Neural Correlates of Chronic Low Back Pain Measured by Arterial Spin Labeling

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ABSTRACT

Background: The varying nature of chronic pain (CP) is difficult to correlate to neural activity using typical functional magnetic resonance imaging methods. Arterial spin labeling is a perfusion-based imaging technique allowing the absolute quantification of regional cerebral blood flow, which is a surrogate measure of neuronal activity.

Methods: Subjects with chronic low back and radicular pain and matched healthy normal subjects, undergoing identical procedures, participated in three sessions: a characterization and training session and two arterial spin labeling sessions. In the first imaging session, CP (if any) was exacerbated using clinical maneuvers; in the second session, noxious heat was applied to the affected leg dermatome, the intensity of which was matched to the pain intensity level of the CP exacerbations for each back pain subject.

Results: The clinically significant worsening of ongoing CP (± 30%, n = 16) was associated with significant regional blood flow increases (6–10 mm/100 g of tissue/min, P < 0.01) within brain regions known to activate with experimental pain (somatosensory, prefrontal, and insular cortices) and in other structures observed less frequently in experimental pain studies, such as the superior parietal lobule (part of the dorsal attention network). This effect is specific to changes in ongoing CP as it is observed during worsening CP, but it is not observed after thermal pain application, or in matched, pain-free healthy controls.

Conclusions: Study findings demonstrate the use of arterial spin labeling to investigate the neural processing of CP, and these findings are a step forward in the quest for objective biomarkers of the chronic pain experience.

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stimuli (although not necessarily specific only to pain), often termed the “pain matrix.” The pain matrix includes the primary and secondary somatosensory (S1 and S2), anterior cingulate, insular, and prefrontal cortices, and the thalamus.

However, significant limitations have hindered the application of neuroimaging to the study of a patient’s own clinical pain. Foremost is the nearly constant and varying nature of the chronic pain experience (including the ongoing process of evaluating salient meaning) that is difficult to correlate to neural activity using typical fMRI methods, such as blood oxygen level-dependent (BOLD) imaging. The strength of BOLD imaging is the ability to correlate fMRI signal changes to stimulus changes that occur over a period of a few seconds. Unlike experimental pain, chronic clinical pain often cannot be switched on and (especially) off at will; for instance, the ongoing levels of pain in patients with chronic low back pain (CLBP) frequently linger above their baseline levels well after the end of a straight leg-raising maneuver. Given this decoupling between stimulation and pain sensation, CLBP (as well as other pain disorders) eludes two-state subtraction design studies with BOLD imaging, because this technique requires multiple on and off alternations to have sufficient statistical power.

Arterial spin labeling (ASL) is a perfusion-based fMRI technique that uses water in arterial blood as a freely diffusible tracer to measure perfusion noninvasively. This allows for the absolute quantification of regional cerebral blood flow (rCBF), which is a surrogate measure of neuronal activity, and it may be superior to BOLD imaging as a proxy measure of regional glucose utilization. Because of its better estimation of brain activity for low frequency experimental designs and its ability to quantify in absolute units rCBF, ASL appears better suited to study some aspects of the chronic pain experience, although few studies have done so to date. An important experimental pain study applied pulsed ASL (pASL) to healthy volunteers undergoing tonic, experimental muscle pain stimulation. More recently, pASL has been applied in a patient during and after an acute migraine headache.

In this study we use pASL to investigate the neural correlates of changes in baseline levels of ongoing CLBP and radicular pain. We hypothesized that the experimentally induced worsening of CLBP and/or radicular pain, but not the control conditions, would be associated with increased rCBF in a widespread network of brain regions, including (but not exclusively) those of the experimental pain matrix.

Materials and Methods

Study Design and Population
This was an institutional review board-approved (Partners Healthcare, Boston, MA) cohort study of 16 right-handed patients with CLBP and radicular pain and a control group of 16 healthy, right-handed subjects with no pain, matched for age and sex. The experimental design included within subject and between subject controls. Subjects participated in three sessions: a characterization and training session and then two fMRI sessions. One fMRI session (“clinical maneuvers session”) included 10-s periods of temporary exacerbation of back and leg pain through clinical maneuvers, such as straight leg raising or pelvic tilt. The other session (“heat pain session”) included periods of heat pain applied to the affected leg dermatome, the intensity of which was matched to the pain intensity level of the clinical pain exacerbation periods (fig. 1). Subjects with CLBP were included if they were: (1) between the ages of 21–65 yr, (2) had ongoing chronic pain that averaged at least 3 on a 0–10 scale of pain intensity, (3) had no back surgery within the past year, (4) were not having pain management procedures during the study period, (5) were not taking opioids or benzodiazepines, (6) had low back pain with radicular pain of at least 6 months’ duration, (7) did not have sensory or motor deficits that precluded participation in the pain procedures, (8) were right handed, and (9) had a significant discogenic component to their pain syndrome, confirmed by lumbar MRI. Eligibility was determined by investigator ADW at the first visit through a review of a history and physical examination and MRI findings confirming disc disease. Patients were included if this evaluation found that a source of pain was at least one degenerated, herniated, or torn lumbar disc with either a minimum grade III disc degeneration, abnormal morphology, or a hyperintense zone. These criteria, used by the authors in previous studies, exclude those with pain resulting from purely nonspecific or myofascial causes and include those with the commonly presenting mixed syndrome of low back pain with underlying disc pathology and possibly spinal stenosis or facet disease.

Characterization Methods

After the physician obtains written, informed consent, the following self-report questionnaires were administered at the start of each of the three sessions. The low back pain subjects completed all of the scales described in the following paragraphs, whereas the healthy subjects only completed the Pain Catastrophizing Scale and the Garsely Box Scales (GBS) because they had no chronic pain.

Brief Pain Inventory. This is a 15-item questionnaire assessing pain location and 0–10 ratings of pain intensity, relief, quality, pain-related quality of life, and function. It has been validated in cancer and noncancer pain conditions. The activity interference items measure separate domains of function, such as pain interference with activity, sleep, or work.

Neuropathic Pain Questionnaire. This validated scale describes the presence or absence of neuropathic pain symptoms using self-rated descriptive terms for neuropathic pain symptoms such as burning or numbness. It has a predictive accuracy for neuropathic pain of 73% and is used to classify the neuropathic components of a pain syndrome (Yes/No).

Oswestry Disability Index. This is an extensively used
10-item scale to describe the level of disability in patients with CLBP.21

Pain Catastrophizing Scale. This 13-item survey assesses beliefs and thoughts about pain that have been shown to have an independent relationship to pain from other psychologic constructs.22 It can be administered to patients with chronic pain and healthy volunteers.

Gracely Box Scales (GBS). These 20-point scales rate perception of the sensory and affective (unpleasantness) components of pain sensations.23 They were administered at the beginning of each pASL scan and throughout both fMRI sessions to characterize the subjects’ level of baseline, ongoing chronic pain and the levels of pain during the acute pain exacerbation periods and thermal pain stimuli. The Gracely scales include descriptors anchored to values from 0 to 20, such as “0—no pain sensation” and “18—extremely intense.” The GBS are exponential ratio scales ranging from $10^0$ to $10^9$, and they are structured to correct for the nonlinearity of the 0–10 or 0–100 numeric or visual analog pain scales. They are particularly suited and sensitive to determining the degree of change in pain within an experimental session.24 The raw change scores can be converted into percent changes in pain. Throughout the imaging sessions, these scales were presented to the subjects in the scanner with EPRIME software (Psychology Software Tools, Sharpsburg, PA), using a mirror to project them onto a screen comfortably in their field of view. Subjects used an MRI compatible button box to rate pain levels. This method and these scales have been used extensively and validated by our group to assess pain during an fMRI scan session.25

Chronic Pain Exacerbation

For each CLBP subject, if the radicular pain was greater than the axial pain component, a bilateral, passive straight leg-raising maneuver was performed, with the height and angle of elevation recorded. Using an fMRI compatible device custom-made for this study, the legs were raised to two levels that when held for 10 s would acutely worsen the pain to either a Moderate level (10–11 on the GBS sensory scale, “moderate pain condition”) or a Strong level (14–15 on the GBS, “high pain condition”). To familiarize subjects with an actual fMRI session, they were placed in a mock scanner and underwent four stimulations (2 moderate and 2 high pain conditions in random order, spaced 110–120 s apart). Before each stimulation, they rated their baseline (current) pain using the GBS sensory scale; after each stimulation they rated pain intensity and unpleasantness experienced during the exacerbation periods using both the GBS sensory and affective scales, presented in random order.

For each subject it was confirmed that the pain returned to a lower level within 30 s after a 10-s exacerbation period, with the understanding that baseline pain rating before each stimulus may or may not rise over time with repeated stimuli. Subjects could not go further in the study if the pain stimulation acutely worsened their ongoing, baseline pain beyond
this time (which was communicated in discussions with potential subjects before enrollment). Thus, this method enables subjects to distinguish between and assess pain exacerbations and ongoing chronic pain.

Each healthy subject underwent identical passive straight leg-raising maneuvers to the same angles of elevation matched to a CLBP patient for the moderate and high pain stimuli. They participated in an identical fMRI mock scanning session and performed the same rating procedures before and after each stimulus as the CLBP group.

For those CLBP subjects whose axial pain was greater than their radicular pain, to exacerbate their pain they performed either a pelvic tilt or lumbar extension maneuver while supine (depending on whichever method most reliably worsened their pain and allowed it to return to a lower level within 30 s). We recorded the distance the hips were raised off the MRI table or the degrees of extension using an inclinometer, for the moderate and high pain stimuli levels. They underwent identical calibrated pain stimuli and rating procedures as those who performed straight leg-raising maneuvers. The healthy normal subjects performed these exact procedures to the same distances or degrees as the CLBP patient to which they were matched.

**Thermal Pain Stimuli**

Noxious heat was applied to the affected lower leg dermatome in the CLBP patients and to the identical area in the matched healthy control subjects using a Medoc TSA-II device (Medoc, Ramat Yishai, Israel). The thermode size is 30 × 30 mm, with a rate of increase in temperature of 5°C/s. During the training session, thermal stimuli were applied to each subject by gradually increasing the temperature to find the level that produced a rating of Moderate (10–11, “moderate pain”) and Strong (14–15, “high pain”) on the GBS sensory scale when applied for 5 s (calibrated thermal pain stimuli). Subjects were then placed into the mock scanner and underwent a trial run of four random stimuli, 2 moderate pain and 2 high pain, spaced 90–110 s apart. The required temperatures were adjusted if needed. For each subject it was confirmed that there was no lingering thermal pain before the subsequent stimulus, and the probe was moved slightly after each stimulus to prevent tissue sensitization. The rating procedures were identical to the back and leg pain exacerbation methods.

**FMRI Sessions**

The scanning bed was modified to maximize comfort for the CLBP subjects so that chronic pain was less likely to increase from simply lying in the scanner. In both sessions (fig. 1), two 6-min pASL scans were collected using the “PICORE-Q2TIPS” MRI labeling method, with a 3 T Siemens TIM Trio MRI System (Siemens Medical, Erlangen, Germany), equipped with a 32-channel head coil (TR/TE/TI1/TI2 = 3000/13/700/1700 ms, voxel size = 3.515*3.515*6.25 mm, number of slices = 16). “Tag” images were acquired by labeling a thick inversion slab (110 mm) proximal to the imaging slices (gap = 21.1 mm). “Control” images were acquired interleaved with the tag images, by applying an off-resonance inversion pulse without any spatial encoding gradient. At the beginning of each pASL scan, an M0 scan (i.e., the longitudinal magnetization of fully relaxed tissue) was acquired for rCBF quantification purposes (see next paragraph). The resting state (eyes closed) pASL scans were acquired before and after 12 clinical maneuvers (session 1) or 12 heat pain stimuli (session 2, both sessions had 6 moderate pain and 6 high pain stimuli, randomized), delivered in three separate runs. At the start of the “clinical maneuvers session” just before scanning, the CLBP subjects had the level of the clinical maneuver “recalibrated” to determine the levels of straight leg raising, pelvic tilt, or lumbar extension capable of eliciting ratings of moderate and high pain. Before and after each of the 6-min pASL scans, subjects rated their level of baseline pain (if any) using the GBS sensory scale. The delivery of pain stimuli, the duration of each run, and the pain rating procedures were performed identically to the training session. The thermal pain fMRI session was conducted in exactly the same fashion: subjects were recalibrated to the appropriate heat pain levels, rated their pain (if any) just before the pASL scan, underwent three runs of four stimuli each, rated their pain just before a stimulus and after the third run, and then had a repeat pASL scan. A high-resolution MPRAGE scan (TR/TE = 2300/3.39 ms, voxel size 1*1*1.33 mm) was also acquired during one of the two sessions, for the purposes of cortical surface reconstruction.

**Statistical Analyses**

For the demographic history factors and baseline pain questionnaire responses, analysis of variance (ANOVA) and chi-square tests were used to describe the groups. The ratings of chronic pain obtained just before and after a pASL scan were averaged. Repeated measures ANOVA was used to characterize the serial ratings of ongoing pain during the fMRI session, to determine the percentage of change in pain over time and to compare groups. Comparisons of each time point to baseline were made using the Dunnett test for multiple comparisons. All CIs were reported at 95% and all testing was two-tailed. ASL data analysis was performed using a combination of analysis packages including FSL* and Freesurfer.†† The tag, control, and M0 scans were first motion-corrected using MCFLIRT. Then, tag and control scans were surround subtracted (i.e., given each tag0 ((controlX + controlY + tagX + tagY)/2 – tag0)), to achieve perfusion-weighted images. All the perfusion-weighted maps were then averaged and scaled by a factor proportional to the M0 scan to obtain rCBF maps in absolute values (mm/100 g tissue/min). Because accurate measurement of rCBF in white matter presents methodologic challenges (particularly given

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the poor signal-to-noise ratio and longer arterial transit time), additional processing and analyses were carried out on the cortical surface level. RCBF maps were registered to the high-resolution anatomic images using FreeSurfer’s boundary-based registration tool, interpolated onto FreeSurfer-reconstructed cortical surfaces, and then smoothed at the surface level with a kernel of 7.03 mm (2 voxel size). Average global CBF was calculated for both ASL sessions and compared using two-way, repeated measures ANOVA. Significant changes in rCBF values (ΔrCBF, i.e., rCBFASL1-rCBFASL2) were computed for each subject, interpolated to a standard surface space (fsaverage), and then group-averaged. The clusters of change in CBF were extracted from the whole brain data. The same calculations were performed on normalized rCBF data. Montecarlo simulations were run on both the raw change and normalized rCBF change analyses to identify the clusters exhibiting a significant ΔrCBF, using a vertex-level threshold of P = 0.01 and a cluster-level threshold of 0.05. We compared within-session and between-session differences between groups for the ΔrCBF calculated for each session (i.e., rCBFASL1-rCBFASL2 for the same session). Montecarlo simulations control for multiple comparisons when reporting P values of the clusters. Cluster-size approaches (as opposed to single voxel level approaches) are based on the assumption that the probability is low that a given number of pixels exceeding threshold due to chance will be contiguous. Montecarlo simulations allow for the estimation of the probability distribution of cluster size as a function of α level, and thus the identification of a cluster-size threshold, by creating high numbers (10,000 in our case) of simulated null datasets.

Linear regression was used to examine whether there were significant linear correlations between changes in pain and changes in rCBF activation patterns in either of the two sessions in the CLBP patients. This was performed on a whole brain and a cluster level of analysis.

**Results**

Twenty-three CLBP patients were enrolled, and seven could not complete the first session because of failure of their pain to return to a lower level with 30 s of pain stimulation. Baseline data for the 16 CLBP subjects and 16 matched healthy control subjects who completed the study are displayed in Table 1. Most CLBP subjects were female, had a duration of pain greater than 5 yr, did not have a clinically significant neuropathic component to their pain, were clinically and statistically more significantly disabled than their healthy counterparts, and had clinically and statistically greater levels of pain catastrophizing. Ten CLBP patients had predominantly right-sided low back and radicular pain and six had predominantly left-sided pain. In 13 patients the L5 dermatome was most affected, in 2 S1 was most affected, and in 1 L4 was most affected.

Nine CLBP subjects did bilateral straight leg raising maneuvers for their clinical pain exacerbations. For the moderate clinical pain condition the average degree of elevation was 17°, and for the high pain condition the average was 22°. Five subjects did pelvic tilt, raising their hips on average 10 cm for the moderate and 14 cm for the high pain conditions. Two subjects did low back arching, extending 14 and 18 degrees on average for the two conditions, respectively. For the thermal pain testing the average temperature across both groups for the moderate pain condition was 45°C and 47°C for the high pain condition.

**Psychophysics.** Figure 2 displays the means of the baseline pain ratings collected during the clinical maneuvers session and heat pain session using the GBS Sensory Scale for the CLBP patients. For the clinical maneuvers session, the mean baseline level of CLBP at the start of the scanning session was 6.4/20 (‘very mild’) versus 4.3/20 (‘very weak’) in the heat pain session (P = 0.006). The CLBP subjects experienced an average 34.3% (CI, 18.9, 49.8) worsening of pain during the clinical maneuvers session and the healthy control subjects reported no pain (P = 0.0001). During the heat pain fMRI session, the CLBP patients had an average 19.4% (CI, 2.1, 36.7) increase in pain, and the healthy control subjects reported no ongoing pain (only transient heat pain due to the thermal stimuli, P = 0.0001).

**ASL Data.** Baseline whole-brain within-subject and between-subject comparisons contrasted prestimulation rCBF maps in both sessions (i.e., ASL1, before clinical maneuvers or heat pain stimuli) and revealed no statistically or clinically significant differences between session or between group in global CBF values (mean = 50.8 mm/100 g tissue/min session 1, 50.8 mm/100 g tissue/min session 2).

### Table 1. Baseline Demographic and Pain History Information

<table>
<thead>
<tr>
<th>Variable</th>
<th>CLBP Patients (N = 16)</th>
<th>Healthy Control Subjects (N = 16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr, CI)**</td>
<td>47.4 (40.0, 54.8)</td>
<td>46.7 (40.1, 53.2)</td>
</tr>
<tr>
<td>Sex (%female)</td>
<td>69</td>
<td>69</td>
</tr>
<tr>
<td>Avg. duration of pain (yr, CI)</td>
<td>6.24 (3.9, 11.8)</td>
<td>–</td>
</tr>
<tr>
<td>Avg. pain (0–10, CI)</td>
<td>4.8 (3.8, 5.9)</td>
<td>–</td>
</tr>
<tr>
<td>Neuropathic pain (NPQ, %yes)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Disability level (ODI, %, CI)</td>
<td>35.8 (30.0, 41.6)</td>
<td>0* (0.0)</td>
</tr>
<tr>
<td>Pain catastrophizing (PCS, mean, CI)</td>
<td>36 (27.8, 42.1)</td>
<td>14.2* (12.2, 16.2)</td>
</tr>
</tbody>
</table>

* P = 0.0001. ** All CIs are 95%.

Avg = average; CLBP = chronic low back pain; NPQ = Neuropathic Pain Questionnaire; ODI = Oswestry Disability Index; PCS = Pain Catastrophizing Scale.
CI = 43.3, 63.3; and 51.7, CI = 45.4, 62.4, for session 2). Table 2 lists the brain clusters demonstrating significant differences in rCBF changes between the first and second ASL scans for the two sessions. Of note, mean baseline rCBF values in each of these clusters were not significantly different statistically between session or group. Figure 3 displays these clusters on inflated cortical surfaces. For the clinical maneuvers session in the CLBP subjects, statistically significant activity increases (ASL₂ vs. ASL₁) were observed in the bilateral medial and dorsolateral prefrontal cortices, the superior parietal lobules, S1/M1 (primary somatosensory and motor cortices), and S2 (secondary somatosensory cortex) after enhancement of endogenous CLBP by clinical maneuvers. The activations in S1/M1 corresponded to the homuncular areas for the low back and leg. Statistically significant unilateral increases were found in the right anterior insula, presupplementary motor area, and supramarginal gyrus. The bilateral occipital cortices serve as a control region, and they did not show any statistically significant changes in rCBF (fig. 4). Similarly, in the normalized analysis activations were found in the previously listed areas as well as in the anterior cingulate cortex and insula bilaterally. In contrast, the healthy control subjects (HCs) did not exhibit any statistically significant changes in rCBF during their clinical maneuvers session in the raw change or normalized data analyses. A comparison between changes across groups in this session (the CLBP(ASL₂-ASL₁) minus HC(ASL₂-ASL₁) interaction) revealed statistically significant clusters consistent with left S1/M1 and superior parietal lobules (fig. 5).

During the heat pain session, the CLBP subjects did not have any statistically significant clusters of change in rCBF in the raw change and normalized analyses. The HCs exhibited

**Fig. 2.** Mean ratings of baseline clinical pain in chronic low back pain (CLBP) patients (0–20 Gracely Box Scale). Ratings were obtained in-between pain stimulations (approximately 1 min after the end of the preceding heat [B] or clinical [A] maneuver). Bars represent group averages (±SEM) of the following: pASL1 and pASL2 = ratings before and after each pASL scan (two ratings per timepoint); Run1–3 = ratings within each run (four ratings per timepoint). pASL = pulsed arterial spin labeling.

**Table 2.** Significant Vertex-level Clusters

<table>
<thead>
<tr>
<th>Anatomic Label</th>
<th>Size (mm²)</th>
<th>Cluster P Value</th>
<th>x_MAX</th>
<th>y_MAX</th>
<th>z_MAX</th>
<th>ASL1 rCBF*</th>
<th>ASL2 rCBF*</th>
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</thead>
<tbody>
<tr>
<td>Patients–Clinical Maneuvers Session</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>L superior parietal lobule</td>
<td>507.55</td>
<td>0.0001</td>
<td>-10.4</td>
<td>-69.4</td>
<td>52.1</td>
<td>40.2</td>
<td>50.0</td>
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<td>L secondary somatosensory cx</td>
<td>186.54</td>
<td>0.0104</td>
<td>-58.1</td>
<td>-17.1</td>
<td>27.4</td>
<td>41.4</td>
<td>49.0</td>
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<td>L superior frontal gyrus</td>
<td>142.65</td>
<td>0.0481</td>
<td>-11.6</td>
<td>25.5</td>
<td>32.9</td>
<td>43.7</td>
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<td>L rostral middle frontal gyrus</td>
<td>496.51</td>
<td>0.0001</td>
<td>-21.2</td>
<td>56.3</td>
<td>16.2</td>
<td>42.4</td>
<td>48.7</td>
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<td>L superior parietal lobule</td>
<td>179.31</td>
<td>0.0135</td>
<td>-35.6</td>
<td>-49.4</td>
<td>59.2</td>
<td>41.9</td>
<td>48.2</td>
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<td>L paracentral gyrus</td>
<td>231.62</td>
<td>0.0017</td>
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<td>66.0</td>
<td>37.1</td>
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<td>40.2</td>
<td>24.2</td>
<td>51.1</td>
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<td>L precentral gyrus</td>
<td>226.56</td>
<td>0.0022</td>
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<td>-18.8</td>
<td>69.5</td>
<td>33.1</td>
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<td>221.09</td>
<td>0.0034</td>
<td>-37.9</td>
<td>6.2</td>
<td>43.0</td>
<td>52.3</td>
<td>58.3</td>
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<td>R insula</td>
<td>214.31</td>
<td>0.0018</td>
<td>28.9</td>
<td>18.6</td>
<td>-5.4</td>
<td>51.3</td>
<td>58.3</td>
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<td>647.65</td>
<td>0.0001</td>
<td>13.3</td>
<td>40.9</td>
<td>20.8</td>
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<td>227.75</td>
<td>0.0012</td>
<td>34.7</td>
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<td>43.2</td>
<td>47.0</td>
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<td>475.42</td>
<td>0.0001</td>
<td>8.8</td>
<td>4.7</td>
<td>49.7</td>
<td>38.6</td>
<td>45.5</td>
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<td>R paracentral gyrus</td>
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<td>0.0001</td>
<td>4.8</td>
<td>-26.5</td>
<td>68.3</td>
<td>31.7</td>
<td>40.9</td>
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<td>R rostralmiddlefrontal gyrus</td>
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<td>0.0006</td>
<td>45.8</td>
<td>23.9</td>
<td>30.4</td>
<td>45.6</td>
<td>51.8</td>
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<td>R postcentral gyrus</td>
<td>139.06</td>
<td>0.0385</td>
<td>60.6</td>
<td>-10.7</td>
<td>31.1</td>
<td>41.5</td>
<td>50.0</td>
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<td>R supramarginal gyrus</td>
<td>182.88</td>
<td>0.0066</td>
<td>52.8</td>
<td>-35.2</td>
<td>46.0</td>
<td>46.4</td>
<td>53.4</td>
</tr>
<tr>
<td>Patients–Heat Pain Session</td>
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<td>No significant clusters</td>
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<td>Control Subjects–Clinical Maneuvers Session</td>
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<td>Control Subjects–Heat Pain Session</td>
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<td>L isthmus cingulate gyrus</td>
<td>88.8</td>
<td>0.0033</td>
<td>-5.6</td>
<td>-36.4</td>
<td>32.3</td>
<td>61.2</td>
<td>69.0</td>
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<tr>
<td>R superior temporal gyrus</td>
<td>115.6</td>
<td>0.0001</td>
<td>49.9</td>
<td>-14.4</td>
<td>-4.4</td>
<td>54.9</td>
<td>48.8</td>
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* mm/100 g. tissue/min; italics indicates ASL₂<ASL₁.

ASL = arterial spin labeling; L = left; R = right; rCBF = regional cerebral blood flow.
small clusters of rCBF change in the left posterior cingulate gyrus and in the right superior temporal gyrus (table 2). However, no clusters were statistically significant when changes were compared across groups in the raw change and normalized analyses. In addition, for each of the clusters listed previously there were no statistically significant within-subject differences in the rCBF values in the baseline ASL scans at the start of each session in each CLBP and healthy normal subject. Figure 4 illustrates the mean changes in rCBF across all sessions. During the clinical maneuvers session, these clusters in CLBP subjects exhibited an average increase in rCBF that ranged between 6 and 10 ml/100 g of tissue/min, corresponding to a 17–25% increase ($P < 0.01$). These statistically significant increases in rCBF were not found in the aforementioned brain regions for the heat session in CLBP subjects, in either session in the healthy normal control subjects, or in the occipital control regions.

A sensitivity analysis examining whether there were statistically significant linear correlations between changes in pain and changes in rCBF activation patterns in either of the two sessions in the CLBP patients indicated that there were no significant clusters that had a linear relationship to changes in chronic pain.

**Discussion**

In this study we were able to characterize on a behavioral level a patient’s ongoing chronic back and leg pain after temporary periods of evoked, acute exacerbation. We were then able to associate the ongoing experience of chronic pain to neural correlates of brain activity using ASL. Using assessment methods particularly suited to detect these changes (the Gracely Box Scales), the CLBP subjects experienced a mean 34% increase in chronic pain after clinical maneuvers versus a mean 19% increase in chronic pain after heat pain application. Given that a minimum 30% increase in pain has been shown to be clinically relevant, the clinical maneuvers session meaningfully worsened chronic pain whereas the heat session did not.

These clinically meaningful increases in endogenous pain ratings were positively associated with statistically significant increases in rCBF in a widespread network of cortical areas, including the bilateral medial and dorsolateral prefrontal cortices, superior parietal lobules, S1 and S2, and unilaterally in the right insula. As noted in previous studies of experimental pain, these areas encompass the sensory-discriminative and affective pain processing regions related to pain. Although many of these regions are well accepted as key areas of the pain matrix, the superior parietal lobules are important as a component of the dorsal attention network, whose functional connectivity to pain matrix areas has also been associated with greater clinical pain in fibromyalgia patients. Although not specific to pain per se, increased activity within the superior parietal lobules may reflect increased vigilance to a salient stimulus. Activation of these areas during the clinical maneuvers session but not the heat pain session in the CLBP patients is additional evidence of the clinical salience of the worsening of CLBP in the clinical maneuvers session. One could argue that the differences in rCBF increases found
for the CLBP group in the sessions may have been because of the different baseline levels of pain measured at the start of the first ASL scan in each session (fig. 2). However, mitigating this concern is that the mean rCBF values recorded in these clusters during the first (baseline) ASL scan in each session were not statistically significantly different from each other.

Overall, the measured changes in rCBF appear to have a specificity for meaningful changes in chronic pain, as statistically significant activations of pain matrix areas only occurred in the clinical maneuvers session in the CLBP subjects and not in their heat pain session or in the healthy normal control group. Moreover, the positive interaction analysis for the comparison of statistically significant changes in rCBF between CLBP and healthy control subjects in the clinical session also indicates that the rCBF increases in these areas are related to changes in clinical pain ratings, after controlling for any possible areas of significant rCBF changes in the healthy normal subjects. Furthermore, the lack of a linear relationship between changes in pain and changes in rCBF in specific clusters can be expected because we did not see significant changes in rCBF when the change in pain was less than 30%. This threshold effect serves as a neural marker for clinically significant changes in pain, i.e., >30%.

As noted, brain areas deemed to be components of the pain matrix are largely derived from studies of acute experimental pain in healthy volunteers, and it is unclear to what extent these findings apply to a clinical pain matrix, the network of brain areas underlying clinical pain processing in chronic pain patients. Our results indicate that previously defined pain matrix brain areas are also activated in worsening CLBP, and our findings provide neural correlates for the

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**Fig. 4.** The mean changes in rCBF in the activation clusters across all sessions. For the clinical maneuvers session in the CLBP subjects, these areas had a 17–25% increase in rCBF. For all comparisons to the clinical maneuvers session, $P < 0.01$. aINS = anterior insula, ASL = arterial spin labeling, BIL = bilateral, CLBP = chronic low back pain, CTRL = control, gr = grams, HC = healthy controls, L = left side, M1 = primary motor cortex, MFG = medial frontal gyrus, MPFC = medial prefrontal cortex, preSMA = presupplementary motor area, R = right side, rCBF = regional cerebral blood flow, S1 = primary somatosensory cortex, S2 = secondary somatosensory cortex, SMG = superior marginal gyrus, SPL = superior parietal lobule.
chronic pain experience. Recent studies have attempted to address this scientific gap between experimental pain and clinical pain. In one study, 13 patients with CLBP rated spontaneous, moment-to-moment fluctuations in their pain while undergoing BOLD fMRI imaging.39 Baliki et al. used an experimental model that isolated the neural correlates of a possible neuropathic component of spontaneous pain (a specific component of the chronic pain experience), which was most highly associated with activity in the medial prefrontal cortex. Other neuroimaging approaches include fMRI studies of resting (intrinsic) brain connectivity in chronic pain conditions, in particular, fibromyalgia in which connectivity of insula cortex was linearly related to greater spontaneous clinical pain at the time of the scan.38 We also found that significant activity in the medial prefrontal and insular cortices was associated with higher ratings of clinical pain in CLBP.

Another recent study by Kobayashi et al.40 used the application of 30 s of pressure (with an air-filled syringe) to painful back areas of 6 CLBP subjects while undergoing BOLD imaging. Although this study reported that the predominant areas of activation were in the right insula, prefrontal cortices, and posterior cingulate cortex, the experimental approach used in this study (evoked pain) suggests that these brain areas are likely related more to acute experimental pain processing in CLBP than to chronic clinical pain. Our experimental model worsened chronic low back and leg pain to a clinically significant level, as would often occur throughout a patient’s typical daily activities.

Several limitations of this study merit discussion. First, the order of the clinical maneuvers and heat pain sessions was not randomized and could be a confounder. However, the lack of differences in baseline rCBF values between sessions for each CLBP or HC subject, using a whole brain map or region of interest level of analyses, speaks against this notion. Second, on average the baseline level of CLBP at the start of each session was significantly different. This difference can be attributed to the recalibration of the painful stimuli required before ASL scanning in session 1, but not required for session 2, the heat pain session. As noted, it is unlikely that this is a significant confounder. Third, we did not find a linear relationship between pain ratings and rCBF changes, which argues against a specificity of this neural marker for chronic pain severity. Fourth, in conducting the thermal pain testing to find the temperatures eliciting a moderate and high pain response, we used the methods of ascending limits and adjustment, but not descending limit methods. Thus, even though these temperatures reliably reproduced the target pain within a fMRI session, they may not be accurate temperatures if used across several sessions.

Conclusions
As a highly subjective experience, development of objective, physiologic correlates of patient reports of chronic pain can significantly improve the practice of pain medicine. Our results suggest that neural correlates of CLBP found during pASL scanning could be developed as biomarkers for detection of pain or as surrogate endpoints in clinical outcome studies. Much has been written on the potential for neuroimaging findings to become surrogate endpoints in drug development.51,52 One unique feature of our study is that we
increased CLBP using a calibrated maneuver, which allowed us to consistently evoke chronic pain to a target level, transiently, and then to track a gradual increase in baseline pain ratings over time. Our study presents results pertinent for phases 0 (assay development) and I (feasibility and clinical relevance) of biomarker development. Of course, much work needs to be done to fulfill the promise of this potential biomarker, such as experiments in phase II (validation and standardization for clinical utility), phase III (independent confirmation of results), and phase IV (impact assessment).

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References
4. Iannetti GD, Mouraux A: From the neuromatrix to the pain matrix (and back). Exp Brain Res 2010; 205:1–12
6. Tracey I, Johns E: The pain matrix: Reloaded or reborn as we image tonic pain using arterial spin labelling. Pain 2010; 148:559–60
measured on an 11-point numerical pain rating scale. Pain 2001; 94:149–58


