Connectional and Architectonic Evidence for Dorsal and Ventral V3, and Dorsomedial Area in Marmoset Monkeys

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The existence of a third visual area, V3, along the outer margin of V2 was originally proposed for primates on the basis of projections from V1. The evidence for V3 was never totally convincing because investigators failed to demonstrate V1 projections to ventral V3, and projections to dorsal V3 could be attributed to the dorsomedial visual area (DM). We have reexamined the issue by placing large injections into both dorsal and ventral portions of V1 and subsequently processing flattened cortex for myelin and cytochrome oxidase so that borders of V1 and V2 could be determined accurately. The injections were in small-brained marmosets, where ventral V1 was most accessible and cortex could be flattened easily. The results indicate that dorsal V1 (representing the lower visual quadrant) projects to a narrow “dorsal V3” located between DM and dorsal V2, whereas ventral V1 (representing the upper visual quadrant) projects to a narrow “ventral V3.” Architectonic borders for these dorsal and ventral strips were clearly apparent. In addition, all parts of V1 project to DM, whereas ventral V1 connections indicate that the dorsolateral area (DL) extends more ventral than has been established previously. We also placed injections within dorsal V2, dorsal and ventral DM, and dorsal, central, and ventral middle temporal (MT) area. Results from these injections were consistent with the proposed retinotopic organizations of V3, DM, and DL. We provide compelling evidence for the existence of areas V3, DM, and DL in marmosets and suggest that these areas are likely to be found in all primates.

Key words: V3; DM; DL; V4; MT; visual cortex; extrastriate cortex

Despite >30 years of intensive research on the organization of visual cortex in primates, there is widespread agreement on the existence of only a few visual areas (Kaas, 1996, 1997). Surprisingly, V3 or VIII, one of the first proposed extrastriate visual areas in primates, has been one of the most questioned. The first evidence for a V3 came from studies of the projection patterns of V1 in macaque monkeys (Cragg, 1969; Zeki, 1969). V1 appeared to project topographically to two successive belt-like surrounding fields, an immediately adjacent V2 and a V3 along the outer border of V2. The proposed areas were continuous and approximately equal in width, and they formed mirror images of each other in retinotopic organization. The concepts of V2 and V3 fit well with expectations stemming from the early cytoarchitectonic descriptions of Brodmann (1909) of a large primary visual cortex, area 17, surrounded by areas 18 and 19, especially because areas 18 and 19 had already been associated with a V2 and a V3 in cats (Hubel and Wiesel, 1965). In fact, V2 and V3 were originally designated as “areas 18 and 19” in macaques (Zeki, 1969) despite the recognition that the V1 projection fields were both well within Brodmann’s proposed area 18.

Although evidence accumulated for the existence of V2, and V2 is now a widely accepted subdivision of visual cortex, convincing evidence for V3 failed to materialize. The first problem was that V3 did not correspond to any of Brodmann’s architectonic subdivisions, and subsequent architectonic attempts to define the field were inconsistent and unconvincing (see Discussion). A second problem was that further studies of V1 projections in primates did not reveal a complete ring-like projection field along the outer border of V2. Projections to cortex just rostral to V2 either were not found in squirrel monkeys (Spatz and Tigges, 1972) or were found only in a small region of cortex along the dorsolateral (DM) rostral border of V2. These V1 projections to a dorsomedial location were interpreted either as within another visual area, DM, in owl monkeys (Lin et al., 1982; Krubitzer and Kaas, 1993) or to a dorsal V3 that differed considerably from the V3 of initial proposals in macaque monkeys (Burkhalter et al., 1986; Van Essen et al., 1986). Some held that V3 existed only along the dorsomedial border of V2, rather than as a long, band-like area. This dorsal V3 was a fragment of its former size, and it had the improbable characteristic of representing only the lower visual quadrant. Other types of evidence, from microelectrode mapping studies, investigations of single neuron response properties, and imaging brain activity, have not resolved the issue, and a range of opinions continues to persist (see Discussion).

Because of the importance of the issue, we reinvestigated the connections and architecture of the V3 region. The results provide evidence for a modified V3, DM, and an enlarged dorsolateral (DL) visual area [Stepniewska and Kaas (1996); for the relationships of DM to V3A and DL to V4 of macaque monkeys, see Beck and Kaas (1999)].
MATERIALS AND METHODS

We related cortical patterns of cytochrome oxidase (CO) and myelin to connections of V3 and DM after multiple injections of fluorescent and bi-directional tracers into retinotopic locations in V1, V2, DM, and middle temporal (MT) areas in a total of three marmosets (Callithrix jacchus) (see Table 1). Of particular interest was whether ventral V1 interconnects with ventral cortical lateral to V2. Because it is difficult to inject all parts of V1 and flattened cortex in macaques, we used the more manageable brains of New World marmoset monkeys. The small, liscencephalic brain of the marmoset enabled us to more easily place large injections of tracers in ventral V1 and more effectively flatten cortex so that areal patterns in the architecture and connections could be evaluated. In addition to V1 injections, we placed injections within V2, MT, and the proposed field DM to help determine further whether V3 and DM are valid areas. Marmosets also served as a useful model because of the existence of detailed retinotopic maps of the areas that we injected (Rosa and Schmid, 1995; Fritsches and Rosa, 1996; Rosa et al., 1997; Rosa and Elston, 1998; Rosa and Tweedale, 2000). Therefore we were able to test the validity of these areas.

Surgery and injections. All procedures were approved by the Vanderbilt animal care and use committee and followed National Institutes of Health guidelines. Marmosets were anesthetized with ketamine hydrochloride (30 mg/kg) and xylazine (5 mg/kg) and placed in a stereotaxic headholder, and different portions of the dorsolateral surface of visual cortex were exposed depending on the intended injection sites (see below).

For V1 injections, a small portion of the skull covering the caudal end of visual cortex in the left hemisphere was removed by trephination and small rongeurs. After removal of the dura, portions of the dorsal lateral surface, the medial wall, and ventral surface of striate cortex were gently aspirated away to allow access to the calcarine sulcus of the right hemisphere (Fig. 1A). Once the dura was removed from the medial wall of the right hemisphere the calcarine sulcus was clearly visible (Fig. 1B). We were able to inject a total of nine neuroanatomical tracers (see below) in two cases along the upper and lower banks of the calcarine sulcus (see Results) without contamination of the opposite hemisphere. Pressure injections of 0.1–0.3 μl were made at ~20° from perpendicular to the cortical surface using a stereotaxic syringe holder and 1.0 μl Hamilton syringes with sealed glass micropipettes. Any leakage of tracer from the injection site was removed with a swab during and after the injection to prevent any inadvertent spread. Injections placed on the lower bank were considered to be in the upper quadrant of V1, whereas injections along the upper bank were considered to be in the lower quadrant (Fig. 2B) [based on retinotopic maps of marmoset V1 by Fritsches and Rosa (1996)]. Injections were placed by estimated depth into the middle layer of cortex. The dense core of injected tracer included Fritsches and Rosa (1996); in V1 and up to 5 mm lateral from the medial lip. After injections, a soft ointment. Each marmoset was then given prophylactic injections of penicillin (60,000 U/kg) and an analgesic, banamine (1 mg/kg). They were carefully monitored during recovery from anesthesia.

Tracers used in this study included 0.1–0.3 μl of fluorochromes diamidino yellow (DY; 2%) (Keizer et al., 1983), Fast Blue (FB; 3%) (Kuypers et al., 1980), fluororuby (FR; 10%) (Schmued et al., 1990), and fluoroemerald (FE; 10%) (Nováková et al., 1997) and the bi-directional tracer, cholera toxin-B subunit (CTB; 5%) (Ericson and Blomqvist, 1988).

Histology and data analysis. Seven to nine days after surgery, the marmosets were first anesthetized with ketamine hydrochloride (30 mg/kg) and xylazine (5 mg/kg) and then a lethal dose of sodium pentobarbital. When areflexive, they were perfused transcardially with 0.9% PBS followed by a solution of 2% paraformaldehyde fixative and subsequently 2% paraformaldehyde with 10% sucrose in PBS.

After removal of the brain, the cortical hemispheres were carefully separated from underlying structures and subsequently flattened in one of two ways. All flattening was similar to previous methods used for macaques (Krubitzer and Kaas, 1990b, 1993) but differed in how the calcarine sulcus was unfolded. We focused on either maintaining the opened calcarine sulcus in one piece (Fig. 2A) or cutting it through the middle (along the fundus of the calcarine), presumably separating upper and lower quadrants of V1 into equal halves. After they were flattened, the hemispheres were submerged overnight in 30% sucrose in PBS while placed under a glass slide. Additional weight was not placed on top of the glass slide because the relatively liscencephalic marmoset cortex was easy to flatten.

The next day, the flattened hemispheres were frozen and cut parallel to the surface in 40 μm sections. A series of every third or fourth section was mounted on glass slides without further processing to observe
fluorescent tracers. If CTB was injected, a second series was put through an immunocytochemical procedure to visualize the transported label (Bruce and Grofova, 1992; Sakai et al., 2000). Two more series were processed for either myelin (Gallyas, 1979) or CO (Wong-Riley, 1979).

The location of retrogradely labeled fluorescent and CTB cells and anterogradely labeled terminals were digitized by using an XY encoder attached to the stage of a high-powered fluorescent and bright-field microscope and plotted with the aid of Igor Pro (WaveMetrics, Lake Oswego, OR). The XY coordinates were imported as EPS files from Igor Pro into Canvas 7 (Deneba) and printed. Borders from CO and myelin stains of adjacent sections were drawn on printed copies of each cortical section with plotted cells, scanned into Canvas 7, and redrawn digitally. Within Canvas 7, labeled cells were aligned with adjacent sections using injection sites, blood vessel patterns, and section contours as landmarks.

The image in Figure 1 was captured using a digital camera (Scion Corp.) and NIH image software and adjusted for contrast within Canvas 7. Other images (Figs. 3, 4, and 5) were captured using a Spot 2 camera mounted on a Nikon E800 microscope and acquired through Adobe Photoshop 5.02 software and adjusted for brightness. None of the images were altered in any other way.

RESULTS

Our results provide architectonic and connectional evidence for the existence of retinotopically organized dorsal and ventral halves of V3 (V3d and V3v, respectively) along the outer border of the dorsal and ventral halves of V2, and the DM visual area rostral to the dorsal portion of V2 near the medial wall but displaced from the V2 border by V3d (Fig. 2A). Furthermore, we identified other visual areas including DL just rostral to central V2 and central V3v and V3d, area MT and its crescent (MTc) rostral to DL, multiple regions within posterior parietal cortex, and a distinct area in inferotemporal (IT) cortex (Fig. 2). These results are presented in several parts. First, we show the relative locations and extents of V1, V2, V3d, V3v, DM, and MT in sections of flattened cortex processed for cytochrome oxidase or myelin. The connection patterns are then presented, beginning with the results of V1 injections in both the upper and lower banks of the calcarine sulcus that revealed retinotopic connections of extrastriate cortex. Results from the placement of tracers in the dorsolateral aspect of V2 are presented next. Finally, the connection patterns after injections in DM and MT are compared and provide further evidence of retinotopy within V3d/v and DM.

Architectonic subdivisions of marmoset visual cortex

We used stains for CO and myelin as a reliable means of identifying visual areas on flattened marmoset cortex. Figure 3 shows an example of a brain section that was processed for CO, covering most of a right hemisphere. A portion of striate cortex is easily distinguished as the most darkly stained region along the caudal end of the brain section (the full extent of V1 is portrayed in Fig. 2). V1 is most densely stained in layer 4. Variations in staining density in V1 reflect variations in the depth of the section through cortex. The characteristic CO blobs of primate V1 are
The CO blobs are surrounded by interblobs that stain dark for myelin where the CO stain is light (data not shown) (Krubitzer and Kaas, 1990a).

Marmoset V2 is characterized by the CO dark and light bands of similar thickness extending out from the border of V1 (Fig. 3) [also see Krubitzer and Kaas (1990b)]. The bands extend 3–4 mm at the dorsal and ventral ends of V2 and only 1–2 mm in dorsolateral cortex where V1 bulges into V2.

Another set of CO dark and light bands occupies strips of cortex along the outer border of the dorsal and ventral halves of V2 (Fig. 3). These strips correspond to V3d (Fig. 3C) and V3v (Fig. 3D, E).

The bands in V3 do not seem to be simple continuations of the V2 bands, and they appear somewhat thicker, darker, and more diffuse than those of V2. They are approximately two-thirds to one-half the length of the V2 bands. The V3 strips of cortex stain more lightly for myelin than adjacent DM (Fig. 4A). Previous reports on the architecture of New World monkey visual cortex may have included these V3 strips of bands as either continuations of V2 or as parts of DM and the ventroposterior (VP) area (Krubitzer and Kaas, 1990a,b, 1993; Beck and Kaas, 1998). As we demonstrate in the next section, V3d represents the lower visual quadrant and V3v represents the upper quadrant.
In New World monkeys, DM is clearly identified architectonically as a darkly myelinated oval of cortex similar in size to MT (Fig. 4B) (Allman and Kaas, 1975; Krubitzer and Kaas, 1990b, 1993; Beck and Kaas, 1998) and contains an orderly retinotopic map of the entire contralateral hemifield [Fig. 2; from connectional data presented here and from microelectrode recordings by Allman and Kaas (1975); Krubitzer and Kaas (1993); Rosa and Schmid (1995)]. As mentioned above, previous descriptions of DM were based on CO bands of V3d (Fig. 3). Thus, we distinguish both a DM that coincided with the CO bands of V3d (Fig. 3) and a DM that is medial to the dimple that comprises the superior temporal “sulcus” (STS) in marmosets. DM lies lateral to the ventral part of MTc. DM is clearly identified architectonically as a darkly myelinated oval of cortex lying lateral to VPP and nearest to LS (Krubitzer and Kaas, 1990b). The patches relate to the modular organization of intrinsic MT connections (Malach et al., 1997) and callosal connections (Krubitzer and Kaas, 1990b). Functionally, the patches may represent separate modules for orientation and direction selectivity (Malonek et al., 1994) or local and global motion selectivity (Born and Tootell, 1992).

Figure 4 shows the locations of auditory cortex (A), primary somatosensory cortex (SI), and surrounding somatosensory areas 1, 3a, S2, and PV, based on CO architecture. All of these areas are also visible in sections stained for myelin (data not shown) (Krubitzer and Kaas, 1990c).

V1 injections
Tracer injections in different parts of V1 revealed intrinsic connections of V1 and retinotopic patterns of connections with V2, V3d/v, DM, and MT as well as connections with other extrastriate areas. Most importantly, injections in the portion of V1 representing the upper visual quadrant demonstrated connections located at the rostral border of MT is less uniformly stained, probably because of uneven flattening near the LS. Areas FSTd and FSTv are visible as moderately myelinated regions of cortex located successively lateral to ventral MTc. The caudal end of the darkly stained auditory core is also visible (A). For both A and B, right is rostral and up is medial. For abbreviations, see Figure 2. Scale bars, 2 mm.
with DM and V3v. We found that the lower bank of the calcarine sulcus, a reliable marker for V1 upper field (Fritsches and Rosa, 1996), can be approached through an aspirated opening in the opposite hemisphere (see Materials and Methods and Fig. 1). Using this approach, we were able to make nine injections, six along the lower bank and three on the upper bank of the calcarine sulcus in two cases (99-48 and 99-64) (Figs. 5-7), which allowed us to compare connection patterns of the peripheral fields of both the upper and lower quadrants (Table 1).

Areal patterns of intrinsic connections after V1 injections in primates have been described previously using wheat germ agglutinin conjugated with horseradish peroxidase injections in squirrel monkeys and macaque monkeys (Rockland and Lund, 1983), and in prosimian galagos (Cusick and Kaas, 1988a). Here we briefly present evidence for a similar pattern in marmosets based on injections of CTB and fluorochromes. The CTB injection in case 99-48 shows an intricate pattern of labeled terminations and cell patches that surround the injection site in a pinwheel-like pattern (Fig. 5). The cell pattern is seen throughout most sections of cortex, including superficial and deep cortical layers (although in deeper sections the pattern was less distinctive). Terminals were primarily found in more superficial sections. Thus, intrinsic connections of V1 arise from neurons in both shallow and deep layers, but terminations are most dense in supragranular layers. Fluorochrome injections showed similar patterns of labeled cells. The FB and DY injections in case 99-48 formed distinct groups of small patches (Fig. 6). Because the needle track went through the lip of the upper bank of the calcarine, the FR injection in case 99-64 resulted in two injection sites with distinct pinwheels of adjacent patches (Fig. 7). Other injections in V1 produced uniform distributions of adjacent label.

Table 1. Summary of injection sites

<table>
<thead>
<tr>
<th>Case number</th>
<th>Injection site</th>
<th>Tracer</th>
<th>Figure</th>
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<tr>
<td>99-64</td>
<td>Ventral V1</td>
<td>CTB</td>
<td>7B</td>
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<tr>
<td></td>
<td></td>
<td>DY</td>
<td>7C</td>
</tr>
<tr>
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<td></td>
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<td>1B,7A</td>
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<td>Dorsal V1</td>
<td>FR</td>
<td>7D</td>
</tr>
<tr>
<td></td>
<td>Dorsal V2</td>
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<td>N.S.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FB</td>
<td>8A</td>
</tr>
<tr>
<td></td>
<td></td>
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<td>8B</td>
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<td></td>
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<td>6</td>
</tr>
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<tr>
<td>97-109</td>
<td>Dorsal V2</td>
<td>DY</td>
<td>N.S.</td>
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Injection site indicates whether a tracer was placed in the dorsal or ventral aspect of the listed visual area. DY, Diamidino yellow; FB, Fast Blue; FE, fluoroemerald; FR, fluororuby; N.S., not shown.

When the intrinsic label was patchy, the patches were about the size of the CO blobs, yet patches did not appear to have any
consistent relationship to blobs. Patches of label fell within both the CO blobs and the myelin-dense interblobs.

The injections in V1 demonstrated interconnections with V2, V3, DM, DI, MT, MTc, MST, FST, DL, and IT. These connections are all demonstrated in case 99-48, with a large CTB injection in the ventral portion of V1 that represented the upper visual quadrant (Fig. 5). The bidirectional tracer labeled axons that were clustered with cells in such high density in some regions that they were easily seen under low magnification (Fig. 5A). Dense patches of labeled terminations were obvious in V2, V3v, DM, and MT. The patches are in parts of V2, DM, and MT known to represent the upper visual quadrant, and thus the projections are retinotopically matched. The patch in V3v is clearly distinct from the one in ventral V2, and it is clearly beyond the architectonic boundary of V2. This patch represents the first clear demonstration that ventral V1 projects to cortex along the rostral border of ventral V2, that is, to V3v.

V1 may project to other areas of extrastriate visual cortex as well, but if so the projections were too sparse to demonstrate as obvious patches of labeled axons. However, retrogradely labeled neurons were obvious both in regions with labeled patches of fibers and in additional regions of cortex (Fig. 5B). In V2, V3v, DM, and MT, the distributions of labeled neurons included regions of labeled axons but were more extensive. Thus, feedback connections may be more broadly distributed than feedforward projections. In addition, many labeled neurons were found in IT cortex, DI, MST, FST, LPP, and VPP. A few labeled neurons were in V3d, and thus these connections were not retinotopic.

Labeled cells were seen in similar locations after other injections into ventral V1 of retrograde fluorescent tracers FE and FR, in case 99-48 (Fig. 6), and FB and DY in case 99-64 (Fig. 7A, C), which included another injection of CTB in ventral V1 (Fig. 7B).

Injections of fluorescent tracers in dorsal V1, also in cases 99-48 and 99-64, labeled cells in the same visual areas (Figs. 6, 7D). However, the fluorescent tracers failed to label cells in MST and FST, possibly reflecting a difference in transport between fluorescent dextran beads and CTB. Labeled cells outside of V1, V2, and V3 were found only in the deepest cortical sections, whereas the labeled terminals (CTB injections only) were seen in middle and superficial sections. This is consistent with the fact that feedforward projections usually terminate in layer IV or III, whereas feedback connections usually arise from cells in the infragranular layers (Van Vliet and Van Essen, 1983). Furthermore, because V2, V3, DM, and MT receive direct projections from V1, they can be considered to be at early stages in the processing hierarchy of visual areas (Felleman and Van Essen, 1991).

Most injections in V1 (Figs. 5, 6, 7) consistently labeled cells in at least roughly matched retinotopic locations in V2, V3, DM, and MT. Injections in ventral V1, corresponding to the upper visual quadrant, labeled large patches of nearby cells that occupied both dark and light CO bands in ventral V2 and V3v (Figs. 5, 6, 7A–C).

According to traditional views, V2 and V3 border each other along a representation of the zero vertical meridian and form mirror image reversals of each other (Fig. 2). Pairs of closely spaced injections in V1 provide support for this view. Figure 6 illustrates a mirror reversal pattern for two injections. The FE injection is farthest from the V1/V2 border and thus is farther from the zero vertical meridian in V1, whereas the FR injection is closer to the border and closer to the zero vertical meridian in V1. The FR-labeled cells (red) are caudal to the FE-labeled cells (green) in V2 and rostral to FE cells in V3v, thus providing clear evidence for a mirror reversal of the topographic pattern in V1 and V2 and between V2 and V3v. A similar pattern of connections is revealed by the three upper field injections in case 99-64 (Fig. 7A–C). The DY injection placed farthest from the zero vertical meridian in V1 labeled cells in V2 away from the V1/V2 border and nearer to the V3v/V2 border within V3v (Fig. 7C). The CTB and FB injections were ~2 and 3 mm closer to the vertical meridian in V1 and resulted in a dense patch of labeled cells adjacent to the V2/V1 border (Fig. 7A, B).

Furthermore, the FB injection that was essentially placed at the V1 vertical meridian labeled a dense patch of cells at the rostrolateral border of V3v (Fig. 7A), consistent with the interpretation that this border represents the vertical meridian of V3v.

The pattern of label from dorsal V1 injections, representing the lower visual quadrant, in V2 and V3v was symmetrical to that of the upper quadrant connections to V2 and V3v, although in some instances the pattern was more difficult to see because of the lower field representation of DM that is partially adjacent to the
rostral border of V3d. The DY and FB injections of case 99-48 were both placed close to the zero vertical meridian near the V1/V2 border on the lip of the upper bank of the calcarine sulcus (Fig. 6). A large cluster of DY-labeled cells (yellow) in V2 was adjacent to the injection site. A smaller DY cluster was more lateral, in a retinotopically matched portion of V2. A few FB-labeled cells (blue) were found exclusively in caudal V2 at the vertical meridian of V2. In V3d, the largest cluster of DY-labeled cells as well as a few FB-labeled cells were found at the rostral border along the representation of the vertical meridian. A third lower field V1 injection was placed farther from the V1/V2 border in a separate case (Fig. 7D). As expected, labeled cells in V2 clustered at the rostral border along the zero horizontal meridian. However, in V3d the cluster of labeled cells stretched across the width of the area.

Cells labeled in DM and MT were also in roughly matched retinotopic locations after the V1 injections. As shown in Figure 2, both areas have retinotopic maps, with the lower field largely medial to the upper field, with the horizontal meridian extending diagonally in a rostromedial direction, and with central vision situated at the caudolateral border [for New World monkeys see Allman and Kaas (1974, 1975); Krubitzer and Kaas (1993); Rosa and Schmid (1995); Rosa and Elston (1998)]. Injections in the representation of the upper quadrant of V1 [Figs. 5, 6 (FE), 7A–C] consistently labeled the lateral halves and thus the representation of the upper visual quadrants in DM and MT, whereas lower quadrant injections in V1 labeled the medial segments of DM and MT. The main cluster of labeled cells and terminals in MT (Fig. 5A) was centered at the matching retinotopic location, whereas additional labeled cells were scattered throughout rostrolateral MT. For most ventral V1 injections, the pattern of labeled cells in DM was more diffuse and extended diagonally throughout much of the upper quadrant representation (Figs. 5B, 7A, B). Likewise, patches of labeled cells were found in dorsal DM after injections in dorsal V1. Thus, the pattern of labeled cells after V1 injections bolsters previous reports of the retinotopic organizations of DM and MT.

Injections in V1 labeled neurons in a number of additional regions. It was difficult to assign some of these labeled neurons to specific cortical areas because of uncertainties about architectonic borders. Injections in V1 consistently labeled neurons in the temporal lobe in a broad expanse of cortex rostral to V3v. The more caudal portion of this zone of labeled neurons probably corresponds to the ventral extension of rostral and caudal division of DL where representations of the upper visual quadrant are expected. However, the boundaries of DL are not yet well established. More rostral portions of this zone correspond to the caudal division of inferotemporal cortex (ITc) (Weller and Kaas, 1987). Neurons in this region appear to be labeled by injections in either ventral or dorsal V1, suggesting that there is little retinotopic organization. Injections in dorsal V1 labeled more dorsal locations in DL corresponding to the lower visual quadrant representation, whereas labeled neurons in more dorsal locations near DM may have been in DI. Labeled neurons were also

Figure 7. Distribution of labeled cells in visual cortex after V1 injections of four tracers in marmoset 99-64. Arrows point to white ovals that represent the injection site and uptake zone of each tracer. Three injections were on the lower bank of the calcarine sulcus in ventral V1 in the upper quadrant (A–C). The FB injection was placed nearest the V1/V2 border (see Fig. 1), whereas CTB and DY were placed progressively further caudal into V1. A fourth injection, FR, was placed on the upper bank of the calcarine, in the lower quadrant of V1 (D). Ventral V1 injections labeled V3v (A–C), whereas the dorsal V1 injection labeled V3d (D). This dorsal and ventral retinotopic pattern of label is also observed for DL, MT, and somewhat for the MTc, whereas the remaining areas with label did not have a clear retinotopic pattern. Cortex was flatted as diagramed in Figure 2. Right is rostral; up is medial. Conventions as in Figure 5.
commonly found in regions of cortex just outside of MT. Many of these labeled neurons were within MTc, and their locations roughly reflected the retinotopic organization of MTc (Rosa and Elston, 1998). The few labeled neurons just rostral to MT appeared to be in MST, whereas a scattering of labeled neurons ventral to MT may have been in FST. A few injections labeled neurons in parts of posterior parietal cortex.

V2 injections
A total of five injections were placed in V2 of three separate cases (Table 1). All injections were in dorsal V2 representing the lower quadrant, and they labeled similar patterns of connections. In case 99-51, FB was injected in caudal V2 near the representation of the zero vertical meridian and labeled cells in immediately adjacent V1 (Fig. 8A). In the same case, the FR injection, more centered in V2 far from the vertical meridian, labeled cells in V1 away from the V1/V2 border (Fig. 8B). FR also labeled a distinct band of cells in V3d, adjacent to the injection site. A patch of cells was labeled in the dorsal half of DM, where the lower quadrant is represented (Fig. 2) (see previous section). FB labeled a small patch of cells dorsally in V3d and a larger cluster extending across the lower quadrant in DM along its dorsal border. Both injections labeled a small cluster of cells in rostomedial MT. Furthermore, both labeled two clusters of cells in what may be caudal DL (DLc) and rostral DL (DLr). A small cluster of FB-labeled cells was in IT cortex, whereas only a few scattered cells were labeled in IT by FR. We conclude that V2 has connections with both V3d and DM.

DM and MT injections
Although the connections of DM and MT in marmosets have been described previously (Krubitzer and Kaas, 1990a,b, 1993; Rosa and Tweedale, 2000), they have not been evaluated for evidence for V3. Here, we present results from two cases in which both DM and MT injections were made and a third case with a single DM injection (Figs. 9, 10). Having multiple injections allowed us to compare lower visual quadrant DM injections with upper visual quadrant MT injections to provide further evidence of the retinotopy in DM as well as V3d/v. Because MT and DM are approximately one-tenth the size of V1 and the retinotopic maps are more compressed and less detailed, the topographic patterns of connections from injections in these areas provide less precise information about the retinotopy of other areas.

A DY injection in case 99-51 was placed at the caudolateral border of DM (Fig. 9). This site was near the expected location of the central visual field representation of DM (Rosa and Schmid, 1995; Beck and Kaas, 1998). This single injection labeled the representations of both upper and lower fields of V2, V3, and MT. Although labeled cells in V1 were expected, only a few were found, possibly because a portion of V1 was lost in the flattening process. Nevertheless, the results provide clear evidence for a V3 that projects to DM, as well as for both ventral and dorsal subdivisions of V3. These and other connections demonstrate further that DM represents both upper and lower visual quadrants.

Ventral V2 and V3v both had clusters of labeled cells forming distinct bands that matched the thickness and separation of CO dark and light bands. These bands of cells overlapped the CO dark bands of V2 and V3v (data not shown). Furthermore, the bands of labeled cells in V3v were thicker and shorter than the V2 bands of labeled cells, similar to the differences seen in CO architecture (Fig. 3). The dorsal part of V2 and V3 had fewer labeled cells, some of which clustered together in a band-like pattern.

The remaining two DM injections were both placed at the dorsal extreme of DM, just lateral to the medial wall (Fig. 10A(F-B),B(FB), blue), which corresponded to the representation of the lower quadrant in DM (Fig. 2) (Krubitzer and Kaas, 1993; Rosa and Schmid, 1995). In the same two cases, MT injections were
placed at the lateral border of MT [Fig. 10A(DY), B(DY), yellow], which corresponds to the upper quadrant. The two DM injections (Fig. 10, blue) showed similar patterns of connections). Both labeled a dense cluster of cells within the dorsal half of DM. A band of label was seen in adjacent V3d. Cells in dorsal V2 and V1 were also labeled, and in a somewhat modular fashion, with noticeable bands of cells in V2 (Fig. 10B, blue) and patchy-like blobs of labeled cells in V1 (Fig. 10, blue). Although the relationship of labeled cells to CO bands and CO blobs in V1 were not analyzed, these CO dense regions tend to provide more of the projection to DM in New World monkeys (Krubitzer and Kaas, 1993; Beck and Kaas, 1998).

The dorsal DM injections also labeled dorsal regions in MT and MTc, cells in VPP, LPP, and frontal cortex. Both cases had labeled cells in the proposed location of DLr.

Cases 97-123 and 98-4 also had injections in MT, at virtually identical locations, situated at the ventral border adjacent to MTc (Figs. 10, yellow). Within MT, labeled cells were concentrated mostly in the ventral half. In addition, there were distinct clusters of labeled cells in ventral DM, V3v, and the ventral halves of V2 and V1. In Figure 10A, labeled cells (yellow) were clustered into band-like arrays in V3v and V2. At least one focus of labeled cells was found adjacent to the lateral border of DM, probably area DI (Fig. 10, yellow). Labeled cells were also in MTc, FST, MST, and frontal cortex in regions that may correspond to the frontal eye field and frontal ventral visual motor areas (Krubitzer and Kaas, 1990b). The important result from the two cases in which injections were placed in the representation of the lower visual quadrant of one of the visual areas, DM, and the upper visual quadrant of another visual area, MT, is that the lower quadrant injections labeled V3d and the upper quadrant injections labeled V3v. Note also that the lower visual quadrant injections (blue) also labeled lower quadrant portions of V2 and V1, whereas the upper visual quadrant injections (yellow) labeled the upper quadrant of V2 and V1 (Krubitzer and Kaas, 1990b).

An injection was placed more dorsally in MT in two other cases (data not shown). In both cases, labeled neurons were found in V3d, whereas a few labeled neurons were in V3v of one of the cases where the tracer uptake zone was partially in both dorsal and ventral MT. Other connections were typical of MT.

**DISCUSSION**

The most important conclusion derived from the present results is that marmosets, and by implication other primates, have a V3. This V3 is not like the originally proposed V3 for monkeys in that it is discontinuous, with a gap in the center, and it is perhaps half the originally proposed size. However, it does have both a dorsal half, representing the lower visual quadrant, and a ventral half, representing the upper visual quadrant. In these respects, the V3 proposed for marmosets closely resembles the V3 proposed by Gattass et al. (1988) for macaques.

The connection pattern provides evidence that this modified V3 (V3 m) represents the visual hemifield as a smaller, cruder, mirror image of V2. Both the connection pattern and the architecture suggest that V3 is modular in internal organization with band-like modules that are wider than in V2. This V3 is interconnected with V1, V2, DM, and MT (Fig. 11). V3 m appears to represent a hierarchical level of processing that is higher than V1 and V2 and possibly lower than DM and MT. Part of the evidence for V3 m is architectonic, in that both dorsal and ventral divisions are apparent in flattened cortex sections processed for CO or myelin. The other part is connectional. Neurons and terminals labeled in V3d and V3v were obviously separate from those in adjoining parts of V2, DM, and DL. This separation was apparent in part because cortex was flattened and cut parallel to the surface. Thus, surface-view patterns of connections could be evaluated directly. In addition, our unique approach to the ventral part of V1 allowed us to make large injections that produced more obvious connection patterns. Thus we were able for the first time to demonstrate V1 connections with a ventral V3. These results and others are related to previous findings (see below).

**A brief history of V3**

The original concept of V3 was that of a visual area along the outer border of V2, equal to V2 in size and shape, and mirroring V2 in a retinotopic organization (for review, see Kaas and Lyon, 2001). This concept was based on evidence from early studies of V1 projections in macaque monkeys (Cragg, 1969; Zeki, 1969) that dorsal V1 projects to both a dorsal V2 and a dorsal V3. Because of the greater difficulty in placing lesions in ventral V1, there was no convincing evidence for V1 projections to a separate ventral V3 in these studies, and later studies in macaques also failed to produce evidence for V1 projections to cortex in the expected location of ventral V3 (for review, see Burkhalter et al., 1986). For this and other reasons, cortex in the location of ventral...
V3 was assigned by some investigators to another visual area, VP (Newsome et al., 1986). This scheme left dorsal V3 as the only V3, representing only the lower visual quadrant, and VP as another incomplete visual area representing only the upper visual quadrant. In New World monkeys, initial studies revealed V1 projections to V2 but no connectional evidence for either a dorsal or a ventral V3 (Spatz and Tigges, 1972). Subsequently, V1 projections to cortex along the dorsomedial border of V2 (Lin et al., 1982; Krubitzer and Kaas, 1993; Beck and Kaas, 1998) were attributed to DM (Allman and Kaas, 1975) rather than V3, partly because these connections included both dorsal and ventral parts of V1, and thus provided evidence for a complete representation of the visual hemifield.

Another approach has been to define V3 architectonically. V3 has been described as a densely myelinated area (Burkhalter et al., 1986). However, because cortex in the region of dorsal V3 was thought to be more densely myelinated than cortex in ventral V3, the difference was seen as further evidence that only a dorsal V3 existed. A further difficulty is that dorsal V3 has been variously defined from a broad myelinated zone of 4–5 mm in width (Girard et al., 1991; Gegenfurtner et al., 1997) to a highly variable strip as narrow as 1–2 mm (Van Essen et al., 1986). Furthermore, DM has been consistently described across primate species as a densely myelinated oval (for review, see Beck and Kaas, 1999), and much of the proposed territory of V3d could include DM. The usefulness of myelin stains in identifying V3 was questioned by Gattass et al. (1988), when they concluded that V3 was difficult to distinguish from V2 and that V3 could not be distinguished from V3A (DM). We conclude that V3 can be identified in myelin, CO, and perhaps other (Hof and Morrison, 1995) preparations, and that V3d and V3v are similar in appearance. Yet, the difference between V3 and adjoining cortex can be subtle and is best seen in sections cut parallel to the cortical surface.

Microelectrode mapping studies of retinotopic patterns have resulted in a number of interpretations, some without any V3 (Allman and Kaas, 1975; Rosa and Tweedale, 2000), some with separate dorsal and ventral halves of V3 (Gattass et al., 1988), and some with a continuous dorsal to ventral V3 (Piñon et al., 1998; Rosa et al., 2000). These various interpretations are possible, in part, because adjoining recording sites tend to have similar receptive fields, and adjoining sites across the borders of areas have similar receptive fields. Thus, it is difficult to recognize whether recording sites have receptive fields belonging to one pattern or the other. In the instance of V3, there is the question of whether the separate representations of upper and lower parts of the visual hemifield belong together, as V3d and V3v, or correspond to separate fields, V3d and VP, or combine with parts of other areas such as DM or caudal DL (V4).

The present results support an interpretation that may apply to all primates. Separate dorsal and ventral narrow strips of cortex along the outer margin of V2 constitute ventral and dorsal halves of a modified concept of V3. Both halves are similar in architecture and connections. In addition, dorsal V3 has a densely myelinated DM along its rostral border. DM resembles V3 by having connections with V1, V2, and MT, but V3 and DM have quite different retinotopic organization. Most notably, V3 is a split area, whereas DM is whole. The split V3 in marmosets is much like the split V3 of Gattass et al. (1988) in macaques in propor-

**Figure 10.** Distributions of labeled cells after FB injections in dorsal DM and DY injections in ventral MT within the same cases: 97-123 (A) and 98-4 (B). Arrows point to black ovals that represent the injection site and uptake zone of each tracer. Note that ventral MT injections (yellow) label cells in V3v, in a band-like pattern, and dorsal DM injections (blue) label V3d. A similar connection pattern was seen for dorsal and ventral DL. FEF, Frontal eye field. Other conventions as in Figure 8.
tions, location, and retinotopy. Previous studies without a dorsal V3 may have included some of V3d in V2 and some in DM. Previous studies with a broad V3d may have included much or all of DM in V3d. Studies with a continuous V3 may have included some of DL (V4) in V3. Much of VP would appear to be within V3v, although a recent portrayal extends Vp dorsolaterally to include caudal DL (Rosa and Tweedale, 2000). Because of long-standing difficulties in defining boundaries of V3d, V3v, VP, and DM, studies of the properties of single-unit neurons in these areas need to be interpreted with caution.

**Other connections of V1, V2, and DM**

Although the focus of this report is on the connectional and architectonic evidence for V3, the injections in V1, V2, and DM did label other regions of cortex. V1 injections, for example, labeled a number of extrastriate regions of cortex. In addition to V3, the major connections are with V2, DM, and MT, and these connections have been described repeatedly in various primates (Krubitzer and Kaas, 1993; for review, see Casagrande and Kaas, 1994). Other connections such as those with MST and IT are not commonly reported, but they have been noted, especially in macaques (for review, see Rockland and Van Hoesen, 1994). What is perhaps surprising is the large number of labeled neurons in IT cortex after injections of V1 in marmosets (Fig. 5). Of course some of this label could be in other areas, because the boundaries of IT are uncertain. Some have proposed (Rosa and Tweedale, 2000) that a VA area borders VP (our V3v), and if such a bordering area exists, it would clearly project to V1. Some of the cortex along the rostral border of V3v would also be occupied, according to our scheme, by DL, and some of the label attributed to IT could be in DL, but we remain uncertain about the ventral extent of DL. Nevertheless, even with these uncertainties, an impressive number of neurons in IT cortex obviously project to V1. Other labeled neurons scattered in posterior parietal cortex and cortex between DM and MT are in the territories of proposed visual areas such as DI, VPP, and LPP, but these areas have not been well defined, and further studies are needed. However, MTC does appear to be a visual area that was clearly identified by architectonic characteristics in the present study, and we present clear evidence of MTC projections to V1. Such connections have been reported for V4t of macaques (Felleman and Van Essen, 1991), corresponding to dorsal MTC.

Many of the neurons labeled by V1 injections were in locations where we had no evidence of reciprocal projections from V1. These include the IT region, MST, FST, MTC, DI, and posterior parietal cortex. This simply could be the consequence of it being more difficult to detect a sparse distribution of labeled fibers in comparison to labeled cell bodies. Yet, others have also noticed that V1 injections label neurons in locations where no fibers are labeled by such injections and have concluded that some regions of feedback connections to V1 do not receive direct projections from V1 (Felleman and Van Essen, 1991; Rockland and Van Hoesen, 1994).

Finally, our V2 and DM injections, in addition to providing evidence for V3, further demonstrated the connections of these two fields. The results are largely consistent with previous descriptions of V2 (for review, see Stepnowska and Kaas, 1996; Gattass et al., 1997) and DM (for review, see Beck and Kaas, 1999) connections. However, we can be fairly certain that our V2 or DM injections did not also include V3d, and thus the results are not contaminated by V3 connections.

**REFERENCES**


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