Machine learning–based prediction of clinical pain using multimodal neuroimaging and autonomic metrics


Abstract
Although self-report pain ratings are the gold standard in clinical pain assessment, they are inherently subjective in nature and significantly influenced by multidimensional contextual variables. Although objective biomarkers for pain could substantially aid pain diagnosis and development of novel therapies, reliable markers for clinical pain have been elusive. In this study, individualized physical maneuvers were used to exacerbate clinical pain in patients with chronic low back pain (N = 53), thereby experimentally producing lower and higher pain states. Multivariate machine-learning models were then built from brain imaging (resting-state blood-oxygenation-level-dependent and arterial spin labeling functional imaging) and autonomic activity (heart rate variability) features to predict within-patient clinical pain intensity states (ie, lower vs higher pain) and were then applied to predict between-patient clinical pain ratings with independent training and testing data sets. Within-patient classification between lower and higher clinical pain intensity states showed best performance (accuracy = 92.45%, area under the curve = 0.97) when all 3 multimodal parameters were combined. Between-patient prediction of clinical pain intensity using independent training and testing data sets also demonstrated significant prediction across pain ratings using the combined model (Pearson’s r = 0.63). Classification of increased pain was weighted by elevated cerebral blood flow in the thalamus, and prefrontal and posterior cingulate cortices, and increased primary somatosensory connectivity to frontoinsular cortex. Our machine-learning approach introduces a model with putative biomarkers for clinical pain and multiple clinical applications alongside self-report, from pain assessment in noncommunicative patients to identification of objective pain endophenotypes that can be used in future longitudinal research aimed at discovery of new approaches to combat chronic pain.

Keywords: Support vector machine, Low back pain, Arterial spin labeling, Primary somatosensory connectivity, Heart rate variability

1. Introduction
Although self-report pain intensity ratings are the gold standard in clinical pain assessment, they are highly variable, inherently subjective in nature, and significantly influenced by multidimensional factors. The lack of objective biomarkers for pain has contributed to suboptimal chronic pain management (eg, opioid public health crisis). Thus, research focused on the development of quantitative, objective biomarkers/predictors alongside self-report to aid diagnosis, estimate prognosis, and predict treatment efficacy is of increasing importance to combat chronic pain.

Growing consensus has suggested that altered central nervous system processing can support and maintain abnormal pain perception in chronic pain, implicating aberrant activity and connectivity of multiple functional brain networks, including default mode, salience, and sensorimotor networks, as well as amplification of sensory input to the brain. In addition, altered processing extends to the autonomic nervous system; a recent meta-analysis found that high-frequency heart rate variability is reduced in patients with chronic pain, suggesting that diminished parasympathetic modulation is also associated with chronic pain.
As brain imaging and autonomic data could potentially capture objective measures of the pain experience, multivariate machine-learning techniques have been gaining attention. Such techniques use "features" extracted from a set of clinically relevant data, allowing computer algorithms to "learn" from those features and form a predictive model. This model can then be applied to new data sets or individuals to diagnostically predict disease states or/and treatment efficacy.15 Towards this goal, multivariate machine-learning techniques have used neuroimaging data to propose a brain signature for evoked experimental pain.39 Neuroimaging-based pain prediction, however, has been in a discovery phase and mostly limited to discrimination of brain activity patterns contrasting noxious stimulus-evoked painful vs nonpainful states in healthy, pain-free individuals7,8,37 and estimation of experimental pain ratings.10,37,38 A few prediction studies attempted to discriminate patients with chronic pain from healthy controls22,34,61, however, patients and controls can differ on much more than pain experience (eg, mood, drug levels, etc.). To the best of our knowledge, no study has attempted to classify clinical pain states (ie, higher vs lower clinical pain within an individual) or predict clinical pain ratings at the time of data acquisition, which could provide greater clinical relevance. Moreover, previous studies have reported only modest accuracy, and to our knowledge, no studies have tried to combine multimodal (eg, central and autonomic) parameters to boost prediction of clinically relevant pain states using machine-learning techniques.

We built a multivariate machine-learning model that learns from central and autonomic features, and then classifies clinical pain states and predicts pain intensity. Importantly, to control clinical pain states, our approach modulated pain in patients with chronic low back pain (cLBP) through physical maneuvers aimed to exacerbate their LBP,20,43 thereby creating experimentally controlled lower and higher clinical pain states. Multimodal features included resting-state functional connectivity of the back representation in primary somatosensory cortex,16 whole-brain regional cerebral blood flow (rCBF),20,26,43 and HF HRV.35 Our novel multimodal combinatorial machine-learning approach was then applied to classify and predict clinical pain intensity.

2. Methods

2.1. Patients

We enrolled 71 patients suffering from cLBP meeting Quebec Task Force Classification System categories I-II (ie, patients were unlikely to have significant nerve root involvement, stenosis, or mechanical instability1,21) as confirmed by the study physician and/or review of medical records. All patients were screened with the following inclusion criteria for eligibility: (1) age between 18 and 60 years with a diagnosis of cLBP (duration >6 months) by physician, (2) average LBP intensity >4/10 during the past 2 weeks before consent (0: no pain and 10: most pain imaginable), (3) fluent English, and (4) ability to provoke or exacerbate clinical back pain by performing physical maneuvers. Eligibility was also assessed with the following exclusion criteria: (1) specific causes of back pain (such as cancer, fractures, spinal stenosis, and infections), (2) radicular pain extending below the knee, (3) complicated back problems such as prior back surgery, intent to undergo surgery at time of the study, and unresolved medical legal/disability/workers compensation claims in relation to cLBP, (4) major systemic or neuropsychiatric disease that might confound interpretation of results (eg, severe fibromyalgia, rheumatoid arthritis, major psychiatric disorders, psychoses, seizure disorder, severe cardiorespiratory or nervous system diseases, etc.), (5) self-reported substance abuse disorder in the past 2 years, (6) contraindications to magnetic resonance imaging (MRI) scanning (eg, cardiac pacemaker, metal implants, claustrophobia, and pregnancy), or (7) use of prescription opioids greater than 60 mg morphine equivalents per day or steroids for pain.

Successful maneuver-related pain increase and available data for all model parameters were limited to 53 patients; therefore, we also limited classification, regression, and parameter evaluation and comparison to these 53 patients only (age = 37.37 ± 11.29 years, mean ± SD, 33 females, pain duration = 7.63 ± 7.42 years, range: 0.5-30). See Fig. S1 under supplementary materials for flowchart outlining data collected, excluded, and analyzed (available at http://links.lww.com/PAIN/A671). All patients were informed of the entire experimental protocol and provided written informed consent. The IRB of Partners Human Research Committee approved this experimental protocol (2011P001364), and this study was performed in accordance with the principles of the Declaration of Helsinki (trial registration number at ClinicalTrials.gov: NCT01598974).

2.2. Study design and clinical pain exacerbation

In this study, back pain exacerbation maneuvers were implemented to increase the endogenous levels of clinical back pain in patients.17,20,43 The maneuvers consisted of individualized dynamic physical procedures that exacerbated patient’s clinical LBP intensity and were implemented based on discussions between patients and a trained experimenter during an initial behavioral session. Patients first listed usual activities that most exacerbated their back pain and performed certain repetitions of one or more of these typical back pain exacerbating maneuvers (such as toe touches, back arches, and facet-joint loading twists). Meanwhile, the experimenter kept detailed records of the number of repetitions, distance, depth, angle, and other respective metrics to allow for reproducibility of the maneuvers during MRI scan session. The experimenter also recorded pain ratings before and after maneuvers; based on our previous publications,17,20,43 the protocol aimed for at least 30% increase in subjective pain ratings without exceeding an intolerable pain level. The most common maneuvers used by patients were toe touches (performed by 57% of patients), back arches (19%), and facet-joint loading twists (19%). A few patients could not exacerbate their back pain with typical maneuvers; hence, we chose to have them perform a painful movement from their daily lives (eg, one patient experienced intense pain while wearing socks; hence, this set of actions was used repeatedly in a controlled manner to exacerbate back pain). The sole goal of this manipulation was to temporarily exacerbate clinical LBP.

During the MRI scan session, all patients were scanned twice, once before and once after patients performed the LBP exacerbation maneuvers (pre- and post-maneuver, respectively), to collect multimodal data associated with different (ie, relatively lower and higher) clinical pain states. Neuroimaging data included resting-state functional MRI (fMRI) with 2 approaches: (1) rCBF using Pseudo-Continuous Arterial Spin Labeling (PCASL) imaging and (2) functional connectivity of the primary somatosensory cortical representation of the back (S1 CONN) using blood-oxygenation-level-dependent (BOLD) imaging. Pseudo-Continuous ASL is a noninvasive perfusion imaging method that provides absolute quantification of rCBF across the brain and is
ideal for capturing brain activity related to ongoing, slowly fluctuating clinical pain. In addition, a different functional imaging modality, BOLD, was used to investigate functional connectivity between the back representation in the primary somatosensory cortex (S1) and the rest of the brain. Because physical maneuvers cause direct nociceptive processing in S1 and several studies have now implicated S1 in chronic pain,16,17,45 we evaluated whole-brain connectivity for this seed region, which was identified with a functional localizer using a separate BOLD fMRI scan (see supplementary materials, Section S1, available at http://links.lww.com/PAIN/A671). Automatic data included HRV in the HF range (HFHRV), a marker for cardiovascular modulation. Thus, each of rCBF, S1CONN, and HFHRV measures a unique dimension of central and autonomic processing for different clinical pain intensity states.

Patients rated clinical back pain intensity (0-100, 0: no pain and 100: most pain imaginable) before and after each fMRI scan run and maneuvers. An omnibus F test was conducted on all post-maneuver clinical pain ratings, demonstrating no significant differences (P = 0.19), suggesting that maneuver-related clinical pain elevation was maintained throughout the post-maneuver period, until the end of the scanning session. In addition, a separate omnibus F test on all pre-maneuver clinical pain ratings showed no significant differences (P = 0.43), suggesting stability in ratings during the pre-maneuver period. Hence, average ratings of respective pre-maneuver and post-maneuver rating periods were used for subsequent analyses.

2.3. Acquisition of multimodal neuroimaging and autonomic parameters

All MRI data were collected at the Athinoula A. Martinos Center for Biomedical Imaging, Massachusetts General Hospital, using a 3.0T Siemens Skyra scanner (Siemens Medical, Erlangen, Germany) equipped with a 32-channel head coil. Patients were asked to rest in a supine position inside the MRI scanner with their eyes open and head still during the data collection. A T1-weighted MP-RAGE pulse sequence (time to repetition [TR]/time to echo [TE] = 2530/1.64 ms, flip angle = 7°, field of view [FOV] = 256 × 256 mm, 176 axial slices, and voxel size = 1 × 1 × 1 mm) was used to collect structural MRI data. Resting-state BOLD fMRI data were collected using a T2*-weighted gradient-echo BOLD echo-planar imaging pulse sequence (TR/TE = 3000/30 ms, flip angle = 90°, FOV = 220 × 220 mm, 44 axial slices, voxel size = 2.62 × 2.62 × 3.12 mm, and total acquisition time = 6 minutes). A PCASL pulse sequence (TR/TE = 3800/15 ms, labeling duration = 1500 ms, post-labeling delay = 1200 ms, flip angle = 90°, FOV = 256 × 256 mm, 25 axial slices, voxel size = 4 × 4 × 5 mm, and total acquisition time = 6 minutes) was used to measure rCBF during ASL runs.

During the resting BOLD fMRI and ASL scan runs, physiological data including finger pulse and respiration were collected to calibrate artifacts from cardiovascular and respiratory movement from the fMRI data and to estimate autonomic, cardiovascal modulation during MRI scans associated with different pain states. Finger pulse was collected from the patient’s left index finger using a piezoelectric pulse transducer (MLT1010 Non-Ferrous Transducer; ADInstruments, Colorado Springs, CO), whereas respiration was measured using a custom-built MR-compatible pneumatic belt. These physiological (cardiac pulse and respiration) signals were recorded at 500 Hz sampling frequency during the fMRI runs using an MRI-compatible system (MP150; Biopac Systems Inc., Goleta, CA).

2.4. Preprocessing and assessment of multimodal neuroimaging and autonomic parameters


Resting-state BOLD fMRI data were preprocessed with physiological artifact correction (mcflirt, FSL), susceptibility-induced distortion correction (topup, FSL), and skull-stripping (bet, FSL). Additional artifacts were then removed using a generalized linear model, modeling nuisance regressors for (1) heart rate and respiratory volume per time convolved with respective cardiorespiratory response functions,6,11 (2) white matter and cerebrospinal fluid data identified with the top 5 principal components using the COMPCOR algorithm4,44 with FAST (FSL) tissue segmentation, (3) head motion correction parameters, and (4) a censoring confound matrix of head motion outliers (fsl_motion_outliers, FSL). The corrected data were coregistered to MNI space (bbregister, FreeSurfer), spatially smoothed (full-width at half-maximum = 6 mm, Fslmaths, FSL), and temporally high-pass filtered (cutoff frequency = 0.006 Hz, 3d Bandpass, AFNI). For resting-state seed connectivity analysis, we used the S1 low back representation (see supplementary materials, Section S1, available at http://links.lww.com/PAIN/A671), which was identified by a separate event-related functional localizer fMRI scan and evoked nociceptive stimulus. The S1 seed was created with a 4-mm radius sphere centered on the peak activation voxel within contralateral S1 (peak x/y/z location in MNI space = ±18/−38/72 mm). We created a bilateral seed (S1back) by mirroring this sphere to the ipsilateral hemisphere, as for clinical pain, patients varied in LBP laterality. Averaged fMRI signal from this S1back seed was used for seed connectivity analysis, and the parameter estimates from each patient were used for further analyses (ie, S1CONN).16

The PCASL data were used to estimate an rCBF map for each patient. Data were processed as follows: First, affine tag-control weighted motion correction was performed (batch realign, ASLtbx, https://cnf.upenn.edu/~zewang/ASLtbx.php); and these head motion realignment parameters were regressed out of the difference maps (rigid-body transformation-based MoCo, SPM).41 After high-pass filtering (cutoff frequency = 0.01 Hz), nonbrain voxels were removed from the data (BET, FSL).30 In addition, physiological noise (driven by cardiovascular fluctuation) was regressed out using a generalized linear model with 6 principal components from anatomically defined cerebrospinal fluid and white matter regions using the COMPCOR algorithm.4,44 Tag and control images were then subtracted, and the average of subtracted pairs was divided by the average of control images to obtain a percent change in rCBF. The percent change maps were converted to absolute values (mm/100 g of tissue/min). These rCBF maps were coregistered to individual structural images (bbregister, FreeSurfer),15 followed by nonlinear transformation into MNI152 common space (FNIRT, FSL). Finally, whole-brain rCBF normalization and spatial smoothing (full-width at half-maximum = 8 mm) were performed.

Cardiac pulse pressure data collected during ASL scan runs were used for HRV analysis to estimate cardiovascular modulation associated with different (ie, low and high) clinical pain states. Inhouse scripts (MATLAB 8.3; MathWorks) were used to annotate the finger pulse signal. Commonly available HRV analysis software (HRV standard, Kubios) was used to calculate HRV.
parameters including HF power (ie, \( {\text{HF}}_{\text{HRV}} \)), which is a well-known marker for cardiovagal modulation. 33,35

2.5. Classification of clinical pain states using a support vector machine algorithm

For classification of clinical pain states (ie, between relatively lower and higher clinical pain), a supervised support vector machine (SVM, scikit-learn 0.18.1, https://scikit-learn.org/stable/) with linear kernel was used, as this algorithm is known to have high accuracy and easy interpretability.24 Briefly, the SVM algorithm segregates 2 classes (relatively lower pain vs higher pain, in this case) of data in feature space by finding an optimal hyperplane/decision boundary that maximally separates them. As our multimodal data were collected at 2 time points for each patient (ie, before and after physical maneuvers), and variability existed in baseline clinical pain level for each patient, we used a paired-SVM approach wherein within-patient differences in the pre- and post-maneuver data (ie, change in pain) were emphasized, and baseline pain levels across patients were accounted for (Fig. 1A). The paired-SVM classifier discriminated “pre — post” vs “post — pre” parameter differences (Fig. 1B), a procedure that has been used in past brain imaging research to discriminate state differences between 2 time points.32 Paired-SVM classification was applied for each parameter (ie, \( {\text{S1}}_{\text{CONN}}, {\text{rCBF}} \), and \( {\text{HF}}_{\text{HRV}} \)) independently, yielding SVM classification weights (ie, contribution of each feature/voxel to the classification) and decision responses (ie, the dot product between SVM weights and feature/voxel values for each modality in each patient, where the classification is determined by the sign of these response values) (Fig. 1C).39 Results of classification for each modality (ie, decision responses) were then used for a combined multimodal parameter classification in an attempt to bolster prediction compared with single-modality prediction models.22

2.6. Prediction of clinical pain intensity using a support vector regression algorithm

Next, we sought to predict clinical pain intensity obtained at the time of data acquisition in a between-patients manner using...
a linear support vector regression (SVR) algorithm. Briefly, linear SVR tries to find a linear function that fits all continuous data points (clinical pain intensity ratings, in this case), with minimized error. Support vector regression is appropriate for predicting continuous variables, as opposed to SVM, which has the goal of separating classes of data.

To prepare the data for SVR analysis, each clinical pain rating was considered an independent sample; in other words, data belonging to an individual patient were used independently in an unpaired manner. This procedure was performed to obtain an effective N = 106 for clinical pain ratings and corresponding multimodal features, allowing us to break-up and allocate the samples into separate training (N = 53) and testing (N = 53) data sets. The training data set allowed for creation of the SVR model and the testing data set allowed for validation of the model.

The allocation of clinical pain ratings into training and testing data sets was done through randomization of these 106 clinical pain ratings (as illustrated in Fig. 2A). After randomization, the distribution of clinical pain intensity was equalized and the dynamic range was no longer discrepant (see Results, section 3.3). Furthermore, we posited that the randomization potentially mitigated any bias of experimental design, ie, for the SVM, the lower pain state (time point 1) always occurred before the higher pain state (time point 2) (pre-maneuver always occurred before post-maneuver). Randomization allowed for the training data set to include an equal number of cases from both time point 1 and time point 2 (same for the testing data set), hence accounting for any potential order effects between a training and testing data set.

After allocation into training and testing, each clinical pain intensity rating and the corresponding multimodal features (rCBF, S1CONN and HFHRV) for each patient were used to calculate decision responses (Fig. 2B), representing the degree to which features corresponding to each clinical pain intensity loads on to the clinical pain state SVM model. In other words, the decision response links the SVM model’s confidence of discriminating lower vs higher pain states with raw features corresponding to a range of clinical pain ratings. To elaborate this further, let us say that a particular voxel in S1CONN SVM map shows a high positive weight (ie, highly confident and robust discrimination of post – pre >> pre – post); the dot product of this SVM weight with S1CONN voxel intensity corresponding to a particular clinical pain intensity will produce a positive decision response with a high magnitude. Similar procedures have been used in past research,22 where weights from a previously created model of experimental heat pain39 were applied (as a dot product) to brain features from assessment of experimental mechanical pain in patients with fibromyalgia. The outcome of the computation was the degree to which brain activity to mechanical pain in fibromyalgia loaded on to the generalized experimental heat pain model, allowing for inferences of hypersensitivity to be made.22

The decision responses and their corresponding clinical pain intensity ratings in the training data set were then used to train/build the SVR model (Fig. 2C, left panel). Next, the trained SVR model was evaluated for consistency using independent multimodal SVM decision responses in the testing data set with corresponding pain ratings, and the predicted pain ratings were compared with true pain ratings reported by the patients (Fig. 2C, right panel).

### 2.7. Statistical analysis

After the within-patient classification of pain intensity states using paired SVM, leave-one-patient-out cross-validation was performed to evaluate classification performance (ie, accuracy, sensitivity, specificity, precision, and area under the curve [AUC]). For the paired-SVM approach, sensitivity, specificity, and precision are identical to the accuracy due to the paired characteristics (ie, number of true positives [TP] = true negatives [TN] and false positives [FP] = false negatives [FN]). Permutation analysis was also performed (N = 5000, P < 0.01) to identify the significant features contributing to the classification of different pain intensity states and to calculate significance of classification measures by comparison with those from a random classifier.

For the between-patients SVR analysis, the correlation coefficient (ie, Pearson’s r) and root mean square error (RMSE) between true pain ratings provided by patients (ie, true LBP) and predicted pain ratings provided by the algorithm (ie, predicted LBP) was calculated to assess prediction model performance, and significance was tested with permutation analysis (N = 5000, P < 0.01). For a case of perfect prediction (ie, predicted pain = true pain), the Pearson’s r value would be 1 and RMSE would be 0.

### 3. Results

#### 3.1. Higher clinical pain elicited by physical maneuvers

Patients suffering from cLBP (N = 53 used in final analyses, Table 1 and supplementary materials, Fig. S1; available at http://links.lww.com/PAIN/A671) were recruited to perform physical maneuvers that exacerbated back pain intensity during an fMRI session. Both brain and autonomic data were collected at 2 different clinical pain states for each patient (ie, pre- and post-physical maneuvers). Patients reported increased average LBP intensity after physical maneuvers (change in LBP: +23.6 ± 12.3/100, mean ± SD, P < 0.00001, pre-maneuver LBP: 31.6 ± 19.0, post-maneuver LBP: 55.2 ± 18.9, 0-100 numerical rating scale, where 0 was “no pain” and 100 was “most pain imaginable”) (Fig. 3A). Variability in baseline clinical pain level (ie, pre-maneuvers) was considerable and ranged from 0 to 72.5 out of 100 (Fig. 3B), supporting our paired-SVM approach for within-subject evaluation.

#### 3.2. Within-patient classification of low and high clinical pain states

Multivariate machine learning–based classification was applied to predict clinical pain intensity states (ie, relatively higher vs lower clinical pain states, as modulated by physical maneuvers in patients with cLBP), using combinations of multimodal neuro-imaging (brain) and autonomic outflow parameters. Patients responded to maneuvers with significant decrease in HFHRV (change: −0.29 ± 0.47 log(ms²), paired t test P < 0.001, pre: 6.02 ± 1.15, post: 5.73 ± 1.21).

A supervised paired-SVM algorithm was used to learn from the aforementioned parameters (rCBF, S1CONN, and HFHRV) (Fig. 1) and to classify relatively lower and higher pain states. Paired-SVM classification found that, independently, all 3 parameters significantly contributed to the within-patient classification between pain intensity states (rCBF: accuracy = 81.13%, AUC = 0.90, TP/TN/FN/FP = 43/10/10/43; S1CONN: 79.24%, 0.85, 42/11/11/42; HFHRV: 67.92%, 0.81, 36/17/17/36). The voxelwise paired-SVM weight map for rCBF classified higher vs lower clinical pain states through increased cerebral blood flow to several subcortical and cortical structures including thalamus, and prefrontal and posterior cingulate cortices, and decreased flow to nonback representation subregions of S1 (ie, outside the
putative location of the somatotopic representation of the back (Fig. 4A, and supplementary materials, Table S1, available at http://links.lww.com/PAIN/A671). For S1CONN, the paired-SVM weight map classified higher vs lower clinical pain through increased S1\textsubscript{back} connectivity to frontoinsular cortex, and decreased connectivity to medial prefrontal cortex and other nonback representation subregions of S1/M1 cortices (Fig. 4B, and supplementary materials, Table S2; available at http://links.lww.com/PAIN/A671).

Moreover, combining multimodal parameters (rCBF + S1\textsubscript{CONN} + HFHRV) (Fig. 1) produced the best classification performance (accuracy = 92.45%, AUC = 0.97, and TP/FP/FN/TN = 49/4/4/49) compared to the classification with individual parameters as noted above (Fig. 5). For this combined model, all 3 multimodal

Figure 2. Prediction of between-subject clinical pain intensity using support vector regression (SVR). (A) Clinical pain ratings for each patient (N = 53) were randomized and equally allocated into TRAIN and TEST data sets (providing total effective N = 106). This randomization into TRAIN and TEST equalized the range of pain ratings, previously found to be discrepant between PRE and POST. Corresponding multimodal features for each clinical pain intensity time point were taken, and (B) decision responses were calculated using a dot product between multimodal features and corresponding SVM classification group weights, resulting in decision responses for each modality. (C) The decision responses for TRAIN and corresponding clinical pain ratings were used to build an SVR model. The trained SVR model was then applied to decision responses from TEST, to produce an output of predicted pain ratings. The true (TEST) and predicted pain ratings were plotted, and a Pearson’s correlation coefficient was computed to evaluate model performance. N.b. HFHRV, high-frequency heart rate variability power; PRE, pre-maneuver; POST, post-maneuver; rCBF, regional cerebral blood flow; S1\textsubscript{CONN}, S1\textsubscript{back}<connectivity; SVM, support vector machine.
parameters significantly contributed (S1CONN and rCBF; \( P < 0.001 \), and HFHRV; \( P = 0.007 \)) to the classification. As head motion may confound neuroimaging findings, we explored the relationship between head motion and within-patient pain state classification. Head motion showed no significant contribution to discriminate relatively lower vs higher clinical pain intensity states (\( P = 0.31 \)) (supplementary materials, Section S2; available at http://links.lww.com/PAIN/A671).

### 3.3. Between-patient prediction of clinical pain intensity ratings

Another goal of our analysis was to form a model that can directly predict clinical pain intensity ratings across patients through the introduction of independent training and testing data sets using multimodal parameters. Hence, we conducted a linear SVR analysis, which is optimized for the prediction of continuous variables such as clinical pain ratings (Fig. 2). Data from different patients and different time points, including corresponding clinical pain intensity ratings, were randomized into separate training (\( N = 53 \)) and testing (\( N = 53 \)) data sets, allowing for the range of clinical pain intensity values to be equalized (Fig. 2A).

After randomization, the distribution of clinical pain intensity was equalized (clinical pain intensity range: training set = 0–92.8/100, testing set = 0–86.7; clinical pain intensity: training set = 43.38 ± 23.76/100, testing set = 43.45 ± 20.90, \( P = 0.98 \)).

Combining all 3 multimodal parameters (ie, decision responses of S1CONN, rCBF, and HFHRV) for prediction of pain intensity ratings demonstrated significant performance in terms of predicted vs true clinical pain intensity ratings for both the independent training (Pearson’s \( r = 0.52 \), RMSE = 20.51) and testing (\( r = 0.63 \), \( P = 0.02 \), RMSE = 16.69, \( P < 0.001 \), Fig. 6) data sets. For this combined model, only the S1CONN parameter significantly contributed (S1CONN; \( P = 0.002 \), rCBF; \( P = 0.31 \), and HFHRV; \( P = 0.41 \)) for the prediction.

Head motion did not significantly predict clinical pain intensity ratings (\( P = 0.30 \)). In fact, when head motion was included in the prediction model as a fourth parameter, the correlation between true LBP and predicted LBP did not show much change (\( r = 0.63 \)) compared with our 3-parameter model where head motion was not included (\( r = 0.63 \)) (supplementary materials, Section S2; available at http://links.lww.com/PAIN/A671).

To explore the degree to which SVR accuracy was driven by maneuver-induced changes, we examined 2 other, new SVR models that did not randomize or reallocate data (ie, would not be driven by a maneuver effect wherein data sets contained some patients at pre-maneuver and some at post-maneuver): (1) SVR with pre-maneuver data only and (2) SVR with post-maneuver data only. Thus, these SVR models allowed us to test whether SVM weights (built from maneuver-induced changes) can be used to predict between-subject pain ratings in a data set with consistent timing relative to physical maneuver performance. Results showed that although the SVR performed with pre-maneuver data only yielded a poor correlation (\( r = 0.63 \)) between actual and predicted pain ratings, the SVR performed with post-maneuver data only yielded a much better correlation between actual and predicted pain ratings (\( r = 0.42 \), \( P = 0.002 \)).

### 4. Discussion

Although pain is inherently a subjective self-reported experience, there is growing need for objective biomarkers for pain. We

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**Table 1**

<table>
<thead>
<tr>
<th>Measures (cLBP, N = 53)</th>
<th>Values/scores</th>
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<tr>
<td>Age (y)</td>
<td>37.4 ± 11.3</td>
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<tr>
<td>Sex (male/female)</td>
<td>20/33</td>
</tr>
<tr>
<td>Pain duration (y)</td>
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<td>% using opioids</td>
<td>5.70%</td>
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<td>BDI</td>
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<td>BPSD</td>
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<td>PROMIS-physical function (T-score)</td>
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<tr>
<td>PROMIS-pain interference (T-score)</td>
<td>58.1 ± 5.5</td>
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<tr>
<td>PCS</td>
<td>11.7 ± 8.4</td>
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<tr>
<td>Back pain bothersomeness</td>
<td>5.0 ± 1.6</td>
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</table>

Back pain bothersomeness was collected on a VAS (0: “not at all bothersome” and 10: “extremely bothersome”). Data are shown as mean ± SD.

BDI, Beck Depression Inventory II (0–63 scale); BPSD, Back Pain-Specific Disability (0–10 scale); cLBP, chronic low back pain; PCS, Pain Catastrophizing Scale; PROMIS, Patient-Reported Outcomes Measurement Information System; VAS, visual analogue scale.

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**Figure 3.** Clinical pain intensity changes due to physical maneuvers. (A) Individually customized physical maneuvers significantly exacerbated low back pain levels in patients with cLBP (\( N = 53 \)). (B) Patients reported a wide range of baseline low back pain levels (pre-maneuvers), and after maneuvers, all included patients reported increased pain levels (post-maneuvers), which were maintained throughout the duration of the post-maneuver scans. N.b. Bar plots in (A) show mean ± SD. Each data point in (B) represents an individual patient. cLBP, chronic low back pain.
address a significant gap in the clinical pain research field by introducing and evaluating candidate predictive, combinatorial biomarkers for clinical pain intensity. Multimodal brain and autonomic physiology data were evaluated with machine learning–based prediction modeling in patients with cLBP. Individualized physical maneuvers were successfully implemented to exacerbate clinical back pain for post-maneuver brain imaging scan runs in the majority of patients with cLBP (87%, 62 of 71 patients). For these patients, maneuvers produced an average increase of 74.8% in clinical pain, allowing us to evaluate patients in a relatively lower vs higher clinical pain state. When combined, multimodal parameters (S1CONN, rCBF, and HF HRV) produced a synergistic effect, resulting in successful within-patient classification between relatively lower and higher clinical pain intensity states with high accuracy. Moreover, this model was successfully applied to predict between-patient clinical pain ratings with minimal prediction model overfitting and with independent training and testing data sets.23

Each putative biomarker targeted a unique physiological dimension of central and autonomic processing supporting pain perception. For instance, rCBF is obtained from ASL fMRI and captures slowly varying state changes in activity across the brain that may be linked to pain exacerbation. S1CONN was obtained from BOLD fMRI data and captured the temporal coherence (whole-brain connectivity) of the S1 representation of the low back, which is the primary encoding node for afferent nociceptive
SVR prediction of clinical pain

Figure 6. Prediction of clinical LBP ratings across patients with cLBP using all multimodal parameters. True LBP plotted against predicted LBP from SVR results demonstrated that our model was successfully able to predict between-subject clinical pain intensity ratings for both training (TRAIN, N = 53, r = 0.52) and independent testing data sets (TEST, N = 53, r = 0.63, shown above). cLBP, chronic low back pain; LBP, low back pain; SVR, support vector regression.

Classification was strongly influenced by the brain imaging features, and the SVM weighting maps inform the brain circuitry that directly supports higher clinical pain intensity. For instance, the rCBF parameter weighting map encompassed positive predictive weights in subcortical regions such as the thalamus, and cortical pain-processing regions such as the prefrontal cortex and ventral posterior cingulate cortex. Regions such as thalamus are known to process nociception and pain salience, whereas prefrontal cortex has been strongly linked to clinical pain perception. Importantly, combining these multimodal putative biomarkers produced a synergistic effect for clinical pain prediction, both within and between patients with cLBP.

The use of a within-patient model in our study through pain exacerbation maneuvers mitigates interindividual differences in how different patients interpret and use a numerical pain rating scale. Furthermore, extension of these multimodal data to a between-patient analysis using training and testing data sets allow for prediction of clinical pain intensity across different patients. The between-patient SVR was conducted through randomized 50% to 50% allocation of pre- and post-maneuver data into respective training and testing data sets, which mitigates effects such as passage of time, as the lower-pain state always occurred before the higher-pain state. When the SVR was performed without this randomized allocation, ie, SVR performed with pre-maneuver data only and a separate SVR performed with post-maneuver data only, the pre-maneuver data SVR demonstrated poor performance (r = −0.07), whereas the post-maneuver data SVR demonstrated significant actual-to-predicted pain correlation (r = 0.42). Our interpretation is that pre-maneuver ratings may have been influenced by a broad array of factors leading to substantial differences in how individuals use the 0 to 100 pain scale we presented to them. On the other hand, after completing the maneuvers, which increased their pain, all patients with cLBP had a consistent anchor as reference for how to rate their back pain—an explicit, physical maneuver–exacerbated back pain. This effectively normalized how patients used the 0 to 100 pain scale after maneuvers, leading to better prediction of clinical pain ratings by our SVR model, which was created using a maneuver-evoked SVM design. Further testing on future, independent samples would of course be needed to assess generalizability of this SVR model, but providing this anchor for ratings seems to be beneficial for prediction, and is in line with previous studies suggesting that clinical pain rating fidelity can be enhanced with training in how to consistently use pain scales, for example, by applying evoked pain stimuli as anchors.

Data fusion techniques for multimodal brain imaging data take advantage of combining unique aspects of each data modality’s contribution to enhance prediction. However, the use of such multimodal combinatorial prediction techniques has remained limited. We demonstrated synergistic performance when parameters from different modalities were combined. Our

cardiovagal modulation did significantly contribute to prediction of relatively higher vs lower clinical pain states. The primary reason behind low accuracy for the HF-HRV metric could be due to the single-feature nature of this metric, leading to a fit of the decision boundary in a unidimensional feature space (Fig. 1B). By contrast, the brain imaging metrics contained thousands of features/voxels. Indeed, future research should incorporate several autonomic metrics (both time-domain and frequency-domain cardiac information, galvanic skin response, and pupillometry) to allow for a multidimensional feature space and better model performance.

To the best of our knowledge, the analyses in this current study represent the first use of multimodal central and autonomic data to directly predict clinical pain states. One recent promising study used multimodal task-evoked brain imaging data to discriminate between patients with fibromyalgia and healthy controls. However, patients with fibromyalgia and healthy adults differ on much more than just the pain experience (eg, mood, cognitive task performance, etc.), and this study linked model prediction to pain intensity assessed at a different time point (an hour before brain imaging acquisition). Our study significantly extends such previous work by building a multimodal predictive model that provides diagnostic and clinical utility by directly predicting concurrent clinical pain states.

Although our results showed less robust classification accuracy for the autonomic HF-HRV metric, decreased

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multimodal approach could be further extended to include other brain imaging (eg, structural MRI, diffusion tensor imaging, positron emission tomography, chemical shift imaging, magnetoencephalography, and electroencephalography), physiological/autonomic activity (including time-domain and other frequency-domain metrics of HRV, galvanic skin response, and pupillometry), behavioral (eg, facial expression and body gesture), mood/affect parameters (eg, state-based pain catastrophizing and anxiety), and quantitative sensory testing measures, as well as experimental tasks related to different (eg, sensory, affective, and cognitive) aspects of pain.

Our research is in line with increased interest in predictive modeling with brain-based biomarkers. The clinical pain classification and prediction model created in this study could eventually lead to applications in clinical practice and could help predict pain intensity for clinical settings without the presence of patient-reported ratings (eg, noncommunicative patients). Furthermore, combining several non-neuroimaging parameters for prediction could lead to a more cost-effective and quicker approach to pain prediction in clinical practice. However, this field is still nascent—the current consensus around such multivariate predictive models is that they should not be used in lieu of subjective clinical pain ratings, but rather in conjunction with and supporting clinical pain ratings. Such neuroimaging approaches can be used to understand the underlying mechanisms of clinical pain. We hope that the model presented in this study will lead to future research to create well-validated predictive models from larger samples and multimodal features, eventually working towards clinical application.

Limitations to our study should also be noted. For instance, because successful maneuver-related pain increase and available data for all model parameters were limited to 53 patients, we also limited classification, regression, and parameter evaluation and comparison to these 53 patients only. Future applications should extend our predictive model to multiple sampling visits within a longitudinal trial framework. In addition, we did not include any control conditions (eg, healthy controls performing maneuvers). However, our previous study found that only patients with chronic back pain (and not healthy controls performing the same maneuvers) showed any changes in rCBF maps between the pre- and post-maneuver periods. Hence, we wanted to allocate resources toward larger sample size for this current study. A further limitation is that we used pulse signal from subjects’ fingers instead of ECG to compute HFHRV. This finger pulse signal was used, as it was not contaminated by MRI scanner noise. Finger pulse-based HFHRV is not as commonly used or evidence-supported as ECG; however, our group has successfully used pulse-based frequency-domain metrics in several recently published studies.

In conclusion, our machine-learning approach with a clinical pain exacerbation model found synergistic effects of using multimodal brain and autonomic markers in classification of clinical pain states and prediction of pain intensity. If the model is generalized across different chronic pain populations and different contexts, this pain signature could have great promise for pain assessment in noncommunicative patients, because successful maneuver-related pain increase and availability of objective pain endophenotypes that can be used in future longitudinal research aimed at discovery of new approaches to combat chronic pain.

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Appendix A. Supplemental digital content

Supplemental digital content associated with this article can be found online at http://links.lww.com/PAIN/A671.

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References


Conflict of interest statement

The authors have no conflict of interest to declare.