INTRODUCTION

As a part of ongoing projects aimed at defining the visual areas of primates, we report on the visuotopic organization of the primary visual cortex (area 17, or V1) and adjacent prestriate cortex in a prosimian primate, the small-eared greater galago (Galago garnetti). Although most of the research on the organization of primate visual cortex has concentrated on the simian (i.e., anthropoid) primates (New World monkeys, Old World monkeys, apes, and humans) (Hoffstetter 1982; Martin 1990), the study of cortical organization in prosimian primates is important for several reasons. The brains of prosimians are smaller relative to body size than those of monkeys (Jerison 1979; Preuss and Goldman-Rakic 1991a,b; Stephan and Andy 1969; Stephan et al. 1981), and it is possible that the number and the complexity of cortical visual areas are reduced in comparison with those of simian primates. This simpler pattern of organization may be easier to unravel. Moreover, by drawing attention to those features of the organization of cortical visual areas shared by different primate groups, a comparative analysis can help understanding of the organization in primates in general, and also clarify which changes occurred in the cortex after the separation of the Order Primates from the lineages leading to related mammals, such as megachiropteran bats, dermopterans, and tree shrews (see Preuss and Goldman-Rakic 1991a; Rosa et al. 1996 for summaries of the phylogenetic relationships between galagos and other primates).

The galago is a good choice for study of the organization of striate and extrastriate visual areas because much work has been done to characterize the anatomy and physiology of other portions of its visual system. For example, it has been demonstrated that the parallel (magnocellular, parvocellular, and koniocellular) pathways from the retina to V1 are distinct and well developed in this genus (Casagrande and Norton 1991; Florence and Casagrande 1987; Florence et al. 1983); in addition, it is one of only three primate species in which the receptive field structures of cells in both the lateral geniculate nucleus and V1 have been compared (DeBruyn et al. 1993; Irvin et al. 1993; Norton and Casagrande 1982; Norton et al. 1988). There is also detailed information on the connectional anatomy of all levels of the visual system in this prosimian (Cusick et al. 1984; Krubitzer and Kaas 1990, 1993; Symonds and Kaas 1978; Tiggges and Tiggges 1970; Tiggges et al. 1973; Wall et al. 1982; Weller and Kaas 1982). Finally, and perhaps most importantly, visual resolution and sensitivity have been determined both behav-
iorally and physiologically in this species (Bonds et al. 1987; Langston et al. 1986).

The primary objective of this study was to describe the representation of the visual field in striate cortex and immediately adjacent extrastriate cortex in Galago. This type of information is likely to be valuable for the interpretation of anatomic and physiological studies of cortical organization in this genus. In the process of recording from V1 and the second visual area (V2), we also obtained many recordings from neurons located in the adjacent visually responsive cortex. Although not sufficient to yield a full picture of the organization of extrastriate cortex in Galago, these exploratory recordings directly address important aspects of the organization of the dorsomedial visual cortex rostral to V2 in prosimians.

METHODS

Four adult galagos (G. garnetti) were submitted to single recording sessions. Although most of the data on V1 and V2 were collected in two experiments, the results from all four cases were in close agreement. Anesthesia was induced with halothane (4%) and maintained during surgery with multiple intravenous doses of brevital (methohexital sodium, 10 mg/kg). Once anesthetized, the animals also received intramuscular injections of atropine (0.15 mg/kg) and dexamethasone (0.4 mg/kg). A tracheotomy was performed, and the trachea was cannulated to allow for artificial ventilation. The skull was fitted with a stainless steel bolt and an acrylic well centered on the midline. An extensive craniotomy was made exposing the occipital lobe as well as adjacent parts of the temporal and parietal lobes, and the dura mater was resected. Once these procedures were completed the well was filled with silicone oil, and a photograph of the cortical surface was taken to be used as reference for the placement of electrode penetrations.

During the recordings, muscular paralysis was induced by the intravenous infusion of pancuronium bromide (Pavulon, 2 mg/kg, followed by 2 mg·kg⁻¹·h⁻¹) in lactated Ringer solution with 5% dextrose. For anesthesia, the intravenous injection also included the opioid sufentanil citrate (average dose 12 µg·kg⁻¹·h⁻¹). In one of the animals, this was supplemented with small intravenous doses of brevital (2 mg/kg) on the 3rd day of the experiment, to suppress episodes of cyclical cortical spontaneous activity. The galagos were also maintained under artificial ventilation with a gaseous mixture of 75% nitrous oxide-23.5% oxygen-1.5% carbon dioxide, which potentiates the effects of the intravenous anesthetics. The respiratory volume and rate were adjusted to keep the percentage of CO₂ in the expired air between 4.0 and 4.5%.

Mydriasis and cycloplegia were induced by the topical application of atropine (1%) eye drops. Appropriate focus was achieved by means of hard contact lenses, which brought into focus the surface of a 57-cm-diam translucent hemispheric screen centered on the eye contralateral to the cerebral hemisphere to be studied. The positions of the area centralis, main blood vessels, and blind spot were projected onto the screen by means of a fiberoptic light source (Pettigrew et al. 1979) and checked every 2–3 h during the experiment. The position of the horizontal meridian (HM) and the vertical meridian (VM) of the visual field were defined according to the relationship between retinal landmarks such as the blind spot and the ganglion cell distribution (DeBruyn et al. 1980).

First, the HM was defined as a line passing through the area centralis and parallel to the “visual streak.” Because the visual streak in Galago cannot be visualized directly, the estimate was a line connecting the center of the area centralis (which was directly visualized) to a point 5° below the blind spot, corresponding to the average distance between the center of this retinal landmark and the axis of maximal ganglion cell distribution in flat-mounted retinas (DeBruyn et al. 1980). The VM was then defined as perpendicular to the HM through the center of the area centralis. As shown in Fig. 3C, these estimates of the axes of coordinates of visual field result in a distribution of receptive fields near the V1/V2 border that resembles the pattern observed in other species with developed vision, including cat, flying fox, and New World monkey (for reviews, see Frische and Rosa 1996; Payne 1990); the receptive fields near the V1 border invade the ipsilateral hemifield to a small extent, and the amount of invasion is similar in the upper and lower quadrants of the visual field.

Parylene-coated tungsten microelectrodes (with an exposed tip of 10 µm) were inserted in parasagittal rows of penetrations. Electrode penetrations were usually separated by 500–1,000 µm along the anteroposterior dimension, and were marked with electrolytic lesions. The response fields of single units and small unit clusters were mapped by correlating the stimulation of specific portions of the visual field with increments of the neural activity. Visual stimuli comprised luminous white spots and bars of variable size and luminance, moved or flashed on the surface of the screen by means of a hand-held projector. Receptive fields were drawn as rectangles parallel to the axis of best orientation, or as ovals if this axis could not be determined. Receptive field size was calculated as the square root of the receptive field area.

RESULTS

In the first part of this section we summarize the cortical architecture of areas V1, V2, and prostriata as revealed by both cytochrome-oxidase- and myelin-stained material. We then describe the extent and general topography of the visual field representation in V1 of Galago and demonstrate the detailed visuotopy of this area in both serial sections and “unfolded” bidimensional reconstructions. We also present a quantitative analysis of the variation of receptive field size and magnification factor in different parts of V1, and use these data to calculate the point image size in striate cortex. Finally, we describe some aspects of the organization of cortex surrounding V1, including the visuotopic organization of V2 and cortex rostral to dorsomedial V2.

Histological characteristics of areas V1, V2, and prostriata

The extent of V1 in galagos was defined both histologically and physiologically, and was consistent between ani-
In V2, the supragranular and granular layers together stain rather homogeneously, and more intensely than the infragranular layers. An infragranular band of slightly increased cytochrome oxidase reactivity is also apparent in V2, due to the presence of intensely stained cells in layer 5 (Fig. 1A). The rostral border of V2 is not clearly defined in cytochrome-oxidase-stained parasagittal sections. Area prostriata presents an overall light staining, being also noticeably thinner than either V1 or V2 (Fig. 1B).

In myelin-stained tissue, V1 is characterized by clearly separated inner and outer bands of Baillarger (Fig. 2A). In contrast, in V2 a dense plexus of coarse bundles of radial fibers runs without interruption between layers 3 and 6, obscuring the cleft between the bands of Baillarger. This characteristic is a reliable criterion for separation of this area from V1 (Fig. 2B). In addition, it allows a distinction between V2 and the visual areas rostral to it in ventral, lateral, and dorsolateral extrastriate cortices. Although the areas rostral to V2 vary in terms of density of myelination (Krubitzer and Kaas 1990, 1993), they can all be distinguished from V2 by the presence of a less myelinated band inserted between the inner and outer bands of Baillarger (Fig. 2C).

**Location and overall organization of V1**

V1 occupies the caudal pole of the dorsolateral surface of occipital cortex, the underlying ventral surface adjacent to the tentorium, portions of the caudal cortex along the midline fissure, and both banks of the calcarine sulcus (Fig. 3, A and B). Within this area there is a single, continuous representation of the entire contralateral hemifield, up to eccentricities of 50–60° along the VM and 100° along the HM (Fig. 3C). The representation of the area centralis occupies the lateral edge of the portion of V1 exposed on the dorsolateral cortex (Fig. 3A). The representation of the visual field around the VM forms most of the border of V1 (Fig. 3, squares), and the representation of the HM (Fig. 3, circles) bisects V1 in such a way that the representation of the lower quadrant is dorsal to the representation of the upper quadrant. Most of the cortex exposed on the surface of the occipital lobe corresponds to the representation of the lower visual quadrant, up to eccentricities of 15–20°. Although parts of the representation of the upper quadrant near the HM are also accessible on the caudal pole of the occipital lobe, our results indicate that most of the paracentral (5–20° eccentricity) upper quadrant representation is located on the tentorial surface of the brain (Fig. 3B). The part of V1 located along the walls of the interhemispheric fissure represents parts of the periphery of the lower quadrant, near the VM, up to ~40°. All of the remaining portions of the visual field periphery are represented along the banks of the calcarine sulcus. Although the ventral bank of this sulcus is entirely dedicated to the representation of parts of the upper visual field periphery, the dorsal bank represents portions of both the upper and lower visual quadrants (Fig. 3B). The most rostral part of V1, in the calcarine sulcus, represents the perimeter of the contralateral visual field (Fig. 3, triangles).

**Visuotopic organization of V1**

Figure 4 illustrates results obtained in one animal in which recordings covered most of the cortex within V1. Figure 4,
A–F, shows the correlation between the positions of recording sites in parasagittal sections (A being the most lateral, F the most medial) and the visual field coordinates (azimuths and elevations) of the corresponding receptive fields. Given that the electrode penetrations were many millimeters long, they rarely could be reconstructed within a single 50-μm section; each of the parasagittal sections illustrated in Fig. 4 is, therefore, a composite view of several adjacent sections.

Considering first the cortex exposed on the dorsolateral and tentorial surfaces of the brain, the analysis of parasagittal sections reveals a clear mediolateral gradient in the representation: the most lateral sites (Fig. 4, A1–A4) correspond to the most central receptive fields, and eccentricity gradually increases as the electrode is moved medially (Fig. 4, B1–B21, C1–C16, and D1–D18). The most peripheral receptive fields of neurons on the dorsolateral and tentorial surfaces (Fig. 4, E1 and D19, respectively) represent eccentricities of ~20°. At the same time, for each mediolateral level, the receptive fields gradually move from the lower to the upper visual field as the electrode tip is moved ventrally (e.g., Fig. 4, B1–B21).

Comparison of receptive fields recorded in electrode penetrations close to the midline (Fig. 4F) reveals a gradual displacement toward the periphery of the lower quadrant as the electrode is moved rostrally. A similar centroperipheral trend characterizes the recording sites located in the calcarine sulcus: at each mediolateral level, the most caudal sites represent the midperiphery of the visual field (30–45°, e.g., Fig. 4, B39, C29, and D35), and the most rostral sites always correspond to the representation of the temporal crescent of the visual field (Fig. 4, B55–B59, C40, and D47–D48). The regions of the visual field represented on the banks of the calcarine sulcus include the entire periphery of the upper quadrant (lateral and ventrally in the sulcus, e.g., Fig. 4, A, B, and C17–C27) as well as the portions of the lower quadrant periphery away from the VM (medial and dorsally in the sulcus, e.g., Fig. 4, C28–C40, D36–D48, E14–E18, and E31–E33).

These trends can be more readily visualized in graphically unfolded views of V1 generated from the contours of individual parasagittal sections. Three steps in the generation of a visuotopic map are illustrated in Fig. 5 (corresponding to the data shown in Fig. 4). In Fig. 5A, the contours of individual parasagittal sections are aligned and properly spaced, to reconstruct the approximate shape of V1. As an aid to orientation, different regions of V1 are indicated by different shading. Note that the anteroposterior axis of the cortex within the calcarine sulcus is inverted relative to Fig. 4, as a consequence of the process of “opening” the banks of this sulcus (the rostral tip of the calcarine sulcus now occupies the far right of the drawings). As shown in Fig. 5A, V1 in Galago is pear shaped, and not as elongated as in simian primates (e.g., Fritsches and Rosa 1996). The reconstructed surface area of V1 was 205 mm² in this animal and 218 mm² in the other animal with extensive recordings in V1 (these estimates are correct only within 20–30% because of the error inherent in the graphic unfolding method) (Gattass et al. 1987; Van Essen and Maunsell 1980). The portion of V1 that is directly accessible on the dorsolateral and caudal surfaces of the occipital lobe only corresponds to ~20% of this area. In Fig. 5, B and C, the recording sites are ...

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**Fig. 2.** Digital video frames illustrating myelin patterns and transitions between V1, V2, and visually responsive ventral cortex rostral to V2 (V2). Caudal is right, dorsal is up. A: myelin pattern of V1. Parts of infragranular layers are sectioned tangentially in ventral part. B: oblique section through border between V1 and V2 (at a level similar to that of section in Fig. 4D), illustrating disappearance of less myelinated “cleft” between inner and outer bands of Baillarger, moving from V1 to V2. C: crossing rostral border of V2, supragranular layers become relatively broader, and “matted” appearance of V2 in myelin stains gives way to areas where bands of Baillarger are well separated. Arrows: approximate location of borders of V2. Scale bar in A: 0.5 mm.
FIG. 3. Summary of visuotopic organization and extent of V1 in Galago garnetti. A and B: dorsolateral and medial views of a left hemisphere in which banks of calcarine sulcus have been separated. Extent of V1 is indicated in white. Visuotopy of this area is illustrated according to following symbols (see inset): black squares, representation of vertical meridian (VM); white circles, representation of horizontal meridian (HM); thin continuous lines, isoeccentricity contours (5, 10, 20, 40, 60, and 80°); dashed lines, 2 additional isopolar contours (−45 and +45°). C: projection of all receptive fields recorded in V1 of 4 animals onto an equatorial-azimuthal chart, illustrating extent of visual field represented in V1 on 1 side of brain. Each division of chart corresponds to a 10° interval of azimuths and elevations. Black circle: location of blind spot.
FIG. 4.  A–F: relationship between receptive fields and recording sites at 6 parasagittal levels (A is most lateral, F is most medial) in V1 of 1 animal ("Piglet"). A–F each consist of a parasagittal section where recording sites are indicated (left), a diagram of central portion of visual field illustrating receptive fields of neurons located on superficial cortex (middle), and a chart of entire hemifield illustrating receptive fields of neurons recorded on banks of calcarine sulcus (right). Portion of visual field magnified in middle is indicated by gray box on right. Arrows and dashed lines on sections: borders of V1. CiS, cingulate sulcus; IpS, intraparietal sulcus; LaS, lateral sulcus; P, prostriata.
indicated according to the symbols used in Fig. 4, together with the observed value of the receptive field center eccentricity (Fig. 5B) and polar angle (Fig. 5C). On the basis of their position in the bidimensional reconstruction, isoeccentricity (Fig. 5B) and isopolar (Fig. 5C) lines are drawn, summarizing the visuotopy of V1.

In the map of eccentricities (Fig. 5B) it can be observed that there is an emphasis on the representation of central
FIG. 5. Reconstruction of visuotopy of V1 in “flat” representations of cortical surface. A: bidimensional reconstruction of shape of V1, based on alignment of contours of layer 4 from a series of parasagittal sections (Van Essen and Maunsell 1980). To aid in localization, different regions of V1 are shaded in different tones of gray. B: map of eccentricity representation. Value of receptive field center eccentricity corresponding to each recording site illustrated in Fig. 4 is indicated (different symbols indicate different parasagittal levels illustrated in Fig. 4). On the basis of these values, isoeccentricity lines (5, 10, 20, 40, 60, and 80°, left to right) were interpolated by eye to fit individual observations. C: map of polar angle representation, illustrating 5 interpolated isopolar lines (−60°, −30°, HM, +30°, and +60°, from dorsal to ventral in cortex).
vision, with approximately half of the surface of V1 being dedicated to the representation of the central 30° of the visual field. The map of V1 in Galago seems to depart significantly from that expected on the basis of a logarithmic-conformal map (Schwartz 1977) in that a geometric progression of isoeccentricity contours does not result in lines that are separated by roughly equal distances in the cortex. For example, the distance between the 10° and 20° isoeccentricity lines is smaller than that between the 20° and 40° lines, and the distance between the 20° and 40° isoeccentricity lines is smaller than that between the 40° and 80° lines. The map of polar angles (Fig. 5C) suggests that for a given eccentricity there is relatively more magnification of those parts of the visual field near the HM than those close to the VM (see also Fig. 8). In addition, the region of V1 dedicated to the representation of the periphery of the upper quadrant is larger than that representing the periphery of the lower quadrant. Very similar results were obtained in the other animals, which had less complete sampling of the visual field representation (a summary map of another case is illustrated in Fig. 8, bottom).

Quantitative analyses

RECEPTIVE FIELD SIZE. It is clear from Fig. 4 that there is a marked increase of multiunit receptive field size from the central to the peripheral representation in V1 of Galago. For example, the sizes of receptive fields within the central 5° ranged from 0.8° to 2.7°, whereas in the far periphery (>60° eccentricity) they ranged from 4° to 20°. This trend is quantified in Fig. 6, which illustrates the receptive field size (square root of receptive field area) as a function of eccentricity for one animal. A power function (described by Eq. 1), calculated on the basis of a model II (principal axis) regression, provides a reasonable description of the relationship between eccentricity and receptive field size in V1

\[
RF \text{ size (degrees) } = 0.57 \times \text{eccentricity}^{0.60}
\]

(1)

It is noteworthy that the receptive field sizes that we find here are comparable to those found in a single-unit recording study (DeBruyn et al. 1993).

CORTICAL MAGNIFICATION FACTOR. The linear cortical magnification factor (CMF) was calculated as the ratio of the distance between two points in the cortex (in mm) and the angular distance between the centers of the corresponding receptive fields (in deg) (Daniel and Whitteridge 1961). We concentrated our analysis on the case that yielded the most complete sample of V1 (Fig. 7). Distances between recording sites were measured in unfolded reconstructions. Only pairs of recording sites separated by >1 mm and <3 mm were used in the analysis. As implied by the surface reconstructions, statistical analysis reveals a clear relationship between eccentricity and CMF in V1 (r = 0.88). Unlike in monkeys (Gattass et al. 1987; Van Essen et al. 1984), no difference was found between the CMF measured along different axes of V1 of Galago (e.g., parallel vs. perpendicular to the VM representation). Equation 2 describes the power function that best fits the data, calculated with the use of a principal axis regression technique

\[
CMF (\text{mm/deg}) = 2.36 \times \text{eccentricity}^{-0.34}
\]

(2)

A power function, although providing a reasonable description of the variation of CMF in the paracentral and peripheral representations, cannot adequately account for the variation of CMF within the central visual field (a map generated on the basis of such a function would have an infinitely large central representation). One proposed strategy to deal with this problem is to add a constant to the
Fig. 7. Cortical magnification factor (CMF), calculated between pairs of recording sites in V1, as a function of mean eccentricity of centers of receptive fields recorded at these sites in 1 animal (Piglet). Thick continuous line: best-fitting power function calculated on basis of a principal axis regression. Coarse dashed line at left: estimate of magnification factor in central representation ( \( \overline{\text{CMF}} \approx 0.7 \) ), which includes a correction factor based on size of representation of area centralis in V1 cortex (see Eq. 3).

**Visuotopic organization of V2**

The visuotopy of the cortex immediately rostral to V1 demonstrated a homologue of V2 similar to that described in other primates (Allman and Kaas 1974; Gattass et al. 1981; Rosa et al. 1988b). Although corresponding in general with the visuotopic organization of V2 described in simians, the organization of this area in galagos (summarized in Fig. 10) does have some distinctive features. First, V2 is relatively small and narrow. In the two animals with extensive recordings, the surface area of V2 was 63 and 68 mm\(^2\) (in both cases corresponding to 31% of the surface area of V1), with a maximum width of \( \approx 3 \) mm. Second, although in simians V2 has been described as wrapping around V1 almost completely (Allman and Kaas 1971; Rosa et al. 1988b), in *Galago* a relatively large portion of the perimeter of V1 in the calcarine sulcus is formed by less myelinated cortex (including area prostriata; Allman and Kaas 1971; Gattass et al. 1987; Sanides 1972). Third, the data suggest that the line of “field discontinuity” in the visual field representation in V2 is tilted relative to the HM. Whereas the dorsal portion of V2 represents most of the lower visual field, the ventral portion represents the entire upper quadrant plus a portion of the lower quadrant adjacent to the HM.

**Point image size.** The point image size, i.e., the linear dimension of the cortical locus of all neurons representing a point of the visual field, measured parallel to the cortical layers (McIlwain 1976), was estimated as the product of the average receptive field size and magnification factor for a given eccentricity (Gattass et al. 1987; Van Essen et al. 1984). Although this calculation only provides a minimum estimate of point image size for a given eccentricity (by virtue of not taking into consideration the receptive field scatter and the sizes of the largest receptive fields), the resulting function nonetheless accurately reflects the variation of point image size in different portions of a visual area (Fiorani et al. 1989; Rosa and Schmid 1995a). As shown in Fig. 9, the point image size in V1 of *Galago* varies little with eccentricity, from \( \approx 1 \) mm (in the representation of the central 10°) to 0.8 mm (in the representation of the far periphery of the visual field).
dorsal bank of the calcarine sulcus to a small extent (Figs. 10 and 11), in close agreement with the architectonic delimitation of dorsal V2 by Preuss and Goldman-Rakic (1991a). There is a clear mediolateral gradient of eccentricity representations, from lateral (central representation) to medial (peripheral representation), and nearly half of the surface area is dedicated to the central 20° of the visual field. The border between V2 and V1 is formed by the representation of the VM (e.g., Fig. 11, fields 6, 14, 21, and 28–30). In general, the recorded receptive fields move away from this meridian as the electrode is moved rostrally. However, the receptive field sequences are rather irregular, and caudal-to-rostral sequences of recording sites often reveal partial reversals near the HM representation (e.g., Fig. 11, fields 16–20 and 24–27). These partial reversals, as well as changes in receptive field size and architectonic pattern, suggest that a large portion of the rostral boundary of V2 with another visually responsive area in dorsal cortex corresponds to a representation of the lower visual field near the HM, rather than corresponding exactly to the representation of the HM as in simians. For example, in Fig. 12 we illustrate results obtained in rows of penetrations that crossed the border between V2 and the presumptive dorsomedial visual area (DM) (Krubitzer and Kaas 1990) (also referred to as the dorsal visual area by Allman et al. 1979) at different mediolateral levels. In each case, the receptive fields of neurons recorded at the most rostral sites within V2 are centered below the HM, rather than coinciding with this meridian (Fig. 12, A8, B5, and C7).

Our observations of ventral extrastriate areas are also consistent with the presence of V2, similar to what has been observed in simians (Fig. 13). The results demonstrate a systematic representation of the entire upper quadrant, extending across the tentorial surface and invading the lower bank of the calcarine sulcus to a small extent. In addition, rows of penetrations crossing from V2 into adjacent areas demonstrate that the rostral border of ventral V2 corresponds to a representation of a portion of the lower quadrant near the HM rather than to the HM itself (e.g., Fig. 13, fields 18–20, 31–32, and 41). The sizes of the receptive fields of neurons located in this narrow strip of lower quadrant representation are similar to those recorded in more caudal portions of dorsal and ventral V2 (Fig. 14). These observations lend support to the suggestion that the line of field discontinuity in V2 is tilted relative to the HM. As shown in Figs. 10 and 13, V2 is narrowest (just under 1 mm) at the representation of the center of gaze. However, sequences of recording sites crossing dorsoventrally at this point yield no evidence for a discontinuity between the dorsal and ventral parts of V2 (Fig. 13, fields 1–7, 8–16, and 21). These data suggest that the field discontinuity in V2 corresponds to an eccentricity of <5° (Fig. 10).

In simian primates, the representation of visual field in V2 is anisotropic: a similar distance in the visual field (in deg) is represented by a distance twice as long in the cortex parallel to the V1/V2 border as perpendicular to this border (Rosa et al. 1988b). This “stretching” of the visual field map has been correlated with the presence of functionally distinct stripes in V2, first revealed by cytochrome oxidase histochemistry, that run perpendicular to the V1/V2 border (Roe and T’so 1995; Tootell et al. 1983). Because cyto-

**FIG. 8.** Summary visuotopic maps of V1 in Galago. Top: diagrammatic map based on power function describing relationship between cortical magnification factor and eccentricity (Fig. 7). Middle and bottom: summaries of data in 2 animals. Animal “Linda” (bottom) lacked recording sites near occipital pole, and therefore there is a gap in representation where visuotopic coordinates could not be estimated. Symbols used in representation of visuotopic coordinates are similar to those employed in Fig. 3, and are indicated in inset.
chrome oxidase stripes are not present in *Galago* V2 (Condo and Casagrande 1990), we were interested in determining whether the representation in V2 of this species would also be anisotropic. Therefore the CMF of V2 was calculated for two animals on the basis of interpolated visuotopic maps of both dorsal and ventral V2 (Fig. 15). Different power functions were fitted to data measured between sites located along imaginary lines parallel to the V1/V2 border (CMF$_{pl}$; Eq. 4) and perpendicular to this border (CMF$_{pd}$; Eq. 5).

$$\text{CMF}_{pl} (\text{mm/deg}) = 2.05 \times \text{eccentricity}^{-0.80}$$

$$\text{CMF}_{pd} (\text{mm/deg}) = 0.83 \times \text{eccentricity}^{-0.80}$$

This analysis reveals that, despite the absence of cytochrome oxidase stripes, there is an anisotropic representation of the visual field in *Galago* V2, which is as marked as that observed in simians. At all eccentricities, the magnification factor parallel to the V1/V2 border is over twice that perpendicular to this border. It should also be noted that the exponents of Eq. 4 and 5 are identical to that of the CMF function of V1 (Eq. 2). Thus, at least to a first approximation, the emphasis on central vision is similar in areas V1 and V2.

**Evidence for DM**

Recordings in dorsomedial cortex immediately rostral to V2 resulted in rows of recording sites corresponding to systematic shifts in receptive field positions (Fig. 12) in the location of the architectonically defined DM (Allman and Kaas 1975; Krubitzer and Kaas 1990). Because of histological artifacts related to the large number of long electrode penetrations in this region, we were unable to reconstruct the myeloarchitectonic borders of DM in the same animals. Nonetheless, the rostral border of DM is well defined physiologically in the anesthetized galago as a transition between neurons with vigorous visual responses and sharply defined receptive fields (in DM) and neurons that are harder to drive and that tend to have larger receptive fields (Fig. 16). The presumptive boundaries of DM illustrated in Figs. 12 and 16, which are based on the physiological criterion, are in good agreement with histological estimates in other animals (Preuss and Goldman-Rakic 1991a). Neurons in the presumptive DM have receptive fields that are on average larger than those for neurons in V2, although there was a considerable overlap between the samples obtained in these areas. Laterally in DM, neurons have receptive fields that include the center of gaze and the central lower quadrant, sometimes invading the ipsilateral hemifield (Fig. 12A). More centrally in this region, receptive fields cover paracentral portions of the lower quadrant, near the border with V2, and of the upper quadrant rostrally (Fig. 12B). Near the midline, the receptive fields extend to $\approx 60^\circ$ from the center of gaze, including portions of both quadrants (Fig. 12C). Thus the cortex within our current estimate of the border of DM encompasses a representation of the entire upper quadrant, plus the lower quadrant near the HM.

We have naturally wondered whether DM is an area with incomplete representation of the visual field, or whether the “missing” portion of the lower quadrant representation (including the periphery near the VM) is located outside the current estimated boundaries. The data clearly indicate that the lower quadrant periphery is not represented medial to the estimated boundary. Instead, the cortex medial to DM along the midline re-represents the upper quadrant, with large receptive fields (Fig. 16, fields 10–15). In addition, the cortex immediately lateral to the presumptive DM, dorsolaterally, has neurons with smaller receptive fields, which systematically represent the central lower quadrant (Fig. 16, bottom right), confirming the results of Allman et al. (1979) in the lesser galago (*Galago senegalensis*). Finally, as mentioned above, most of the cortex along the rostral border of DM has neurons with very different response properties in comparison with DM neurons (including large receptive fields, Fig. 16, fields 5–9 and 16–20). The exception, illustrated in Fig. 16 (fields 1–4), is a small region of cortex.
located rostrolateral to the presumptive DM border. This region, which was observed in one animal, includes a compressed representation of the lower quadrant periphery that nearly exactly complements the visual field representation in DM, with receptive fields of appropriate size for DM. Thus, although this result demands confirmation with additional experiments, it is possible that DM in galagos forms a second-order representation of the entire visual field, with a field discontinuity running ventral to the HM.

**Area prostriata**

As mentioned above, a relatively large proportion of the perimeter of V1 (25% and 27%, in the two most extensively studied animals; see Figs. 11 and 13) was formed by agranular, poorly myelinated cortex. In terms of both location and architectonic appearance, this cortex resembles area prostriata (Allman and Kaas 1971; Sanides 1972) and the splenial visual area of cats (Kalia and Whitteridge 1973). Similar to the findings of MacLean (1966) in the squirrel monkey, we were able to evoke visual responses from some neurons in this area (3 of 12 recording sites). Although the sample was too small to allow any conclusions regarding visuotopy, we found that the receptive fields of neurons in this area include both the central and the peripheral visual field and are large (4–6 times the diameter) in comparison with those at a comparable eccentricity in V1.

**DISCUSSION**

Our main finding is that the visuotopic organization of V1 and V2 in the prosimian galago shares many of the features described for simian primates, suggesting that these characteristics existed in the last common ancestor of all primates. Both V1 and V2 are continuous areas, each containing a single representation of the entire visual field, with the VM at their common border. As in other primates, V1 forms a continuous (1st-order) representation with an expanded representation of central vision, whereas V2 contains a split (2nd-order) representation of the visual field. The
Fig. 11. Correlation between positions of recording sites and receptive fields in dorsal V2. Top left: bidimensional reconstruction of caudal cortex in 1 animal, illustrating extent of V2, MT, and area prostriata (estimated borders are indicated in gray). Bottom left: region enclosed by box, shown magnified, plots location of 39 recording sites in dorsal part of V2. To facilitate presentation, these sites were connected to form 7 arbitrary sequences, from lateral (1–5) to medial (37–39) in cortex. Other plots: location of receptive fields recorded from neurons at each of these sites, in diagrams of central (top row) and peripheral (right: bottom row) visual field. STS: superior temporal sulcus.
FIG. 12. A–C: sequences of receptive fields and recording sites crossing border between V2 and presumptive dorsomedial visual area (DM) at different mediolateral levels. A–C each illustrate a dorsal view of brain (left) indicating recording sites and estimates of borders of V1, V2, and DM. Corresponding receptive fields in V2 (middle) and DM (right) are illustrated relative to HM and VM (thick lines) and blind spot (gray).
results also provide evidence for a homologue of area DM located anterior to V2. In addition to these findings, our data reveal some differences in the organization of V1 and V2 of galagos in comparison with simian primates. The significance of these similarities and differences in visual field representation between galagos and simians is considered in more detail below.

V1: quantitative analyses

One of the main discussions in the current literature regarding the visuotopy of V1 refers to whether or not the extent of the representation of different sectors of the visual field in this area is directly proportional to the ganglion cell density in different parts of the retina. Although several studies have proposed that such a direct relationship exists (e.g., Schein and De Monasterio 1987; Wässle et al. 1989), many others have concluded that there is a preferential representation of the fovea (or area centralis) in V1, such that the number of V1 cells representing the central visual field is much larger than that expected on the basis of the distribution of ganglion cells (Myerson et al. 1977; Perry and Cowey 1985, 1988; Rosa and Schmid 1995a; Silveira et al. 1989, 1993). The difficulty in answering this question is compounded by two main factors: the presence of a population of displaced amacrine cells in the ganglion cell layer, whose density varies less steeply with eccentricity than that of the ganglion cells, and the displacement of ganglion cells toward the foveal rim, with the consequent difficulty in determining the numbers of ganglion cells that are connected to foveal photoreceptors (Silveira et al. 1993; Wässle et al. 1989; Wilder et al. 1996). As pointed out previously (Silveira et al. 1993), nocturnal primates may be good animal models in which to study this question, given that the absence, or...
rudimentary nature, of the fovea in these species greatly facilitates the quantification of ganglion cell densities in the central visual field. A previous study in G. garnetti (DeBruyn et al. 1980) (note that in that study this species was referred to as Galago crassicaudatus) has demonstrated that the peak density of neurons in the ganglion cell layer is ~11,200 cells per mm², and that there are <500 cells per mm² in the nasal periphery of the retina (90°–100° eccentricity). These estimates yield a centroperipheral gradient of 22.4:1, which contrasts sharply with a gradient of 250:1 for the areal CMF (estimated as the square of the linear magnification factor) in V1 in the same range of eccentricities. Such a large discrepancy between the two gradients is difficult to reconcile with the view of a proportional visuo-tectal relationship along the retinogeniculocortical pathway. The presence of displaced amacrine cells was not recognized at the time of the study of DeBruyn et al. (1980), and, as demonstrated by others (Silveira et al. 1993; Wässle et al. 1989; Wilder et al. 1996), the density profile of such cells may be very different from that of ganglion cells. Nonetheless, even allowing for a hypothetical increase in the proportion of amacrine cells from center to periphery, it is unlikely that the visuotopy of Galago V1 will directly reflect the retinal ganglion cell distribution. For example, assuming that

FIG. 15. Cortical magnification factors in V2, measured between points in cortex located along imaginary lines parallel (□; thick continuous line) and perpendicular (▲; thick dashed line) to V1/V2 border.
FIG. 16. Receptive fields of neurons recorded in areas surrounding DM in 2 animals. Top row: dorsal views of caudal neocortex, illustrating recording sites within presumed borders of DM (●) and outside DM (○). Second row: extent of visual field represented within estimated boundaries of DM. Third row: receptive fields of neurons recorded in sites immediately rostral to estimated border of DM. Four receptive fields that may be part of peripheral lower quadrant representation of DM (fields 1–4) are indicated in gray. Bottom left: receptive fields of neurons recorded in sites immediately medial to DM. Bottom right: extent of visual field represented in cortex immediately lateral to DM.
in the area centralis displaced amacrine cells form ~5% of the neuronal population of the ganglion cell layer (as in both diurnal and nocturnal simians) (Silveira et al. 1993; Wässle et al. 1989; Wilder et al. 1996), then they would need to form >90% of the cell population in peripheral retina to yield a direct proportional relationship between retina and cortex. Thus our data suggest that galagos resemble simian primates in having a preferential representation of the area centralis in V1, in agreement with Azzopardi and Cowey’s (1993) anatomic demonstration based on transneuronal transport of retrograde tracers from the cortex to the retina. It does remain possible, however, that the visuotopy of V1 will reflect the density of a particular type of ganglion cell (Malpeli et al. 1996; Schein and de Monasterio 1987).

The analysis of the cortical point image size (Fig. 9) reveals that the steep decrease of CMF with increasing eccentricity is almost exactly counterbalanced by an increase in the average receptive field size. Thus in Galago, as in the flying fox (Rosa et al. 1993a), a nearly constant number of V1 neurons is used to process stimuli located anywhere in the visual field. One interpretation of these data is that V1 is formed by physiologically modular units of similar size throughout the visuotopic map; these are likely to include columns responsive to both eyes and at least a full cycle of orientation columns (Hubel and Wiesel 1974). The actual point image size curve obtained for one individual with extensive mapping of V1 suggests a slight decrease (from 1.0 to 0.8 mm) of the point image size between the central and peripheral representations. It is interesting to observe that this is paralleled by changes in the distribution of cytochrome oxidase blobs in the upper layers. In Galago, as in most other primates (see Preuss and Kaas 1996 for review), V1 is characterized by a histologically defined regular modular organization, evidenced by the cytochrome oxidase blobs of the upper cortical layers (Fig. 1). There are more blobs per mm² within the calcarine sulcus than on the dorsolateral surface (2.9 vs. 2.6 blobs/mm²) (Condo and Casagrande 1990), and this may indicate that the histologically defined modules are slightly smaller in the peripheral representation in comparison with the central representation. Thus the physiological and histological data converge to support the notion of a slight decrease in module size from central to peripheral V1. The slight variation of module size in Galago V1 is reminiscent of the organization demonstrated in New World primates such as the Cebus monkey and squirrel monkey (Horton and Hocking 1996; Rosa et al. 1991). In the macaque, early reports have suggested a marked variation of module size with eccentricity on the basis of both blob distribution and ocular dominance hypercolumn width (Le Vay et al. 1985; Livingstone and Hubel 1984). However, these findings have been challenged (Florence and Kaas 1992; Gattass et al. 1995), and it is presently unclear to what extent the macaque departs from the pattern observed in other primates.

V1: comparative analysis

On the basis of both paleontological data and studies of molecular evolution, primatologists have suggested that the lineages leading to present-day lorisiform prosimians (which include Galago) and simians have been separated at least since the Palaeocene epoch (55–66 million years ago) (Martin 1990). To put these estimates in context, it is worth remembering that the first evidence of simians of modern aspect has only been recovered from Eocene formations (~40 million years ago) (Beard et al. 1994; Godinot and Mahboubi 1992; Simons 1995). Yet, despite their remote common ancestry, previous studies have demonstrated that several aspects of the organization of V1 in galagos are similar to those observed in simians, including the pattern of termination of geniculocortical afferents relative to the cortical layers (Diamond et al. 1985; Florence and Casagrande 1987; Florence et al. 1983; Lachica and Casagrande 1992; Lachica et al. 1993), the architectonic distribution of cytochrome oxidase and calcium-binding proteins (Condo and Casagrande 1990; Johnson and Casagrande 1995), neuronal response properties (DeBruyn et al. 1993), and the pattern of projections to extrastriate cortex (Krubitzer and Kaas 1990, 1993; Symonds and Kaas 1978; Tigges et al. 1973). These data converge to suggest that the features of V1 organization common to lorisiform prosimians and simians were established early during primate evolution. The present study demonstrates that the visuotopy of Galago V1 is consistent with that observed in other mammals as far as the relative locations of central/peripheral and upper/lower visual field representations are concerned. Nonetheless, the data also demonstrate some characteristics that are intermediate between those of nonprimate mammals on one hand and simian primates on the other. These include the size of V1 relative to body size, the degree of emphasis on representation of central vision, and the topographic relationship between areas V1, V2, and prosstriata.

Many authors have suggested that primates as a group, and simians in particular, are highly dependent on vision for tele receptor. Attempts to define primates as a natural group distinct from other mammals commonly mention that “the visual sense is greatly emphasised” (Martin 1990; see also Polya 1957) and contain references to an “elaboration and perfection of the visual apparatus” (Le Gros Clark 1959). In agreement with this notion, a previous study (Rosa et al. 1993a) has demonstrated that the surface area of V1 of both simian and prosimian primates is significantly larger than that of nonprimates of similar body mass. The present data indicate that the surface area of V1 in G. garnetti (~200 mm²) is 2–3 times as large as would be expected in comparison with nocturnal nonprimates of similar body size, such as ferrets (Law et al. 1988) and hedgehogs (Kaas et al. 1970). Yet, despite the similarities in functional architecture, it is also apparent that V1 in the galago is relatively small in comparison with that of simians.

To what extent is the “intermediate” condition of V1 in galagos a reflection of adaptations to a nocturnal niche? A relevant comparison can be drawn with the nocturnal owl monkey, which is similar to G. garnetti in terms of body mass and density of retinal ganglion cells (DeBruyn et al. 1980). As in Galago, the striate cortex of owl monkeys differs from that of diurnal simians in a number of aspects, including a relatively small surface area (286–400 mm²) (Myerson et al. 1977; Tootell et al. 1985), which corresponds to about half of that in the similarly sized but diurnal squirrel monkey (Cowey 1964). Thus it is likely that adaptations to nocturnal lifestyles are a major factor determining the relative size of visual areas. For example, the size of V1 might be correlated with the number of ganglion cells projecting centripetally and with the convergence ratio along
the visual pathway, factors that may differ in nocturnal versus diurnal species. As demonstrated by studies comparing the retina and lateral geniculate nucleus of feral and domestic animals of the same genus, these characteristics can be regulated by developmental processes in such a way that vast quantitative changes can occur in a relatively small number of generations (Peichl 1992; Williams et al. 1993).

Given the possibility that the volume of visual areas is a labile character, in evolutionary terms, is there any reason to suspect that the relatively small size of V1 (and other areas) in Galago reflects a lesser degree of differentiation in comparison with simians? Although nocturnality is clearly an important element, the fact remains that presently published figures suggest that the surface area of V1 in the owl monkey is still \(\approx 40\%\) larger than that in the galago. Both the owl monkey and the galago are present-day representatives of what seem to be long-standing but independent lineages of nocturnal primates that can be traced back at least 15 million years (McCrosin 1992; Setoguchi and Rosenberger 1987). In view of the relative rapidity with which quantitative changes in the visual pathway can occur, it is unlikely that the differences between the owl monkey and the galago could be solely attributed to the fact that the owl monkey probably descended from diurnal ancestors (e.g., Hoffstetter 1982). In theory, it is possible that in the course of the evolutionary process leading to present-day simians further quantitative changes in the retinogeniculostriate pathway occurred, possibly related to the reduction in the olfactory apparatus and increasing reliance on vision among simians (Cave 1973; Le Gros Clark 1959; see Martin 1990 for a review), or that new neuronal circuits were added to simian striate cortex after the divergence from the prosimian lineages. Alternatively, given that the brains of simians tend to be larger overall than those of size-matched prosimians, the larger surface of visual areas may be pleiotropic, being the result of a developmental regulatory process affecting the entire brain rather than targeting the visual apparatus in particular. Morphometric studies of the retina and other visual structures in diurnal prosimians could help clarify these questions.

Analysis of the data also demonstrates that the emphasis on central vision representation in V1 of Galago is less marked than that observed in simians. One simple index for comparing this across different species is the proportion of the surface area of V1 dedicated to the representation of the central 10° of the visual field. Whereas in diurnal simians ranging from marmosets to humans this portion of the visual field occupies 50–60% of the surface area of V1 (Fritsches and Rosa 1996), in G. garnetti it corresponds to only 20–22% of the total V1 surface (Fig. 8). The proportion observed in Galago is even smaller than that reported in the nocturnal owl monkey (30%) (Allman and Kaas 1971), but approaches that found in nonprimates with well-developed vision such as cats (20%) (Tusa et al. 1978). It is interesting to observe that the peak resolution of the visual system of galagos is very similar to the cat’s, and about half of that in the owl monkey ( Bonds et al. 1987; Langston et al. 1986).

In primates (including galagos) the unfolded surface of V1 forms an ellipse with the major axis parallel to the representation of the HM (e.g., Daniel and Whitteridge 1961; see Fritsches and Rosa 1996 for a review). In these species, the representation of the VM in V1 bows out markedly, forming an “apex” that corresponds to the representation of the fovea or area centralis (e.g., Dow et al. 1985; Tootell et al. 1988). Although similar to other primates, the present data suggest that these characteristics are less marked in Galago than in simians, both diurnal and nocturnal (Fig. 17). In marked contrast, in most (if not all) nonprimates the longer axis of V1 is parallel to the VM, and the representation of the area centralis does not protrude toward V2 (Choudhury 1978; Hughes 1971; Kaas et al. 1970, 1972a,b, 1989; Law et al. 1988; Löwel 1987; Picanço-Diniz et al. 1991; Rosa et al. 1993a; Sousa et al. 1978; Tusa et al. 1978; Wagor et al. 1980). Several authors have correlated the overall shape of V1 with the functional architecture for ocular dominance, suggesting that the map tends to be elongated perpendicular to the preferred orientation of ocular dominance stripes (e.g., Anderson et al. 1988; Löwel et al. 1988; Rosa et al. 1988a, 1992; Tootell et al. 1982, 1988). Information on the layout of ocular dominance columns in galagos is still sketchy (Glendenning et al. 1976). However, given the shape of striate cortex and the isotropic CMF we observed in this area, one expects that a more complete study of these animals will reveal a system of bands less regularly oriented than that observed in simian primates.

The border relationship of V1 with V2 and area prostriata also differs markedly between simian primates and nonprimates (Fig. 17). In both nocturnal and diurnal simians, V2 wraps around V1 almost completely, with a relatively small area prostriata, located at the rostral end of the calcineurine sulcus, forming only \(\approx 10\%\) of the perimeter of V1 (Allman and Kaas 1971; Gattass et al. 1981, 1987; MacLean 1966; Rosa et al. 1988b; Van Essen et al. 1984). In contrast, in nonprimates, visually responsive “limbic” agranular cortex located medial to V1 may form over half of the perimeter (e.g., Kaas et al. 1989; Kalì a and Whitteridge 1973; McConnell and LeVay 1986; Rosa et al. 1995). Here again, the pattern revealed by our experiments suggests an intermediate condition in Galago: V2 forms about three-quarters of the perimeter of V1, with the remaining quarter being formed by area prostriata. Studies of interhemispheric connections in galagos reveal that projecting cells are distributed well within V1, much as in nonprimates, rather than being restricted to the immediate vicinity of the border of this area, as in simians (Beck and Kaas 1994; Cusick et al. 1984; Kennedy et al. 1986). This anatomic organization may be reflected in the presence of a representation of the ipsilateral hemifield near the VM, as revealed by our experiments (Fig. 3C).

**Organization of V2**

We propose that the visuotopic map in Galago V2 is a second-order representation of the visual field, featuring a field discontinuity that does not exactly correspond to the HM. Given the existence of other plausible interpretations of the data, we acknowledge that the issue of the exact location of the field discontinuity and anterior border of V2 should be considered a hypothesis to be tested by further studies. One reasonable alternative is that, as in simians (e.g., Allman and Kaas 1974; Gattass et al. 1981), the rostral border of V2 in galagos is coincident with the representation of the HM. Purely on the basis of the progression of the receptive field centers (Figs. 11–13), one might argue that...
the border of dorsal V2 could be placed more rostrally (e.g., around Fig. 12, A10, B8, and C10), leaving V2 directly abutting the upper quadrant representation in DM, and that the border of ventral V2 could be placed more caudally, therefore excluding the sector of lower quadrant representation. Although the myeloarchitecture of ventral cortex suggests that the sector of lower quadrant representation near the rostral border of V2 is part of V2, this is a narrow strip of cortex, and there is a degree of uncertainty in determining the border. Thus the histological evidence alone is not definitive. However, the sizes of the receptive fields of neurons recorded in this transitional region of lower quadrant representation are well within the range of those found in other parts of V2 (Fig. 14), and they are much smaller than the ones recorded in the more rostrally located area (which may be a homologue of the ventral posterior area). Furthermore, sequences of recording sites extending rostral to this transitional region reverted to re-represent the upper quadrant, rather than advancing further into the lower quadrant. Excluding the lower quadrant transitional region from ventral V2 would therefore leave us with a narrow area that only represents a sector of the lower visual field near the HM.

In view of these arguments, we believe that the balance of evidence in ventral cortex favors the interpretation that there is some lower quadrant representation forming the rostral border of ventral V2. This, combined with the (admittedly subtle) transitions in receptive field size between dorsal V2 and DM tending to occur below the HM, and the evidence on the organization of DM itself (see below), suggest the model of organization of V2 summarized in Fig. 10. The “split representation” tilted relative to the HM is reminiscent of the proposal of Gattass et al. (1988) for macaque V4. One feasible experiment to test these hypotheses would be to combine, in the same animals, electrophysiological recordings across the rostral border of V2 and examination of transitions in the laminar patterns of connections after tracer injections in other visual areas. If the hypothesis depicted in Fig. 10 is correct, one would expect that the sector of lower field representation that we propose to be part of ventral V2 will show the same laminar patterns of labeled cells and terminals as other parts of V2, and that the connectional boundary between V2 and DM will correspond to the representation of a point below the HM.

Another possibility is that the impression of a tilted field discontinuity at the rostral border of V2 is due to error in measurement, specifically, in our estimates of the location of the area centralis and the HM. For example, the eyes of cats rotate during paralysis, and a correction is necessary when measuring the polar angles of receptive fields relative to the screen (Cooper and Pettigrew 1979). Fortunately, there are good reasons to believe that this was not a serious factor in our experiments. Although there is some degree of error in interpolating the HM on the basis of the position of the area centralis and optic disk, any error would be similar for all visual areas. Yet the receptive fields recorded in V1 (often in the same penetrations) result in a visuotopic map in which the center of the visual field coincides well with the apex of the curvature of the border of V1 in lateral cortex and 2) there is no evidence of a large asymmetry between the representations of the upper and lower quadrants. If anything, the mild asymmetry we observed in V1 (the periphery of the upper quadrant being more magnified than that of the lower quadrant; Figs. 5 and 8), which may be due to rotation of the stimulated eye, would point to an error in the opposite direction. Furthermore, given the extent of the receptive fields recorded in V1 (Fig. 3C), a rotation of our frame of coordinates to bring the HM into alignment with the most ventral receptive fields recorded in ventral V2 (Fig. 13, fields 20, 31, and 41) would result in a large invasion of the ipsilateral hemifield in the upper quadrant representation and a lack of representation of the VM in the lower quadrant.

It is interesting to note that a field discontinuity oblique relative to the HM was also demonstrated in V2 of megachiropteran bats (Rosa et al. 1994), animals that are considered by many authors to be members of one of the possible sister groups of Order Primates (Pettigrew et al. 1989; Rosa and Schmid 1994; see also Fig. 1 of Preuss and Kaas 1996). A similar visuotopic pattern, with ventral V2 representing the entire upper quadrant plus parts of the lower quadrant, may also exist in other nonprimates, such as tree shrews and squirrels (for discussion, see Rosa et al. 1994). Thus the pattern whereby the upper and lower quadrants are almost perfectly segregated into ventral and dorsal V2, respectively, may be unique to simian primates (Allman and Kaas 1974; Cowey 1964; Gattass et al. 1981; Rosa et al. 1988b).
data in galagos also suggest a partial overlap between the sectors of the visual field represented in dorsal and ventral V2, as demonstrated for capuchin and marmoset monkeys (Rosa and Schmid 1995b; Rosa et al. 1988b).

Our results suggest that V2 in galagos is formed by a continuous belt of variable width, as in other primates. These electrophysiological data are consistent with previous architectonic studies in this and other prosimians (Krubitzer and Kaas 1990; Preuss and Goldman-Rakic 1991a; Preuss et al. 1993), but do not support the view of Sereno and Allman (1991), who proposed that V2 in Galago is formed by two islands of cortex separated by another area that represents both the upper and lower visual quadrants. Although the brief nature of the latter report makes a comparison of data difficult, we can propose at least two reasons why the impression of a physically segregated V2 may arise. First, V2 in Galago is very narrow at the representation of the area centralis (just 1 mm wide, or slightly less), and it is difficult to obtain a number of recording sites from this region unless long penetrations tangential to the lateral cortex are used (as in Fig. 13). This introduces an element of uncertainty in the interpretation of the data. Second, because of the fact that the field discontinuity in Galago V2 may be tilted relative to the HM, recordings from neurons in ventrolateral cortex, just over 1 mm from the V1 border, show evidence of receptive fields with centers below the HM. It would be natural to interpret these recordings as evidence of an additional visual area in ventrolateral cortex, except for the fact, as argued above, that this strip of cortex only represents a small part of the lower quadrant adjacent to the HM.

Analysis of CMF in Galago V2 revealed an anisotropy similar to that described in simian primates (Roe and T'so 1995; Rosa et al. 1988b). An elongation of the visuotopic map parallel to the V1/V2 border has also been demonstrated in other species lacking cytochrome oxidase stripes, such as the cat and flying fox, and therefore this seems to be a general feature of mammalian V2 (see Rosa et al. 1994 for review). It is possible that the anisotropy in V2 of nonsimian species is due to the presence of regular striplike functional modules that are not revealed by the cytochrome oxidase stain. For example, in the cat, transneuronal transport of anterograde tracers injected in the eye reveals a pattern of appropriately oriented stripes in V2 (Anderson et al. 1988).

**Homologue of DM?**

DM is a subdivision of extrastriate cortex first described in owl monkeys (Allman and Kaas 1975). Microelectrode recordings allowed a systematic map of the contralateral hemifield to be related to a strip of relatively heavy myelination located rostral to dorsomedial V2. More recently, the visual topography of DM has been demonstrated in a diurnal New World monkey, the marmoset (Rosa and Schmid 1995b), and anatomic criteria such as cortical architecture and the pattern of connections with V1 and the middle temporal area (Krubitzer and Kaas 1990, 1993; Vogt-Weisenhorn et al. 1995; Wager et al. 1975; Weller et al. 1991) have been used to support the hypothesis that DM is present in Old World monkeys and prosimians as well (Stepniewska and Kaas 1996).

A previous study of the cortex rostral to V2 in *G. senegalensis* (Allman et al. 1979) has proposed the existence of a dorsal visual area (area D) in the approximate location of DM. In agreement with this report, we found that the dorsomedial cortex includes representations of both the upper and lower contralateral quadrants (Fig. 12), and that the rostral border of this area is formed by a representation of the VM. However, in contrast with these previous results, our data suggest that much of the lower quadrant representation is located caudal to the upper quadrant representation, rather than being entirely lateral to it. Another difference is that the cortex along the banks of the interhemispheric fissure was included in the dorsal area of *G. senegalensis*, whereas our data indicate that this region forms a representation of the upper quadrant that is distinct from that of DM. At least in part, these discrepancies could be related to the different species used (note, for example, that *G. senegalensis* is 3–4 times smaller, in terms of body mass, than *G. garnetti*). On the basis of the fact that their recordings suggested an area where the lower quadrant representation was lateral to the upper quadrant representation, Allman et al. (1979) suggested that area D might be homologous to the medial visual area (area M) of owl monkeys. Given that subsequent research has demonstrated that there are significant interspecies differences in the visuotopic organization of DM (Rosa and Schmid 1995b; see Rosa 1997 for a review), the now-extensive anatomic evidence for a homologous DM in many primate species (see above), and the fact that we found another visually responsive area medial to the presumptive DM, it seems more likely that both DM and area M are present in galagos. If so, the argument is strengthened that DM is a visual area that emerged early in primate evolution and has been retained in most primates, including humans.

We also found that recording sites rostral to V2 in lateral cortex yield neurons with receptive fields centered in the lower quadrant, and that neurons at recording sites rostral to V2 in ventral cortex yield receptive fields centered in the upper quadrant (Figs. 13 and 16). Although it is clear that more data are needed, these results are compatible with the previous suggestion, made on the basis of anatomic connections, that there is a homologue of the dorsolateral area in lorisiform prosimians (Krubitzer and Kaas 1990; Preuss et al. 1993).

**Area prostriata**

The term “prostriata” was originally used by Sanides (1972) to describe a part of the cortex bordering V1 that is less granular and less myelinated than area 18. The marked difference in appearance of the two fields (Allman and Kaas 1971; Gattass et al. 1987) helped dispel the idea that V2 completely surrounds V1. Area prostriata of primates has long been thought to have visual functions (e.g., MacLean 1966; Sanides 1972), but compelling evidence has been difficult to obtain. Recent anatomic tracing experiments have demonstrated that prostriata projects to visual areas, including V1 and the medial superior temporal area (Rosa et al. 1993b; Sousa et al. 1991). Present results add to this story in that neurons in only a few sites in area prostriata respond to visual stimuli. Yet, visual responses were obtained, and the receptive fields did not correspond to the visuotopy of V2. In particular, neurons had receptive fields that included central vision, a feature incompatible with the visuotopy of
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