RESEARCH ARTICLE

Altered Modulation of Silent Period in Tongue Motor Cortex of Persistent Developmental Stuttering in Relation to Stuttering Severity

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Abstract

Motor balance in developmental stuttering (DS) was investigated with Transcranial Magnetic Stimulation (TMS), with the aim to define novel neural markers of persistent DS in adulthood. Eleven DS adult males were evaluated with TMS on tongue primary motor cortex, compared to 15 matched fluent speakers, in a "state" condition (i.e. stutterers vs. fluent speakers, no overt stuttering). Motor and silent period thresholds (SPT), recruitment curves, and silent period durations were acquired by recording tongue motor evoked potentials. Tongue silent period duration increased in DS, especially in the left hemisphere (P<0.05; Hedge’s g or Cohen’s d_unbiased = 1.054, i.e. large effect size), suggesting a “state” condition of higher intracortical inhibition in left motor cortex networks. Differences in motor thresholds (different excitatory/inhibitory ratios in DS) were evident, as well as significant differences in SPT. In fluent speakers, the left hemisphere may be marginally more excitable than the right one in motor thresholds at lower muscular activation, while active motor thresholds and SPT were higher in the left hemisphere of DS with respect to the right one, resulting also in a positive correlation with stuttering severity. Pre-TMS electromyography data gave overlapping evidence. Findings suggest the existence of a complex intracortical balance in DS tongue primary motor cortex, with a particular interplay between excitatory and inhibitory mechanisms, also in neural substrates related to silent periods. Findings are discussed with respect to functional and structural impairments in stuttering, and are also proposed as novel neural markers of a stuttering “state” in persistent DS, helping to define more focused treatments (e.g. neuro-modulation).
Introduction

Developmental Stuttering (DS) is a disruption in the normal rhythm of speech: persons are unable to utter a fluent speech. It begins during childhood and, in some cases, persists in adulthood. DS symptoms are syllable repetitions and blocks, usually at the beginning of sentences/words and in the majority of cases it affects males. It is accompanied by secondary, associated movements/spasms, mainly of facial muscles [1,2]. The cause of DS is not completely clear. It is considered a complex/multifactorial disorder [3,4,5], arising from genetic factors [6], which alter neurologic function [7]. DS may be related to heritable impairments of late fibers myelination [8,9], which may cause the abnormal connectivity observed in DS [10,11]. DS is characterized by neural markers that represent differences in brain anatomy/functioning in comparison to fluent speakers [5,12,13]. Speech-related structures show greater activity in the right hemisphere with respect to homologue areas of the left hemisphere [12], but also greater neural activation in supplementary motor regions [5]. It is not clear if the majority of DS neural abnormalities are a prerequisite for its appearance, or if they are mainly the result of long-term stuttering (comprising attempts to overcome dysfluencies) [14,15]. Overt dysfluency is not essential to differentiate DS brain from fluent individuals (stuttering “state”-i.e. DS vs. controls- in comparison to a stuttering “trait”-i.e. fluency vs. dysfluency-) [5,13]. However, DS remains an incompletely understood neurologic problem. Speech is a task that requires rapid motor control and its demands may easily be disrupted [16]: thus, dysfluency may be a symptom of a more complex motor syndrome also involving abilities not directly involved in speech [17,18,19,20]. Importantly, DS is characterized by a dopamine-related abnormal functioning of basal ganglia [4,21,22], and it may share characteristics with other basal ganglia-related motor disturbances, such as Tourette's Syndrome [1] or Parkinson's Disease, especially when considering timing-related skills [23]. DS improves after administration of antidopaminergic, serotoninergic or GABAergic drugs [24,25,26,27,28,29,30], i.e. substances acting on the excitatory/inhibitory ratio of motor networks that rely also on basal ganglia. Interestingly, previous work suggests slight neural differences between DS males and females [31,32,33], likely also in relation to hormone variability [34]. Neural characteristics of DS have been extensively studied but the potential of non-invasive brain stimulation, and thus the possibility to directly investigate excitatory/inhibitory ratio of neural networks [5], has been underestimated until now. Transcranial magnetic stimulation (TMS) has been used only recently [25,31,35,36,37,38,39,40,41,42,43], especially when considering speech-related motor effectors [37,39,43]. As a consequence, the present work aims to further investigate abnormalities in the excitatory/inhibitory ratio of the primary motor cortex representation of speech muscles (tongue) in persistent DS by using TMS. Considering that neural abnormalities in DS are also present when stutterers are not realizing a speech-specific motor task [38], and/or during rest [31,35,42], in the present work we aimed to investigate the “basic” motor activity of speech muscles, in DS males, by measuring TMS indexes, such as motor thresholds, motor evoked potentials (MEPs), and durations of silent periods, in comparison to fluent speakers (some of which will be investigated for the first time in tongue muscles of DS, such as silent period). Measures were recorded aiming to further characterize the aberrant neural functioning in speech-related motor effectors of persistent DS. More specifically, the main objective of the present work is to individuate novel neural markers of a stuttering “state”, to better clarify the existing ratio between excitatory and inhibitory motor circuits in DS. As a consequence, data were also correlated with stuttering severity and behavioral/cognitive indexes of DS.
Materials and Methods

Participants
A total of 30 participants were recruited for the study, including 14 DS adult males and 16 matched fluent speakers. All DS participants had been developmental stutterers since childhood. Undetected stuttering (and no previous history of DS) was excluded in fluent speakers. All participants had no history of major neurological and/or psychiatric problems. All participants were right-handed Italian native speakers and none were under treatment with psychotropic drugs, such as antidepressants or antipsychotic drugs. Four participants (three DS and one fluent speaker) dropped out due to difficulties in maintaining electrodes on the tongue, TMS discomfort and/or impossibility to evoke reliable responses [44]. Thus, 11 DS adult males (age 24–47 years) and 15 adult fluent speaker males (age 22–42 years) were evaluated and considered in experimental analyses. Of this sample, one DS participant and one fluent speaker were unable to complete the entire procedure due to peripheral stimulation discomfort. Procedures were approved by the Unique Regional Ethical Committee (CERU) of Friuli-Venezia Giulia (referring to the University Hospital of Trieste), and were in accordance with the Declaration of Helsinki (and with recent TMS guidelines) [45]. A screening for evaluating the possible risks related to TMS delivery was also conducted before the experiments [46]. Participants signed a written informed consent before the experiments. Groups were controlled for variables such as age, sex, education, handedness [47], smoking habits [48], migraine [49], musical expertise [50], depressive symptoms (Beck Depression Inventory-II- BDI-II-) [51], and physical activity habits [52,53].

TMS settings and experimental design
TMS (Medtronic MagPro R30; eight-shaped coil C-B60; biphasic stimulation; antero-posterior direction of the first phase of the current in the coil) was administered to obtain MEPs from tongue muscles on its primary motor cortex representation, in every hemisphere. Participants sat in a relaxed position and they were asked to keep their eyes open during stimulations. Murdoch et al. [54] suggested that an optimal coil direction should be individuated in every participant when investigating tongue motor cortex, but this may add experimental variability. As a consequence, after verifying the presence of reliable MEPs, TMS coil was always maintained on the head by the experimenter, normally at 45° with respect to the inter-hemispheric fissure (coil handle pointing backward). Muscular activity was recorded by four surface Ag/AgCl electrodes on the tongue dorsal part. They were symmetrically placed: two electrodes were about 0.5 cm from the midline, close to the tongue tip, on its right and left side. Remaining electrodes were placed at a distance of about 2 cm, toward the posterior part of the tongue, on external sides. All electrodes were connected to an amplifier, with ground electrode on the right forearm. Positioning of electrodes was checked visually and by electromyography (EMG), considering that a complete rest of tongue is difficult to obtain. A tissue cap was placed on the participants’ head, to individuate the position of the hot-spot on the scalp: it was normally placed about 2 cm ahead and 4 cm laterally, with respect to hand muscles representation) [44]. Tongue representation in primary motor cortex is able to reach both the left and right side of this muscular district [55], and thus measures obtained from contralateral and ipsilateral sides were considered. TMS was delivered by randomized blocks of stimulation to obtain indexes such as motor threshold (MT; asking to maintain 10–20% of tongue maximal muscular activation, visually verified during stimulations), active motor threshold (AMT) and silent period threshold (SPT), asking to maintain 60–70% of tongue maximal muscular activation (dorsiflexion), visually verified on EMG. MT was considered as the stimulation intensity able to evoke MEPs of at least 50–100 µV in half of stimulations. AMT was defined as the stimulation...
intensity able to evoke a MEP of at least 200 μV in half of the stimulations. SPT [56,57] was defined as the stimulation intensity that was able to evoke a visible and reliable silent period (measured in tens of ms) in half of the stimulations. Motor thresholds were determined by modifying TMS intensity at 1% steps. Silent period durations and latency, recorded maintaining about 60–70% of tongue maximal muscular activation (dorsiflexion) were also bilaterally measured, stimulating at 130% SPT. Silent period duration was measured from the appearance of the MEP until the reappearance of muscular activity [58]. Recruitment curve was recorded by stimulating at 110%, 125%, and 140% MT, maintaining lower tongue activation (see MT). Tongue MEPs were considered reliable when responses showed latencies comprised between about 6–13 ms [44]. Two adjutive EMG channels were used to evaluate artifacts: the first channel evaluated peripheral TMS responses, such as jaw displacement due to peripheral nerve stimulation (Ag/AgCl electrodes placed on the jaw), while the second was used to evaluate TMS-evoked eye blinks (Ag/AgCl electrodes placed around eyes). We also registered MEPs from first dorsal interosseous (FDI) muscle in DS, by stimulating the left and the right primary motor cortex and recording contralateral responses using Ag/AgCl electrodes placed in a tendon-belly montage. AMT and SPT were recorded, asking participants to maintain a muscular contraction of about 30–40% of maximal activation, as well as silent period durations (measured as above indicated), stimulating at 150% SPT. This was realized to compare silent period data obtained from hand muscles of persistent DS males [31], and data obtained from tongue, with stuttering severity in the same sample of participants, considering the existence of neurophysiological heterogeneity in DS [14,15]. TMS coil was maintained by the experimenter at 45° with respect to the inter-hemispheric fissure, with the handle pointing backward. EMG data before TMS delivery were also analyzed to verify if the groups were homogeneous when considering spontaneous tongue EMG activity, bilaterally. In this case, 20 ms before TMS delivery were considered in recruitment curve data, and 60 ms in silent period (duration and latency) data. EMG was acquired by using a sampling rate of 8000 Hz, visualized by a digital band-pass filter of 20–2000 Hz. Tongue silent period (duration and latency) was obtained by averaging about six consecutive stimulations, in every hemisphere. The same was done for every stimulation intensity (in every hemisphere) for recruitment curve. When considering FDI silent periods in DS, about six stimulations were averaged, in every hemisphere. Experimental design was implemented to minimize muscular fatiguing; TMS was delivered by applying random inter-stimulus intervals of about 2–6 seconds.

Behavioral measures
Severity of stuttering was evaluated in DS by using the Stuttering Severity Instrument-4 (SSI-4) [59]. A speech sample was acquired from DS participants, measuring percentages of stuttered syllables, the severity of physical concomitants, and the duration of the longest blocks. Both groups were administered an Italian version of the BigCAT Questionnaire [60], a self-evaluation of speech-associated attitudes (negative or positive) adapted to DS (35-items). The Italian version of the Cognitive Behavioral Assessment (CBA) 2.0 scale [61] was also administered: it is a battery of self-administered questionnaires useful for investigating personality characteristic, emotional adjustment, and psychological status, evaluating whether groups differ in behavioral/cognitive states/traits. Data were correlated with neurophysiologic indexes.

Statistical analysis
Motor thresholds were expressed in percentages with respect to maximal TMS output. Recruitment curve data were expressed as peak-to-peak amplitudes (μV), latencies (ms), and areas under the curve (V/sec). Silent period durations and latencies were expressed in ms. EMG data
(pre-TMS), were evaluated considering areas under the curve (V/sec). Raw EMG data (obtained from TMS and pre-TMS recordings) were always considered for analyses. i.e. data were not rectified and/or averaged between conditions before calculations, in order to limit data handling. Data from cognitive/behavioral measures (SSI-4, BigCAT, CBA 2.0) were reported as interval measures. Statistical analyses were performed by mixed model analysis [62,63]. When considering TMS, factors were groups (stuttering vs. fluent speakers), stimulated hemispheres (left vs. right), and, when considering silent period durations, silent period latencies, and recruitment curve, also side of the tongue (left vs. right). When considering recruitment curve, also stimulation intensities were considered (110%, 125%, 140% MT). Main effects and interactions among factors were investigated. Suitable degrees of freedom for the mixed model analysis were approximated by considering sample sizes, and subtracting free parameters. Pre-TMS EMG data were analyzed by considering groups, stimulated hemispheres, tongue sides, and stimulation intensities (when appropriate). In post-hoc analyses, cognitive/behavioral variables, and analyses of homogeneity (i.e. age, handedness, education, musical expertise, smoke habits, migraine, and physical activity), data normality was evaluated by the Shapiro-Wilk Test. Homogeneity of variance was also verified in between-groups comparisons. Differences in normally distributed data were assessed by Student's t-test (Welch's t-test in non homogenous data; t), while non-parametrical tests were performed in not normally distributed data (Mann-Whitney or Wilcoxon rank sum test; Z). Categorical data (physical activity) were evaluated by using a Chi-square test (Yates correction for low frequencies). Neef at al. [37] suggested that neurophysiologic measures may not always be independent with an unknown real degree of correlation, making classical corrections for multiple comparisons unfeasible. Moreover, no formal consensus has been reached on this topic [64,65,66]. As a consequence, we decided to report here raw, uncorrected, values, applying a false discovery rate method on a family-wise basis (i.e. behavioral data, motor thresholds data based on MEPs, recruitment curves, silent period data, pre-TMS EMG data; significant values that did not survive are accompanied by (*) [67], on pairwise, between/within groups post-hoc comparisons. An estimate of the effect sizes was also performed in significant two-means comparisons, depending on data normality and homogeneity, and between/within-participants comparisons (absolute values; Hedge’s g or Cohen’s d_unbiased, Glass’s delta, Φ, d: 0.2<d_unbiased, delta, d<0.5 small effect size; 0.5<d_unbiased, delta, d<0.8 medium effect size; 0.8<d_unbiased, delta, d>0.8 large effect size; 0.1<r<0.3 small effect size; 0.3<r<0.5 medium effect size; r<0.5 large effect size) [68,69,70,71,72,73,74]; in non-parametric comparisons both r and the parametric counterparts are reported, to allow a more complete evaluation of effects. In this context, "a priori" statistical power (about 80%) calculations, mainly based on hypothesized differences between groups in silent periods and recruitment curves, justified sample sizes (considering also possible deviations from normality in data). Finally, a correlation analysis was also performed to evaluate relations between SSI-4, TMS indexes, and behavioral/cognitive data, in DS and fluent speakers, by Pearson's correlation (r; in normally distributed data), and by Spearman's correlation (in not normally distributed data; in alternative, Gamma statistic was used when more appropriate -i.e. when tied observations were also present; γ). A P<0.05 was considered significant (two-tailed assumption; P<0.1: trend toward significance).

Results
Behavioral measurements

No significant differences were evident between groups when considering measures evaluating group homogeneity in factors such as age, education, handedness, depressive symptoms (BDI-II; this was confirmed also when separately evaluating cognitive/somatic symptoms,
Table 1. Summary of the characteristics of the DS and fluent speakers groups.

| Group characteristic/Exp. Group | Stuttering | Fluent speakers | Statistics | P |
|---------------------------------|------------|----------------|------------|---|
| Age                             | 32.8 (9.0) | 29.5 (5.6)     | Z = 0.628  | 0.55 |
| Education                       | 17.5 (3.9) | 15.7 (2.3)     | Z = 1.079  | 0.29 |
| Handedness                      | 83.2 (12.8)| 85.9 (12.3)    | t_{94} = 0.562 | 0.58 |
| Beck Depression Inventory-II    | 5.1 (5.0)  | 3.1 (4.0)      | Z = 1.079  | 0.29 |
| Smoking                         | 0.29 (0.5) | 0.20 (0.4)     | Z = 0.805  | 0.48 |
| Musical Expertise               | 0.31 (0.5) | 0.18 (0.4)     | Z = 1.244  | 0.24 |
| Migraine                        | 0.02 (0.1) | 0 (0)          | Z = 1.677  | 0.42 |
| Physical Activity               | 5/6        | 12/3           | $\chi^2$ (Yates correction) = 1.6 | 0.2 |
| BigCAT                          | 24.9 (10.3)| 37.3 (3.4)     | Z = 4.154  | <0.001 |
| EPQ/R-N scale (CBA 2.0)         | 60.4 (24.4)| 36.3 (28.0)    | t_{94} = 2.277 | 0.031 |
| QPF/R scale (CBA 2.0)           | 63.4 (26.5)| 26.4 (27.0)    | Z = 2.889  | 0.003 |
| IP/PH scale (CBA 2.0)           | 51.8 (20.2)| 25.3 (14.2)    | Z = 3.120  | 0.001 |

Data obtained from the main characteristics evaluated to match experimental groups. Data regarding smoke habits, musical expertise, migraine have been evaluated on a scale basis, while physical activity has been evaluated on a categorical basis. Data obtained from speech attitudes evaluation (BigCAT) and data resulted significantly different in cognitive/behavioral evaluation (CBA 2.0) are also reported. Data are reported indicating mean and standard deviation in brackets. Significant differences between groups are marked in bold.

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Statistics not reported), smoking habits, musical expertise, migraine, and physical activity (see Table 1). SSI-4 showed that DS ranged from “very mild” to a “severe” level. BigCAT showed a higher negative attitude toward speech situations in DS than controls (effect size: $r = 0.815$ -Hedge’s g or Cohen’s d_{unbiased} = 2.876; see Table 1 for significance). When considering CBA 2.0, DS group resulted significantly different with respect to fluent speakers in subscales such as the EPQ/R-N scale (Hedge’s g or Cohen’s d_{unbiased} = 0.878; higher levels of emotional lability in DS), the QPF/R scale (r = 0.567; Hedge’s g or Cohen’s d_{unbiased} = 1.337; higher tendency of psychophysiological disturbances in DS), and in the IP/PH scale (r = 0.612; Hedge’s g or Cohen’s d_{unbiased} = 1.514; higher levels of phobia in DS). Scores obtained from DS and fluent speakers always resulted under the threshold for psychopathological disturbance ($\geq95^{th}$ percentile). On a singular participant level, two fluent speakers had values $\geq95^{th}$ percentile, while six DS had values $\geq95^{th}$ percentile. The main findings are reported in Tables 1 and 2.

Motor thresholds

When considering TMS data and, more specifically, motor thresholds comparisons, statistics showed a marginal difference between groups and stimulated hemispheres in MT (overall significance of the model, $P = 0.063$; groups x stimulated hemispheres: $t_{12} = -1.845$, $P = 0.079$). This was supported by the fact that the fluent speakers’ left hemispheres had lower MT with respect to their right one ($t_{14} = 2.414$, $P = 0.030$; $d = 0.475$), while this difference was not highlighted in DS. When considering AMT, there was a marginal difference between groups and stimulated hemispheres (overall significance of the model, $P = 0.065$; groups x stimulated hemispheres $t_{12} = -1.836$, $P = 0.081$). More specifically, left hemisphere AMT resulted higher with respect to the right hemisphere AMT in DS ($t_{9} = 1.958$, $P = 0.082$; $d = 0.411$). Finally, when considering SPT, there was a significant difference between DS and fluent speakers when considering the interaction between groups and stimulated hemispheres (overall significance of the model, $P = 0.026$; groups x stimulated hemispheres: $t_{12} = -2.374$, $P = 0.027$). Findings suggest higher SPT in the left hemisphere of DS participants with respect to their right one ($t_{9} = 2.388$, $P = 0.041$; $d = 0.466$). The main findings are summarized in Table 3 and Fig 1.
Recruitment curves

When considering recruitment curves, MEPs amplitudes (\(P<0.001\)) show a general effect related to intensity of stimulation (t22 = 6.395, \(P<0.001\); t22 = 4.260, \(P<0.001\)). An effect was evident in the interaction between groups, stimulated hemispheres, and intensities of stimulation (t19 = 2.111, \(P = 0.048\)), showing marginal difference in DS, when comparing MEPs obtained in the two hemispheres and stimulating at 110% MT (left hemisphere MEPs higher of right hemisphere MEPs; Z = 1.956, \(P = 0.05^{(*)}\); \(r = 0.590 - d = 0.588\)). When considering MEPs areas (significance of the overall model: \(P<0.001\)), an effect related to intensity of the stimulation was evident (t22 = 6.040, \(P<0.001\); t22 = 3.760, \(P = 0.001\)). A marginal effect related to the interaction between groups and intensity of stimulation was also evident (t22 = -1.745, \(P = 0.096\)), suggesting a possible difference between DS and fluent speakers when stimulating at 125% MT (Z = 1.946, \(P = 0.054^{(*)}\); \(r = 0.382 - Hedge's g or Cohen's d_{unbiased} = 0.389\); higher MEPs areas in DS). Fluent speakers showed a significant negative correlation of MEPs areas (obtained when stimulating the right hemisphere at 125% MT) with physical activity (\(\Gamma = -0.78\)). No differences were evident in MEPs latencies. Pre-TMS EMG analyses related to recruitment curve are reported in the Supporting Information (S1 File). The main findings are summarized in Table 3, Table A and Fig A in S1 File.

Silent period durations

When considering silent period durations, statistics resulted significant (overall significance of the statistical model, \(P = 0.044\)), suggesting differences in the interaction between groups and stimulated hemispheres (t22 = -2.257, \(P = 0.034\)). More specifically, silent period durations were longer when stimulating tongue motor cortex of the left hemisphere in DS, with respect to fluent speakers (t22 = 2.651, \(P = 0.014\); Hedge's g or Cohen's d_{unbiased} = 1.054). Analysis of silent period latencies did not revealed significant differences. When considering pre-TMS EMG data related to silent period, recorded side of the tongue resulted significantly different (t22 = 2.403, \(P = 0.025\)), suggesting that the right side is generally more activated, with respect to the left side, during spontaneous and sustained contractions before TMS delivery, in both groups (Z = 1.841, \(P = 0.066^{(*)}\); \(r = 0.361 - d = 0.211\)). The interaction between groups and recorded side of the tongue resulted marginally significant (t22 = 2.023, \(P = 0.055\)), suggesting a greater difference in DS (with respect to fluent speakers) in the spontaneous and sustained pre-

**Table 2. Stuttering Severity Instrument-4 evaluation of DS participants.**

| Stuttering participant/SSI-4 indexes | SSI-4 value | SSI-4 percentile | SSI-4 category |
|-------------------------------------|------------|-----------------|----------------|
| a                                   | 18         | 12–23           | Mild           |
| b                                   | 31         | 61–77           | Moderate       |
| c                                   | 25         | 41–60           | Moderate       |
| d                                   | 30         | 61–77           | Moderate       |
| e                                   | 32         | 78–88           | Severe         |
| f                                   | 23         | 24–40           | Mild           |
| g                                   | 12         | 1–4             | Very mild      |
| h                                   | 32         | 78–88           | Severe         |
| i                                   | 31         | 61–77           | Moderate       |
| j                                   | 36         | 89–95           | Severe         |
| k                                   | 13         | 5–11            | Very mild      |

Stuttering Severity Instrument-4 scores, in every DS participant, are reported.

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Table 3. Summary of the main findings obtained by TMS.

| Neurophysiologic index/Exp. Group | Stuttering LH | Stuttering RH | Fluent speakers LH | Fluent speakers RH |
|----------------------------------|--------------|--------------|--------------------|-------------------|
| MT (%)                           | 58.3 (11.3)  | 57.8 (11.4)  | 50.8 (11.5)        | 55.9 (8.1)        |
| AMT (%)                          | 49.1 (10.8)  | 44.7 (11.9)  | 43.6 (13.4)        | 45.6 (7.1)        |
| SPT (%)                          | 50.2 (9.9)   | 45.4 (11.7)  | 44.1 (12.9)        | 47 (7.7)          |
| 110% MT amplitude (μV)           | 184.5 (140.6)| 103.5 (59.7) | 148.2 (109.3)      | 152.0 (171.4)     |
| 125% MT amplitude (μV)           | 153.3 (127.5)| 110.8 (62.9) | 138.9 (122.4)      | 147.4 (129.5)     |
| 140% MT amplitude (μV)           | 336.0 (248.0)| 315.7 (230.0)| 298.3 (249.0)      | 205.8 (144.7)     |
| 110% MT area (V/sec)             | 1495.8 (910.7)| 1555.1 (996.7)| 1472.1 (1609.9)     | 1054.9 (1004.1)   |
| 125% MT area (V/sec)             | 1724.1 (977.8)| 1735.1 (1036.3)| 1319.0 (1551.1)     | 1092.7 (985.4)    |
| 140% MT area (V/sec)             | 2388.5 (1356.8)| 2392.9 (1919.1)| 2794.4 (2328.3)     | 1940.7 (1331.1)   |
| 110% MT latency (ms)             | 10.0 (1.7)   | 10.2 (0.9)   | 9.5 (1.1)          | 9.9 (1.1)         |
| 125% MT latency (ms)             | 10.1 (1.6)   | 10.1 (1.8)   | 10.1 (1.4)         | 10.3 (1.1)        |
| 140% MT latency (ms)             | 10.1 (1.5)   | 9.8 (1.5)    | 9.8 (1.5)          | 10.0 (1.5)        |
| Silent period duration (ms)      | 51.6 (7.6)   | 50.2 (9.1)   | 44.1 (8.5)         | 46.7 (9.6)        |
| Silent period latency (ms)       | 51.1 (4.9)   | 48.6 (11.3)  | 42.8 (7.6)         | 47.3 (13.7)       |

Mean values are accompanied by standard deviations in brackets. Data are reported for right/left side of the tongue when considering amplitudes, areas and latencies of recruitment curve data, silent period durations and latencies. Significant comparisons are reported in bold, marginal differences in italic; LH = left hemisphere, RH = right hemisphere.

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TMS EMG activity, when comparing the tongue right side versus the left one (Z = 2.845, P = 0.004; r = 0.858 - d = 0.429-). The main findings are summarized in Table 3, Table A in S1 File, Figs 2 and 3, and Fig B in S1 File.

Correlation analysis

Here only significant results (P<0.05) related with the principal aim of the manuscript will be reported (main relations between TMS, SSI-4 and BigCAT are here reported; remaining correlations are reported in S1 File), without reporting irrelevant correlations (e.g. correlations between motor thresholds and TMS data) or marginal correlations (P<0.1). When considering SSI-4, a positive relation with BigCAT is evident (r = 0.63; participants with higher DS severity perceive more negative attitudes toward speech situations). When considering TMS data, SSI-4 showed a positive relation with tongue AMT and SPT of the left hemisphere (AMT: r = 0.68; SPT: r = 0.69). Similarly, silent period durations of the right FDI muscle (i.e. when stimulating the left hemisphere) were positively related with SSI-4 (r = 0.68), and a positive relation was highlighted between SSI-4, bilateral AMT and SPT of FDI muscles (left hemisphere AMT: r = 0.62; right hemisphere AMT: r = 0.75; left hemisphere SPT: r = 0.62; right hemisphere SPT:
A positive relation was evident in all participants (DS and fluent speakers), between bilateral tongue SPd, obtained stimulating the left hemisphere, and the BigCAT (tongue right side: $r = 0.41$; tongue left side, $r = 0.43$). The main findings are summarized in Fig 4.

Discussion
Summary of findings
In the present work, the main findings suggest that longer silent period durations may be evident in DS, when recording from tongue muscles, even when no overt dysfluency is present and during no speech tasks, especially when stimulating the left hemisphere motor cortex. Moreover, DS had higher SPT in the left hemisphere with respect to the right one. A similar but less defined pattern is evident when considering AMT (see also preliminary cases report of Barwood et al. [36]) while MT (obtained during lower muscular activation) had a higher asymmetry in fluent speakers (left hemisphere more excitable than the right one), which was not evident in DS (compare with [39]). TMS data (tongue and hand muscles), such as motor thresholds and SPd, showed positive correlations with SSI-4, as well as with DS behavioral/cognitive indexes (some of which resulted more elevated in DS). A positive relation was evident in DS between left hemisphere AMT and SPT, recorded from the tongue, and SSI-4, as well as between left hemisphere silent period durations of the tongue and BigCAT (in all participants; BigCAT was positively related to SSI-4). Silent period durations of the right FDI (i.e. stimulating the left hemisphere) were positively related with SSI-4, and a positive relation was evident...
Fig 2. Silent period durations obtained in the stuttering and fluent speakers groups. Significant differences are indicated with an asterisk. Data are reported by considering also the right/left side of the tongue. LH = TMS administered on the left hemisphere; RH = TMS administered on the right hemisphere.

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Fig 3. Pre-TMS EMG data obtained in the two groups during sustained tongue contractions requested for the silent period TMS protocol. About 60–70% of maximal muscular activation was requested. Marginally significant comparisons are indicated with a circle. Data are reported by considering also the right/left side of the tongue. LH = TMS administered on the left hemisphere; RH = TMS administered on the right hemisphere.

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The delicate (dis)equilibrium between neural networks in DS

The present findings suggest that persistent DS is related to a delicate interplay between different cortical/intracortical mechanisms involved in the functioning of tongue motor cortex, thus influencing cortico-bulbar responses, even when no overt stuttering is evident. Motor thresholds and recruitment curve data point toward the confirmation of the presence of lower left hemisphere cortico-bulbar excitability in DS, in contrast to higher activity in homologue structures of the right one [12,13], with respect to fluent speakers. On the other hand, lower inhibitory responses in the left hemisphere of DS, when considering SPT, was also evident. This
index represents the functionality of intracortical networks (see below, silent period discussion section), and suggests the presence of a continuous interplay between excitatory and inhibitory motor mechanisms in DS, influencing the final speech and not strictly speech related motor output, likely looking for homeostasis (in this case, trying to avoid speech dysfluencies). Pre-TMS EMG differences (see also S1 File) may also be interpreted as being due to the presence of a lower (and more variable) activity in tongue motor regions of the left hemisphere with respect to the right hemisphere, in stuttering. Previous evidence on this is conflicting, possibly also showing no differences in EMG activity between DS and fluent speakers [37,39]. DS is related to an aberrant language brain lateralization, especially of motor outputs needed for correct speech execution [32,75,76,77]. These neural patterns may be related to mechanisms that help compensate for an aberrant transmission among inferior frontal regions and speech sensori-motor cortices [10,11]: this is supported by evidence showing an inverse correlation between right hemisphere neural activity and indexes of stuttering severity [78,79,80], as well as no evidence of this augmented activity in DS children [81]. In the end, this may be related to imbalanced wiring in DS [5]. As a consequence, motor planning/execution of motor tasks in speech muscular areas need the punctual integration of excitatory and inhibitory neural signals. Indeed, DS shows less and/or different neural changes during motor planning in both speech- and non-speech-related tasks in frontal, temporal, and sub-cortical regions [82,83]. The correct functioning of this system may be related to signaling and connections from left frontal regions [10,11,84], and also from the sub-cortical basal ganglia system [3,85].

**DS and impairments in white matter and in the basal ganglia system: relations with present findings and insights from previous TMS studies**

White matter abnormalities are indicated as one of the fundamental neural markers of DS [5,8,10,11,13]: they are widespread and evident in cortical and sub-cortical networks, comprising structures such as corpus callosum, superior longitudinal fasciculus, fibers of chorona radiata, cortico-spinal and cortico-bulbar tracts, frontal regions (e.g. premotor and sensorimotor regions), left angular gyrus, arcuate fasciculus, and thalamo-cortical circuits [5,8,81,85,86,87,88,89,90,91]. They are preferably evident in the left hemisphere [8,10,11,80,84,86,92]. With respect to the present findings, Sommer et al. [10] reported less white matter in the left operculum of DS in correspondence of the tongue and larynx sensori-motor representation (see also [81]). Similarly, Connally et al. [89] reported weaker connectivity in left cortico-bulbar and cortico-spinal tracts in DS (versus augmented connections in the right hemisphere), and reduced white matter in the bilateral arcuate fasciculus; higher stuttering severity was related with higher connectivity in the left cortico-bulbar fibers. Jäncke et al. [93] showed increases in white matter in right hemisphere networks in DS adults, comprising the inferior frontal gyrus and face/mouth motor representations. Watkins et al. [11] showed less white matter in bilateral premotor/motor regions in DS, as well as in fibers of inferior frontal regions that are involved in speech motor aspects (see also [80]); lower neural activity in sensorimotor and inferior frontal regions in DS was also reported. Chang et al. [80,81] showed that DS adults and children have lower white matter levels in the left superior longitudinal fasciculus, which transmits information to motor structures and left inferior frontal regions. Interestingly, Kemerdere et al. [94] suggest that the stimulation of the left frontal aslant tract induce stuttering in fluent speakers: the bilateral frontal aslant tract has an increased mean diffusivity in persistent DS, and a negative correlation between fluency and mean diffusivity in the left tract was evident. The frontal aslant tract connects inferior frontal regions with the supplementary motor complex, playing a role in speech motor organization [90,95]. DS related neural fiber impairments are also in direct relationship with sub-cortical neural targets, such as
basal ganglia. In fact, DS could be also viewed as a movement disorder related to basal ganglia dysfunction [3,4,23], and to dopaminergic hyperactivity in these regions [21,22]. Intracortical neural circuits, and consequently cortico-spinal/cortico-bulbar activity, may be influenced by dopamine modulation [96,97,98]. In fact, basal ganglia dysfunction in DS may influence cortical functioning by means of cortico-striatal-thalamo-cortical loops [19]. Stuttering severity and dysfluencies have been showed to be related with basal ganglia activity [5,11,99,100,101], before but not after fluency-inducing treatments [99]. Supplementary motor area (part of the cortico-striato-thalamo-cortical loop) [19] hyperactivity is consistently found in DS [5,13], which likely influenced motor cortex activity [102]. Considering its relation with basal ganglia dysfunction, stuttering may be considered as a timing impairment in neural networks that support volitional, self-paced (preferably speech-related) motor acts, relying on dysfunction of left hemisphere motor structures, with a sub-optimal connectivity of basal ganglia-thalamo-cortical and motor neural networks [83,92], also indirectly involving speech-related muscular districts.

Interestingly, present and previous TMS findings in DS are consistent with the above reported white matter abnormalities. Only few studies investigated DS by TMS. Moreover, previous work has often concentrated on non-speech-related aspects of DS motor system, because of the challenging methods needed to record TMS-evoked potentials from speech muscles. This led to the definition of DS as a more general motor impairment [25,31,35,38,42]. DS abnormal motor functioning is also evident when no overt stuttering is present, helping to disentangle “basic” neural excitability in DS, and its modulation during speech- or non-speech-related tasks [25,31,35,37,38,39,42]. Abnormal ratios of neural activity have been shown between left and right hand motor cortex [35] and with respect to fluent speakers, both during rest [31,42] or rhythm-related motor tasks [38]. Intracortical excitability seems to be normal in DS when considering hand motor representations [41,42], even if it responds to selective serotonin re-uptake inhibitors aimed at reducing stuttering [25]. On the other hand, abnormal patterns of excitability affect intracortical (and cortico-bulbar) networks of speech muscles in DS [36,37,39,40]: a weaker inhibition in the right hemisphere (associated with delayed inhibition of intracortical circuits in both hemispheres), and a bilaterally reduced facilitation was evident in DS, accompanied by steeper stimulus-response curves (preferably during muscular activation of the same districts), especially in the right hemisphere [36,39,40]. Moreover, the lack of left hemisphere facilitation of tongue MEPs during speech was evident, suggesting an asymmetry controlling speech motor planning in motor cortex [37]. Parts of these findings correlated with stuttering severity. The lack of a left hemisphere facilitation of MEPs during speech [37], as well as the evidence of lower neural activity in regions such as larynx motor cortex [5], may be related with pre-existent abnormal excitatory/inhibitory ratios in intracortical circuits modulating DS motor outputs, demonstrated here by findings related to silent period durations and SPT. Indeed, the present findings may be also viewed as the missing link of a more complex picture in which intracortical motor circuits play a central role, even in a more general stuttering “state” condition. Present correlations between stuttering indexes, AMT, SPT, and silent period durations of both hemispheres in speech- and not strictly speech-related muscular districts (see also [31]) support these suggestions. These observations may help to understand how different neural substrates may be mutually related to maintain the disorder, also in adulthood. One of the possible mechanisms of this relationship is highlighted in the next section.

**DS and cortical silent period: possible relations with basal ganglia and white matter abnormalities**

DS motor thresholds and recruitment curves have also been investigated in previous TMS studies in both speech (and not strictly speech-related) muscles with a certain level of agreement
Here, we extend those observations, considering that silent periods have been not extensively studied in DS, especially in DS speech muscles, such as the tongue. Silent period is considered a temporary suppression of voluntary muscle activity, after depolarization of motor neuronal cells by TMS. Mechanisms underlying silent period are still not completely understood (see [57]) and, thus, interpretation of results may be not simple. Silent period depends on both spinal and cortical mechanisms (likely interruptions of the cortical drive), resulting from the influence of intracortical inhibitory cells on the motor cortex [103,104]: it is an index of intracortical inhibition modulated by GABAergic activity of interneurons that synapse with neurons of pyramidal tracts [105]. In DS, it has been reported that not directly speech-related silent period may have a role in the effects of drugs that help to manage stuttering [25]. Rogić Vidaković et al. [40] showed differences in the silent period of DS hand muscles in both hemispheres. The present, and previous, findings [31] suggest that silent period may be differently related with indexes of stuttering severity in both speech- and not strictly speech-related muscles. As a consequence, this TMS index may have an important role in DS neurophysiology, likely representing a neural marker of DS intracortical inhibition and functioning, especially in the motor structures of left hemisphere (please note, that the different intracortical TMS indexes until now reported in DS have mainly been obtained from paired pulse protocols, with different evidence, such as for example the lacking of a clear correlation with stuttering severity; see [39,41,42]). Speculatively, the here reported lengthened silent period can be related, for example, to the consequences of a decrease in tonic excitation modulated by afferent pathways to motor cortex, as a result of the widespread white matter impairments evident in DS, favoring a prolonged GABA-mediated inhibition on pyramidal cells. It might be related to a general lack of inhibition of neurons in motor cortex, again resulting in an over-activation of GABAergic interneurons (also as a consequence of the DS abnormalities in basal ganglia-thalamo-cortical circuits), favoring plastic neural mechanisms looking for a sustainable equilibrium between excitatory/inhibitory signals in the brain. Interestingly, a lengthened SP in the affected hemisphere of stroke patients was associated with motor deficits such as movement initiation and inability to maintain constant force levels, fitting with present evidence, while clinical improvements were related to shortened silent periods [106], also in DS [25]. Classen et al. [106] suggested that motor dysfunctions may be related to the hyperactivity of inhibitory mechanisms in the cortex, and an increased silent period may result from damage to different input pathways of the motor cortex, as it seems to be the case in DS. A diminished and/or augmented activity of inhibitory interneuronal systems may result in abnormalities of the (motor) neuronal networks, considering that they should target and modulate (motor) neural activity (by means of lateral inhibition), as seems to be the case in a series of other basal ganglia related motor disturbances such as Parkinson’s Disease [96], Tourette’s Syndrome [107], and dystonia [108].

DS sub-groups from a neurophysiologic point of view?

Present correlation analyses showed that pre-TMS EMG data (see S1 File) were poorly correlated in DS with respect to fluent speakers, confirming higher variability in DS when a motor task is required [109,110,111]. In general, differences with previous reports may be due to methodological issues or related to the presence of different neurophysiological profiles in persistent DS, i.e. in adults that modeled their brain patterns during attempts to manage dysfluencies [14,15]. Indeed, when recording from FDI muscles, a positive relation is evident between stuttering severity, left hemisphere silent period duration, and bilateral AMT/SPT: higher stuttering degree was associated with higher activity in intracortical inhibition networks of left hemisphere and with particular excitatory/inhibitory ratio of neural activity also in motor
regions not directly involved in speech control. This supports Busan et al.'s [31] observations that DS males showed a negative correlation between silent period durations obtained when stimulating right hemisphere FDI representation, and stuttering severity. This supports the idea that different DS groups may show slight differences in neurophysiologic profiles, pointing toward a generalized left hemisphere motor inhibition, counter-parted by higher right hemisphere activity in homologue brain regions. Moreover, it may sustain the idea that DS may be the only overt symptom of a more general motor disorder (see [31]). In general, different possible DS sub-divisions have been proposed, on the basis of neural and genetic factors [112], effect of pharmacological agents (see [3]), neural and genetic differences between DS males and females [31,32,33,113], anxiety levels, and co-morbidities [114]. Indeed, the present findings sustain the idea that DS may be related to higher levels of emotional lability [115,116], higher tendencies toward psychophysiological disturbances, and higher levels of phobia, likely related to DS social implications (see [117,118]; see also further additional correlations reported in S1 File). In fact, here we report that a negative attitude toward speech situations is evident in DS (please note also the positive correlation between SSI-4 and BigCAT scores; see [119]); BigCAT (negative speech attitudes in DS) also positively correlated with bilateral AMT and SPT obtained from hand muscles (see S1 File).

Methodological issues and limitations of the study

The present work has some limitations. For example, recordings from the tongue are methodologically challenging and uncomfortable, and as a consequence a limited amount of data can be obtained. In fact, TMS indexes, especially when registering from speech related muscles, may be influenced by peripheral activity and muscular activation. Moreover, the here reported qualitative differences in correlations between groups (for example in pre-TMS EMG; see S1 File), sometimes have to be cautiously interpreted, considering the different sample sizes. Indeed, difficulties in recruiting a homogenous sample of DS participants should always be considered [120]. When considering DS, especially in adulthood, it is always difficult to disentangle between causal neural mechanisms (perhaps related to factors that predispose to the development of the disturbance), long-life stuttering (and its neural consequences), and compensatory neural patterns, both at cortical and spinal levels. This is quite common in movement disorders, such as Parkinson’s Disease or dystonia. In fact, neural activations may be related to mal-adaptive plasticity as a consequence of a life of stuttering. This might be especially true when considering that abnormal/defective sensory inputs (as for example those related to stuttering episodes [121]) may be related to deviant neural motor activations [122], acting also on neural plasticity. This may finally result in a cascade of neural changes that help to maintain the disorder. For these reasons, the present findings cannot automatically be generalized to DS in childhood.

Conclusions and Future Perspectives

The present work contributes to defining the aberrant balance between motor excitatory and inhibitory mechanisms in tongue representation of the primary motor cortex of DS adults, even when no overt dysflueny is occurring (i.e. in a stuttering “state” condition) [5,13]. It generally confirms that motor (excitatory/inhibitory) balance is a fundamental issue when considering neural functioning in motor disorders [123]. It has been conducted during no concurrent speech tasks, to better investigate some previously undefined indexes of intracortical motor excitability in DS (tongue silent period). The use of a “basic” functional modality opens the way toward more complex TMS studies aimed at investigating modulations of aberrant indexes during complex speech and motor tasks. The present findings confirm that DS is a motor
disturbance characterized by an abnormal ratio between excitatory and inhibitory neural circuits, especially in the speech motor networks of the left hemisphere, but suggest tongue silent period as a novel neural marker for persistent DS, also in relation to here reported relations with stuttering severity. In fact, silent period may reflect neural oscillations in cortical (motor) systems, reflecting excitatory/inhibitory ratios [124]. Dysfluency could be only the more evident symptom of a subtle motor syndrome, characterized by the constant presence of an aberrant excitability of motor system and an aberrant interplay between its components. The present data will be useful to help define more focused neural targets for disentangling new treatment options, such as non-invasive neuro-modulation or more focused pharmacological interventions, targeting and modulating here reported indexes of abnormal functioning in persistent DS.

Supporting Information
S1 File.
(PDF)

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