INTRODUCTION

Complex regional pain syndrome (CRPS) is a chronic pain disorder of the limbs, most often affecting the upper limbs (Marinus et al., 2011). The development of CRPS is usually precipitated by an injury such as a fracture but can also occur spontaneously and is characterized by pain, sensory disturbances, motor dysfunction, and autonomic dysregulation (Harden et al., 2007; Marinos et al., 2011). Motor dysfunction in CRPS can involve a decreased range of motion, muscle weakness, tremor, and dystonia (Harden et al., 2007). Symptoms of CRPS can persist for many years, and the most persistent sign is motor dysfunction (Bean et al., 2014).

To date, cortical circuits have been the primary focus of studies examining motor dysfunction in CRPS, with functional magnetic resonance imaging (fMRI) studies showing altered activation...
patterns in the primary motor cortex (M1) and supplementary motor area (SMA) during finger tapping or action observation (Hotta et al., 2017; Mihöhner et al., 2007). Subcortical structures, such as the basal ganglia, are also crucial in motor function, including action selection and voluntary movement (Alexander et al., 1986; Redgrave et al., 2011). Basal ganglia motor loops, via the putamen, are somatotopically organized (Nambu et al., 2002), with thalamocortical projections determining whether a movement in a particular body region is facilitated or suppressed (Albin et al., 1989). It is well established that basal ganglia pathology results in movement disorders (e.g., Parkinson’s disease), and specific putamen lesions can lead to dystonia (Neychev et al., 2011), as well as changes in pain perception (Borsook et al., 2010; Starr et al., 2011).

Altered thalamocortical rhythm is associated with neuropathic pain and burst firing of infraslow oscillations (ISOs) (Gerke et al., 2003; Iwata et al., 2011; Sarntchein et al., 2006). In a recent series of studies, we reported differences in resting infraslow oscillations (ISOs) in pain processing regions in individuals with CRPS and other chronic neuropathic pain conditions (Alshelh et al., 2016; Di Pietro et al., 2020). We suggested that these differences may underpin altered thalamocortical rhythm and ultimately in the persistence of pain. Altered thalamocortical rhythm has not only been reported in neuropathic pain. Putamen projections mediate thalamocortical rhythm regulating movement (Opri et al., 2019) and altered thalamocortical rhythm is also found in Parkinson’s disease (Vanneste et al., 2018). Furthermore, in Parkinson’s disease, increased ISO power is associated with increased motor dysfunction (Hou et al., 2014; Wang et al., 2020). Interestingly, in CRPS subjects, we also found increased ISOs in the region of the putamen with extension into the insula cortex contralateral to the affected limb (Di Pietro et al., 2020). The frequency range of ISO increase in CRPS as well as Parkinson’s disease is indicative of astroglisis as it coincides with the range of astrocytic calcium waves (0.03–0.06 Hz) and release of gliotransmitters (Crunelli et al., 2002; Henderson & Di Pietro, 2016).

Indeed, increased ISO and increased astrocyte activation have been found in the same brain regions in neuropathic pain (Alshelh et al., 2016; Okada-Ogawa et al., 2009). Furthermore, optimal astrocytic calcium levels are needed as either attenuating or elevating astrocytic calcium levels lead to motor impairments (Aguilhon et al., 2013; Padmarshri et al., 2015). Thus, a detailed investigation of ISOs in multiple putamen and caudate seeds in CRPS may reveal underlying astrocytic calcium differences compared to controls that may be related to altered motor function.

Several studies have reported structural and functional connectivity changes in the putamen in adult and pediatric cases of CRPS (Azqueta-Gavaldon et al., 2020; Becerra et al., 2014; Linnman et al., 2013), however, these studies did not evaluate the putamen’s somatotopic organization. In addition, functional connectivity changes in non-motor cortico-basal ganglia re-entrant loops that regulate cognition, reward/motivation, visual processing, and sensory integration (Alexander et al., 1986; Choi et al., 2012; Da Cunha et al., 2012; Middleton & Strick, 1996; Redgrave et al., 2010) have been reported in CRPS (Becerra et al., 2014; Geha et al., 2008; Lebel et al., 2008). However, again, these studies did not systematically evaluate each basal ganglia functional loop, nor did they determine if changes were related to sensory, motor, or non-motor disturbances.

The aim of this resting-state fMRI study was to systematically investigate if adult CRPS subjects have altered ISOs and functional connectivity in motor (putamen) and non-motor basal ganglia loops, compared to pain-free control subjects. We hypothesized that CRPS subjects would have altered basal ganglia ISOs and connectivity, and that changes in motor loop activity and connectivity would be associated with motor dysfunction of CRPS.

## 2 | MATERIALS AND METHODS

This study was approved by the Human Research Ethics Committee of the University of Sydney (HREC reference number 2018/073) and was conducted in accordance with the Declaration of Helsinki. Due to the uncommon nature of CRPS and difficulty in recruiting eligible and willing upper limb CRPS participants, no sample size calculation was performed, hence the study recruited a convenience sample. Prior to study participation, informed written consent was obtained from all participants. Data from the CRPS participants and a subset of the pain-free healthy control participants have been published in prior studies (Di Pietro et al., 2020; Lee et al., 2020).

### 2.1 | Study participants

Sixteen eligible individuals with CRPS gave consent to participate in the study. Imaging data were not obtained from one CRPS subject due to claustrophobia in the MRI scanner. Thus, imaging data from 15 upper limb CRPS subjects (11 females; mean±SEM age: 47.5±3.2 years) and 45 age- and sex-matched pain-free healthy controls (33 females; 47.3±1.9 years) were collected. CRPS subjects were diagnosed following the International Association for the Study
of Pain “Budapest” diagnostic criteria (Harden et al., 2007) and had ongoing pain for at least 3 months. CRPS subjects were eligible for the study if they reported other regions of pain or CRPS, but upper limb CRPS was required to be their primary complaint. Exclusion criteria included any MRI contraindications such as cardiac pacemakers and metal implants, or any significant mental health disorders, developmental delays, or neurological disorders that would prevent safe participation.

2.2 | CRPS assessment

The researcher assessed CRPS signs in both upper limbs of each CRPS subject. (i) Sensory: Hyperalgesia and allodynia were assessed by pinprick on the dorsal webspace of the hand and light brush strokes on the dorsum of the hands, respectively. (ii) Vasomotor: Skin temperature asymmetry was assessed through touch, and skin color changes/asymmetry was assessed visually. (iii) Sudomotor/edema: Sweating (sudomotor) was assessed by touching the subject’s palms of both hands. Edema was assessed through measurement of the circumference of the wrist and proximal phalanx of the middle finger with a tape measure. Signs of sudomotor function and edema were recorded as present if there was asymmetry between the upper limbs. (iv) Motor/Trophic: We observed for motor signs such as tremor and dystonia. Motor weakness was assessed through a power grip test on the researcher’s index and middle fingers. Hair, nail, and skin changes/asymmetry between the upper limbs (trophic changes) were assessed visually.

2.3 | Questionnaires

2.3.1  Each CRPS subject completed several questionnaires

Pain
CRPS participants rated their pain intensity on the day of the study (“day pain”) on a 10 cm visual analogue scale (VAS) (0 = no pain to 10 = worst imaginable pain). Using the 10 cm VAS, CRPS subjects were also asked to record their ongoing pain intensity three times a day for 7 days before or following the scanning session. The mean “diary pain” score was obtained by averaging the 21 pain intensity scores.

Functional assessment
The patient-rated Wrist and Hand Evaluation (PRWHE) assessed task-associated pain intensity and functional difficulty of the CRPS affected limb (MacDermid, 1996). The PRWHE is divided into pain and function subscores, as well as total score. The PRWHE pain score ranges from 0 to 50, the function score 0 to 50, and the total score 0 to 100. Higher scores indicate more pain and functional disability of the CRPS affected limb. The shortened 11-item Disabilities of the Arm, Shoulder and Hand (QuickDASH) Outcome Measure (Beaton et al., 2005) assessed the overall function of the upper limbs irrespective of CRPS affected side, and the score ranges from 0 to 100. A higher QuickDASH score indicates greater disability of the upper limbs.

Body perception and motor dysfunction
The Bath CRPS Body Perception Disturbance Scale (Lewis & McCabe, 2010) and Foreign limb Feelings (FLF) questionnaire (Galer & Jensen, 1999) assessed self-perception of the CRPS affected limb. The Bath and FLF questionnaires scores range from 0 to 57 and 0 to 20, respectively. In addition, the FLF questionnaire assesses aspects of motor dysfunction such as involuntary movement. For both scales, a higher score indicates greater disturbance to self-perception of the CRPS affected limb.

2.4 | Tactile acuity (two-point discrimination)

Tactile acuity data of CRPS participants in this study have previously been published elsewhere (Di Pietro et al., 2020; Lee et al., 2020). Tactile acuity measures were not obtained from 3 of the 15 CRPS subjects due to extreme hand pain. The 2-point discrimination (TPD) test assessed tactile acuity and was performed immediately following the MRI. TPD is the ability to discriminate two distinct points touching the skin as two points and not one. The researcher applied a TPD wheel (Exacta™, CA) longitudinally to the distal pulp of the subject’s index finger and asked the subject to report if they felt 1 point or 2 points touching their skin with each stimulation. The distances of 0 (i.e., 1 point), 2, 3, 4, and 5 mm between two points were applied seven times in a pseudo-randomized order, resulting in 35 trials per hand. The percentage of correct two-point perception versus distance between the points was fitted by a binary logistic regression (IBM SPSS Statistics for Windows, Version 24.0. Armonk, NY: IBM Corp.) to obtain individual psychometric functions of TPD for each subject’s hand. The discrimination distance was determined at a 50% (chance) threshold of correct two-point perception. A higher discrimination distance indicates poorer tactile acuity.

2.5 | MRI data collection

All MRI data were collected in a 3 Tesla MRI scanner (Achieva TX, Philips Medical Systems) at Neuroscience Research Australia in Sydney, Australia. Each subject lay supine on the MRI scanner bed with their head immobilized in a padded 32-channel head coil. A high-resolution T1-weighted anatomical image of the whole brain in the sagittal plane was obtained (repetition time = 5600 ms; echo time = 2.5 ms, flip angle = 8°, raw voxel size 0.87 mm³). Subjects were then asked to relax with their eyes closed as resting-state fMRI (rsfMRI) images were collected (series of 180 fMRI image volumes, gradient-echo planar sequence with blood oxygen level-dependent contrast; repetition time = 2000 ms; echo time = 30 ms, flip angle = 90°, 37 axial slices, raw voxel size 3 x 3 x 4 mm).
MRI data analysis

Image analysis was performed using SPM12 (Friston et al., 1995) and custom software. The T1 and rsfMRI images of the CRPS subjects with pain restricted to the left upper limb (or the more intense pain in the left upper limb) were left-right reflected across the midline on the y-axis before data processing. Hence, the brain’s left hemisphere was contralateral to the CRPS affected limb in all 15 subjects. All fMRI images were then slice-time corrected, realigned and the Dynamic Retrospective Filtering (DRIFTER) toolbox (Särkkä et al., 2012) used to model and remove cardiac (frequency band of 60–120 beats per minute +1 harmonic) and respiratory (frequency band of 8–25 breaths per minute +1 harmonic) noise. LMRP detrending was used to remove movement-related signal changes. The linear model of the global signal (LMGS) method (Macey et al., 2004) was then used to remove global drifts in fMRI signal intensity. Each subject’s fMRI images were then co-registered to the subject’s own T1-weighted anatomical image. The T1 images were spatially normalized to Montreal Neurological Institute (MNI) space, and the normalization parameters were applied to the fMRI images sets to place them into MNI space. The resulting fMRI images were then smoothed using a 6 mm full-width at half-maximum (FWHM) Gaussian filter.

Seeds

To explore regional specificity, we created multiple basal ganglia seeds (Figure 1a). First, we refined a left putamen seed (putamen ISO seed) from a previous study where we showed that ISO power was significantly greater in CRPS compared with control participants (Di Pietro et al., 2020). The putamen ISO seed for this current study was refined to only include the putamen portion of a larger seed that originally encompassed the putamen and extended into the insular cortex. Second, we created four putamen seeds, each a 3 mm radius sphere and representative of: the motor hand (center X, Y, Z MNI coordinates: −28, −7, 2), motor foot (−28, −8, 6), motor tongue (−32, −8, −5), and SMA hand (−25, −7, 5) (Choi et al., 2012). Third, to elucidate the involvement of non-motor basal ganglia circuitry, we created six more seeds (3 mm radius spheres) representing the left caudate tail (visual processing loop; −29, −8, −17), caudate body (oculomotor loop; −15, −1, 18), ventrolateral (vl) caudate head (lateral orbitofrontal loop; −9, 10, 1), dorsolateral (dl) caudate head (dorsolateral prefrontal loop; −12, 10, 8), ventral putamen (default network loop; −29, −11, −10), and ventral striatum (limbic loop; −12, 11, −8) (Alexander et al., 1986; Choi et al., 2012; Middleton & Strick, 1996). In total, we created and assessed 11 basal ganglia seeds. In addition, to investigate basal ganglia functional connectivity, the fMRI analysis was restricted to cortical areas that receive input from the basal ganglia seeds (Alexander et al., 1986; Middleton & Strick, 1996) by applying a mask of the frontal lobe, anterior cingulate cortex (ACC), middle temporal gyrus, and parietal lobe (Figure 1b). All seeds were created in the left hemisphere; hence all seeds are contralateral to the CRPS affected limb.

ISO analysis

We used the SPM Data Processing Assistant for Resting-State fMRI (DPARSF) toolbox (Chao-Gan & Yu-Feng, 2010) to calculate the amplitudes of low-frequency fluctuations (ALFFs), that is, power, of three standard ISO frequency ranges (slow-5: 0.01–0.027 Hz, slow-4: 0.027–0.073 Hz and slow-3: 0.073–0.198 Hz) defined by Buzsaki and Draguhn (2004) on the preprocessed fMRI image sets of control participants.
and CRPS subjects. The analysis was restricted to the 11 basal ganglia seeds to investigate ISO differences within the basal ganglia. Two-sample $t$ tests were used to determine any differences between controls and CRPS subjects’ slow-5, slow-4, and slow-3 ALFF power ($p<0.05$, two-tailed). The slow-5, slow-4, and slow-3 mean±SEM ALFF power for controls and CRPS subjects were plotted for each seed. For seeds that were significantly different between controls and CRPS subjects, Pearson’s correlations ($p<0.05$, two-tailed) were performed between CRPS subjects’ ALFF power and pain duration, pain intensity, questionnaire scores, and tactile acuity, using GraphPad Prism 9.1 (GraphPad Software Inc., San Diego, CA). The D’Agostino-Pearson normality test confirmed the normal distribution of pain, questionnaire, and tactile acuity data before correlation analysis. To account for multiple comparisons for each seed and each ISO frequency range, the Benjamini and Hochberg method (Benjamini & Hochberg, 1995) was used to calculate FDR-adjusted $p$ values at 5% FDR.

2.9 Seed-based connectivity analysis

For each of the 11 basal ganglia seeds, the mean signal intensity at each time point was calculated and averaged over the entire seed. Relationships between signal intensity changes and the signal intensity changes in each voxel of the brain were then determined for each seed in each subject. The resultant resting-state connectivity strength brain maps were then smoothed using a 6mm FWHM Gaussian filter and second-level random-effects analysis was performed to determine significant differences between CRPS and control groups ($p<0.05$, FDR corrected for multiple comparisons, minimum contiguous cluster size of 20 voxels). For the resting-state connectivity analysis of the putamen ISO seed, the beta value (connectivity strength effect size) of each significant cluster was extracted, and the mean±SEM plotted. In addition, for the CRPS subjects, linear relationships between these beta values and pain duration, pain intensity, questionnaire scores, and tactile acuity data were determined using Pearson’s correlations ($p<0.05$, two-tailed) and the Benjamini and Hochberg method (Benjamini & Hochberg, 1995) was used to calculate FDR-adjusted $p$ values at 5% FDR. One-sample one-sided $t$ tests were performed within groups for each basal ganglia seed (see Supporting Information).

3 RESULTS

3.1 CRPS subject characteristics

Individual CRPS subject characteristics are reported in Table 1. The CRPS subjects’ average pain duration was $4.6±0.9$ years (mean±SEM). All 15 CRPS subjects reported ongoing pain in the upper limb, with 12 of the 15 subjects reporting upper limb pain restricted to one side (10 right, 2 left). Three subjects reported bilateral upper limb pain (2 reporting greater pain in the right upper limb and 1 on the left). Seven CRPS subjects also reported pain in the lower limb. CRPS subjects’ motor signs, questionnaire scores, and tactile acuity data are reported in Table 2. Thirteen CRPS subjects displayed motor signs on the day of the study; 10 CRPS subjects presented with weakness, 2 with dystonia, 1 with tremor, and 1 with rigidity. CRPS subjects reported an average pain intensity of $5.3±0.5$ for day pain and $4.6±0.6$ for diary pain.

3.2 ISOs

As expected, for the putamen ISO seed, the CRPS group displayed significantly greater slow-4 ALFF (0.027–0.073Hz) power than controls (mean±SEM, CRPS: $1.06±0.06$, controls: $0.85±0.03$, $p<0.001$) (Figure 2a). The putamen ISO seed also displayed significantly greater slow-3 ALFF (0.073–0.198Hz) power in the CRPS group (CRPS: $1.11±0.09$, controls: $0.88±0.03$, $p=0.003$). This result was regionally specific; only 3 of the remaining 10 basal ganglia seeds displayed significant differences in ALFF power between groups. Namely, the putamen SMA hand, motor hand, and motor tongue area seeds demonstrated greater CRPS slow-4 ALFF power in CRPS than controls (SMA hand: CRPS: $0.93±0.06$, controls: $0.81±0.03$, $p=0.046$; motor hand: CRPS: $1.06±0.08$, controls: $0.86±0.03$, $p=0.005$; motor tongue: CRPS: $1.03±0.07$, controls: $0.84±0.03$, $p=0.004$) (Figure 2b). There were no significant differences in slow-4 ALFF power for the remaining seven seeds or for either slow-3 or slow-5 (0.01–0.027Hz) frequency bands in any basal ganglia seed, apart from the putamen ISO seed.

In CRPS subjects, increased ISO power of the putamen was correlated with pain and functional disability (Table 3). A lower PRWHE function score, indicating better wrist function of the CRPS affected hand, correlated with increased slow-4 ALFF power of the putamen ISO seed (Figure 3a) and putamen motor hand (Figure 3b). A lower QuickDASH score, indicating less functional disability of the upper limbs, correlated with increased slow-4 putamen motor hand (Figure 3c), putamen motor tongue, and slow-3 putamen motor hand (Figure 3d) ALFF power.

3.3 Functional connectivity

For each of the 11 basal ganglia seeds, significant differences in resting-state functional connectivity strengths between CRPS and control groups were determined. In no region was connectivity strength greater in controls than in CRPS subjects. For the putamen ISO seed, CRPS subjects displayed significantly greater connectivity strengths compared with controls in the ipsilateral and contralateral primary motor cortices (M1) in the region innervating the lower limbs (mean±SEM beta values: ipsilateral M1: CRPS $0.13±0.02$, controls $0.02±0.02$; contralateral M1: CRPS $0.15±0.02$, controls $0.03±0.01$), contralateral M1 in the upper limb and hand region (CRPS: $0.13±0.02$, controls $0.02±0.01$), contralateral cingulate motor area (CMA; CRPS $0.16±0.02$, controls $0.05±0.02$).
| Subject | Age (years) | Sex | Pain duration (years) | CRPS affected region(s) | Inciting event | Medications | Comorbidities |
|---------|-------------|-----|-----------------------|-------------------------|----------------|-------------|--------------|
| 1       | 49.2        | M   | 7.0                   | **R UL, R LL, L LL, face, abdomen** | Pain in R hand | Turmeric tablets | None |
| 2       | 56.4        | F   | 4.2                   | **L UL, R UL, R LL, R and L chest** | L humerus fracture | Duloxetine, gabapentin, oxycodone, quetiapine, tapentadol | L radial nerve palsy, triangular fibrocartilage complex degeneration of R hand |
| 3       | 55.8        | F   | 0.9                   | **R UL, R neck, R chest** | Spontaneous onset | Ashwagandha, budesonide, cannabis, codeine, formoterol, oxycodone, paracetamol, salbutamol | Back pain, fibromyalgia, osteoarthritis, radiculopathy, spinal disc herniation |
| 4       | 61.7        | F   | 6.2                   | **L UL, R UL** | R hand tendon release surgery | Amitriptyline, cannabidiol drops, codeine, levothyroxine, magnesium, paracetamol, topiramate, tramadol, valerian | None |
| 5       | 58.1        | F   | 8.7                   | **R UL, R LL, R face** | R arm surgery | Codeine, duloxetine, linagliptin, meloxicam, metformin, paracetamol | None |
| 6       | 66.6        | F   | 9.5                   | **R UL, L UL, R LL, L LL** | R radius fracture | Amlodipine, gabapentin, ketamine, in lipoderm cream, metformin, metoprolol, pantoprazole, salbutamol | Osteoarthritis |
| 7       | 46.8        | M   | 1.5                   | **R UL** | Spontaneous onset | Amlodipine, atorvastatin, ibuprofen, paracetamol, perindopril, pregabalin | None |
| 8       | 34.2        | F   | 5.3                   | **R UL, R LL, R hip** | R wrist fracture | Amitriptyline, buprenorphine patch | Migraine, R hip bursitis |
| 9       | 25.9        | F   | 1.3                   | **R UL, L and R neck, spine, L LL** | R hand nerve damage | None | Endometriosis |
| 10      | 45.2        | M   | 1.2                   | **R UL** | R wrist injury | Codeine, ibuprofen, magnesium, pregabalin, topical rub (containing copaiba, frankincense, peppermint, coconut oil), vitamin B, vitamin C | Bell's palsy, bulging discs |
| 11      | 45.9        | F   | 3.9                   | **L UL** | L hand carpal tunnel release surgery | Amitriptyline, betahistine, duloxetine, naproxen, pantoprazole, rizatriptan, tapentadol, valaciclovir | Carpal tunnel of R hand, fibromyalgia, migraine |
| 12      | 23.8        | F   | 2.6                   | **R UL, L UL** | Overload | Amitriptyline, gabapentin, levothyroxine | None |

*Comorbidities: Pain-related, Non-pain-related*
ipsilateral primary somatosensory cortex (S1) in the hand region (CRPS 0.11±0.02, controls 0.004±0.01), ipsilateral supramarginal gyrus (CRPS 0.17±0.02, controls 0.04±0.01), ipsilateral pars opercularis (CRPS 0.19±0.02, controls 0.04±0.02), ipsilateral orbitofrontal cortex (CRPS 0.14±0.02, controls 0.02±0.01), and in the contralateral middle temporal gyrus (CRPS 0.13±0.03, controls 0.01±0.01) (Figure 4 and Table 4). In CRPS subjects, the connectivity between the putamen ISO seed and the ipsilateral OFC was positively correlated with day pain ($r = 0.60$) and putamen ISO and contralateral M1 hand area was positively correlated with a higher foreign limb feelings (FLF) questionnaire score ($r = 0.62$), however, neither correlations were significant after multiple comparisons adjustment.

Analysis of the remaining 10 basal ganglia seeds revealed that 5 seeds displayed significant connectivity increases in the CRPS group, 4 of which were putamen motor loop seeds. Similar to the putamen ISO seed, the putamen motor hand seed in CRPS subjects displayed greater resting connectivity strengths than controls with: the ipsilateral M1 lower limb region (mean±SEM beta values: CRPS 0.14±0.02, controls 0.04±0.01), ipsilateral and contralateral M1 hand region (ipsilateral: CRPS 0.16±0.02, controls 0.05±0.01; contralateral: CRPS 0.16±0.03, controls 0.04±0.01), contralateral CMA (CRPS 0.17±0.03, controls 0.05±0.01), ipsilateral OFC (CRPS 0.13±0.03, controls 0.02±0.01), and the ipsilateral angular gyrus (CRPS 0.16±0.02, controls 0.04±0.01) (Figure 5a, Table 4). Similarly, for the putamen motor foot seed analysis, CRPS subjects also displayed significantly greater connectivity strengths in the ipsilateral M1 hand region (CRPS 0.17±0.03, controls 0.04±0.01), contralateral M1 hand region (CRPS 0.17±0.04, controls 0.04±0.02), ipsilateral M1 face region (CRPS 0.08±0.01, controls 0.01±0.01), ipsilateral and contralateral CMA (ipsilateral: CRPS 0.13±0.03, controls 0.01±0.01; contralateral: CRPS 0.20±0.03, controls 0.06±0.01), ipsilateral angular gyrus (CRPS 0.18±0.04, controls 0.01±0.02), and in the ipsilateral supramarginal gyrus (CRPS 0.18±0.03, controls 0.04±0.02) (Figure 5b, Table 4). In contrast, connectivity strength for the putamen motor tongue seed was only greater in CRPS subjects compared with controls in a discrete region of the ipsilateral frontal cortex (CRPS 0.18±0.02, controls 0.01±0.02) (Figure 5c, Table 4). For the putamen SMA hand seed, CRPS subjects displayed greater connectivity than controls in the contralateral CMA (CRPS 0.14±0.02, controls 0.03±0.01), ipsilateral OFC (CRPS 0.11±0.03, controls 0.004±0.01), contralateral M1 hand area (CRPS 0.15±0.03, controls 0.05±0.01), ipsilateral M1 tongue area (CRPS 0.13±0.03, controls 0.01±0.01), ipsilateral angular gyrus (CRPS 0.16±0.03, controls 0.05±0.01), ipsilateral and contralateral supramarginal gyrus (ipsilateral: CRPS 0.16±0.02, controls 0.03±0.01; contralateral: CRPS 0.13±0.03, controls 0.01±0.02), contralateral frontal cortex (CRPS 0.11±0.03, controls 0.01±0.02), and the contralateral middle temporal gyrus (CRPS 0.11±0.02, controls 0.03±0.01) (Figure 5d, Table 4).

The caudate body seed was the only non-motor region in which resting functional connectivity was significantly different between
| Subject | Motor signs       | Pain intensity | Questionnaire scores | TPD (mm) |       |       |
|---------|------------------|----------------|----------------------|----------|-------|-------|
|         |                  | Day pain       | PRWHE                |          | Affected hand | Unaffected hand |
|         |                  | Diary pain     | Pain score           | Function score | Total score | QuickDASH | Bath | FLF |       |       |
| 1       | Tremor, dystonia | 4.0            | 32                   | 30.5      | 62.5          | 81.8      | 34   | 18  | 3.8   | 1.6  |
| 2       | Rigidity         | 7.8            | 45                   | 30.5      | 75.5          | 75.0      | 29   | 16  | -     | -    |
| 3       | Weakness         | 7.9            | 46                   | 84        | 75.0          | 39        | 12   |     | 3.7   | 3.3  |
| 4       | Weakness         | 4.3            | 42                   | 88        | 54.5          | 3         | 6    |     |       |      |
| 5       | Weakness         | 4.1            | 43                   | 83.5      | 75.0          | 32        | 19   |     | 4.1   | 2.3  |
| 6       | Weakness         | 5.0            | 26                   | 52.5      | 56.8          | 10        | 5    |     | 4.1   | 3.3  |
| 7       | Weakness         | 7.1            | 41                   | 78        | 75.0          | 12        | 4    |     | 2.1   | 1.8  |
| 8       | Weakness         | 2.4            | 28                   | 33.2      | 61.2          | 31.8      | 19   | 15  | 3.7   | 2.4  |
| 9       | Weakness         | 6.8            | 40                   | 36.5      | 76.5          | 75.0      | 18   | 2   | 2.6   | 2.2  |
| 10      | Weakness         | 3.0            | 38                   | 76        | 79.5          | 18        | 11   |     |       |      |
| 11      | -                | 5.6            | 5                    | 8.5       | 13.5          | 25.0      | 29   | 3   | 2.3   | 1.9  |
| 12      | Weakness         | 4.4            | 28                   | 16.5      | 44.5          | 47.7      | 21   | 0   | 2.5   | 2.7  |
| 13      | -                | 3.5            | 28                   | 15.5      | 43.5          | 54.5      | 30   | 11  | 4.1   | 2.7  |
| 14      | Weakness         | 5.9            | 23                   | 21        | 44            | 54.5      | 25   | 7   | 2.0   | 1.9  |
| 15      | Dystonia         | 7.6            | 46                   | 93        | 88.6          | 26        | 16   |     |       |      |

Mean (±SEM) | 5.3 (±0.5) | 4.6 (±0.6) | 34.1 (±2.9) | 31.0 (±2.9) | 65.1 (±5.6) | 63.3 (±4.8) | 23.0 (±2.5)  | 9.7 (±1.6) | 3.1 (±0.2) | 2.3 (±0.2) |

Abbreviations: Bath, bath CRPS body perception disturbance scale; FLF, foreign limb feelings; PRWHE, patient-rated wrist and hand evaluation; QuickDASH, shortened disabilities of the arm, shoulder and hand; SEM, standard error of mean; TPD, two-point discrimination.
CRPS subjects and controls (Figure 6, Table 4). CRPS subjects displayed greater connectivity strength than controls in several brain regions, including the ipsilateral OFC (CRPS 0.13 ± 0.03, controls 0.02 ± 0.01), ipsilateral and contralateral dorsolateral prefrontal cortex (dIPFC; ipsilateral: CRPS 0.15 ± 0.01, controls 0.04 ± 0.01; contralateral: CRPS 0.17 ± 0.02, controls 0.06 ± 0.02), ipsilateral and contralateral mid-cingulate cortex (MCC; ipsilateral: CRPS 0.17 ± 0.01, controls 0.07 ± 0.01; contralateral: CRPS 0.15 ± 0.02, controls 0.02 ± 0.01), ipsilateral and contralateral superior parietal cortex (ipsilateral: CRPS 0.16 ± 0.04, controls 0.03 ± 0.02; contralateral: CRPS 0.17 ± 0.01, controls 0.05 ± 0.02).
contralateral: CRPS 0.19 ± 0.02, controls 0.08 ± 0.02), contralateral superior frontal cortex (CRPS 0.15 ± 0.02, controls 0.04 ± 0.01), contralateral anterior cingulate cortex (ACC; CRPS 0.17 ± 0.02, controls 0.06 ± 0.01), contralateral supramarginal gyrus (CRPS 0.17 ± 0.04, controls 0.05 ± 0.02), ipsilateral and contralateral M1 (ipsilateral: CRPS 0.13 ± 0.03, controls 0.02 ± 0.02; contralateral: CRPS 0.13 ± 0.03, controls 0.01 ± 0.02), ipsilateral and contralateral angular gyrus (ipsilateral: CRPS 0.17 ± 0.03, controls 0.05 ± 0.02; contralateral: CRPS 0.16 ± 0.04, controls 0.03 ± 0.02), ipsilateral and contralateral piriform cortex (ipsilateral: CRPS 0.14 ± 0.02, controls 0.04 ± 0.01; contralateral: CRPS 0.12 ± 0.02, controls 0.02 ± 0.01), ipsilateral and contralateral middle temporal gyrus (ipsilateral: CRPS 0.14 ± 0.02, controls 0.02 ± 0.01; contralateral: CRPS 0.13 ± 0.02, controls 0.03 ± 0.01). No significant functional connectivity strength differences were found for the non-motor regions, caudate tail, dl caudate head, vl caudate head, ventral striatum, and ventral putamen seeds.

**DISCUSSION**

Consistent with our hypothesis, we report significant differences in both basal ganglia ISO power and basal ganglia functional connectivity between individuals with CRPS and pain-free controls. These differences are primarily restricted to the motor basal ganglia loops, particularly regions that represent the upper limb, with one exception being the caudate body oculomotor loop. More specifically, we found increased ISO power in CRPS subjects in putamen divisions representative of the SMA hand, motor hand, and motor tongue seeds. In addition to the putamen SMA hand, motor hand, motor tongue seeds, the putamen motor foot, and caudate body seeds displayed significantly greater (than controls) resting connectivity strength to multiple basal ganglia-cortical input areas such as M1, SMA, CMA, and OFC in CRPS subjects. These results highlight alterations in basal ganglia function at rest in CRPS and likely underpin the alterations in motor control commonly seen in this condition.
We found greater putaminal ISO power, primarily of the slow-4 bandwidth, which includes the 0.03–0.06 Hz range, a range we have previously reported on (Di Pietro et al., 2020). This range coincides with the astrocytic calcium wave propagation frequency range and subsequent astrocytic gliotransmitter release (Cornell-Bell et al., 1990; Crunelli et al., 2002) and we have hypothesized that increased ISOs found in CRPS and other forms of chronic neuropathic pain may be due to chronic astrogliosis (Henderson & Di Pietro, 2016). While no investigation has explored chronic astrogliosis in the putamen of CRPS patients, a postmortem study reported chronic astrogliosis in the spinal cord dorsal horn in CRPS (Del Valle et al., 2009). Furthermore, two human positron emission tomography studies have reported increased putaminal translocator protein binding in CRPS, a marker that binds to microglia and potentially also astrocytes (Jeon et al., 2017; Seo et al., 2021). Given this, we now propose that astrogliosis may also contribute to motor dysfunction in CRPS. Indeed, in mice, artificial elevation of calcium through activation of GFAP+ glial cells (astrocytes) in the brain and spinal cord resulted in motor coordination impairment (Agulhon et al., 2013).

From the results of this current study, we speculate that astrogliosis within the basal ganglia is specific to the areas of greater slow-4 oscillations found in CRPS subjects, namely putamen SMA hand, motor hand, and motor tongue areas contralateral to the CRPS affected limb. Supporting this idea is evidence of elevated pro-inflammatory cytokine levels in the striatum of rats with chronic neuropathic pain (Al-Amin et al., 2011; Apkarian et al., 2006; Fiore & Austin, 2016). We also identified greater CRPS slow-3 oscillations in the putamen and motor hand area of the putamen contralateral to the CRPS affected limb. This suggests that in addition to calcium wave elevation and astrogliosis, there are other aberrant mechanisms that contribute to the motor dysfunction seen in CRPS.

In addition to altered resting oscillation patterns, we report significant increases in resting putamen connectivity. The putamen is the input region of the basal ganglia motor loop; it is organized somatotopically and has segregated inputs from M1 and SMA (Alexander et al., 1986; Nambu et al., 2002). Activity fluctuations in

![Figure 3: Increased infraslow oscillations (ISO) power in CRPS subjects correlated with less disability. Scatter plots showing Pearson correlations; the line represents the best fit for the correlations. The Pearson’s correlation coefficient (r) and the FDR-adjusted significance of correlation (p) are displayed on the plots. Patient-rated wrist and hand evaluation (PRWHE) correlated with slow-4 ALFF power of the (a) putamen ISO seed and (b) putamen motor hand area. Shortened disabilities of the arm, shoulder and hand (QuickDASH) outcome measure correlated with (c) slow-4 ALFF power, and (d) slow-3 ALFF power of the putamen motor hand area.](image-url)
the putamen motor hand, motor foot and SMA hand areas displayed greater synchrony with cortical areas in CRPS subjects. Overall, these areas showed stronger coupling with other motor related cortical regions such as the M1 region innervating the body as well as the CMA. Our findings of altered bilateral putamen-M1 connectivity are consistent with the bilateral nature of M1 changes in CRPS (Di Pietro et al., 2013a; Maihöfner et al., 2007). Contralateral changes are hypothesized to be due to altered functioning of the affected limb and ipsilateral changes related to compensatory use of the unaffected limb (Azqueta-Gavaldon et al., 2020). Consistent with this, we found that increased contralateral putamen-M1 hand functional connectivity was correlated with higher FLF scores in CRPS, indicating a greater self-perceived disturbance in motor function and neglect-like feelings of the CRPS affected limb. In Parkinson's disease and CRPS, greater putamen-M1 connectivity predicts poorer motor performance in a pegboard task that requires fast and accurate motor coordination (Azqueta-Gavaldon et al., 2020; Simioni et al., 2015). Furthermore, in CRPS, decreased putamen-M1 functional connectivity correlated with reduced forearm range of motion (Azqueta-Gavaldon et al., 2020). Thus, the increased putamen-M1 hand functional connectivity in CRPS subjects may underlie motor dysfunction.

As well as altered connectivity to well-described motor cortical regions, we found increased putamen connectivity with frontal and parietal association cortices, consistent with putamen anatomical connectivity to those areas (Jarbo & Verstynen, 2015). We found increased connectivity of the putamen ISO and putamen SMA hand seeds with the OFC, an important brain area in emotion and reward. Increased OFC connectivity is associated with non-reward, punishment, and depression (Rolls et al., 2020) and the increased putamen-OFC connectivity found in the current study may underpin the aversion to movement commonly seen in CRPS. Our study demonstrated increased putamen motor hand, motor foot, and SMA hand connectivity to the parietal cortex, mainly the inferior parietal lobule (IPL; formed by the supramarginal and angular gyri). The IPL's role involves sensorimotor integration, motor planning, spatial and nonspatial attention, and motor preparation (Caspers et al., 2013). In CRPS, finger tapping on the CRPS affected side increased IPL activation compared to the unaffected cerebral hemisphere and also to healthy controls (Schwenkreis et al., 2009), and in CRPS...
TABLE 4 Montreal neurological institute (MNI) coordinates, t-scores, cluster size, and beta values for regions with significant functional connectivity differences between CRPS subjects and healthy controls

| Region | Side | MNI coordinates | t-score | Cluster size | Beta values (mean ± SEM) | CRPS | Controls |
|--------|------|-----------------|---------|--------------|--------------------------|------|----------|
|        |      | X   Y  Z        |         |              |                          |      |          |
| Putamen ISO seed CRPS > Controls |      |     |     |     |     |     |     |
| Primary motor cortex (M1) leg area | ipsi | 8   −28  64 | 3.75   | 21            | 0.13 ± 0.02              | 0.02 ± 0.02 |
| Primary motor cortex (M1) leg area | contra | −10  −24 58  | 4.33   | 53            | 0.15 ± 0.02              | 0.03 ± 0.01 |
| Primary motor cortex (M1) hand area | contra | −30  −18 54  | 4.37   | 59            | 0.13 ± 0.02              | 0.02 ± 0.01 |
| Cingulate motor area (CMA) | contra | −4   −8  54  | 3.95   | 31            | 0.16 ± 0.02              | 0.05 ± 0.02 |
| Primary somatosensory (S1) hand area | ipsi | 24   −42 48  | 3.83   | 23            | 0.11 ± 0.02              | 0.004 ± 0.01 |
| Supramarginal gyrus | ipsi | 46   −42 38  | 4.86   | 555           | 0.17 ± 0.02              | 0.04 ± 0.01 |
| Pars opercularis | ipsi | 48   10   8   | 4.30   | 156           | 0.19 ± 0.02              | 0.04 ± 0.02 |
| Orbitofrontal cortex (OFC) | ipsi | 30   34   0   | 4.66   | 26            | 0.14 ± 0.04              | −0.003 ± 0.02 |
| Orbitofrontal cortex (OFC) | ipsi | 40   42  −12  | 4.60   | 38            | 0.14 ± 0.02              | 0.02 ± 0.01 |
| Middle temporal gyrus | contra | −46  −4   −30 | 4.84   | 23            | 0.13 ± 0.03              | 0.01 ± 0.01 |
| Putamen (motor hand) seed CRPS > Controls |      |     |     |     |     |     |     |
| Primary motor cortex (M1) leg area | ipsi | 6    −30  62  | 4.25   | 26            | 0.14 ± 0.02              | 0.04 ± 0.01 |
| Primary motor cortex (M1) hand area | ipsi | 36   −8   50  | 4.75   | 35            | 0.16 ± 0.02              | 0.05 ± 0.01 |
| Primary motor cortex (M1) hand area | contra | −32  −10 48  | 4.50   | 93            | 0.16 ± 0.03              | 0.04 ± 0.01 |
| Cingulate motor area (CMA) | contra | −8   −8  50  | 4.34   | 59            | 0.17 ± 0.03              | 0.05 ± 0.01 |
| Angular gyrus | ipsi | 34   −70 22   | 4.75   | 82            | 0.16 ± 0.02              | 0.04 ± 0.01 |
| Angular gyrus | ipsi | 42   −62  8   | 4.01   | 81            | 0.18 ± 0.03              | 0.05 ± 0.02 |
| Orbitofrontal cortex (OFC) | ipsi | 38   42  −14  | 4.32   | 32            | 0.13 ± 0.03              | 0.02 ± 0.01 |
| Putamen (motor foot) seed CRPS > Controls |      |     |     |     |     |     |     |
| Primary motor cortex (M1) hand area | ipsi | 22   −16  62  | 3.74   | 41            | 0.16 ± 0.03              | 0.02 ± 0.02 |
| Primary motor cortex (M1) hand area | ipsi | 38   −8   50  | 4.58   | 82            | 0.17 ± 0.03              | 0.04 ± 0.01 |
| Primary motor cortex (M1) hand area | contra | −32  −10 46  | 3.57   | 26            | 0.17 ± 0.04              | 0.04 ± 0.02 |
| Primary motor cortex (M1) face area | ipsi | 62   10  18   | 3.50   | 21            | 0.08 ± 0.01              | 0.01 ± 0.01 |
| Cingulate motor area (CMA) | ipsi | 20   −4   54  | 3.81   | 26            | 0.13 ± 0.03              | 0.01 ± 0.01 |
| Cingulate motor area (CMA) | contra | −10  −8  50  | 4.60   | 89            | 0.20 ± 0.03              | 0.06 ± 0.01 |
| Supramarginal gyrus | ipsi | 54   −36 30   | 4.01   | 64            | 0.18 ± 0.03              | 0.04 ± 0.02 |
| Angular gyrus | ipsi | 34   −70 22   | 4.21   | 54            | 0.19 ± 0.03              | 0.05 ± 0.02 |
| Angular gyrus | ipsi | 54   −66 18   | 4.10   | 24            | 0.12 ± 0.03              | −0.003 ± 0.02 |
| Angular gyrus | ipsi | 46   −50 10   | 4.23   | 81            | 0.18 ± 0.04              | 0.01 ± 0.02 |
| Putamen (motor tongue) seed CRPS > Controls |      |     |     |     |     |     |     |
| Frontal cortex | ipsi | 26   8   48  | 5.76   | 25            | 0.18 ± 0.02              | 0.01 ± 0.02 |
| Putamen (SMA hand) seed CRPS > Controls |      |     |     |     |     |     |     |
| Cingulate motor area (CMA) | contra | −10  −10 48  | 4.31   | 33            | 0.14 ± 0.02              | 0.03 ± 0.01 |
| Superior frontal cortex | contra | −6   48  38  | 3.76   | 24            | 0.11 ± 0.03              | 0.01 ± 0.02 |
| Primary motor cortex (M1) hand area | contra | −30  −10 46  | 4.13   | 33            | 0.15 ± 0.03              | 0.05 ± 0.01 |

(Continues)
TABLE 4 (Continued)

| Region                                      | Side   | MNI coordinates | t-score | Cluster size | Beta values (mean ± SEM) |
|---------------------------------------------|--------|-----------------|---------|--------------|-------------------------|
| Primary motor cortex (M1) tongue area       | ipsi   | X: 50 Y: -6 Z: 30 | 3.94    | 42           | 0.13 ± 0.03 0.01 ± 0.01 |
| Supramarginal gyrus                        | contra | X: -40 Y: -60 Z: 26 | 3.60    | 20           | 0.13 ± 0.03 0.01 ± 0.02 |
| Supramarginal gyrus                        | ipsi   | X: 52 Y: -36 Z: 22 | 4.98    | 294          | 0.16 ± 0.02 0.03 ± 0.01 |
| Angular gyrus                              | ipsi   | X: 36 Y: -70 Z: 24 | 4.08    | 35           | 0.16 ± 0.03 0.05 ± 0.01 |
| Orbitofrontal cortex (OFC)                  | ipsi   | X: 32 Y: 18 Z: -18 | 4.04    | 39           | 0.11 ± 0.03 0.004 ± 0.01 |
| Middle temporal gyrus                      | contra | X: -52 Y: -2 Z: -32 | 4.21   | 38           | 0.11 ± 0.02 0.03 ± 0.01 |

**Caudate body seed CRPS > Controls**

| Region                                      | Side   | MNI coordinates | t-score | Cluster size | Beta values (mean ± SEM) |
|---------------------------------------------|--------|-----------------|---------|--------------|-------------------------|
| Superior parietal cortex                    | ipsi   | X: 22 Y: -54 Z: 68 | 3.18    | 51           | 0.10 ± 0.03 0.01 ± 0.01 |
| Superior parietal cortex                    | ipsi   | X: 4 Y: -52 Z: 62 | 3.36    | 31           | 0.16 ± 0.03 0.04 ± 0.02 |
| Superior parietal cortex                    | ipsi   | X: 18 Y: -56 Z: 58 | 2.97    | 25           | 0.11 ± 0.03 0.01 ± 0.02 |
| Superior parietal cortex                    | contra | X: -28 Y: -56 Z: 40 | 3.07    | 80           | 0.19 ± 0.02 0.08 ± 0.02 |
| Superior parietal cortex                    | contra | X: -14 Y: -78 Z: 34 | 3.15    | 27           | 0.17 ± 0.03 0.06 ± 0.02 |
| Superior parietal cortex                    | ipsi   | X: 18 Y: -82 Z: 20 | 4.40    | 337          | 0.16 ± 0.04 0.03 ± 0.02 |
| Middle cingulate cortex (MCC)               | contra | X: -14 Y: -10 Z: 42 | 4.84    | 6866         | 0.15 ± 0.02 0.02 ± 0.01 |
| Middle cingulate cortex (MCC)               | ipsi   | X: 12 Y: 26 Z: 42 | 3.31    | 93           | 0.17 ± 0.01 0.07 ± 0.01 |
| Superior frontal cortex                     | contra | X: -20 Y: 30 Z: 32 | 3.18    | 67           | 0.19 ± 0.03 0.07 ± 0.02 |
| Superior frontal cortex                     | contra | X: -12 Y: 56 Z: 32 | 3.12    | 48           | 0.17 ± 0.03 0.07 ± 0.01 |
| Superior frontal cortex                     | contra | X: -16 Y: 66 Z: 6 | 3.71    | 135          | 0.15 ± 0.02 0.04 ± 0.01 |
| Anterior cingulate cortex (ACC)             | contra | X: -12 Y: 32 Z: 28 | 3.45    | 131          | 0.17 ± 0.02 0.06 ± 0.01 |
| Supramarginal gyrus                        | contra | X: -46 Y: -36 Z: 38 | 3.22    | 60           | 0.17 ± 0.04 0.05 ± 0.02 |
| Primary motor cortex (M1)                   | contra | X: -40 Y: -6 Z: 20 | 3.74    | 84           | 0.13 ± 0.03 0.01 ± 0.02 |
| Primary motor cortex (M1)                   | ipsi   | X: 50 Y: 6 Z: 10 | 3.18    | 124          | 0.13 ± 0.03 0.02 ± 0.02 |
| Angular gyrus                              | contra | X: -26 Y: -88 Z: 14 | 3.88    | 216          | 0.16 ± 0.04 0.03 ± 0.02 |
| Angular gyrus                              | ipsi   | X: 48 Y: -52 Z: 14 | 3.30    | 80           | 0.17 ± 0.03 0.05 ± 0.02 |
| Angular gyrus                              | contra | X: -38 Y: -66 Z: 8 | 2.97    | 37           | 0.14 ± 0.04 0.02 ± 0.02 |
| Dorsolateral prefrontal cortex (dlPFC)      | contra | X: -36 Y: 36 Z: 6 | 3.38    | 21           | 0.17 ± 0.02 0.06 ± 0.02 |
| Dorsolateral prefrontal cortex (dlPFC)      | ipsi   | X: 44 Y: 48 Z: -2 | 4.00    | 837          | 0.15 ± 0.01 0.04 ± 0.01 |
| Orbitofrontal cortex (OFC)                  | ipsi   | X: 22 Y: 56 Z: -16 | 4.32    | 88           | 0.13 ± 0.03 0.02 ± 0.01 |
| Piriform cortex                            | ipsi   | X: 20 Y: 4 Z: -16 | 3.75    | 66           | 0.14 ± 0.02 0.04 ± 0.01 |
| Piriform cortex                            | contra | X: -22 Y: 8 Z: -24 | 4.15    | 76           | 0.12 ± 0.02 0.02 ± 0.01 |
| Middle temporal gyrus                      | contra | X: -48 Y: -8 Z: -26 | 3.62    | 91           | 0.12 ± 0.02 0.03 ± 0.01 |
| Middle temporal gyrus                      | ipsi   | X: 46 Y: -4 Z: -30 | 4.70    | 173          | 0.13 ± 0.02 0.04 ± 0.01 |

Note: All seeds are contralateral to the CRPS affected limb.

subjects with dystonia, imagined movements of the affected limb resulted in reduced IPL activation compared to control subjects (Gieteling et al., 2008). In Parkinson’s disease subjects with tremor, IPL-M1 connectivity was increased compared to controls (Vervoort et al., 2016). While our study did not test the connectivity between IPL and M1, both IPL and M1 had increased connectivity to the putamen seeds contralateral to the CRPS affected limb. Thus, it is probable that increased contralateral putamen-IPL connectivity indicates dysfunction in IPL in CRPS subjects and may underlie motor dysfunction such as tremor.

Most basal ganglia seeds that displayed significant connectivity differences were motor areas; however, we also found alterations in the caudate body, which represents the striatal portion of the oculomotor loop (Alexander et al., 1986). A recent study identified this basal ganglia region as a convergence zone of corticostriatal projections with integration of reward, executive control and spatial attention during spatial reinforcement learning thought to be associated with the caudate body (Jarbo & Verstynen, 2015). In our study, the caudate body seed displayed the greatest degree of increased functional connectivity to cortical areas including M1,
prefrontal and parietal association cortices in CRPS subjects compared to controls. Perceptual disturbances such as neglect-like syndrome and a foreign feeling of the CRPS affected limb are common features of CRPS (Galer & Jensen, 1999; Kuttikat et al., 2016). The caudate body showed increased connectivity with cortical regions governing visual attention (middle temporal gyrus), spatial attention (superior frontal cortex) and visuospatial perception (superior parietal cortex) and thus, may relate to the visuospatial issues in CRPS. Indeed, lesions to the caudate and the IPL can lead to deficits in visuospatial perception that present as neglect (Karnath et al., 2005; Kumral et al., 1999). However, in the current study, we did not perform correlation analysis of caudate body functional connectivity to...
body perception. Thus, it is unknown to what degree visuospatial perception may be related to caudate body functional connectivity in CRPS.

There are several limitations to this study. This study had a limited sample of 15 CRPS subjects. However, given the rare nature of CRPS, it is comparable to previous studies (Di Pietro et al., 2013a, 2013b). It is noted, however, the sample size of this study is a statistically limits correlation interpretations (even when adjusted for FDR), and a much larger sample size is needed for greater reliability on correlation interpretations. The longitudinal following of subjects recruited before CRPS development would identify whether brain differences are true changes, and whether they are pathophysiologically or adaptive. Furthermore, it is unknown whether the findings in our study are specific to CRPS or are also present in other pain or motor dysfunction conditions. Future studies should aim to compare changes in sensory and motor disabilities between different conditions. Nonetheless, it is clear that the sensory and motor disturbances are a result of reorganization of the sensory and motor cortices and related cortico-basal ganglia loops and are part of the CRPS disease pathology. The resting-state nature of our scanning and the lack of quantitative assessment of motor dysfunction are also weaknesses. Future studies should include quantitative assessment of motor dysfunction (e.g., pegboard task, Unified Parkinson Disease Rating Scale) or examine functional connectivity during movement tasks (e.g., finger tapping, wrist extension)—although this would present a practical challenge. Medications may have had a potential effect on results as 13 of the 15 CRPS subjects had taken medication before the study; however, it is not easy or necessarily ethical to exclude medicated CRPS subjects or include a washout period before the study. Ideally, the study would have recruited CRPS participants with no comorbidities, however, given the rare nature of CRPS and multi-system dysfunction following CRPS (Schwartzman, 2012), this was difficult. Hence, it cannot be fully excluded that multiple comorbidities may have also had a potential effect in this study’s findings. However, all CRPS participants had been diagnosed with CRPS, and the list of comorbidities varies greatly between each CRPS participant. Given the great variability, collectively the multiple comorbidities are unlikely to contribute significantly to the overall changes reported in this study.

5 | CONCLUSION

Our study is the first to systematically evaluate infraslow oscillations and resting-state functional connectivity in motor and non-motor basal ganglia functional loops in CRPS subjects. Compared to controls, we identified increased ISOs in the putamen contralateral to the CRPS affected side, but not in non-motor basal ganglia regions in CRPS subjects. Moreover, putaminal ISOs correlated with wrist function and disability scores. We demonstrated increased functional connectivity in the contralateral putamen (especially the motor hand region) and caudate body seeds, with a network of cortical structures including the M1 hand region, SMA, and other frontal
and parietal association cortices. There were no changes in striatal seeds representing cognitive and limbic basal ganglia loops. In summary, the recruitment of anatomically specific motor basal ganglia-cortical networks likely underlies motor symptoms, such as dystonia and tremor, while the caudate body network may be related to altered visuospatial integration, in CRPS.

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CONFLICT OF INTEREST
The authors have no conflict of interest to declare.

AUTHOR CONTRIBUTIONS
All authors had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Conceptualization, F.D.P., L.A.H., and P.J.A.; Methodology, F.D.P., L.A.H., and P.J.A.; Formal Analysis, B.L., L.A.H., and P.J.A.; Writing – Original Draft, B.L., L.A.H., and P.J.A.; Writing – Review & Editing, B.L., F.D.P., L.A.H., and P.J.A.; Visualization, B.L., F.D.P., L.A.H., and P.J.A.; Supervision, F.D.P., L.A.H., and P.J.A.; Funding Acquisition, F.D.P.

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of the article at the publisher’s website.

**FIGURE S1** One-sample t-test within group results for putamen seeds; a: putamen ISO, b: putamen (motor hand), c: putamen (motor foot), d: putamen (motor tongue), and e putamen (SMA hand). Figures were masked using frontal lobe, anterior cingulate cortex (ACC), middle temporal gyrus and parietal lobe and visualised at a threshold of $p = 0.05$ FDR and 20 voxel minimum cluster size

**FIGURE S2** One-sample t-test within group results for non-motor basal ganglia seeds; a: caudate body, b: caudate tail, c: ventrolateral (vl) caudate head, d: dorsolateral (dl) caudate head, e: ventral striatum, and f: ventral putamen. Figures were masked using frontal lobe, anterior cingulate cortex (ACC), middle temporal gyrus and parietal lobe and visualised at a threshold of $p = 0.05$ FDR and 20 voxel minimum cluster size. The ventral striatum CRPS (e) did not have any contrast at $p = 0.05$ FDR, 20 voxel minimum cluster size when masked with the frontal lobe, anterior cingulate cortex (ACC), middle temporal gyrus and parietal lobe mask

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