Fast temporal dynamics and causal relevance of face processing in the human temporal cortex

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We measured the fast temporal dynamics of face processing simultaneously across the human temporal cortex (TC) using intracranial recordings in eight participants. We found sites with selective responses to faces clustered in the ventral TC, which responded increasingly strongly to marine animal, bird, mammal, and human faces. Both face-selective and face-active but non-selective sites showed a posterior to anterior gradient in response time and selectivity. A sparse model focusing on information from the human face-selective sites performed as well as, or better than, anatomically distributed models when discriminating faces from non-faces stimuli. Additionally, we identified the posterior fusiform site (pFUS) as causally the most relevant node for inducing distortion of conscious face processing by direct electrical stimulation. These findings support anatomically discrete but temporally distributed response profiles in the human brain and provide a new common ground for unifying the seemingly contradictory modular and distributed modes of face processing.
Studies using lesion methods, functional imaging tools, or scalp encephalography (EEG) and magnetoencephalography have offered invaluable causal, spatial, and temporal information about the neural mechanisms of face processing in the human brain. Work in non-human primates has also provided important and novel insights. However, despite great progress, the long-lasting controversy between modular versus distributed models of face processing has persisted in the literature. Some studies have revealed face-selective responses in anatomically discrete regions of the temporal cortex (TC), and other observations have shown that the pattern of responses to face stimuli can be discerned from sampled data from non-selective regions of the TC, suggesting that face information is anatomically distributed. Both theories have unfortunately relied on the information with limited temporal resolution averaged over multiple seconds or from methods using regions of interest and averaging across subjects, or direct recordings from a single or a pair of recording sites. Thus, the fast temporal dynamics of face processing across a large extent of the cerebral cortex within individual brains remains poorly explored.

Intracranial recordings in neurological subjects with a large number of electrodes spread over a relatively large extent of the cortical surface—a method known as electrocorticography (ECoG)—offers a new opportunity for acquiring fast temporal information from precisely localizable sources of signal. This method offers millisecond temporal resolution and millimeter anatomical precision in the subject’s own native brain space. Unlike the uniform spatial coverage of imaging methods, intracranial EEG (iEEG) relies on sampling from a limited number of implanted areas and leaves behind regions outside the coverage zones. While this leads to limited anatomical sampling, it may provide sufficient coverage for recording simultaneously from many sites within each individual brain, in order to explore the spatiotemporal dynamics of activity across different cortical areas. In addition, the intracranial approach allows delivering electrical pulses to discrete neuronal populations while causal changes in the subjective experience of the participant can be probed.

Using simultaneous recording across a relatively large area of the human brain one could test the hypothesis that face information is first processed within the most face-selective sites that are anatomically discrete and localizable within individual brains, which then is distributed to less selective sites.

While recent intracranial recordings and stimulation studies, including our own, have addressed the neurophysiological underpinnings of face processing in the human brain, to our knowledge, these studies have yet to address the notion of anatomical selectivity and temporal distribution of face information. The current study was designed to rely on a multiprong approach using univariate and multivariate methods of recording, as well as causal probing with electrical stimulation to test our proposed hypothesis.

We would like to emphasize that the aim of the study was not to decipher the complex computational code of face perception in the human brain. Specifically, the study was not designed to address the nature of face processing in face-selective or non-selective sites. Studies in primate brains are perhaps better suited for that purpose. The goal of our work was twofold: (1) to explore the timing of response across different regions of the TC and test if there is distribution of face information in time, and (2) whether the stimulation of face-responsive sites cause the same effect in conscious viewing of faces. Here and in the remainder of the text, we will use the term “site” as the cortical region underneath a given electrode (~2 mm diameter). We use the term “face-selective site” to imply that neuronal responses in that site are significantly higher to face stimuli than non-face stimuli. We use the term “task-active site” to refer to sites in which visual objects induce significant activations compared to baseline, but responses to faces are not significantly different than responses to non-face stimuli.

Simultaneous recordings over a relatively large mantle of the human TC allowed us to address our two overarching questions. Here, we report that about half of recording sites show non-selective responses to at least one category of stimuli and only 10% of the sites have selective activations to face stimuli. Responses to face stimuli in the “task-active sites” are weak and correlate to the face responses present in face-selective sites. Both face-selective and task-active sites display a posterior to anterior gradient of selectivity as well as timing, suggesting that face information is distributed anatomically and in time. Finally, we report that only the stimulation of the posterior fusiform site (pFUS) region of the fusiform face areas (FFAs) appears to affect face perception.

Results

Data and design. We recruited eight neurosurgical patients implanted with ECoG electrodes as part of their presurgical invasive evaluation for medication-resistant focal epilepsy. Subjects had unilateral electrode implantation in the right (five subjects) or left (three subjects) hemisphere (Supplementary Table 1). Electrodes across all subjects (n = 357) provided suitable coverage over the ventral and lateral TC.

We recorded from each of the 357 implanted electrodes with high temporal resolution (>1000 samples per second) while the subjects performed a visual task, in which they viewed images of faces (human, mammal, bird, and marine), and non-faces, including bodies without faces (same four categories), limbs (human), objects, and places. Randomly, a red hashtag sign appeared on the screen (without any other superimposing images) at which time the subjects were instructed to press a key.

Encoding of face information in the TC. We quantified face information (i.e., the ability to discriminate faces from non-faces stimuli) in narrow bands of frequencies (Θ: 4–7 Hz, α: 8–12 Hz, β1: 13–29 Hz, β2: 30–39 Hz, and γ: 40–69 Hz) as well as High-Frequency Broadband (HFB, 70–177 Hz) signals using univariate and a multiple kernel learning (MKL) method configured for iEEG signals. Compared to the power of any other frequency band, the signal in the HFB showed the strongest findings, we focused further analyses on the HFB power and its profile of response across anatomical sites and task conditions.

We are mindful that the richness of the intracranial electrophysiological signal could have been explored by analyzing the power or phase of slower frequencies, or their coupling with higher frequencies. However, the HFB signal is well suited for the purpose of testing our predictions not only because of our MKL findings, but also because of the large body of evidence from other human and non-human studies (as summarized in ref. 18) that have confirmed HFB power as a reliable correlate of hemodynamic signal and averaged single and multiunit activity of a population of neurons in a given cortical site. More importantly, HFB has a more precise anatomical source compared to lower frequencies. The HFB signal hence provides a suitable marker for the engagement of a given cortical site in a given function. Similarly, the time of onset and the power of HFB provide valuable information about the time and level of engagement of a population of neurons adjacent to the recording electrode.
Using HFB activity, we found a heterogeneous profile of responses across recordings sites (Supplementary Table 3). Of the recorded sites ($n = 357$), 53.22% ($n = 190$) had significant responses to at least one category of stimuli relative to baseline (i.e., “active” sites). A total of 13.45% recording sites ($n = 48$) showed selective activations to human faces compared to any other stimuli (“human face-selective” sites, Fig. 1a). Only 10.64% of the recording sites ($n = 38$) showed face-selective responses (comparing all subcategories of faces to all non-faces; i.e., “face-selective” sites). The (human) face-selective sites were clustered in the fusiform gyrus or lateral occipital gyrus. While the group-level visualizations are presented in Montreal Neurological Institute (MNI) space, for our anatomical claims, we rely on the precise location of the electrodes in each subject’s own brain space (Supplementary Figs. 1 and 2). Interestingly, while there was clear overlap between the “face-selective” and “human face-selective” sites, a few sites (17 out of 357 sites) showed selective responses to human faces while lacking significant responses to other face stimuli. In further analyses, we refer to the sites that were assessed as “active” and that were neither face nor human face selective as “task-active” sites.

To explore whether biological similarity of the face stimuli (humans and mammals versus birds or marine) influences the neural responses in the face-selective sites, we compared responses to faces of different categories. This analysis showed that human faces induced the strongest activations on face-selective sites (median $= 5.21$ dB, $n = 1579$) followed by mammal faces (median $= 4.06$ dB, $n = 1609$), bird faces (median $= 3.48$ dB, $n = 1666$), and marine faces (median $= 3.48$ dB, $n = 1706$, Fig. 1c left). All pairs of face subcategories showed a significant difference in HFB power (permutation test, $p < 0.05$ after Bonferroni correction), except for the comparison of bird and marine faces (human-mammal: $p < 0.0001$, human-bird: $p < 0.0001$, human-marine: $p < 0.0001$, mammal-bird: $p = 0.0012$, mammal-marine: $p = 0 < 0.0001$, bird-marine: $p = 0.9896$, permutation tests). In comparison, the amplitude of responses to subcategories of faces in the task-active sites showed lower HFB power for the face categories (human faces = $0.73$ dB, $n = 5761$, mammal faces = $0.91$ dB, $n = 5497$, bird faces = $0.67$ dB, $n = 5837$, marine faces = $0.79$ dB, $n = 5808$, permutation tests, Fig. 1c right). Only mammal faces elicited significantly higher responses than did human and bird faces (mammal–human: $p < 0.001$, mammal–bird: $p < 0.0001$). Other pairwise comparisons did not show differences in terms of HFB amplitude across the four subcategories of faces (permutation test, $p > 0.05$ after false discovery rate (FDR) correction). Please note that these univariate results were not driven by physical differences in the stimuli (Supplementary Table 4, Supplementary Discussion 4).

Decoding of face information in the TC. Given that the HFB responses to human faces had the highest signal-to-noise ratio (SNR), further analyses focused on the processing of human face stimuli. To address the earlier imaging reports of distributed face processing, we investigated the decoding of face information across human face-selective and task-active (i.e., non-selective)
sites. In particular, we investigated whether task-active sites included face information that was not correlated with the face information present on selective sites. Such information could be too weak to be detected by our (stringent) univariate test or have another pattern than the one we assume in our test (e.g., a sinusoidal variation of the HFB signal for faces could lead to a mean of 0 dB over the considered time window). Instead of relying solely on decoding performance, we referred to a recently developed framework\(^ {15}\) to infer causal relationships between the stimuli and the observed EEG activity in decoding (here machine learning-based modeling) settings. This approach allows to isolate the causal contribution of task-active sites following the presentation of face stimuli in face processing. Please note that in this analysis, the term “causal” relates to the relationship between the presentation of a stimulus (cause) and subsequent changes in brain activity (effect), and as such may not imply causality in terms of lesion studies or electrical stimulation experiments. The considered framework assesses the “relevance” of both face and task-active sites in discriminating between human face and non-face epochs instead of referring to decoding performance. A feature is assessed as “relevant” in decoding settings if, when removed, the performance of the model is significantly affected. This is similar to the procedure used in a recent publication\(^ {46}\), in which the face information shared across regions of interest was taken into account. Features assessed as (respectively not) relevant in both encoding settings (corresponding to our univariate tests) and decoding settings are (respectively not directly) causally related to the stimuli.

To address face processing in face-selective versus task-active sites, we ran three machine learning models discriminating between human face and non-face (i.e., human, mammal, bird, and marine body parts, places, objects, and human limbs pooled) stimuli, using signals from different sets of sites. The results from all models are displayed in Supplementary Table 5 and Fig. 2a. For each subject, model I incorporated data from all TC sites. For all subjects, model I was able to significantly discriminate between human faces and non-faces (permutation test \(p < 0.05\), FDR corrected). Model II used the same classification but excluding the face-selective sites. Significant discrimination between faces and non-faces was only observed in two subjects out of eight (permutation test, \(p < 0.05\) FDR corrected). In these two subjects, the significant decoding accuracy suggests that at least one site contains information about the discrimination at hand. This could however be related to selective responses to non-face stimuli, as we had not excluded sites selective to one or more subcategories of non-face stimuli. This first result suggests that face information on task-active sites is weak in the considered time window. Comparing models I and II revealed that there is a significant decrease in accuracy when excluding face-selective sites from the classification across subjects (Wilcoxon signed-rank test, \(p = 0.0078, n = 8\)). This result suggests that face sites are relevant in decoding settings for face processing\(^ {45}\) and hence causally related to face stimuli. To test whether random subsets of task-active sites are relevant in decoding settings, we removed all task-active sites from the classification model (model IIIa). This led to an average decrease in model performance of 4.9% (Supplementary Table 5, Fig. 2a), with a large variability across subjects (min: +0.86%, max: −11.66%). While these results display a significant effect of task-active sites on model performance (Wilcoxon signed-rank test, \(p = 0.0234, n = 8\)), they were built using only ~13.45% of the features. To disentangle a decrease in feature space from a decrease in (face) signal, we built 499 models that included the same number of sites as model II by randomly selecting task-active and face-selective sites, but including at least one face-selective site. This means that across the 499 models (referred to as “random set” models), the proportion of face-selective sites varied, but was non-null. This procedure guarantees the same number of features as in model II.

**Fig. 2 Nature and amount of face information on TC sites in decoding settings.** a Dark grey triangles represent the performance of model I, i.e., including all TC sites. Performance of model II is represented by red triangles. Red circles represent the performance of model IIIa. Each colored dot represents one of the “random set” models (499 models per subject, model III), their color representing the proportion of face sites included in the model (dark blue is one face site, light green is all face sites). Violin plots represent the distribution of the “random set” model performances compared to models I and II. b Sparse models perform as well or better than distributed models. Bar plot representing the balanced accuracy for model I compared to the model accuracy of the sparse model IV, for each subject. c Site contributions to the sparse model, plotted across subjects. The contribution of the site (in %) is represented by a color-coded fill. Black diamonds have a perfectly null contribution to the model while circles had a positive contribution to the model. Sites assessed as human face selective by the univariate analysis are highlighted by a pink rim.
and asserts whether removing random subsets of task-active sites affects performance. Across subjects, removing random sets of task-active and face-selective sites did not affect the model performance significantly (Wilcoxon signed-rank, $p = 0.5781$, Fig. 2a), nor did it within subjects for seven out of eight (Supplementary Table 5). For subject 2, a multimodal distribution is observed reflecting whether a specific face-selective site is included in the model (leading to accuracies > 79%) or not (accuracy < 70%, 83 models out of 499). There is a significant difference in model performance when this site is included or excluded from the model ($p < 0.0001$). The results from models IIIa and III suggest that task-active sites had a limited effect on the decoding model and were at best weakly relevant. Notably, we found a significant relationship between the proportion of face sites included in the analysis (i.e., from one face site to all sites, randomly selected in the 499 models) and the model performance for six out of seven subjects (Supplementary Table 5, $n = 499$). This result could arise from two scenarios (or a combination of the two): either face information on face sites is not redundant, i.e., each face site brings unique face information and/or including more sites leads to better SNR of the face pattern. In both cases, this result suggests that the pattern can be more easily identified when more human face sites are included.

The degree of inter-subject variability (both for model II and the “random set” model III) could not be explained by the number of trials ($\rho$(model II—#trials) = 0.073, $p = 0.1994$, $n = 8$; $\rho$(median random sets—#trials) = 0.0068, $p = 0.9872$, $n = 8$) or of sites included in the analysis ($\rho$(model II—#sites) = −0.1221, $p = 0.7734$, $n = 8$; $\rho$(median random sets—#sites) = 0.00156, $p = 0.9708$, $n = 8$). The inter-subject variability, however, could be explained by various factors, such as SNR, amount of correlated noise or placement of the electrodes, which are complex to quantify. Please note that due to the limited numbers of trials per subject, we were not able to perform the univariate and multivariate analyses on different partitions of the data. See additional analyses in Supplementary Table 6 and Supplementary Discussion 6.

To further investigate face processing in task-active sites, we performed the same classification as for model I, except that the considered algorithm enforces sparsity at the site level, i.e., it automatically selects a subset of sites to perform the classification. Comparing sparse and non-sparse modeling techniques is a common machine learning strategy to investigate data properties: Support Vector Machine (SVM) (i.e., model I) assumes that the information is fully distributed across features, by construction. If this assumption is correct, it should perform better than a sparse model. In contrast, if the sparse model performs better than or as well as model I while including only a subset of sites, it suggests that the information contained in the selected subset of sites is sufficient to perform the classification and that the non-selected features do not bring further relevant information. As in previous publications, we used the simple MKL algorithm to perform the sparse modeling. Please note that the basis algorithm for the simple MKL is an SVM, hence the effect of implementation on the results is limited. The results are displayed in Fig. 2b and Supplementary Table 7. Across all subjects, the sparse model performs slightly better than the distributed SVM model (Wilcoxon signed-rank test, $p = 0.25$, $n = 8$, average increase in performance of 2.28%). In five subjects out of eight, there is a clear increase in performance, of up to 8.95% for subject S1. In subjects 5, 6, and 8, the difference in model performance between model I (SVM) and model IV (sparse MKL) is around −2%. Of note, in two subjects the accuracy of the model was over 90%, which left little room for improvement (S5 and S8). This accuracy was reached with 35% of the sites being included, displaying that most sites did not contribute to the model. We however preclude from relating a contribution to the model to face processing as some sites can act as “noise canceller”.

In addition, the sparse model weight maps significantly overlapped with the univariate maps of human face selectivity across subjects ($\rho = 0.4708$, $p < 0.0001$, Fig. 2c) and within subjects for seven out of eight subjects (except subject 8, Supplementary Table 7, Supplementary Fig. 2). This result shows that MKL relies heavily on face-selective sites for the classification. For subject 8, the classifier seems to rely on non-face information, as the site with highest contribution contains higher amplitude signals for non-face categories than for human faces (pooled non-face > face: $p < 0.05$, permutation test). These results suggest that the potential face information present in task-active sites is correlated with the information present in face-selective sites. Please note that this analysis does not suffer from circularity as it does not rely on the previous univariate results.

### Temporal distribution of face processing

Thus far, our univariate and multivariate results display that face sites are “relevant” for human face processing. However, we also showed that including more face sites in the model increases model performance (random set models). We then explored the relationship between selectivity, anatomy, and timing of the HFB responses on face and task-active sites during human face processing.

While the face sites displayed significant responses to human face stimuli, the selectivity of their responses decreased from posterior to anterior sites as demonstrated by correlating the MNI y-coordinate with selectivity ($\rho = 0.06097$, $p = 5.39e−06$, $n = 47$). For task-active sites, selectivity to human faces increased from posterior to anterior sites ($\rho = 0.2699$, $p = 0.0013$, $n = 139$). In addition, our fast event-related paradigm combined with simultaneous recordings across selective and task-active sites with high sampling rate allowed us to compare the latency of neuronal population responses to faces within the first 500 ms of stimulus presentation. We measured the response onset latency (ROL) to human face stimuli across human face-selective sites and task-active sites based on unsmoothed, normalized signals (Figs. 3, 4). A recent study revealed a temporal lag between the posterior fusiform gyrus (pFG, $y < −45$) and medial fusiform gyrus (mFG, $−45 < y < −35$) across face-selective sites. In line with these anatomical boundaries, our data indicates an average temporal delay of 33 ms across the two regions and a further delay of 13 ms with the more anterior face-selective region ($y > −35$). Furthermore, ROL values across human face-selective sites were significantly correlated with the y-coordinate of the corresponding site ($\rho = 0.5207$, $p = 9.5399e−04$, $n = 37$). Interestingly, the posterior-to-anterior lag was also significant for task-active sites ($\rho = 0.6563$, $p = 7.3554e−07$, $n = 46$, Fig. 4). Importantly, the group-level findings presented here were also present at the individual subject level for human face-selective sites, when calculating the ROL for each site relative to the most human face-selective site within subject (Fig. 3b–d): $p$(ROL−y) = 0.7593, $p = 1.7911e−06$, $n = 29$ and $p$(selectivity−y) = −0.5680, $p = 0.0013$, $n = 29$. In other words, the posterior to anterior ROL gradient was not driven by one or a few subjects who happened to have coverage over a specific region of the TC. For task-active sites (Fig. 4b, c), the correlation between ROL and y-coordinate was also significant: $\rho = 0.6718$, $p = 3.2032e−07$, $n = 46$. On the other hand, there was no significant correlation between anatomical position and selectivity when compared to the most face-selective site: $\rho = 0.2269$, $p = 0.1294$, $n = 46$. We remind the reader that task-active sites showing non-selective responses can be of two types: those in the earlier visual cortices (located more posterior than the first face-selective sites) and engaged much earlier in presumably low-level processing of visual information and those
located in the anterior temporal lobe that are engaged much later in presumably high-level processing of visual information.

We explored the timing of events across task-active and face-selective sites (both including posterior and anterior sites), and compared the ROL values between the two groups, but this comparison was not significant \((p = 0.9452, \text{Wilcoxon rank sum test})\). We argued that the lack of difference could be due to large variability in ROL values within each group. To mitigate this confounding effect, we matched face-selective sites to their closest task-active sites in terms of \(y\)-coordinate (up to 2 mm difference), and found that the time of activation in face-selective sites trended earlier than anatomically matched adjacent non-selective sites but this difference did not reach statistical significance (paired Wilcoxon signed-rank, \(p = 0.1752, n = 20\) face sites matched to 16 unique task-active sites). Figure 4d shows the distribution of ROL values across the two groups after matching.

Importantly, signal amplitude or slope did not affect our ROL method (Supplementary Fig. 4, Supplementary Discussion 8).

**Effect of electrical brain stimulation on face perception.** Lastly, we explored the effect of electrical perturbation of face-selective sites on the subjective processing of faces. We hypothesized that the stimulation of face-selective sites in the posterior TC would cause more salient effects than the stimulation of non-selective sites in the anterior TC. This hypothesis has not been addressed in prior to electrical stimulation studies showing distortion in conscious viewing of faces\(^{26-28,30,51}\) or naming of famous faces\(^{52,53}\). Subjects were instructed to view a human face at the bedside while we performed active or sham (zero current) stimulations of selective, task-active, or non-responsive sites. Subjects then reported whether the face remained the same or was distorted. We emphasize that this anecdotal report departs from the well-controlled experimental procedures that have been performed in non-human primates and human subjects due to limitations in our equipment and clinical time constraints.

Figure 5 summarizes the results of electrical stimulations and details of the procedure in each patient is provided in Supplementary Table 9. Our findings indicate that distortion of human face perception was reported only with real (but not sham) stimulation of some (but not all) face-selective sites. Among 52 stimulated sites in the fusiform area, 24 were face selective and 28 were not. Among the 24 face-selective sites (17 pFG sites and 7 mFG sites), 6 caused distortions in face perception when stimulated while 18 did not; all 6 sites that caused distortions in face perception when stimulated were located in the posterior fusiform gyrus \((y < -45)\). Among the 28 non-selective sites (14 pFG sites and 14 mFG sites), 3 caused distortions in face perception when stimulated while 25 did not; again, all 3 sites that caused distortions in face perception when stimulated were located in the pFG. We found a significant effect of anatomical site in predicting the subjective responses to electrical stimulation: one-tailed \(z\)-score tests for two population proportions showed that the proportion of sites whose electrical stimulation induced a face distortion in the pFG \(28/31\) versus the same proportion in the mFG area \(0/21\) was significantly different \((z = 1.833, p = 0.034)\). This trend held true when comparing the proportion of pFG versus mFG sites that distort

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**Fig. 3 Temporal distribution of human face information on face-selective sites.** \(a\) Average traces and ROL according to \(y\)-coordinate. Mean \(\pm\)standard error HFB traces across face-selective sites in the posterior \((-45, n = 24)\), middle \((-45\) to \(-35, n = 7)\), and anterior \((-35, n = 6)\) TC sites for human faces (pink) and non-faces (grey) stimuli. \(b\) Response onset latency (ROL) over human face-selective sites for the human face category, as represented by a purple color scale fill on the MNI cortex with the best (i.e., most human face selective) site in each individual chosen as the point of reference (triangles). Here only relevant electrodes are shown. \(c\) ROL in each subject, when compared to MNI \(y\)-coordinate with the best site in each individual chosen as the point of reference. Latency is related to anatomical location of the sites. \(d\) Selectivity in each subject, when compared to MNI \(y\)-coordinate with the best site (i.e., most human face selective) in each individual chosen as the point of reference. The best site is represented as a black circle, at the crossing of the two axes. Selectivity is related to anatomical position.
face perception when stimulated within face-selective sites only (pFG: 6/17; mFG: 0/7; $z = 1.815$, $p = 0.035$) and within non-selective sites only (pFG: 3/14; mFG: 0/14; $z = 1.833$, $p = 0.034$). These results indicate that electrodes that cause distortions in face perception when stimulated are greater in proportion in the pFG than in mFG.

Moreover, among the nine pFUS sites that caused face-perception distortion, six were face-selective sites and three were sites immediately abutting the face-selective sites within the likely area of distribution of electrical charge. In fact, two of the abutting task-active sites displayed weak face selectivity (permutation test, $p < 0.05$ before FDR correction) during face processing. Based on our previous quantification of the spread of electrical charge in the human brain$^{34}$, we believe that the electrical fields generated by the stimulation of the three task-active sites likely spread to the abutting face-selective sites.

Of note, face distortion occurred only with right hemisphere stimulations except in one left hemisphere case (S1) who happened to be left handed. We have previously discussed the lateralized effect of stimulation in a separate report$^{35}$. As such, we find that stimulation of posterior face patches, in the language non-dominant hemisphere, causes face specific distortions.

In one subject (subject 8), we probed the effect of stimulations across two patches of the FFA (sites numbered as 1, 2, and 3 in Fig. 5b). Using bipolar (adjacent pair of sites stimulated together) or unipolar stimulations (site was paired with a distant reference), we elicited distortions when the subject looked at his own face in the mirror, looked at a cartoon face drawn on a piece of paper, and focused on parts of a face (e.g., eyes or lips). He also reported induced perceptions of a face during electrical stimulation while his eyes were closed. The subject’s verbal reports after stimulation include “one side of the face changed”; “facial features [turned] into a cartoon”; one eye “became someone else’s”; “face wiggled a little bit”; and “face looked familiar” (Supplementary Movie 1).

**Discussion**

Our study addresses the spatiotemporal distribution of face information based on univariate measures, machine learning-based modeling, timing analysis, and direct cortical stimulation. Using this multipronged approach and by leveraging the temporal resolution of the ECoG method, we confirmed the following results: the majority of recording sites in the human TC do not show any significant change of activity in response to the visual presentation of faces; a minority of sites respond non-selectively to at least one category of visual stimuli; and only fewer sites respond selectively to face stimuli, and that these are clustered in the fusiform gyrus or lateral occipital gyrus across individuals.

We found that face information was represented to a greater degree in face-selective regions than in task-active sites. Removing face-selective sites from machine learning-based classification significantly dropped the decoding accuracy of the model. In addition, classification model suggested that task-active sites had a limited effect on the decoding model and were at best weakly relevant (in the technical sense as used in Weichwald et al.$^{45}$).

**Fig. 4 Temporal distribution of human face information on task-active sites.**

- **a** Mean (±standard error) HFB traces across task-active sites in the posterior ($<-45$, $n = 24$), middle ($-45$ to $-35$, $n = 7$), and anterior (after $-35$, $n = 6$) sites for human faces (pink) and non-faces (grey) stimuli.
- **b** Selectivity in each subject, when compared to MNI $y$-coordinate with the best (i.e., most human face selective) site in each individual chosen as the point of reference. The best site is represented as a black circle, at the crossing of the two axes. Selectivity is not related to anatomical position.
- **c** ROL in each subject, when compared to MNI $y$-coordinate with the best site in each individual chosen as the point of reference. Latency is related to anatomical location of the sites.
- **d** Histograms (in probabilities) of ROL values for human face-selective sites ($n = 32$) and their matched task-active sites in terms of $y$-coordinate ($n = 19$). N.s. refers to a non-significant difference between the two distributions.
This was supported by an improvement in model performance when considering a sparse approach. Overall, our results suggest that face information in task-active sites is weak, highly correlated with the face information that is present (earlier and stronger) in face-selective sites, and not causally necessary nor sufficient (in terms of our machine learning analysis) for face processing. We confirm the presence of temporally distributed information within task-active and face-selective sites that may be functionally relevant and even necessary for downstream processing of faces (e.g., associating faces with memory or name information).

Comparing our results to prior neuroimaging studies, we would like to emphasize the methodological differences including the number of recording sites, the time scales considered, and the recorded signals used in the analysis. For instance, the number of anatomical samples in each subject is limited with the ECoG method, which can reduce the power of the algorithm in detecting weak, distributed patterns. To maximize the detection of small effects, we used Support Vector Machine classifiers, which are known for their ability to detect subtle, distributed patterns and we focused our analysis on the high SNR of HFB responses induced by human face versus non-face stimuli. While we could not relate the number of sites to model performance (Results section), it is possible that the signal in task-active sites is too weak to be detected across a few tens of anatomical samples in each subject. On the other hand, the novelty of our data is in part due to the high temporal resolution of ECoG, which enabled us to measure the fast temporal dynamics of face processing in the human brain. Specifically, our electrophysiological analysis relied on responses elicited within 500 ms of stimuli presented for 300 ms with an inter-stimulus interval of 400 ms. This is in stark contrast to some of the classic neuroimaging studies whose temporal window included >10 s of signal processing (e.g., 24 or 16 s long blocks of visual stimuli for each category).

Another methodological caveat that needs to be considered is that the task-active areas in humans may be variable in their relative size and primary cytoarchitectonic composition. Therefore, hubs of activity in posterior to anterior TC may have different sizes or shapes of physiological responses or locations on the surface of gyri versus depth of sulci which could have influenced the properties of recorded electrophysiological signals as well as the subjective effect of electrical stimulation. As such, the results reported here could represent an idiosyncrasy of our iEEG method. However, it is still noteworthy that responses to faces were significantly faster in the posterior sites than in anterior face-selective sites. By using ECoG recordings, our results confirm our overarching hypothesis that face information is anatomically localized but temporally distributed. The posterior to anterior gradient was observed within both face-selective and task-active sites, with no statistically significant time differences between task-active and face-selective sites—though the time of activations in face-selective sites trended earlier (Fig. 4d). The lack of statistical significance in terms of ROL can be explained by several factors. First, there is a posterior-to-anterior temporal lag in activation, i.e., anterior face-selective sites may respond later than posterior non-selective task-active sites. Second, there are task-active sites in the more posterior regions that respond...
significantly earlier than any other sites. For instance, one cannot assume that face-selective activations precede the ones in the primary visual cortex. Third, given these two confounds, the number of electrodes in each subject may be too small for statistically differentiating a subtle temporal lag between selective and adjacent non-selective sites. Finally, despite the trending results reported here, we would caution against any conclusions from it. Given our knowledge of brain anatomy, we would anticipate that task-active and face-selective sites would be engaged in bilateral information exchange at multiple levels along the posterior—antero axis of signal propagation and, as such, the ROI values across sparse recording sites in each individual brain may not be able to address this issue and addressing it with IEEG may be ill-advised to begin with.

Additionally, our findings support the causal link between some, but not all, face-selective sites of the fusiform gyrus and conscious processing of faces as suggested previously by our group. Our findings from electrical stimulation procedures (1) replicate our prior findings in a new cohort of subjects that the stimulation of the right (non-language dominant) hemisphere is more important for eliciting subjective reports of face distortion; (2) that some of the posterior face-selective sites (i.e., y > 45 in MNI space) are more critical than anterior sites for conscious processing of faces, and (3) that the pFUS (but not the mFUS) region may serve as a key node in the conscious processing of faces. While the low numbers of anterior site stimulations preclude any definite conclusions, the third finding is intriguing and needs to be replicated in a larger sample of subjects with a task-based and stimulus-locked electrophysiological procedures. As demonstrated in non-human primates, and also in the human lateral occipital and fusiform cortices, we acknowledge that the face-selective sites outside the fusiform gyrus also play an important role in face processing. For example, a recent study in non-human primates showed that the micro-stimulation of the anterior face-selective patch (area anterior medial) severely distorted the monkey’s percept of facial identity, such that faces depicting the same identity appeared to depict different identities.

Our findings may provide neurophysiological explanations for some of the classic behavioral observations. The preferential responses to human faces compared to non-human faces is in line with a recent human functional magnetic resonance imaging (fMRI) study, in which multivariate patterns of activity throughout the ventral TC were found to correlate with behavioral judgments of biological similarity of the same stimuli (i.e., monkeys and mammals versus insects and birds). Moreover, two electrophysiological studies in the macaque brain have shown faster responses of neurons to primate faces compared to non-primate animal faces, or a change in species selectivity across different patches of face-selective areas. Our findings along with these studies may explain the behavioral observations of conspecific advantage in face recognition; that is, humans and other primates recognize members of their own species more readily than faces of other species. For instance, chimpanzees raised in a human environment are better at discriminating pictures of unknown human faces than unknown chimp faces.

In closing, our data suggest that a few anatomically discrete sites play a crucial role in processing face information and that there is a time delay in their processing of the same information. However, our study does not suggest that the conscious perception of faces solely depends on the operation of these isolated patches of face-selective sites. We acknowledge that different facets of face processing may occur in different patches of face-responsive sites and that these neuronal patches are embedded in a larger network of visual and other association areas of the brain, and their function should not be seen in isolation from this wider integrated brain network. Future studies with simultaneous recording across face-selective and other association areas are needed to determine how different facets of face information is encoded in each of the face patches and how their function is embedded and relayed to the rest of the brain for serving human cognition and behavior.

Methods

Demographics and recordings. Eight subjects (six males, two females, aged 23 and 68 years) were implanted with intracranial electrodes to localize the source of drug-resistant seizures. The procedure was approved by the Stanford Institutional Review Board and the subjects provided written informed consent to participate in the study. The location of the grids was determined by clinical needs (Supplementary Fig. 1, three left hemisphere implantations, five right). Data were obtained at 1.5 T (83.88 Hz through a 128-channel recording system (Tucker Davis Technologies) for the first seven subjects while a Nihon Kohden Technology system with simultaneous video monitoring was used to perform 1 kHz recordings in subject 58. Each electrode was a platinum plate, either 2.3 mm or 1.15 mm in diameter (exposed recording area) with center-to-center spacing of 4–10 mm between adjacent electrodes on the grid or strip. Electrodes containing artifacts or pathological activity were discarded from further analyses.

Anatomical localization of electrodes. Structural MRIs were acquired with a GE 3-Tesla Sigma scanner at Stanford University equipped with a head coil of a T1-weighted SPCRG pulse sequence. The images were AC–PC (anterior commissure–posterior commissure) aligned and were resampled to 1 mm isotropic volume65. Electrodes were visualized on the subject’s own brain volume and reconstructed onto a three-dimensional cortical surface allowing for accurate anatomical localization of electrodes. The electrode positions were also transposed into the MNI space and displayed on a MNI cortex file for visualization of results across subjects. The coordinates of the left hemisphere sites were mirrored such that all sites could be displayed on a single hemisphere.

Experimental paradigm. The experiment was administered using psychtoolbox running on Mac OSX. The laptop was placed ~70 cm from the subject’s eyes at chest level. Screen resolution was 1280 × 800. Each image was subtended five visual degrees in its longest dimension. Each subject underwent a visual task during which images of different categories were presented at the center of the screen for 300 ms, with an ISI of 400 ms. The categories included ten human face, human body, mammal face (one monkey face), mammal body, bird face, bird body, marine face, marine body, human limbs, object, and place. The image backgrounds were phase scrambled at 3% in order to reduce visual artifact. The visual dimensions of the image plus its scrambled background were 11.10 × 11.10 cm, and the visual angle was 9 degrees. Each category comprised 25 images, presented twice. This hence leads to 50 stimuli per category and 550 visual stimuli in total. During image presentation, the subject was asked to press a key (‘press 1’) when the pattern appeared in red at the center of the screen (further referred to as a “Response Block”). The onset of each stimulus was recorded by a photodiode signal generated by a luminance change in the display at image onset.

Signal preprocessing. All preprocessing steps were performed using Matlab (The MathWorks, Inc., Natick, Massachusetts) and the SPM (www.fil.ion.ucl.ac.uk/spm) toolbox in custom routines (https://github.com/LBCN-Stanford preprocessing pipeline). The data were first down sampled to 1000 Hz and filtered for power-line noise (band-stop between 57 and 63 Hz) and harmonics (117–123 Hz and 177–183 Hz). Sites underwent an automatic quality assessment: sites with variances five times larger or smaller than the average variance across all sites were labeled as pathological and excluded. Sites with three times lower average signal (defined as changes in the signal due to the signal from adjacent sites compared to the averaged across sites were considered as spiky and excluded. The signal was then re-referenced to the average of the signal over all selected sites. Each event was extracted (i.e., epoched) in the −200 to 700 ms time window around its onset and baseline correction was performed (using the [−200 to 0]ms time window around onset as baseline). Events were marked as artifacts if they contained spikes of >100 μV and were discarded from further analyses. At this stage, there were no further checks for interictal activity. A time-frequency decomposition was then computed using a seven-cycle Morlet wavelet, with frequencies ranging from 70 to 177 Hz (steps of 1 Hz, avoiding discarded frequencies from Notch filtering). A similar five-cycle Morlet wavelet time-frequency decomposition was performed for frequencies ranging from 1 to 69 Hz. The power in each frequency and time bin was rescaled using the log-power of the [−100 to 0]ms window around stimulus onset. Six frequency bands were considered in this work: θ (4–7 Hz), α (8–12 Hz), β1 (13–29 Hz), β2 (30–39 Hz), γ (40–69 Hz), and HFB (70–177 Hz). The signal was finally averaged across frequency bins within each band considered and smoothed with a 50 ms width Gaussian window. The HFB power in the [−100 600]ms window around stimulus onset was considered for...
Considered as relevant in encoding settings for face processing if they displayed a significant HFB response in the [150 500]ms after stimulus onset for at least one category, when compared to the event baseline ([−100 0]ms before onset). Paired non-parametric permutation tests (50,000 permutations) assessed the significance of the response \( p < 0.05 \), FDR corrected for the number of sites tested. Out of the active sites, "face-selective" sites were identified as sites where significantly higher responses were seen to the four face categories (human, mammal, bird, and marine) pooled compared to all other stimuli pooled (non-parametric permutation tests, FDR corrected) in the time window [150 500]ms after onset. Finally, "human face-selective" sites displayed significantly higher responses to human faces compared to all non-face stimuli pooled (non-parametric permutation tests, FDR corrected) in the time window [150 500]ms after onset. Sites that were not assessed as face selective or human face selective are further referred to as "task-active" sites. Sites assessed as (human) face selective were considered as relevant in encoding settings for face processing if they displayed a significant HFB response in the [150 500]ms time window after onset. Permutation tests assessed the significance of potential differences between subcategories (10,000 permutations, FDR corrected for the number of tests \( n = 12 \), 6 binary comparisons for face categories) and \( 6 \) for (task-active sites). Please note that there is no circularity in this analysis as the contrast to select sites (i.e., faces versus non-faces) is different from the effect investigated (i.e., human faces versus mammal faces versus bird faces versus marine faces).

### Frequency information for faces

For each frequency band, a univariate analysis assessed active, face-selective, and human face-selective sites in addition to the four face categories (human, mammal, bird, and marine) pooled compared to all other stimuli pooled (non-parametric permutation tests, FDR corrected) in the time window [150 500]ms after onset. Sites that were not assessed as face selective or human face selective are further referred to as "task-active" sites. Sites assessed as (human) face selective were considered as relevant in encoding settings for face processing if they displayed a significant HFB response in the [150 500]ms time window after onset. Permutation tests assessed the significance of potential differences between subcategories (10,000 permutations, FDR corrected for the number of tests \( n = 12 \), 6 binary comparisons for face categories) and \( 6 \) for (task-active sites). Please note that there is no circularity in this analysis as the contrast to select sites (i.e., faces versus non-faces) is different from the effect investigated (i.e., human faces versus mammal faces versus bird faces versus marine faces).

### Low-level image features

To ensure that low-level features in the stimulus images did not drive our results, we performed multiple control analyses. First, spatial frequency power spectrum with rotational average was calculated for each stimulus and averaged across categories. Averaged spatial frequency power spectrum were compared using the one-way ANOVA and post hoc analysis was performed using the "Tukey-Kramer" method. Power spectral analysis returned 153 values for each image, which were averaged within each category (\( n = 153 \)). We also computed the mean luminance across pixels in each category, pooling all non-faces together.

We then investigated the effect of mean luminance on the univariate results (i.e., on sites defined as face selective). To this end, we plotted the histogram of mean luminance in the face and in the non-face categories. We defined as low (resp. high) luminance faces, face stimuli with a mean luminance smaller (resp. larger) than 120 (threshold defined arbitrarily). The neural signals in each category was compared on all face sites (high luminance faces: \( n = 45 \), human:15, mammal:12, bird:7, and marine:1; low luminance faces: \( n = 55 \), human:10, mammal:13, bird:18, and marine:14), using permutation tests.

### Relevance of TC sites in encoding

We then assessed the relevance of face and task-active sites of sites in decoding settings. To this end, a machine learning algorithm was trained using the comparison of human face and non-face epochs as the target. However, the proportion of face sites for modeling was kept consistent with models I, II, and IV (see "Relevance of TC sites in decoding"). In addition, a second MKL model combined the power in HFB and the ERP (i.e., before time-frequency decomposition) as a recent study showed that ERPs could bring complementary decoding information to HFB signals. These analyses were performed in version 3.0 of the Pattern Recognition Toolbox (PhoNTo).

### Sparse versus distributed decoding information

Assessing relevance in encoding and decoding settings assumes the interpretation of negative results. As negative results are by nature inconclusive, we here investigate the effect of priors on model performance. For one model (model I), the algorithm used assumes a "distributed" prior (SVM, L2-regularization), i.e., all features contribute to the model. If this prior is appropriate, model performance should be high, and potentially higher than for other types of prior. We test this hypothesis by comparing model I to a model that enforces sparsity on the sites. Thereby, if the sparse model performs better or as well as a distributed model, the face information is likely captured by another feature category bringing in formation. The sparse model, further referred to as model IV, is based on the sparse MKL method. The main difference between model I is that sparsity is enforced at the site level. In practice, one linear kernel is built per site (i.e., number of features = \( 1 \times 351 \) time points) and those kernels are combined through a sparse prior (L1-norm penalty) for each category. In this section, we perform univariate analyses on another visual task recorded on the same subjects that includes human faces and another visual task recorded on the same subjects that includes human faces and non-faces. As negative results are by nature inconclusive, we here investigate the effect of priors on model performance. For one model (model I), the algorithm used assumes a "distributed" prior (SVM, L2-regularization), i.e., all features contribute to the model. If this prior is appropriate, model performance should be high, and potentially higher than for other types of prior. We test this hypothesis by comparing model I to a model that enforces sparsity on the sites. Thereby, if the sparse model performs better or as well as a distributed model, the face information is likely captured by another feature category bringing in formation. The sparse model, further referred to as model IV, is based on the sparse MKL method. The main difference between model I is that sparsity is enforced at the site level. In practice, one linear kernel is built per site (i.e., number of features = \( 1 \times 351 \) time points) and those kernels are combined through a sparse prior (L1-norm penalty) for each category. In this section, we perform univariate analyses on another visual task recorded on the same subjects that includes human faces and non-faces.

Circular analysis: With a non-null null hypothesis, we here investigate the effect of priors on model performance. For one model (model I), the algorithm used assumes a "distributed" prior (SVM, L2-regularization), i.e., all features contribute to the model. If this prior is appropriate, model performance should be high, and potentially higher than for other types of prior. We test this hypothesis by comparing model I to a model that enforces sparsity on the sites. Thereby, if the sparse model performs better or as well as a distributed model, the face information is likely captured by another feature category bringing in formation. The sparse model, further referred to as model IV, is based on the sparse MKL method. The main difference between model I is that sparsity is enforced at the site level. In practice, one linear kernel is built per site (i.e., number of features = \( 1 \times 351 \) time points) and those kernels are combined through a sparse prior (L1-norm penalty) for each category. In this section, we perform univariate analyses on another visual task recorded on the same subjects that includes human faces and non-faces.
sites. It is important to note that this visual task comprises a different contrast from our main task, as the visual task mostly comprises images of words and numbers, and not pictures. This explains why we chose not to include this task in our main analyses.

All subjects underwent a visual localizer comprising images of human faces, animal (faces with bodies), places, objects, logos, false fonts, English and Spanish words, numbers and Persian numbers. Each image was presented for 400 ms, with an inter-stimulus time interval of 300 ms. During the stimulat presentation, the patient was asked to pay attention to the center of the screen where a dot that would randomly change color (from red to blue) was displayed. The patient was asked to respond using a keyboard (“press 1”) when the dot changed color. This was a median over-trial trial-to-trial ROL estimates in order to assign singular ROL values for specific sites. Sites for which a ROL value could not be obtained in 50% of the trials or more were discarded from the analysis.

To estimate the effect of signal slope on the ROL results, we performed the same simulation while normalizing the amplitude of the signal in each trial before ROL detection. We estimated potential group differences in ROL between face-selective and task-active sites. The amplitude of the signal in the ramp window was varied by modifying the SNR between trials A and B. Hence, varying signal amplitude is strongly correlated with varying the selectivity of sites to condition A. The amplitude of the signal in the ramp window was computed based on a desired SNR on each site:

\[ X_{\text{eff}} = X_b + \text{SNR}_b \times \text{std}(X_b) \]

where \( X_{\text{eff}} \) represents the amplitude of the effective simulated signal for condition A trials, \( X_b \), the amplitude of the signal for trials A, \( \text{SNR}_b \) a fixed number representing the desired SNR, and \( \text{std}(X_b) \) the average trace of “B” trials. SNR was varied from 2 to 10 by steps of 0.5. In our real dataset, the distribution of estimated SNR varies from −2 to 17 on human face-selective sites, with only three sites with SNR > 10.

To estimate the effect of signal slope on the ROL results, we performed the same simulation while normalizing the amplitude of the signal in each trial before ROL detection (\( \text{L}_\text{norm} \), i.e., dividing by the maximum amplitude). This simulated dataset hence varies the slope of the signal, but not the amplitude (set to maximum of 1).

For each SNR level, ROL detection is performed at the trial level, for the “A” trials on both each “good” site. All parameters are identical to the technique reported above. Trials with <50% of detected offsets were excluded from the results. This analysis was performed on both the un-normalized (i.e., varying amplitude at fixed slope) and the normalized (i.e., varying slope at fixed amplitude) simulated data. The results are presented in Supplementary Discussion 8 and Supplementary Fig. 4.

Causal importance of TC sites for face perception. A set of electrical brain stimulations (EBS) was performed on seven of the nine subjects in addition to the experimental task (but not concurrently). The sites of stimulations were chosen based on a priori knowledge about their HBF responses to face and non-face stimuli, as assessed by the univariate analysis of a localizer task. During the procedure, face-selective and task-active sites were stimulated with electrical charge where several parameters were kept fixed, such as varying the stimulus frequency (per sessions at the bedside). Across the seven subjects, the instructions given included (A) looking at the face, (B) looking at the lips, (C) looking at the nose, (D) looking at self in the mirror, (E) looking at a cartoon face on a sheet of paper, or (F) close eyes and imagine a face. Please note that these clinical procedures differ from stimulus-locked electrical perturbation experiments. Between 3 to 8 mA (depending on the excitability of the stimulated site) were delivered at a duration ranging from 1 to 3 s at 50 Hz frequency and 200 μs pulse width, of a square wave electrical waveform in unipolar (subject SB) or bipolar montage (SB and other subjects). In unipolar montage, a TC site and a remote cortical reference site were stimulated whereas in the bipolar montage, a pair of adjacent electrodes were stimulated. Sham stimulations were also administered at 0 mA. Continuous EEG monitoring showed no after discharges or epileptic activity during the sessions. Verbal reports were collected following each stimulation. We recognized an electrode as a catalyst in face perception change if (A) the stimulation of this electrode yielded a face-specific change; (B) the stimulation of this electrode paired with any other electrode yielded a face-specific change; (C) the result of the stimulation and its resulting face-perception change was replicable across multiple stimulation trials. In addition, in subject SB, the estimated cortical area affected by the stimulation on electrode 3 was calculated using the relationship between the estimated charge per trial and the cortical area. The charge delivered per trial (μC) was calculated as a product of the pulse width (ms), current (mA), frequency (Hz), and duration (s) of stimulation for each trial; then, we estimated the cortical area (mm²) affected by the stimulation as a function of the charge deposited per trial (μC) according to the methods described in a previous publication. Furthermore, using a one-tailed z-test, we tested whether each of the sites in face perception where significantly different in the posterior versus mid fusiform gyrus across all subjects.

**Effects of signal amplitude and slope on ROL.** Our method performs response onset detection at the trial level, based on unsmoothed data. However, different parameters of the signal could affect the obtained ROL values, including noise, signal amplitude, and slope. In this section, we evaluate the effect of signal amplitude and slope on the obtained ROL values.

The level of noise is the one that is naturally present in ECoG data. The semi-smoothed data used in this work have been designed for other work and are described in detail below. The data and code for generating the simulation are available open-source (https://github.com/jessicaSchroff/Processed_ECoG). The original data was recorded from subject S1, during a 5-min wakefulness rest period, with eyes closed. Sites assessed as “pathological” by medical doctors were discarded from further analysis. A fake experimental design was simulated: two conditions: “A” and “B”, presented at random every 1.9 s. The stimuli are further assessed by signal processing. Signal preprocessing was performed with specific ECoG routines (github/LBCN/Preprocessing_Pipeline) using Matlab (www.mathworks.com) and SPM12 (www.fil.ion.ucl.ac.uk/spm). First, the data were converted to SPM format and downsampled to 1 kHz. The continuous signal was filtered for line noise and harmonics at 50–133 Hz. The data were band-pass filtered from 0 to 100 Hz. The data were then linearly detrended and band-pass filtered from 1 to 200 Hz. The data were then downsampled to 400 Hz. The data were then linearly detrended and band-pass filtered from 1 to 200 Hz. The data were then downsampled to 400 Hz. The data were then linearly detrended and band-pass filtered from 1 to 200 Hz.
Statistical testing. Throughout this work, statistical testing was performed using non-parametric permutations. When suited, the tests were paired (e.g., when comparing two conditions in terms of ROL value on the same set of sites, or when comparing the HFB response of a stimulus category to its baseline). A minimum of 1000 permutations was performed, to ensure a good estimation of the null distribution. FDR or Bonferroni correction was applied when multiple comparisons tested for the same effect (e.g., testing for human face selectivity on each site). Significance was determined at p < 0.05, after correction if applicable. Population statistics for the decoding models I–IV was performed using Wilcoxon signed-rank tests.

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

Data availability
Due to the presence of patient identifying information in the data, we cannot release the presented recordings.

Code availability
All analyses were performed in Matlab (www.mathworks.com). Preprocessing and univariate analyses were performed based on SPM (http://www.fil.ion.ucl.ac.uk/spm/) and in-house routines available at https://github.com/LBCN-Stanford/Preprocessing_pipeline. ROL in-house codes are available on Github at https://github.com/LBCN-Stanford/. Multivariate analyses were performed using a development version of pFReTo⁶⁹. This code will be released as pFReTo v3 and be available at http://www.mlnl.cs.ucl.ac.uk/pronto/. The code to build semi-simulated data is available at https://github.com/JessicaSchroff/Simulated_ECoG, along with the rest data from subject S1 used to generate the noise structure.

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References
1. Damasio, A. R., Damasio, H. & Van Hoesen, G. W. Prosopagnosia: anatomic basis and behavioral mechanisms. Neurology 32, 331–341 (1982).
2. Meadows, J. C. The anatomical basis of prosopagnosia. J. Neurol. Neurosurg. Psychiatry 37, 489–501 (1974).
3. Sergent, J., Ohta, S. & MacDonald, B. Functional neuroanatomy of face and object processing. A positron emission tomography study. Brain 115(Pt 1), 15–36 (1992).
4. Puce, A., Allison, T., Gore, J. C. & McCarthy, G. Face-sensitive regions in human extrastriate cortex. J. Cogn. Neurosci. 4, 388–451 (1992).
5. Vida, M. D., Nestor, A., Plaut, D. C. & Behrmann, M. Spatiotemporal dynamics of similarity-based neural representations of facial identity. Proc. Natl. Acad. Sci. USA 114, 388–393 (2017).
6. Yang, Y. et al. Exploring the spatio-temporal neural basis of face learning. J. Vis. 17, 1 (2017).
7. Bruce, C., Desimone, R. & Gross, C. G. Visual properties of neurons in a polysensory area in superior temporal sulcus of the macaque. J. Neurophysiol. 46, 369–384 (1981).
8. Hendrick, K. M. & Baldwin, B. A. Cells in temporal cortex of conscious sheep can respond preferentially to the sight of faces. Science 236, 448–450 (1987).
9. Umegaki, I., Tsuchiya, A., Nakata, K., Hasegawa, T. & Hayashi, T. Neural correlates of similarity-based facial identity processing. Neuroimage 113, 50–58 (2015).
10. Afzaal, M. A., Kiani, R. & Estokh, H. Microstimulation of inferotemporal cortex influences face categorization. Nature 422, 692–695 (2003).
11. DiCarlo, J. J., Zoccolan, D. & Rust, N. C. How does the brain solve visual object recognition? Neuron 73, 415–434 (2012).
12. Behrmann, M. & Plaut, D. C. Distributed circuits, not circumscribed centers, mediate visual recognition. Trends Cogn. Sci. 17, 210–219 (2013).
13. Haxby, J. V. et al. Distributed and overlapping representations of faces and objects in ventral temporal cortex. Science 293, 2425–2430 (2001).
14. Plaut, D. C. & Behrmann, M. Face-selective regions in the human fusiform gyrus. Cereb. Cortex 24, 1879–1893 (2014).
15. Keller, C. J. et al. Tuning face perception with electrical stimulation of the fusiform gyrus. Hum. Brain Mapp. 38, 2830–2842 (2017).
16. Hamarane, C. M. et al. Functional selectivity in the human occipitotemporal cortex during natural vision: evidence from combined intracranial EEG and MEG tracking. Neuroimage 95, 276–286 (2014).
17. Murphey, D. K., Maunsell, J. H., Beauchamp, M. S. & Yoshor, D. Perceiving electrical stimulation of identified human visual areas. Proc. Natl Acad. Sci. USA 106, 5389–5393 (2009).
18. Jacoby, C. et al. Corresponding ECoG and fMRI category-selective signals in human ventral temporal cortex. Neuropsychologia 83, 13–28 (2016).
19. Ranganujan, V. & Parvizi, J. Functional asymmetry between the left and right human fusiform gyri explored through electrical brain stimulation. Neuropsychologia 83, 29–36 (2016).
20. Ranganujan, V. et al. Electrical stimulation of the left and right human fusiform gyrus causes different effects in conscious face perception. J. Neurosci. 34, 12828–12836 (2014).
21. Parvizi, J. et al. Electrical stimulation of human fusiform face-selective regions distort face perception. J. Neurosci. 32, 14915–14920 (2012).
22. Schroot, J., Mourao-Miranda, J., Phillips, C. & Parvizi, J. Decoding intracranial EEG data with multiple kernel learning method. J. Neurosci. Methods 261, 19–28 (2016).
23. Canolty, R. T. & Knight, R. T. The functional role of cross-frequency coupling. Trends Cogn. Sci. 14, 506–515 (2010).
24. Crone, N. E., Miglioretti, D. L., Gordon, B. & Lesser, R. P. Functional mapping of human sensorimotor cortex with electrocorticographic spectral analysis. II Event-related synchronization in the gamma band. Brain 121, 2301–2315 (1998).
25. Nir, Y. et al. Coupling between neuronal firing rate, gamma LFP, and BOLD fMRI is related to interneuronal correlations. Curr. Biol. 17, 1275–1285 (2007).
26. Nir, Y. et al. Interhemispheric correlations of slow spontaneous neuronal fluctuations revealed in human sensory cortex. Nat. Neurosci. 11, 1100–1108 (2008).
27. Mukan, R. Coupling Between neuronal firing, field potentials, and fMRI in human auditory cortex. Science 309, 951–954 (2005).
28. Niesen, J. et al. Hemodynamic signals correlate tightly with synchronized neuronal oscillations. Science 309, 948–951 (2005).
29. Manning, I. R., Jacobs, J., Fried, I. & Kahana, M. J. Broadband shifts in local field potential power spectra are correlated with single-neuron spiking in humans. J. Neurosci. 29, 13631–13632 (2009).
30. Ray, S., Crone, N. E., Niebur, E., Franaszczuk, P. J. & Hsiao, S. S. Neural correlates of high-gamma oscillations (60–200 Hz) in macaque local field potentials and their potential implications in electrocorticography. J. Neurosci. 28, 11526–11536 (2008).
31. Logothetis, N. K., Pauls, J., Augath, M., Trinath, T. & Oeltermann, A. Physiological investigation of the basis of the fMRI signal. Nature 412, 150–157 (2001).
32. Boege, J. A. & Logothetis, N. K. Physiologically of the BOLD fMRI signal in awake monkeys. Curr. Biol. 18, 631–640 (2008).
33. Kreiman, G. et al. Object selectivity of local field potentials and spikes in the macaque inferior temporal cortex. Neuron 49, 433–445 (2005).
34. Liu, J. & Newsome, W. T. Local field potential in cortical area MT: stimulus tuning and behavioral correlations. J. Neurosci. 26, 7777–7790 (2006).
35. Ray, S. & Maunsell, J. H. Different origins of gamma rhythm and high-gamma activity in macaque visual cortex. PLoS Biol. 9, e1000610 (2011).
36. Mazzotta, J. C., Toga, A. W., Evans, A., Fox, P. & Lancaster, J. A Probabilistic Atlas of the Human Brain: Theory and Rationale for Its Development. Neuroimage 2, 89–101 (1995).
37. Weichwald, S. et al. Causal interpretation rules for encoding and decoding models in neuroimaging. Neuroimage 110, 48–59 (2015).
38. Hessdahl, Z. & McCarthy, G. Category representations in the brain are both discretely localized and widely distributed. J. Neurophysiol. 119, 2256–2264 (2018).
39. Schroff, J. et al. Embedding anatomical or functional knowledge in whole-brain multiple kernel learning models. Neuronformatics 16, 117–143 (2018).
40. Haufe, S. et al. On the interpretation of weight vectors of linear models in multivariative neuroimaging. Neuroimage 87, 96–110 (2014).
49. Li, Y., Richardson, R. M. & Ghuman, A. S. Posterior fusiform and midfusiform contribute to distinct stages of facial expression processing. *Cereb. Cortex* **29**, 3209–3219 (2018).
50. Jonas, J. et al. Self-face hallucination evoked by electrical stimulation of the human brain. *Neurology* **83**, 336–338 (2014).
51. Schalk, G. et al. Facephemes and rainbows: causal evidence for functional and anatomical specificity of face and color processing in the human brain. *Proc. Natl Acad. Sci. USA* **114**, 12385–12390 (2017).
52. Allison, T. et al. Face recognition in human extrastriate cortex. *J. Neurophysiol.* **71**, 821–825 (1994).
53. Puce, A., Allison, T. & McCarthy, G. Electrophysiological studies of human face perception. III: Effects of top-down processing on face-specific potentials. *Cereb. Cortex* **9**, 445–458 (1999).
54. Wimawer, J. & Parvizi, J. Linking electrical stimulation of human primary visual cortex, size of affected cortical area, neuronal responses, and subjective experience. *Neuron* **92**, 1213–1219 (2016).
55. Rangarajan, V. et al. Electrical stimulation of the left and right human fusiform gyrus causes different effects in conscious face perception. *J. Neurosci.* **34**, 12828–12836 (2014).
56. Weiner, K. S., Natu, V. S. & Grill-Spector, K. On object selectivity and the anatomy of the human fusiform gyrus. *Neuroimage* **173**, 604–609 (2018).
57. Weiner, K. S. et al. The cytoarchitecture of domain-specific regions in human high-level visual cortex. *Cereb. Cortex* **27**, 146–161 (2017).
58. Moeller, S., Crapse, T., Chang, L. & Tsao, D. Y. The effect of face patch microstimulation on perception of faces and objects. *Nat. Neurosci.* **20**, 743–752 (2017).
59. Connolly, A. C. et al. The representation of biological classes in the human brain. *J. Neurosci.* **32**, 2608–2618 (2012).
60. Kiani, R., Esteky, H., Mirpour, K. & Tanaka, K. Object category structure in high-level visual cortex. *J. Neurophysiol.* **97**, 4296–4309 (2007).
61. Tsao, D. The macaque face patch system: a window into object representation. *Cold Spring Harb. Symp. Quant. Biol.* **79**, 109–114 (2014).
62. Malpass, R. S. & Kravitz, J. Recognition for faces of own and other race. *J. Pers. Soc. Psychol.* **13**, 330–334 (1969).
63. Kravitz, D. J., Saleem, K. S., Baker, C. I. & Mishkin, M. A new neural framework for visuospatial processing. *Nat. Rev. Neurosci.* **12**, 217–230 (2011).
64. Martin-Malivel, J. & Okada, K. Human and chimpanzee face recognition in chimpanzees (Pan troglodytes): role of exposure and impact on categorical perception. *Behav. Neurosci.* **111**, 1145–1155 (2007).
65. Mesulam, M. M. Large-scale neurocognitive networks and distributed processing for attention, language, and memory. *Am. Neurol.* **28**, 597–613 (1990).
66. Hermes, D., Miller, K. J., Noordmans, H. J., Vansteensel, M. J. & Ramsey, N. F. Automated electrocorticographic electrode localization on individually rendered brain surfaces. *J. Neurosci. Methods* **185**, 293–298 (2010).
67. Miller, K. J., Schalk, G., Hermes, D., Ojemann, J. G. & Rao, R. P. Spontaneous decoding of the timing and content of human object perception from cortical surface recordings reveals Supplementary information in the event-related potential and broadband spectral change. *PLoS Comput. Biol.* **12**, e1004660 (2016).
68. Schrouff, J. et al. PRoNTo: pattern recognition for neuroimaging toolbox. *Neuroinformatics* **11**, 319–337 (2013).
69. Pereira, F., Mitchell, T. M. & Botvinick, M. Machine learning classifiers and fMRI: a tutorial overview. *Neuroimage* **45**, 5199–5209 (2009).
70. Baldassarre, L., Pontil, M. & Mourao-Miranda, J. Sparsest with stability: combining accuracy and stability for model selection in brain decoding. *Front. Neurosci.* **11**, 62 (2017).
71. Raccach, O., Daitch, A. L., Kuci, A. & Parvizi, J. Direct Cortical Recordings Suggest Temporal Order of Task-Evoked Responses in Human Dorsal Attention and Default Networks. *J. Neurosci. Res.* **38**, 10305–10313 (2018).
72. Tang, H. et al. Spatiotemporal dynamics underlying object completion in human ventral visual cortex. *Neuron* **83**, 736–748 (2014).
73. Schrouff, J. & Mourao-Miranda, J. in *International Workshop on Pattern Recognition in Neuroimaging (PRNI)* 1–4 (2018).

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**Author contributions**

J.P. and V.R. contributed to the design of experiments and data acquisition; J.S., O.R., J.M.M., and A.L.D developed analysis tools; J.S., O.R., S.S., V.R., and Z.H. performed data analyses and all authors contributed to writing of the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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