Color signals in retina and lateral geniculate nucleus of marmoset monkeys

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Abstract
Marmosets show sex-linked polymorphism of color vision, whereby all males and some females show dichromatic (“red-green color-blind”) vision based on two classes of photoreceptor sensitive to short or medium wavelength bands. Most female marmosets by contrast express three photoreceptor classes, one sensitive to short wavelengths and two classes in the medium-long wavelength sensitivity band. We used this ‘natural knock-out’ to study the organization of color vision pathways in primates. We review here results from our and other laboratories showing how the primordial dichromatic blue-yellow pathway is characterized by selective connections to short wavelength sensitive cones in the retina and that signals for blue-yellow color vision travel through an ancient part of the subcortical visual pathway called the koniocellular system. Signals serving red-green color vision by contrast are tightly linked to retinal circuits serving high-resolution spatial vision at the fovea and show little sign of specific patterns of connections with medium- and long-wavelength sensitive cones. Routine trichromatic color vision thus is based on converging signals from two quite distinct retinal and subcortical pathways. Keywords: color, neurophysiology, neuroanatomy, primate, evolution.

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Introduction
The collected papers in this volume are presented in honor of our long-standing mentor, friend and collaborator Barry B. Lee. We thank him sincerely for his valued and continued contributions to the fields of color science and visual neuroscience. This review paper concerns the nerve pathways carrying signals that serve color vision in primates with emphasis on data from experiments on marmoset monkeys.

Our concentration here is on a single species of New World monkey (Callithrix jacchus) that our laboratories have studied for over 20 years. In common with many other New World monkeys, marmosets show sex-linked polymorphism of cone photoreceptor opsin expression. Such species provide a “natural knock-out experiment” to study how variation in cone opsin expression changes the function of visual pathways. The methods used include light- and electron microscopy, immunocytochemistry, retrograde labelling and tract-tracing, and extracellular recording from single neurones in the afferent visual pathways. Recommended reviews in related topics are as follows: genetics and evolution of mammalian color vision (Jacobs, 1993; Nathans, 1999; Neitz & Neitz, 2011), retinal anatomy and physiology (Dacey & Packer, 2003; Field & Chichilnisky, 2007; Lee, Martin, & Grünert, 2010; Silveira et al., 2004; Wässle, 2004), and cortical mechanisms processing color (Conway et al., 2010; Gegenfurtner, 2003; Shapley & Hawken, 2011).

Basis of marmoset polymorphic color vision
Color vision requires photoreceptors sensitive to distinct spectral bands and on a brain that can use these photoreceptor signals to recover information about spectral reflectance of objects in the environment. Figure 1 illustrates the variety of photoreceptor types exhibited by marmosets (Tovée, Bowmaker, & Mollon, 1992; Travis, Bowmaker, & Mollon, 1992; Williams, Hunt, Bowmaker, & Mollon, 1992). All male marmosets show the common mammalian pattern, whereby two cone receptor pigments are expressed and can support dichromatic (“red-green color blind”) discriminations, roughly corresponding to the blue-yellow axis of routine trichromatic (“normal”) human color vision (Jacobs, 1993; Mollon, 1991). The short wavelength-sensitive (S or “blue”) pigment is coded on chromosome 7 and has a spectral peak close to 423 nm. One of three allelic gene variants on the X chromosome codes for opsin with peak sensitivity near 543 nm, 556 nm or 563 nm in
the medium-long (ML) wavelength range of the visible spectrum. Female marmosets carrying the same allele on each X chromosome show, in common with male marmosets, dichromatic color vision. Female marmosets carrying distinct alleles on each X chromosome express two ML pigments with spectral peaks separated by 7 nm (for the 556 nm/563 nm combination), 13 nm (for the 543 nm/556 nm combination), or 20 nm (for the 543 nm/563 nm combination) and can make trichromatic color discriminations (Tovée et al., 1992; Williams et al., 1992). Thus, in addition to studying differences between dichromatic and trichromatic color vision, the effects of varying degree of spectral separation between M and L pigments can be examined in marmosets.

Retinal pathways for colour signals: S cone pathways

The retinal pathways thought responsible for transmitting S cone signals to the marmoset brain are summarized in Figure 2. Signals from cone photoreceptors are transmitted to retinal output neurons (ganglion cells) through multiple classes of bipolar cells, which contact cones in the outer plexiform layer and contact ganglion cells in the inner plexiform layer (Boycott & Wässle, 1991). The major functional division of bipolar cells is into ON type bipolar cells (which are depolarized subsequent to increased photon absorption in photoreceptors), and OFF type bipolar cells, which are depolarized by decreased photon absorption in photoreceptors (Nelson & Kolb, 2003).

In whole mount retina preparations, the S cones and the blue cone bipolar cells can be labelled and their pattern of connections traced (Figure 2A, modified from Luo, Ghosh, Martin, & Grüner, 1999). Such preparations reveal that dendritic processes of blue cone bipolar cells (stippled outlines, Figure 2A) extend horizontally across the outer plexiform layer to reach individual S cone pedicles (blue outlines, Figure 2A). In this way, signals from the S cone array are transmitted to the blue cone bipolar array (Ghosh, Martin, & Grüner, 1997; Kouyama & Marshak, 1992; Kouyama & Marshak, 1997; Schein, Sterling, Ngo, Huang, & Herr, 2004).

In common with other ON-type bipolar cells, the blue cone bipolar cells express the sign-inverting metabotropic glutamate receptor MgluR6 and thus increased photon absorption in S cones leads to depolarization of blue cone bipolar cells (Nomura et al., 1994; Vardi, Duvoisin, Wu, & Sterling, 2000). Additional to the selective connections with the S cone array, the blue cone bipolar cells make a small number of “blind ending” dendrites with unknown connectivity. Some of these endings may contact ML class cones, but the exact connectivity has not been established (Kouyama & Marshak, 1992; Luo et al., 1999).

There is agreement that the vast majority of ON-type connections between bipolar cells and S cones are made with blue cone bipolar cells (Herr, Klug, Sterling, & Schein, 2003; Kolb, Goeke, Roberts, McDermott, & Gouras, 1997; Kouyama & Marshak, 1992; Luo et al., 1999). Thus, the spectral sensitivity of blue cone bipolar cells should be weighted to the S wavelength band. The pattern of connections between S cones and other bipolar cell classes is less well established but some progress has been made, as summarized below.

The ML class cones in primates make contact with multiple parallel bipolar channels (Boycott & Wässle, 1991; Calkins, Tsukamoto, & Sterling, 1996; Haverkamp, Grüner, & Wässle, 2000). The majority of bipolar contacts to M and L cones are from midget bipolar...
Figure 2. Selective and random connections in retinal color pathways. (A) Drawing from a double-labelled preparation of whole mount marmoset retina showing blue cone bipolar cells (stippled) and S cone pedicles (blue plaques). The M and L cone pedicles are not shown. Note that blue cone bipolar dendrites are selectively associated with S cone pedicles. (B) Drawing of a marmoset retina showing the dendritic terminals of diffuse-ON DB4 bipolar cells (black puncta) are selectively associated with ML class cones (yellow plaques) and make little or no contact with S cones (starred blue plaques). (C) Selective contact patterns can contribute to blue-ON/yellow-OFF response property of small bistratified (blue-ON) ganglion cell. The blue cone bipolar cells connections are selective for S cones, whereas the diffuse-OFF bipolar (most likely, DB2 and DB3) connections are selective for ML cones. These cone selective signals confer blue-ON/yellow-OFF properties by convergence to the ganglion cell dendrites. (D) Midget-OFF bipolar cells and diffuse DB6 bipolar cells are labelled by antibodies to CD15 in this vertical section of marmoset retina. Note the palisade of single-cone contacting midget dendrites (two are enclosed by the white box). (E) Drawings of midget bipolar cells in marmoset fovea showing points of single cone connections (black clusters) and axons that contact midget-PC pathway ganglion cells (grey clusters). (F) Drawings of two midget bipolar cells at 3.6 mm from the fovea. These cells get convergent input from three and four cones, respectively (black clusters). (G,H,I) Evidence for synaptic (OFF-type) contact of midget bipolar cells at ML cone pedicles: (G) midget bipolar terminal, (H) OFF-type (AMPA) glutamate receptor clusters, (I) overlay. The dendritic tips are associated with glutamate receptors at points of contact with the cone pedicle. Panel A modified from Luo et al. (1999); panel B modified from Lee & Grünert (2007); panel D modified from Chan et al. (2001); panels E and F modified from Telkes et al. (2008).

cells (Kolb, 1970); the remaining contacts are divided among seven or more diffuse bipolar cell types that each make relatively small numbers of synapses with multiple cones (Boycott & Hopkins, 1991; Boycott & Hopkins, 1993; Boycott & Wässle, 1991; Calkins, Tsukamoto, & Sterling, 1998; Chun, Grünert, Martin, & Wässle, 1996; Hopkins & Boycott, 1997). There is evidence for a bias against diffuse bipolar cell connections with S cones (Figure 2B, Lee & Grünert, 2007; Lee, Jusuf, & Grünert, 2004; but see also Joo, Peterson, Haun, & Dacey, 2011; Puthussery, Gayet-Primo, Taylor, & Haverkamp, 2011). Thus, the spectral sensitivity of diffuse bipolar cells should be weighted to the medium-long wavelength band. The differential spectral weighting of bipolar inputs to (ON type) blue cone bipolar and (OFF type) diffuse bipolar classes DB2/DB3 is the best-supported explanation for the characteristic “blue-ON/yellow-OFF” receptive field property of small bistratified ganglion cells in the retina (Figure 2C) and their thalamic target cells in the lateral geniculate nucleus (Chichilnisky & Baylor, 1999; Crook et al., 2009; Dacey & Lee, 1994). Blue-ON cells typically respond poorly to achromatic contrast (Tailby, Solomon, & Lennie, 2008a; Tailby, Szmajda, Buzás, Lee, & Martin, 2008b); this
fact is consistent with *in vitro* recordings from macaque retina showing that both the blue-ON and yellow-OFF receptive field subunits have inherited center-surround structure from their bipolar and horizontal cell inputs (Crook et al., 2009; Field et al., 2007).

The blue-OFF pathway is less well understood than the blue-ON pathway. To date, an S cone-selective blue-OFF bipolar cell has not been described for primate retina. In marmoset retina, only very sparse connections between S cones and OFF midget bipolar cells are present (Lee, Telkes, & Grünert, 2005; Telkes, Lee, Jusuf, & Grünert, 2008). In macaque fovea, an electron microscopic study found that OFF midget bipolar cells made contacts with S cones (Klug, Herr, Nго, Sterling, & Schein, 2003), but blue-OFF receptive fields reported in primates are substantially larger than midget-parvocellular fields at the same eccentricity (Dacey & Packer, 2003; Dacey, Peterson, Robinson, & Gamlin, 2003; Szmajda, Buzás, FitzGibbon, & Martin, 2006; Tailby et al., 2008a,b; Tailby, Dobbie, Hashemi-Nezhad, Forte, & Martin, 2010). Signals from S cones contribute to midget-OFF receptive fields in peripheral macaque retina (Field et al., 2010) but these signals are also non-opponent (i.e., they have the same sign as M and L inputs). These apparent inconsistencies between anatomy and physiology are an important unsolved puzzle. Recent demonstration of a blue-on type amacrine cell in rodent retina (Chen & Li, 2012) raises the possibility that a homologous pathway in primate retina could invert blue-on signals from bipolar cells to supply blue-off type responses in ganglion cells.

**ML cone pathways in the fovea**

In contrast to the selective connections between the S cone array and the blue cone bipolar array, there is to date no compelling evidence that midget or diffuse bipolar cells discriminate between M and L cones (Boycott & Wässle, 1991; Chan, Martin, Clunas, & Grünert, 2001; Silveira, Grünert, Kremers, Lee, & Martin, 2005; Silveira, Lee, Yamada, Kremers, & Hunt, 1998; Telkes et al., 2008). Figure 2D shows a vertical section of marmoset retina processed with antibodies to CD15; in marmosets this antibody labels OFF-type midget bipolar cells and the diffuse bipolar class DB6 (Chan et al., 2001; Telkes et al., 2008). In the fovea, the midget bipolar cells form a dense array with each cell contacting a single cone photoreceptor. Each midget bipolar cell can thus signal spatial contrast over a receptive field corresponding to that of a single cone and will also inherit the spectral signature (M or L) of the contacted cone. Each midget bipolar cell makes dominant contact with a single midget ganglion cell in the inner plexiform layer. Again in the fovea there is one-to-one connectivity giving a ‘private line’ for signals from each foveal cone (Jusuf, Martin, & Grünert, 2006a).

The ‘private line’ connections extend to about 10 degrees in macaque monkey, cebus monkey, marmoset, and human retina (Silveira et al., 1998); at greater eccentricities in macaque retina the one-to-one connections between cones and midget bipolar cells persist, but each ganglion cell starts to draw input from multiple bipolar cells (Goodchild, Ghosh, & Martin, 1996; Wässle, Grünert, Martin, & Boycott, 1994). Some private line connections may persist up to 50 degrees in human retina (Kolb, Linberg, & Fisher, 1992). Beyond ~10 mm (40 degrees) in macaque retina the midget bipolar cells contact two or more cones. In marmoset retina the one-to-one pattern breaks down even closer to the fovea (Telkes et al., 2008; Yamada, Silveira, Gomes, & Lee, 1996): below 1 mm eccentricity each midget bipolar cell contacts a single cone (Figure 2E) but already at 1 mm the midget bipolar dendrites branch to form multiple clusters contacting 3–4 cone pedicles (Figure 2F).

Apart from qualitative differences in convergence, nearly all anatomical evidence points to common functional mechanisms for retinal processing in New World (e.g., marmoset) and Old World (e.g., macaque, human) primates (Kremers, Weiss, & Zrenner, 1997; Kremers, Zrenner, Weiss, & Meierkord, 1998). For example, OFF bipolar cells express ionotropic AMPA/kainate receptors (Figure 2G–H) in both marmoset (Puller, Haverkamp, & Grünert, 2007) and macaque retina (Haverkamp, Grünert, & Wässle, 2001). There may, however, be a species difference with respect to expression of AMPA/kainate receptors postsynaptic to S cone pedicles. Haverkamp et al. (2001) found GluR1 receptors opposed to all cones including S cones in macaque retina, but in marmosets GluR1 (AMPA) receptors were not associated with S cones (Puller et al., 2007). This difference may be related to the potential differences in S cone connections of OFF midget ganglion cells outlined above, but other possibilities such as antibody cross-reactivity cannot be ruled out.

**ML cone pathways outside the fovea**

The presence of one-to-one connections in the fovea gives the only (so far) demonstrated substrate for preserving M and L cone signals in primate retina. But how then could the psychophysically demonstrated (albeit weak) persistence of red-green color vision in peripheral visual field be supported? Figure 3 summarizes the problem. In the fovea (Figure 3A), each M and L cone provides input through one-to-one connections to the receptive field center of ON and OFF type midget ganglion cells: in other words, there is divergence of cone signals to ganglion cells. Further from the fovea (Figure 3B; for simplicity, only connections of ON type ganglion cells are shown) the cones are more widely spaced (i.e., their spatial density is lower), some midget bipolar cells contact multiple cones, and midget ganglion cells receive input from multiple bipolar cells: in other words, there is convergence of cone signals to ganglion cells.

Outside the fovea, spectral mixing of inputs to ganglion cells can occur in the outer plexiform layer where midget bipolar cells may contact cones of different (M or L) types or in the inner plexiform layer where bipolar cell inputs converge to ganglion
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Figure 3. Random wiring model for transmission of red-green signals in the midget-parvocellular (midget-PC) pathway. (A) Schematic vertical section of foveal retina. Each medium (M) and long (L)-wavelength sensitive cone makes contact with midget-ON and midget-OFF bipolar and ganglion cells, thus preserving the spatial and chromatic signals in the foveal array. (B) Schematic of the mid-peripheral retina. For simplicity, only ON-type connections are shown. Here, some bipolar cells contact multiple cones and there is also convergence of bipolar to ganglion cells, thus spectral mixing of cone signals to midget-PC pathway ganglion cell centers becomes increasingly likely with increasing distance from the fovea. (C,D) Estimates of the numbers of M and L cones providing convergent input to midget-PC ganglion cells (via midget bipolar cells) at different distances from the fovea. Drawings in (D) show midget-PC ganglion cells labelled by intracellular injection or following retrograde injections in lateral geniculate nucleus parvocellular layers. (E) Numerical estimates of convergence and predicted cone mixing to midget-PC pathway receptive field centers at different distances from the fovea.

We know that outside the fovea the receptive field center area of midget cells roughly corresponds to the area of its dendritic arbour (Buzás, Blessing, Szmajda, & Martin, 2006; Dacey, 1999, see also Figure 7 below; Wässle & Boycott, 1991). The potential cone inputs to midget ganglion cells at different eccentricities can thus be estimated by comparing local cone density (Figure 3C) with ganglion cell dendritic arbors (Figure 3D). The resulting estimates for marmoset retina (Figure 3E) show that the midget receptive field center receives input from ~15 cones at 10 degrees eccentricity and ~90 cones at 40 degrees (Goodchild et al., 1996). The binomial probabilities that >80% of cones will have the same (M or L) type are 0.03 at 10 degrees and 8 x 10^-9 at 40 degrees, so it is obvious that outside the fovea, spectrally pure cone centers would be exceedingly rare if connections were random. We return to this issue in a later section.

Segregation of color signals in the lateral geniculate nucleus

Figure 4 summarizes the outputs of primate retina and distribution of cone signals in the main dorsal thalamic target of ganglion cell axons, i.e., the dorsal lateral geniculate nucleus (LGN). Thalamocortical relay cells in the LGN get dominant input from one or (less frequently) more ganglion cells and each ganglion cell action potential triggers a relay cell action potential with probability about 0.5 (Kaplan, Purpura, & Shapley, 1987; Lee, Virsu, & Elefant, 1983). It is important to remember that the LGN is not just a simple relay for visual signals but participates in local interneuron circuits, sends collaterals to and receives inhibitory feedback from the thalamic reticular nucleus, receives reciprocal projections from layer VI of primary visual cortex, and is also innervated by neuromodulating systems in divisions of the midbrain reticular formation (Casagrande & Norton, 1991; Sherman & Guillery, 2006). Nevertheless, in anesthetized animals, the dominant excitatory drive to relay cells comes from the retina and the receptive field properties of relay cells are a good reflection of the properties of the afferent ganglion cells. Thus, monitoring extracellular action potentials from LGN relay cells is a way to monitor the first site of interaction between the eye and the brain, with emphasis in anesthetized animals on signals coming from the eye.
Figure 4. Transmission of color signals through the lateral geniculate nucleus (LGN) in marmosets. (A) Schematic view of marmoset eye and brain showing connections of a retinal ganglion cell (RGC) in the eye to a relay cell in the dorsal lateral geniculate nucleus (LGN). The relay cell signals are transmitted to primary visual cortex (V1); extracellular action potentials (AP) can be monitored at the relay cell to characterize receptive field properties. (B) Anatomical distinction of midget-parvocellular (PC), parasol-magnocellular (MC) and small bistratified-koniocellular (K-bon) pathways. Dendritic field diameter (a rough proxy for receptive field center diameter) of PC cells is smaller than that of MC or KC cells at a given distance from the fovea. (C) Drawing from a coronal section through marmoset LGN showing arrangement of the PC, MC and KC layers. (D) Summary of receptive field encounter position in a ‘flattened’ schematic of the LGN. The blue-yellow opponent receptive fields are largely, but not completely, segregated to the KC layers, whereas red-green opponent responses are segregated to PC layers. By contrast, the red-green opponent responses are segregated to PC layers (Blessing, Solomon, Hashemi-Nezhad, Morris, & Martin, 2004; Martin et al., 1997; Szmajda et al., 2006). Consistent results were also obtained from recordings in LGN of macaque monkeys (Dreher, Fukada, & Rodieck, 1976; Roy et al., 2009; Schiller & Malpeli, 1978; Wiesel & Hubel, 1966). In summary, the pathways serving blue-yellow and red-green axes of color vision have quite distinct retinal substrates and remain anatomically segregated at least as far as the geniculocortical relay.

Multiplex color and acuity signals in PC pathway receptive fields.

As outlined above, the low convergence (one-to-one connections) of midget-PC pathway bipolar and ganglion cells in the fovea give a plausible explanation for high spatial acuity as well as chromatic selectivity in the receptive field center of foveal PC cells. Theory and observation also show that red-green opponent responses in PC cells can arise from the spatial opponent (center-surround) structure of PC cell receptive fields provided that spatial pooling in the surround is extensive enough to draw mixed M and L cone inputs (Crook, Manookin, Packer, & Dacey, 2011; Diller et al., 2004; Lennie, Haake, & Williams, 1991; Paulus & Kröger-Paulus, 1983; Solomon, Lee, White, Rüttiger, & Martin, 2005). This “random wiring” explanation for transmission of red-green signals is, however, incomplete; two main problems are as follows. First, red-green opponent responses are consistently recorded from PC cells projecting to the LGN are midget, parasol and small bistratified cells. Midget cells project to the parvocellular (PC) layers of the LGN and parasol cells project to the magnocellular (MC) layers (Leventhal, Rodieck, & Dreher, 1981; Perry, Oehler, & Cowey, 1984; Rodieck & Watanabe, 1993; Szmajda, Martin, & Grünert, 2008). Anatomically the midget pathway constitutes the smallest dendritic fields and the PC layers make up roughly 80% of the volume of the LGN (Ahmad & Spear, 1993; Connolly & Van Essen, 1984; White, Goodchild, Wilder, Sefton, & Martin, 1998). At a given distance from the fovea the midget-PC projecting cells have about 1/3 the diameter of parasol-MC projecting cells (Figure 4B), and the MC layers make up roughly 10% of LGN volume. Small bistratified cells likely project dominantly to the koniocellular/interlaminar (KC) division of the LGN: they are preferentially labelled following retrograde tracer injections targeted to KC layers (Szmajda et al., 2008) and blue-ON/yellow-OFF receptive fields are commonly (but not exclusively) encountered in layer K3 between the PC and MC layers (Martin, White, Goodchild, Wilder, & Sefton, 1997). Figure 4C shows a schematic coronal section through marmoset LGN to illustrate the relative size and position of the PC and MC layers with the intercalated KC layers. The encounter frequency of blue-yellow and red-green opponent responses (recorded in trichromatic female marmosets) is shown relative to the borders of PC, MC and KC layers in Figure 4D. Here the anatomical segregation of blue-yellow opponent responses becomes clear. Although the KC layers make up only ~10% of LGN volume, the majority of blue-yellow receptive fields is encountered in KC layers. By contrast, the red-green opponent responses are segregated to PC layers (Blessing, Solomon, Hashemi-Nezhad, Morris, & Martin, 2004; Martin et al., 1997; Szmajda et al., 2006). Consistent results were also obtained from recordings in LGN of macaque monkeys (Dreher, Fukada, & Rodieck, 1976; Roy et al., 2009; Schiller & Malpeli, 1978; Wiesel & Hubel, 1966). In summary, the pathways serving blue-yellow and red-green axes of color vision have quite distinct retinal substrates and remain anatomically segregated at least as far as the geniculocortical relay.
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Figure 5. Characterization of red-green opponent responses (A–D). Each column shows a different hypothetical pattern of sampling from the extra-foveal ML cone mosaic to midget-parvocellular pathway (PC) cells. Top row shows excitatory (“center”) inputs, center row shows inhibitory (“surround”) inputs. Lower row shows predicted spatial frequency tuning for L cone selective and M cone selective gratings. Mixed inputs of either cone type to both excitation and inhibition lead to bandpass spatial frequency tuning, and selective inputs lead to lowpass frequency tuning. (E,F) Frequency tuning curves of two PC cells in marmoset lateral geniculate nucleus. Dashed lines show responses to L-cone selective (Lt) stimuli, solid lines show responses to M-cone selective (Mt) gratings. Curves in E are consistent with type II receptive field organization, the receptive field in F is dominated by M cone inputs. (G) Population responses of PC cells showing wide variation in relative response amplitude for Lt and Mt gratings, and cells with approximately balanced cone weights show high responsivity for red-green gratings. Panels E,F,G modified from Martin et al. (2011).

outside the fovea where spatial pooling from M and L cones to the receptive field center should, on average, substantially degrade chromatic selectivity. The resultant question then is whether red-green opponent PC cells outside the fovea are a specialized population or just edges of a functional continuum of chromatic and spatial opponent properties. Second, the signals from ‘randomly wired’ PC cells are ambiguous in the sense that changes in discharge rate could be produced by luminance or chromatic variation in the environment (Dreher et al., 1976; Ingling & Martinez, 1985; Lee, 1996; Shapley & Perry, 1986). Can the brain disentangle the responses of PC cells to get high-acuity spatial signals or are PC cell responses only useful for color vision?

We have attacked these questions by studying receptive field properties of PC cells at different distances from the fovea and by comparing responses of PC cells in LGN of dichromatic and trichromatic marmosets.

Figure 5A–D shows alternative ways that cone inputs to PC cell receptive fields could be arranged in the peripheral (extrafoveal) visual field of dichromatic and trichromatic marmosets. The anatomical convergence sketched here is appropriate for receptive fields at 20–30 degrees eccentricity in the marmoset retina (Buzás et al., 2006; Goodchild et al., 1996; Jusuf, Martin, & Grünert, 2006b). The location of individual cones that...
could provide input (via retinal interneurons) to PC pathway ganglion cells are sketched as small colored points; cones showing strong functional connections are filled with saturated color. Each column shows a different receptive field arrangement. The upper row shows excitatory input to the receptive field; the center row shows inhibitory input. The lower rows shows the predicted shape of spatial frequency tuning curves for gratings that modulate only M cones ("M selective", Ms) or only L cones ("L selective", Ls). Because PC cells get negligible functional input from S cones in marmosets (Tailby et al., 2008b), the S cones have been omitted from the sketch diagrams. In a dichromatic marmoset expressing only M cones (Figure 5) there is no response to Ls gratings and the frequency response to Ms gratings shows the bandpass characteristic of center-surround receptive field organization. Three alternative (but not mutually exclusive) hypothetical arrangements in trichromatic marmosets are shown. For simplicity, high spatial frequency aliasing due to fine structure of L and M cone expression is not considered. In a pure “random wiring” arrangement (Figure 5B), spatial tuning is band-pass for Ms and Ls gratings because both cone types contribute to center and surround. Alternatively, if the functional weight of inputs is biased (Figure 5C), in this example, to L cones in the center and M cones in the surround, the Ms response would be weaker than the Ls response and show more lowpass tuning. Figure 5D shows a canonical “Type II” receptive field arrangement where there is functional segregation and spatial overlap of cone mechanisms; this receptive field is specialized to signal chromatic contrast as for luminance (M+L) modulation the M and L signals would cancel. We evaluated the response patterns of PC cells in dichromatic and trichromatic marmoset against these model receptive fields.

Figure 5E and F shows for two PC cells spatial frequency tuning curves consistent with pattern of cone inputs shown in Figure 5C,D. One cell (Figure 5E) shows overlapping and lowpass tuning for Ms and Ls gratings as predicted for “Type II” receptive field organization. The second cell (Figure 5F) shows responses consistent with biased input whereby M cones contribute to both center and surround (yielding bandpass frequency tuning). L cones make only a weak contribution biased to the surround. These two cells were recorded from the same animal showing there is variability in receptive field properties. Population data from seven marmosets are shown in Figure 5G (Martin, Blessing, Buzás, Szmajda, & Forte, 2011). Here the relative weight of M and L inputs (X axis) is estimated from response amplitude at low spatial frequencies (which activate both center and surround). The response amplitude for out-of-phase modulation of M and L cones (M–L, “red-green”) is shown relative to amplitude for in-phase (M + L, “luminance”) modulation on the Y axis. It is clear that cells with approximately balanced inputs from M and L cones show higher chromatic selectivity. The important point is that there are not distinct clusters of luminance- or color-selective receptive fields, but variation of cone weights naturally yields some cells that are more color selective and others that are less so. Comparable reports of relative cone weights in macaque ganglion cells and LGN cells show relatively well-balanced M and L cone inputs in parafoveal (<15 degrees) receptive fields with increased variability at higher eccentricities (Derrington, Krauskopf, & Lennie, 1984; Solomon et al., 2005). Different observations in two species may be reconciled by the fact that spatial pooling of cone inputs at a given eccentricity in marmoset retina is two to three times greater than in macaque retina; for example, ~50 cones feed to a PC cell center at 20 degrees in marmoset retina, but this level of convergence is only present at 55 degrees in macaque retina. Thus increased cone convergence may lead to greater variability in cone weight.

A further example of variability in PC cell selectivity is shown in Figure 6. Responses of two simultaneously recorded PC cells are shown. The spatial frequency amplitude tuning of these two cells for in-phase modulation of M and L cones (Figure 6A) is indistinguishable; they differ only in response phase (ON vs. OFF) as shown by the PSTH histograms above the tuning curves. Exploration with Ls and Ms gratings reveals, however, a dramatic difference in cone inputs: the L cones contribute to both center and surround of one cell (yielding bandpass tuning) but only to the center of the second cell, which shows lowpass tuning for Ls gratings (Figure 6B). The M cones contribute only weakly to one cell but provide strong inhibitory (surround) input to the second cell (Figure 6C); consistent with the population data shown above (Figure 5) these balanced opponent inputs yield vigorous responses to “red-green” (M – L) modulation (Figure 6D). This example shows directly that expression of M and L cones can generate red-green opponent responses in PC cells without dramatic effects on frequency tuning for luminance modulation.

Color and acuity signals in dichromatic and trichromatic marmosets

Further lines of evidence for multiplex color and acuity signals in PC pathway cells are assembled in Figure 7. Dendritic field diameter of a typical midget-PC pathway ganglion cell at 13 degrees (example from Figure 3) is close to 0.25 degrees (red circle). This value is consistent with the optimum aperture for low spatial frequency luminance modulation in a typical LGN PC cell (Figure 7B, upper). In other words, the receptive field center diameter of PC cells outside the fovea is consistent with the dendritic field diameter. Responses of the same LGN PC cell to red-green chromatic modulation (Figure 7B, lower) show that as expected from the examples and population responses in Figures 5 and 6, the red-green response is dependent on center-surround interactions (Buzás et al., 2006). In other words, at this point in the visual system, red-green opponency and spatial opponency are manifest in the same cell type. Figure 7C shows the position of
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the example cell (red circle) in the population of PC cell center sizes. Here the PC cells (measured with achromatic gratings) are compared with blue/yellow opponent cells (blue-ON and blue-OFF, measured with S cone selective gratings) to show the higher resolving capacity of PC cells (Szmajda et al., 2006). Finally, the spatial resolving capacity of PC cells in dichromatic and trichromatic marmosets shows heavy overlap (Figure 7D,E), and in trichromatic marmosets the center radius of cells showing red-green opponent responses is indistinguishable from that of cells dominated by a single cone type (Martin et al., 2011; but see also Kilavik, Silveira, & Kremers, 2003). Thus, the PC cells in trichromats carry a red-green signal but show the same spatial acuity for luminance variation as do PC cells in dichromats.

The peripheral puzzle

The foregoing data and analysis show that on comparing dichromatic and trichromatic marmosets, red-green signals appear as an additional response dimension on spatial opponent receptive (center-surround) PC cell receptive fields. Furthermore, the spatial resolving capacity of the PC cell array for luminance change is not degraded by the manifestation of red-green opponent responses. The only remaining puzzle in this otherwise consistent “random wiring” model for PC cell receptive fields is the improbably large proportion of cells showing red-green opponent responses in peripheral retina of marmoset and macaque monkeys (Buzás, Szmajda, Hashemi-Nezhad, Dreher, & Martin, 2008; Rodieck, 1991; Solomon et al., 2005; Zrenner & Gouras, 1983). As outlined above, in peripheral retina purely random convergence of M and L cones to PC cell centers should yield spectral mixing and non-opponent response properties in the vast majority of PC cells. One possible explanation is that during development the clonal identities of midget-PC pathway cells in the retina are somehow linked to preserve the pattern of lyonization (X-chromosome inactivation) of M or L cones in the allelic array. Alternatively the correlated changes in activity among M and L cone arrays might produce neuroplastic alteration of retinal connections. However, neither of these possibilities has been tested, and fine-scale studies of retina anatomy have as yet revealed no sign of selective connections in the midget-PC pathway (Dacey, 1993; Goodchild et al., 1996; Jusuf et al., 2006b; Yamada, Silveira, & Perry, 1996).

Conclusion and further directions

The data support a model of color signal processing where signals from blue cones are transferred mainly through the evolutionary ancient part of the visual system called the koniocellular pathway. Red-green color signals by contrast appear as an extra response dimension of the parvocellular pathway, which likely evolved to signal fine image details. Although the role of the magnocellular pathway in setting the limits of spatial vision at low contrast is established, the ubiquitous presence and fine resolving capacity of the PC cell array in dichromatic marmosets (and, as far as can be determined, dichromatic humans) is de facto proof for a role of PC cells in spatial vision. Because the parvocellular and koniocellular pathways show distinct patterns of connections within the brain, an outstanding question is where and how the input signals are recombined to yield coherent color perception.

Figure 6. Variability in red-green opponent response. Example tuning curves of two simultaneously recorded parvocellular (PC) cells in marmoset lateral geniculate nucleus show. (A) For luminance (L+M) modulation the frequency response of these cells is indistinguishable. They differ only in response phase (PHSTHs at low and optimum spatial frequency, indicated by vertical lines). (B) For L cone selective (Ls) gratings cell 1 (thin red lines, solid symbols) shows lowpass tuning but the cell 2 (thick grey lines, open symbols) shows bandpass tuning indicating input from L cones to center and surround. (C) Responses to M cone selective (Ms) gratings show strong M cone inputs to cell 1 surround but only feeble M cone inputs to cell 2. (D) Cell 1 shows vigorous response to red-green chromatic (M–L) gratings as result of synergistic activation of L excitation and M inhibition. Cell 2 is insensitive to the same stimulus.
Figure 7. Spatial acuity of color opponent cells in marmosets. (A) Example of midget-parvocellular pathway (PC) cell in marmoset peripheral retina. The red circle encompasses the dendritic field. (B) Aperture tuning curves for luminance (upper panel) and red-green modulation (lower panel) recorded from a PC cell in marmoset LGN. The shaded red bar shows diameter of the red circle from panel A, indicating the optimum aperture (receptive field center) is similar to the dendritic field diameter at similar eccentricity. Responses to red-green modulation are restricted to apertures where both center and surround are recruited. (C) Receptive field size recovered from spatial frequency tuning curves. In comparison to blue-ON and blue-OFF cells, PC cells have relatively small receptive fields. The red circle indicates the radius of a typical midget-PC pathway dendritic tree at 13 deg. (D,E) Comparison of receptive field radius of PC cells measured with achromatic or luminance (L+M) gratings. There is heavy overlap between PC cell acuity in dichromats and trichromats and between opponent and non-opponent cells in trichromats. Panel C modified from Szmajda et al. (2006). Panels D and E modified from Martin et al. (2011).
Likewise, the question how signals from the PC cell array are disentangled to yield information about red-green color difference as well as fine spatial detail has not been solved. Comparative functional analysis of cortical visual pathways in primates may in the future contribute to this big challenge for understanding color vision.

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**Abbreviations**

- AMPA: α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid
- S: short wavelength sensitive (“blue”)
- M: medium wavelength sensitive (“green”)
- L: long wavelength sensitive (“red”)
- ML: medium/long wavelength sensitive
- LGN: lateral geniculate nucleus
- PC: parvocellular
- MC: magnocellular
- K: koniocellular
- mGluR6: metabotropic glutamate receptor type 6