

Corticospinal Tract Wiring and Brain Lesion Characteristics in Unilateral Cerebral Palsy: Determinants of Upper Limb Motor and Sensory Function.

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1 **Corticospinal tract wiring and brain lesion characteristics in**
2 **unilateral cerebral palsy: determinants of upper limb motor and**
3 **sensory function**

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14 **Abstract**

15 Brain lesion characteristics (timing, location, and extent) and the type corticospinal tract
16 (CST) wiring have been proposed as determinants of upper limb (UL) motor function in
17 unilateral cerebral palsy (uCP), yet an investigation of the relative combined impact of these
18 factors on both motor and sensory function is still lacking. Here, we first investigated
19 whether structural brain lesion characteristics could predict the underlying CST wiring, and
20 we explored the role of CST wiring and brain lesion characteristics to predict UL motor and
21 sensory function in uCP.

22 Fifty-two participants with uCP (mean age (SD): 11y3m (3y10m)) underwent a single-pulse
23 Transcranial Magnetic Stimulation session to determine CST wiring between the motor
24 cortex and the more affected hand (n=17 contralateral; n=19 ipsilateral; n=16 bilateral) and
25 an MRI to determine lesion timing (n=34 periventricular (PV) lesion; n=18 cortico-
26 subcortical (CSC) lesion), location, and extent. Lesion location and extent were evaluated
27 with a semi-quantitative scale. A standardized protocol included UL motor (grip strength,
28 unimanual capacity, bimanual performance) and sensory measures.

29 A combination of lesion locations (damage to the PLIC and frontal lobe) significantly
30 contributed to differentiate between the CST wiring groups, re-classifying the participants in
31 their original group with 57% of accuracy. Motor and sensory function were influenced by
32 each of the investigated neurological factors. However, in a multiple regression analysis,
33 motor function was predicted by the type of CST wiring (more preserved in individuals with
34 contralateral CST wiring ($p<0.01$)), lesion extent and damage to the basal ganglia and
35 thalamus. Sensory function seemed to be best predicted by the combination of a large and
36 later lesion, and an ipsilateral or bilateral CST wiring, which led to increased sensory deficits
37 ($p<0.05$).

38 These novel insights contribute to a better understanding of the underlying pathophysiology
39 of UL function and may be useful to delineate individualized treatment strategies.

40 **Introduction**

41 Upper limb (UL) function is commonly impaired in individuals with unilateral cerebral palsy
42 (uCP), negatively impacting on daily life activities [1]. The large variability in the clinical
43 presentation of UL function, but also in treatment response, has resulted in increasing interest
44 in understanding the underlying neural mechanisms that determine UL function, and its
45 contribution to further optimize therapy planning for the individual with uCP. A number of
46 neurological factors have been put forward as potential predictors of UL function, i.e. the
47 structural brain lesion characteristics (i.e. lesion timing, location, and extent) and the type of
48 corticospinal tract (CST) wiring [2–6].

49 The timing of the lesion during gestation is closely related to the type of the damaged tissue,
50 and can be classified into three categories: malformations (1st and 2nd trimester of pregnancy),
51 periventricular lesion (PV, early 3rd trimester), cortico-subcortical lesions (CSC, late 3rd
52 trimester and around birth) [7]. Previous studies investigating the impact of lesion timing on
53 UL function have shown that individuals with a later lesion (i.e. CSC lesions) present with
54 poorer UL motor and sensory function [2, 3, 5]. Besides lesion timing, lesion location and
55 extent have shown to play an important role in determining UL function, whereby damage to
56 the posterior limb of the internal capsule (PLIC) and the basal ganglia, and a larger lesion
57 extent are related to worse UL motor and sensory function [2, 3] However, there is still large
58 variability in UL function that remains unexplained based on these factors.

59 The unilateral brain damage in individuals with uCP can also result in a partial or complete
60 reorganization of the CST towards the non-lesioned hemisphere [8]. This reorganization of
61 the CST wiring is unique in uCP and refers to the efferent motor input to the affected hand.
62 Researchers have identified three types of CST wiring, i.e. contralateral (CST_{contra}, the
63 affected hand receives input from the crossed CST, originating in the lesioned hemisphere),
64 ipsilateral (CST_{ipsi}, the affected hand receives input from the uncrossed CST, originating in
65 the non-lesioned hemisphere) and bilateral (CST_{bilat}, the affected hand receives input from
66 both the crossed and uncrossed CST, originating in the lesioned and non-lesioned
67 hemisphere, respectively) [8, 9]. It has been suggested that the type of CST wiring is the main
68 factor influencing UL function, whereby individuals with CST_{contra} present with more
69 preserved UL function compared to the other groups [6, 10–13]. Nevertheless, assessing the
70 underlying CST wiring with Transcranial Magnetic Stimulation (TMS) in young children

71 might become challenging. Therefore, the identification of either behavioural or brain lesion
72 features that relate to the underlying CST wiring could be useful to define tailor-made
73 interventions in a clinical setting.

74 Whilst the role of lesion timing, location and extent has been well investigated [2, 3, 14],
75 only a few studies examined the impact of the CST wiring on UL function, and they often
76 have several limitations (i.e. small sample sizes, ordinal scoring of impairments, limited to
77 motor deficits) [5, 10, 15]. Moreover, studies thus far focused on each factor independently,
78 whereas only one study described the impact of the CST wiring and lesion timing on UL
79 function in uCP [10], and only one study reports the impact of CST wiring and lesion extent
80 in children with PV lesions [4]. Although the authors suggested the relevance of both lesion
81 timing and type of CST wiring in predicting UL function, the small sample size, the lack of a
82 standardized evaluation of motor function, and the merely descriptive nature of the study,
83 hampered the possibility of drawing strong conclusions. Furthermore, it has been shown that
84 an intact sensory function is essential to develop an adequate motor function in other
85 neurological disorders (such as adult stroke) [16, 17]. Also in individuals with uCP, sensory
86 and motor function are highly related [1], although the impact of the CST wiring on this
87 relationship remains unknown.

88 In this study, we investigated the impact of CST wiring and structural brain lesion
89 characteristics on UL motor and sensory function in a large group of individuals with uCP,
90 using a systematic and comprehensive evaluation. Our first hypothesis is that the type of CST
91 wiring pattern in unilateral CP can be predicted based on a linear combination of measures of
92 lesion timing, location and extent. Second, we hypothesize that the combination of these
93 predictors together with the CST wiring has a stronger predicting value for UL motor and
94 sensory function than any of these factors alone. Last, we speculate that the relation between
95 motor and sensory function is disrupted by the type of CST wiring.

96

97 **Materials and Methods**

98 **Participants**

99 Children and adolescents with uCP aged between 5-21 years old were recruited via the CP
100 reference center of the University Hospitals Leuven between 2014 and 2017. They were
101 excluded if they (1) received UL botulinum toxin injections six months prior to the
102 assessment, (2) had UL surgery two years prior to the assessment and/or (3) had other
103 neurological or genetic disorders. All individuals assented to participate, all parents signed
104 the informed consent (participants younger than 18 years old), and participants older than 12
105 years also signed the informed consent, in accordance with the declaration of Helsinki. This
106 study was approved by the Medical Ethical Committee of the University Hospital Leuven
107 (S55555 and S56513).

108 Participants with contraindications for the MRI (e.g. metal implants) or the Transcranial
109 Magnetic Stimulation (TMS; ventricular-peritoneal (VP) shunt, seizure two years prior to the
110 study) did not undergo the respective assessment. All TMS measurements were conducted by
111 two experienced physiotherapists (CSM and EJ) and UL function was evaluated by four
112 experienced physiotherapists (LM, CSM, JH and EJ) at the Clinical Motion Analysis
113 Laboratory of the University Hospitals Leuven (campus Pellenberg, Belgium).

114 **Upper limb evaluation**

115 *Motor function*

116 Grip strength, unimanual capacity and bimanual performance composed the motor
117 evaluation. Maximum *grip strength* was assessed using the Jamar® hydraulic hand
118 dynamometer (Sammons Preston, Rolyan, Bolingbrook, IL, USA). The less-affected hand
119 was measured first and the mean of three maximum contractions was calculated per hand.
120 The ratio between hands was used for further analyses to cancel out the effect of age (grip
121 strength ratio = grip strength less-affected hand/grip strength affected hand, whereby a lower
122 score (closer to 1) indicates a grip strength in the affected hand similar to that of the less-
123 affected hand). *Unimanual capacity* was assessed with the Jebsen-Taylor hand function test
124 (JTHFT). The JTHFT reliably measures movement speed during six unimanual tasks [18,
125 19]. Similar to other studies, we used a modified version for children and adolescents with
126 uCP in which the writing task was removed, and the time to carry out each task was reduced

127 from 3 to 2 minutes to avoid frustration [19, 20]. The time to perform every task was summed
128 up and the ratio between hands was used for further analyses to cancel out the effect of age
129 (JTHFT ratio = JTHFT affected hand/JTHFT less-affected hand, whereby a lower score
130 (closer to 1) indicates movement speed in the affected hand similar to that of the less-affected
131 hand). *Bimanual performance* was evaluated with the Assisting Hand Assessment (AHA),
132 which assesses how effectively the affected hand is used in bimanual activities [21–23]. The
133 spontaneous use is evaluated during a semi-structured play session with standardized toys
134 requiring bimanual handling. Given the age range of the participants of this study, the School
135 Kids AHA and the Ad-AHA were administered [22, 24]. The AHA was scored by certified
136 raters (LM and CSM), using the 5.0 version which includes 20 items that are scored from 0
137 (‘does not do’) to 4 (‘effective use’), resulting in a final score between 0-100 AHA units.

138 *Sensory function*

139 Sensory assessments comprised measures of exteroception (tactile sense), proprioception
140 (movement sense), two-point discrimination (2PD, Aesthesiometer®) and stereognosis
141 (tactile object identification), which have been shown to be reliable in this population [25].
142 Tactile and movement sense were classified as normal (score 2), impaired (score 1) or absent
143 (score 0). 2PD was classified according to the width between the two points that the
144 participants could discriminate: normal (0-4mm, score 2), or impaired (>4mm, score 1) [26].
145 Tactile object identification was used as the number of objects that the children could
146 recognize (0-6). In addition, a kit of 20 nylon monofilaments (0.04g - 300g) (Jamar®
147 Monofilaments, Sammons Preston, Rolyan, Bolingbrook, IL, USA) was used to reliably
148 determine threshold values for touch sensation [27, 28]. Touch sensation was categorized as
149 normal (0.008-0.07g), diminished light touch (0.16-0.4g), diminished protective sensation
150 (0.6-2g), loss of protective sensation (4.19-180g) and untestable (300g), according to the
151 manual (Jamar® Monofilaments, Sammons Preston, Rolyan, Bolingbrook, IL, USA).

152 **Structural MRI**

153 Structural images were acquired using three-dimensional fluid-attenuated inversion recovery
154 (3D FLAIR) [321 slices, slice thickness = 1.2 mm, slice gap = 0.6 mm, repetition time = 4800
155 ms, echo time = 353 ms, field of view (FOV) = 250 x 250 mm², 1.1 x 1.1 x 0.56 mm³ voxel
156 size, acquisition time = 5 minutes]. In addition, magnetization prepared rapid gradient echo
157 (MPRAGE) was acquired [182 slices, slice thickness = 1.2 mm, slice gap = 0 mm, TR =

158 9.7ms, TE = 4.6ms, FOV = 250x250mm², voxel size = 0.98x0.98x1.2, acquisition time = 6
159 minutes]. The structural MRI was used to provide a detailed description of the lesion location
160 and extent and to classify the timing of the lesion, which was conducted by a paediatric
161 neurologist (EO).

162 **Timing of the brain lesion** was classified according to the predominant pattern of damage as
163 described by Krägeloh-Mann and Horber (2007) [7]: malformations (1st and 2nd trimester of
164 pregnancy), periventricular lesion (PV, early 3rd trimester), cortico-subcortical lesions (CSC,
165 late 3rd trimester and term), or acquired brain lesions (between 28 days and two years
166 postnatally).

167 **Lesion location and extent** was determined using a semi-quantitative scale recently
168 developed by Fiori et al (2014) [29]. The scale consists of a graphical template with six axial
169 slices of the brain, and an extra template for the basal ganglia (lenticular and caudate),
170 thalamus, posterior limb of the internal capsule (PLIC), brainstem, corpus callosum, and
171 cerebellum. Firstly, the slices corresponding to the template slices are to be found and the
172 lesion is drawn onto the template. Next, the damage to the periventricular, middle and
173 cortico-subcortical layers of each lobe are scored for both hemispheres separately. The sum
174 of the damage to each lobe results in the lobar score, ranging from 0-3 for each lobe. Damage
175 to the basal ganglia (lenticular and caudate), thalamus, PLIC, and brainstem directly is
176 binarily scored from the MRI (affected or non-affected). Damage to the corpus callosum is
177 scored from 0-3, based on the involvement of the anterior, middle and posterior thirds of the
178 corpus callosum on a sagittal view. Last, the involvement of the cerebellum is based on
179 damage to the vermis (0-1) and each of the hemispheres (0-2), resulting in a total score
180 ranging from 0-3. A total ipsilesional score is calculated based on the damage to the lobes (0-
181 3 for each lobe, i.e. total of 0-12) and damage to the subcortical structures (0-5; ranging from
182 0-17). More detailed information about the scale and its scoring procedure can be found in
183 the respective study [29]. This semi-quantitative scale has been shown valid and reliable in
184 children with uCP [29, 30].

185 In the present study, lesion **location** was indicated by the damage to the frontal and parietal
186 lobes (0-4), damage to the basal ganglia and thalamus (0-3), and damage to the PLIC (0-1).
187 These locations were chosen based on their relation to the sensorimotor system [31]. Lesion
188 **extent** was indicated by the total ipsilesional score (0-17).

189 **Transcranial Magnetic Stimulation**

190 Single-pulse TMS was conducted to assess CST wiring. TMS was applied using a MagStim
191 200 Stimulator (Magstim Ltd, Whitland, Wales, UK) equipped with a focal 70mm figure-
192 eight coil and a Bagnoli electromyography (EMG) system with two single differential surface
193 electrodes (Delsys Inc, Natick, MA, USA). A Micro1401-3 acquisition unit and Spike
194 software version 4.11 (Cambridge Electronic Design Limited, Cambridge, UK) were used to
195 synchronize the TMS stimuli and the EMG data acquisition. Motor Evoked Potentials
196 (MEPs) were bilaterally recorded from the muscles opponens pollicis brevis. During the TMS
197 assessment, participants wore a cap that allows creating a grip with a coordinate system to
198 identify the optimal point to stimulate (hotspot) in a standardized and systematic way. The
199 hotspot and the resting motor threshold (RMT, defined as the minimum intensity required to
200 obtain 5/10 MEP of at least 50 μ V in the corresponding muscle) were identified by starting
201 the stimulation intensity at 30% with an incremental increase of 5% [4]. For each
202 hemisphere, stimulation started from the assumed “motor hotspot”, which is located 5cm
203 lateral and 1cm anterior from the scalp middle point (Cz), at 30%. After approximately 2-3
204 pulses, the stimulation intensity was increased 5% for another 2-3 pulses, until MEPs were
205 found. If no MEP can be elicited after increasing up to 60 to 80%, the coil would be moved to
206 a different location on the scalp grid, and the procedure would be repeated until an MEP was
207 elicited. Stimulation up to 100% of the maximum stimulator output was continued until an
208 MEP was elicited. The non-lesioned hemisphere was always stimulated first and allowed to
209 identify contralateral CST projections to the less-affected hand. Stimulation in the non-
210 lesioned hemisphere was continued up to 100% of the maximum stimulator output to search
211 for possible ipsilateral CST projections to the affected hand. Next, the lesioned hemisphere
212 was stimulated to identify possible contralateral CST projections to the affected hand. If only
213 contralateral MEPs from each hemisphere were found, the child was categorized as having a
214 CST_{contra} wiring. If MEPs in the affected hand were evoked from both hemispheres, the child
215 was categorized as having a CST_{bilat} wiring. Lastly, if MEPs in the impaired hand were only
216 evoked when stimulating the non-affected hemisphere, the child was categorized as having a
217 CST_{ipsi} wiring. TMS measures have been shown to be reliable in adults [32, 33] and in
218 children [34]. In this study, the TMS assessment was used for diagnostic purposes. In cases
219 when high intensities were not tolerated, the stimulation intensity was increased up to at least
220 80% of the maximum stimulator output and children were asked to hold a pen to ensure pre-
221 contraction of the evaluated muscle, and thereby facilitate the CST and MEP detection. This

222 allowed us to rule out the possibility of mis-categorizing the child regarding their CST wiring
223 pattern.

224 **Statistical analyses**

225 First, descriptive statistics were used to document the distribution of brain lesion
226 characteristics according to the CST wiring. Next, we investigated the differences in
227 occurrence of lesion timing, location, and extent between the CST wiring groups by using
228 analysis of contingency tables (Chi-Square and Fisher's exact test), Kruskal-Wallis test
229 (ordinal data), and ANOVA (lesion extent). Lastly, we used discriminant analysis to explore
230 whether the type of CST wiring would differ depending on the linear combination of lesion
231 timing, location, and extent, in a multivariate way. Cross-validation procedure was included
232 to investigate the accuracy of the model in reclassifying the participants in the original CST
233 wiring groups. Variables related to lesion timing, lesion location (damage to the frontal lobe,
234 parietal lobe, PLIC, and basal ganglia and thalamus), and extent (ipsilesional extent of the
235 lesion) were included in the model, which was fitted using the stepwise selection method.

236 To investigate the impact of the type of CST wiring and brain lesion characteristics on UL
237 function, we first used linear simple regression and then multiple regression analysis to
238 investigate the combined impact of these factors on UL motor and sensory function. For the
239 continuous variables related to motor function, normality was first verified by inspecting the
240 histograms and with the Shapiro-Wilk test, showing a normal distribution only for the AHA.
241 For the JTHFT ratio and the grip strength ratio, a logarithmic transformation was applied (y'
242 $= \log_{10}(y)$). To investigate the impact of the type of CST wiring and brain lesion
243 characteristics on UL motor function, we computed a multiple regression analysis. Similarly,
244 for UL sensory function, we conducted a simple ordinal logistic regression for stereognosis
245 and thresholds for touch sensation, and a simple logistic regression for 2PD to investigate the
246 impact of each individual neurological factor on the sensory function. Next, we performed
247 multiple regression analyses (ordinal and logistic) to investigate the combined impact of the
248 neurological predictors on the sensory deficits. The predictors included in the multiple
249 regression model were the type of CST wiring, lesion timing, location (damage to the frontal
250 lobe, parietal lobe, PLIC, and basal ganglia and thalamus), and ipsilesional extent of the
251 lesion. To predict both motor and sensory function, interaction terms were built between the
252 CST wiring and (i) lesion timing, and (ii) lesion extent, and included in the model. The

253 multiple regression models were fitted with the backward elimination method until a set of
254 variables significantly contributing to the model was identified.

255 Lastly, to investigate the relation between sensory and motor function for the whole group
256 and within CST wiring groups, Spearman rank correlation coefficients were used between
257 each of the motor function variables and deficits in stereognosis. Correlation coefficients
258 were considered as little or no correlation (<0.30), low ($0.30-0.50$), moderate ($0.50-0.70$),
259 high ($0.70-0.90$) and very high correlation (>0.90) [35].

260 In addition, effects sizes were calculated for the comparisons and interpreted according to
261 Cohen, depending on the computed test: η^2 (partial eta squared) for the prediction models
262 (small 0.01, medium 0.06, large 0.14) [36, 37]. Statistical significance was set at $\alpha < 0.05$ for
263 main tests with Bonferroni correction for post-hoc tests. All statistical analyses were
264 computed with SPSS Statistics for Windows version 24.0 (IBM Corp. Armonk, NY: IBM
265 Corp.).

266 **Results**

267 **Participants**

268 Seventy-five children and adolescents with uCP participated in this study (mean age (SD):
269 11y1m (3y6m); 33 girls; 39 left uCP). According to the Manual Ability Classification System
270 (MACS), 25 individuals were classified as MACS I, 25 as MACS II and 25 as MACS III.
271 Sixteen participants did not have CST wiring data ($n=1$ panic attack, $n=2$ hemispherectomy,
272 $n=3$ VP shunt, $n=2$ epilepsy, $n=1$ tumor, $n=4$ refusals to participate, $n=3$ inconclusive TMS
273 results), resulting in a total of 59 participants. The TMS assessment identified 20 individuals
274 with CST_{contra} , 18 with CST_{bilat} and 21 with CST_{ipsi} . For the analyses in this study,
275 participants with malformations ($n=1$), acquired lesions ($n=4$) or no visible lesions ($n=2$)
276 were excluded due to the very small sample size of these sub-groups, resulting in a total
277 group of 52 participants (mean age (SD): 11y4m (3y10m); 22 girls; 28 left uCP) with
278 available CST wiring ($n=17$ contralateral; $n=19$ ipsilateral; $n=16$ bilateral) and data related to
279 the timing, location, and extent of the lesion. A summary of the lesion locations and extent
280 according to the lesion timing is provided in Supplementary materials (Table 1). Thirty-four
281 individuals had a PV lesion and 18 had a CSC lesion. Clinical motor and sensory data was
282 missing in one participant (boy, 19y7m, PV lesion, and CST_{contra} wiring) and sensory data

283 was evaluated in a subsample of participants (see sensory function results section for more
284 details).

285 **CST wiring and brain lesion characteristics**

286 Table 1 displays the distribution of lesion timing, location and extent variables according to
287 the three CST wiring groups. Except for the damage to the parietal lobe, all variables were
288 significantly different between the CST wiring groups ($p < 0.05$) (Table 1).

289 In the discriminant analysis, we found that the combined value of the damage to the PLIC
290 and the damage to the frontal lobe could significantly discriminate between the type of CST
291 wiring (Wilks' $\lambda = 0.611$, Chi-square = 23.88, $df = 4$, Canonical correlation = 0.602, $p <$
292 0.001). The two functions extracted accounted for nearly 57% of the variance in the type of
293 CST wiring. The standardized discriminant function coefficients of the two extracted
294 functions indicated the contribution of each retained independent variable (damage to the
295 PLIC and damage to the frontal lobe) to each function, showing how strongly the
296 discriminant variables affect the score. These coefficients can be then used for the
297 classification of a single individual (Function 1 = $0.81 * \text{damage to the PLIC} + 0.50 * \text{damage}$
298 $\text{to the frontal lobe}$; Function 2 = $-0.60 * \text{damage to the PLIC} + 0.88 * \text{damage to the frontal}$
299 lobe).

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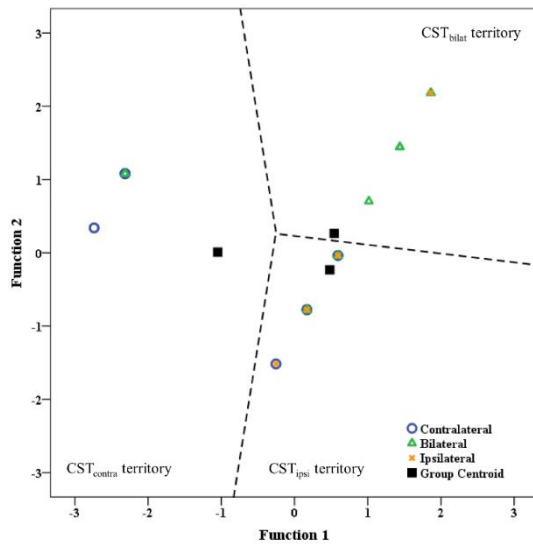
308 Table 1. Contingency table (count and percentage, descriptive statistics) of the occurrence of
 309 lesion timing, location, and extent according to the CST wiring.

			CST wiring			p-value
			Contralateral	Bilateral	Ipsilateral	
Timing						
Lesion timing [¥]	PV	N (%)	15 (88.2%)	8 (50%)	11 (57.9%)	0.04
	CSC		2 (11.8)	8 (50%)	8 (42.1%)	
Location						
PLIC [¥]	Not affected	N (%)	8 (47%)	1 (6%)	0 (0%)	<0.001
	Affected		9 (53%)	15 (94%)	19 (100%)	
Basal ganglia and thalamus [◇]		Me (p25-p75)	0 (0-1)	1.50 (0 (2.50)	1 (1-2)	0.006 ^{a,b}
Frontal Lobe [◇]		Me (p25-p75)	1 (1-1)	1.50 (1-2.25)	1 (1-1.50)	0.004 ^{a,b}
Parietal Lobe [◇]		Me (p25-p75)	2 (1-2)	2 (1.25-3)	2 (2-2.50)	0.09
Extent						
Ipsilesional extent [○]		X (SD)	5.18 (3.07)	8.38 (3.95)	9.05 (3.27)	0.004 ^{a,b}

310 CST, corticospinal tract; PV, periventricular; CSC, cortico-subcortical; PLIC, posterior limb
 311 of the internal capsule. [¥]Chi-Square statistic, [§]Fisher's exact test, [◇]Kruskal-Wallis test,
 312 [○]ANOVA. ^aContralateral vs. Ipsilateral; ^bContralateral vs. Bilateral.

313

314 Cross-validated reclassification of cases based on the new canonical variables was successful
 315 in 57.7% of the cases: 89.5% were correctly classified in the CST_{ipsi} group, 47.1% in the
 316 CST_{contra} group, and only 31.3% in the CST_{bilat} group (Fig 1).



317

318 Fig 1. Territorial map showing the relative location of the boundaries of each CST wiring
 319 category and the location of each of the participants. The group centroids are indicated with a
 320 black filled square [CST_{contra} (-1.05, 0.01); CST_{ipsi} (0.48, -0.23); CST_{bilat} (0.54, 0.26)].

321

322 CST wiring, brain lesion characteristics and UL function

323 *Motor function*

324 Descriptive statistics of the motor function according to the type of CST wiring, lesion
 325 timing, location, and extent are presented in Supplementary materials (Table 2). The simple
 326 linear regression analyses to predict motor function based on a single neurological factor
 327 showed that every factor had an influence on motor function (grip strength, $p < 0.04$; JTHFT,
 328 $p < 0.004$; AHA, $p < 0.01$; see Supplementary materials Table 2 for detailed information).

329 When all the neurological factors were included in the same model in a multiple regression
 330 analysis, the backward elimination method identified the variables that were significantly
 331 contributing to the model. Table 2 documents the estimated marginal means, which represent
 332 the mean response in each CST wiring group adjusted by the covariates that significantly
 333 contribute to the model. The multiple regression model to predict **grip strength** deficits only
 334 retained the type of CST wiring, explaining 46% of the variance ($F(2, 51) = 20.90$; $p < 0.001$;
 335 $\eta^2 = 0.47$). For the **JTHFT**, 54% of the variance was explained by the type of CST wiring
 336 ($F(2, 51) = 12.20$; $p < 0.0001$; $\eta^2 = 0.34$, $R^2 = 46\%$) and the total extent of the lesion ($F(1,$

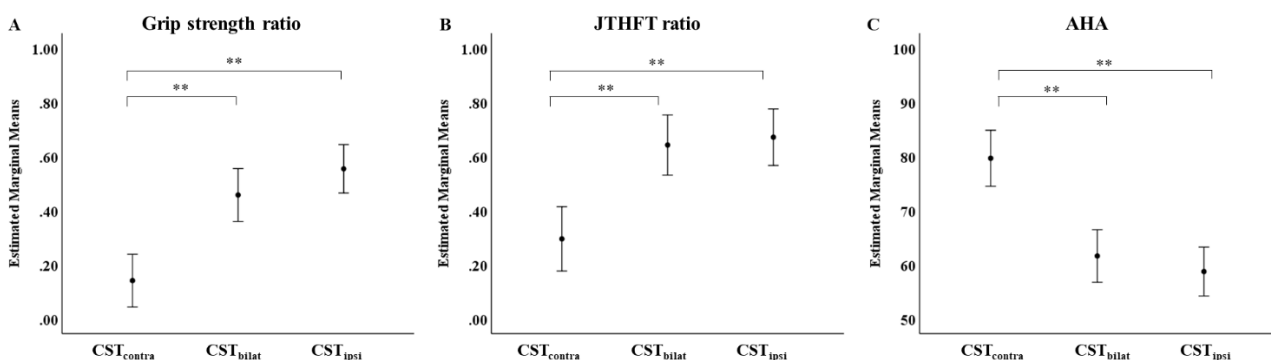
337 51)=8.05; $p=0.007$; $\eta^2=0.15$, $\Delta R^2=8\%$). For **bimanual performance** (AHA), the regression
 338 model explained 61% of the variance, with the type of CST wiring ($F(2, 51)=19.03$;
 339 $p<0.0001$; $\eta^2=0.45$, $\Delta R^2=52\%$), the total extent of the lesion ($F(1, 51)=10.65$; $p<0.001$;
 340 $\eta^2=0.19$, $\Delta R^2=5\%$), and the damage to the basal ganglia and thalamus ($F(1, 51)=4.90$;
 341 $p=0.03$; $\eta^2=0.10$, $\Delta R^2=4\%$) significantly contributing to the model (Fig 2). No interaction
 342 effects were identified for any of the motor outcome variables.

343 Table 2. Descriptive statistics of the observed and estimated marginal means of upper limb
 344 motor function according to the CST wiring groups.

Estimated marginal means and SD			
	CST _{contra} (n=16)	CST _{ipsi} (n=19)	CST _{bilat} (n=16)
Grip strength ratio (log)	0.14 (0.13) ^a	0.55 (0.20) ^a	0.46 (0.24) ^a
JTHFT ratio (log)	0.30 (0.24) ^b	0.67 (0.23) ^b	0.64 (0.22) ^b
AHA (0-100)	79.66 (10.28) ^c	58.70 (9.81) ^c	61.58 (9.67) ^c

345 CST, corticospinal tract; JTHFT, Jebsen-Taylor Hand Function test; AHA, Assisting Hand
 346 Assessment; SD, standard deviation. ^a The values coincide with the observed values, as there
 347 is not significant covariate in the model. ^b Adjustments based on ipsilesional lesion extent
 348 mean = 7.67. ^c Adjustments based on ipsilesional lesion extent mean = 7.67, and damage to
 349 the basal ganglia and thalamus mean = 1.12.

350



351

352 Fig 2. Upper limb motor function differs in individuals with CST_{contra} compared to those with
 353 CST_{bilat} or CST_{ipsi} wiring. Estimated Marginal Means and 95% CI per CST wiring type and
 354 lesion timing group for A) Grip strength (log ratio, i.e. closer to zero indicates preserved grip
 355 strength), B) JTHFT (log ratio, i.e. closer to zero indicates preserved manual dexterity,
 356 measured by speed) and C) AHA.

357 AHA, Assisting Hand Assessment; JTHFT, Jebsen Taylor Hand Function Test; CST,
 358 corticospinal tract. * $p<0.01$; ** $p<0.001$. Estimated Marginal Means are adjusted according to
 359 the significant covariates (see Table 2 for details).

360 *Sensory function*

361 Descriptive information of sensory function according to each neurological factor is
362 summarized in Table 3 of Supplementary materials. Sensory function data (tactile sense,
363 movement sense, stereognosis and 2PD) and thresholds for touch sensation, as assessed with
364 the monofilaments, were available in 46 and 35 individuals, respectively. Due to the lack of
365 variation in the tactile sense and movement sense modalities, the predictive model was only
366 applied to the stereognosis, 2PD and the thresholds for touch sensation.

367 The simple linear analyses to predict sensory function based on a single neurological
368 predictor indicated that every predictor impacted on stereognosis ($p < 0.032$). In contrast, 2PD
369 was influenced by all neurological predictors ($p < 0.04$) except the damage to the PLIC
370 ($p < 0.17$), and touch sensation could be significantly predicted by all factors ($p < 0.01$) except
371 damage to the PLIC ($p = 0.99$) and type of CST wiring ($p = 0.42$).

372 When all the neurological factors were included in the same model in a multiple regression
373 analysis, the backward elimination method identified predictors that were significantly
374 contributing to the model. For **stereognosis**, the retained main effects were the CST wiring
375 (Wald Chi-square (2) = 9.09, $p = 0.011$), lesion timing (Wald Chi-square (1) = 4.34, $p = 0.04$)
376 and ipsilesional extent of the lesion (Wald Chi-square (1) = 7.15, $p = 0.008$) (Table 3A). These
377 results show that the odds of having better stereognosis function were 5.56 times higher in
378 the group with PV lesions than in the CSC group ($p = 0.04$). Similarly, individuals with a
379 CST_{contra} wiring show 10.23 and 9.7 times higher probability of having better scores in the
380 stereognosis test compared to those with a CST_{ipsi} or CST_{bilat}, respectively ($p = 0.02$), whilst
381 there was no difference between the last two ($p = 0.34$). Lastly, the odds of having higher
382 stereognosis scores decreases by 0.74 for every unit change in the ipsilesional extent of the
383 lesion ($p = 0.01$). No interactions were found between the CST wiring and the brain lesion
384 characteristics to predict deficits in stereognosis ($p > 0.05$)

385 The logistic multiple regression to predict **2PD** showed lesion timing (Wald Chi-square (1) =
386 10.62, $p = 0.001$) and ipsilesional extent of the lesion (Wald Chi-square (1) = 3.75, $p = 0.05$) to
387 be significant contributors ($p > 0.05$) (Table 3B). The odds of having an impaired 2PD is 31
388 times higher in the group with CSC lesions than in the PVL group ($p = 0.001$). Secondly, the
389 odds of having impaired 2PD increase by 1.34 for every unit change in the ipsilesional extent

390 of the lesion ($p=0.05$). No interactions were found between the CST wiring and the brain
 391 lesion characteristics to predict deficits in 2PD ($p>0.05$)

392 Table 3. Descriptive statistics of the sensory function (3A, stereognosis (number of correctly
 393 recognized objects; 3B, two-point discrimination; 3C, touch sensation) according to each of
 394 the variables significantly contributing to each prediction model.

Table 3A

		Stereognosis (number of correctly guessed objects)						
		0	1	2	3	4	5	6
Lesion timing								
PV	N (%)	0 (0%)	0 (0%)	1 (25%)	0 (0%)	5 (71%)	6 (67%)	17 (44%)
CSC	N (%)	5 (100%)	2 (100%)	3 (75%)	1 (100%)	2 (29%)	3 (33%)	1 (6%)
CST wiring								
Contralateral	N (%)	0 (0%)	0 (0%)	1 (25%)	0 (0%)	0 (0%)	1 (11%)	13 (72%)
Bilateral	N (%)	4 (80%)	0 (0%)	2 (50%)	0 (0%)	3 (43%)	3 (33%)	3 (17%)
Ipsilateral	N (%)	1 (20%)	2 (100%)	1 (25%)	1 (100%)	4 (57%)	5 (56%)	2 (11%)
Lesion extent								
Ipsilesional	Me (IQR)	13 (2.07)	13 (-)	10 (3.88)	-	6 (3.50)	6 (5.25)	5.25 (3.75)

Table 3B

		Two-point discrimination	
		Normal ($\leq 4\text{mm}$)	Impaired ($> 5\text{mm}$)
Lesion timing			
PV	N (%)	26 (93%)	3 (17%)
CSC	N (%)	2 (7%)	15 (83%)
Lesion extent			
Ipsilesional	Me (IQR)	5.25 (3.88)	12 (5.25)

Table 3C

		Threshold of touch sensation				
		Normal	Diminished light touch	Diminished protective sensation	Loss of protective sensation	Untestable
Lesion extent						
Ipsilesional	Me (IQR)	6 (4.50)	-	10.50 (11.25)	13 (2.41)	12.50 (-)

395 PV, periventricular lesion; CSC, cortico-subcortical lesion; CST, corticospinal tract; N,
 396 number of cases; Me, median; IQR, interquartile range.

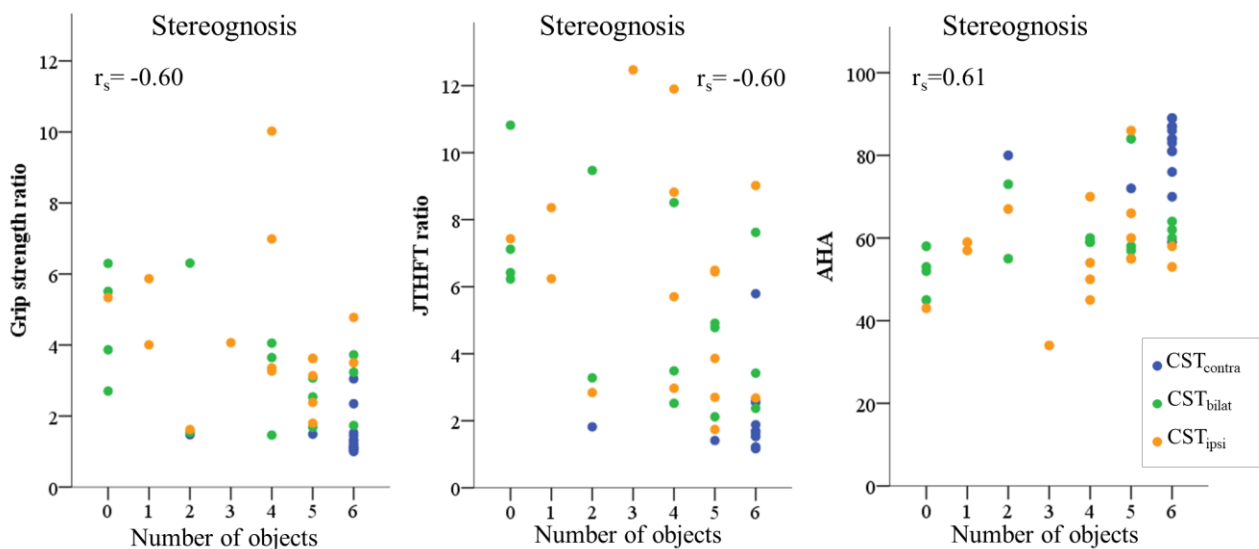
397

398 The ordinal logistic multiple regression for **touch sensation**, as measured by the
 399 monofilaments, indicated that only the lesion extent significantly contributed to the deficits in
 400 touch sensation (Wald Chi-square (1) = 10.75, $p=0.001$) (Table 3C). The odds of having
 401 better touch sensation decreases by 0.66 for every unit change in the ipsilesional extent of the
 402 lesion. No interactions were found between the CST wiring and the brain lesion
 403 characteristics to predict deficits in touch sensation ($p>0.05$).

404 *Impact of CST wiring on the relation between motor and sensory function*

405 The correlation analyses between the motor and sensory function for the whole group
 406 indicated a moderate association between the stereognosis score and grip strength ratio ($r_s = -$
 407 0.60 , $p<0.001$), JTHFT ratio ($r_s = -0.60$, $p<0.001$) and AHA ($r_s = 0.61$, $p<0.001$).

408 After group division according to CST wiring, there was no to low correlation between motor
 409 function and stereognosis in the CST_{contra} and CST_{ipsi} groups (r_s (range) = -0.31 - 0.36 , $p>0.05$).
 410 Interestingly, in the CST_{bilat} group, moderate correlations were found with the JTHFT ratio
 411 ($r_s = -0.48$, $p=0.07$) and the AHA ($r_s=0.65$, $p<0.01$), despite a low correlation with grip
 412 strength ratio ($r_s = -0.31$, $p=0.2$). An illustration of the individual data points regarding these
 413 results can be found in Fig. 3.



414
 415 Fig 3. The relation between motor and sensory function seem to vary depending on the CST
 416 wiring. Individuals with a CST_{contra} and CST_{ipsi} showed no to low correlations, whereas those
 417 with CST_{bilat} showed moderate correlations. Each dot represents an individual child, with
 418 CST_{contra} (blue), CST_{bilat} (green), and CST_{ipsi} (orange). Correlations between stereognosis

419 with grip strength ratio (ratio, i.e. closer to one indicates preserved grip strength), JTHFT
420 ratio (ratio, i.e. closer to one indicates preserved grip strength), and AHA. Correlation
421 coefficients correspond to the analysis for the whole group.

422

423 **Discussion**

424 In this study, we explored the predictive value of brain lesion characteristics on the type of
425 CST wiring as well as the impact of these factors on UL motor and sensory function. A
426 comprehensive and standardized evaluation of both motor (grip strength, unimanual capacity
427 and bimanual performance) and sensory function was used to predict UL function in a large
428 cohort of individuals with uCP.

429 Our first research question examined the discriminant ability of lesion timing, location and
430 extent to predict the type of CST wiring. A simple linear analysis demonstrated that lesion
431 timing, location and extent were significantly different between the CST wiring groups. Our
432 results showed that a CST_{contra} was only seen in 2 out of 18 children with a CSC lesion,
433 compared to 15 out of 34 children with a PV lesion. Current results suggest that damage to
434 cortical and/or subcortical structures (i.e. CSC lesion), reduces the potential of the CST to
435 develop according to its typical contralateral trajectory. We hypothesise that this is likely
436 driven by the reduced neural activity in the motor cortical areas after a CSC lesion, which are
437 crucial for the development of the CST during the postnatal period [38]. However, a
438 contralateral development of the CST is still possible in CSC lesions, and it may occur
439 differently depending on lesion location and extent.

440 Once all predictors were simultaneously entered in a multiple linear analysis, we found that
441 the combination of the damage to the PLIC and the frontal lobe significantly discriminated
442 between the CST wiring groups. Half of the children in the CST_{contra} group showed damage
443 to the PLIC, in contrast to the 94% and 100% in the CST_{bilat} and CST_{ipsi} group who showed
444 damage to this white matter bundle. Furthermore, the frontal lobe was also more damaged in
445 the CST_{bilat} and CST_{ipsi} groups, compared to the CST_{contra}. Although it is not unexpected that
446 the PLIC and the frontal lobe are the two significant predictors in the model, due to their
447 undoubtable relation with the motor cortex and the performance of actions, this is the first
448 time that this interaction with the type of CST wiring is shown. Contrary to the importance of
449 the location, Staudt et al (2002) postulated that the type of CST wiring depended on the

450 lesion extent [4]. However, as they only included children with a PV lesion, their results
451 cannot be extended to all the uCP population. Further efforts should be made to underpin
452 whether structural damage of the brain lesion may serve as a biomarker of the underlying
453 CST wiring.

454 Next to the predictive model, we also investigated how accurate the two functions derived
455 from the discriminant analysis would be to reclassify the individuals in their original
456 categories. Despite the significant contribution of the PLIC and the frontal lobe to the
457 discriminant model, the classification accuracy only reached 57%, suggesting that timing,
458 location and extent of the lesion (as included in the model) do not provide sufficient accurate
459 information to predict the underlying type of CST wiring. Notwithstanding the validity and
460 reliability of the semi-quantitative scale that was used to investigate lesion location and
461 extent, we acknowledge that the semi-quantitative character of the scale may have
462 underestimated the predictive value of the structural brain damage. Therefore, these results
463 should be replicated in the future with volumetric measures of the different brain structures.
464 For example, the projections to the PLIC have been shown to be topographically organized
465 with reduced microstructural integrity in children with uCP [39] by using diffusion measures.
466 Investigating the volumetric damage to the frontal lobe and the microstructural integrity of
467 the PLIC may provide with further insights in determining the type of CST wiring in uCP.

468 For our second research question, we investigated the impact of CST wiring and brain lesion
469 characteristics (timing, location, and extent) on motor and sensory function. Regarding *motor*
470 *outcome*, simple linear regression analyses indicated that the CST wiring and all brain lesion
471 characteristics had an influence on the grip strength, manual dexterity and bimanual
472 performance, which confirmed what previous studies have shown [5, 6, 10]. However, in the
473 multiple linear regression analysis, we found that the underlying CST wiring plays a major,
474 but not unique, role in determining UL motor function, as lesion location and extent also
475 significantly contributed to increasing the explained variance for the JTHFT and for the
476 AHA. Specifically, the type of CST wiring explained 46% and 52% of the JTHFT and the
477 AHA variances, respectively, which was increased up to 54% and 61% by including lesion
478 extent and damage to the basal ganglia and thalamus into the model. In general, our results
479 show that a CST_{ipsi} or CST_{bilat} lead to poorer UL motor function compared to CST_{contra} for all
480 motor outcomes, even when controlling for the significant contribution of lesion extent and
481 location. The importance of the underlying CST wiring is an expected result, as the CST is

482 the main motor drive and its damage causes vast disturbances on voluntary motor control,
483 drastically reducing motor capabilities [38]. Whilst lesion timing, location, and extent have
484 been put forward as a predictor of UL function [2, 3] and was also confirmed in our linear
485 regression analysis, the huge variability in motor function reported by previous studies seems
486 to be mainly explained by the underlying CST wiring. Staudt et al. (2004) were the first to
487 report on the relation between CST reorganization potential at different gestational ages and
488 UL motor function [10]. These authors also found that, along with the CST wiring, UL motor
489 function further worsened in later lesions (CSC lesions) [10]. Linear regression analysis also
490 showed that later lesions led to poor motor outcome, but multiple regression analysis
491 revealed that lesion location and extent were key factors, next to the type of CST wiring.
492 Although later lesions seem to be associated to a larger extent [3], it seems that the lesion
493 extent itself plays a more important role in motor outcome, i.e. children with a PV lesion with
494 large extent will also present with poorer hand function. Interestingly, the damage to the
495 basal ganglia and thalamus explained an extra 4% of the variability in the AHA. In
496 accordance with our results, previous studies have reported the negative impact of these
497 subcortical structures on UL motor outcome [2, 5].

498 It is important to note that we still found large variability in the three motor outcome
499 measures within both the CST_{ipsi} and CST_{bilat} groups, whereas the variability in the CST_{contra}
500 group was rather small (Fig 2, see also Table 2 Supplementary materials for observed
501 means). In other words, some individuals with a CST_{ipsi} and CST_{bilat} had good motor
502 function, similar to those with a CST_{contra} wiring. This variability could not be completely
503 explained by the location and extent of the lesion, and other factors may play a role. In the
504 CST_{ipsi} group, this large variability may be explained by the amount of overlap of the hotspot
505 within the non-lesioned hemisphere to evoke MEPs in the affected and less-affected hand.
506 Vandermeeren et al. (2009) showed that dexterity indeed varies in individuals with ipsilateral
507 wiring depending on the location of the hotspot of the CST innervating the affected hand and
508 less-affected hand: overlapping hotspots resulted in poorer dexterity, whereas distinct non-
509 overlapping hotspots resulted in a preserved dexterity [40]. Conversely, in the CST_{bilat} group,
510 the large variability may be explained by a predominant contralateral or ipsilateral projection
511 that controls the affected hand, as Jaspers et al. (2016) proposed in their theoretical
512 framework [9]. Altogether, this seems to point toward a distinct underlying pathophysiology
513 of the UL motor impairments in these two CST groups (CST_{ipsi} or CST_{bilat}), suggesting that
514 individuals with either a CST_{bilat} or CST_{ipsi} pattern should be treated as two separate groups

515 for future research. To further unravel the underlying mechanisms of the pathophysiology of
516 motor control and motor capabilities in uCP, additional functional measures should be
517 included such as excitatory and inhibitory intracortical circuits based on TMS (e.g. cortical
518 silent period or paired-pulse paradigms) [15, 41], or functional connectivity of the
519 sensorimotor network based on resting-state functional MRI [42, 43].

520 We also investigated the impact of the CST wiring and brain lesion characteristics on *sensory*
521 *function*, based on the fact that CST projections also extend from the primary sensory cortex
522 and mediate several sensory functions at the level of the spinal cord (control of nociceptive,
523 somatosensory, and somatic motor functions) [44, 45]. Although our simple linear regression
524 analyses suggested that all neurological factors individually played a role in determining
525 sensory function, the multiple prediction model showed that a larger lesion extent, a later
526 lesion (i.e. CSC lesion) and a CST_{ipsi} or CST_{bilat} led to higher chances of developing sensory
527 deficits. Our results are in agreement with a recent study by Gupta et al (2017), who showed
528 that more than 80% of the children with larger extent and later lesions (CSC) had disrupted
529 somatosensory anatomy and physiology (lack of ascending sensory tracts and lack of
530 somatosensory evoked potentials), consequently leading to a loss of sensory function [6]. If
531 the sensory tracts are present, there is evidence suggesting that their main compensatory
532 mechanism is an intra-hemispheric reorganization, i.e. the sensory system reaches the
533 original cortical destination on the post-central gyrus, regardless of lesion timing (PV or CSC
534 lesion) or CST wiring [11, 46, 47]. Current study results suggest that lesion extent best
535 predicts the sensory deficits in individuals with uCP, although lesion timing and CST wiring
536 also play an important role. Future research focussing on the pathophysiology of the sensory
537 system based on non-invasive neurophysiological techniques (e.g. short latency afferent
538 inhibition [48] or sensory evoked potentials [11]), as well as functional connectivity
539 measures, may contribute to increase our understanding of the underlying sensory pathways
540 in uCP.

541 Lastly, we investigated whether the relationship between motor and sensory function was
542 disrupted by the type of CST wiring. We first confirmed previous study results indicating a
543 significant relation between the motor and sensory outcomes in the total group [1, 25].
544 However, this association was disrupted by the type of CST wiring, whereby no to little
545 association was shown in the CST_{ipsi} and CST_{contra} groups, but a moderate association was
546 found for the CST_{bilat} group. In the CST_{contra} group, the lack of a significant (or high)

547 correlation seems to be due to the fact that these participants show both adequate motor and
548 sensory function, with little variation in the sensory scale, due to its ordinal nature. This scale
549 used to evaluate sensory function may not be sensitive enough to detect subtle sensory
550 deficits, leading to a possible ceiling effect in the CST_{contra} group. By measuring with more
551 quantitative techniques and devices, e.g. KINARM End-point Lab (BKIN Technologies)
552 [49], we may be able to discern the potential sensory problems that these individuals may
553 present with. Secondly, the sensorimotor dissociation found in the CST_{ipsi} group may be
554 explained at two different levels of the central nervous system. At the level of the spinal cord,
555 the descending CST fibres entering the dorsal horn play an important role in presynaptic
556 inhibition of primary sensory afferent fibres [45, 50], ensuring smooth execution of a
557 movement. A CST_{ipsi} wiring may have consequences in the presynaptic inhibition at the level
558 of the spinal cord and could, consequently, affect the relation between motor and sensory
559 function. On the other hand, at the level of the brain, the intra-hemispheric communication
560 between M1 and S1 has been shown to be very relevant for adequate processing of
561 sensorimotor information [51–53]. As such, the lack of intra-hemispheric cortico-cortical
562 connections may affect the processing of sensory information, having a negative impact on
563 the motor command. On the contrary, the CST_{bilat} group seems to preserve the relation
564 between motor and sensory function, as shown by the stereognosis modality. This may be
565 potentially explained by the predominant behaviour that those with a CST_{bilat} hypothetically
566 show [9]. A relation between adequate sensory and adequate motor function, as seen in the
567 CST_{contra} group may indicate a more ‘contralateral’ behaviour, whilst a disparate relation may
568 be indicative of rather an ‘ipsilateral’ behaviour. However, this needs further confirmation
569 with neurophysiological tools. Although current data do not allow drawing strong
570 conclusions regarding sensorimotor integration, our results highlight the importance of
571 investigating these aspects in the future to better understand the mechanisms of sensorimotor
572 information processing in uCP. By using more advanced techniques to unravel the coupling
573 between the sensory and motor system, we will be able to determine the impact of such
574 dissociation on motor control and motor performance. For instance, short latency afferent
575 inhibition has been put forward as a valuable indicator of the process of bilateral
576 sensorimotor integration [48] and may potentially aid in measuring the reorganization of
577 sensorimotor pathways in uCP.

578 There might be some important clinical implications based on the results of this study. A
579 better understanding of the underlying mechanisms of motor and sensory impairments will

580 surely contribute to developing new treatment approaches, specifically targeting the
581 individual pathophysiological deficits. First, the type of CST wiring has been investigated as
582 a potential biomarker of treatment response. Although motor improvement does not seem to
583 be CST-type dependent after bimanual training [12, 54], there are conflicting results
584 regarding unimanual training [55–57]. Furthermore, our results highlight the importance of
585 considering the sensory system together with the available motor execution paradigms during
586 UL training. Preliminary results of recent studies have shown the effectiveness of bimanual
587 and sensory training on both motor and sensory function in uCP [58, 59]. To further support
588 interventions targeting sensory deficits, there is evidence in healthy adults suggesting that
589 sensory input can modulate the excitability in both motor cortices simultaneously, as well as
590 the communication between hemispheres [60]. In this line, it seems relevant to combine
591 bimanual and sensory training to enhance the excitability of both motor cortices, which may
592 increase intra- and inter-hemispheric connections between the sensory and motor systems,
593 potentially resulting in long-lasting neuroplastic changes.

594 Next to the training approaches, it is also important to identify clinically feasible measures to
595 infer the CST wiring and the sensory system. As these assessments are not always pleasant in
596 young children nor practical in a clinical setting, there is a necessity to find tools that are
597 more applicable to daily practice than neurophysiological techniques. To probe the motor
598 system, mirror movements have been put forward as a valid clinical assessment tool that may
599 reflect the underlying individual CST wiring [9, 61]. On the other hand, it seems very
600 challenging to develop an accessible and simple tool to clinically probe the sensory system in
601 uCP. Further research in this field is required to develop quantitative and valid measures of
602 sensory function (e.g. perceptual threshold of touch with electrical stimulation [62] or robotic
603 measures of proprioception [49, 63]) and to link these measures to the underlying
604 mechanisms of the sensory system in uCP.

605 There are some limitations to be considered for the current study. First, we used scales for the
606 evaluation of lesion location and extent, as well as for assessing sensory function that were
607 based on an ordinal scoring. Although they have been shown to be reliable in uCP [25, 29],
608 such scales may lack sensitivity. Second, our study lacked a neurophysiological technique to
609 probe the sensory system (i.e. sensory evoked potentials), that may contribute to better
610 understand the underlying mechanisms of sensory function in individuals with uCP. Third,
611 the main limitation of the TMS assessment itself lays in the maximum stimulator output

612 intensity that can be reached. This intensity may not have been sufficient to elicit a MEP
613 from either the lesioned or the non-lesioned hemisphere, as the resting motor thresholds are
614 normally higher in children and may be even higher in individuals with uCP. This limitation
615 might have prevented us from finding a CST projection to eventually diagnose the individual
616 as CST_{bilat} or CST_{ipsi} wiring. Furthermore, the MEP data were not analysed, which may
617 provide with useful insights in future studies. Lastly, although our sample size was large and
618 covers the most common lesion timing groups, our results cannot be completely extended to
619 those children with malformations or postnatally acquired brain injuries, as these were not
620 included in the analyses.

621 **Conclusions**

622 CST wiring mainly determines UL motor function, although also lesion extent and damage to
623 the basal ganglia and thalamus significantly contributed to the prediction of UL motor
624 deficits. For sensory function, lesion extent, timing, and the type of CST wiring pattern seem
625 to be important to develop adequate sensory function. The underlying CST wiring seems to
626 disrupt the association between sensory and motor function, pointing toward different
627 mechanisms of sensorimotor integration in uCP. The results of our study contribute to a
628 better understanding of the underlying pathophysiology of motor and sensory function and
629 highlight the importance of investigating sensorimotor integration in future studies.
630 Subsequently, these insights will aid in developing new intervention strategies tailored to the
631 specific deficits of the motor and sensory system of the individual child with uCP.

632 **Data Availability**

633 All data concerning this study is available within the manuscript. Detailed data is available
634 upon request to the first author.

635 **Conflicts of Interest**

636 The authors declare that there is no conflict of interest regarding the publication of this paper.

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647 **Supplementary materials**

648 Table 1. Descriptive information of the distribution of the lesion location and extent
649 according to the lesion timing groups.

650 Table 2. Descriptive statistics (X (SD)) and univariate analysis of upper limb motor function
651 according to the CST wiring and the brain lesion characteristics.

652 Table 3. Descriptive statistics (Me (IQR)) and univariate analysis of upper limb sensory
653 function (3A, stereognosis and 3B, two-point discrimination and thresholds of touch
654 sensation) according to the CST wiring and the brain lesion characteristics.

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