Electrophysiological evidence for pre-attention information processing improvement in patients with central hemiplegic after peripheral nerve rewiring: a pilot study

Tie Li1, Xu-Yun Hua1, Mou-Xiong Zheng1, Yu-Lan Zhu2, Yan-Qun Qiu3, Yun-Dong Shen1, Jian-Guang Xu1, Yu-Dong Gu1 & Wen-Dong Xu1,3

Central neurologic injury (CNI) causes dysfunctions not only in limbs but also in cognitive ability. We applied a novel peripheral nerve rewiring (PNR) surgical procedure to restore limb function. Here, we conducted a prospective study to develop estimates for the extent of preattentive processes to cognitive function changes in CNI patients after PNR. Auditory mismatch negativity (MMN) was measured in CNI patients who received the PNR surgery plus conventional rehabilitation treatment. During the 2-year follow-up, the MMN was enhanced with increased amplitude in the PNR plus rehabilitation group compared to the rehabilitation-only group as the experiment progressed, and progressive improvement in behavioural examination tests was also observed. Furthermore, we found a significant correlation between the changes in Fugl-Meyer assessment scale scores and in MMN amplitudes. These results suggested that PNR could affect the efficiency of pre-attention information processing synchronously with the recovery of motor function in the paralyzed arm of the chronic CNI patients. Such electroencephalographic measures might provide a biological approach with which to distinguish patient subgroups after surgery, and the change in MMN may serve as an objective auxiliary index, indicating the degree of motor recovery and brain cognitive function.

Central neurologic injury (CNI) is a major contributor to physical disability and can affect both adults and children. In addition to motor-sensory functional impairments, patients frequently suffer from severe neurocognitive dysfunction, including damaged motor execution abilities and decreased automatic information processing of the injured cortex. The disorder mainly affects patients’ muscle tone, movement, and daily living skills with varied levels of limb dysfunction and cognitive impairment1. For central hemiplegic patients, dysfunctions of sensory and motor information processing accompany limb dysfunction2,3. These deficits of cognitive ability are potentially related to the difficulty of motion execution4, which may be an important obstacle that influences rehabilitation after CNI.

For hemiplegic patients, we applied a new surgical procedure of peripheral nerve rewiring to restore the motor function of the affected upper extremity. During the procedure, the nerve fibres that originally innervated the intact upper extremity were severed and transferred to the C7 nerve of the paralyzed arm5. This crossed peripheral nerve (contralateral C7, CC7) then connected the paralyzed upper extremity with the contra-lesional (ipsilateral to the paralyzed arm) hemisphere. The control of the contra-lesional hemisphere over the paralyzed arm led to a significant improvement in motor function.

Received: 1 November 2016
Accepted: 28 June 2017
Published online: 31 July 2017

1Department of Hand Surgery, Huashan Hospital, Shanghai Medical College, Fudan University, Shanghai, China.
2Department of Rehabilitation, Huashan Hospital, Shanghai Medical College, Fudan University, Shanghai, China.
3Department of Hand and Upper Extremity Surgery, Jing’an District Central Hospital, Shanghai, China. Tie Li and Xu-Yun Hua contributed equally to this work. Correspondence and requests for materials should be addressed to W.-D.X. (email: wendongxu@fudan.edu.cn)
upper extremity was enhanced, promoting brain plasticity. In the latest follow-up visit 2 years after surgery in the preliminary cases, the results were quite encouraging. However, the relationship between motor recovery and cognitive changes remains a matter of debate, and more specifically, it is also unclear whether the brain plasticity process after peripheral nerve rewiring affects cognitive function.

Event-related potential (ERP) testing is a useful tool for the on-line examination of both normal and impaired information processing, and such research may contribute to the development of theoretical models of interaction between motor and cognitive function. The adequate processing of task-irrelevant deviant events is an important cognitive function for human survival and can be reflected by the mismatch negativity (MMN) component of ERPs. The precise neural function that is reflected by the MMN is controversial. Early studies suggested that the MMN reflects a preattentive, memory-based auditory discrimination process. However, it has recently been suggested that the MMN may reflect a much simpler neural adaption process.

Particularly relevant to the present study are a growing number of CNI (stroke, cerebral trauma, and cerebral palsy) studies that have investigated preattentive automatic processing by recording the auditory MMN. For example, Deouell et al. used the MMN to investigate the discrimination of different sound features in patients with right hemisphere damage that caused a neglect to the left hemi-field and found that an early deficit in detecting changes in the environment hampers the involuntary triggering of attention in these patients. Ilvonen et al. investigated cognitive function in left-hemisphere-stroke patients and found a significant correlation between the change in the duration of MMN amplitude and speech-comprehension performance after stroke, suggesting that the MMN can be used as an index for the recovery of auditory discrimination.

To improve the cognitive function of CNI patients, psycho-physiological and cognitive behavioural training methods have been widely used. For example, there is evidence that frequent exposure to music is able not only to increase evoked potentials and gating in the auditory cortex but also to enhance learning, memory and neural plasticity in the animal brain. Moreover, multimodal sensory stimulation has also been shown to reduce lesion volume and to improve cognitive and motor recovery after brain injury in rats. Similarly, it has been reported that merely listening to music and speech after neural damage can induce long-term plastic changes in early sensory processing, which may facilitate the recovery of higher cognitive functions. However, it is unclear whether the cortical plastic changes that occur after peripheral nerve rewiring are able to affect cognitive function. As described above, in humans, MMN can be used to measure crucial cortical plastic changes caused by training or remediation programmes. In the present study, during the recovery process of central hemiplegic patients after PNR, brain plasticity is an important component of functional restoration as well as peripheral nerve regeneration. We hypothesize that the coordination of the whole paralyzed upper extremity is improved in well-recovered patients. This improvement of motor control must inevitably involve the recovery of cognitive function. To test this hypothesis, we analysed the auditory MMN in patients after PNR with a follow-up period of 2 years.

Results
Attenuation in the MMN amplitude was observed in both the PNR plus rehabilitation and the rehabilitation-only central hemiplegic patients. Figure 1 shows the averaged MMN waveforms of the Fz, Cz and Pz electrodes. The mean amplitudes of the MMNs are presented in Table 1.

**MMN Significance.** After visual inspection of the waveforms, we performed three-way repeated measures analysis on the individual Group (PNR, rehabilitation and control) * Time (pre, post 6 months, post 1 year, post 18 months, post 2 years) * Electrodes (Fz, Cz and Pz), and the analysis yielded the following conclusions regarding the main effects of the three factors.

In the ANOVA analysis of MMN amplitudes, there was a significant main effect of Group $F(2, 54) = 871.2, p < 0.001, \eta^2 = 0.970$, revealing that the MMN in normal healthy controls ($−1.2\mu V$) was larger than that in the PNR plus rehabilitation group ($−0.62\mu V; p < 0.01$), which was larger than that in the rehabilitation-only group ($−0.50\mu V; p < 0.05$). These results show that the condition of the PNR plus rehabilitation group was significantly better than that of patients treated with rehabilitation only, but both groups were still different from the normal control group.

In the ANOVA analysis of MMN amplitudes, there was a significant main effect of time $F(2, 53) = 1691.4, p < 0.001, \eta^2 = 0.985$. Further analyses revealed that the amplitude of MMN was significantly higher at 18 months ($−0.83\mu V; p < 0.01$) and 2 years ($−0.85\mu V; p < 0.01$) after PNR than before the procedure ($−0.75\mu V; p < 0.01$). In the ANOVA analysis of MMN amplitudes, there was a significant main effect of electrode site $F(4, 51) = 10.7, p = 0.000, \eta^2 = 0.456$, showing that the MMN in Fz ($−1.2\mu V$) was larger than that in Cz ($−0.8\mu V; p < 0.01$), which was larger than that in Pz ($−0.3\mu V; p < 0.01$).

Interestingly, we found a significant two-way interaction of Group * Time, $F(8, 216) = 4.2, p < 0.001, \eta^2 = 0.134$. Further analysis of this interaction showed that there was a significant time effect for the PNR plus rehabilitation group ($p < 0.05$), whereas the time effect was not significant for either normal controls ($p = 0.17$) or rehabilitation-only group ($p = 0.86$). Although the group effect was evident for each time point ($p < 0.005$ in all cases), the MMN found in the PNR plus rehabilitation group was larger than that in the rehabilitation-only group only in the 4th ($−0.71\mu V$ and $−0.54\mu V$ for PNR plus rehabilitation and rehabilitation groups, respectively; $p < 0.001$) and 5th ($−0.75\mu V$ and $−0.54\mu V$ for PNR plus rehabilitation and rehabilitation groups, respectively; $p < 0.01$) time points. No other effects reached significant level ($p > 0.1$ in all cases).

For the MMN components, the MMN peak was not easily discernible at all sites and in each condition in many participants, the analysis was based on the mean amplitude calculated between control and oddball MMNs.
Behavioural Assessments. The Fugl-Meyer (FM) Assessment Scale scores for the two latest sessions (18 months and 2 years) showed significantly better performance in the PNR plus rehabilitation group (43.95 ± 4.98, and 46.68 ± 6.84) than the rehabilitation only group (P ≪ 0.05; group t-test). Further analyses revealed that the scores were significantly higher at 18 months (29.68 ± 4.27, 29.79 ± 4.04, P ≪ 0.05; paired t test) and 2 years (P < 0.05; paired t-test) than at pretreatment (29.95 ± 3.06, baseline measurement) and other times in the PNR plus rehabilitation group (Fig. 2C). The motor and sensory function of the unaffected hands remained normal.

The average Modified Ashworth Scale (MAS) score in the PNR plus rehabilitation group was significantly lower than that of the rehabilitation-only group. It suggested that spasticity in all joints was reduced to a lower level in the PNR plus rehabilitation group. However, further investigation of this observation is not included in the present study.

In addition to the pairwise comparisons, we calculated the Pearson correlation coefficients between the behavioural assessment results and the amplitudes of the MMN components. There was a significant correlation between the changes in the Fugl-Meyer Assessment Scale scores and in the amplitude of the MMN (r = 0.73, P < 0.05). No correlations were identified between the FM scores and the latencies of the MMN components (p > 0.1 in all cases). Additionally, there were no correlations between the MAS scores and MMNs (p > 0.1 in all cases).

Nerve Conduction Tests of the Regenerated Contralateral C7. At the 18 months to 2-year follow up after surgery, compound muscle action potentials (CMAP) from forearm extensor on the affected side could be elicited by stimulating the graft nerve at the neck in all 19 subjects of PNR plus rehabilitation group. The mean
The latency of CMAP of forearm extensor muscle was 9.83 ± 0.61 ms, and the mean amplitude was 0.67 ± 0.11 mV. The result reflected a connection of peripheral neural pathway between the unaffected and affected upper extremities.

### Table 1

|                | PNR plus rehabilitation group | Rehabilitation-only group | Control group |
|----------------|------------------------------|---------------------------|---------------|
| Fz Baseline    | −0.77(±0.17)                 | −0.75(±0.23)              | −1.99(±0.40)  |
| 6 months       | −0.81(±0.14)                 | −0.78(±0.25)              | −1.99(±0.54)  |
| 1 year         | −0.84(±0.12)                 | −0.78(±0.25)              | −1.96(±0.46)  |
| 18 months      | −1.12(±0.28)*                | −0.80(±0.14)              | −2.00(±0.45)  |
| 2 years        | −1.18(±0.32)**               | −0.85(±0.15)              | −2.02(±0.47)  |
| Cz Baseline    | −0.53(±0.16)                 | −0.55(±0.12)              | −1.33(±0.37)  |
| 6 months       | −0.57(±0.23)                 | −0.57(±0.18)              | −1.26(±0.20)  |
| 1 year         | −0.66(±0.15)                 | −0.51(±0.23)              | −1.24(±0.25)  |
| 18 months      | −0.75(±0.11)*                | −0.59(±0.21)              | −1.31(±0.18)  |
| 2 years        | −0.81(±0.19)**               | −0.53(±0.24)              | −1.42(±0.25)  |
| Pz Baseline    | −0.24(±0.04)                 | −0.24(±0.04)              | −0.34(±0.12)  |
| 6 months       | −0.26(±0.03)                 | −0.24(±0.05)              | −0.32(±0.09)  |
| 1 year         | −0.24(±0.04)                 | −0.25(±0.03)              | −0.32(±0.12)  |
| 18 months      | −0.26(±0.11)                 | −0.24(±0.05)              | −0.33(±0.10)  |
| 2 years        | −0.25(±0.08)                 | −0.24(±0.05)              | −0.33(±0.10)  |

Table 1. Mean Amplitudes (µV) (M ± SD) of mismatch negativity (MMN) at Fz, Cz, and Pz in central neurological injury (CNI) Patients and Control Subjects. The superscript (*) (P < 0.05) denotes significant differences between the surgery and rehabilitation-only groups. The superscript (**) (P < 0.05) denotes significant differences between the pretreatment and posttreatment values.

Figure 2. (A). After modification, the motor fibers from the contralateral C7 nerve (health side) allows external information input to the paralysed hand. This particular artificial neuropathway could induce interhemispheric plasticity from motor-sensory feedback during the recovery period after the peripheral nerve rewiring (PNR) procedure. (B) The preattentive information processing reflected by the mismatch negativity (MMN) in central neurological injury (CNI) patients after PNR was nearly normal. (C) Percentiles of Fugl-Meyer assessment scale scores with mean and SD. *Significant differences between PNR and rehabilitation groups, P < 0.05; **Significant differences between pretreatment and posttreatment values, P < 0.05.
Discussion

The major finding of this study was that the generation of the auditory MMN is clearly abnormal in central hemispheric patients, indicating cognitive dysfunction in all our patients irrespective of whether they received the surgical treatment. The patients who underwent PNR surgery showed a gradual gain over the rehabilitation-only patients in the peak MMN amplitude during the 2-year follow-up, suggesting improved cognitive function together with the observed improvement in behavioural assessments.

The MMN has been used in cognitive neuroscience research and clinical science research. This rapidly measurable response is highly stable over time and may reflect a predominantly automatic process, allowing investigators to interrogate the earliest stages of information processing free of conscious effort and motivational artefacts that may confound the assessment of higher cognitive operations in clinical populations such as traumatic brain injury (TBI) patients. To our knowledge, the present study is the first to directly measure auditory MMN as a reflection of cognitive function in central hemispheric patients after peripheral neural pathway transposition. Compared with healthy normal control participants, the amplitudes of MMN were significantly lower in CNI patients, both before and after PNR surgery. Consistent with the present findings, MMN decreases also occurs in brain injury and stroke patients. For example, Ilvonen et al. found that patients showed attenuated MMN amplitudes for tone duration and tone frequency changes to harmonically rich tones presented to the right ear at 4 and 10 days following the onset of left-hemisphere stroke. These investigators found that the attenuation of MMN amplitude was significantly correlated with impairment of cognitive ability after CNI. Even if these patients experienced plastic changes due to training or remediation programmes, their cognitive function could hardly be restored to the normal level. A classical idea is that damage to the cerebral cortex could indirectly lead to lesions of fornix, amygdala, hippocampus, mammillary bodies, and cingulate gyres, as well as to frontal cortex nerve cell degeneration, neuronal depigmentation, and apoptosis. However, in the brain regions include strong connections with recognition, information storage, learning and integration functions.

Importantly, we observed enhanced MMN in the PNR plus rehabilitation group compared with the rehabilitation-only group. Similarly, a study by Kaipio et al. also showed changes in MMN amplitude in closed head injury and stroke patients. There was evidence that changes in MMN were correlated with the recovery of speech comprehension in aphasic stroke patients. Furthermore, Ilvonen et al. determined that the recovery of cortical auditory discrimination in aphasic, left-hemisphere-stroke patients, was significantly correlated with the change in the duration of MMN amplitude and speech comprehension performance from 10 days to 3 months after stroke, suggesting that the MMN can be used as an index of the recovery of auditory discrimination. In contrast to the MMN results of the follow-up, enhanced amplitude and decreased latency were observed in the present surgical interventional group and are well documented in the literature. Recently, evidence has emerged suggesting that external environmental stimuli play an important role in shaping our brain. Särkämö et al. reported that merely listening to music and speech after neural damage can induce long-term plastic changes in early sensory processing, which may facilitate the recovery of higher cognitive functions. In that study, the MMN frequency increased significantly for the patients exposed to both music and audio books than in the control group during the 6-month post-stroke period. Moreover, changes in the frequency MMN amplitude correlated significantly with the behavioral improvement of verbal memory and focused attention induced by music listening. However, most of the above studies are based on external stimuli, whereas, in the present study, the enhanced MMN in the PNR plus rehabilitation group suggests that, with the recovery of motor function after surgery, cognitive ability could also be enhanced synchronously, as reflected by the enhanced MMN related to preattentive change detection.

Although the above data provides evidence for the improvement of cognitive function after PNR, the mechanism underlying this improvement is easily explained. As mentioned above, peripheral nerve rewiring artificially establishes a direct peripheral-central connection from the internal neuropathways to the contralateral motor cortex, and could effectively rescue the motor and sensory function of the affected upper extremity after CNI by harnessing the potential of the contralateral cortex. The regeneration of the donor nerve fiber from the contralateral C7 nerve could promote the restructuring of the neural network around the motor areas and other functional region in the brain, and could also stimulate the brain to generate synapses from neurons on the healthy side of the brain for the functional reconstruction of brain cells.

Following PNR, an additional neural pathway was established between the contra-lesional hemisphere and the paralyzed upper extremity (Fig. 2A). Conceptually, it seems reasonable to assume that the motor and sensory feedback to both hemispheres was dramatically promoted, facilitating reorganization of the brain. From a cognitive point of view, the reorganization generates synergy between the motor cortex and the surrounding areas, which can significantly improve the processing of information from external events. In this process, the contralesional hemisphere could mobilize more effective resources, allowing improvements in information recognition, coding, analysis, and integration. Most importantly, additional efficient sensory-motor projections from the paralyzed hand to the ipsilesional hemisphere were reconstructed, as indicated by the enhanced behavioural performance of the paralysed hand at the 2-year follow up (Fig. 2C). Moreover, the increase of motor fibers from the contralateral C7 nerve subsequently promoted behaviour and increased the external information input from the paralysed hand. This particular artificial neuropathway clearly has the potential to induce interhemispheric plasticity from motor-sensory feedback during the recovery period after PNR surgery. It might improve the automatic information processing of external input by the brain (Fig. 2A), as in the case of the enhanced MMN in the present study (Fig. 2B).

It is generally accepted that the potential mechanism of cortical reorganization after stroke is the enhancement of functional connectivity within the brain network. For instance, Schulz et al. investigated task-related effective connectivity between ipsilesional parietal regions and key frontal motor areas using fMRI in well-recovered stroke patients. It showed that in addition to excellent motor recovery, the coupling pattern of the parietofrontal network was near-normal. Similarly, by electroencephalographic (EEG) examination, Wu et al. demonstrated a
Table 2. Group Sample Sizes at Follow-ups. Education: the average number of years of education; disease course: time from the onset of central hemiplegia until recruitment to this study to receive an intervention of PNR or rehabilitation; LHD: left hemisphere damage; RHD: right hemisphere damage; others: other disease of central neural injury, e.g. encephalitis, intracranial tumour.

| Group                        | Gender | Education (year, M ± SD) | Disease course (year, M ± SD) | Lesion side | Pathogenesis |
|------------------------------|--------|--------------------------|-------------------------------|-------------|--------------|
| PNR plus rehabilitation group (n = 19) | 13 | 9.8 ± 3.3 | 14.1 ± 8.2 | 12 | Stroke |
|                             | 6      | 6.1 ± 3.9 | 15.7 ± 8.7 | 5 | Cerebral palsy |
|                             | 11     | 11.1 ± 1.6 | 15.7 ± 8.7 | 4 | Trauma |
|                             | 11     | —           | —             | — | Others |
| Rehabilitation-only group (n = 19) | 12 | 10.2 ± 2.1 | 15.7 ± 8.7 | 11 | Stroke |
|                             | 7      | —           | —             | 5 | Cerebral palsy |
|                             | 1     | —           | —             | 7 | Trauma |
|                             | 8      | —           | —             | 3 | Others |
| Control group (n = 19)      | 11     | 11.1 ± 1.6 | 15.7 ± 8.7 | — | Stroke |
|                             | 11     | —           | —             | — | Cerebral palsy |
|                             | 11     | —           | —             | — | Trauma |
|                             | 11     | —           | —             | — | Others |

consistent relationship between ipsilesional M1-premotor coherence and Fugl-Meyer score at each of four examinations spanning the 28 days of therapy. Such research has increased our understanding of the parietofrontal network of the ipsilesional hemisphere as a prominent circuit involved in plastic changes after stroke. Therefore, cortical reorganization may be the outcome of network changes and the brain network between the ipsilateral hemisphere and surrounding areas might play an important role in the motor recovery of CNI patients after PNR surgery. Further study of this issue is warranted.

Although this work is based on a small sample of subjects, our findings have garnered the attention of palliative care experts. However, the small sample size may also explain some of our surprising results, namely, the apparent lack of correlation of event-related potentials and clinical functional recovery with cognitive function.

The present findings indicate that PNR affects the efficiency of preattentive information processing synchronously with the recovery of motor function in the paralysed arm. It also supports the hypothesis that interhemispheric information processing is required to detect a novel incoming external stimulus, providing insight into the pathophysiology of motor recovery, cognition, and behavioural symptoms. Additionally, the present data demonstrates the use of auditory MMN recordings for CNI patient assessment.

Methods

Subjects. The experimental group subjects in this study were all clinically diagnosed with central hemiplegia and were in the chronic stage of motor function recovery (more than 8 months after the onset of CNI). The lesions were documented by magnetic resonance imaging, which was separate from this study. Excluded from the study were patients with neurological diseases other than stroke or with neuropsychiatric or neuropsychological deficits that could potentially compromise their ability to provide informed consent or ensure compliance during the experiments.

The experimental group in this study comprised two different treatments. Nineteen hemiplegic subjects in the PNR plus rehabilitation group received the PNR surgical procedure and post-operative rehabilitation; the other 19 patients were recruited from the rehabilitation department at the same hospital based on the above inclusion criteria and received conventional rehabilitation therapy only. There were no significant differences in the age, sex, disease course, or disease severity of the subjects in these two groups (Table 2). In addition, the study included a normal control group that consisted of 19 healthy age- and gender-matched hospital personnel and patients’ family members who volunteered to participate in this study. The control group did not significantly differ from the patients in nationality, gender, or age. The follow-up duration was 6–24 months after the recruitment. The disease course of the subjects was 14.1 ± 8.2 years in the PNR plus rehabilitation group and 15.7 ± 8.7 in the rehabilitation group. The demographic, clinical and lesion data of each group are presented in Table 2.

All subjects were right-handed. Audiometric testing was used to ensure that all participants had normal hearing and could detect 40-dB tones at 1000 Hz (mean ± SD = 12.8 ± 6.1 dB). This research was approved by the Institutional Review Board (IRB) of Huashan Hospital, Fudan University, China. All methods were performed in accordance with the approval guidelines and regulations, and the participants signed informed consent in accordance with the IRB approval stipulations.

Stimuli. Ten sinusoidal tones were generated, starting at 500 Hz and increasing in frequency by 10% with each step, resulting in a tonal succession of 500 Hz, for 500, 550, 605, 666, 732, 805, 886, 974, 1072, and 1179 Hz. The tones were 50 ms long including 5-ms rise and 5-ms fade-out times. The intensity of all stimuli (standard and deviants) was 70 dB (sound pressure level).

Procedure. The subjects were comfortably seated and instructed to watch a silent movie on a computer monitor positioned 110 cm in front of them and to ignore the auditory stimulation. Tones were sequentially presented binaurally over headphones at a sound pressure level (SPL) of 70 dB with a stimulus onset asynchrony of 500 ms. There were three blocked conditions, (a) descending deviant (500 Hz) with a 550 Hz standard, (b) ascending deviant (1179 Hz) with a 1072 Hz standard and (c) control sequence, consisting of all 10 tones. Block sequence was counterbalanced across participants. Each block consisted of 1,500 trials. In oddball blocks, deviants occurred among the standards with a relative frequency of 0.1 in a pseudorandomized fashion with the constraint that two deviants could not occur in direct succession. Deviants occurred at the endpoints of the tonal succession, the standards being their closest neighbors. In the control block, stimuli were presented equiprobably (relative frequency = 0.1) in a pseudorandomized sequence such that the set of 10 tones was multiply presented in sequence, each time in a random order, and the repetitions of tones was avoided (Fig. 3).
This procedure resulted in physically identical tones used as controls, presented in the control block, and deviants, presented in the oddball blocks, each with an equal frequency of occurrence within a block. Thus, the only difference between the two presentations was that neurons that specifically responded to the frequency of controls and deviants were less refractory for controls than for deviants, because they were presented among nine other control stimuli with a greater frequency separation in the control block, than the standard-deviant separation in the oddball block. Each experimental session lasted approximately 1 hr, including data acquisition and electrode application and removal.

EEG recording. The EEG signal was recorded while the participants were watching a self-selected, subtitled silent film. Participants were instructed to ignore the acoustic stimuli. To control the level of attention, they were asked to answer some questions about the content of the film at the end of the experiment, including the dramatis personae, scene, and content of incident. Participants were to be excluded if they were unable to answer the question, as this indicated a lack of focus on the film. No participants were excluded for this reason.

Continuous EEG activity was collected with a Synamps 2 Amplifier for 9 electrode sites from 64 channels, F3/z/4, C3/z/4, and P3/z/4, according to the extended international 10–20 system. The reference electrode was placed on the tip of the nose. A vertical electrooculogram (EOG) was recorded from the right eye by supra- and infra-orbital electrodes, and horizontal EOG was recorded from electrodes on the outer canthi of both eyes. EEG and EOG signals were amplified from 0.1 Hz to 100 Hz at a sampling rate of 500 Hz. The electrode impedance was less than 10 kΩ throughout the experiment.

After EOG artifact correction, the EEG was segmented in epochs of 500 ms, timed from stimulus onset, and also included a 100-ms pre-stimulus baseline. Epochs contaminated with artifacts greater than ±100 µV were rejected before the data were averaged. The EEG segments (at least 100 trials for each condition) were averaged separately for 150-ms and 50-ms stimulus conditions, and the averaged ERPs were smoothed through a low-pass digital filter at 30 Hz (24 dB/oct).

Statistical analyses were performed using the software Statistical Package for the Social Sciences, version 18.0 for Windows. The mean amplitudes of auditory MMNs were measured for the 100–250 ms time windows after stimulus onset. The amplitude differences between the groups and within the patient group between sessions were compared using repeated measures ANOVAs. Electrodes Fz, Cz, and Pz, where the MMN is largest, were included in the analyses. Statistical probability from the ANOVAs was corrected using the Greenhouse–Geisser procedure. A p-value of 0.05 was considered as statistically significant.

Behavioural Assessments. Several measures of motor recovery (reflex activity, flexor synergy, extensor synergy, movement combining synergies, movement out of synergy, normal reflexes, wrist, hand, coordination and speed) were assessed on the Fugl-Meyer (FM) Assessment Scale. The Modified Ashworth Scale (MAS) was used to assess spasticity. The correlations between the changes in the MMN amplitude and in the behavioural performance were assessed using the Pearson correlation coefficient.

Nerve Conduction Test of the Regenerated Contralateral C7 Nerve. At the 18 months to 2-year follow up after surgery, a nerve conduction test was performed to confirm the regeneration of the contralateral nerve.
C7 nerve. The stimulator was placed on the graft nerve at the neck, on the side of the unaffected upper extremity. Recording surface electrodes were placed over the forearm extensor on the affected side. The width of the stimulation pulse was 0.5 to 1.0 ms and the intensity varied from 70 to 100 mA.

References
1. Odding, E., Roebroeck, M. E. & Stam, H. J. The epidemiology of cerebral palsy: incidence, impairments and risk factors. Disability and rehabilitation 28, 183–191, doi:10.1080/09638280500158422 (2006).
2. Marshall, S. C. et al. Predictors of driving ability following stroke: a systematic review. Top Stroke Rehabil 14, 98–114, doi:10.1310/txr1401-98 (2007).
3. Denes, G., Semenza, C., Stoppa, E. & Lis, A. Unilateral spatial neglect and recovery from hemiplegia: a follow-up study. Brain: a journal of neurology 105(Pt 3), 543–552 (1982).
4. Deouell, L. Y., Bentin, S. & Soroker, N. Electrophysiological evidence for an early(pre-attentive) information processing deficit in patients with right hemisphere damage and unilateral neglect. Brain: a journal of neurology 123(Pt 2), 353–365 (2000).
5. Xu, W. D., Hua, X. Y., Zheng, M. X., Xu, J. G. & Gu, Y. D. Contralateral C7 nerve root transfer in treatment of cerebral palsy in a child: case report. Microsurgery 31, 404–408, doi:10.1002/micr.28877 (2011).
6. Hua, X. Y. et al. Contralateral peripheral neurotization for hemiplegic upper extremity after central neurologic injury. Neurosurgery 76, 187–195, discussion 195, doi:10.1227/neu.0000000000000590 (2015).
7. Nyman, G. et al. Mismatch negativity (MMN) for sequences of auditory and visual stimuli: evidence for a mechanism specific to the auditory modality. Electroencephalogr Clin Neurophysiol 77, 436–444 (1990).
8. Ritter, W. et al. Event-related potentials to repetition and change of auditory stimuli. Electroencephalogr Clin Neurophysiol 83, 306–321 (1992).
9. Schröger, E. & Wolff, C. Fast preattentive processing of location: a functional basis for selective listening in humans. Neuroscience letters 232, 5–8 (1997).
10. Nätänen, R., Schröger, E., Karakas, S., Tervaniemi, M. & Paavilainen, P. Development of a memory trace for a complex sound in the human brain. Neuroreport 4, 503–506 (1993).
11. Ivonen, T. M. et al. Auditory discrimination after left hemisphere stroke: a mismatch negativity follow-up study. Stroke; a journal of cerebral circulation 34, 1746–1751, doi:10.1161/01.stra.0000087836.26328.3b (2003).
12. Ivonen, T. M. et al. The processing of sound duration after left hemisphere stroke: event-related potential and behavioral evidence. Psychophysiology 38, 622–628 (2001).
13. Engineer, N. D. et al. Environental enrichment improves response strength, threshold, selectivity, and latency of auditory cortex neurons. Journal of neurophysiology 92, 73–82, doi:10.1152/jn.00020.2004 (2004).
14. Percaccio, C. R. et al. Environmental enrichment increases paired-pulse depression in rat auditory cortex. Journal of neurophysiology 94, 3590–3600, doi:10.1152/jn.00433.2005 (2005).
15. Angelucci, E. et al. Investigating the neurobiology of music: brain-derived neurotrophic factor modulation in the hippocampus of young adult mice. Behav Pharmacol 18, 491–496, doi:10.1097/FBP.0b013e3282d28f50 (2007).
16. Maegle, M. et al. Reversal of neuromotor and cognitive dysfunction in an enriched environment combined with multimodal early onset stimulation after traumatic brain injury in rats. Journal of neurotrauma 22, 772–782, doi:10.1089/neu.2005.22.772 (2005).
17. Särkämö, T. et al. Music and speech listening enhance the recovery of early sensory processing after stroke. Journal of cognitive neuroscience 22, 2716–2727, doi:10.1162/jocn.2009.21376 (2010).
18. Kraus, N. et al. Central auditory system plasticity associated with speech discrimination training. Journal of cognitive neuroscience 7, 25–32, doi:10.1162/089892995585506 (1995).
19. Lundborg, G. Brain plasticity and hand surgery: an overview. The Journal of head trauma rehabilitation 16, 2716–2727, doi:10.1097/00009089-200112000-00010 (2001).
20. Hua, X. Y. et al. Long-term ongoing cortical remodeling after contralateral C-7 nerve transfer. Journal of neurosurgery 118, 725–729, doi:10.3171/2012.2.jns12207 (2013).
21. Nätänen, R. Mismatch negativity: clinical research and possible applications. International journal of the International Organization of Psychophysiology: official journal of the International Organization of Psychophysiology 48, 179–188 (2003).
22. Michie, P. T. What has MMN revealed about the auditory system in schizophrenia? International journal of psychophysiology: official journal of the International Organization of Psychophysiology 42, 177–194 (2001).
23. Light, G. A. & Braff, D. L. Mismatch negativity deficits are associated with poor functioning in schizophrenia patients. Arch Gen Psychiatry 62, 127–136, doi:10.1001/archpsyc.62.2.127 (2005).
24. Pekkonen, E., Rinne, T. & Nätänen, R. Variability and replicability of the mismatch negativity. Electroencephalogr Clin Neurophysiol 96, 546–554 (1995).
25. Kujala, T., Kallio, J., Tervaniemi, M. & Nätänen, R. The mismatch negativity as an index of temporal processing in audition. Clin Neurophysiol 112, 1712–1719 (2001).
26. Braff, D. L. & Light, G. A. Preattentional and attentional cognitive deficits as targets for treating schizophrenia. Psychopharmacology 174, 75–85, doi:10.1007/s00213-004-0184-0 (2004).
27. Keren, O., Ben-Dror, S., Stern, M. J., Goldberg, G. & Grososser, Z. Event-related potentials as an index of cognitive function during recovery from severe closed head injury. The journal of head trauma rehabilitation 13, 15–30 (1998).
28. Kaipio, M. L. et al. Fast vigilance decrement in closed head injury patients as reflected by the mismatch negativity (MMN). Neuroreport 12, 1517–1522 (2001).
29. Kim, A. S., Binnis, M. A. & Alain, C. Neuroelectric evidence for cognitive association formation: an event-related potential investigation. PloSone 7, e34856, doi:10.1371/journal.pone.0034856 (2012).
30. Maegle, M. et al. Multimodal early onset stimulation combined with enriched environment is associated with reduced CNS lesion volume and enhanced reversal of neuromotor dysfunction after traumatic brain injury in rats. Eur j Neurosci 21, 2406–2418, doi:10.1111/j.1460-9588.2005.04070.x (2005).
31. Särkämö, T. et al. Music listening enhances cognitive recovery and mood after middle cerebral artery stroke. Brain: a journal of neurology 131, 866–876, doi:10.1093/brain/awr013 (2008).
32. Schulz, R. et al. Enhanced Effective Connectivity Between Primary Motor Cortex and Intraparietal Sulcus in Well-Recovered Stroke Patients. Stroke; a journal of cerebral circulation 47, 482–489, doi:10.1161/strokeaha.115.011641 (2016).
33. Wu, J. et al. Connectivity measures are robust biomarkers of cortical function and plasticity after stroke. Brain: a journal of neurology 138, 2359–2369, doi:10.1093/brain/awv156 (2015).
34. Kaji, R. et al. Botulinum toxin type A in post-stroke upper limb spasticity. Current medical research and opinion 26, 1983–1992, doi:10.1185/03007995.2010.497103 (2010).

Acknowledgements
This study was supported by a grant from the Chinese High-tech R&D (863) Program (SS2015AA020501).
Author Contributions
T.L. and H.X.Y. designed the study, recorded the data and helped perform the surgery procedure, analyzed the electrophysiology data, wrote the main manuscript, prepared the Figs 1,2A,2B,3. M.X.Z. recorded the data of the behavior examination tests and helped draft the manuscript, prepared the Fig. 2C. Y.L.Z. performed the rehabilitation treatment. Y.Q.Q. helped recorded and analyzed the electrophysiology data, prepared the Fig. 2A. Y.D.S. helped perform the surgery procedure. J.G.X. and Y.D.G. helped design the study, review the data critically. W.D.X. designed the study, perform the surgery procedure, review the data critically and approve the final version of the manuscript.

Additional Information
Competing Interests: The authors declare that they have no competing interests.

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2017