Anatomical correlates of face patches in macaque inferotemporal cortex

Michael J. Arcaro, Theodora Mautz, Vladimir K. Berezovski, and Margaret S. Livingstone

*Department of Psychology, University of Pennsylvania, Philadelphia, PA 19104; and *Department of Neurobiology, Harvard Medical School, Boston, MA 02115

Contributed by Margaret S. Livingstone, October 12, 2020 (sent for review September 10, 2020; reviewed by Jon H. Kaas, Pasko Rakic, and Xiaomin Yue)

Primate brains typically have regions within the ventral visual stream that are selectively responsive to faces. In macaques, these face patches are located in similar parts of inferotemporal cortex across individuals although correspondence with particular anatomical features has not been reported previously. Here, using high-resolution functional and anatomical imaging, we show that small “bumps,” or buried gyri, along the lower bank of the superior temporal sulcus are predictive of the location of face-selective regions. Recordings from implanted multielectrode arrays verified that these bumps contain face-selective neurons. These bumps were present in monkeys raised without seeing faces and that lack face patches, indicating that these anatomical landmarks are predictive of, but not sufficient for, the presence of face selectivity. These bumps are found across primate species that span taxonomy lines, indicating common evolutionary developmental mechanisms. The bumps emerge during fetal development in macaques, indicating that they arise from general developmental mechanisms that result in the regularity of cortical folding of the entire brain.

Significance

This study reports the discovery that buried gyral folds along the superior temporal sulcus in macaque lateral temporal cortex predict the locations of regions selectively responsive to faces. These findings demonstrate a link between cortical topology and the functional organization of higher-order visual cortex in macaques. We show that these anatomical landmarks are established prenatally and are likely the result of general developmental mechanisms such as molecular signaling gradients, topographic maps, and the formation of cortical areas. These folds are found across primate species. Identification of such anatomical landmarks may provide insight into evolutionary changes in the functional organization of high-level visual cortex.

Anatomical correlates of face patches in macaque inferotemporal cortex

Prior research has demonstrated a relationship between cortical folding and the functional organization of primary sensory areas (1–4). Within the visual system, not only does the calcarine sulcus serve as a macroanatomical landmark for primary visual area V1 (5, 6), but also the folding patterns within the sulcus are predictive of the retinotopic organization (7, 8). Given the complexity and variability in cortical size and shape across individuals, traditionally, it has been thought that there is little correspondence between cortical folding and visual areas beyond V1. However, research over the past decade has revealed a surprising degree of structure-function correspondence across the cortical surface (9–13).

Primates typically have several regions within the inferotemporal cortex (IT) that are selectively responsive to faces. Three spatially distinct sets of face patches—the posterior lateral (PL), middle lateral (ML), and anterior lateral (AL)—have been identified along the lower bank of the superior temporal sulcus (STS) in macaques (14, 15). Two other patches, the middle fundal (MF) and anterior fundal (AF), lie farther down the sulcus at the same anterior-to-posterior (AP) location as ML and AL, respectively. A sixth patch, the anterior middle (AM), is located on the ventral surface of anterior IT. These six face patches are located in similar parts of IT across individuals, although correspondence with particular anatomical features has not been observed previously.

Here, we performed high-resolution anatomical neuroimaging on 18 rhesus macaques. Seven monkeys were raised with normal visual experience of faces and developed face patches identified by functional magnetic resonance imaging (fMRI) within the first year of life. A topological analysis of each monkey’s cortical surface revealed that the face patches PL, ML, and AL were localized to focal convex convolutions along the STS, which we refer to as “bumps.” These convolutions are morphologically similar to the “buried” or “annectant” gyri previously described in human and monkey brains, but are probably too subtle to qualify as gyri. Neural recordings from five monkeys confirmed that each bump contained face-selective neurons and demonstrate that targeted recordings of face-selective neurons in the macaque brain can be achieved with high success without requiring fMRI. These bumps were morphologically similar in a separate group of monkeys raised with abnormal visual experiences of faces and that lacked face patches. Thus anatomical landmarks may predict the location of functional specializations but are not sufficient to indicate their presence. Using publicly available datasets, we identified these bumps in utero in rhesus macaques as well as postnatally in several other primate species. This suggests that bump formation emerges from general mechanisms ubiquitous across the primate order. These general mechanisms may underlie the organization of maps, areas, and patterns of architectonics, which in turn may influence functional specializations.

Results

Localization of Face Selectivity to STS Bumps. To probe the relationship between face selectivity and the anatomical topology of IT, we performed functional and anatomical MRI on seven macaque monkeys reared with normal (laboratory) visual experience. Functional scans were aligned to high-resolution (0.5 mm isotropic) T1 anatomical MR images. Regions preferentially responsive to images of faces compared to objects were identified along the lower bank of the STS in the right (Fig. 1) and left

Anatomical correlates of face patches in macaque inferotemporal cortex

www.pnas.org/cgi/doi/10.1073/pnas.2018780117

PNAS | December 22, 2020 | vol. 117 | no. 51 | 32667–32678
(SI Appendix, Fig. S1) hemispheres of each monkey. Face-selective responses covaried with the topology of the STS. The lower bank of the STS is not a uniformly smooth sulcus; rather, there are several focal convolutions along the posterior-to-anterior axis, which we refer to as bumps, where the cortical surface bulges. These bumps are most apparent in parasagittal sections of the STS. When viewed coronally, the bumps are less prominent relative to the broader convexity of the temporal gyrus. These bumps were visible on both high-resolution T1 images and lower-resolution functional echo-planar imaging (EPI) images (SI Appendix, Figs. S2 and S3). Across monkeys, face selectivity consistently fell on three bumps along the posterior-to-anterior extent of the STS.

Fig. 1. Anatomical localization of face selectivity in right-hemisphere STS. Preferential activity for faces vs. objects was identified along the lower bank of the STS in all seven monkeys reared with normal face experience ($P < 0.0001$, FDR-corrected). (Top) Seven sequential sagittal slices from medial (Left) to lateral (Right) cortex at 1-mm spacing in the right hemisphere of monkey 1. (Middle) Single sagittal slices showing localization of face-selective activity to bumps in the right hemispheres of the other normally reared monkeys. (Bottom) Group-average face selectivity falls on anatomical bumps in the STS. Seven sagittal slices from medial (Left) to lateral (Right) cortex at 1-mm spacing in the right hemisphere of the NMT brain. Group-average faces vs. objects contrast map threshold to show only voxels where at least three monkeys showed significant activity. See SI Appendix, Fig. S1, for left-hemisphere counterpart.
which we refer to as the posterior (Fig. 1, green line), middle (pink line), and anterior (blue line) bumps. More broadly, there was no consistent relationship between surface curvature and the magnitude of the response to faces compared to object contrast along the entire STS ($t_{[13]} = 2.0268, P = 0.0637$). The lack of a positive relationship is likely due to the presence of additional bumps along the STS that are not face selective as well as the prominent mediolateral convexity of the temporal gyrus. Similarly, neither cortical thickness nor sulcal depth were significantly related to face selectivity within the STS ($t_{[13]} < 0.78, ps > 0.45$). Together, this indicates that face selectivity was specific to these three anatomical landmarks and does not reflect a broader relationship between convex cortical folding and face selectivity.

The extent of each bump was identified manually based on cortical curvature (Fig. 2 and SI Appendix, Fig. S4). Convexity (curvature) maps were derived from cortical surface reconstructions of the segmented gray matter of T1 images for each monkey. Along the lateral bank of the STS, each bump was constrained laterally by the gyral crown and medially by the fundus. Along the AP axis, each bump was constrained to all adjacent cortex surrounding the peak convexity (highest point of bump; red color in the convexity map) terminating in the local

![Fig. 2. Identification of STS bumps in each monkey. (A) Convexity maps are shown along the lower bank of the STS in seven sagittal slices from medial (Left) to lateral (Right) (1.5-mm spacing) for the right hemisphere of monkey 2. Enlarged views of single sagittal slices show the bump peaks (red) and troughs (blue). (B) Smoothed white matter and inflated cortical-surface reconstructions showing the convexity maps of the STS in the right hemisphere of the same monkey. Posterior (green), middle (pink), and anterior (blue) bumps are shown in both hemispheres. (C) Group-average convexity maps with outlines of the three bumps are shown on the right and left hemisphere surfaces of the NMT brain.](image-url)
minimum (troughs; blue color in the convexity map) of the convexity map (Fig. 2 and SI Appendix, Fig. S4)). Local troughs surrounding the bumps were clearest within the sulcus of the STS just lateral to the gyral crown. Points near and along the gyral crown had uniformly large positive convexities (red color in the convexity map; Fig. 2 and SI Appendix, Fig. S4). In these regions, the boundaries of each bump were identified based on the troughs within the sulcus and the relative low points of the gyral convexity. Across monkeys, the median surface areas for the posterior, middle, and anterior bumps were 83.19 mm² (+/− 5.37 SEM), 75.66 mm² (+/− 6.81 SEM), and 60.23 mm² (+/− 3.32 SEM). The posterior and middle bumps bordered each other in every hemisphere, but there was a gap between the middle and anterior bumps. The AP length of this gap varied across monkeys and always lacked face selectivity (Fig. 1 and SI Appendix, Fig. S1). Additional bumps were identified within the STS. In each monkey, one or two bumps were present in the posterior-most part of the STS just anterior to the lunate sulcus where the STS arches superiorly toward the parietal cortex. In a few hemispheres, another small bump was also present between the middle and anterior bumps (e.g., see SI Appendix, Fig. S1, monkey 2 [M2]). These additional bumps were not face selective and were not evaluated further.

Along the posterior-anterior axis, the bumps were localized to similar regions of the STS relative to other anatomical folds. The posterior bump (SI Appendix, Fig. S4, green outlined region) was localized to where the inferior occipital sulcus (IOS) intersects the STS and just posterior to the posterior middle temporal sulcus (PMTS). The middle bump (pink outlined region) was identified anterior to the posterior bump directly medial and dorsal to the PMTS. The anterior bump (blue outlined region) was found near the anterior tip of the STS directly medial and dorsal to the anterior middle temporal sulcus (AMTS). Across monkeys, the middle bump’s surface area tended to scale with the surface area of PMTS (r = 0.64, P < 0.001), even when controlling for total surface area in each hemisphere (partial correlation: r = 0.61, P < 0.001), suggesting a relationship between local morphology of the STS and neighboring sulci. However, such a relationship was unclear for the other two bump-sulcal pairs (rs < 0.28, ps > 0.15).

Face selectivity was strongest and most consistent within the three group-average bumps. Across the cortical surface, the magnitude of face selectivity was strongest in the middle and anterior bumps (Fig. 3, Top). Furthermore, face selectivity was most consistently found across monkeys within these bumps as compared to the rest of the STS (and brain for that matter; Fig. 3, Bottom). In the right hemisphere, the only regions to show significant face selectivity in each monkey were found along the lateral half of the middle and anterior bumps. In the left hemisphere, the most consistent face selectivity (six of seven monkeys) was found along the lateral side of the middle bump. In general, face selectivity was most consistently found on the lateral half of each bump. Together, these data illustrate a tight correspondence between face selectivity and STS bumps and demonstrate that the most likely cortical location to find face-selective neurons is in the lateral half of the middle and anterior bumps.

Each STS bump corresponded to an individual lateral face patch and was aligned along the AP dimension with fundal face patches. Functional activations were mapped to each monkey’s cortical surfaces (SI Appendix, Fig. S5), and regions of face selectivity corresponding to previously reported PL, ML, and AL face-selective patches were identified (14, 16, 17). Each face patch was identified in both hemispheres of all monkeys except for AL in the left hemisphere of monkey 2 and PL in the right hemisphere of monkey 4 (SI Appendix, Fig. S5). The median surface areas for PL, ML, and AL were 18.78 mm² (+/− 3.07 SEM), 48.25 mm² (+/− 7.81 SEM), and 33.33 mm² (+/− 6.79 SEM). Face-selective regions PL, ML, and AL were localized to the STS bumps. In both hemispheres of each monkey and in the group average data, PL consistently fell on the posterior bump, ML on the middle bump, and AL on the anterior bump (SI Appendix, Fig. S6). Overall spatial correspondence between face regions and STS bumps was quantified by a Dice overlap index that compares the overlap of two regions to their total area and results in values ranging between 0 (no overlap) to 1 (perfect overlap) (Methods – Overlap analysis). For each face patch, a Dice index was calculated with each of the bumps. For PL, ML, and AL, the largest overlap was with the posterior bump (0.31), the middle bump (0.58), and the anterior bump (0.41), respectively (SI Appendix, Fig. S6). Although surface area and Dice overlap measures are dependent on the face-localizer threshold, it is clear from these data that 1) face patches were smaller than the bumps, and 2) each face patch covered only a portion of the corresponding bump. Although face patches extended laterally in a few hemispheres across monkeys, each face patch was most consistently localized to the lateral half of the corresponding bump (Fig. 3). Although not previously noted, the localization of PL, ML, and AL face patches to the lateral portions of these bumps is apparent in several prior studies (SI Appendix, Fig. S7). Together, these data emphasize a correspondence between each lateral face patch and an individual bump in the STS.

How does knowing the location of these bumps compare with alternative approaches for localizing face patches? When functional mapping is not an option, the alternative gold standard for localizing a functional region is to use a probabilistic atlas where the predicted location of an area is calculated from a separate group of individuals. For each monkey, we created probabilistic atlases of each face patch in both hemispheres using the remaining six monkeys. Consistent with prior work (18), the probabilistic map for each face patch fell within a focal region of the STS. However, there was variability in the spatial overlap of any two monkeys’ face patches. For example, when projected onto the National Institute of Mental Health macaque template (NMT) brain, the entire right ML face patch of monkey 5 fell anterior to the ML face patch of monkey 4. This spatial offset along the AP axis mirrored monkey 5’s middle bump falling anterior to monkey 4’s middle bump on the NMT brain.

![Fig. 3.](#) Localization of face selectivity to STS bumps. (Top) Outlines of the three group-average bumps are shown overlaid on group-average face vs. object activations. The group-average map thresholds were set to show only surface nodes where bumps or face selectivity were present in at least three monkeys. Across monkeys, the most face-selective regions along the lower bank of the STS fell within the middle and anterior bumps. (Bottom) Outlines of the three group-average bumps are shown overlaid on an overlap map that shows how many monkeys had significant face-selective activity in a given cortical region. Throughout the STS, overlap was strongest in the middle and anterior bumps.
At a finer scale, variability in where ML fell on the middle bump was apparent when looking at face selectivity on the native EPI images of individual monkeys (SI Appendix, Figs. S2 and S3). ML falls on the anterior part of monkey 5’s middle bump, but on the posterior part of monkey 4’s middle bump. Furthermore, few (or no) parts of each probabilistic face patch had 100% overlap across monkeys. On average, the overlap between any individual monkey’s face patch and the corresponding probabilistic atlas (SI Appendix, Figs. S8; PL, ML, and AL Dice indices = 0.30, 0.46, 0.35, respectively) was worse than the overlap of individual bumps and face patches (SI Appendix, Fig. S6). Thus, while probabilistic atlases are useful, similar to the bumps, such approaches are also imperfect at predicting the location of face regions in an individual subject. To directly compare the predictability of bumps with functional atlases, we evaluated the distances between the centroids of each face patch and its corresponding bump/probabilistic map. Although each face patch at least partially overlapped the corresponding bump, a centroid analysis provides additional information on how well aligned these regions are. The centroids of face patches were clustered near the centroid of corresponding bumps, particularly along the AP axis, although most face patch centroids fell closer in the medial-to-lateral axis to the gyral crown of the STS’s lower bank than the bump centroids (Fig. 4, Top). Consistent with the Dice overlap analysis, the distance along the cortical surface was shortest between each face-patch centroid and the centroid of the corresponding bump (PL and posterior bump = 4.28 mm; ML and the middle bump = 2.72 mm; AL and the anterior bump = 5.42 mm). Furthermore, for each face patch, the variance in centroid distances across monkeys was smallest for the corresponding bumps. This indicates that each bump was not just the closest anatomical landmark for its corresponding face patch but also the most predictive of the face patch’s location. Across monkeys, the overlap was highest and the centroid distances

![Fig. 4. Distances between face regions and STS bumps.](image-url)
were shortest between the ML face patch and middle STS bump, indicating that this pair had the strongest structure-function correspondence of the three lateral face regions. The distances between the posterior bump and PL tended to be smaller than the distances between the probabilistic PL and individual monkey's PL (t [12] = −2.93, P = 0.0127). Similarly, the distances between the middle bump and ML tended to be smaller than the distances between the probabilistic ML and individual monkey's ML (t [13] = −3.16, P = 0.0076). There were minimal differences in these distances for AL (t [13] = 0.43, P = 0.68). Overall, this suggests that identifying the location of these bumps is comparable to, and in most cases better than, probabilistic atlases.

**Using Bumps to Target Neural Recordings.** Each bump contained face-selective neurons. We performed computed tomography (CT) imaging to anatomically target multielectrode array implantation along the lower bank of the STS in five monkeys. Each array was successfully implanted within one of the STS bumps. In each monkey, the vast majority of channels were selectively responsive to faces vs. hands, bodies, and inanimate objects (Fig. 5 and SI Appendix, Fig. S9). Although we used fMRI in each monkey to verify that face selectivity fell on bumps, we used the bumps as anatomical markers during surgery for our array implantation. Given that the centroids of each face patch tended to fall on the lateral extent of each bump (Fig. 4 and SI Appendix, Fig. S6), we targeted the lateral half of each bump just below the gyral crown of the lower bank of the STS for implantation. The long dimension of each PL and ML array was oriented posterior-to-anterior along the STS and covered the majority of each bump. Furthermore, all arrays targeting face-selective regions have been within bumps, and we have yet to implant an array into a bump and not find the majority of channels to be face selective. Acute recordings from single-electrode penetrations should also benefit from targeting the bumps as a chamber centered on a bump will guarantee coverage of the corresponding face patch, although multiple penetrations may be required. Together, this indicates that the bumps are sufficient for targeting recordings from face-selective neurons.

**Anatomical Landmarks Are Not Sufficient for Face Selectivity.** Although the location of face patches can be predicted by the STS bumps, this anatomical feature of the STS does not necessarily indicate the presence of face selectivity. We collected high-resolution T1 anatomical images in seven monkeys with abnormal early visual experience of faces. Four of these monkeys were raised without seeing faces for the first year of life and did not develop face patches (three of these monkeys were originally reported in ref. 19). Two other monkeys were raised under conditions of binocular visual form deprivation (not specific to faces) for the first year of life and also did not develop face patches. The seventh monkey was raised in an environment where he had excess exposure of faces to his peripheral visual field during early development (in contrast to the typical foveally biased visual experience of faces). Although this monkey saw faces and developed face patches, his experience was atypical and serves as a test case for whether the type of face experience affects bump anatomy. In each monkey, the posterior, middle, and anterior bumps were present despite abnormal early face experience.

![Fig. 5. Multielectrode recordings from STS bumps. Multielectrode arrays were implanted in the posterior, middle, and anterior bumps of three macaque monkeys. In each array, the majority of channels were selectively responsive to human and monkey faces vs. hands, bodies, and inanimate objects. In each brain, the array base and wires from CT imaging are in blue. See SI Appendix, Fig. S9, for additional monkeys/arrays.](image-url)
experience (SI Appendix, Figs. S10 and S11). Relative to nearby anatomical landmarks such as the IOS, PMTS, and AMTS, the bumps were in similar parts of the STS as compared to the seven control monkeys reared with normal face experience. Furthermore, the group average bumps from this abnormal monkey group were in good spatial correspondence with the group average bumps from control monkeys (SI Appendix, Fig. S12). There was a high degree of Dice overlap in the extent of the posterior (0.81), middle (0.73), and anterior (0.77) bumps between the two monkey groups. The distances between the centroids of the posterior (2.42 mm +/- 0.44), middle (2.17 mm +/- 0.37), and anterior (2.01 mm +/- 0.56) bumps in each monkey with abnormal early visual experience and the mean centroids in the control group were, on average, within a few millimeters of each other. The distances between centroids of each bump in the abnormal early visual experience group and centroids of the probabilistic face patches were comparable to the centroid distances between bumps and face patches in normally reared individuals (SI Appendix, Fig. S12, Right), demonstrating that these bumps are useful for targeting regions of the STS where face patches normally develop such as in baby monkeys younger than 6 mo (20) and in monkeys raised without seeing faces (19). The Dice overlap and centroid distances were comparable between the following three subgroups of abnormal monkeys: 1) monkeys raised without seeing faces, 2) monkeys raised with a binocular visual form deprivation, and 3) monkeys raised with abnormal constant exposure to faces across the visual field (SI Appendix, Fig. S13). The only anatomical difference found between the control and abnormal monkey groups was cortical thickness. The average cortical thickness was greater in control monkeys for the posterior (1.97 +/- 0.03 vs. 1.87 +/- 0.05), middle (2.15 +/- 0.03 vs. 1.97 +/- 0.05), and anterior (2.10 +/- 0.02 vs. 1.95 +/- 0.04) bumps. These small differences were likely due to the abnormal monkey group being younger on average (1.43 +/- 0.24 y old) than the control monkey group (3.51 +/- 1.5 y old). Together, these data suggest that the STS bumps are present in individuals that lack face selectivity and that their macroanatomical organization is similar to that found in monkeys that have face patches. 

Developmental and Evolutionary Origins of STS Bumps. STS bumps appear prenatally and are evolutionarily preserved across pri- mate species. Although the STS bumps are not clearly formed by gestation day (GD) 110 the sulcus has already begun to bend near where bumps emerge (Fig. 64; black dashed circle). By GD 135, the posterior, middle, and anterior bumps are all present (Fig. 64; green, pink, and blue arrows); this is after the cortical plate has formed and the geniculate has innervated visual cortex (21). Thus, the bumps emerge prior to seeing faces and well before the development of face patches. The presence of these bumps in utero indicates that they form from general principles of cortical folding and cortical expansion. Do these bumps manifest in other primate species? Despite the cortical surface in New World Monkeys being relatively smooth, there are asymmetrical folds and gyral regions hypothesized to differ in their myeloarchitecture (28) and areal connectivity (29) and, by extension, their functional properties (30). The identification of these structure-function corre- spondences in multiple species provides insight for testing the functional correlates of cortical folding. 

Our results add to a growing body of literature characterizing the morphological features of cortical folding including sulcal pits and annectant gyri (10, 22, 31, 32). Although less studied in Old World monkeys, such features are associated with functional specializations in humans (31, 33–35). Of particular note, several plis de passage (annectant gyri) in the human STS are thought to reflect distinct structural connectivity bridging superior and middle temporal gyri and functional specializations (35–37). Similar to these plis de passages, the macaque STS bumps 1) are convolutions buried in the main furrow of the STS, 2) are distributed along the posterior-to-anterior axis, 3) are consistent in their location across individuals, and 4) emerge early in develop- ment. However, there are notable differences. Superficial human plis de passages typically span both sulcal banks and form a clear interruption of the sulcus. In contrast, the middle and anterior macaque STS bumps extend into the fundus but are not contiguous with convolutions along the upper bank of the STS in most hemispheres. This discontinuity may be similar to “deep” plis de passages reported recently in the human STS that are partially disconnected in the fundus (36). A continuous bump extending to the upper bank was apparent only for the middle
Fig. 6. Bump phylogeny and ontogeny. (A) The bump appears early in development. The bumps are not clearly identifiable at GD 110 but are present by GD 135. Green, pink, and blue arrows indicate posterior, middle, and anterior regions of the STS with bumps. MRI images are part of the Oregon National Primate Research Center Fetal Macaque Brain Atlas. (B) Bumps along the STS are present in several primate species spanning New World Monkeys, Old World Monkeys, gibbons, and baboons. The gorilla STS has bumps similar to macaques, but all other Great Apes have a more complicated folding structure with several bumps along the STS. All MRI images except for the gibbons come from T2W scans. Gibbons are from T1 scans. Dashed circle shows the STS in Great Apes. Gibbon MRI data are taken from the National Chimpanzee Brain Database (https://braincatalogue.org/). Human MRI data are taken from the T1 group average from the HCP 1200 Young Adult dataset (https://www.humanconnectome.org/study/hcp-young-adult). All other MRI data are taken from the Brain Catalogue (https://braincatalogue.org/).
bump and only in 3 of 14 hemispheres in our control group. Although anatomical connections exist between face patches along the lower and upper banks of the STS (38), the correspondence of these tracts to the convolutions of the STS remain to be resolved. Instead of being homologous to the human STS, the macaque bumps may correspond to convolutions along the fusiform gyrus where putatively homologous face patches exist. Although annexant gyri have not been documented for human face patches along the fusiform, the morphology of a plis de passage in the visual word form area along the fusiform has been shown to be correlated with reading skills (39).

What is the possible developmental relationship between the three bumps distributed along the STS and face patches? A consistent structure-function relationship could be taken as evidence that the location where face patches develop in IT is predetermined and that the bumps represent circuitry genetically specialized for processing faces. However, morphologically similar bumps were apparent in monkeys that lacked face patches, demonstrating that STS bumps are not sufficient to produce face selectivity in the absence of face experience. Furthermore, these bumps are present in utero prior to the onset of vision, and face patches do not appear until 200 d of age (20). The prenatal formation of these bumps and the consistency in their anatomical location across individuals are doubtless the result of the same general developmental mechanisms that result in the regularity of cortical folding of the entire brain. These factors must include molecular signaling gradients that can indirectly influence folding by limiting the expandability, stiffness, or thickness of the cortical surface (40) and mechanical pressures such as axonal tension (29) or differential growth rates of superficial and deep cortical layers (41–43). In particular, regionally specific growth applied to a species-specific initial geometry may suffice for producing species-specific folds (44). The presence of these bumps in late stages of gestation prior to visual experience is consistent with recent observations that gyri form only after the completion of neurogenesis and are driven by the rapid growth of intracortical neuropil and the enlargement of subjacent white matter consisting of cortico-cortical connections (32). Together, our data provide additional support for the protomap hypothesis (33), which postulates that intrinsic patterning mechanisms establish the initial arealization of cerebral cortex. Given that sulcal and gyral regions are hypothesized to differ in their laminar architecture (28) and areal connectivity (29), these bumps comprise part of the early cortical architecture that constrains and guides postnatal development.

The precise location where each face patch fell on a given bump varied across individuals. Face patches typically fell on the lateral half of the bumps although face-selective activity was found medial to the lateral patches (Fig. 2) and likely corresponds to the MF and AF face patches. More notably, face patches varied in their localization along the AP axis of the bumps. This variability was evident in the EPI images (SI Appendix, Figs. S2 and S3) and thus cannot be attributed to misalignment with the high-resolution anatomical images. It is also unlikely that any differences in vasculature across individuals could account for such variability since the peak of face-selective fMRI activity corresponds to the highest concentration of face-selective neurons (16, 45). Instead, this likely reflects true variability in the precise anatomical location of face patches on bumps across individuals. Given that face-patch development depends on face experience (19) and adheres to a retinotopic proto-organization across IT (46), such local variability may simply reflect idiosyncrasies of each monkey’s visual experience with faces and how it mapped onto an intrinsic functional architecture. Indeed, a recent study in humans showed a correspondence between individual face-viewing behavior and the receptive field properties of face-selective regions (47). It remains to be seen whether macaque STS bumps undergo morphological changes across postnatal development and if such changes have any relationship to functional maturation. Thus, it is plausible to have a consistent structure-function correspondence across individuals where the presence and precise localization of function is dependent on how experience interacts with an intrinsic architecture.

Is there a mechanistic explanation for the association between face patches and bumps? Like Gaul, inferotemporal cortex is...
monkeys (M1 to M8) participated in both anatomical and functional neuroimaging experiments. Monkey 1 and the other four monkeys (M8–M11, M13, M17) were raised with abnormal visual experience of faces. As previously reported (57), functional imaging, monkeys were alert, and their heads were immobilized using a foam-padded helmet with a chinstrap that delivered juice. The monkeys were scanned in a primate chair that allowed them to move their bodies and limbs freely, but their heads were restrained in a forward-looking position by the padded helmet. The monkeys were rewarded with juice for maintaining a central fixation within a 2° window. Gaze direction was monitored using an infrared eye tracker (SCAN).

Electrophysiology Display. We used MonkeyLogic to control experimental workflow (https://monkeylogic.nimh.nih.gov). Visual stimuli were displayed on a 19-inch Dell LCD screen monitor with a 60-Hz refresh rate and a 4:3 aspect ratio positioned 54 cm in front of each monkey.

Materials and Methods

Monkeys. Functional and anatomical MRI studies were carried out on 24 Macaca mulatta, 5 female and 19 male. All procedures were approved by the Harvard Medical School Animal Care and Use Committee and conformed with NIH guidelines for the humane care and use of laboratory animals. Seventeen monkeys (M12–M17) were raised with abnormal visual experience of faces. As previously reported (57), the other four monkeys (M8–M11, M13, M17) were hand reared by humans for the first several months and then cohabited with other juveniles, also in a room with other monkeys. Seven of these monkeys (M12–M17) were raised by laboratory staff wearing welders’ masks that prevented the monkey from seeing the staff member’s face. The only visual experience that they had with faces of any kind was by indirect viewing experiences (i.e., reflected in a mirror) 1–2 h per week, with the face exposure being a minor fraction of that time. The other two hand-reared monkeys (M14 and M15) were raised under conditions of binocular-visual-form deprivation via eye lid suturing for the first year. Finally, one monkey (M18) was raised with his mother in an environment with large posters of faces on the walls such that he had an abnormal, constant exposure to faces in his peripheral visual field. As previously reported (57), functional imaging, monkeys were alert, and their heads were immobilized using a foam-padded helmet with a chinstrap that delivered juice. The monkeys were scanned in a primate chair that allowed them to move their bodies and limbs freely, but their heads were restrained in a forward-looking position by the padded helmet. The monkeys were rewarded with juice for maintaining a central fixation within a 2° window. Gaze direction was monitored using an infrared eye tracker (SCAN).

Electrode Implantation. Multielectrode arrays were implanted within the STS of five male Macaca mulatta. Each array was implanted to target face-selective patches. In monkey 1, a floating microelectrode array (FMA) (32-channel Microprobes FMA; https://microprobes.com/products/multichannel-arrays) was implanted within the middle of the STS corresponding to his PL face patch. The internal dimensions of the FMA array were 3.5 × 1.5 mm with a spacing of 370 to 400 μm between electrodes (across the width and length of the array, respectively). In monkey 8, two FMAs were implanted in the left hemisphere—one within the anterior part of the posterior bump corresponding to his PL face patch and a second array within the anterior part of the middle bump corresponding to his ML face patch. In monkey 9, one FMA was implanted in the left hemisphere centrally within the middle bump corresponding to his ML face patch. In monkey 10, one FMA was implanted within the anterior part of the middle bump corresponding to his ML patch. In monkey 11, a 64-channel 12.5-μm NiCr microwire array (58) was implanted centrally within the anterior bump corresponding to his AL face patch. Monkeys were trained to perform a fixation task. Neural recordings were performed on a 64-channel Plexon Omniplex Acquisition System.

Anatomical Imaging. As previously reported (57), a whole-brain structural volume was acquired while the animals were anesthetized with a combination of Ketamine (4 mg/kg) and Dexdomitor (0.02 mg/kg). Scans were acquired in each monkey using a 3 T Siemens Skyra, using a 15-channel transmit/receive knee coil. Monkeys were scanned using a magnetization-prepared rapid gradient echo (MPRAGE) sequence; 0.5 × 0.5 × 0.5 resolution; field of view = 128 mm; 256 × 256 matrix; repetition time (TR) = 2,700 ms; echo time (TE) = 3.35 ms; inversion time (TI) = 859 ms; flip angle = 9°. Three whole-brain T1-weighted anatomical images were collected from each animal.

Reconstruction of cortical surfaces. Each animal’s T1 images were coregistered to derive an average anatomical volume image for each monkey. Each monkey’s average anatomical volume underwent semi-automated cortical surface reconstruction using FreeSurfer. To ensure high accuracy, skull stripping and white-matter masks were first manually segmented by an expert slice-by-slice along coronal, axial, and sagittal planes and then passed into FreeSurfer’s autorecon pipeline. Pial and white-matter surfaces were inspected to ensure accurate segmentation. If poor segmentations were detected, the white-matter mask and control points were edited, and surface reconstruction was rerun until corrected. For several monkeys, FreeSurfer’s autosegmentation had trouble with the calcaneus and highly vascularized regions such as the insula. To fix these segmentation errors, average anatomical volumes were manually edited to improve the gray/white-matter contrasts and remove surrounding nonbrain structures (e.g., sinuses, arachnoid, and dura mater).

Generation of anatomical feature maps. For each monkey, FreeSurfer’s automated algorithm was used to obtain sulcal depth and convexity maps for the pial and white-matter surfaces. Sulcal depth (in mm) is measured as the distance between the inflated surface and pial surface (59) at each vertex. Convexity maps along the pial and white-matter surfaces were obtained using Analysis of Functional Neuroimaging (AFNI)/SUMA’s automated algorithm (part of SurfaceMetrics). The pial and smooth white-matter convexity maps were averaged to produce a mean convexity map.
Stimuli. Visual stimuli were projected onto a screen at the end of the scanner bore. Static images. Responses to image categories of faces and inanimate objects were probed as previously reported (57). Each scan was composed of blocks of each image category; each image subtended 20° × 20° of the visual angle and was presented for 0.5 s; block length was 20 s, with 20 s of a neutral gray screen between image blocks. Blocks and images were presented in a counterbalanced order. All images were centered on a pink noise background. All images were equated for spatial frequency and luminance using the SHINE toolbox (60).

Functional Imaging. As previously reported (57), monkeys were scanned in a 3-T TimTrio scanner with an AC88 gradient insert using four-channel surface coils (custom made by Azma Maryam at the Martinos Imaging Center, Charlestown, MA). Each scan session consisted of 10 or more functional scans. We used a TR of 2 s, TE of 13 ms, flip angle of 72°, integrated parallel acquisition techniques = 2, 1-mm isotropic voxels, matrix size 96 × 96 mm, and 67 contiguous sagittal slices. To enhance contrast (61, 62), we injected 12 mg/kg monocrystalline iron oxide nanoparticles (Feraheme, AMAG Pharmaceuticals) in the saphenous vein just before scanning.

General fMRI Preprocessing. As previously reported (57), functional scan data were analyzed using AFNI [Research Resource Identifier (RRID): nif-0000-00304] (63, 64), Freesurfer (RRID:nif-0000-00304) (65, 66), JIP Analysis Toolkit (67), and MATLAB (Mathworks, RRID:nlx_153890). Each scan session for each monkey was analyzed separately. Using AFNI, all images from each scan session were aligned to a single time point for that session, detrended, and motion corrected. Data were spatially filtered using a Gaussian filter of 2 mm full width at half-maximum to increase the signal-to-noise ratio while preserving spatial specificity. Each scan was normalized to its mean. Data were registered using a two-step linear and then a nonlinear alignment approach (JIP Analysis Toolkit) to a high-resolution (0.5 mm) anatomical image for each monkey. First, a 12-parameter linear registration was performed between the mean EPI image for a given session and a high-resolution anatomical image. Next, a nonlinear, diffeomorphic registration was conducted. To improve registration accuracy of the ventral cortex, we manually drew masks that excluded the cerebellum for both EPI and anatomical volumes prior to registration.

fmRI Stimulus Category Analysis. As previously reported (57), a multiple regression analysis [AFNI’s 3DDeconvolve (63)] in the framework of a general linear model was performed on the category experiments for each monkey separately. Each stimulus condition was modeled with a MION-based hemodynamic response function (61). Additional regressors that accounted for variance due to baseline shifts between time series, linear drifts, and head-motion-parameter estimates were also included in the regression model. Due to the time-course normalization, beta coefficients were scaled to reflect the percentage of signal change. Since MION inverts the signal, the sign of beta values were inverted to follow normal fMRI conventions of increased activity being represented by positive values. Brain regions that responded more strongly to monkey faces than familiar objects were identified by contrasting presentation blocks of each of these image categories. Maps of beta coefficients were clustered (>10 adjacent voxels), and the threshold was at $P < 0.0001$ (false discovery rate [FDR]-corrected).

Lateral Face-Patch Identification. Functional activations were mapped to each monkey’s cortical surfaces (SI Appendix, Fig. 55), and regions of face selectivity along the lower bank of the STS corresponding to previously reported PL, ML, and AL face-selective patches were identified (14, 16, 17). In some monkeys, face-selective activity extended medially into what were likely the fundal face patches (e.g., ML and MF in monkey 7, SI Appendix, Fig. 55). For these cases, the border between adjacent face patches was identified as a trough in the gradient of face selectivity.

Anatomical Analyses. Surface area (in square millimeters) was estimated along the pial surface using AFNI’s SurfMeasures. Distances between surface nodes were calculated along the cortical surface using AFNI’s SurfDist.
