 Hz and 1 Hz respectively with pneumatic movement actuators. Regions-of-interest were used to quantify BOLD-responses from the primary somatosensory (SM1) and secondary (SII) somatosensory cortices and were compared across the groups. Associations between responses strengths and sensorimotor performance measures were also examined.

Proprioceptive responses were stronger for the individuals with CP compared to their TD peers in SM1 (p < 0.001) and SII (p < 0.05) cortices contralateral to their more affected index finger. The ankle responses yielded no significant differences between the groups. The CP group had worse sensorimotor performance for hands and feet (p < 0.001). Stronger responses to finger stimulation in the dominant SM1 (p < 0.001) and both dominant and non-dominant SII (p < 0.01, p < 0.001) cortices were associated with the worse hand sensorimotor performance across all participants.

Worse hand function was associated with stronger cortical activation to the proprioceptive stimulation. This association was evident both in adolescents with CP and their typically-developed controls, thus it likely reflects both clinical factors and normal variation in the sensorimotor function. The specific mechanisms need to be clarified in future studies.

1. Introduction

Cerebral palsy (CP) is a group of permanent disorders in the development of movement and posture that are attributed to non-progressive disturbances in the developing brain (Rosenbaum et al., 2007; Mac Keith et al., 1958). The disorder is often accompanied by disturbances of sensation, perception, cognition, communication, and behavior in addition to co-occurring epilepsy and secondary musculoskeletal problems. The disorder has also relatively high prevalence of 2–2.5 out of 1000 births (Hagberg et al., 2001; Ashwal et al., 2004). Thus, better

\textbf{Abbreviations:} fMRI, functional magnetic resonance imaging; CP, cerebral palsy; HP, hemiplegic; DP, diplegic; BOLD, Blood-Oxygen-Level-Dependent signal; ROI, regions of interest; SI cortex, primary somatosensory cortex; SII cortex, secondary somatosensory cortex; TD, typically-developed; GMFCS, Gross Motor Function Classification System; TR, repetition time; TE, echo time; EPI, echo planar imaging; SIFT, Sensory Integration and Praxis Tests; GLM, General Linear Model; SPM, Statistical Parametric Mapping; EM, expectation maximization; MANCOVA, Multivariate analysis of covariance; PSC, percent signal change.

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understanding of the neurophysiological effects of the disorder and the related specific neuronal mechanisms are important for enhancing the effectiveness and targeting of rehabilitation and treatments in CP. CP can be classified by topography, for instance, as hemiplegic (one side affected more) or diplegic (both sides are affected). Classification systems of motor function such as 5-point-scale Gross Motor Function Classification System (GMFCS; Palisano et al., 1997) are also widely used.

Smooth motor function depends largely on somatosensory afference from the body. More specifically, continuous somatosensory (primarily proprioceptive) feedback about the state of the locomotor system to the brain supports central nervous system motor circuits in controlling and adjusting on-going movements. Somatosensory afference has therefore a key role in motor learning and development of gross (e.g. gait) and fine (e.g. writing) motor skills (Metcalfe et al., 2005; Soechting and Flanders, 2008; Cascio, 2010). Proprioceptive afference provides information about the limb and body position, movement and forces (Frosoke and Gandevia, 2012). Proprioception is known to be impaired in CP (Goble et al., 2009; Wingert et al., 2009). This suggests that impaired proprioceptive afference has a role in the impaired motor function in CP.

Proprioceptive afference is processed widely in the neocortex, but majorly of the fast direct thalamocortical cortical input is directed to the primary somatosensory (SI) cortex. In addition to the SI cortex, the secondary somatosensory (SII) cortex is important in processing the somatosensory afference. SI cortex seems to process early, lower-level stimulus features whereas SII cortex appears to integrate multimodal sensory information in a more bilateral manner (Inoue et al., 2002; Eichhoff et al., 2007). The primary motor (M1) cortex also receives direct, short-latency input from the proprioceptors and cortico-cortical input from the somatosensory cortices (Goldring and Ratcheson, 1972; Rosén and Asanuma,1972; Asanuma and Arissian, 1984; Brovelli et al., 2004; Kocak et al., 2009). SI and M1 cortices are therefore often studied as a single functional unit, the primary sensorimotor (SM1) cortex and together with SII cortex form the most central part of the proprioceptive system when studying cortical proprioception.

Cortical proprioception can be studied using passive naturalistic movements of specific joints using different brain imaging methods (Müller-Putz et al., 2007; Sasaki et al., 2017, 2018; Alary et al., 2002; Druschky et al., 2003; Lange et al., 2001; Lolli et al., 2019; Nurmi et al., 2018; Piitulainen et al., 2015, 2018; 2021; Seiss et al., 2002; Vallinioja et al., 2021). For the continuous index finger movement, strongest BOLD-fMRI responses have been shown to be elicited by 3–6 Hz movements when using a blocked design (Nurmi et al., 2018). Cortical somatotopy and somatosensory processing are altered in CP patients evidenced by MEG and fMRI (Nevalainen et al., 2012; Papadelis et al., 2014, 2018; Riquelme et al., 2010; 2014). These cortical alterations appear to have functional significance, as they have been associated to worse motor performance in several motor disorders (Rusovska et al., 1982; Nociti et al., 2008).

Our primary aim was to clarify whether proprioceptive responses in the sensorimotor cortices of the fingers and ankles differ between adolescents with CP and their typically-developed (TD) peers. Secondly, we aimed to examine whether sensorimotor performance of the hand (motor skill) or foot (balance) were associated with the respective magnitudes of the cortical proprioceptive responses. Therapies and treatments in CP are most effective when started as early age as possible. For this reason, adolescents were chosen as participants. Our exploratory results will potentially offer basis for formulating new hypotheses in clinical research of CP.

2. Materials and methods

2.1. Participants

Controls. In total, 35 TD healthy controls were recruited for the study. Three of them quit prior or during the fMRI recording due to discomfort in the scanner or fear entering to it. In addition, nine participants were excluded from the fMRI analysis due to insufficient data quality such as absence of the activations and/or clear movement artifacts (2 participants), head movement exceeding 6 mm (5 participants), being on migraine medication during the measurement (1 participant) or having tic-symptoms (1 participant).

The remaining 23 healthy controls (15 females, age: mean ± standard deviation, 14.2 ± 2.4 years) were included in the final analysis. Majority of the TD controls were right-handed (21 out of 23, Edinburgh Handedness Inventory mean score: 67.8; range: −87–100) and right-footed (20 out of 23, mean: 50.0; range: −60–100). The TD group was intellectually within normal variation (Wechsler Adult Intelligence Scale / Wechsler Intelligence Scale for Children mean score: 107.6, SD: 15.2; range: 77–132; three participants missing a test score). See Table 1 for TD participant demographics.

Patients. In total 31 participants with CP (GMFCS classification of 1–2) were recruited to the study. Six of them quit due to discomfort in the scanner or fear entering to it, two were excluded due to absence of activations and/or clear movement artifacts, and five of them were excluded due to head movement exceeding 6 mm.

The remaining 18 participants were included in the final analysis (12 hemiplegic and 6 diplegic participants; 12 females, age: mean ± standard deviation, 13.8 ± 2.3 years). Seven CP participants were right-

| Participant demographics. |
| Group / CP type | Dominant side (Hand) | Dominant side (Foot) | Sex | Age (years) |
|-------------------|----------------------|----------------------|-----|-------------|
| 1 TD Right Right Male 13 |
| 2 TD Right Right Male 12 |
| 3 TD Right Right Female 12 |
| 4 TD Right Right Female 15 |
| 5 TD Right Right Female 16 |
| 6 TD Right Right Female 12 |
| 7 TD Right Right Female 12 |
| 8 TD Right Left Male 17 |
| 9 TD Right Right Female 17 |
| 10 TD Left Left Male 17 |
| 11 TD Right Right Female 17 |
| 12 TD Right Right Male 13 |
| 13 TD Right Right Female 16 |
| 14 TD Right Right Male 13 |
| 15 TD Right Right Female 14 |
| 16 TD Right Right Female 14 |
| 17 TD Right Right Male 11 |
| 18 TD Left Left Female 12 |
| 19 TD Right Right Female 10 |
| 20 TD Right Right Male 12 |
| 21 TD Right Right Female 10 |
| 22 TD Right Right Female 16 |
| 23 TD Right Right Female 16 |
| 24 CP/HP Left Left Female 17 |
| 25 CP/HP Left Left Female 10 |
| 26 CP/HP Left Left Male 13 |
| 27 CP/HP Left Left Female 11 |
| 28 CP/HP Right Right Female 15 |
| 29 CP/HP Left Left Male 10 |
| 30 CP/HP Right Right Female 16 |
| 31 CP/HP Left Left Female 15 |
| 32 CP/HP Left Left Female 17 |
| 33 CP/HP Right Right Female 13 |
| 34 CP/HP Left Left Female 10 |
| 35 CP/HP Left Left Female 14 |
| 36 CP/DP Right Right Male 10 |
| 37 CP/DP Left Left Male 12 |
| 38 CP/DP Left Left Male 11 |
| 39 CP/DP Right Right Female 15 |
| 40 CP/DP Right Right Female 15 |
| 41 CP/DP Right Right Male 15 |
handed (three hemiplegic and four diplegic) and 11 were left-handed (mean: –16.7, range: –100–100) and seven of them were right-footed (three hemiplegic and four diplegic; mean: –20.2; range: –100–89). For the hemiplegic participants, handedness and footedness were congruent with the clinical definition of their more affected side. The CP participants were cognitively mostly within normal variation (Wechsler Adult Intelligence Scale / Wechsler Intelligence Scale for Children mean score: 90, SD: 20.7, range: 43–117; five participants missing a test score).

The study was conducted according to declaration of Helsinki. All participants were 10–18 years of age. All participants gave their written consent, and in the case of minors, also their custodian gave a written consent. The protocol was approved by the ethics committee of Aalto University and Hospital District of Helsinki and Uusimaa (HUS). See Table 1 for detailed CP participant demographics and Table 2 for their lesion information.

2.2. Handedness and footedness

The hand and foot dominance were defined according to the Edinburgh Handedness Inventory (Oldfield 1971) and Waterloo Footedness Questionnaire Revised (Elias et al., 1998) test scores for healthy controls and diplegia patients. For the hemiplegia participants, the contralateral side was always defined as the dominant side (i.e. the functionally less affected side; the less affected side was always also the dominant side according to the handedness and footedness tests).

2.3. MRI equipment and parameters

Structural and functional imaging were carried out using a 3 T-MATRIGEM Skyra whole-body scanner (Siemens Healthcare, Erlangen, Germany). A 32-channel head coil was used. The measurements were carried out in Advanced Magnetic Imaging (AMI) Centre of Aalto Neuroimaging infrastructure in Aalto University, Espoo, Finland.

An anatomical T1 (MPRAGE) was obtained with 176 slices with slice thickness of 1 mm without an interslice gap and a 256x256 matrix with a field-of-view (FOV) of 256x256 mm, yielding a voxel size of 1x1x1 mm. Orientation of the structural image was sagittal. Repetition time (TR) was 2.53 s and echo time (TE) was 3.33 ms. Flip angle was 7°.

The functional images were obtained using a standard ECHO-Planar Imaging (EPI) spin-echo sequence with a TR of 2.5 s and a TE of 30 ms. The functional volumes consisted of 44 slices with slice thickness of 3 mm without an interslice gap whose FOV was 192x192 mm with a 64x64 matrix yielding a voxel size of 3x3x3 mm. Flip angle was 90°.

2.4. Proprioceptive stimulation in fMRI

Four custom-made pneumatic movement actuators were used to evoke flexion-extension movements of the right and left index finger (for a detailed description of the actuators, see Nurmi et al. 2018; Pitulainen et al., 2015) and sagittal plane rotations of right and left ankle joints. These proprioceptive stimulators consisted of a plastic frame and an artificial pneumatic muscle (Fig. 1 a, b). The fingertips were attached to the pneumatic muscle with a surgical tape, and the feet were attached to the footrests using elastic straps. Surgical tape was also wrapped around participants’ index fingers to minimize tactile activation during stimulation. Participants’ hands and distal part of the arms rested on the upper plate of the finger-movement-actuator.

The artificial muscle moved along its longitudinal axis when its internal air pressure (1–5 bar) changed. The pressure was regulated by a solenoid valve (SYS220-6LOU-01F-Q, SMC Corporation, Tokyo, Japan) that was controlled by computer-generated trigger pulses. The solenoid valve was placed outside the MRI room and a 3.5 m non-elastic tube (internal diameter 2.5 mm) conveyed the airflow to the artificial muscle. The artificial muscle shortened in response to the increasing air pressure (opening of the valve), thereby flexing the finger or dorsiflexing the ankle joint, and then returned back to the initial position when the air pressure was released (closing of the valve).

For the index fingers, continuous 3-Hz movement was used. This movement frequency was found to be efficient in eliciting BOLD-responses in the somatosensory cortices (Nurmi et al., 2018). Ankle joint rotations were evoked at lower 1-Hz frequency, which lies well

Table 2

| CP type | Lesion type | Side of lesion | Lesion location (MNI coordinates) | Lesion size (mm³) | MRI description |
|--------|-------------|----------------|----------------------------------|------------------|-----------------|
| 24 HP  | infarction  | Left           | −54, −30, 57                     | 17,573           | Large lesion left parietal lobe |
| 25 HP  | infarction  | Left           | −62, 8, 17                       | 19,470           | Large lesion left middle frontal lobe |
| 26 HP  | infarction  | Left           | −63, −46, 31                     | 67,544           | Very large lesion covering left parietal and parts of superior temporal and posterior frontal lobes |
| 27 HP  | PVL         | Left/Right     | −14, −16, 26                     | 1826 + 444       | Slightly enlarged posterior left lateral ventricle; small lesion in front of right anterior lateral ventricle |
| 28 HP  | PVH/infarction, HBI, PVL | Right | 14, 2, 23                        | 1295             | Slightly enlarged posterior right ventricle |
| 29 HP  | IHI, IVH, WMI, PVL, WD | Left | −18, −5, 42                      | 37,054           | Extensively enlarged left ventricle |
| 30 HP  | infarction  | Right          | −10, −5, 22                      | 6654             | Slight enlargement of right lateral ventricle |
| 31 HP  | unknown     | Left           | −15, −20, 25                     | 2764             | Slight enlarged left lateral ventricle |
| 32 HP  | IHI, mild PVL | Right | −16, −23, 25                     | 1006             | Slightly enlarged left lateral ventricle |
| 33 HP  | infarction  | Right          | 40, −10, 17                      | 65,833           | Very large lesion in anterior-posterior axel in the lateral regions covering frontal, temporal and parietal regions and making contact with an enlarged ventricle |
| 34 HP  | infarction  | Left           | −11, −11, 22                     | 18,613           | Large enlargement of left ventricle |
| 35 HP  | perinatal injury | Left | −26, −58, 3                      | 675              | Slightly enlarged ventricle |
| 36 DP  | PVL, local ischemia | None | (no visible lesion) | 0                | No visible lesion |
| 37 DP  | HBI, PVL    | Left           | −24, −67, 4                      | 381              | Very slight enlargement of left inferior lateral ventricle |
| 38 DP  | PVL         | None (no visible lesion) | 0               | No visible lesion |
| 39 DP  | unknown, normal MRI | None (no visible lesion) | 0                | No visible lesion |
| 40 DP  | unknown     | Right          | 15, −10, 25                      | 2566             | Slight enlargement of right lateral ventricle |
| 41 DP  | unknown     | Right          | 28, 24, 53                       | 336              | Very small lesion in right anterior frontal lobe |

CP participants’ lesion information (type, side, location, size, MRI description). Lesion type was diagnosed by a clinician whereas MRI description was evaluated by a researcher (author of this article). Abbreviations: PVL, periventricular leukomalacia; PVH, periventricular hemorrhage; HBI, Hypoxic-ischemic brain injury; IHI, local hypoxia–ischemia; WMI, white matter injury; WD Wallerian degeneration.
within the range of physiological foot actions and produces negligible head movement.

The stimulus runs with passive movement consisted of 20-s stimulation blocks with movement of one limb at a time, alternating with 20-s rest blocks without movement. Thus, the experiment consisted of four different experimental conditions: movement of (1) right index finger, (2) left index finger, (3) right ankle and (4) left ankle.

The stimulus runs with passive movement consisted of block lists (see Fig. 1e for an example of the first run; each row represents a block list in the figure). These block lists were created in order to present different conditions in semi-random order while still ensuring that the same condition is not presented several times in a row but different conditions are presented with similar incidence in every part of the stimulus run. A block list was defined to consist of alternating stimulation and rest blocks. Each of the experimental conditions in a stimulation block was presented in random order exactly once per a block list. When a block list is finished, another block list follows where the order of the presented stimulation blocks was again re-randomized. The first run consisted of four block lists and the second run consisted of eight block lists yielding 12x20-s stimulus blocks per condition in total, i.e., 4 min of stimulation per condition.

2.5. MRI protocol

During the (f)MRI scan, participants were laying on the MRI table with their limbs attached to the movement actuators (Fig. 1a). A pillow was placed under the participants' calf and hamstrings. Two pillows were placed under the participant’s triceps, shoulders and elbows to give support for the arms. Respiration belt and pulse oximeter were attached to participant’s chest and middle or ring finger, and the respective signals were recorded throughout the fMRI scans using BIOPAC MP150 system (BIOPAC Systems Inc., Goleta, USA).

The MRI session consisted of four parts. First, a one-minute head-localizer scan was performed followed by a six-minute T1 anatomical scan. Then, 11 min long stimulus run was conducted. Second stimulus run lasted 22 min. The whole MRI session lasted 45–50 min. During the stimulus runs, a video of slowly changing pictures was shown to help the participant to remain alert.

2.6. Sensorimotor performance

Hand. Fig. 1c illustrates the sensorimotor tests. (1) A two-point discrimination test (range: 2–15 mm apart) to measure tactile discrimination ability of the hands in millimeters. Each digit in both hands was placed under the participant’s chest and middle or ring finger, and the respective signals were recorded throughout the fMRI scans using BIOPAC MP150 system (BIOPAC Systems Inc., Goleta, USA).

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quantify unilateral gross-manual dexterity as number of blocks moved from one box to another in 60 s. (5) Nine-Hole Peg test (see Mathiowetz et al., 1985) to quantify the unilateral fine-motor dexterity as the time (s) to place 9 pegs into 9 holes and back to the container. (6) Stereognosis test from the Sensory Integration and Praxis Test (SIPT; WPS, Torrance, California, US) test pattern to quantify unilateral shape recognition perception test of the hand measured by the number of shapes recognized correctly and the mean time (s) it took to recognize those shapes (two separate variables). (7) Bimanual bottle-opening test to quantify the ability to perform a bimanual task classified into four different performance levels based on the degree of involvement by the affected hand. (8) Unilateral kinesthetic accuracy test from the SIPT where the participant had to repeat finger position from initial position to end position on table surface measured as the distance. The movement was first guided by researcher and was then repeated by the participant. Five different directions and distances for each hand were performed and error was calculated in centimeters (average of all trials was used).

Foot. Fig. 1d illustrates tests for the lower limb (foot) bilateral sensorimotor performance. Standing postural stability was quantified using four standing balance tasks. The participant stood as still as possible on a platform recording plantar pressure distribution (0.5 m Hi-End Footscan® system, RSscan international, Brussels, Belgium) for 30 s per task. The four tasks were: (1) standing eyes open and feet shoulder width apart, (2) standing eyes closed and feet shoulder width apart, (3) standing with eyes open and feet together and (4) standing with eyes closed and feet together. Unit of measurement was center-of-force velocity (mm/s). Dynamic stability (standard deviation of step duration) was assessed using inertial measurement unit (NGIMU, x-io Technologies Limited, Bristol, UK) recordings during three walking tasks (for details, see Piitulainen et al., 2020a) (5) normal walking, (6) carrying a tray with a cup on it and (7) listing words in a given category during walking. Mean standard deviation in step duration was used (s).

Sum variables of sensorimotor performance. Sum variables were constructed for hand and foot sensorimotor performance separately. Each measure was first normalized across the whole studied population to scale from 0 (value of the worst performance) to 1 (value of the best performance). Then, the test values were averaged for each participant to obtain the sum variable. In case of unilateral tests, the left and right hand/foot measures were pooled together. The internal consistency of the sum variables items were estimated using Cronbach’s alphas and ensuring the values were 0.7 or above, indicating acceptable internal consistency (Cronbach, 1951; Nunnally, 1978).

Comparison of sensorimotor performance between groups. All of the single sensorimotor test values were compared between the CP and TD groups using MANOVA (hand and foot tests were tested separately) and post-hoc tests. Note that no imputation was used for these values.

Likewise, the overall sensorimotor performance of hands and feet were compared between the CP and TD groups using MANOVA (hand and foot tests were tested separately) and post-hoc tests. Imputation was used for these the overall sensorimotor performance values.

2.7. MRI preprocessing

The (f)MRI data of each participant was preprocessed using Matlab (R2016b, Mathworks, Natick, Massachusetts, United States) with a custom script using SPM12 functions (Wellcome Department of Imaging Neuroscience, University College London, UK). At first, both the structural and functional volumes were converted from Dicom to Nifti format.

The functional volumes were slice-time-corrected, motion-corrected by realigning to the last functional volume, functional volumes co-registered to anatomical volume. A tool called Drifter (Särkkä et al., 2012) was used to remove respiration and pulse artefacts using the physiological signals (pulse and respiration) recorded. Then, the data was smoothed with a kernel of 6 mm with the functions in SPM. A temporal high-pass filters of 334 s and 658 s were applied to each of the runs respectively in order to compensate for the signal drift.

Next, the timings and durations of the stimulation blocks were extracted, and a design matrix was constructed accordingly. The design matrix contained also six movement regressors that were used to reduce the confounding effect of movement artefacts. A general linear model (GLM) analysis was applied to obtain the beta-values of each voxel in response to the different stimulus blocks. This was done by having a canonical haemodynamic response function convolved with the stimulus columns of the design matrix. Finally, contrast images of each participant were constructed.

2.8. FMRI analysis

All analysis steps (time-courses, responses strength analyses) were performed in participants’ native space except when region of interest (ROI) locations were compared in the common MNI space and were projected into MN152 template for visualization purposes.

ROI construction and response time-courses. Functional ROIs were constructed from the contrast images using the Marsbar toolbox (MARSellie Boîte À Région d’Intérêt; Marseille, France; version: 0.44). First, for the finger, SM1 cortex ROI was defined as activation posterior to the hand knob of the precentral M1. For the ankle, SM1 cortex ROI was defined as activation in the postcentral mesial wall of paracentral lobule (foot area). The SII cortex ROI was defined as activations within an/or proximity to the posterior insula for contra- and ipsilateral SII cortex separately. In case of the participant having more anterior activation only at M1 cortex, more anterior region, likely covering parts of M1 was used. In case of the CP group participant having lesion near or at the anatomical region of SM1 or SII cortices, primary activation in proximity of these regions was selected.

Next, a sphere of contralateral SM1 or SII cortex and 6 mm (leading to a ROI size of approximately 900 mm² with slight variation between the participants due to discrete spatial data) radius was constructed. This sphere was positioned around the main activation center. After the ROIs were constructed, Marsbar functions were used in a custom Matlab script to extract time-courses for percent-of-signal-change. The value at each time-point was estimated using Marsbar’s finite impulse response model. Then, the value at the peak-response time-point was extracted. Time-courses correspond the average value across all voxels within a given ROI.

After the individual time-courses were constructed, a group-level mean time-course was computed by averaging the individual-level time-courses separately for TD and CP groups. The 95% confidence interval was also calculated for the group-level time-courses.

BOLD response strength. The response strength was quantified using percent-of-signal change of the BOLD signal. Two Multivariate Analysis of Covariance (MANCOVA; R package: jmv-1.2.5) tests were used for the index fingers and ankles separately to test whether group (CP or TD; factorial variable), hand performance or foot performance (covariates) had a main effect on any of the BOLD-responses in the somatosensory cortices. These tests were false discovery rate (FDR)-corrected according to the number of tests in a test family involving BOLD-responses (i.e. 2).

In addition, we controlled the effect of total cumulative head movement (i.e. the sum of head movement over each time point) during stimulation, handedness/footness score and age as potentially confounding covariates and sex as potentially confounding factor. MANCOVA assumptions were tested (see supplementary material / MANCOVA assumptions for more details) and Pillai’s trace (Pillai, 1955) selected as a robust test statistic.

Equivalence testing (R package: equivUMP, function: equiv.test;
Lesion locations. Lesion locations were determined by a semimanual procedure where lesion ROIs were first constructed by visually inspecting CP participants’ T1 MRI images. Then, voxels were manually marked as part of the lesion ROI when white or gray matter was missing using tools of the MRicron software (version: 1.0.20190902). In the case of enlarged ventricles, the enlarged ventricle was compared to the ventricle in the other hemisphere and lesion ROI was manually determined to be approximately the size difference of the ventricles between the hemispheres. The size of the lesion ROI was used to report the size of the lesion in mm$^3$. After the manual determination of the lesion ROI voxels, procedures identical to building the sensorimotor cortex ROIs and projecting them to the MNI space was used to obtain MNI coordinates of the ROIs.

3. Results

The fMRI measurements were successful for all but one CP participant due to strong spasticity of the non-dominant hand. This participant was excluded from the time-series data. The missing response strength value of the spastic hand of this participant was filled in by imputation to the response strength data of 18 CP patients and 23 TD participants.

If the participant did not perform each sensorimotor-performance test, the values were filled in by imputation (4–5 TD and 3–5 CP participants for hand tests, and 2–3 TD and 1 CP participant(s) for foot tests). Only two TD and one CP participant had all of their sensorimotor test scores imputed.

3.1. Proprioceptive BOLD responses

Fig. 2 shows the group-average BOLD time-courses for the proprioceptive stimuli. The response strength (i.e. percent-of-signal change) reached its peak ~ 5–10 s after onset of the proprioceptive stimuli and returned to baseline at ~ 30–35 s.

The proprioceptive BOLD-response strengths of the index fingers differed between the groups (main effect: $p < 0.05$; FDR-corrected for tests related to fMRI-BOLD-responses). Post-hoc tests detected stronger responses for the non-dominant (more affected side in CP) index finger both for SM1 ($p < 0.001$, Cohen’s $d$: 1.0) and SII ($p < 0.05$, Cohen’s $d$: 0.60) cortices. No group differences were significant for the ankles (main effect: $p = 0.50$).

3.2. Sensorimotor performance

Main between-group effect (CP vs. TD) was observed for both hand ($p < 0.001$) and foot ($p < 0.01$; FDR-corrected for behavioral only-related tests) sensorimotor tests. Table 5 presents the performance results and between group post-hoc test values for each test. The sensorimotor performance of CP group was significantly weaker compared to TD group in majority of the applied tests.

Main effect was also observed for the overall sensorimotor performance sum variables ($p < 0.001$). Post-hoc tests revealed that group (CP vs. TD) had a main effect to hand ($p < 0.001$) and foot performance ($p < 0.001$). The TD group had better overall hand and foot performance indicated by the sum variables of the overall sensorimotor performance. Internal consistency of the sum variables was acceptable indicated by Cronbach’s alphas of 0.86 for hand and 0.87 for foot tests respectively.

3.3. Association between proprioceptive BOLD response strength and sensorimotor performance

Fig. 3 shows Spearman correlation coefficients between the proprioceptive BOLD-response strength and sensorimotor performance. Significant negative correlations were observed only for the hand across the entire population (main effect $p < 0.001$). Post-hoc tests revealed that dominant hand function correlated significantly both with SM1 and SII cortex response strengths ($p < 0.001$), and non-dominant hand function correlated with the SII cortex response strength ($p < 0.01$), but not with SM1 cortex response strength ($p = 0.11$). No significant associations were detected for the foot performance and ankle responses. The negative correlations indicate that the worse sensorimotor performance of the hand was associated with stronger proprioceptive BOLD-response strength in the SM1 and SII cortices.

Table 6 shows how each individual sensorimotor performance test contribute to the BOLD response strength variance in each of the cortical ROI both in the case of the linear univariate model and Spearman correlation.

3.4. Lesion and ROI locations

Table 2 shows lesion locations in MNI coordinates in addition to lesion side, type and description.

Finger ROI locations. According to the AAL labels (see supplementary material/ AAL labels tables), 16/23 of TD and 13/18 of CP participants had their dominant SM1 ROI in the SI cortex (postcentral regions). 2 of both TD and CP participants in M1 cortex (precentral regions). Other
regions included inferior parietal (1 TD participant). Four TD and 3 CP participants had their ROIs in undefined regions.

For the non-dominant SM1, 17/23 of TD and 9/17 (one participant had no ROI and response strength was imputed) of CP participants had their SM1 ROI in SI cortex, 5 of TD and 2 of CP participants had their SM1 ROI in M1. Other regions included superior frontal regions (2 CP participants). One TD and 4 CP participants had their ROIs in undefined regions.

For the SII cortex of the dominant finger, 8/23 of TD and 5/18 of the CP participants in the supramarginal regions, 6 of TD and 10 of the CP participants had their ROIs in Rolandic opercular regions 5, of TD and 1 CP participant had their ROIs in the postcentral regions and 4 of TD and 2 of CP participants in the superior temporal regions.

For the non-dominant SII cortex, 17/23 TD and 4/17 (one participant had no ROI and response strength was imputed) CP participants had their ROIs in Rolandic opercular regions, 3 TD and 4 CP participants had their ROIs in defined regions.

For the SII cortex of the dominant finger, 8/23 of TD and 5/18 of the CP participants in the supramarginal regions, 6 of TD and 10 of the CP participants had their ROIs in Rolandic opercular regions 5, of TD and 1 CP participant had their ROIs in the postcentral regions and 4 of TD and 2 of CP participants in the superior temporal regions.

For the non-dominant SII cortex, 17/23 TD and 4/17 (one participant had no ROI and response strength was imputed) CP participants had their ROIs in Rolandic opercular regions, 3 TD and 4 CP participants had their ROIs in undefined regions.

Fig. 2. Group-average BOLD time-courses. Individual participants’ ROI time-courses (an average of the voxel time-courses of the ROIs) were average to yield group-average-time course. Percent-of-signal-change in BOLD-signal over time is used as the measure of the time-course. The grey background indicates the stimulation period. Confidence intervals of 95% are shown in colored shadings. Note that SI and SII cortices are always contralateral to the stimulated limb. * denotes \( p < 0.05 \) and ** denotes \( p < 0.01 \) in peak-BOLD-response amplitude between the groups (TD \( n = 23 \) and CP \( n = 18 \)). Note the difference between the mean time-course sample size (CP group \( n = 17 \)) and statistical test sample size (CP group \( n = 18 \)) in CP as different sample sizes were used by using imputation for statistical testing.

Fig. 3. BOLD-responses versus overall sensorimotor performance. Associations between cortical response strength and sensorimotor performance in SI and SII cortices for hand and foot. Sensorimotor performance is the mean of each performance measure where each performance measure was normalized to a linear scale from 0 (lowest performance value among the participants) to 1 (highest performance value among the participants). * denotes \( p < 0.05 \), ** denotes \( p < 0.01 \) and *** denotes \( p < 0.001 \) for the whole examined population.
in supramarginal regions and 1 TD and 4 CP in postcentral regions. Other regions included superior temporal (1 TD and 1 CP participant), inferior temporal (1 CP participant), insula (1 CP participant), Hescl region (1 CP participant). One TD and CP participant had their ROIs in undefined regions.

Ankle ROI locations. The dominant SM1 ROI was located in the paracentral lobule in 11/23 of TD and 9/18 of CP participants and in precuneus in 5/23 of TD and in 4/18 of CP participants. Other regions included postcentral regions (4 TD and 2 CP participants), superior parietal regions (2 TD participants) and supplementary motor regions (2 CP participants). One TD and 1 CP participant had their ROIs in undefined regions.

For the non-dominant SM1 ankle ROIs, 12/23 TD and 11/18 CP participants had their ROI in paracentral lobule and 9/23 TD and 4/18 had their ROI in the postcentral regions. Other regions included superior parietal regions (1 TD participant), superior motor area (1 TD participant), precuneus (2 CP participants) and precentral regions (1 CP participant).

The dominant SII ROIs were located in supramarginal gyrus in 10/23 of TD and 8/18 of CP participants, superior temporal regions in 5/23 of TD and 3/18 of CP participants and in Rolandic operculum in 4/23 of TD and 6/18 of CP participants. Four TD and 1 CP participant had their ROIs in undefined region.

**Fig. 4.** Finger peak locations projected on group brain. Participants’ finger peak locations of the ROIs are projected on a MNI152 template. Brain is shown from left-lateral, right-lateral and superior views. Single coordinate marked by a small cube indicating the centre-of-mass of the ROI is used.
The non-dominant SII ROIs were located in supramarginal gyrus in 7/23 of TD and 6/18 of CP participants and in Rolandic operculum in 10/23 of TD and 2/18 of the CP participants. Other regions included superior temporal (1 TD and 3 CP participants), middle temporal regions (2 CP participants) and insula (2 CP participants). Five TD participants and 3 CP participants had their ROIs in undefined regions.

**ROI information and visualization.** Figs. 4 and 5 show finger and ankle ROI locations respectively, projected on a MNI152 template. Tables 3 and 4 show finger and ankle ROI sizes and locations in MNI space. AAL labels were also obtained and can be seen in supplementary material tables 1 and 2 (supplementary_material_AAL_labels.docx).

### 3.5. Confounding covariates

None of the confounding factor were significant. The association between BOLD-response strength and cumulative head movement, age or sex or handedness were not significant for the finger (p = 0.36–0.93) or ankle covariates (p = 0.21–0.94).

![Ankle regions-of-interest locations](image)

*Fig. 5.* Ankle peak locations projected on group brain. Participants’ ankle peak locations of the ROIs are projected on a MNI152 template. Brain is shown from left-lateral, right-lateral and superior views. Single coordinate marked by a small cube indicating the centre-of-mass of the ROI is used.
Head movement equivalence. Mean cumulative head movement was 49.5 mm for right hand, 47.3 mm for left hand, 49.0 mm for right ankle and 46.2 mm for left ankle stimulation. Equivalence test revealed that mean head movement did not differ significantly between the different conditions (p < 0.001 for all; null hypothesis reversed).

4. Discussion

The main observation was that adolescents with CP showed higher sensorimotor cortical responsiveness to proprioceptive stimulation of their more affected index finger compared to healthy controls. Moreover, stronger proprioceptive responses were predominantly associated with worse sensorimotor performance of the hands across both groups — with the exception of the SM1 cortex of the non-dominant finger whose association was not significant. Notably, these findings were not replicated in the ankle joint, and this was not explained by head movement during the fMRI scanning but may be related to the sensorimotor control and function of the hands and feet that differ fundamentally, i.e., for example the cortex is more directly involved in the fine motor control of the distal hand whereas spinal and subcortical control is more pronounced in the coarser control of the lower limbs. Altogether, it seems that stronger responsiveness of the sensorimotor cortices of the index finger is associated with worse sensorimotor function of the hand in general whether clinical or non-clinical in essence.

4.1. Responsiveness of the sensorimotor cortices to proprioceptive stimulation in cerebral palsy

It is challenging to conclude how well our findings are in-line with the research literature, as the available prior evidence is limited and highly divergent between the studies. We are aware of only a single study using passive movements of the fingers in fMRI directly comparing CP and TD participants. In contrast to our results, Van de Winckel et al. (2013) reported no evidence of difference in response strength of the SM1 activation between CP and TD groups using passive movements of...
observed weaker somatosensory responses in patients with CP than line with our results, Riquelme and Montoya (2010) reported enhanced movements in fMRI vs. using single movements in MEG. (i.e. active top-down versus passive bottom-up). Electric stimulation and naturalistic passulates also proprioceptive afference, these discrepancies to our study are not necessarily contradicting. In the cutaneous tactile domain, Wingert et al. (2010) demonstrated weaker BOLD-responses of the SM1 cortex to active tactile discrimination task in fMRI when diplegia patients were compared to healthy participants, but no difference between CP and TD children (Van de Winkel et al., 2013). These results are in contrast with our findings. In line with our results, Riquelme and Montoya (2010) reported enhanced early evoked MEG potentials and excitability in the SI cortex to tactile stimulation in participants with CP compared to healthy peers. Furthermore, Maitre et al. (2012) reported stronger contralateral late responses of the more affected hand when compared to the less affected hand in patients with CP. Peripheral electrical stimulation of the median and tibial nerves activate a mixture of tactile and proprioceptive afferents, and thus is adequately relevant comparison to our proprioceptive stimuli. In contrast to our study, Guo et al. (2012) and Trevorrow et al. (2021) observed weaker somatosensory responses in patients with CP than healthy peers using MEG. Even though electric stimulation likely stimulates also proprioceptive afference, these discrepancies to our study are not necessarily contradicting. Electric stimulation and naturalistic passive movements might still activate the sensorimotor cortices differently. The aforementioned partly contradictory findings may arise from different (1) imaging methods (i.e. fMRI versus MEG), (2) lesions types, (3) CP types, (4) modality (e.g. tactile vs proprioceptive vs electric stimulation), (5) stimulation protocol (e.g. single vs. repetitive movements), and/or (6) difference in the attentional engagement of the task (i.e. active top-down versus passive bottom-up). The sensorimotor performance was associated with the strength of the proprioceptive responses. As expected, the CP group had significantly worse hand and foot sensorimotor performance. Majority of the specific performance tests indicated large effect size between the CP and TD participants. CP participants showed worse performance in tasks relying largely on proprioception, such as gross and fine motor skill, stereognosis and balance, or muscle force production (grip-strength test). Thus, these tests appear sensitive enough to identify sensorimotor impairment in the context of CP. Moreover, the dynamic stability of the gait in CP in partly same participants was impaired when compared to controls (see Piitulainen et al., 2020b). However, tactile sensitivity (2-point discrimination and monofilament test) was not significantly affected in our adolescents with CP when compared to the TD peers. The intact cutaneous tactile sense was somewhat surprising, although we did expect more problems in the proprioception-based tasks. Our patients with CP expressed distinct proprioceptive or kinesthetic impairments although they were relatively well functioning, e.g., were able to move without assistive devices. Furthermore, the hemiplegic and diplegic patients showed similar sensorimotor performance, but the hemiplegic patients showed more impaired gross and fine motor skills, albeit tested unilaterally. The sensorimotor performance was associated with the strength of the proprioceptive responses. Stronger cortical responses to proprioceptive finger stimulation indicated worse sensorimotor performance of the hand, in both CP patients and their healthy controls, with the exception of the non-dominant SM1 of the finger. In contrast to this result, Harrach et al. (2020) reported that stronger response strength in the sensorimotor cortices during an active hand motor task in fMRI was somewhat surprising, although we did expect more problems in the context of CP. However, our results are in-line with the research literature on impaired gross and fine motor skills, albeit tested unilaterally.

### Table 5: Sensorimotor Performance and its Associations to Proprioceptive Responses

| Test                                             | TD (n = 23) | CP (n = 18) | HP (n = 12) | DF (n = 6) | MANOVA (CP vs. TD) | Effect size (Cohen's d) |
|--------------------------------------------------|-------------|-------------|-------------|------------|--------------------|------------------------|
| **Hand tests**                                   |             |             |             |            |                    |                        |
| Box and Block (blocks/min)                       | 72.2 ± 7.0  | 53.1 ± 12.1 | 51.3 ± 11.0 | 56.9 ± 14.6| p < 0.001          | -1.4                   |
| Stereognosis (correct answers)                   | 9.6 ± 0.7   | 8.3 ± 2.1   | 7.9 ± 2.4   | 9.1 ± 0.6  | p = 0.09           | -0.8                   |
| Stereognosis (s)                                 | 6.2 ± 2.6   | 12.6 ± 8.4  | 14.5 ± 9.9  | 8.8 ± 1.9  | p < 0.05           | 1.0                    |
| Nine-Hole Peg (s)                                | 17.7 ± 1.8  | 31.3 ± 18.8 | 37.3 ± 20.4 | 19.3 ± 5.6 | p < 0.001          | 1.0                    |
| 2-point discrimination (accuracy mm)             | 2.4 ± 0.3   | 2.7 ± 0.6   | 2.8 ± 0.8   | 2.5 ± 0.3  | p = 0.58           | 0.6                    |
| Monofilaments (thicknes mm)                      | 1.2 ± 0.1   | 1.2 ± 0.4   | 1.3 ± 0.4   | 1.1 ± 0.2  | p = 0.73           | 0.3                    |
| Kinesthesia (accuracy cm)                        | 2.3 ± 0.5   | 2.8 ± 0.6   | 2.7 ± 0.6   | 2.8 ± 0.6  | p = 0.13           | 0.9                    |
| Grip strength (kg)                               | 21.2 ± 6.4  | 16.7 ± 5.3  | 16.2 ± 4.0  | 17.6 ± 6.9 | p < 0.05           | -1.0                   |
| Bottle opening (level)                           | 1.0 ± 0.0   | 1.8 ± 1.0   | 1.8 ± 0.8   | 1.6 ± 1.3  | p < 0.05           | 1.1                    |
| **Foot tests**                                   |             |             |             |            |                    |                        |
| Standing postural stability                       |             |             |             |            |                    |                        |
| Standing eyes open (mm/s)                        | 6.6 ± 3.2   | 9.6 ± 3.1   | 9.0 ± 2.7   | 11.0 ± 3.7 | p < 0.05           | 0.9                    |
| Standing eyes closed (mm/s)                      | 7.5 ± 2.8   | 10.4 ± 3.2  | 9.4 ± 2.3   | 12.8 ± 4.0 | p < 0.05           | 0.9                    |
| Feet together eyes open (mm/s)                   | 8.6 ± 3.2   | 12.3 ± 3.1  | 12.6 ± 3.5  | 11.5 ± 2.0 | p < 0.01           | 1.0                    |
| Feet together eyes closed (mm/s)                 | 12.5 ± 4.1  | 17.2 ± 8.8  | 18.2 ± 9.8  | 14.7 ± 6.1 | p = 0.06           | 0.7                    |
| Dynamic gait stability                           |             |             |             |            |                    |                        |
| Normal gait                                      | 0.04 ± 0.02 | 0.07 ± 0.03 | 0.08 ± 0.03 | 0.06 ± 0.02 | p < 0.001          | 1.2                    |
| Motor task constrained gait                      | 0.04 ± 0.02 | 0.07 ± 0.02 | 0.07 ± 0.03 | 0.06 ± 0.02 | p = 0.01           | 1.1                    |
| Cognitive task constrained gait                  | 0.04 ± 0.02 | 0.07 ± 0.03 | 0.08 ± 0.03 | 0.06 ± 0.03 | p = 0.01           | 1.0                    |

Group and CP subtype differences between single sensorimotor tests. Values are given as mean ± SD. MANOVA was performed between groups (CP vs. TD). Post-hoc test significance values are shown for specific sensorimotor tests. Effect sizes are also shown for the between group difference for specific sensorimotor tests.
used active motor tasks simultaneous with functional imaging.

The association between worse sensorimotor performance and stronger cortical response strength supports our main observation that the CP participants had stronger sensorimotor cortical responses to the proprioceptive stimulation. This was observed in their more affected hemisphere, i.e. in the regions controlling the more affected hand. The same association was detected also in the less affected hemisphere of CP and also for the healthy peers. Thus, it seems unlikely that the between-group differences are due to a confound, e.g., ROI selection bias or spasticity, but reflect physiologically valid effect.

Finally, the lack of association between the response strength and sensorimotor performance at the lower limb might partly be due to the homogeneity of the lower limb performance tests (postural and dynamic balance). Whereas the upper limb performance tests covered a wide variety of upper limb functionality. Thus, the aspects of lower limb function measured might not have covered comprehensively the aspects being coupled with the function of the somatosensory cortices. There is a lack of standardized lower limb test battery that would assess the lower limb function comprehensively, especially in the proprioceptive domain.

### 4.3. Lesion and ROI locations

Enlargement of the lateral vertices was the most common apparent anatomical finding in our CP participants. This was expected as ventricular enlargement is found to be a significant risk factor for CP (Pinto-Martin et al., 1995). In our participants with clearly enlarged ventricles, the cortical responses were mostly in the expected locations. The enlarged ventricles often altered the morphology in vicinity or within the SII cortex, which made ROI identification more challenging in such patients. However, we were able to define the ROI adequately for all CP patients.

The ROI locations appeared to be spatially more variable in participants with CP than in their healthy peers, both in native and common MNI spaces, and especially in the more affected hemisphere. For this reason, the ROI localization was typically more challenging in the CP patients. For example, one patient (participant 3) had a large lesion covering nearly the entire SII cortex, and therefore the ROI of the more affected finger was identified posterior and inferior to a typical SII cortex location. Another similar example was participant 31, whose SII cortex location was identified superior to typical SII cortex location. For the healthy participants, both SM1 and SII cortices of the fingers were typically located in the expected post-central sulcus and posterior insula respectively. However, in few healthy participants the ROI in the SM1 cortex was identified in a more anterior M1 cortex. This is plausible as proprioceptive afference is directed also directly to the M1 cortex (Goldring and Ratcheson, 1972). In the common MNI space, some SM1 cortex ROIs (2-4 cases) were slightly misplaced with 1–2 mm outside from the cortex, possibly because of the smaller head size in our child participants compared to the MNI template (MNI152) constructed from adult data. In the native space, all the ROIs were located within the cortex. It is important to note that all analyses regarding the main hypothesis were performed in native space. Therefore, a failure to correctly map participant into the common MNI space is insignificant, since it was used only for visualization purposes and to provide response locations in the common coordinate system. We were able to use the valid anatomical and functional (i.e. activation) criteria at the individual anatomy (i.e. in native space), however, the SM1 cortex activations in some of the participants’ were located near the skull surface but still within the gray matter of the SM1 cortex.

| Table 6 | Bold response variance explained by sensorimotor tests of the hand. |
|---------|---------------------------------------------------------------|
|         | SM1 cortex | SII cortex | SM1 cortex | SII cortex |
|         | Dom. hand | Non-dom. hand | Dom. hand | Non-dom. hand |
| TD group | Variance expl. (%) | p | Variance expl. (%) | p | Variance expl. (%) | p | Variance expl. (%) | p |
| Box and Block | 5 | -0.24 | 24 | -0.06 | 14 | -0.46 | 12 | -0.28 |
| Stereognosis (correct answers) | 4 | -0.05 | 3 | 0.03 | 10 | -0.28 | 6 | -0.05 |
| Stereognosis (s) | 11 | -0.24 | 3 | -0.06 | 12 | -0.52 | 10 | -0.25 |
| Nine-Hole Peg | 14 | 0.04 | 3 | 0.11 | 22 | -0.02 | 24 | -0.27 |
| 2-point discrimination | 5 | 0.07 | 6 | 0.21 | 2 | -0.04 | 2 | 0.07 |
| Monofilaments & | 1 | 0.01 | 14 | 0.25 | 14 | -0.39 | 2 | -0.02 |
| Kinetesthia | 18 | 0.03 | 4 | -0.08 | 7 | -0.13 | 2 | -0.05 |
| Grip strength | 39 | -0.37 | 35 | -0.37 | 7 | -0.40 | 7 | -0.29 |
| Bottle opening | 4 | -0.05 | 8 | 0.14 | 13 | -0.19 | 36 | -0.46 |
| CP group |  |  |  |  |  |  |  |  |
| Box and Block | 10 | -0.52 | 50 | -0.60 | 8 | -0.51 | 43 | -0.60 |
| Stereognosis (correct answers) | 19 | -0.70 | 3 | -0.31 | 15 | -0.52 | 3 | -0.19 |
| Stereognosis (s) | 24 | -0.65 | 5 | -0.41 | 35 | -0.78 | 7 | -0.29 |
| Nine-Hole Peg | 2 | 0.11 | 9 | -0.19 | 10 | 0.4 | 6 | -0.18 |
| 2-point discrimination | 19 | -0.45 | >1 | -0.18 | 10 | -0.49 | >1 | -0.16 |
| Monofilaments | 9 | -0.22 | 5 | -0.33 | 11 | -0.47 | 7 | -0.45 |
| Kinetesthia | 2 | -0.17 | 8 | 0.16 | 2 | -0.24 | 2 | -0.05 |
| Grip strength | 11 | -0.51 | 14 | -0.45 | 7 | -0.44 | 25 | -0.49 |
| Bottle opening | 4 | -0.07 | 5 | 0.12 | 2 | 0.02 | 7 | -0.06 |

The univariate linear model sums up to 100% of relative variance. **Spearman’s correlation coefficients (p) to BOLD response strength. Positive correlation values are indicated as bold.**

### 4.4. Limitations

As expected for a CP population, the lesion location and size varied substantially within our CP participants from indiscernible or slightly enlarged ventricles to large lesions covering almost the entire SM1 and SII cortices of the affected hemisphere(s). The anatomical and symptomatic heterogeneity in CP populations is a common challenge for convergent conclusions about the disorder. The heterogeneity and limited availability of the patients are likely reasons for diverging observations in the research literature regarding the cortical sensorimotor function in CP. The number of CP participants was low also in the current study, and thus we could not estimate how lesion type or developmental onset affects the cortical proprioceptive responses. Our focus was on the somatosensory cortices, in which the laterality shifts are rare compared to the M1 cortex in CP (Thickbroom et al., 2001). Typically, the laterality shifts of the somatosensory cortices are associated with severe motor impairments (Lemée et al., 2019), whereas our study was focused to the adolescents with spastic CP who were able to walk without assistive devices (GMFCS I–2).
Differences in handedness and footedness can also be a concern when interpreting the between-group results. The CP group was predominantly left-handed and footed (i.e. their more affected side was right) whereas opposite was true for the healthy controls. The sensorimotor performance and BOLD-response strengths were grand-averaged according to the dominance. However, this was the only reasonable approach, because our primary aim was to examine the effect of the brain lesion on the cortical proprioceptive processing in the more affected hemisphere. Moreover, pooling the participants according to limb dominance for the association analysis between the proprioceptive responses and sensorimotor performance might be another potential concern. It is not clear whether the typically-developed, non-dominant side of the healthy participants is directly commensurable to the non-dominant lesion-side of the CP participants. However, we assumed that the non-dominant side of the healthy participants is the best analogue for the lesion-side of the participants with CP when comparing the groups.

We examined patients with spastic CP, which could potentially enhance the proprioceptive BOLD-response strength. The spastic activity could sensitize the muscle spindles via gamma neuron activity. However, the muscle spindles are extremely sensitive to even to tiny length changes (as low as 5 µm during vibration) of their parent muscle even in perfectly relaxed passive conditions (Brown et al., 1967). Therefore, it is unlikely that the mild spasticity in some individuals would have significantly influenced the current results. This view is also supported by our observation that the worse sensorimotor performance was associated with the stronger cortical proprioceptive responses regardless of the participant having CP or not. We further mitigate this potential concern by not including CP participants with severe spasticity in their limbs.

Seven participants were excluded from the data due to excessive head movements during fMRI scanning. Rejection due to excessive head motion is typical in fMRI studies in children and adolescents (Rajagopal et al., 2014; Vanderwal et al., 2015). We attempted to minimize the head motions by presenting the participants a video of slowly changing pictures during the scanning. It has been shown that presentation of video during the fMRI scanning effectively reduced head movements in children and adolescents (Vanderwal et al., 2015). Furthermore, a rather liberal movement rejection threshold (6 mm) was applied and imputation of data in some participants when applicable. This was unavoidable, but the reader should be aware of these potential confounds when making interpretations of this and other studies with populations with elevated risk of excessive head movements.

Functional localizer was not used to identify the ROIs. The use of anatomical ROIs was not possible due to cortical lesions in the CP participants, and thus they did not always follow the normal somatotopic organization. To avoid excessive head movements, the scanning duration was limited as short as possible. For these reasons, the same fMRI was used in the selection of the ROIs locations and extracting the mean BOLD-signal in them. This approach can potentially bias the results (for details, see Poldrack, 2007). However, since robust significant effect of the lesion on the cortical responses was observed with significant correlation with the behavioral sensorimotor performance, it can be assumed that the aforementioned potential confounds negligibly affected the results.

4.6. Conclusions

Individuals with spastic CP showed stronger BOLD-responses to proprioceptive stimulation of the index finger in their more affected (by the brain lesion) sensorimotor cortices compared to healthy peers. The possible neurological mechanism may include (1) impaired efficiency of the cortical proprioceptive processing due to altered inhibition-excitation balance (2) and/or compensatory recruitment of additional neuronal resources to accomplish adequately the proprioceptive processing. Moreover, the stronger cortical proprioceptive responses were predominantly associated with worse sensorimotor performance of the hands across the studied CP and healthy population. Future studies on cortical proprioceptive processing in CP could attempt to test the aforementioned hypotheses potentially explaining the stronger proprioceptive responses in CP.
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CRediT authorship contribution statement
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Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data
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