



Neurophysiological correlates of color vision: a model

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Abstract

The tree-receptor theory of human color vision accounts for color matching. A bottom-up, non-linear model combining cone signals in six types of cone-opponent cells in the lateral geniculate nucleus (LGN) of primates describes the phenomenological dimensions hue, color strength, and lightness/brightness. Hue shifts with light intensity (the Bezold-Brücke phenomenon), and saturation (the Abney effect) are also accounted for by the opponent model. At the threshold level, sensitivities of the more sensitive primate cells correspond well with human psychophysical thresholds. Conventional Fourier analysis serves well in dealing with the discrimination data, but here we want to take a look at non-linearity, i.e., the neural correlates to perception of color phenomena for small and large fields that span several decades of relative light intensity. We are particularly interested in the mathematical description of spectral opponency, receptive fields, the balance of excitation and inhibition when stimulus size changes, and retina-to-LGN thresholds. **Keywords:** human color vision, opponent theory, three-color theory, three-receptor theory, perception, neuroscience.

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Introduction

Current neuroscience has a good understanding of the three-receptor theory of color vision, but the opponent-color theory (Hering, 1920) still lacks important neural correlates. Even though one still searches for neural processes that underlie the perception of the elementary colors (yellow, blue, red, green, white and black), the opponent color theory is as relevant as before. The chromatic unique hues help us orient ourselves in color space more or less like four compass directions help orientation on the map. Scientific investigations imply that the elementary colors are associated with measurable physiological processes or states of the brain. How this comes about is unknown and poses a fundamental philosophical question, e.g., how can a perceived quality like a certain color be related to a physical-chemical state or process? Color qualities are different from such processes, and clarification of their status within natural science must deal with the physical conditions for perception as well as thorough knowledge of correlations between the qualitative properties and neural activity in the visual pathway.

By the 1960s, visual neuroscience had made great advances. Thorsten Wiesel and David Hubel (1966),

and Russell De Valois (1965) in the U.S. started recording the activity of single neurons in the primate visual pathway. These recordings largely confirmed the idea behind zone theories (e.g., Müller, 1930) and Schrödinger's (1925) transformations, although it later became clear that some modifications were necessary. Further quantitative data were gathered at the end of the millennium in Otto Creutzfeldt and Barry B. Lee's laboratories at the Max Planck Institute for Biophysical Chemistry in Göttingen, Germany. Around 1980, one of us (A.V.) was invited to join this team. Our recordings from opponent cells in the retina and lateral geniculate nucleus (LGN) of the macaque monkey (*Macaca fascicularis*) led to a physiological model of color vision that accounts for several color phenomena and psychophysical data (Valberg et al., 1986a; Lee et al. 1987).

In this paper we provide an overview of the model by listing its main features and their relation to the phenomenology and neurophysiology of color vision. The main purpose is to point at some factors that need incorporation in this and all similar physiological color vision models. In addition to spectral opponency, relations that must be considered in a revised version are as follows:

- receptive fields and the response of opponent neurons to changing size of the stimulus
- the excitatory/inhibitory balance within the opponent receptive field that must be independent of stimulus size (Seim and Valberg, 2013)
- the newly discovered (Seim et al., 2012) substantial threshold of the retinal input to an LGN neuron

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A physiological model of color vision

Recent studies of color vision (see Shapley and Hawken, 2011 for a review) have mainly dealt with the properties of the neural channels that lead from the retina via the lateral geniculate nucleus to the higher visual centers of the brain. Anatomical and morphological studies have been complemented with neurophysiological recordings, the latter often using system-theoretical methods like mathematical Fourier analysis of sensitivity and response (Lee et al., 1990; Smith et al., 1992). Within this mechanistic framework the nerve cells are treated as filters of optical information. Cells are characterized by their behavior to threshold contrasts and frequencies within the temporal and spatial domains. This approach is, ideally, limited to linear systems. Therefore, in visual science such analysis is adequate for neural activities at or near the threshold of visibility. It gives a rapid and useful overview of functional importance. This method is a useful supplement but is perhaps less intuitive than more traditional studies of neural correlates to perception of color phenomena that spans several decades of relative light intensity. Under normal conditions on a sunny day, for a given adaptation, the activity of receptors and nerve cells must cope with at least five logarithmic units of intensity, that is a ratio of more than 1:100,000 between the lowest and the highest luminance, and non-linear behavior is the rule.

Experiments have shown that psychophysical detection of color thresholds of humans, e.g., spectral sensitivity to small spots projected upon a white background, corresponds well with the threshold sensitivity of the most sensitive opponent LGN cells of the macaque monkey (Lee et al., 1987). Also, the achromatic interval between detection and the discrimination of hue corresponds nicely with the sensitivity difference of non-opponent magnocellular cells (MC cells) and of opponent cells (of either the parvocellular (PC cells) or the koniocellular types (KC cells)) (Valberg and Lee, 1989). Moreover, at the suprathreshold level, the same size color differences and color scaling was found to correlate with relative firing rates in a network model combining six different types of opponent cells. The same applies to the change of hue with intensity (the Bezold-Brücke phenomenon) and saturation (the Abney effect) (Valberg et al., 1991). In these experiments, large and small chromatic and achromatic stimuli of different relative intensities were alternated with a 4 x 5 deg² white adaptation field of constant luminance.

However, after the recent finding of a substantial threshold in the afferent prepotential activity that elicits a LGN response (particularly for small test fields; Seim et al., 2012), improvements can be expected in the modelling of LGN cell responses. Below we shall take a closer look at the consequences of this new finding for the modelling approach.

The first step is the excitation of cones by light absorption. For cone absorption spectra we use used

the Stockman-Sharpe cone fundamentals L, M and S (Stockman & Sharpe, 2000, generally accepted by CIE). The receptor hyperpolarization, V_M , evoked by light can be described by the hyperbolic Naka-Rushton formula of receptor excitations M :

$$V_M = M^n / (M^n + \sigma^n) \quad \text{Eq. (1)}$$

for the M-cone and similarly for responses V_L and V_S of the two other cone types. The exponent n is allowed to vary between 0.7 and 1.0 (Valeton & van Norren, 1983). σ is the half-saturation constant (a parameter that is interpreted as a sensitivity measure) with a different value for each cone type. It was allowed to vary freely during simulation. This formula is sometimes called the Michaelis-Menton function but can also be traced back to Archibald Hill (1910). The exponent n is often called the Hill coefficient (Hsien-Che Lee, 2005). When plotted on a logarithmic luminance x-axis, this equation gives the characteristic sigmoid I-R curve for the cone response.

Let us use an 'L-M' opponent PC cell to illustrate the simulation procedure. The cell receives its (prepotential) input from a retinal ganglion cell G. To model the firing rate N_G of the retinal ganglion cell in imp/s, we use the equation

$$N_G = N_L - N_M, \quad \text{Eq. (2)}$$

where $N_L = A_L V_L$ and $N_M = A_M V_M$ and A_L and A_M are amplitude scalars for the L and M cone contributions, respectively. The eye was adapted to a large achromatic field A that was shortly replaced by the test stimulus. The recorded ganglion cell input (the prepotential N_p) was the difference between the response to the test field N_G and the adaptation value N_{GA} . The response is then:

$$N_p = N_G - N_{GA} = N_L - N_M - N_0, \quad \text{Eq. (3)}$$

where the L and M responses to the adaptation field are included in a single constant N_0 . The response of the 'L-M', LGN cell is then described by the equation

$$N_{LGN}(L-M) = A_L V_L - A_M V_M - N_0 \quad \text{Eq. (4)}$$

where A_L and A_M are amplitude weighting constants. In the early version of the model (Valberg et al., 1987), N_0 represented the anticipated activity of the LGN cell for zero luminance (when the adaptation field was replaced by a black field). Later, due to the discovery of a relatively high prepotential threshold spike frequency (a relative high frequency of the retinal ganglion cell response was required before the LGN cell started to fire, see Seim et al., 2012), it became necessary to give N_0 the new interpretation that it represents the anticipated activity of the prepotential (from the retinal ganglion cell) for a zero luminance test field. In the optimization routine, N_0 was therefore allowed to have a negative value relative to the LGN cell response. This new role of N_0 leads to

improved response simulations with values of n usually larger than 0.7.

The general version of Eq. (4) is given in Eq. (5) below with opponent combinations (differences) of cone output signals, V_S , V_M and V_L :

$$N_{LGN} = \pm A_L V_L \pm A_M V_M \pm A_S V_S - N_0 \quad \text{Eq. (5)}$$

For each cell, the optimization of the parameters in Eq. (5) was achieved by minimizing the root mean square differences (*rms*) in firing rates between the measured responses in an I-R series and those predicted by the model. Figure 1A shows an example of the simulation according to the “Pre-2010” version of the model (that did not work well for small test field sizes as seen here for the 699 nm stimulus (filled triangles)). Figure 1B displays the much improved results when N_0 is allowed to go negative (relative to the LGN cell response).

Figure 1A

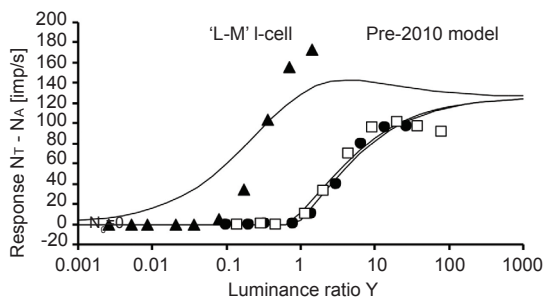


Figure 1B

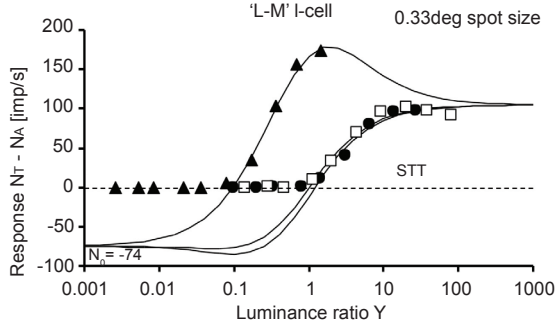


Figure 1. (A) The data points show the recorded response to a 0.33 deg stimulus for three wavelengths. The cell was a parvocellular (PC) LGN cell with ‘L-M’ opponency (filled triangles: 699 nm; open squares 529 nm; filled circles: 500 nm). The curves show the best simulation of the responses by the “Pre-2010 model” using $N_0 = 0$ imp/s, and the exponent $n = 0.7$ in the response equation. The result is worse for the 699 nm stimulus. (B) Simulation of the same data as in (A) but using the revised model described in the text, with a negative $N_0 = -74$ and an exponent $n = 1.0$. STT is a signal transfer threshold (of the preopthal firing) that must be reached by the retinal input before the LGN cell starts to fire. The root mean square (*rms*) deviation between data and model is 2.5% of maximum response.

When the adaptation state of the cells was changed by projecting a steady white surround of different luminance around the test spot, the I-R curves were

shifted horizontally without changing their shape (Valberg et al., 1985). This is consistent with the half-saturation constants σ in Eq. (1) being multiplied by the same constant factor for both the L- and M-cones. Increasing the test stimulus size to exceed that of the receptive field center usually reduced the responsiveness (amplitude) of the LGN cell, but equally for excitatory and inhibitory processes. This attenuation of the modelled I-R curve is achieved by reducing the amplitudes A_L and A_M by the same factor (see Eq. (4)).

Interpretations

As a result of this extended and improved model, new insight was gained as to the role of each cell type in processing color information (Valberg & Seim, 2008; Valberg, 2005). For instance, Increment (I cell) and Decrement (D cell) cells (the ON- and OFF-types) can be shown to have opposite roles in discriminating graded luminance and chrominance (see Figure 2). Let us summarize:

- There are mainly four LGN parvocellular cone-opponent cell types that combine L- and M-cones; the ‘L-M’, I cells and the ‘L-M’, D cells, the ‘M-L,’ I cells and the ‘M-L’, D cells (Kaplan & Shapley, 1982). S-cones are connected to the ‘M (+L) –S’ cells and ‘S-L (+M)’ cells in the konicellular pathway (Martin et al., 1997).
- After an initial threshold, the firing frequency of most opponent parvocellular LGN cell types to large fields can be mathematically simulated by means of opponent inputs of excitatory and inhibitory cone signals (proportional to receptor potentials).
- Whereas for a given adaptation, linearity between stimulus strength and response magnitudes applies only over a relatively narrow stimulus range, non-linear hyperbolic functions (Eq. (1)) work well for the cones over 5-6 logarithmic units of stimulus intensity.
- For opponent cells or a combination of opponent cells, the firing rate in response to a chromatic stimulus—relative to that for a white stimulus of the same luminance—corresponds to color strength or chroma (as, for instance, in the Munsell system).
- The magnitude of the responses (firing rates) of these cells can be regarded as vector-lengths in an opponent color diagram.
- None of the color-space axes defined by the responses of these six cell types (see above) corresponds to the directions defined by the unique hues.
- A constant ratio of firing rates (responses) between cells with different opponencies corresponds closely to a perception of constant hue. This also accounts for the Abney effect (the change of hue with saturation).
- As the luminance of a chromatic stimulus increases, the magnitude and ratio of firing rates of orthogonal opponent cells change as expected

for the Bezold-Brücke phenomenon (when hue and chroma are combined) (Valberg et al., 1991).

- Brightness is well accounted for by the combined responses of L/M, I cell types. Similarly, blackness (the opposite of lightness and brightness) is well accounted for by the combined responses of L/M, D cell types (Figure 2; Valberg and Seim, 2008).
- The balance of cone inputs to an opponent cell is stable and does not change with test field size (Seim et al., 2012; Seim and Valberg, 2013).

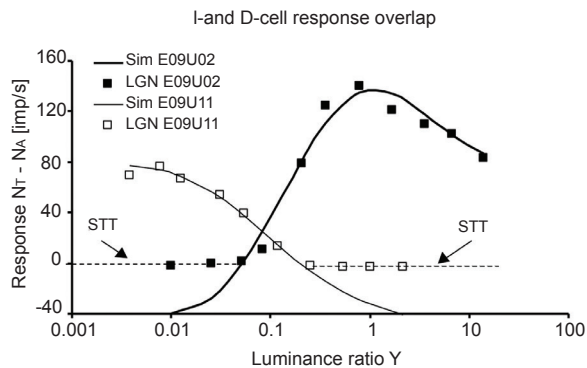


Figure 2. The distribution of information about light increments and light decrements on two opponent parvocellular cell types; the Increment and the Decrement cells (ON and OFF cells). The signal transfer threshold STT is somewhat higher than -40 imp/s in both cases. As in Figure 1, only a model with a high threshold value is able to account for the recorded data.

Comments

Obviously, conscious perception of color qualities arises neither in the photoreceptors nor in the opponent cells of the retina and the LGN. However, *constancy* of several color percepts (such as a constant hue, constant chroma (color strength), constant chromaticity, etc.) seem to correlate with constancy in the relative responses of opponent neurons. Newton said that “*the rays are not coloured.*” With due respect to Helmholtz, we must add that “*colours neither reside in the receptors nor in the cone-opponent cells of the retina and LGN*” (see Valberg, 2001). However, the *same* perceptual attribute (as, for example, a certain orange hue independent of purity) seems to correlate with a *constant ratio* of response magnitudes between cone-opponent neurons. *Constant* color strength (chroma) in an equiluminous chrominance diagram correlates well with a constant vector length from the white point in a vector space based on neural responses.

PC- and KC-cone opponent channels have important roles in color vision, whereas the retinal parasol cells that projects to the LGN magnocellular cells, sum L- and M-cone inputs and do not contribute directly to color vision. Their second harmonic $|L-M|$ rectified opponent response correlates with border distinctness between equiluminant chromatic stimuli (Valberg et al., 1992). These transiently responding magnocellular cells have an unusually high luminance contrast sensitivity and a spectral sensitivity that corresponds closely to the human CIE luminous efficiency function $V(\lambda)$ (Lee et

al., 1988). Because stimuli that have a luminance above the magnocellular threshold, but below the chromatic threshold of PC and KC cells, have a colorless, achromatic appearance, it has been speculated that magnocellular cells contribute to the surface perception of white (Hofer et al., 2005). However, their transient firing of impulses speaks against such a possibility.

As indicated by the word “bistratified,” S-cone activated cells have peculiar dendritic fields different from those of the midget ganglion cells. Their dendritic field branches in two layers corresponding to the ON and OFF layers of other retinal cells (Dacey and Lee, 1994). Until recently there were so few findings of cells with excitatory L/M input combined with inhibitory S-cone input that there were reasonable doubts that they could play a role in color coding, but today this doubt has been removed (Valberg et al., 1986; Dacey and Packer, 2003; Tailby et al., 2008). Cells with excitatory L/M cone input and inhibitory S-cone input discriminate well between stimuli along a white-yellow dimension in color space (but not in the white-blue direction), contrary to PC-cells. Cells with an excitatory S-cone input discriminate well between blue and white stimuli.

LGN and beyond

The discovery of a substantial prepotential threshold spike frequency (the STT; Seim et al., 2012) has led to an improved simulation of LGN cell responses. This threshold leads to I-R curves that span a larger stimulus range, resulting in a larger Hill coefficient, n , in Eq. (1). The important fact that color vision is stable and largely independent of stimulus size and adaptation finds its theoretical support in the simulations where the ratio of half saturation constants, σ_L/σ_M , and of the opponent response components, N_L/N_M are independent of stimulus size and light level (Seim et al., 2012; Seim and Valberg, 2013).

The primate receptive and perceptive fields (Spillmann, 1971) associated with the dendritic fields of opponent cells are still discussed. The current debate is concerned with the organization of cone opponency in simple concentric and double opponent cells. For the retina and LGN, the classical view is one of a center and surround organization that for I-center cells form a “Mexican hat” spatial sensitivity profile with an excitatory cone type in the center and another inhibitory cone type in the surrounding annulus. This spatial sensitivity profile has commonly been modelled as a Difference of Gaussian (DoG) function. However, due to some inconsistencies in the DoG modelling of the response of an opponent cell as a function of stimulus size, it appears that power functions would be better (Seim et al. 2011; 2012). The relative size of center and surround seems to vary among cell types, and cells with S-cone inputs seem to have coextensive or overlapping excitatory and inhibitory receptive fields. Higher up, in area V1 of the brain, one frequently finds so-called double opponent receptive fields with one cone opponency in the center (e.g., L–M) and the opposite

(M–L) in the classical surround (Shapley and Hawken, 2011).

The receptive fields of cells in V1 seem to be different from those in the LGN. Most cells in V1 respond to achromatic contours and contrasts and to movement and orientation and are relatively insensitive to changes in color. However, there seems to be a higher concentration of color selective cells in the so-called “blobs” that make the center of orientation selective columns in V1. In agreement with Livingstone and Hubel’s early findings (1984), these blob cells project to the thin stripes in area V2 (Sincich et al., 2007). The color selective cells of V1 have many preferred hue directions compared to the few cardinal directions of the six cell types in the LGN (Wachtler et al., 2003; Solomon and Lennie, 2007; Conway, 2009; Valberg et al., 1986). This is not easily explained if cortical receptive fields are constructed by convergence from the LGN.

A few studies claim to have proven correspondence between, for example, the sequence of hues in a color circle and a spatial organization of what appears to be color-specific neurons in the thin stripes of area V2 (Xiao et al., 2003). Zeki (1983) claimed that area V4 is specialized for color perception, whereas newer investigations find color-specific cells in an area somewhat larger than V4. Here the color-sensitive cells are arranged in “globs”, i.e., in areas with a high density of color-specific cells that can be viewed using cytochrome oxidase marking, in analogy to the color-specific cells in the blobs of V1 (Conway, 2009). These glob cells seem to project further on to the inferotemporal cortex (IT), and this area is regarded by some as pivotal in the development of perceived color categories (Komatsu, 1997; see also Gegenfurtner and Kiper, 2003).

Conclusion

Color matches, color thresholds, and the laws of additive color mixture are empirical facts that can be satisfactorily summarized by the three-receptor hypothesis. Together with chromatic adaptation, these relations belong to G. S. Brindley’s class A experiments (Brindley, 1960). Class A experiments deal with the identity or non-identity of two sense impressions and, in our case, the physical and physiological conditions that give rise to the same color.

Beyond color matching, we need to deal with color appearance including the ordering and scaling of colors within different categories and in color systems (Hering’s approach). Attributes like hue, saturation, and lightness/brightness, chromatic strength, white and black, are the properties of color that are in focus here as well as scaling of these attributes in color systems (e.g., Ostwald, Natural Colour System NCS, Munsell, etc.). Scaling belongs to Brindley’s class B experiments as does the search for correlations between phenomenological properties of light and color and neural processes.

In our investigation of neural correlates to color perception, i.e., of the relationship between brain activity and the uniqueness of qualia (e.g., unique hues, the redness of red, etc.), we have replaced perceptive qualities by symbols in a mathematical language. In doing so, we attempt to bridge class A and class B experiments in a unifying, quantitative model. More directly, we attempt to bridge the gap between matching in the three-receptor theory and perception in the opponent colors theory (see also Valberg, 2013).

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