



# Individual variability in brain representations of pain

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**Characterizing cerebral contributions to individual variability in pain processing is crucial for personalized pain medicine, but has yet to be done. In the present study, we address this problem by identifying brain regions with high versus low interindividual variability in their relationship with pain. We trained idiographic pain-predictive models with 13 single-trial functional MRI datasets ( $n = 404$ , discovery set) and quantified voxel-level importance for individualized pain prediction. With 21 regions identified as important pain predictors, we examined the interindividual variability of local pain-predictive weights in these regions. Higher-order transmodal regions, such as ventromedial and ventrolateral prefrontal cortices, showed larger individual variability, whereas unimodal regions, such as somatomotor cortices, showed more stable pain representations across individuals. We replicated this result in an independent dataset ( $n = 124$ ). Overall, our study identifies cerebral sources of individual differences in pain processing, providing potential targets for personalized assessment and treatment of pain.**

The multidimensional experience of pain arises in the brain from highly distributed processes, including both serial and parallel processing of nociceptive input that ascends to the cerebral cortex via multiple pathways<sup>1–3</sup>. The distributed nature of pain processing is reflected in the fact that some brain regions have been reported consistently across pain studies (for example, primary and secondary somatosensory cortices (S1, S2), midcingulate cortex (MCC), thalamus and insula), whereas some have been reported less consistently (for example, prefrontal regions, cerebellum and basal ganglia)<sup>3,4</sup>. These regions probably play different, complex functional roles in pain<sup>2</sup>. Therefore, viewing a set of brain regions as one fixed core pain system is an oversimplification<sup>5</sup>. For example, lesions of brain areas known to be important for pain processing, such as the S2, anterior and posterior insula or anterior cingulate cortex (ACC), often do not affect the ability to perceive pain<sup>6,7</sup>, suggesting that pain processing in the brain has the feature of degeneracy: pain may arise as a function of activity in multiple pathways, which may differ across individuals. If so, different combinations of brain systems and pathways can be involved in constructing and processing an individual's experience of pain<sup>1</sup>. Furthermore, pain can be modulated by multiple factors, including attention<sup>8</sup>, emotional state<sup>9</sup>, social context<sup>10</sup>, past experiences<sup>11</sup>, sex<sup>12</sup> and others<sup>13</sup>. Such external and internal factors—along with individual differences in personality, self-regulation and coping ability, and more—may render some brain systems differentially important for pain in different individuals, and thus contribute to the

interindividual variability in pain processing and variable effects of treatment. Understanding this variation is critical for understanding the causes and implications of pain at the individual person level.

Although it is important to understand the individual variability of the brain representations of pain for both basic and clinical science, it remains unclear which brain regions have more consistent versus more variable representations of pain across individuals. There is a growing interest in personalized (or idiographic) brain mapping—previous studies have revealed individuals' distinct brain features that may be lost in group-level analyses<sup>14,15</sup>. This personalized mapping approach is particularly important for pain research because it will help not only to understand the neural mechanisms of pain perception and regulation, but also to identify personalized targets for intervention and make better decisions about treatment planning and selection<sup>16</sup>. However, most of the existing neuroimaging-based biomarkers of pain have employed a population-level predictive modeling approach that relies heavily on multivariate pattern information well conserved across individuals<sup>17,18</sup>. Although these brain markers are also useful for understanding pain on an individual basis, for example, as part of multidimensional pain assessments, they are not sufficient by themselves to capture the complexity of the neural mechanisms and representations of pain for each person<sup>1</sup>. For example, a population-level brain measure related to pain may capture pain in some individuals better than others or respond in the same way to two different brain maps from two individuals, but the underlying reasons for the response could be different.

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The present study encompasses two main objectives. First, we aimed to take a step toward personalized brain mapping of pain by employing idiographic predictive modeling. We trained a pain-predictive model for each individual from functional magnetic resonance imaging (fMRI) data of 404 individuals from 13 single-trial thermal pain datasets collected by 2 independent laboratories. Among these, 11 datasets have been used in previous publications and 2 are unpublished (Supplementary Table 1). The second objective was to identify pain-predictive brain regions that show high versus low interindividual variability in their pain representations. Previous studies mainly focused on brain regions that showed activation or deactivation patterns consistent across individuals. These studies, by design, could not identify brain regions that are important for pain processing but have idiosyncratic patterns (that is, the 'A' category in Fig. 1a). Thus, they provide an incomplete picture. To overcome this issue, we first identified pain-predictive brain regions based on their importance for within-individual pain prediction regardless of the between-individual consistency of the predictive weights (both 'A' and 'B' categories in Fig. 1a). We then quantified the interindividual variability in regional weight patterns. We replicated our findings in an independent dataset ( $n=124$ ) and showed that they were not driven by variation in study-level task or acquisition parameters or varying signal quality across brain regions.

Together, we provide a comprehensive list of important pain-predictive brain regions and characterize them in terms of their interindividual variability, elucidating which brain systems contribute to the interindividual variability in pain processing. The present study paves a path toward personalized brain mapping of pain and clinical interventions.

## Results

**Identifying brain regions important for pain prediction.** To find important pain-predictive brain regions (analyses 1–3 in Fig. 1a), we fitted linear support vector regression (SVR) models to data from each of the 404 participants in our dataset, creating 404 individualized pain-predictive maps (prediction-outcome correlation =  $0.34 \pm 0.01$  (mean  $\pm$  s.e.m.) with fivefold cross-validation), and identified important predictive features using bootstrap tests with 5,000 iterations for each predictive map. Unlike the population-level (or group-level) predictive modeling approach, which can identify only brain regions with multivariate pattern information consistent across individuals, our approach allows us to identify brain regions important for pain prediction within each individual, even if the brain regions have distinct patterns across individuals. With the  $P$  values from the bootstrap tests, we computed the mean ( $-\log(P)$ ) values for the whole brain across all individualized maps, and for further analyses we selected the top 10% voxels, of which the threshold was the mean ( $-\log(P)$ ) = 1.549. The selected voxels are displayed in Fig. 1b. We then parcellated the voxels into regions based on multiple anatomical atlases, identifying 21 regions as important for pain prediction (Fig. 1c). For further reference to other commonly used parcellations and brain signatures of pain, we calculated the overlap of the neurological pain signature<sup>17</sup> (NPS), stimulus intensity independent pain signature-1<sup>19</sup> (SIIPS1) and large-scale resting-state functional networks<sup>20</sup> with the selected voxels. We found larger overlap with our regions in SIIPS1, NPS, ventral attention and somatomotor networks than in other networks (Extended Data Fig. 1a). In addition, a preliminary analysis, in which we compared the prediction performance of individualized models and group-based models on three-session data from a single individual and data from study 14, suggested that individualized models achieve higher prediction accuracy than group-based models (Supplementary Fig. 1).

The 21 pain-predictive regions included (in alphabetical order): (1) anterior MCC (aMCC), (2) anterior middle insula (AMIns), (3) anterior middle operculum (AMOp), (4) basal ganglia (BG), (5)

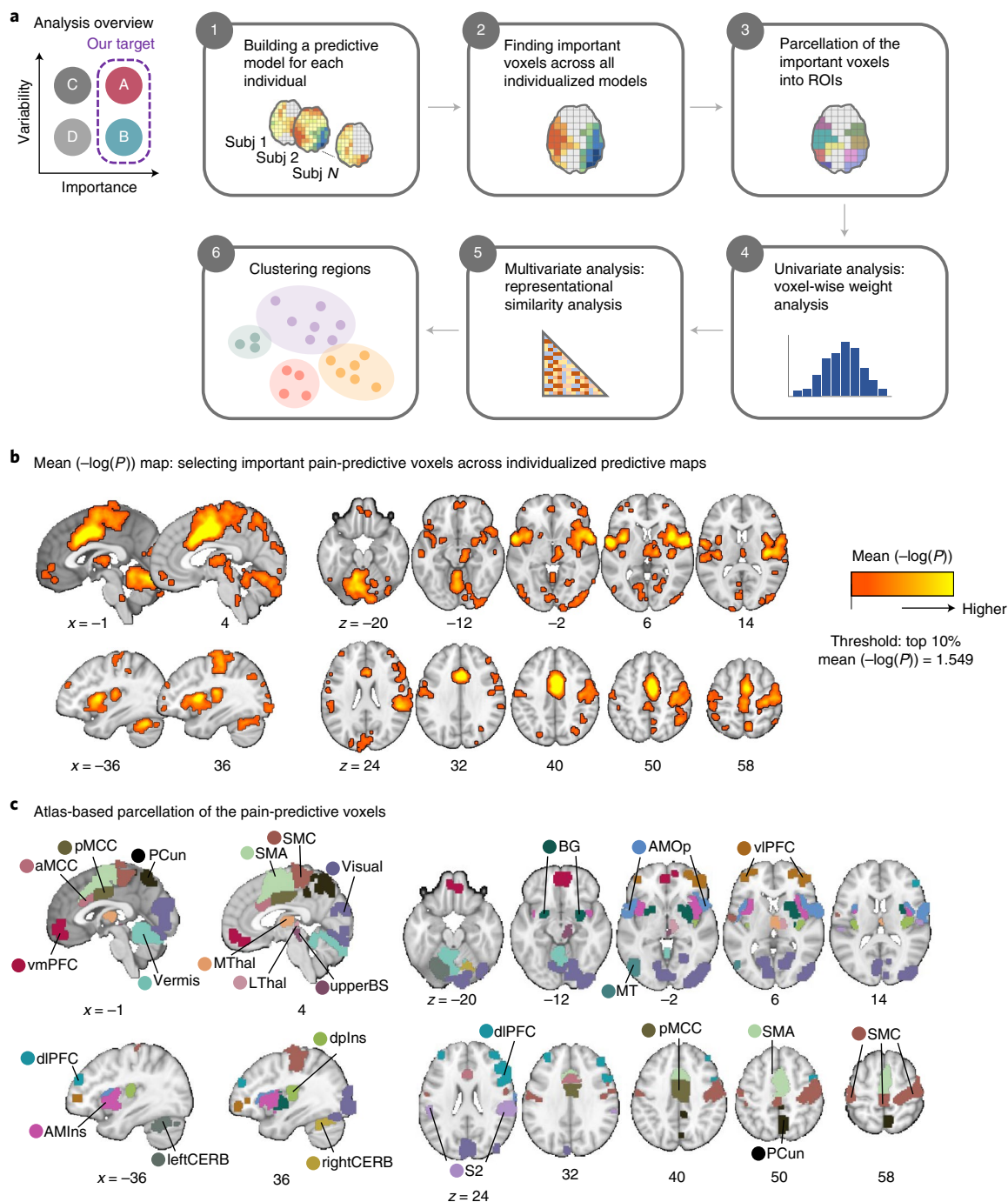
dorsal lateral prefrontal cortex (dlPFC), (6) dorsal posterior insula (dpIns), (7) left cerebellum (leftCERB), (8) lateral thalamus (LThal), (9) middle temporal area (MT), (10) middle thalamus (MThal), (11) precuneus (PCun), (12) posterior MCC (pMCC), (13) right cerebellum (rightCERB), (14) secondary somatosensory cortex (S2), (15) supplementary motor area (SMA), (16) sensorimotor cortex (SMC), (17) upper brain stem (upperBS), (18) vermis, (19) visual cortex (visual), (20) ventrolateral prefrontal cortex (vlPFC) and (21) ventromedial prefrontal cortex (vmPFC). We applied masks of these regions to the individualized predictive maps and used only the masked voxel data for further analyses. The individualized maps, region masks and other data and code are available at [https://github.com/cocoonlab/individual\\_var\\_pain](https://github.com/cocoonlab/individual_var_pain).

**Univariate analysis of the individual variability.** We first examined the voxel-wise individual variability of the predictive weights for the important pain-predictive regions using variance across all individualized maps (analysis 4). Before calculating the voxel-wise variance, we normalized each predictive map by dividing the weights by the spatial s.d. across the whole brain, rendering their scales comparable across images. The variance of the individual voxel weights is visualized in Fig. 2a. The region-level summary shown in Fig. 2c indicates that the vmPFC and MThal showed relatively high weight variance, whereas the MT, dpIns and pMCC showed low weight variance. For additional reference, we examined the voxel-wise variance in the large-scale, resting-state, functional networks<sup>20</sup>, NPS<sup>17</sup> and SIIPS1 (ref. <sup>19</sup>), finding the highest variance in the limbic network and the lowest variance in the dorsal attention network (Extended Data Fig. 1d).

The signs of averaged voxel weights were largely consistent with those in the previous literature. For example, brain regions such as the insula, S2, MCC and thalamus were, on average, positively predictive of pain, whereas the PFCs and PCun were negatively predictive (Fig. 2b,c)<sup>17,19</sup>. This result may seem to suggest that the brain regions have consistent roles in pain prediction across people. However, when examining the signs of median voxel weights in individuals, we found a large individual variability in the proportion of positive versus negative signs across people, indicating that regions were not strictly positive or negative in all subjects (Extended Data Fig. 2a). For example, the vmPFC consisted of negative weights, on average, but 36.1% of the subject median weights were positive, indicating variability in the sign of the multivariate pattern. Similarly, the dlPFC, MT, PCun and vlPFC showed negative average weights, although their median weights were positive in 40.8%, 44.8%, 35.4% and 48.5% of individuals, respectively. The most variable weight sign appeared in the visual cortex with 49% positive and 51% negative median weights (Extended Data Fig. 2b). This indicates substantial individual variability in the functional role and/or functional topography in relation to pain in these areas. Figure 2d depicts the relationship between the region-level variance and region importance.

**Multivariate analysis of the individual variability.** Although the univariate approach can provide useful voxel-level information about the individual variability of predictive weights, the multivariate patterns may convey more critical information about neural population codes of pain processing for each individual<sup>17,21</sup>. In addition, multivariate analyses have been shown to be more stable and less influenced by noise than univariate approaches<sup>22</sup>. Therefore, as a next step, we assessed the variability of the regional multivariate patterns across individuals using representational similarity analysis<sup>23</sup>, which allowed us to quantify the similarity between the spatial patterns of corresponding regions across different individuals (analysis 5).

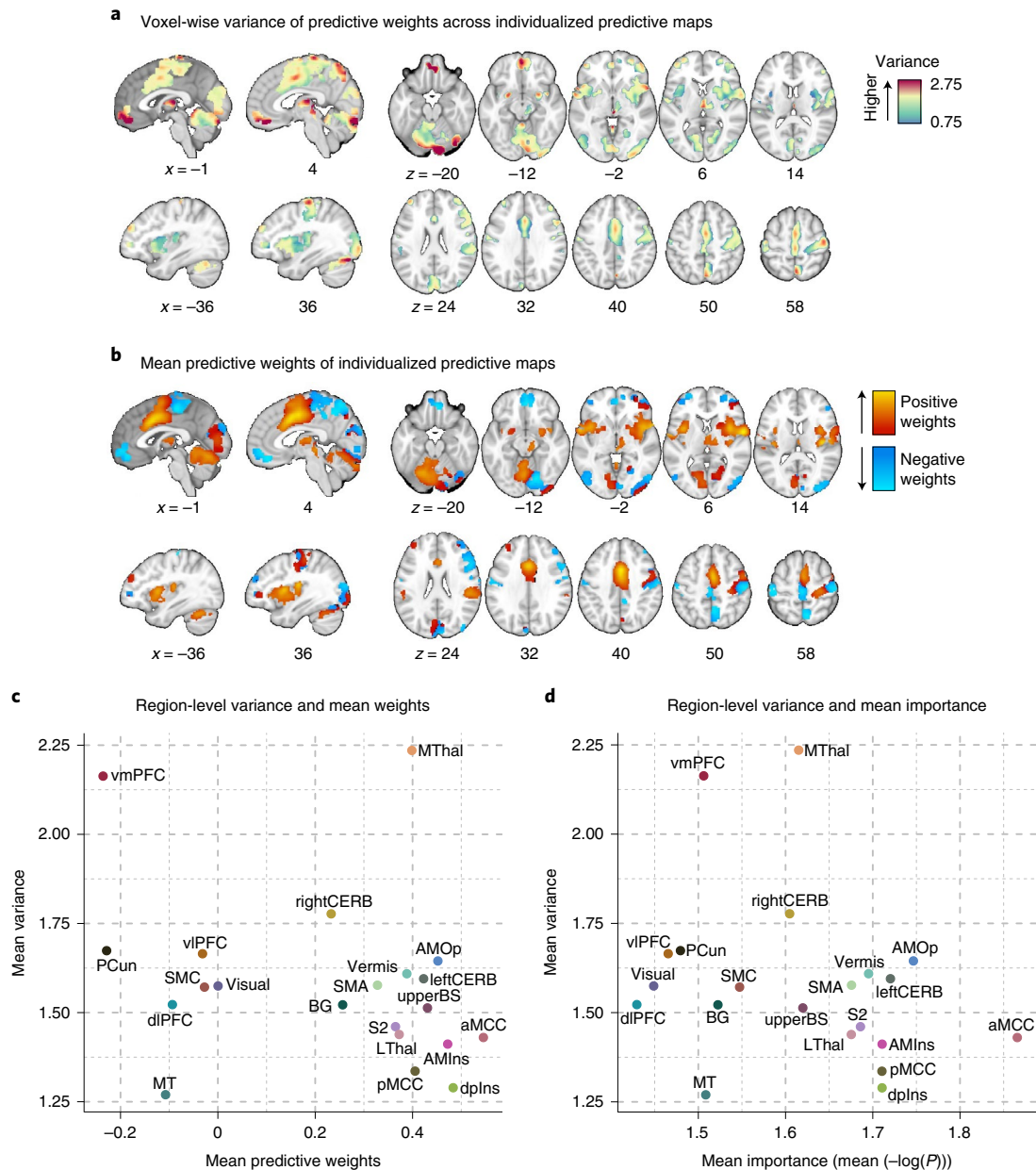
As shown in Fig. 3a, we first calculated the interindividual representational dissimilarity matrices (RDMs) as a measure of the interindividual distance for the 21 pain-predictive regions, using



**Fig. 1 | Analysis overview and important pain-predictive regions.** **a**, Analysis overview. We identified brain regions that were important for pain prediction and examined whether they showed variable or stable pattern representations across individuals ('A' and 'B' categories; top left). To this end, a series of analyses was conducted that could be divided into six steps. Detailed explanations about the analysis steps can be found in Results and Methods. ROIs, regions of interest. **b**, Selected voxels important for pain prediction in step 2 analysis. We operationalized the importance in terms of mean  $(-\log(P))$ , where the two-tailed  $P$  values were calculated from the bootstrap tests performed in each individualized predictive map. The selected voxels had mean  $(-\log(P))$  values higher than the top 10% mean  $(-\log(P))$  value ( $=1.549$ ). **c**, The important voxels parcellated into 21 regions based on the cortical and cerebellar atlases (see Methods for more details).

regional multivariate pattern information extracted from the 404 individualized pain-predictive maps (from 404 participants), resulting in 21  $404 \times 404$  RDMs. To ensure fair comparisons across different regions that have different locations and sizes, we normalized the RDMs (that is, obtained  $z$ -scores) using null RDM baselines for each region resulting from a permutation test (see Methods for more details).

Figure 3b displays the averages of the lower triangles of the normalized RDMs (that is, mean  $z$ -scores) against the number of voxels of the pain-predictive regions, suggesting a strong negative linear relationship between the mean representational distance and region size even after permutation-based normalization (Pearson's  $r = -0.537$ ,  $P = 0.012$ ). Thus, to further account for the effects of region size, we regressed out the effects from the mean



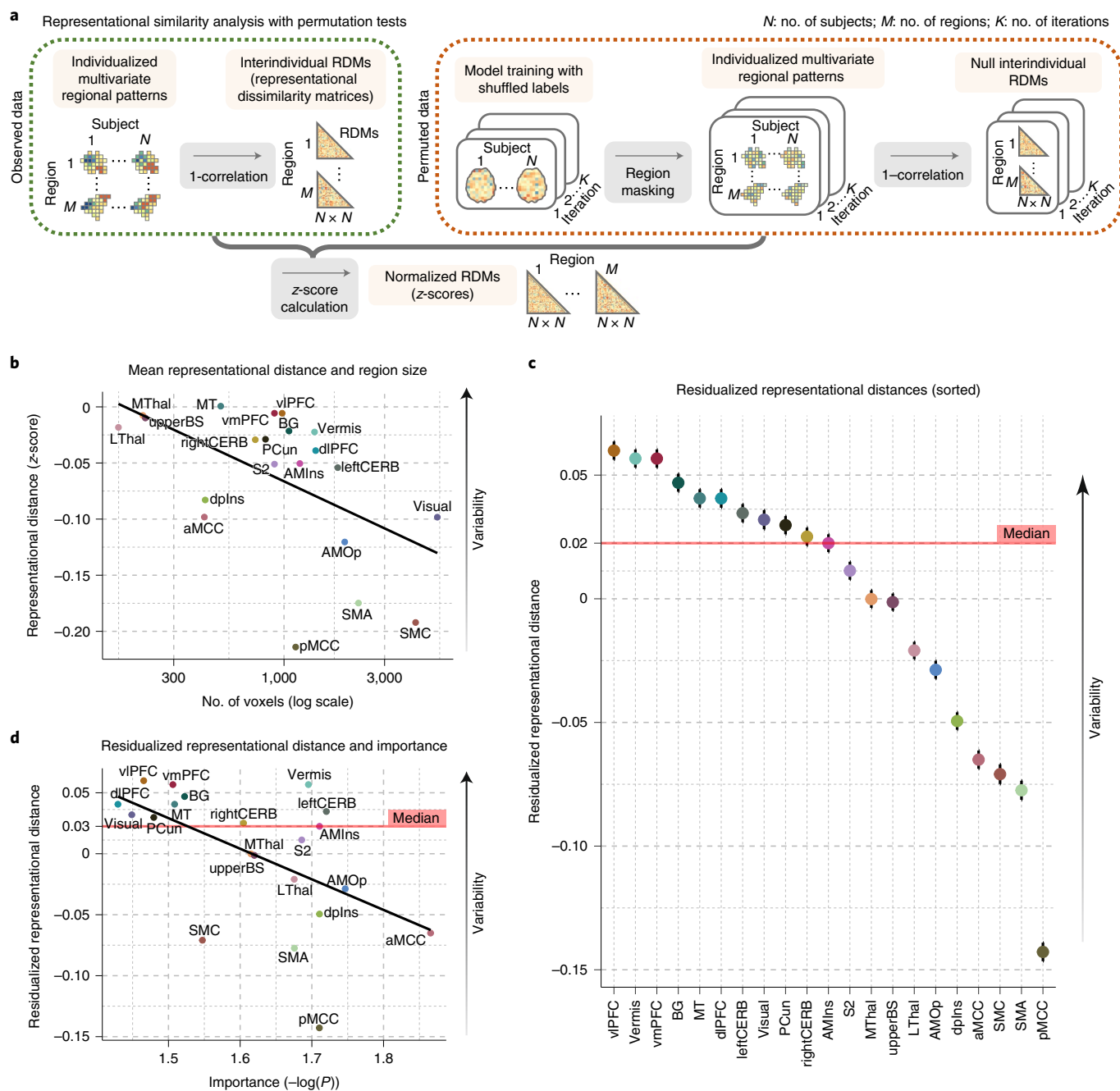
**Fig. 2 | Univariate analysis of the individual variability of predictive weights.** **a**, The voxel-wise variance of voxel weights across all individualized pain-predictive maps. **b**, The voxel-level mean predictive weights across the individualized maps. The positive weights were shown in warm colors (that is, positively predictive of pain) and the negative weights in cool colors (that is, negatively predictive of pain). **c**, The scatter plot shows the region-level summary with the mean predictive weights on the x axis and the mean weight variance on the y axis. **d**, The scatter plot displays the mean weight variance against the mean importance as  $-\log(P)$  (based on two-tailed  $P$  values).

representational distances. The residualized mean representational distance for each region is displayed in Fig. 3c, indicating that the vlPFC, vermis and vmPFC showed the highest individual variability whereas the pMCC, SMA and SMC appeared to be the most stable across individuals. Additional results without residualizing the region size are shown in Supplementary Fig. 2. For further reference, we performed the analysis in the NPS<sup>17</sup>, SIIPS1 (ref.<sup>19</sup>) and large-scale functional networks<sup>20</sup>. The limbic and visual networks manifested the highest interindividual variability, whereas the somatomotor and ventral attention networks and NPS showed the lowest variability (Extended Data Fig. 1e). Last, Fig. 3d shows the relationship between the residualized mean distance and mean importance (the mean  $-\log(P)$  values) of each region with Pearson's  $r = -0.542$ ,  $P = 0.011$ , suggesting that highly important

regions for individualized pain prediction also have more stable multivariate patterns across people.

We performed several analyses to assess the robustness of the results. Results from analyses run on a reduced dataset of individualized models with significant prediction performance only were significantly correlated with the results based on the full dataset at Spearman's  $\rho = 0.96$ ,  $P = 0.00001$  (Extended Data Fig. 3). We additionally tested whether each study had a significant effect on the final results (that is, the variability ranking across 21 regions) by removing one study from all analyses at a time (that is, a study-level jackknife), repeating this whole analysis for all 13 studies and comparing the results of a reduced set of studies with the original results of the full set of studies. Both the univariate and the multivariate analyses showed stable results even after removing each





**Fig. 3 | Multivariate analysis of the individual variability of predictive weights using a representational similarity analysis.** **a**, A representational similarity analysis for assessment of the interindividual variability of the regional multivariate patterns<sup>23</sup>. A detailed description of the analysis can be found in Multivariate representational similarity analysis. **b**, The scatter plot showing the relationship between the mean representational distance based on the permutation tests (y axis) and region size (displayed in the logarithmic scale on the x axis). The two variables showed strong negative correlation ( $r = -0.537$ ,  $P = 0.012$ , two-tailed). Higher representational distance values indicate higher pattern-level variability across people. **c**, Results of the representational similarity analysis after controlling for the region size. To account for the effects of region size on the representational distance, the region size effects were regressed from the representational distance. The plot shows the residualized representational distance, sorted from the highest distance, and the s.e.m. across all pair comparisons of individuals, that is,  $C(404, 2) = 81,406$ . **d**, The scatter plot showing the relationship between the residualized representational distance and the mean importance measured by mean ( $-\log(P)$ ) with Pearson's  $r = -0.542$ ,  $P = 0.011$ , two-tailed.

study. Spearman's correlation coefficient  $\rho$  ranged from 0.86 to 0.98 for the univariate results and from 0.98 to 0.99 for the multivariate results (Extended Data Fig. 4), suggesting that each study and study-specific parameters had minimal effects on the results. We also tested the effects of context manipulations present in some studies in our dataset, such as placebo or cognitive regulation. To assess the

impact of these manipulations, we performed the representational similarity analysis separately on subsets of studies with and without context manipulations. Results from studies with and without context manipulation were significantly correlated with the results based on the whole dataset with Spearman's  $\rho = 0.92$ ,  $P = 4.2 \times 10^{-6}$  and  $\rho = 0.83$ ,  $P = 4.3 \times 10^{-7}$ , respectively (Extended Data Fig. 5). In

addition, we investigated the effects of study-related variables on the interindividual variability within regions. We performed a multiple regression analysis with independent variables ( $x$  variables) being study-related factors such as the field strength of the scanner, data collection site, stimulated body site, stimulus duration, number of trials available for the participant and presence of pain-predictive cues, with the dependent variable ( $y$  variable) being the mean interindividual distance in each region of interest (Supplementary Table 2). Although some regression models showed sensible significant effects of some variables, the main findings presented in the present study were not affected after controlling for these effects (Spearman's  $\rho = 1.0$ ) because the analysis was conducted within each region and our main findings compared the interindividual variability across regions. Finally, we tested whether the pattern of results was influenced by excluding studies with low within-study prediction performance. We performed the analyses excluding three studies with the lowest average prediction performance and found that the overall pattern of results presented was preserved (Extended Data Fig. 6). Thus, in sum, the results are robust across studies that vary in psychological context and not driven by individual studies or studies with poorer overall prediction performance.

**Replication in a large-scale independent dataset.** To further validate and show that our findings are not a mere consequence of different experimental designs across multiple studies and varying signal quality across brain regions, we replicated our findings with a large-scale independent dataset ( $n = 124$ ) acquired at one location with a single experimental design. As in the discovery dataset, we fitted an SVR model to the single-trial data and obtained 124 individualized predictive maps (prediction-outcome correlation =  $0.68 \pm 0.02$ , mean squared error =  $0.014 \pm 0.001$  with fivefold cross-validation). We then applied the region masks, performed a multivariate representational similarity analysis on the masked regional data and regressed out the effects of region size on the mean representational distance. Figure 4a shows the residualized distance indicating that the highest interindividual variability was found in the dlPFC, visual areas and vmPFC, whereas the pMCC, leftCERB and SMC manifested the lowest variability. To compare this result with the result from the discovery dataset, we assigned a rank to each region based on the residualized distance in both datasets and measured the rank correlation between them. The rank correlation coefficient, Spearman's  $\rho$ , was significant at a value of 0.57,  $P = 0.008$  (Fig. 4b). We also evaluated whether the important pain-predictive voxels identified in the discovery dataset also replicate in this replication dataset. We found a significant phi correlation,  $\phi = 0.502$ ,  $P < 2.2 \times 10^{-16}$ , suggesting high similarity between the two binarized voxel importance maps (Supplementary Fig. 3). In addition, we performed a preliminary test of within-individual reliability of the predictive patterns by splitting the data for each individual into two folds (that is, the first four and the last four runs), training models on both folds and evaluating the similarity of the whole-brain and regional predictive patterns. The pattern similarity showed mostly medium-to-large correlations, providing preliminary evidence of within-individual reliability of the predictive patterns (Supplementary Fig. 4a). We also performed the same analysis on a preliminary four-session dataset from a single participant, obtaining further support for within-individual reliability (Supplementary Fig. 4d).

As the signal quality in the fMRI data may vary across brain regions, which in turn may affect the regional variability, we examined whether the temporal signal:noise ratio (tSNR) of our data correlated with our results. To this end, we first calculated the tSNR for each participant as the mean of images acquired at each repetition time divided by the s.d.. Subsequently, we obtained a group-average tSNR map (Extended Data Fig. 7) and calculated the mean tSNR for each region. We then assigned ranks to regions based on the tSNR

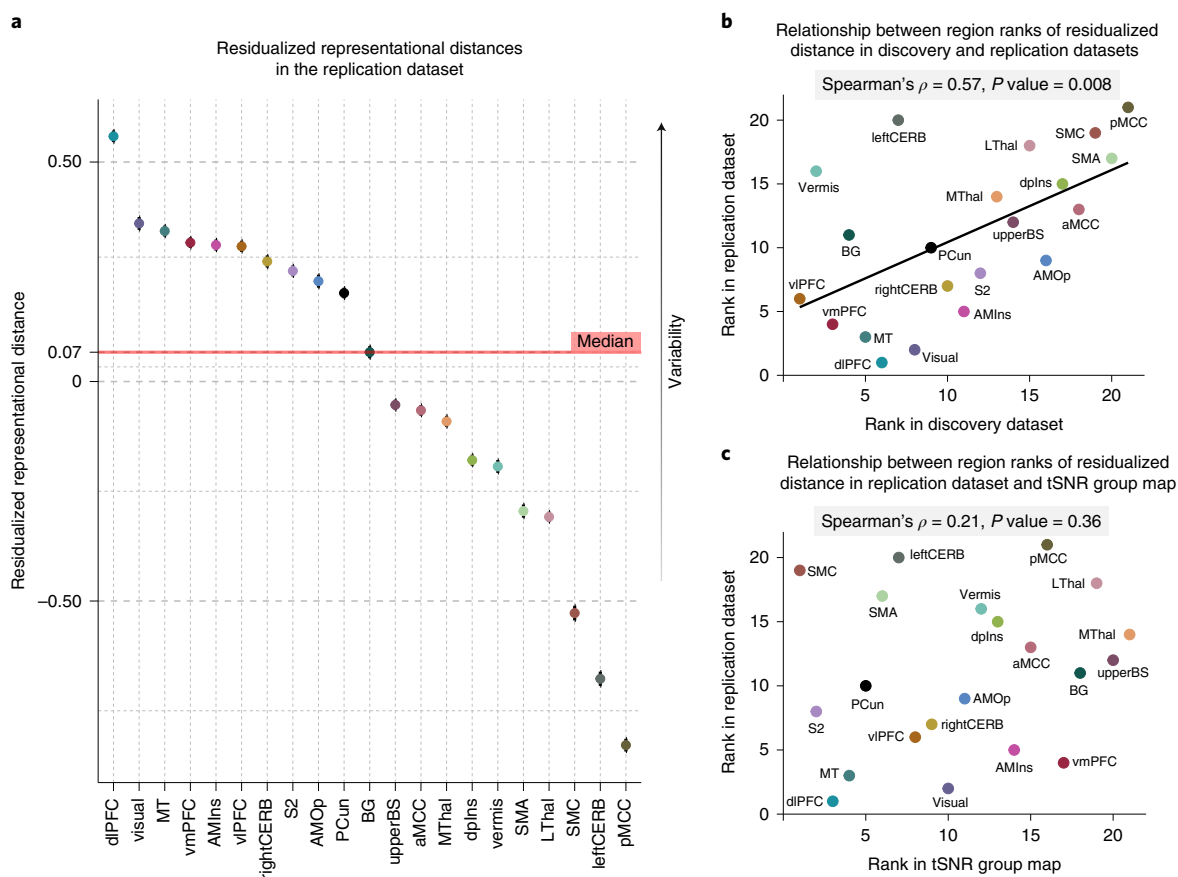
and compared them with the ranks based on the residualized representational distance using rank correlation. The comparison suggested that the regional variability cannot be explained by varying tSNR across regions (Spearman's  $\rho = 0.21$ ,  $P = 0.36$ ; Fig. 4c).

**Clustering pain-predictive brain regions.** Next, we compared the regional interindividual variability profiles using the nonparametric rank correlation, Kendall's  $\tau_A$ , to identify clusters of brain regions that exhibited similar patterns of the representational distance across individuals (analysis 6). This correlation matrix provides the representational connectivity (as termed by Kriegeskorte et al.<sup>23</sup>) across 21 pain-predictive regions, which quantifies the representational information shared across regions. This approach can be regarded as a type of multivariate connectivity that allows us to group multivariate pattern representations using clustering analysis. The Kendall's  $\tau_A$  values ranged from  $-0.005$  to  $0.17$  (Fig. 5a), with higher values indicating higher similarity in the individual variability patterns for a pair of regions. For example, relatively high similarity was found between the AMOp and AMins with Kendall's  $\tau_A$  of 0.17, between the SMA and pMCC with the value of 0.155 or between the MThal and LThal with the value of 0.112.

To cluster the 21 pain-predictive regions based on the patterns of interindividual representational similarity, we first transformed Kendall's  $\tau_A$  into a distance matrix and performed nonmetric multidimensional scaling (NMDS). NMDS scores were then used as input features for hierarchical clustering. To select the optimal number of clusters ( $k$ ), we compared the clustering quality scores (silhouette values) against the null-hypothesis baseline obtained from a permutation test for  $k = 2-15$  solutions (see Methods for details). The optimal  $k$  was 10 clusters, which performed at the mean silhouette score of 0.59 (from the range 0–1, where 1 indicates perfect separability of clusters) with  $z = 3.72$ ,  $P = 0.0002$  over the null baseline clustering quality (for more details, see Extended Data Fig. 8). The identified clusters of regions included: the aMCC, pMCC, SMA and SMC for cluster 1; the leftCERB, rightCERB, vermis and visual cortex for cluster 2; the dpIns and S2 for cluster 3; the AMIns and AMOp for cluster 4; the vlPFC and dlPFC for cluster 5; and the MThal, LThal and upperBS for cluster 6. Also, the BG, MT, PCun and vmPFC each constituted a stand-alone cluster.

For display, we performed  $t$ -distributed stochastic neighborhood embedding ( $t$ -SNE) on the NMDS scores and plotted it with Kendall's  $\tau_A$  values transformed into weights connecting the regions in color-coded region clusters (Fig. 5b). With thresholding to show only the top 25% of connection weights, the vmPFC was not connected to any other region, suggesting that it showed a unique pattern of individual variability of predictive weights distinct from other brain regions (bottom panel of Fig. 5b). After computing the mean residualized representational distance of the clusters, the vmPFC and d/vlPFC clusters showed the highest distance among the region clusters, and the aMCC/pMCC/SMA/SMC and S2/dpIns clusters showed the lowest distance (Fig. 5b). For further illustration of the interindividual variability in the region clusters, we applied the predictive patterns of the region clusters from each individual to the activation maps of all other individuals to predict the corresponding pain ratings (Extended Data Fig. 9). The results show that the cross-individual prediction performance correlated with the interindividual variability of the clusters, with the former being more reliable in the less variable clusters. In other words, along with our results in Supplementary Fig. 4 in which we showed that predictive weights are reliable within individuals, these results support the idea that the interindividual variability of the regional predictive weight patterns is driven by individual variability in brain representations of pain, not by mere degeneracy properties common in high-dimensional data modeling or noise.

We also explored the relationship between these clusters and cortical hierarchy levels identified using unimodal-to-transmodal



**Fig. 4 | Replication of the multivariate representational similarity analysis in an independent dataset and tSNR in the dataset.** To validate the previous results, we employed an independent replication dataset acquired under the same experimental settings in the same location. **a**, Results of a representational similarity analysis performed in the replication dataset after regressing out the effects of region size on the mean representational distance. The plot depicts the residualized representational distance (a measure of interindividual variability in neural patterns) in each region, with higher values representing higher regional variability. **b**, Comparison of the results in discovery and replication datasets. Based on the residualized distance, we assigned ranks to all regions in both the discovery and the replication datasets. The scatter plot visualizes the statistically significant relationship between the results in the two datasets. **c**, The relationship between the region ranks based on the residualized distance and mean tSNR. To further inspect whether our results in the replication dataset are influenced by a different tSNR in different regions, we calculated the mean tSNR in all regions. The scatter plot compares the region ranks based on the tSNR and residualized representational distance, suggesting that the regional variability in the replication dataset cannot be explained by varying tSNR in the regions.

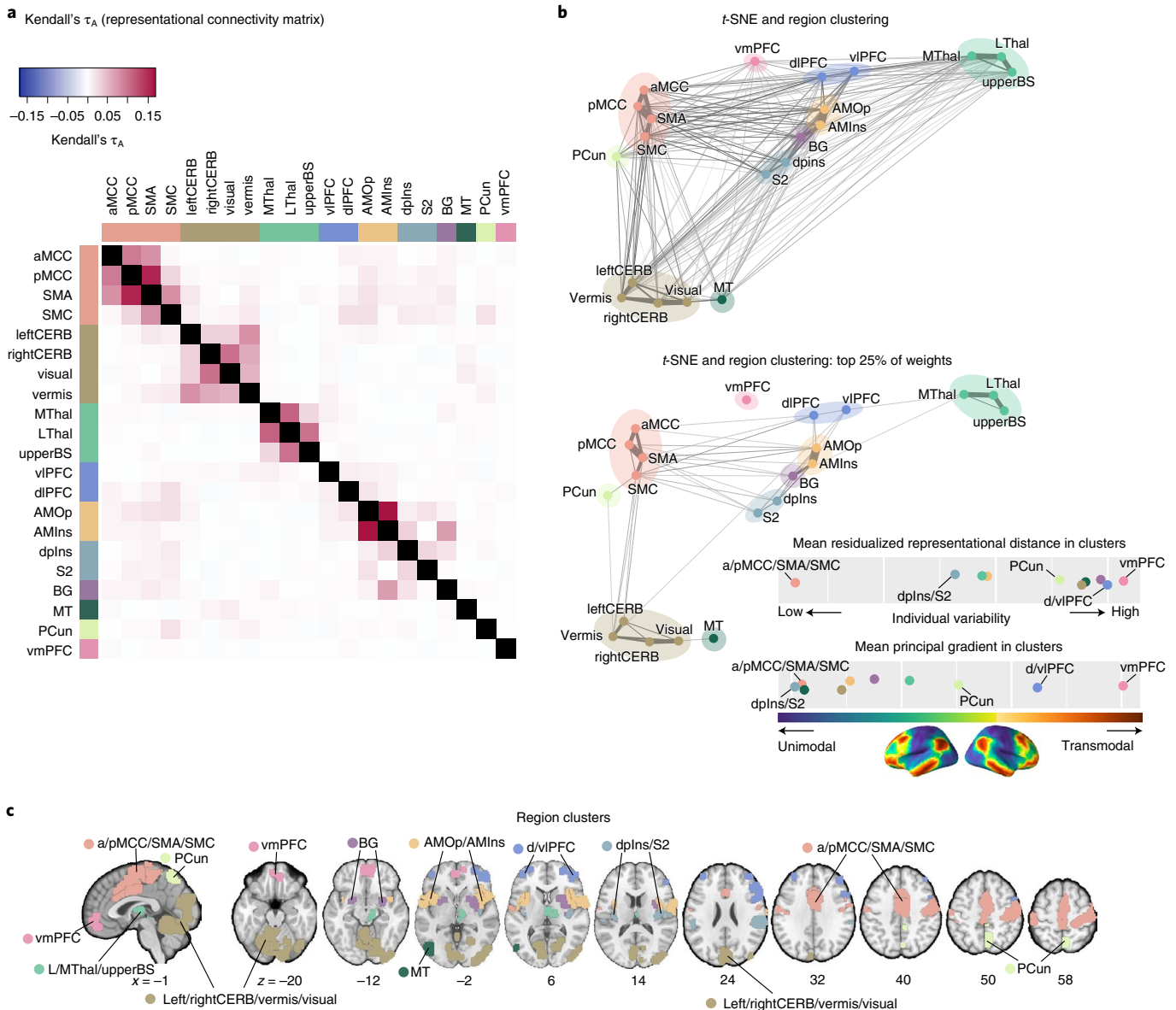
connectivity gradients<sup>24</sup>. In the present study, the principal connectivity gradient captures the diversity of functional connectivity patterns on a spectrum ranging from unimodal to higher-order transmodal brain structures<sup>24</sup>. We calculated the mean principal gradient in our region clusters based on a principal gradient map derived from an independent resting-state fMRI dataset ( $n=59$ ). The results showed that the mean residualized distances were significantly correlated with the mean principal gradients (Spearman's  $\rho=0.68$ ,  $P=0.04$ ; Extended Data Fig. 10), suggesting that the areas in which pain representations are more variable across individuals are higher-order transmodal regions located at the top of the principal gradient, whereas those that are stable across individuals are unimodal sensory regions located at the bottom of the cortical hierarchy.

## Discussion

Our study investigated the heterogeneity of individualized brain representations of pain based on fMRI data from a large sample of  $n=404$  for discovery and  $n=124$  for replication (total  $n=528$ ). Across all individualized pain-predictive maps, we first identified 21 predictive regions. Among these, the prefrontal and cerebellar regions, including the vmPFC, vlPFC, dlPFC, vermis and leftCERB, showed high interindividual variability in both univariate and

multivariate analyses. In contrast, the cingulate, somatomotor and insular cortices, including the a/pMCC, SMA, SMC and dpINS, showed lower interindividual variability. Importantly, this finding was successfully replicated in an independent dataset of  $n=124$ , demonstrating (along with multiple supplementary analyses) that the pattern heterogeneity was not a mere effect of various experimental settings or noise. Clustering regions based on interpersonal representational distance in multivariate predictive patterns revealed ten groups of regions, with more interpersonally variable groups lying on the transmodal end of the spectrum of the principal gradient of cortical organization and more stable groups on the unimodal end. Overall, the present study aims to step toward individualized pain brain mapping by providing brain targets that hold the potential to be a basis for further development of individualized clinical interventions and brain-based subtyping of individuals in pain representations.

Most of the important pain-predictive brain regions identified in the present study have also been reported in previous studies<sup>3,4</sup>. In addition, the signs of group-level predictive weights (that is, whether a brain region is positively or negatively correlated with pain ratings) were largely consistent with previous studies that used experimental acute pain stimuli<sup>25,26</sup>. However, we also found large



**Fig. 5 | Clustering of the pain-predictive brain regions based on the patterns of representational distance. a**, Heatmap showing the representational connectivity matrix (Kendall's  $\tau_A$ ) across 21 pain-predictive regions. Higher Kendall's  $\tau_A$  values (shown in darker red) indicate higher similarity between regions. **b**, A hierarchical clustering analysis of brain regions using the ten dimensional NMDS scores based on the representational connectivity matrix, resulting in ten region clusters. The plot shows the t-SNE map based on the ten dimensional NMDS scores and the ten region clusters are shown using different colors. The line widths indicate the relative connectivity strengths across the brain regions. The top panel shows all the connections, whereas the bottom panel shows only the top 25% of the connections. The upper bottom right chart displays the mean residualized representational distance of the region clusters (color coded). Furthermore, in the region clusters we calculated the mean principal gradient that ranges from unimodal to higher-order transmodal areas as suggested by the literature<sup>24</sup>. The lower chart, then, shows the mean principal gradient in the region clusters, suggesting that more variable clusters are located at the top of the principal gradient spectrum, whereas more stable clusters are located at the bottom. The bottom cortex map visualizes the principal gradient map obtained from an independent dataset ( $n = 59$ ). **c**, The region clusters visualized on a brain underlay.

individual variability in the predictive weight signs of the individualized pain-predictive maps (for example, Extended Data Fig. 2), suggesting that each region's functional role in pain processing varies across individuals. Therefore, our findings indicate that the individualized functional relationships between brain regions and pain perception are more complicated than how they are usually summarized at the group level. Although group-level summaries certainly have their own merits and strengths, such as canceling out noise and identifying findings that are most reproducible across individuals, the idiosyncratic patterns of pain-predictive weights that will be crucial to understanding the personal pain experience are

often ignored. However, our approach allowed each individualized pain-predictive map to have different signs and patterns of weights by focusing on group-level averages of the 'importance' measure. For example, the visual cortex might not often appear in pain neuroimaging studies, perhaps because of its mixed relationship with pain perception, as shown in Extended Data Fig. 2. However, our results suggest that the visual cortex is also an important contributor to pain prediction consistently across people. The functional role of the visual cortex could be manifold, for example, multisensory integration<sup>27</sup>, task related<sup>28</sup> or enhanced attentional state as a consequence of a salient event<sup>29</sup>.



Combining results from both the univariate and the multivariate analyses, we found that the brain regions that showed stable patterns across individuals were mostly those receiving direct inputs from the spinothalamic pathways, including the MCC (aMCC and pMCC), SMC, SMA and insula (dpIns and AMIns)<sup>30,31</sup>. In contrast, the brain regions that showed variable patterns across individuals were the prefrontal and cerebellar regions, including the vlPFC, vmPFC, vermis and rightCERB. Among these regions, the MCC appeared to have the most stable predictive weights in both the univariate and the multivariate analysis results. The MCC is known to work as a hub for processing of the affective–motivational components of pain and behavioral control of pain-related responses<sup>32,33</sup>. In particular, the MCC receives direct inputs from the medial thalamus<sup>30</sup>, and it has been recognized as part of the medial pain pathway, which has been associated with processing of the affective–motivational component of pain<sup>34</sup>. However, whether the medial pathway is specialized in processing this particular pain component has been challenged<sup>33</sup>. It has also been shown that the MCC contains neurons responding specifically to nociception<sup>35</sup> and exhibits generalizable multivariate pain-specific representations<sup>36</sup>. Another brain region with a stable representation across individuals was the dpIns, which is among the regions that best track the perceived intensity of pain<sup>37</sup> and are most consistently reported in pain neuroimaging studies<sup>38</sup>. Supporting the role of the dpIns in nociception, research in monkeys has revealed direct nociceptive-specific projections from the spinothalamic system to the posterior part of the insula encoding modality, location and intensity of the incoming stimulus<sup>39</sup>.

On the other hand, the vlPFC and vmPFC had the most variable representations across individuals. These brain regions are known to be important for cognitive pain modulation, such as reappraisal and placebo<sup>40,41</sup> and pain catastrophizing<sup>42</sup>. In particular, the vmPFC is important for autobiographical memory representations, valuation<sup>43</sup> and endogenous cognitive and affective processes<sup>44,45</sup>, supporting the possibility that these prefrontal regions subservise individually unique representations of pain. High between-study variability in frontal areas, probably stemming from different study contexts, was also reported in a recent meta-analysis on placebo analgesia<sup>46</sup>. In addition to the prefrontal regions, striatal and cerebellar regions also showed variable representations across individuals. Given that the cerebellar contributions to pain have only recently become of interest<sup>47</sup>, further research will be required to fully understand our findings in the cerebellum.

Importantly, these findings were replicated using an independent dataset. Unlike the discovery dataset, the replication dataset was collected under a single experimental setting; thus, it provided an appropriate basis to test whether the idiosyncrasy is a mere effect of the heterogeneity of the discovery dataset or reflects actual characteristics of the regions. Along with the evidence that the regional tSNR is not related to individual variability, study-level jackknife and other supplementary analyses, the replication results provide supporting evidence for our findings in the discovery set.

There was also a slight discrepancy between the results of the univariate and multivariate analyses. In particular, the MThal had the most variable representation in the univariate analysis, whereas it was below the median (that is, less variable) in the multivariate analysis. In addition, the MT was stable in the univariate analysis, but more variable in the multivariate analysis. These discrepancies may stem from the different nature of the analyses. The univariate analysis considers individual voxels separately and, therefore, becomes more vulnerable to outliers. However, the multivariate analysis considers the whole pattern of voxel activations and thus should be more stable than the univariate analysis and capture representational features lost in the univariate analysis.

We also identified ten region clusters based on the patterns of interindividual variability. The region clusters were mostly composed of regions that are anatomically and functionally adjacent,

for example, S2 and dpIns, thalamus and brain stem. However, the vmPFC, PCun, MT and BG each constituted a stand-alone cluster, which may reflect their high individual variability and unique representations of pain. It is interesting that we also observed a relationship between the interindividual variability of the clusters and the principal gradient of cortical organization, based on functional connectivity patterns<sup>24</sup>. The gradient spans from unimodal sensory cortical regions that process concrete direct percepts to higher-level transmodal regions that integrate and process abstract information<sup>48</sup>. Along with this notion, region clusters with lower interindividual variability were found in the unimodal end of the gradient spectrum, including dpIns/S2 and aMCC/pMCC/SMA/SMC clusters that receive direct spinothalamic nociceptive inputs. Many of these (particularly dpIns and S2) are unimodal, in that they are highly selective for somatosensory input or even nociceptive input (as with dpIns) more specifically. On the other hand, region clusters that showed higher variability, such as vmPFC, dlPFC/vlPFC or PCun clusters, were in the transmodal end of the spectrum, suggesting that regions in the higher end of the cortical hierarchy show more individually specific representations. It is noteworthy that this finding is consistent with the findings of Finn et al.<sup>49</sup>, in which the higher-level transmodal regions including the frontoparietal and default mode networks performed best at identifying functional connectivity fingerprints.

The region clusters provided in the present study hold a potential to be, ultimately, used as targets for brain-based biotyping for pain or personalized clinical interventions. Interest in precision medicine and individualized brain mapping has recently increased<sup>14,15,49</sup> because such individualized approaches could potentially improve the effects of treatment by tailoring it to the needs of each individual and provide evidence for disease subtypes. Therefore, as an example of a benefit of our approach, knowing which brain regions are important for pain processing and whether their pain representations are variable or consistent across individuals could help in making decisions about the deep brain stimulation or transcranial magnetic stimulation locations for chronic pain treatment<sup>50,51</sup>. The knowledge of which regions show high or low interindividual variability could also be useful for biomarker development and decoded neurofeedback<sup>1,52</sup>. As the outcomes of regions with different levels of interindividual variability could vary, these strategies could be informed of which brain regions could be more reliable targets across a population and which are better suited for personalized approaches. Moreover, a preliminary analysis suggested that there may not be one universal set of biotypes across the whole brain, but, in fact, different brain regions may provide different individual biotyping (Supplementary Fig. 5). Thus, it is imperative to understand the complex relationship between the regional (or network-level) brain representations and individual differences. Furthermore, the source of idiosyncrasy in these regional pain representations remains an intriguing and important question. There could be numerous underlying causes such as genetic factors<sup>53</sup>, individual differences in pain coping strategies<sup>54</sup> or pain sensitivity<sup>55</sup> and personality traits<sup>56</sup>, among others. However, to probe all possible explanations of the interindividual variability, large quantities of data, including detailed phenotypic data, genetic variants and other psychophysical and psychological measures, will be necessary.

The present study has several limitations. First, although we performed nonlinear spatial normalization, there are substantial interindividual differences in functional topography even when brains are anatomically aligned. Such variability could contribute to the higher level of individual variability we observed in prefrontal and other regions. To further address this issue, future studies could use functional alignment methods such as hyperalignment<sup>57</sup>. In addition, the amount of data per subject is relatively small and varies across studies, which could potentially negatively affect the robustness and reliability of the individualized models. Although our

preliminary results suggest that most of the important pain-predictive regions showed reliable weights within individuals across multiple sessions, it is also possible that some regions show ‘state-like’ variability over a short period of time, if they are highly context sensitive. Disentangling which regions capture intersession variability (‘states’) versus which regions manifest high stability across sessions (‘traits’) is an important research question, which should be addressed in the future. Future studies with a dense sampling design (that is, small *n*, and many repetitions within person) may be able to help provide additional answers to the questions raised by the present study. Also, this study concerns only one dimension of pain experience, which is pain intensity. Although pain intensity usually shows high correlations with other dimensions, such as unpleasantness<sup>58</sup>, the present study did not explicitly consider other dimensions of pain. Therefore, caution is needed when applying the current findings to other dimensions of pain experience. Furthermore, due to the relatively low temporal resolution of fMRI, some processes such as anticipation of the stimulus may not be well resolved and thus they may influence interindividual variability. Therefore, future studies may benefit from using other modalities, such as electroencephalography, electrocorticography or magnetoencephalography. Finally, due to a lack of more comprehensive phenotypic and genetic data, we could not fully test the origin and meaning of the intersubject variability observed in the present study. This poses an important question that should be addressed in future studies by gathering a large quantity of detailed phenotypic data and other measures that could be related to individual variability.

In conclusion, our study characterized the cerebral contributions to individual variability in pain processing by identifying brain regions that show high versus low interindividual variability in pain representations. Our results shed light on the personalized mechanisms of pain processing and could be potentially used in further classification and biotyping of individuals in pain processing, creating a path toward personalized pain medicine.

### Online content

Any methods, additional references, Nature Research reporting summaries, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41593-022-01081-x>.

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## Methods

**Datasets.** In the present study, we employed datasets from a total of 14 previous thermal pain studies using fMRI, of which studies 1–13 were used as a discovery dataset with a total  $n=404$  and study 14 as a large-scale independent replication dataset of  $n=124$ . Detailed information on the datasets is provided in Supplementary Tables 1, 2 and 3. Among these, 11 datasets have been used in previous publications and 3 (studies 7, 12 and 14) are unpublished (Supplementary Table 1). The institutional review board of Columbia University, the University of Colorado Boulder and Sungkyunkwan University, and the Ethics Committee of the Medical Chamber Hamburg approved all the studies. All participants provided written informed consent and were financially compensated for their participation.

**Participants.** The present study included a total of 528 healthy, right-handed participants from 14 independent studies, with sample sizes ranging from  $n=17$  to  $n=124$  per study. Descriptive statistics on age, sex and other details of the studies are provided in Supplementary Table 1. Participants were recruited from New York City and Boulder/Denver Metro Areas in the USA, Hamburg in Germany and Suwon in South Korea. The preliminary eligibility of the participants was determined through an online questionnaire, pain safety-screening form or an MRI safety-screening form. Participants with psychiatric, physiological or pain disorders, neurological conditions and MRI contraindications were excluded before enrollment.

**Procedures.** In each study, participants experienced a series of contact heat stimuli and rated their pain experience after each stimulus. The stimulation sites, number of intensity levels and corresponding temperature, stimulus duration, rating scales and number of trials used for analyses are provided in Supplementary Table 3. Each study also included psychological manipulation, such as predictive cues or placebo manipulation. However, in the present study we focused only on within-subject pain prediction across all trials regardless of any study-specific psychological manipulation.

**Thermal stimulation.** In all studies, thermal stimulation was delivered to multiple skin sites using a TSA-II Neurosensory Analyzer or Pathways system (Medoc Ltd) with a 16-mm Peltier thermode endplate (32 mm only in study 10). In every trial, after the offset of stimulation, participants rated the magnitude of their sensation of warmth or pain on a visual analog scale or a labeled magnitude scale. Other thermal stimulation parameters varied across studies, with temperatures ranging from 40.8 °C to 49.3 °C and the duration of the stimulation ranging from 1.85 s to 20.16 s. The stimulation parameters of all studies are provided in Supplementary Table 3.

**Preprocessing of fMRI data.** Preprocessing of functional images was performed using Statistical Parametric Mapping (SPM) software (<http://www.fil.ion.ucl.ac.uk/spm>). In addition to SPM, study 14 also used FMRIB Software Library (FSL) (<https://fsl.fmrib.ox.ac.uk>) and Independent Component Analysis-based strategy for Automatic Removal Of Motion Artifacts (ICA-AROMA)<sup>59</sup> software (<https://github.com/maartenmennes/ICA-AROMA>) for distortion correction and removal of motion-related artifacts, respectively. Except for studies 3, 5 and 7, which used SPM v.5, and study 14, which used SPM v.12, all other studies used SPM v.8 (see Supplementary Table 4). In SPM, structural T1-weighted images were co-registered to the mean functional image for each subject using the iterative mutual information-based co-registration method and then normalized to the Montreal Neurological Institute space. In studies 3 and 5, additional normalization to the group mean was performed after the SPM normalization using a genetic algorithm-based normalization<sup>60–62</sup>. To stabilize the image intensity, multiple initial volumes were removed in every functional dataset (see Supplementary Table 4 for the number removed in each study). To identify the outliers in the signal, we computed the Mahalanobis distances for the matrix of the concatenated slice-wise mean and s.d. of the intensity values by functional volumes (over time). Values with a significant  $\chi^2$  value (after correction for multiple comparisons with either false discovery rate (FDR) or Bonferroni's correction) were considered to be outliers. In practice, <1% of the volumes were outliers. The timepoints identified as outliers were later included as nuisance covariates in the first-level models. Functional images were then corrected for differences in slice timing and motion (realignment). Finally, the images were warped into the SPM's normative atlas, interpolated to  $2 \times 2 \times 2$  mm<sup>3</sup> voxels, and smoothed with an 8-mm full width at half-maximum (FWHM) Gaussian kernel in studies 1–13 and 5-mm FWHM in study 14. The fMRI data distribution was assumed to be normal. Although this was not formally tested, we conducted a detailed data quality check using the `fmri_data.plot.m` function in <https://github.com/canlab/CanlabCore>.

**Single-trial analysis: studies except for studies 2 and 5.** In each study, we modeled the data using a single-trial, or 'single-epoch', design and analysis approach. The single-trial response magnitude for each voxel was estimated using a general linear model design matrix with separate regressors for each trial, as in the 'beta series' approach. First, boxcar regressors convolved with the canonical hemodynamic response function (HRF) were constructed to model the cue, pain and rating periods in each study. Then, a regressor for each trial and nuisance

covariates, such as head motion parameters  $x$ ,  $y$ ,  $z$ , roll, pitch and yaw, were included. In study 14, motion-related artifacts were removed using ICA-AROMA; thus, only nuisance covariates, such as the top five principal components of white matter and cerebrospinal fluid signals and linear trend, were included. As trial estimates could be strongly affected by acquisition artifacts that occur during the trial, trial-by-trial variance inflation factors (VIFs) were calculated, and trials with VIFs >2.5 were excluded from further analyses. In study 3, we also excluded global outliers, that is, trials exceeding 3 s.d. above the mean, and used a principal component-based denoising step.

**Single-trial analysis: studies 2 and 5.** In the case of studies 2 and 5, fitting a set of three basis functions was used instead of the standard HRF. This procedure allowed for flexible variation of the shape of the HRF across trials and voxels. This is also consistent with the procedures used in the original publication<sup>60</sup>. The pain period basis set comprised three curves shifted in time, and it was customized for thermal pain responses based on previous studies<sup>60,63</sup>. To estimate responses evoked by cues in study 5, the pain-anticipation period was fitted with a canonical HRF. The period was then truncated at 8 s to ensure that modeled anticipatory responses were not affected by the activity evoked by the noxious stimuli. As in other studies, a regressor for nuisance covariates was included and trials with VIFs > 2.5 were excluded. In addition, in study 5, global outliers were excluded. To estimate trial-level pain period activity, the fitted basis functions from the flexible single-trial approach were reconstructed to compute the area under the curve (AUC). The trial-by-trial AUC values were then used as estimates of the trial-level pain period activity.

**Model building, importance calculation and parcellation.** Using the discovery dataset, we first built individualized SVR models on the whole-brain, single-trial pain data of each participant, resulting in 404 individualized predictive maps (Fig. 1a). For each model, we ran the bootstrap tests with 5,000 samples (resampling with replacement) to obtain the two-tailed uncorrected  $P$  values for each voxel for each individual based on the sampling distribution (that is, converting  $z$ -scores to  $P$  values calculated from the mean and s.d. of the sampling distribution). We then calculated the mean negative logarithmic  $P$  values (mean  $(-\log(P))$ ) and selected voxels corresponding to the top 10% of the mean  $(-\log(P))$  values for further analyses (Fig. 1b). We then parcellated the selected voxels into anatomical subregions using a combination of cerebral and cerebellar atlases<sup>64,65</sup>, which together provided a reliable brain parcellation with a sufficient number of brain structures for our purpose, and additionally divided the thalamus into medial and right lateral regions (Fig. 1c). Nevertheless, it is also important to note that multiple choices of parcellation exist. The impact of the selected parcellation could be explored in future studies. Some of the region masks, namely BG, dlPFC, MT, pMCC, PCun, visual, vlPFC and vmPFC, were smoothed with a 1-mm FWHM Gaussian kernel to make the region smooth and large enough for the following pattern-based analyses. After normalizing the individualized predictive maps by dividing each voxel weight by the s.d. of the weights across the whole-brain map, we applied the region masks to each map and used the masked data for further analyses.

**Univariate voxel-wise analysis.** To examine the individual variability of predictive weights at the voxel level, we calculated the variance of each voxel weight across all individualized maps. We also computed the mean weights across all predictive maps and summarized the values in each region as the mean regional weight and weight variance (Fig. 2). In addition, we inspected how variable the sign of weights was across subjects in individual regions (Extended Data Fig. 2). We first determined the median weight for each subject and each region. We then calculated the proportion of positive and negative median weights across the subjects for each region.

**Multivariate representational similarity analysis.** To compare the regional multivariate pattern representations across subjects, we performed a representational similarity analysis<sup>72</sup>. We first calculated the interindividual RDM for each brain region between the individualized multivariate patterns using 1 – correlation as a measure of interindividual distance. To take possible region-specific effects into consideration, such as the region sizes and spatial locations, we normalized the distance measures based on the null distance distribution generated with permutation tests with 1,000 iterations. More specifically, in each iteration, we permuted the trial labels (that is, pain ratings) for each subject's data and fitted a predictive model using the permuted data. We then computed the  $z$ -scores (normalized representational distance) with the observed interindividual distance ( $d_{\text{cor,obs}}$ ) and the permuted distance data ( $d_{\text{cor,null}}$ ) using the following equation:

$$Z_{\text{cor},i} = \frac{d_{\text{cor,obs}} - \bar{d}_{\text{cor,null}}}{\sqrt{\frac{\sum_{i=1}^N (d_{\text{cor,null}} - \bar{d}_{\text{cor,null}})^2}{N-1}}}$$

where  $i$  is the region index and  $N$  the number of iterations. This procedure is illustrated in Fig. 3a. We summarized the normalized RDMs by taking the mean of the lower triangles of the matrices.



To remove the effects of region size on the mean representational distance, we applied linear regression to residualize the mean representational distance. To calculate the s.e.m. for the mean residualized distance, we first used the parameters of the linear regression for residualizing all elements of the lower triangle of each RDM. We then calculated the s.e.m. as the s.d. of the matrix divided by the number of elements.

**Replication using study 14.** For further validation of our results, we employed study 14 as an independent large-scale replication dataset. As in the discovery dataset, we fitted SVR models to the data of each of the 124 participants in the study, creating 124 individualized pain-predictive maps. We applied region masks defined in the discovery dataset to each map and used only the masked region data for further analyses. We performed a representational similarity analysis in the replication dataset in the same manner as described above in the discovery dataset. Briefly, we calculated the interindividual RDMs based on the predictive patterns in all regions and compared the matrices with the null RDMs obtained from the permutation tests. This process resulted in normalized regional RDMs. We calculated the mean of the lower triangles of the matrices and regressed out the effects of the region size to obtain the residualized distance in the regions. To compare the results with those obtained in the discovery dataset, we assigned ranks to the regions in both the discovery and the replication datasets according to the residualized distance values. We then compared the two vectors of ranks using Spearman's rank correlation coefficient.

**Calculation of tSNR.** To examine the quality of the signal across the functional scans, we calculated the tSNR of the images obtained by fMRI in the replication dataset. For each run, we computed the run-level tSNR map as the voxel-wise mean divided by the s.d. of the signal. Subsequently, we obtained an individual tSNR map for each participant by averaging the run-level tSNR maps. Finally, the group-level tSNR map was obtained as the mean of all individual tSNR maps.

**MDS and region clustering.** To cluster pain-predictive brain regions based on the patterns of the representational distance across individuals, we first vectorized the normalized RDMs acquired in the multivariate analysis of the discovery dataset and used them as inputs for the representational connectivity analysis<sup>23</sup>. By calculating Kendall's  $\tau_A$  among the brain regions, we obtained a  $21 \times 21$  representational connectivity matrix, which was used as the basis for the region clustering. For clustering, we transformed the Kendall's  $\tau_A$  matrix into a distance matrix by calculating  $(1 - \text{Kendall's } \tau_A)/2$ , where values ranged from 0 to 1, with 0 indicating identity and 1 the maximum distance. Then, we applied the NMDS to Kendall's  $\tau_A$  distance matrix. To select the appropriate number of NMDS dimensions, we evaluated the stress metric of the NMDS solutions ranging from 1 dimension to 20 dimensions and selected the final number of dimensions, which was 10, based on the scree test (Extended Data Fig. 8a). Finally, we performed hierarchical clustering with average linkage on the selected NMDS results and used permutation tests to choose the number of clusters  $k$ . In particular, for each possible solution ranging from 2 clusters to 15 clusters, we first evaluated the mean clustering quality  $q$  in the observed data using the silhouette values computed as:

$$q = \frac{1}{i} \sum_i \frac{d_{i_{\text{nn}}} - d_{i_w}}{\max(d_{i_w}, d_{i_{\text{nn}}})}$$

where  $i$  denotes the region number,  $d_{i_{\text{nn}}}$  is the distance from a region to the nearest neighboring cluster and  $d_{i_w}$  is the Euclidean distance from a region to the center of its own cluster. Then, we permuted the NMDS scores, applied the same clustering algorithm and evaluated  $q$  of the clustering solutions based on the permuted data. We ran 1,000 iterations of the process to create a null-hypothesis distribution for the clustering quality  $q$ , which allowed us to assess the  $z$ -scores for  $q$  of each clustering solution of the observed data as follows:

$$Z_k = \frac{q_{\text{obs}} - \bar{q}_{\text{null}}}{\sqrt{\frac{\sum_i (q_{\text{null}} - \bar{q}_{\text{null}})^2}{N-1}}}$$

where  $k$  denotes the cluster number,  $q_{\text{obs}}$  is the clustering quality of the original data,  $\bar{q}_{\text{null}}$  is the clustering quality of the permuted data and  $N$  is the number of iterations (Extended Data Fig. 8d). See refs.<sup>66,67</sup> for other examples of use of the analysis.

**Principal gradient analysis.** To examine where our region clusters were on the spectrum of the principal gradient of the cortical organization that ranges from unimodal to transmodal areas, as suggested by a previous study<sup>24</sup>, we calculated the mean principal gradient in the clusters. We first masked a principal gradient map derived from an independent resting-state fMRI dataset ( $N=59$ ) with the region clusters. Then, we calculated the mean value in each cluster. For comparison with the mean residualized representational distance, we assigned ranks to the region clusters based on the principal gradient values and mean representational distance and calculated the rank correlation between them.

**Statistics and reproducibility.** The analyses were performed on 14 studies including 11 published studies. The total sample size was  $N=528$ , which was much larger than sample sizes common in experimental fMRI studies<sup>68</sup>. For this reason, we did not

conduct formal statistical tests to predetermine the sample size. In one study (study 8), four participants were excluded due to the unsuccessful extraction of the single-trial data. In all studies, the stimuli were fully randomized within subjects. Data collection and analysis were not performed blind to the conditions of the experiments.

In Fig. 1b, the top 10% of the mean ( $-\log(P)$ ) values across all 404 individualized maps are displayed. To obtain the  $P$  values, we first ran bootstrap tests on each of the 404 individualized maps (with 10,000 samples), and then calculated the two-tailed uncorrected  $P$  values from  $z$ -scores based on the distribution of weights resulting from the bootstrap tests. In Fig. 3b,d, we used Pearson's correlation coefficient  $r$ , with a subsequent Student's  $t$ -test to determine two-tailed  $P$  values ( $n=21$  regions). In both Figs. 3c and 4a, we show the mean residualized representational distances and their s.e.m. The s.e.m. was calculated across all pair comparisons of individuals, that is,  $C(404, 2) = 81,406$ . In Fig. 4b,c, we used Spearman's rank correlation coefficient  $\rho$  with a subsequent Student's  $t$ -test to determine two-tailed  $P$  values. In Extended Data Fig. 3, to select individualized models with significant prediction performance, we evaluated Pearson's correlation coefficient between the predicted and actual outcome values and conducted a Student's  $t$ -test with the correction for multiple comparisons using an FDR at  $q < 0.05$ . In Extended Data Figs. 3c ( $n=21$ ), 5c–e ( $n=21$ ) and 9c ( $n=10$ ), we used Spearman's rank correlation coefficient  $\rho$  with a subsequent Student's  $t$ -test to determine two-tailed  $P$  values. The mean and s.e.m. are shown in Extended Data Figs. 3a (based on all pair comparisons of  $n=248$ ), 5a,b (based on all pair comparisons of  $n=229$  and  $n=175$ , respectively), 6c,d (based on all pair comparisons of  $n=285$  in both) and 9d (based on all pair comparisons of  $n=124$ ). In Extended Data Fig. 8c, we show the mean and 95% confidence interval for clustering quality obtained from a permutation test of 1,000 samples. In Extended Data Fig. 8d, we calculated the  $z$ -scores for the clustering quality based on the permutation test and the observed clustering quality. Further details on the statistical analyses are reported in the relevant sections.

**Reporting Summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

## Data availability

The individualized pain-predictive maps from the discovery dataset, region masks and regional normalized RDMs are available at [https://github.com/cocoonlab/individual\\_var\\_pain](https://github.com/cocoonlab/individual_var_pain). The cerebral and cerebellar atlases used in the present study are available at [https://github.com/canlab/CanlabCore/tree/master/CanlabCore/canlab\\_canonical\\_brains/Combined\\_multiatlas\\_ROI\\_masks](https://github.com/canlab/CanlabCore/tree/master/CanlabCore/canlab_canonical_brains/Combined_multiatlas_ROI_masks). The data from the replication dataset are available upon request.

## Code availability

In-house Matlab codes for fMRI data analyses used in the present study are available at <https://github.com/canlab/CanlabCore>. An example code for the multivariate representational similarity analysis is available at [https://github.com/cocoonlab/individual\\_var\\_pain](https://github.com/cocoonlab/individual_var_pain).

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### Author contributions

L. Kohoutová and C.-W.W. conceptualized the study, analyzed the data, interpreted the results and wrote the manuscript. M.R. and T.D.W. contributed to study 1 data. A.K. and T.D.W. contributed to study 2 data. L.Y.A. and T.D.W. contributed to data from studies 3, 5, and 8. M.J. and T.D.W. contributed to data from studies 4 and 9. L.S. and T.D.W. contributed to study 6 data. J.T.B. and T.D.W. contributed to study 7 data. L. Koban and T.D.W. contributed to study 10 data. S.G. and C.B. contributed to study 11 data. S.M.S. and T.D.W. contributed to study 12 data. C.-W.W., L. Koban and T.D.W. contributed to study 13 data. D.H.L., S.L. and C.-W.W. contributed to study 14 data. All authors revised the manuscript.

### Competing interests

The authors declare no competing interests.

### Additional information

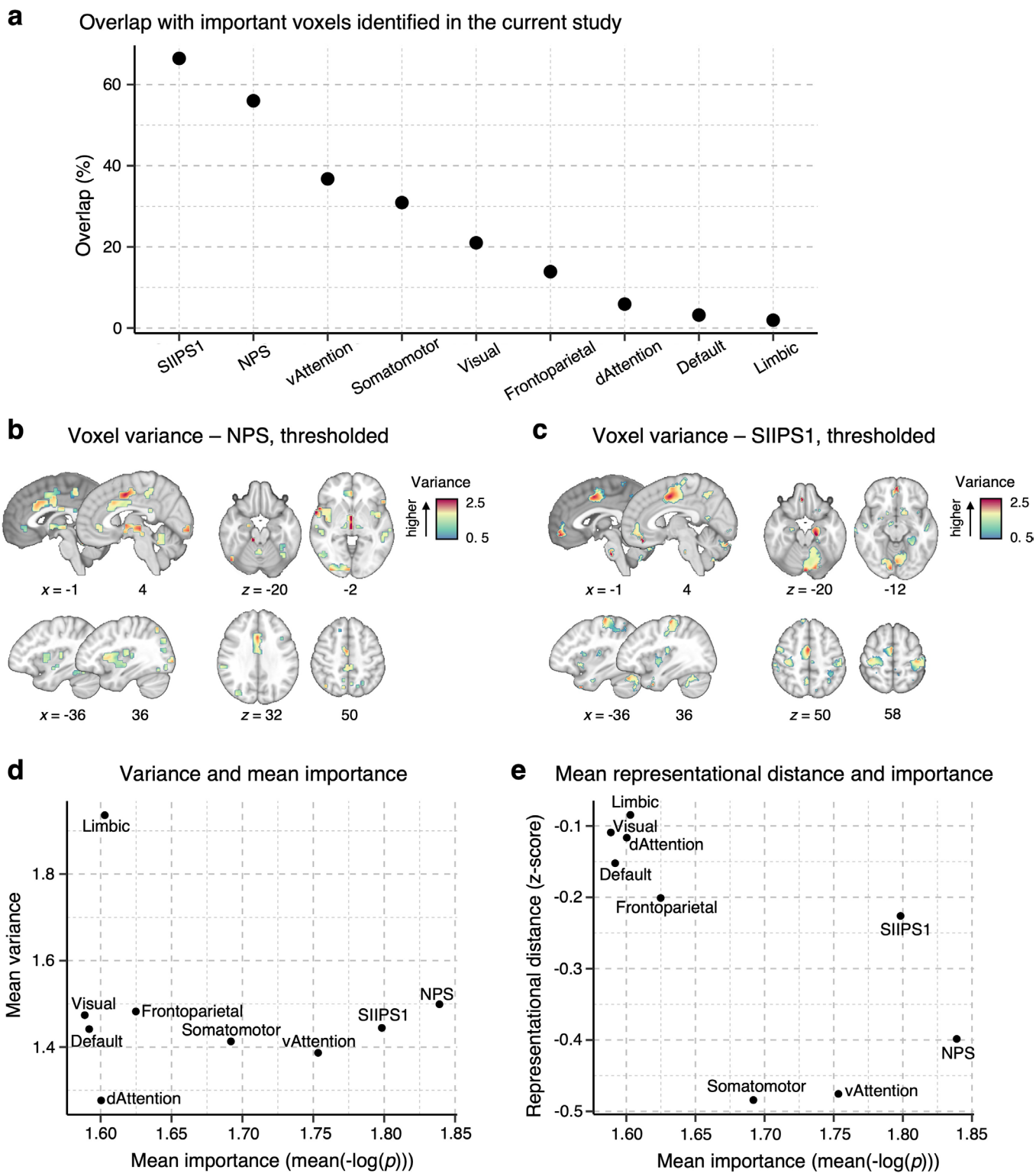
**Extended data** Extended data are available for this paper at <https://doi.org/10.1038/s41593-022-01081-x>.

**Supplementary information** The online version contains supplementary material available at <https://doi.org/10.1038/s41593-022-01081-x>.

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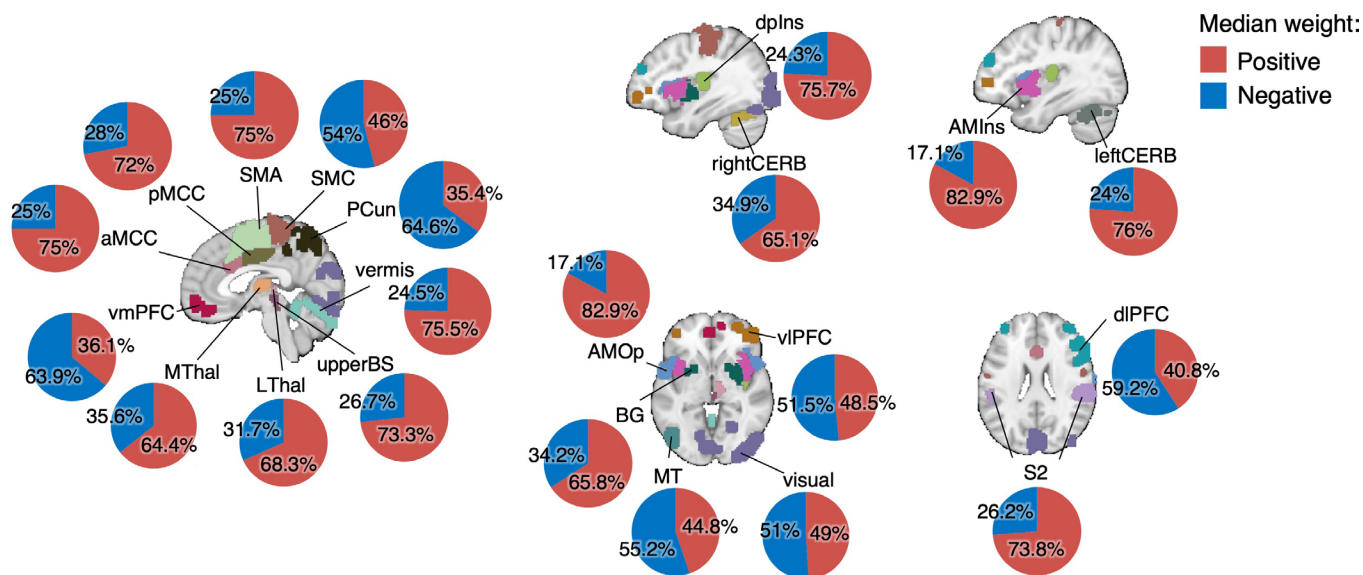


Extended Data Fig. 1 | See next page for caption.

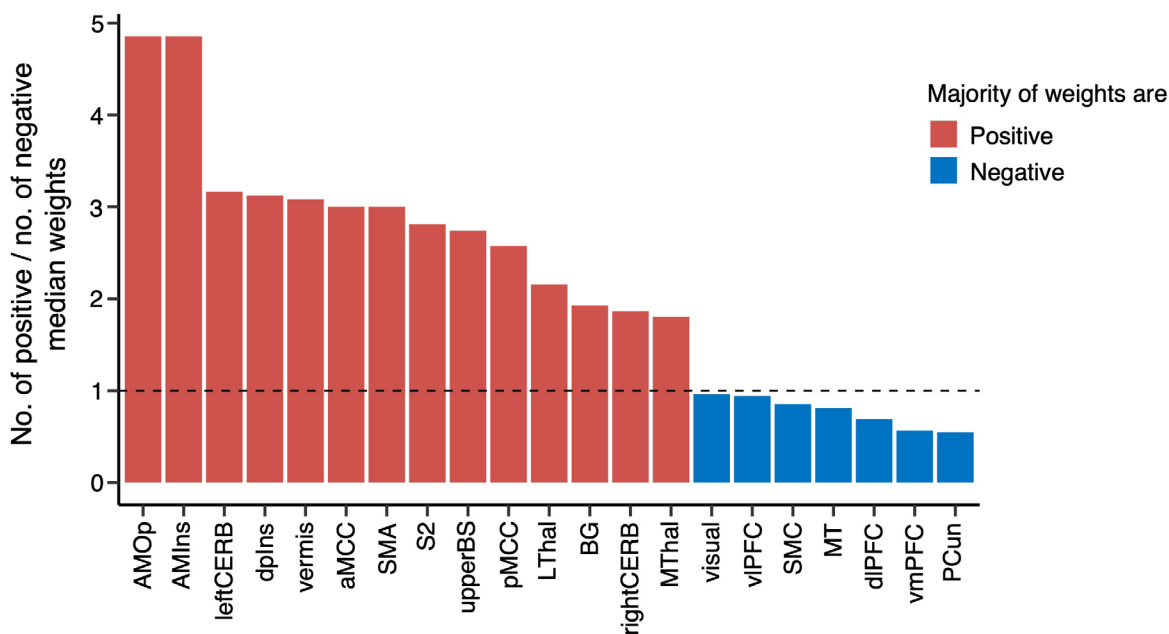
**Extended Data Fig. 1 | Reference results based on pain signatures and large-scale functional networks.** To provide a reference to other commonly used brain parcellations and existing pain signatures, we performed the analyses presented in the manuscript with the NPS<sup>1</sup>, SIIPS1<sup>2</sup> (thresholded at  $q < 0.05$ , false discovery rate [FDR] correction), and seven large-scale resting-state functional networks<sup>3</sup> as masks. **(a)** The plot shows the proportions of the overlapping voxels of the pain signature and network masks with the area of the important voxels identified in the current study. **(b)** and **(c)** show the voxel-wise variance across the individuals from the discovery dataset in the thresholded NPS and SIIPS1 masks, respectively. **(d)** In each signature and network mask, we calculated the mean importance with  $\text{mean}(-\log(p))$  (based on two-tailed  $p$ -values) and the mean voxel-wise variance. The results suggest that the limbic network showed the highest mean variance, while the dorsal attention network showed the lowest mean variance **(e)** We also performed the multivariate analysis. we calculated the inter-individual representational dissimilarity matrix (RDM) using the correlation-based distance for each masked area, performed the permutation tests with 1,000 samples, and calculated the normalized RDMs (z-scores), as we did in the main analysis (see Fig. 3a in the main manuscript). The results suggest that the limbic and visual networks showed the highest mean normalized representational distance (that is, highest inter-individual variability), while the somatomotor and ventral attention networks showed the lowest distance (that is, lower inter-individual variability). dAttention, dorsal attention network; vAttention, ventral attention network.



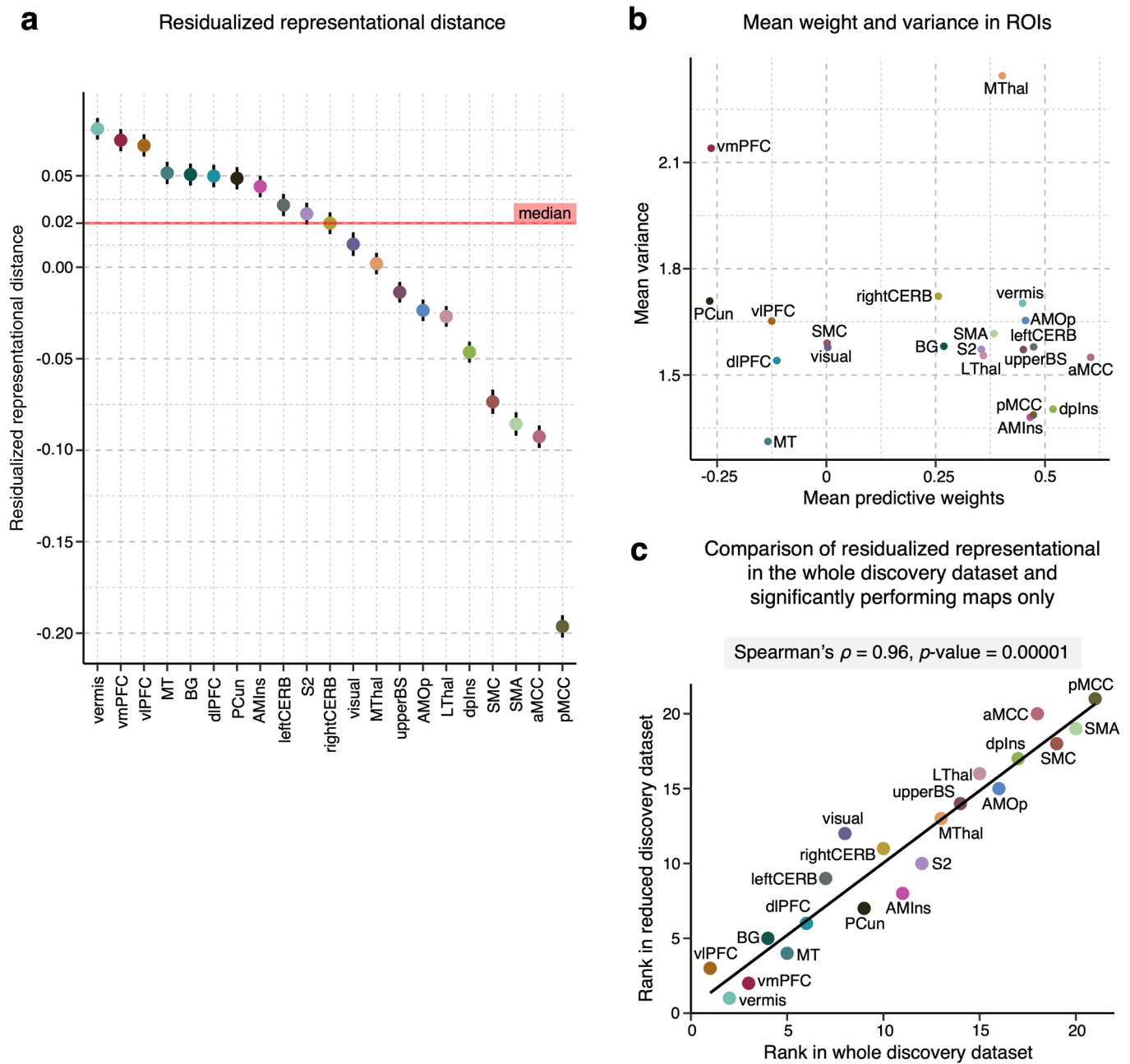
**a** Proportions of positive and negative median weights across individual predictive maps



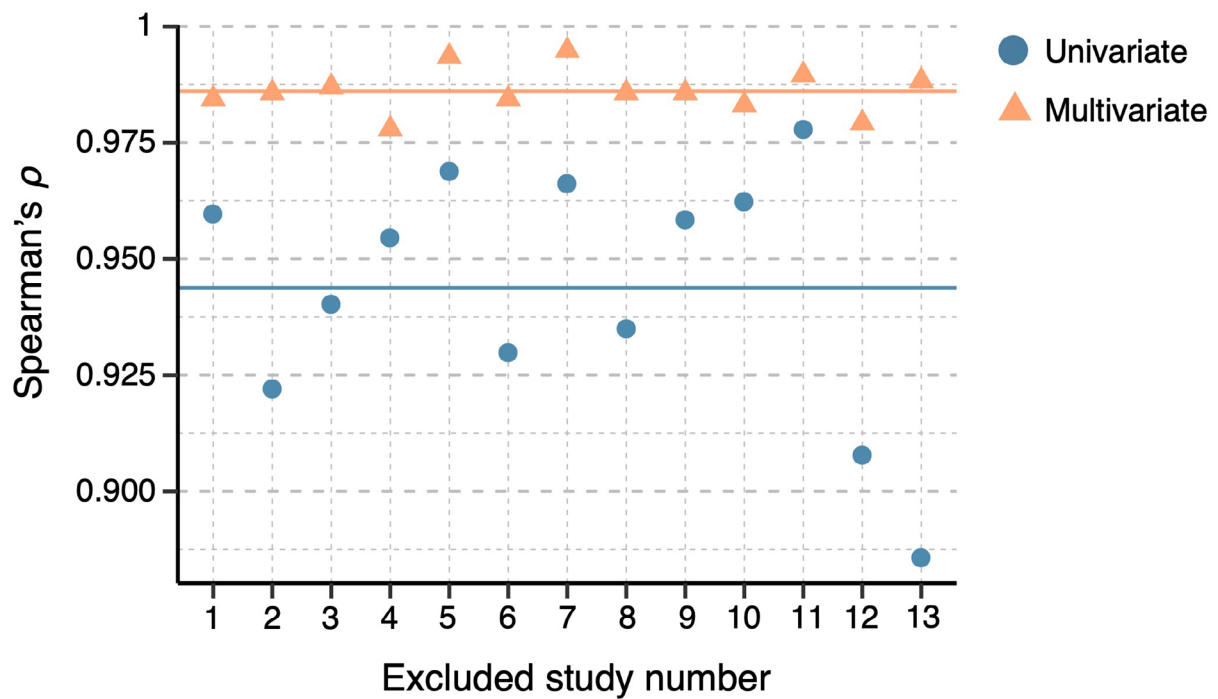
**b** Sorting the brain regions from positive to negative (ratio of median weight signs)



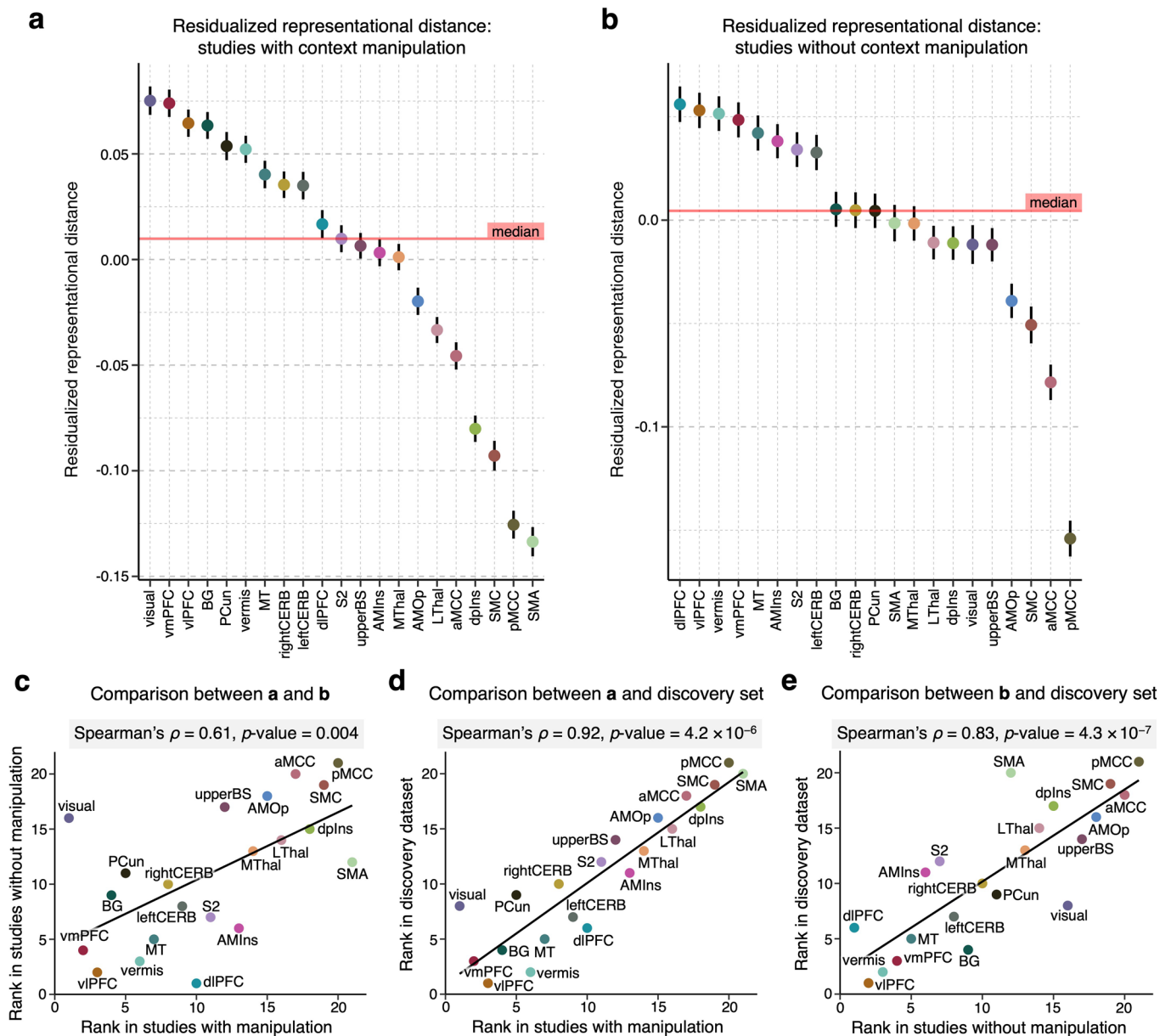
**Extended Data Fig. 2 | Proportions of the signs of median predictive weights.** (a) We found the median weight across voxels for each participant in each region and calculated the proportion of positive and negative median weights across all subjects. The pie charts display the percentage of median positive weights in red and negative weights in blue. (b) The bar plot shows the ratio of the number of positive to the number of negative median weights in each region. The red bars depict the regions with more positive median weights, and the blue bars mark the regions with more negative median weights.



**Extended Data Fig. 3 | Results after removing predictive maps with non-significant prediction performance.** To examine the effects of individuals with poor prediction performance on the inter-individual variability, we conducted the same analyses only with the individualized models with significant performance after correction for multiple comparisons using FDR correction at  $q < 0.05$ . All analyses shown here were performed on a reduced dataset of  $n = 248$  after removing  $n = 156$  with non-significant prediction performance. **(a)** The plot shows the mean representational distance after regressing out the effects of the region size. The error bar indicates the standard error of the mean. This corresponds to Fig. 3c of the main manuscript, which used the whole discovery dataset. **(b)** The scatter plot depicts the univariate analysis result, that is, the mean voxel weight and variance in each region. This corresponds to Fig. 2c of the main manuscript. **(c)** We assigned ranks to each region based on the residualized representational distance in both the original result based on the whole dataset and in the result based on the reduced dataset presented here. The two sets of results were significantly correlated at Spearman's  $\rho = 0.96$ ,  $p = 0.00001$ , two-tailed.

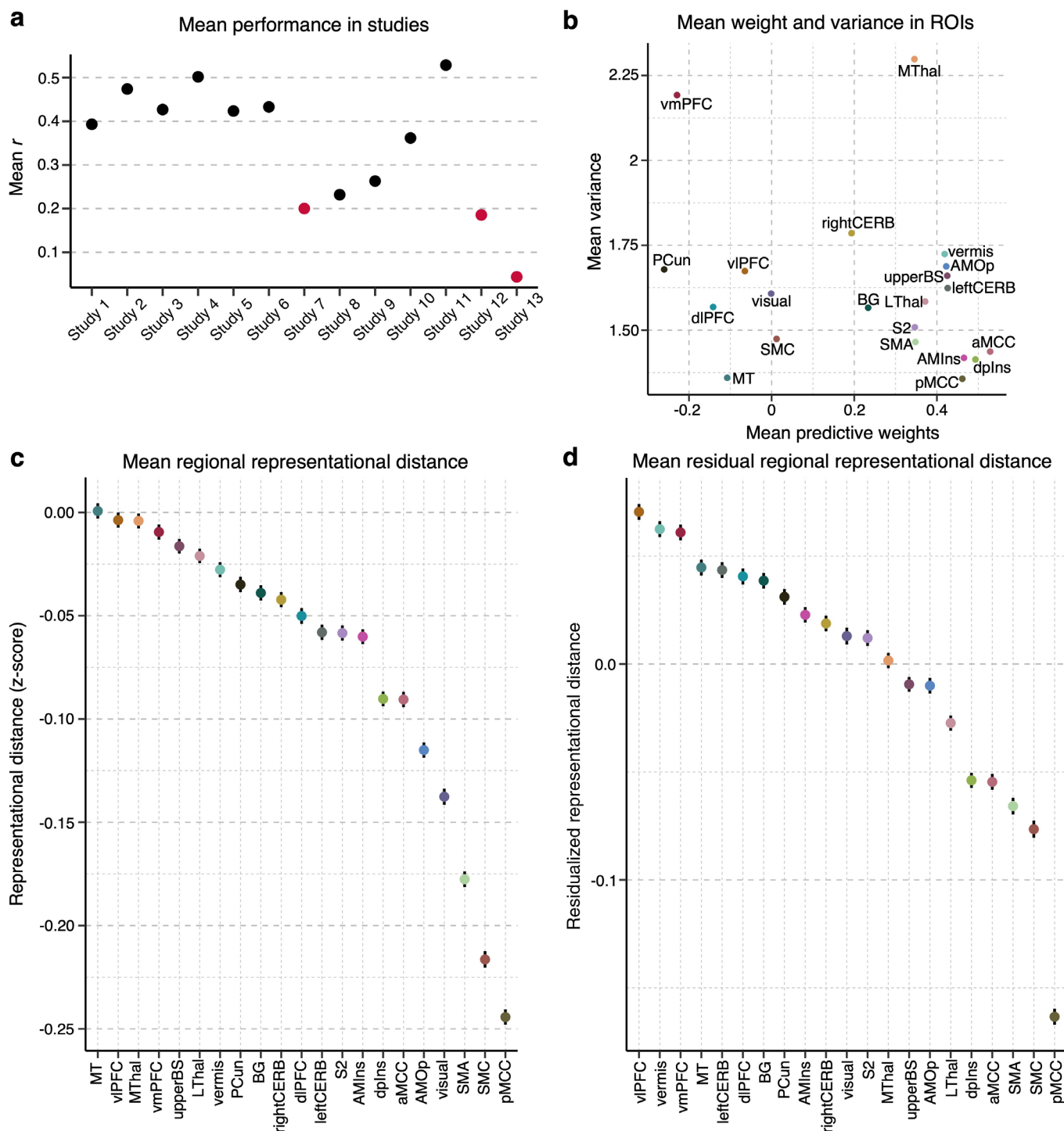


**Extended Data Fig. 4 | Inspection of potential study-specific effects on results.** To evaluate any potential study-specific effects on our results, we compared the final results (both univariate and multivariate) with the results with one study removed. For the comparisons, we calculated Spearman's  $\rho$  using the rank orders of the brain regions' individual variability between the results from the full versus reduced datasets. The blue dots show the results of the univariate analysis, ranging from 0.86 to 0.98, while the orange triangles are the results of the multivariate analysis, ranging from 0.98 to 0.99. The straight lines mark the mean Spearman's  $\rho$  for both cases with respective colors.

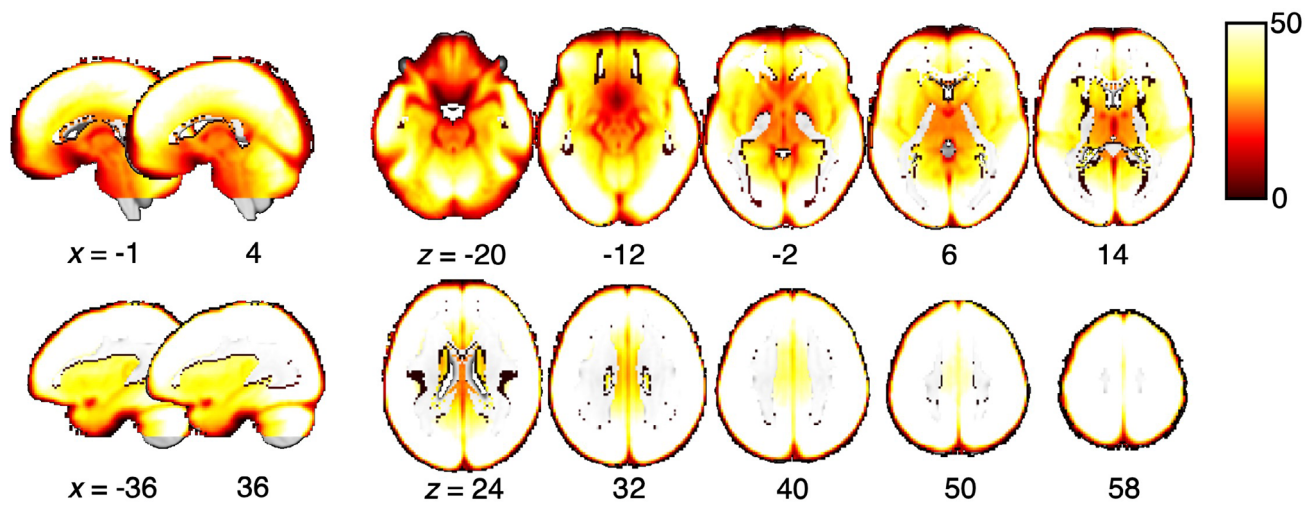


**Extended Data Fig. 5 | Results of the representational similarity analysis for studies with and without context manipulation.** We performed the representational similarity analysis and controlled for the region size in the discovery dataset divided into studies **(a)** with context manipulation (for example, placebo and cognitive regulation; studies 1, 4, 6, 7, 8, 11, and 12;  $n = 229$ ) and **(b)** without context manipulation (studies 2, 3, 5, 9, 10, and 13;  $n = 175$ ). The figures show the mean residualized representational distance and the standard error of the mean. **(c)** We found a significant correlation between region ranks in **(a)** and **(b)** of Spearman's  $\rho = 0.61$ ,  $p = 0.004$ , two-tailed. When compared with the region ranks in the discovery set, both **(d)** result in studies with context manipulation and **(e)** result in studies without context manipulation showed significant correlations of  $\rho = 0.92$ ,  $p = 4.2 \times 10^{-6}$ , and  $\rho = 0.83$ ,  $p = 4.3 \times 10^{-7}$ , respectively, all two-tailed.

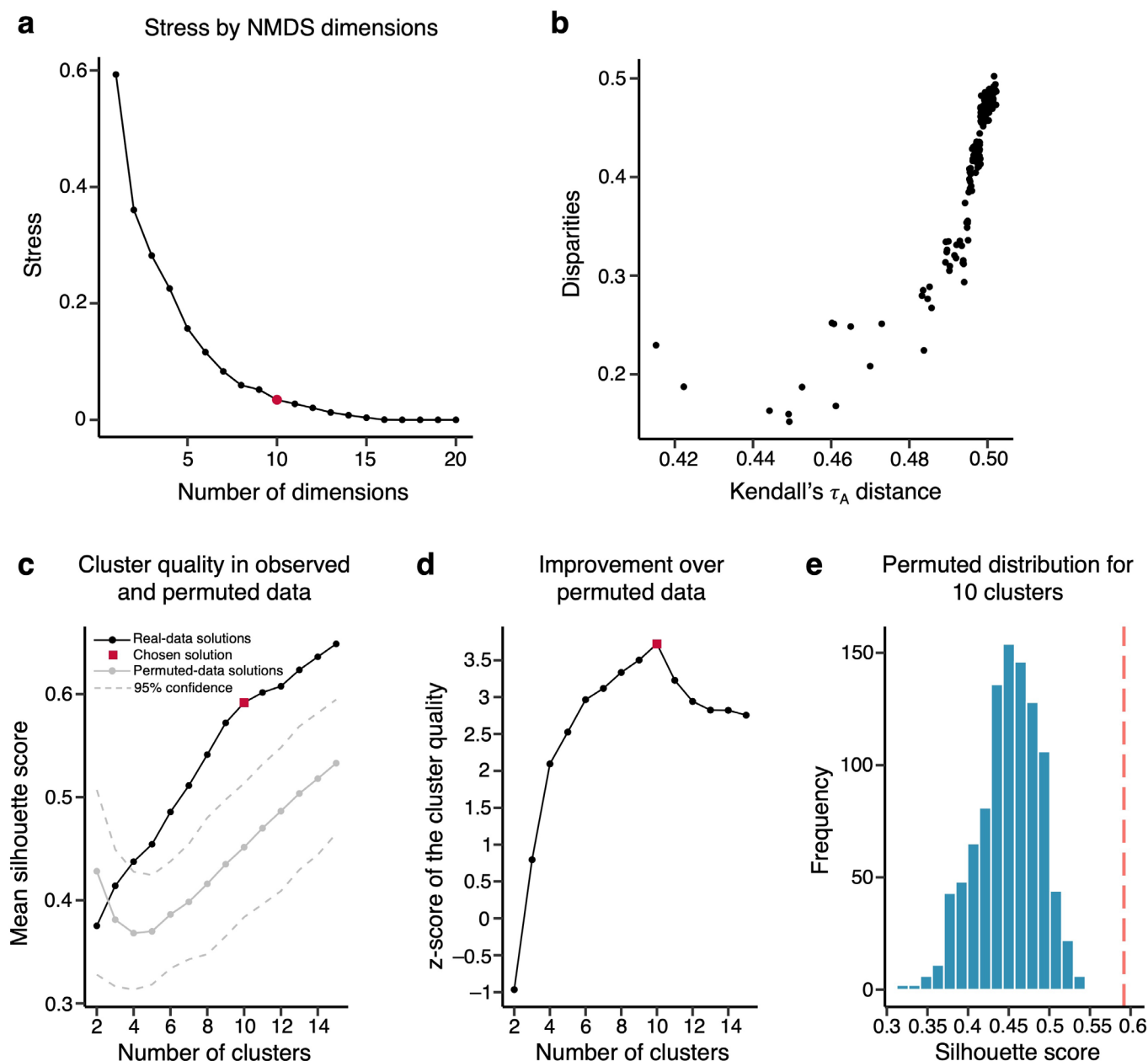




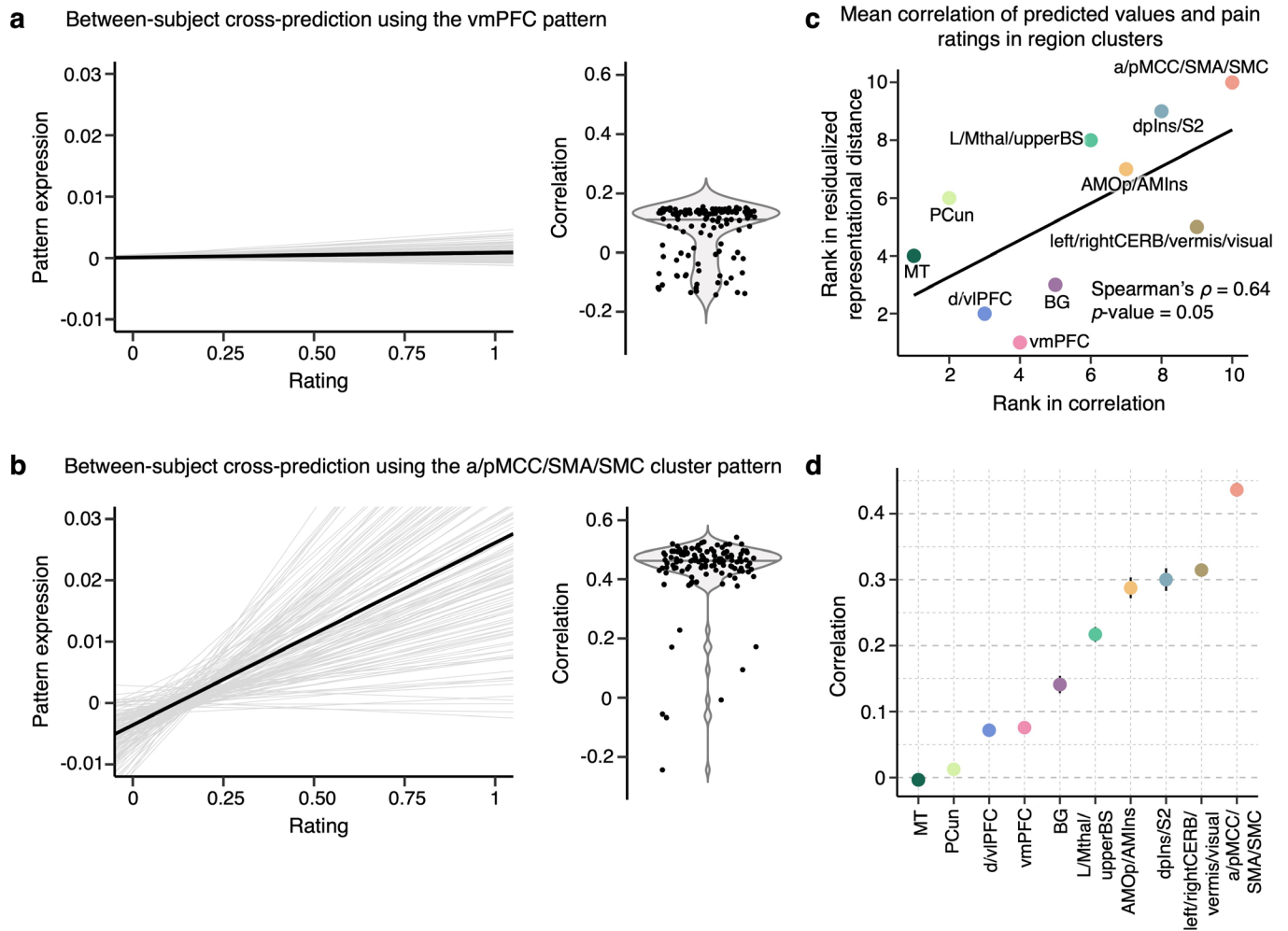
**Extended Data Fig. 6 | Results after excluding the studies that showed low prediction performance.** (a) The plot shows the average prediction performance of the individualized whole-brain SVR models across 13 studies. Studies 7, 12 and 13 (marked in red) had the lowest performance, mean  $r=0.20$ ,  $0.19$ , and  $0.04$ , respectively. To test whether these studies with low performance affected the results, we re-did the analysis without these studies, that is, on  $n=285$  individuals. (b) The scatter plot shows the mean predictive weight and variance across the individualized maps for each region after the exclusion of the three studies. (c) The plot displays the mean representational distance (z-scores) and standard error of the mean in each region based on all pair comparisons of individuals, that is,  $C(285, 2) = 40,470$ . (d) The residuals of the representational distance after removing the effects of the region size and the standard error of the mean based on all pair comparisons of individuals, that is,  $C(285, 2) = 40,470$ , are plotted.



**Extended Data Fig. 7 | tSNR map.** The group average of tSNR is visualized on a brain underlay with brighter colors depicting higher tSNR values, that is, better tSNR.

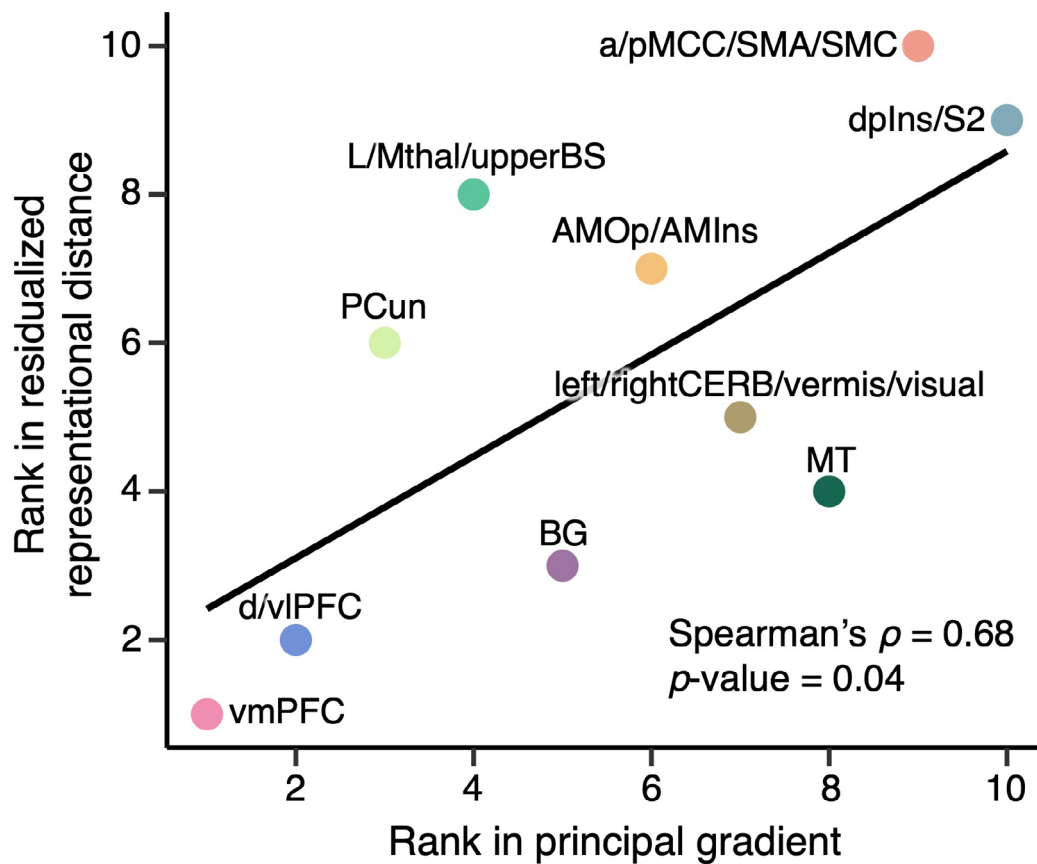


**Extended Data Fig. 8 | Nonmetric multidimensional scaling-based hierarchical clustering analysis.** (a) For the clustering of pain-predictive regions, we first ran the nonmetric multidimensional scaling (NMDS) on the Kendall's  $\tau_A$  distance matrix, which was calculated as  $(1 - \text{Kendall's } \tau_A)/2$ . Based on the stress metric and scree method, we selected 10 dimensions (marked in red). (b) The x-axis of the scatter plot shows the input Kendall's  $\tau_A$  distance between regions, and the y-axis shows the Euclidean distance between the regions scaled into 10 dimensions after the NMDS. (c) We performed the hierarchical clustering with average linkage on the selected NMDS results and used permutation tests to choose the final number of clusters,  $k$ . For the permutation tests, we shuffled the NMDS scores, applied the clustering algorithm, and assessed the clustering quality of the permuted data at each iteration. We ran a total of 1,000 iterations, and the plot shows the mean cluster quality of both the observed (solid black line) and permuted (solid gray line) data, as well as the 95% confidence interval (gray dashed lines) for the permuted cluster quality. The red square marks the selected solution with a Silhouette score of 0.59. (d) The plot shows the z-scores that indicate an improvement of the cluster quality of the observed data compared to the permuted null data. The highest improvement was achieved with the 10 cluster solution (shown as the red square) with a z-score of 3.72,  $p=0.0002$ , two-tailed. (e) The histogram depicts the observed cluster quality of the 10 cluster solution (red dashed line) versus the null distribution from the permutation test (blue histogram).



**Extended Data Fig. 9 | Cross-individual pain prediction.** To further illustrate the inter-individual variability in pain representations across different region clusters, we conducted cross-individual prediction of pain using pain predictive patterns of region clusters in Study 14. The panels **(a)** and **(b)** show examples of the cross-prediction using the vmPFC (the most variable region cluster) and a/pMCC/SMA/SMC (the most stable region cluster) cluster patterns, respectively. The gray lines in the line plots show the mean regression lines of pain prediction in others using an individual's predictive map (that is, each line indicates the prediction using one participant's pain prediction model). The black lines show the global average of all the individual regression lines. The violin plots show the mean correlation between the predicted and actual pain ratings in cross-individual pain prediction. Each dot represents mean prediction-outcome correlation using one participant's pain prediction model. **(c)** We calculated the global cross-individual prediction performance of each region cluster using prediction-outcome correlations. The top panel shows the relationship between the rank in the mean residualized distance (y-axis), where clusters are ordered from the most variable to the least variable cluster, and the rank in the correlation values (x-axis), where the clusters are ordered from the lowest to the highest cross-individual prediction performance. Together with the examples in **(a)** and **(b)**, this plot suggests that the cross-individual prediction is more reliable in the clusters with lower inter-individual variability. **(d)** The plot displays the mean correlation values with the standard error of the mean for each region cluster based on  $n=124$ .





**Extended Data Fig. 10 | Relationship between the principal gradient of functional connectivity and mean residualized representational distance.**

To compare the principal gradient spectrum and mean residualized distance in clusters, we first calculated the principal gradient map using our own resting-state fMRI dataset ( $n=56$ ; 7-min resting scan) to create a volumetric principal gradient image and to include the subcortical regions. We assigned ranks to the region clusters based on both the principal gradient value (x-axis) and mean residualized distance (y-axis) and compared them using Spearman's rank correlation coefficient. We found a significant relationship at Spearman's  $\rho=0.68$ ,  $p=0.04$ , two-tailed.

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|-------------------------------------|-------------------------------------|--|
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | The exact sample size ( $n$ ) for each experimental group/condition, given as a discrete number and unit of measurement  |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly  |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | The statistical test(s) used AND whether they are one- or two-sided<br><i>Only common tests should be described solely by name; describe more complex techniques in the Methods section.</i>   |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | A description of all covariates tested   |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons  |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals) |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | For null hypothesis testing, the test statistic (e.g. $F$ , $t$ , $r$ ) with confidence intervals, effect sizes, degrees of freedom and $P$ value noted<br><i>Give <math>P</math> values as exact values whenever suitable.</i>                            |
| <input checked="" type="checkbox"/> | <input type="checkbox"/>            | For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings   |
| <input checked="" type="checkbox"/> | <input type="checkbox"/>            | For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes   |
| <input type="checkbox"/>            | <input checked="" type="checkbox"/> | Estimates of effect sizes (e.g. Cohen's $d$ , Pearson's $r$ ), indicating how they were calculated   |

*Our web collection on [statistics for biologists](#) contains articles on many of the points above.*

### Software and code

Policy information about [availability of computer code](#)

Data collection In data collection, softwares E-prime and Matlab were used.

Data analysis Preprocessing of functional images was performed using Statistical Parametric Mapping (SPM; versions 5, 8 and 12) software (<http://www.fil.ion.ucl.ac.uk/spm/>), FMRIB Software Library (FSL) (<https://fsl.fmrib.ox.ac.uk>) and Independent Component Analysis-based strategy for Automatic Removal Of Motion Artifacts (ICA-AROMA) software (<https://github.com/maartenmennes/ICA-AROMA>). Analyses of the preprocessed data in this study were performed using in-house Matlab (Matlab 2019b) codes for fMRI data analyses available at <https://github.com/canlab/CanlabCore>. An example code for the multivariate representational similarity analysis is available at [https://github.com/cocoanlab/individual\\_var\\_pain](https://github.com/cocoanlab/individual_var_pain).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research [guidelines for submitting code & software](#) for further information.

### Data

Policy information about [availability of data](#)

All manuscripts must include a [data availability statement](#). This statement should provide the following information, where applicable:

- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

The individualized pain-predictive maps from the discovery dataset, region masks and regional normalized representational dissimilarity matrices are available at [https://github.com/cocoanlab/individual\\_var\\_pain](https://github.com/cocoanlab/individual_var_pain). The cerebral and cerebellar atlases used in this study are available at <https://github.com/canlab/CanlabCore/>

## Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

Life sciences       Behavioural & social sciences       Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](https://www.nature.com/documents/nr-reporting-summary-flat.pdf)

## Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample size	N = 528, of which n = 404 was included in the discovery dataset (13 previous studies), and n = 124 in the replication dataset (1 study). 11 of the studies used in the current study have been published in peer-reviewed journals. Aggregating all data from these and some additional studies provided a significantly large sample size than in previous studies and, thus, was deemed a sufficiently large for our analyses.
Data exclusions	We used datasets from 14 previous studies, out of which in one study (Study 8) 4 participants were excluded due to the unsuccessful extraction of the single-trial data.
Replication	We performed one replication of the main findings of our study using an independent large-scale dataset of n = 124 that was collected in a different location with different equipment than the other 13 studies (discovery dataset).
Randomization	In all studies, the stimuli were fully randomized within subjects.
Blinding	N/A. In each we study we used here, all participants underwent the same study-specific experimental paradigm, i.e., there was no group assignment.

## Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

### Materials & experimental systems

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> Antibodies
<input checked="" type="checkbox"/>	<input type="checkbox"/> Eukaryotic cell lines
<input checked="" type="checkbox"/>	<input type="checkbox"/> Palaeontology and archaeology
<input checked="" type="checkbox"/>	<input type="checkbox"/> Animals and other organisms
<input type="checkbox"/>	<input checked="" type="checkbox"/> Human research participants
<input checked="" type="checkbox"/>	<input type="checkbox"/> Clinical data
<input checked="" type="checkbox"/>	<input type="checkbox"/> Dual use research of concern

### Methods

n/a	Involved in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> ChIP-seq
<input checked="" type="checkbox"/>	<input type="checkbox"/> Flow cytometry
<input type="checkbox"/>	<input checked="" type="checkbox"/> MRI-based neuroimaging

## Human research participants

Policy information about [studies involving human research participants](#)

Population characteristics	528 healthy, right-handed participants (245 females - sex of one participant unknown; mean age in studies ranged from 20.8 to 28, with one study in which the age of participants is unknown)
Recruitment	Participants were recruited via advertisements posted on Facebook and Craigslist, and flyers from New York City and Boulder/Denver Metro Areas in the United States, Hamburg in Germany, and Suwon in South Korea. The recruitment may have been potentially biased towards younger, educated population as the majority of the participants were university students. In addition, there could be a sampling bias related to the willingness to participate in pain research.
Ethics oversight	All studies used in the current manuscript were approved by the institutional review board of Columbia University, the University of Colorado Boulder and Sungkyunkwan University, and the Ethics Committee of the Medical Chamber Hamburg. All participants provided written informed consent and were financially compensated for their participation.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

# Magnetic resonance imaging

## Experimental design

Design type	Block design
Design specifications	In all 14 studies, participants experienced a series of contact heat stimuli rated their pain experience after each stimulus. The number of trials varied across studies from 16 to 97 trials. The duration of the stimulus varied from 1.8 s to 20.16 s. Each study also included a psychological manipulation which were not of interest in this study.
Behavioral performance measures	After each stimulus, the participants rated their pain experience on a visual analogue scale or labeled magnitude scale. The scale used was different in each study, with majority of studies using the visual analogue scale with the range of 0 (no pain) to 100 (worst imaginable pain).

## Acquisition

Imaging type(s)	Both structural and functional images were acquired.
Field strength	10 studies used 3 T, while 4 studies used 1.5 T
Sequence & imaging parameters	Gradient-echo, 13 studies used EPI sequence, while one study used spiral in-out sequence; repetition time (TR) varied from 460 ms to 2,000 ms; echo time (TE) varied from 2.43 ms to 40 ms; field of view was 220 mm in 8 studies, 224 mm in 5 studies, and unknown in one study; flip angle varied from 44 to 84, unknown in 4 studies; number of slices varied from 24 to 56; voxel size varied from 2.7 x 2.7 x 2.7 mm <sup>3</sup> to 3.5 x 3.5 x 4 mm <sup>3</sup>
Area of acquisition	A whole brain scan was acquired.
Diffusion MRI	<input type="checkbox"/> Used <input checked="" type="checkbox"/> Not used

## Preprocessing

Preprocessing software	Nine studies used SPM8 software, three studies used SPM5 software, one study used both SPM5 and SPM8, and one study used a custom pipeline consisting of SPM12, FSL and ICA-AROMA.
Normalization	Structural images were normalized to the Montreal Neurological Institute (MNI) space using SPM. In 2 studies, additional normalization to the group mean was performed after the SPM normalization using a genetic algorithm-based normalization. Functional images were warped into the SPM's normative atlas.
Normalization template	MNI
Noise and artifact removal	To identify the outliers in the signal, we computed the Mahalanobis distances for the matrix of the concatenated slice-wise mean and standard deviation of the intensity values by functional volumes (over time). Values with a significant chi-square value (after correction for multiple comparisons with either false discovery rate or Bonferroni correction) were considered outliers. In practice, less than 1 % of the volumes were outliers. The time points identified as outliers were later used as nuisance covariates in the first-level models.
Volume censoring	N/A

## Statistical modeling & inference

Model type and settings	General Linear Model
Effect(s) tested	Multivariate pattern-based prediction and representational similarity analysis
Specify type of analysis:	<input type="checkbox"/> Whole brain <input type="checkbox"/> ROI-based <input checked="" type="checkbox"/> Both
Anatomical location(s)	We defined our regions of interest based on cortical and cerebellar atlases (Shattuck et al., 2008 and Diedrichsen et al., 2009).
Statistic type for inference (See <a href="#">Eklund et al. 2016</a> )	Voxel-wise inference
Correction	False discovery rate correction was used.

## Models & analysis

n/a	Involvement in the study
<input checked="" type="checkbox"/>	<input type="checkbox"/> Functional and/or effective connectivity
<input checked="" type="checkbox"/>	<input type="checkbox"/> Graph analysis
<input type="checkbox"/>	<input checked="" type="checkbox"/> Multivariate modeling or predictive analysis



