ABSTRACT
To study the circuitry related to the ventral stream of visual information processing and its relation to the cytochrome oxidase (CytOx) modules in visual area V2, we injected anterograde and retrograde cholera toxin subunit B (CTb) tracer into nine sites in area V4 in five Cebus apella monkeys. The injection site locations ranged from 2° to 10° eccentricity in the lower visual field representation of V4. Alternate cortical sections, cut tangentially to the pial surface or in the coronal plane, were stained for CTb immunocytochemistry or for CytOx histochemistry or for Nissl. Our results indicate that the V4-projecting cells and terminal-like labeling were located in interstripes and thin CytOx-rich stripes and avoided the CytOx-rich thick stripes in V2. The feedforward projecting cell bodies in V2 were primarily located in the supragranular layers and sparsely located in the infragranular layers, whereas the feedback projections (i.e., the terminal-like labels) were located in the supra- and infragranular layers. V4 injections of CTb resulted in labeling of the thin stripes and interstripes of V2 and provided an efficient method of distinguishing the V2 modules that were related to the ventral stream from the CytOx-rich thick stripes, related to the dorsal stream. In V2, there was a significant heterogeneity in the distribution of projections: feedforward projections were located in CytOx-rich thin stripes and in the CytOx-poor interstripes, whereas the feedback projections were more abundant in the thin stripes than in the interstripes. J. Comp. Neurol. 522:3091–3105, 2014.

INDEXING TERMS: primates; extrastriate cortex; cytochrome oxidase modules
thick/pale stripes that avoided thin stripes. Cells in the thick stripes receive inputs from layer IVB of V1 and respond to depth cues and to moving stimuli. Lu et al. (2010) found robust direction maps, occupying several restricted regions of V4 and tend to overlap with orientation- and color-prefering domains. Single-cell recordings targeting these direction-prefering domains also showed a clustering, as well as a columnar, organization of V4 direction-selective neurons. These data suggest that, in contrast to the classical view, motion information is also processed in ventral pathway regions such as area V4.

The laminar distributions of labeled cells and terminals have been used as criteria for distinguishing feedforward projections, which originate from cells in the superficial layers and mostly terminate in layer IV of a higher order target area, from feedback projections, which originate from cells in the infragranular layers and mostly terminate outside of layer IV of lower order target areas and have been used to build a hierarchical model of extrastriate cortex. However apparent, this laminar segregation of ascending and descending inputs is not absolute or can be complicated, in part due to the presence of translaminar intrinsic axons and dendrites (Spatz and Tigges, 1972; Rockland and Pandya, 1979; Maunsell and Van Essen, 1983; Weller and Kaas, 1985; Ungerleider and Desimone, 1986; Boussaoud et al., 1991; Felleman and Van Essen, 1991; Shipp et al., 2009; Markov et al., 2014). It becomes vital to fully understand feedforward and feedback connectivity between V4 and other areas such as V2 and how this might contribute to an understanding of the hierarchical structural organization of the visual cortex.

Over the past 30 years, we have been studying the visual system of the New World Cebus monkey as a model for the human visual system, replacing in many aspects the need for using Old World monkeys (Gattass et al., 1990). Cebus apella is a diurnal monkey with brain size and sulcal pattern similar to that of Macaca fascicularis. We propose that it serves as an adequate alternative model for study of the primate visual system. The aim of this study is to gather additional data about the modules of V2 and its connections with V4 in New World monkeys.

We have previously described the feedforward modular V2 projections to V4 in Cebus monkeys (Nascimento-Silva et al., 2003). Here, we were interested in determining how feedforward and feedback projections are distributed through the cortical layers and organized in relation to the CytOx modules of V2. We used cholera toxin subunit B (CTb), which has been used as a bidirectional tracer to visualize cell soma and axon terminals (Angelluci et al., 1996). We report on the arrangement of projections in the region of the representation of the central lower field of V4, which links V2 and V4 in C. apella, in relation to the CytOx-rich and -poor modules of V2 and reveal the laminar and tangential distributions of the feedforward and feedback projections of V2. The use of injections of CTb into V4 allowed us to specifically label the V2 modules and
provided an efficient method for elucidating the modular circuitry of V2. The distributions of projections were significantly heterogeneous; feedforward projections originated in the thin stripes and interstripes (as did the feedback projections from V4) but were richer in the thin stripes than in the interstripes.

MATERIALS AND METHODS

All experimental protocols were conducted following NIH guidelines for animal research and were approved by the Committee for Animal Care and Use of the Institute of Biophysics Carlos Chagas Filho, Federal University of Rio de Janeiro.

Five adult *C. apella* monkeys ranging in weight from 2.8 to 4.0 kg were used. These animals were also used for other anatomical projects. Each animal was anesthetized with ketamine hydrochloride (30 mg/kg IM Ketalar; Parke-Davis, São Paulo, SP, Brazil) and received atropine (0.15 mg/kg IM Atropina; Hypofarma, Rio de Janeiro, RJ, Brazil) and benzodiazepine (0.5 mg/kg IM Valium; Roche, Rio de Janeiro, RJ, Brazil). Anesthesia was maintained with infusions of sodium thiopental (10 mg/kg/h IV Tiopental; Cristália, São Paulo, SP, Brazil) in 5% glucose or with regularly repeated IM injections of a 1:5 mixture of 6% ketamine hydrochloride and 2% dihydrotiazine hydrochloride (Rompun, Bayer, Rio de Janeiro, RJ, Brazil). Level of expired CO2, rectal temperature, and heart rate were monitored throughout the experiment and maintained within physiological ranges. A craniotomy was made over the intended injection sites under sterile conditions, and the dura mater was sectioned to expose the cortical surface. Injections were made under visual guidance. Following the injections, the wound was closed in anatomical layers, and the animals received IM injections of penicillin G (300,000 IU Benzetacil; Eurofarma, São Paulo, SP, Brazil).

Volumes ranging from 0.2 to 1 μl of 1% CTb (Sigma-Aldrich, St. Louis, MO) were injected into the cortical surface of the prelunate gyrus under direct observation using a 1-μl 27-gauge needle Hamilton (Reno, NV) syringe (Fig. 1). As a reference, we used the superficial blood vessel that runs over the superior temporal gyrus from the inferior occipital sulcus to the superior temporal sulcus. Injections were made into one or both hemispheres in each animal for a total of seven hemispheres.

We inferred the eccentricity of the injections based on the location of the back projections of V2, using the visuotopic map described by Rosa et al. (1988). The distance from the fovea of V2 to the projection was confronted with the intended location in V4, using the integration of the magnification factor function for V2.

After of 7–21 days of survival, the animals received lethal doses of sodium pentobarbital (90 mg/kg IV) and were transcardially perfused. Four animals underwent a flattening procedure, and the remaining animal was perfused, fixed, and cut in the coronal plane.

The four animals were perfused with 0.9% saline followed by 0.1 M phosphate buffer at a pH of 7.4. To obtain flattened cortical preparations for the first four animals, the brains were removed from the skull, and blocks that included areas V1, V2, and V4 were separated from the rest of the brain and then flattened. To separate the block, we cut along the posterior tip of the superior temporal sulcus (st), and the cortex was separated from the white matter from the st tip to the lunate sulcus (lu) superior and inferior to the occipital sulcus (io) below, which included part of area V4. Following the occipital operculum, the cortex was dissected along the fundus of the lu and the fundus of the io at the striate cortex far from the V1/V2 border to the large belt that included the dorsal parts of V3, V2, and V1. Next, this tissue block was gently unfolded and interposed between glass slides, pressed, and submersed in a fixation solution. This preparation was fixed with a solution of paraformaldehyde (4%) in 0.1 M phosphate buffer for 12 hours and then stored overnight in phosphate buffer/glycerol (10%). Frozen sections (60 μm) were cut in a plane tangential to the pia mater. Alternate sections were designated for CTb immunohistochemistry (Angelucci et al., 1996) or CytOx enzyme histochemistry, which was performed according to the modified Silverman and Tootell (1987) method.

The remaining animal was perfused with 0.9% saline followed by 4% paraformaldehyde in phosphate-buffered saline (PBS), 4% paraformaldehyde PBS with 2.5% glycerol, PBS with 5% glycerol, and PBS with 10% glycerol overnight. The next day, the brain was frozen and sectioned into 40-μm-thick coronal sections using a cryostat. Adjacent sections were stained for cell bodies using cresyl violet, CO histochemistry, or immunocytochemistry for CTb.

For immunocytochemical reactions, the sections were incubated overnight in a polyclonal cholera toxin antibody (1:1,500–2,000; Sigma-Aldrich) in a solution containing 0.05% bovine albumin and 0.3% Triton X-100 in 0.01 M PBS at a pH of 7.4. Sections were then incubated for an additional hour in biotinylated anti-rabbit secondary antibody and then processed by the avidin–biotin method with ABC kits (Vector, Burlingame, CA) and nickel-enhanced dianimobenzidine. The sections were rinsed in PBS, mounted on gelatin-coated slides, dehydrated, and coverslipped. Control sections were prepared by omitting the primary antibody in the incubation solution. These sections showed no specific staining.

CTb sections were scanned for the distribution of the labeled cells and terminals by using a Leitz (Wetzlar,
Germany) Diaplane microscope interfaced to a Minnesota Datamatrix X-Y system (A) or a Zeiss (Thornwood, NY) microscope interfaced to a Neurolucida system (B; MBF Bioscience, Williston, VT). The labeled neurons were plotted onto drawings of the sections’ contours and photographed. Each section stained for CO was used for the reconstruction of the stripe pattern in V2. Labeled cells were projected onto the photographic reconstruction of the CO compartments of V2 to allow for analysis of the relationships of the labeled cells in V2 with the CO-modules using Adobe Photoshop (San Jose, CA). In the coronal sections, we used the adjacent Nissl sections to identify the layers in which the labeled cells and terminals were located. The contours of the sections, the layers, and the landmarks were drawn by using the Neurolucida system and then transferred to NeuroExplorer (Nex Technologies, Madison, AL) for data analysis.

We used Neurolucida to capture, align, and organize the sections used for the evaluation of cell and fiber density of all animals. The Neurolucida system was used to capture whole slide images from the entire sections (digital scans) and to make serial section reconstructions of V2 and surrounding areas. We used its tools to align the section and to obtain a final aligned serial section reconstruction for each animal. The serial section reconstructions of V2 were used to trace and analyze the CytOx bands and to segregate the cortical layers through multiple sections. After aligning the sections, we transferred the limits of the CytOx-rich label to the sections stained for CTb for quantification.

Quantification was done using all sections. Tangential sections were used for the analysis of width of CytOx band, and distribution of cells and fibers in these compartments. Oblique and coronal sections were used for both the analysis of the distribution in the compartments of V2 and also for distribution in the different layers.

We used coronal section series to draw the limits of the cortical layers by cytoarchitecture. In these series,
we determined the relative depth of these transitions of the cortical layers to use in other series stained for CytOx or CTb.

When 40-μm tangential sections are used, the depth of the sections is an indication of the cortical layer. Cutting from the pial surface to the white matter, section 1 corresponds to the top of layer I, whereas the last section (usually section 37) corresponds to the bottom of layer VI. When we had oblique sections identified by serial section reconstruction, we used these sections to study both the distribution of cells and fibers in the CytOx bands of V2 and the distribution in different cortical layers. For example, we selected oblique sections that were cut around layer IV at the V1/V2 border and would reach the pial surface (layer I) close to the anterior border of V2 to evaluate the distribution in both the CytOx band and the supragranular layers.

The cell counts of each compartment were determined by using NeuroLucida. We used a Matlab (MathWorks, Natick, MA) density program to measure the optical density of fibers from images in slabs of predetermined sizes. CTb-stained sections were cut in rectangles of predetermined sizes (200 × 50 μm). Matlab measures the average optical density of individualized pixels. Densely stained clusters of pixels larger than 20 μm in diameters were not computed. This procedure avoids the interference of cell bodies in fiber density measurements. The measurements made in oblique sections were divided into larger rectangles.

Data from each section from each animal were used to feed a database (summarized in Figs. 8–11). The data were transferred to a Systat 8 file (Systat Software, Chicago, IL) for statistical analysis.

RESULTS

We analyzed data from nine CTb injections into area V4 in five animals. We will first present the connections of V4 with the CytOx modules in the flattened preparations of V2. The laminar distributions of labeled cells and terminals are described as the feedforward and feedback V2 projections from area V4. By analyzing the distributions of labeled cells and terminals in the coro-nal sections, we illustrate and quantify the laminar distribution of the labeling. We then describe the cell types and spatial organization of cells and terminals.

Injections of CTb in V4

Figure 1 shows the CTb injection sites in V4 for each of the five cases. Case 1 received two injections of 0.5 μl CTb into the dorsal part of V4 ranging from 2° to 5° eccentricity. Case 2 received two injections of 0.5 μl CTb in the dorsal part of V4 (5°). Case 3 received two injections of 1.0 μl CTb in the right hemisphere and one injection of 0.2 μl CTb in the left hemisphere; both injections were in the dorsal part of V4. One injection into the right hemisphere was located in the representation of the peripheral visual field (10°), and the other was located in the representation of the central lower visual field in V4 (2°). Case 4 received an injection of 0.2 μl CTb in the representation of the central lower visual field in V4 (2°). In both cases (i.e., cases 3 and 4), we detected similar labeling in V2; however, in case 3 (injection into the representation of the peripheral visual field), the tracer spread into area TEO, and the injection into the representation of the central visual field in the opposite hemisphere invaded the white matter. Case 5 received one injection of 0.5 μl CTb in the dorsal part of V4 in the representation of the pericentral lower visual field at 5° eccentricity. Coronal sections of the occipital cortex in front of the V1/V2 border and behind the lunate sulcus revealed regular arrangements of labeled cells and terminals in different cortical layers (Fig. 1).

Projections in V2

After injections of CTb into V4, labeled cells and terminals were located in the stripes perpendicular to the V1/V2 border in V2. The projections from 2° to 5° were located in dorsal lateral V2 (Fig. 1), whereas those from 6° to 10° eccentricity were located in dorsal medial portions of V2 (not shown). In all four cases, the flattened preparations of V2 showed clusters of projections that were organized into bands that were orthogonal to the V1/V2 border (Figs. 2 and 3).

Figure 2 shows the injection sites in V4 and the projections in V2 in the flattened sections of two animals (cases 1 and 2). The injections were restricted to visual area V4 and produced regularly spaced projections to V2 in both animals. These projections are composed of groups of cells and fibers that were clustered along bands that were orthogonal to the V1/V2 border. Outside V2, in V3, only one cluster was found in case 1.

The CTb-labeled patches of cells represent the origins of the feedforward projections to V4, whereas the terminals constituted feedback projections from V4 to V2.

Projections and CytOx modularity

Figure 3 shows the CytOx bands and CTb-labeled cells and terminals in V2 from case 3. These data allowed comparison of the locations of the CytOx bands of V2 with the patches of projections from V4. One CytOx-stained oblique section of V2 (Fig. 3A), a filtered montage of CytOx staining in layer IV from a set of
neighboring sections (Fig. 3B), and an alternating neighboring CTb-stained section (Fig. 3C) show the correspondence of labeled cells and terminals with the CytOx-rich thin bands and the CytOx-poor interbands of V2. The montage (middle sections) was based on stacked segments of layer IV from neighboring CytOx-stained sections of V2. The collage (Fig. 4B) was filtered (Gaussian filter \( \alpha = 100 \, \mu m \) [30 pixels]) using Photoshop 7 to generate a pictorial representation of the bands of V2. The patches of CTb-stained cells and terminals in the upper layers of V2 were located in thin CytOx-rich modules and intermodules of V2. Lightly stained cells were clustered mainly in the upper layers, whereas dense fibers were clustered in the lower layers (Fig. 3C).

Figure 4 shows the clusters of labeled cells and terminals located in the supragranular layers II and III that corresponded to the CytOx-rich thin stripes and CytOx-poor interstripes in V2 after injections of CTb into V4 of case 3. No labeled cells were found in layers I and IV in V2 (Fig. 4). Most of the neurons were located in layer III, in the CytOx-rich thin stripes. The cells in this layer were larger than the labeled cells in other layers. The distribution in the infragranular layers was concentrated in layer VI. Strong labeling of terminals was found in the supragranular (layers I and II) and infragranular layers (bottom of layer V and layer VI) (Fig. 4).

Figure 5 shows CTb-stained cells and fibers from the thin CytOx-rich band and interbands in V2 of case 4. Low-power photomicrographs from sections of V2 stained for CTb (Fig. 5A, left) and CytOx (Fig. 5A right) show the correspondence of the location of the CytOx modules of V2 with the clusters of labeled cells and fibers. Figure 5B shows high-power photomicrographs obtained from subsequent tangential sections (10, 08, 06, and 02) with CTb-labeled cells and terminals. The overlay of these data shows that the CTb-labeled cells and terminals were related to one CytOx-rich thin stripe that was interposed by two CytOx-poor interstripes of V2 and extended approximately 2 mm, avoiding the thick stripes (Fig. 5A, right). Overall, the clusters of labeled cells appear to be distributed in the interbands and thin bands, although the terminal-like labeling appears to be restricted to interstripes. Thus, the clusters of cells that were in the thin bands and interbands avoided the domains of the thick CytOx-rich bands.

**Laminar distribution of labeled cells and terminals in V2**

Figure 6 shows the location of the CTb-labeled cells in the cortical layers of V2 in coronal sections from case 5. Nissl-stained (Fig. 6A) and neighboring CTb-immunoreacted sections (Fig. 6B and 6C) show that the labeled cells were mainly located in layer III and were sparsely located in layer V. In this figure, approximately 91% of the labeled cells were located in the supragranular layers, and 9% were located in the infragranular layers. The patches of labeled cells encompass approximately 600 \( \mu m \), and they were separated by roughly 2 mm (Fig. 6B).
Figure 7 shows the labeled cells and terminals in two enlarged bright- and darkfield photomicrographs of V2 after injections of CTb in V4 of case 1. The brightfield photomicrographs show the distribution of cells and fibers in different cortical layers of V2 (Fig. 7A). The darkfield photomicrograph emphasizes the distribution of labeled fibers in the infragranular layers of V2 (Fig. 7B). Figure 7B shows complex meshes of fibers in lower and upper layers and patchy vertical bundles of fibers at the thin CytOx-rich modules of V2. The cells predominantly occupied the supragranular layers, mainly layer III, whereas the terminals were located in the infragranular layers, particularly layer V.

The majority of labeled cells in area V2 were pyramidal cells with ascending vertical dendrites, and large stellate cells containing large nuclei composed a small population (Fig. 8A,D). These cells were densely organized in clusters in the thin stripes and sparsely organized in the interstripes. These clusters spread to one CytOx-rich thin stripe and one or two CytOx-poor interstripes in the flattened preparations. Figure 8D shows clusters of CTb-labeled cells in six enlarged microphotographs of flattened sections of V2.
Morphometric analyses

CytOx bands

To evaluate the correspondence of the locations of the patches and the CytOx modules of V2, we measured the width of the bands and interbands. Figure 9 shows the average width of the CytOx-rich thin and thick bands and CytOx-poor interbands of V2 in three animals. On average, the thin bands were 1,300 μm wide, and the thick bands were 1,600 μm wide. The interbands were, on average, 350 μm wide.

Laminar distribution

We also measured the number of cells in each cortical layer and in each CytOx compartment. Figure 10 shows...
the relative frequency of labeled cells in each CytOx compartment and in cortical layers II, III, and V after injections of CTb into V4. Layer III had the greatest number of cells, followed by layer V, and very few cells were located in layer II. In every layer, there were more cells in the thin bands than in the interbands and very few cells in the thick bands.

Figure 11 shows the variations in the numbers of labeled cells and fiber densities throughout the cortical layers of V2 at the location of a set of modules composed of one CytOx-rich thin band and two CytOx-poor interbands of V2 after injections of CTb into V4. The density of labeled cells peaked in layer III (5 cells/mm²), whereas the density of fibers was greatest in layer V.

DISCUSSION
CytOx modules and pathways
The occipitotemporal cortical pathway is called the ventral stream of visual processing and is necessary for visual recognition in monkeys (Ungerleider and Mishkin, 1982; Mishkin et al., 1983; Ungerleider, 1985; Maunsell and Newsome, 1987; Weller, 1988; Desimone and Ungerleider, 1989; Felleman and Van Essen, 1991; Van Essen et al., 1991, 1992). This pathway includes successively ordered areas that begin in the striate cortex (visual area V1), and project to the extrastriate visual areas V2 and V4, and then further on to the inferior temporal cortex (areas TEO and TE). Along this stream, the complexity of the physiological properties of neurons and their responses to the visual stimuli increase, and receptive field sizes also increase at each stage (Gross et al., 1972; Desimone and Gross, 1979; Gattass et al., 1981, 1988; Tanaka et al., 1991; Pasupathy and Connor, 1999; Orban, 2008). Kravitz et al. (2011) reviewed the anatomical and functional evidence in primates and suggested that the dorsal stream actually gives rise to three distinct, major pathways: a parieto-prefrontal pathway, a parieto-premotor pathway, and a parieto- medial temporal pathway.

Within at least the early stations of these pathways, there are functional modules that make differential
contributions to color, form, and motion processing. In V1 and V2, modules can be identified histologically in cytochrome oxidase (CO)-stained sections (Horton and Hubel, 1981; Livingstone and Hubel, 1982, 1984, 1987; Humphrey and Hendrickson, 1983; Tootell et al., 1983; Carroll and Wong-Riley, 1984; Horton, 1984; Wong-Riley and Carroll, 1984; DeYoe and Van Essen, 1985; Hubel and Livingstone, 1985, 1987; Shipp and Zeki, 1985; Cusick and Kaas, 1988; Zeki and Shipp, 1989; Gattass et al., 1990; Krubitzer and Kaas, 1990).

Area V2 has both thick and thin stripes of dense CO staining situated deep in layer III and layers IV and V, and both of these stripes are bordered by CO-poor interstripe regions. The CO-rich thin stripes of V2 receive inputs from the blobs of V1, the interstripe regions receive inputs from the interblob regions of V1, and the CO-rich thick stripes receive inputs from layer IVB of V1 (Livingstone and Hubel, 1984, 1987). However, Federer et al. (2013) have proposed that the pale stripes in V2 are anatomically two distinct compartments receiving different projections from layers II and III in V1, with only one set receiving a significant projection from layer 4B, whereas the other set receives few or no layer 4B projections. Consistent with these inputs, although the CO-dense stripes are not consistently distinguishable as thick or thin, they can be identified by functional differences, with neurons in the thick stripes sensitive to binocular disparities and stimulus orientation and direction, neurons in the thin stripes sensitive to luminance and color, and neurons in pale stripes sensitive to stimulus orientation (Livingstone and Hubel, 1984; DeYoe and Van Essen, 1985; Hubel and Livingstone, 1985, 1987; Shipp and Zeki, 1985; Lu and Roe, 2007, 2008; Felleman, 2008, Kasnak et al., 2009; Lim et al., 2009). Lu et al. (2010) found robust direction maps that exhibited pinwheel or linear organizations in V2 thick/pale stripes that avoided thin stripes. The presence of motion maps in V2 points to a newfound prominence of V2 in motion processing, for contributing to motion perception in the dorsal pathway and/or for form perception dependent of motion.

Area V4, which receives inputs from both CO-rich thin stripes and CO-poor interstripe regions of V2, has both color- and form-selective cells and many cells that are sensitive to both features (Desimone et al., 1985, 1992; DeYoe and Van Essen, 1985; Shipp and Zeki, 1985, 1989; Desimone and Schein, 1987; Zeki and Shipp, 1988, 1989; Schein and Desimone, 1990; DeYoe and Sisola, 1991; Van Essen et al., 1991). V4 was initially characterized as a color-processing area, but subsequent studies revealed that it contains a diverse complement of cells, including those with preferences for color, orientation, and disparity, as well as higher order feature preferences. V4 also appears to have a modular organization (DeYoe and Sisola, 1991; DeYoe

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**Figure 8.** Types of cells that were CTb-labeled in oblique sections of V2. **A:** Enlarged colored photomicrograph of CTb-labeled cells in layer III in V2. **B,C:** CTb-immunoreacted sections showing clusters of labeled cells in the upper layers of V2. Asterisks indicate corresponding blood vessels. **D:** Cell types exhibiting feedforward projections in layer III of V2. Enlarged photomicrographs show the labeled cells in V2 after an injection of CTb in V4. Scale bar = 200 µm in A, C (applies to B,C), and D.
et al., 1994; Yoshioka et al., 1992); the geometry of the modules and their relations to color and form analyses remain unclear. Tanigawa et al. (2010) suggested that V4 consists of multiple, alternating, band-like structures for color and orientation. Such band-like organization was also found in the pattern of afferent connections from V2 thin stripes and pale stripes (Xiao et al., 1999). More recently, Li et al. (2013) found that V4 even contains direction-preferring domains that are preferentially activated by stimuli moving in one direction, suggesting that, in contrast to the classical view, motion information is also processed in ventral pathway regions such as area V4. Although neurons within the stations along the occipitotemporal pathway appear to play a significant role in the analyses of color and form information, it should be emphasized that this pathway is not equivalent to the “parvo” system. First, the blobs in V1, which contribute to this pathway, receive magnocellular and parvocellular inputs (Lachica et al., 1992). Second, deactivations of both the magnocellular and parvocellular layers of the LGN affect the activity of V4 neurons (Ferrera et al., 1992). In Cebus monkeys, we found that the sources of the projections from V2 to V4 are located in the CytOx-rich thin stripes and CytOx-poor interstripe regions, which corroborates earlier reports in the macaque (Nakamura et al., 1993; Felleman et al., 1997).

Yau et al. (2013) revealed a dynamic transformation of shape information occurring in area V4. Such dynamics might reflect influences from earlier (V1, V2) and later (central and anterior IT) processing stages in the ventral pathway and reflect a sequential hierarchical process of shape synthesis.

Pan et al. (2012) suggested that V4 is also a key cortical locus for integration of local features into global contours for the representation of illusory contours that are achieved by a progressive integration of local cues through the reciprocal circuits of V1, V2, and V4, with V4 as a key player mediating the feedforward and feedback processing streams. In addition, according to Maunsell and Treue (2006), the existence of feature segregation in V4 would have significant implications for studies on the role of V4 in visual perception and feature-based attention. An understanding of the feedback and feedforward interplay between V2 and V4 is also important to an understanding of how attention modulates early stages of visual processing. Attentional influences have now been demonstrated throughout the visual cortex, and their influence on the processing of visual information is profound (Desimone, 1998). Reynolds et al. (2000) and Cohen and Maunsel (2011) have shown that attention increases sensitivity of V4 neurons.

Feedback and feedforward connections and laminar distribution

The laminar location of the cell bodies and terminals of interareal connections has been seen as a criterion
for the hierarchical structural organization of the cortex and has been intensively studied. It has been proposed that feedforward projections from lower order cortical visual areas originate mainly in layer III and terminate predominantly in layer IV, whereas feedback projections from higher order areas to lower order areas originate mainly in layers V and VI and avoid layer IV. In this previous study, anterograde labeling in the remainder of the areas was always heaviest in layer IV. Based on the distinctions mentioned above, Ungerleider et al. (2008) were able to categorize the connections of V4 in macaque as feedback, intermediate, or feedforward. In both V2 and V3, retrogradely labeled cells were located predominantly in the supragranular layers, and the anterogradely labeled terminals were located in layers I and VI and avoided layer IV. Our results in *Cebus* emphasize the origin of the V4 feedforward connections in layer III and identify the target of the feedback connections as the supra- and infragranular layers, but mainly layer V.

**Figure 11.** Variation in the numbers of labeled cells and the densities of fibers in different cortical layers through a CytOx-rich thin band of V2 after injection of CTb into V4. The density of labeled cells peaks in layer III (5 cells/mm²), whereas the density of fibers is greater in layer V. The measurements were made in an oblique section divided into 21 rectangles of $534 \times 70 \mu m$. Scale bar = 200 μm.

Information conveyed along the occipitotemporal pathway

In addition to the major anatomical routes into the temporal lobe, there are other, smaller, "side routes" whose significances are currently unclear. Two of the most intriguing side routes are the direct projection from the foveal representation of V1 to area V4 that bypasses V2 (Kuypers et al., 1965; Cragg and Ainsworth, 1969; Zeki, 1978; Yukie and Iwai, 1985; Steele et al., 1991), and the direct projection from V2 to TEO that bypasses V4 (Morel and Bullier, 1990; Baizer et al., 1991; Distler et al., 1993; Weller and Steele, 1992). These bypass pathways may allow for a certain amount of parallel processing across the areas of the occipitotemporal projection system.

The intermingling of V2 cells projecting to areas V4 and TEO suggests that similar information is relayed to both areas. Although there is still some question about the relative proportions of color- and orientation-selective cells in the CO-rich thin stripes and CO-poor interstripe regions of V2, it is commonly accepted that these two stripe systems, taken together, contain many color- and form-selective cells and some cells that are selective for both (Livingstone and Hubel, 1984; DeYoe and Van Essen, 1985; Hubel and Livingstone, 1985, 1987). The thin stripe and interstripe regions lack large numbers of directionally selective cells, which instead are mainly located in the CO-rich thick stripes that project to area MT (DeYoe and Van Essen, 1985; Shipp and Zeki, 1985; Hubel and Livingstone, 1987).
CONCLUSIONS

We found that the sources of the projections from V2 to V4 are located in the CytOx-rich thin stripes and CytOx-poor interstripe regions, which confirms earlier reports (Nakamura et al., 1993; Felleman et al., 1997; Nascimento-Silva et al., 2003). Although some authors have reported some intermingling between the visual pathways (Sincich and Horton, 2005; Nasti and Callaway, 2006; Nasti et al., 2006; Ninomiya et al., 2011), our data reinforce the idea that some degree of segregation seems to be maintained between the dorsal and ventral streams. So, the parallel inputs from the LGN can be recombined in areas V1 and V2, where the dorsal and ventral streams originate, and conveyed to areas MT and PO as well as other parietal areas for the processing of motion and depth and to V4 and IT for the processing of form and color (Gattass et al., 2005; Orban, 2008; Nasti and Callaway, 2009). We also found that the majority of feedforward projections were from pyramidal cells with ascending dendrites located predominantly in the supragranular layers and sparsely in the infragranular layers, whereas the feedback projections (i.e., terminal-like labels) were located in the supragranular and infragranular layers.

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CONFLICT OF INTEREST STATEMENT

We declare that there is no conflict of interest, in the form of either financial, personal, or other relationships with other people or organizations within 3 years of beginning the submitted work that could inappropriately influence the results or interpretation of the data of the manuscript.

ROLE OF AUTHORS

All authors had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. Study concept and design: SNS, RG. Acquisition of data: SNS, RG, JGMS, CP. Analysis and interpretation of data: SNS, RG, JGMS. Drafting of the manuscript: SNS. Critical revision of the manuscript for important intellectual content: RG, JGMS, CP. Statistical analysis: SNS. Obtained funding: RG. Study supervision: RG.

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