Color vision in the comb frequency domain

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ABSTRACT

In 1982, Horace Barlow considered the question of human trichromacy in the context of information theory: according to the Sampling Theorem, three types of receptors covering the visible spectrum (400–700 nm) might be sufficient to reconstruct the color signal. Although Barlow was led to reject the direct application of the Sampling Theorem to explain color dimensionality, the theoretical framework offers a fresh point of view for analyzing the color system in conjunction with the physical characteristics of natural color signals. This review aims to illustrate that if the strict mathematical reconstruction (as implied by the Sampling Theorem) is replaced by a pragmatic approximation of color signals, then trichromacy, with its subsequent opponent-color process, could be regarded as an optimization of color constancy abilities in the spectral environment of primates. Higher dimension systems (tetrachromacy) found in other species can also serve the purpose of color constancy optimization in environments where color signals exhibit a finer spectral structure.

Key terms: Color vision, color dimensionality, comb-filtered spectra, color constancy, Fourier analysis.

COMB-FILTERED SPECTRA

Isaac Newton experimented with prisms breaking down the composite light of the sun into its monochromatic components circa 1666. In one experiment he placed a comb behind the prism so that regular spectral intervals were intercepted while others were transmitted. After recombining the colored beams by means of a lens, Newton could use a white sheet of paper to observe a small circle of light formed at the focal distance (Fig. 1A). The color of the circle changed successively from red to violet as the comb was slid slowly along a parallel axis. When the comb was moving faster, colors were not seen individually, and the sensation of white arose instead. This observation demonstrated yet again the fact that the sensation of white results from the mixture of all primary colors, as Newton concluded:

“… the Impression therefore of all the successive Colors are at once in the Sensorium…the commix’d Impressions of all the Colors do stir up and beget a Sensation of white, that is, that Whiteness is compounded of all the Colors” (from Newton’s text Opticks published in 1730:141).

In 1982, Horace Barlow rekindled our interest in the use of comb-filtered spectra in the study of human color vision, considering them in strict analogy with the spatial frequency gratings used in spatial vision to measure the spatial analogy with the spatial frequency gratings used in spatial vision to measure the spatial resolution of the visual system. In this approach, visual stimuli are characterized by extended sinusoidal modulations in space, and defined by their frequency, amplitude, phase and mean level of energy (Fig. 1B). One can then determine the smallest spectral intervals between two consecutive maxima of amplitude or teeth of the comb that can be resolved by the chromatic system. In Newton’s experiment, if the teeth of the comb covered 100 nm and were spaced out by an interval of the same size (i.e. equivalent to 1 c / T, where T is the extent of the visible spectrum 400-700 nm), the change of color produced when the comb slid over the spectrum (equivalent to a change of phase, Figure 2, top panel) would have been dramatic. With a reduction in
the size of teeth and intervals, equivalent to 3 c / T for example (Figure 2, middle panel), the change in color would be more subtle or even non-perceptible for higher comb frequencies. This is due to the fact that the spectral power distribution (energy per wavelength) of the light is not measured by an array of sensors; rather the continuous spectrum is encoded by the photon absorption rates in a limited number of mechanisms (three for a trichromatic system) that are maximally responsive at different locations in the spectrum (Figure 2, bottom panel). Considering the large absorption curves of the photoreceptors, the photon absorption rate in each type of photoreceptors for a comb-filtered spectrum of high frequency is similar at all phases; a shift of phase is barely detected, and since each type of photoreceptors is similarly stimulated, the comb-filtered spectrum is indistinguishable from a continuous spectrum. The comb frequency at which this confusion occurs is taken as the spectral resolution or frequency cut-off of the color system.

Band Limitation and the Sampling Theorem

Barlow’s original analysis relied on estimating the modulation transfer function (MTF) of color mechanisms, which indicates the level of demodulation imposed on comb-filtered spectra of different frequencies. Due to their broad spectral sensitivity curves, human photoreceptors act as low-pass filters, as can be seen from their response in the comb frequency domain where frequencies greater than 6 c / T are filtered out (Fig. 3). After receptor filtering, the effective signal is band or frequency limited (low frequency components are also demodulated) and can be reconstructed by a finite number of samples (i.e. type of photoreceptors) within a defined spectral interval (i.e. the visible spectrum). The number (N) of samples can be deduced from the application of the Sampling Theorem, which states that to recover a signal of extent T (i.e. wavelength domain) and of frequency cut-off B (i.e. comb-frequency domain), N = 2BT (see Appendix A). Such a relationship could offer an explanation for human trichromacy.
C. The light from the source (S) is located at the focal point of a Fresnel lens (F1), and focused by a second lens (F2) to form a real image S’ of S at the aperture of the integrating sphere (IS). The LCD is mounted in the collimated beam directly after the interference wedge (IF) which gives a continuous linear spectrum from 400 to 700 nm. The output of the integrating sphere, viewed through a lens (L1), produces a homogeneous spot. The figure on the right shows examples of comb-filtered spectra achieved by applying electronic masks to the spectrum (symbolized by the color initials). The masks presented here have $f = 4 \ c/T$ and $p = 0 \ deg$ with 3 different amplitudes. When $m \times 0$ the spot appears colored, and for constant amplitude its hue depends on the combination of frequency and phase, that is, the number and position of the spectral bands transmitted.
If the number of photoreceptors satisfies the equation, adding another sampling dimension (e.g. a fourth type of photoreceptors) would be redundant. For a trichromatic system, the equation becomes $3 = 2BT$, yielding a maximum value of $B = 1.5 \text{ c/T}$ to satisfy the Sampling Theorem’s strict limit. However, the actual frequency cut-off of the photoreceptors ($4 - 6 \text{ c/T}$) would require 8 to 12 types of photoreceptors for the accurate reconstruction of a signal with a similar frequency limit. Barlow was thus led to reject the direct application of the Sampling Theorem as an explanation of human trichromacy and his analysis reveals the existence of signal aliasing occurring at the receptoral level, producing what is known as color metamerism.

This review aims to re-examine the comb-frequency approach to color vision by considering the filtering properties of the color system in relation to the neural processing of natural color signals. Barlow’s original analysis will be extended to encompass current knowledge of the characteristics of natural color signals encountered in primates’ environment. Linear models of color constancy that assume signal approximation will then be introduced. This analysis must indeed be based on color constancy, as this establishes the need for the chromatic system to approximate the incoming signal; color discrimination would not require such process. We will next consider the transfer characteristics of the chromatic system beyond the receptor level. From this analysis we conclude that if an exact mathematical reconstruction of the color signal could not be achieved by a trichromatic system, a pragmatic

Figure 2
The top two sections show comb-filtered spectra of 1 and 3 c/T, with two phases 180° apart. The lowest section shows the three cone-fundamentals (B G R) of Smith and Pokorny (1975) plotted in relative units. The cone-fundamentals represent the sensitivity of the photoreceptors at the corneal level. The color of the 1 c/T comb-filtered spectra will change dramatically with a shift of phase (from violet to greenish-yellow) and a large demodulation of amplitude will be necessary to make this change undetectable. The change in color will be far less prominent for a comb frequency of 3 c/T, even with large modulation of amplitude. (From H. Barlow, 1982).

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Figure 3
Fourier transform of Smith and Pokorny (1975) cone fundamentals. At 3 c/T (indicated by the leftmost vertical dotted line), the demodulation is approximately 25% for the short-wavelength (B) and approximately 12 and 4 % for the medium- and long- wavelength cone fundamentals (G and R), respectively. The abscissa axis is expressed in $c/B$ where 1 cycle corresponds to 0.33 c/T.
approximation carried out by well-designed opponent-color channels could have taken place to secure a suitable degree of color constancy. This conclusion leads to the conjecture that tetrachromacy, in addition to its adaptation to a wider visible spectrum, can also serve the purpose of color constancy optimization in spectral environments of greater complexity than those of trichromats.

Band limitation of natural color signals

Color signals are physically constrained to be band-limited or low-pass functions—functions that do not have frequency components beyond a certain limit. This claim, which dates back to Stiles et al. (1977), was made from the mere observation that natural color signals generally have smooth spectral profiles.

The color signal $I(\lambda)$ results from two intertwined functions: the spectral power distribution of the illuminant $e(\lambda)$ and the spectral reflectance of the object surface $s(\lambda)$, so that: $I(\lambda) = e(\lambda) \times s(\lambda)$. To estimate the color signal frequency cutoff, the two components need to be considered in turn. Maloney (1986) provided the first direct estimate obtained for spectral surface reflectances from stained color samples used in industry (Nickerson-Munsell chips) and surfaces of natural formations (measured by Krinov in 1947). At least ninety-nine percent of the energy in the Fourier transform power spectra was recovered for frequencies as low as 1.5 c / T, and all energy was recovered for frequencies of less than 6 c / T. A second analysis performed by van Hateren (1993) on the spectra of various natural objects showed an identical frequency cut-off. However, these studies could be criticized because they originate from stained samples (Nickerson-Munsell chips) and a heterogeneous collection of objects (Krinov and van Hateren’s samples), and therefore the results cannot be generalized to spectral reflectance surfaces present in the natural environment or attributed to a specific category of natural colors.

More recently, Bonnardel and Maloney (2000) performed a Fourier analysis of the spectral reflectance functions arising from fruits, flowers, and leaves. These natural surfaces, described as biochrome colors (Nassau, 1983), owe their coloration to the selective absorption of incident light by pigments. This process of light reflection is commonly referred to as ‘body reflection’ as opposed to ‘interface reflection’ (also known as specular reflection) (these two processes are described in the Dichromatic Reflection Model proposed by Shafer (1985)). Body reflection is diffuse as the light is reflected in many different directions, and the spectral composition of the light reflected does not depend upon the viewing geometry but is determined by the property of the pigment particles and thus is indicative of the intrinsic color of the material. In contrast, interface reflection is concentrated in one direction, is dependent upon the viewing geometry and for many types of materials the reflection is non-selective, and can thus be indicative of the spectral properties of the illuminant. As we will see later in this paper, the explanation of the light reflection process in terms of physico-chemical properties is important for a proper characterization of the color signal.

The FFT performed on 1695 biochrome samples indicates that more than 99% of the energy is recovered for frequencies below 4 c / T (Bonnardel and Maloney, 2000). This limit is somewhat higher than that obtained in Maloney’s analysis; however, only a residual amount of energy (<0.005) is found at frequencies higher than 6 c / T. The latter result confirms the low-pass hypothesis of the reflectance functions of biochrome surfaces. Due to the lack of availability of empirical measurements, estimates of the cutoff frequency for natural illuminants are far less numerous. However, in the same study a second FFT was performed on a small sample of 348 daylight spectral power distributions provided by Romero et al (1997). Results show that more than 99% of power density is recovered below 1 c / T.

From these two estimates, we can deduce the color signal frequency cut off. The
product of the two functions $e(\lambda) \times s(\lambda)$ when transposed into the frequency domain corresponds to a convolution of their Fourier transforms. It can be shown that the frequency limit of the product is at least as high as either the frequency limit of the illuminant ($b$) or the surface reflectance ($b'$), but less than the sum of the two frequency limits ($b + b'$) (Bracewell, 1965). It follows that the upper limit of the frequency cut off of the color signal reflected by biochrome surfaces under daylight illuminant for the samples considered was estimated at $5 \, c/T$ for the criterion of 99%.

**Linear models of color constancy**

Color constancy can be described as ‘the tendency to see colors as unchanging even under changing illumination condition’ and computational models are designed to recover the invariant spectral reflectance properties of object surfaces from the color signal (Hurlbert, 1998). Given that color signals are band-limited functions, as indicated by the previous analysis, linear models of color constancy offer a computational method for recovering the descriptors of the surface reflectances. Band-limited functions can be adequately approximated by a small number of basis functions whose number, as stated by the Sampling Theorem, is proportional to the frequency limit of the color signal. Thus, a set of illuminants will be approximated by a limited number of illuminant basis functions $E_i(\lambda)$:

$$e(\lambda) \approx \varepsilon_1 E_1(\lambda) + \ldots + \varepsilon_m E_m(\lambda) \quad [1]$$

Likewise, a set of surface reflectances will be approximated by a limited number of surface reflectance basis functions $S_j(\lambda)$:

$$s(\lambda) \approx \sigma_1 S_1(\lambda) + \ldots + \sigma_n S_n(\lambda) \quad [2]$$

the $\varepsilon_i$ and $\sigma_j$ are the weights or the descriptors of the illuminant and surfaces and the quality of the approximation increases with the number ($n$, $m$) of basis functions\(^\dagger\). For the color system to solve the color constancy, the descriptors of the illuminant and surface need to be recovered. The response $Q$ of $p$ types of receptors with response function $q_k(\lambda)$ to the color signal is:

$$Q_k = \sum_{\lambda} e(\lambda)s(\lambda)q_k(\lambda) \quad [3]$$

If $e(\lambda)$ and $s(\lambda)$ are replaced by their linear models expressed in eq. [1] and [2], the receptor response becomes:

$$Q_k = \sum_{i=1}^{m} \sum_{j=1}^{n} \varepsilon_i \sigma_j \sum_{\lambda} E_i(\lambda)S_j(\lambda)q_k(\lambda) \quad [4]$$

In conditions where the illuminant can be known, i.e. the $\varepsilon_i$ can be recovered\(^\dagger\) to find a solution to eq. [4], the number of unknown descriptors $\sigma_j$ of surface reflectance should not exceed the number of type of receptors. In practical terms, this number should not exceed 3 for a trichromatic system, hence the importance of verifying the quality of the approximation of natural reflectance surfaces with low-dimension linear models.

It is possible to derive a set of basis functions from a set of measured reflectance functions using Principal Component Analysis (PCA) and to determine which contributes the most to the total variance of that particular set. Cohen (1964) applied the PCA to 433 Munsell chips and found that 99.18% of the variance can be accounted for by the first three basis functions, although Parkkinen et al. (1989) estimated that the variance could be accounted for by as many as 8 basis

\(^\dagger\) Illuminants are probably approximated by a linear model of smaller dimension compared to that of surface reflectances. From their own daylight measurements Romero et al. (1997) confirmed Judd, MacAdam and Wyszecki (1964) earlier result that three basis functions are suitable to describe natural daylight and reported a percentage as high as 99.97 of the total variance recovered. This value is superior to any value obtained for natural surface reflectances (see text).

\(^\dagger\) The illuminant can be known through specular reflection for instance, but see Maloney (1999) for a detailed review of the different methods and assumptions used in computational approaches to solve this question.
functions for the complete set of Munsell chips (1257) when accuracy of the approximation is used as a criterion instead of the percentage of the total variance. Similarly, using the accuracy criteria, Vrhel et al. (1994) found that 3 to 7 basis functions were necessary to describe a set of heterogeneous spectra (Munsell chips, Du Pont paint chips and natural and man-made objects). Finally, 98% of the total variance of forest and coral scenes is also recovered with the first three basis functions in a study recently done by Chiao et al. (2000). Basis functions determined for a given sample would not necessarily provide a satisfactory fit with data from another sample, but they do share the same characteristics in all the studies cited. While the first basis function is an all-positive function with a generally flat profile, the second has one zero-crossing and the third, two zero-crossings along the wavelength axis (Fig. A). The profile of the basis functions does not correspond to that of the photoreceptors, and in linear models of color constancy the receptors should first

Figure 4
A: Second and third basis functions determined from the PCA of heterogeneous spectral reflectance functions by Vrhel et al. (1994).

1 However, the coral spectra were smoothed out with a low-pass filter of $6 \text{ c} \times \text{T}$ (which corresponds to a sampling of 25 nm) thus limiting the estimation of the percentage of variance at this frequency.
undergo a linear transformation into a new set corresponding to the basis functions in order to achieve color constancy (Hurlbert, 1998). This new set of functions is actually found in the visual system and corresponds to the opponent color stage that follows receptor response. The two post-receptoral opponent color channels also exhibit one or two zero-crossings somehow shifted towards the short wavelengths, compared to the computed basis functions (Figure 4B). Although not identical to the basis functions they are comparable.

*Filtering properties of the human color system*

The two stages of color processing mentioned above were the subject of the conflict between Young’s trichromatic theory (1802), which proposed the existence of three distinct types of receptors, and Hering’s opponent color theory (1878), which was developed to account for sensation phenomena left unexplained by the former theory. The controversy was later resolved by integrating the two processes in multi-stage models of color vision (e.g. Hurvich and Jameson, 1957).

The two opponent-color channels result from a linear combination of the three types of photoreceptors: A ‘yellow-blue’ channel receives combined inputs from the medium- and long-wavelength receptors (M-and L-cones) in opposition to inputs from the short-wavelength receptors (S-cones) and exhibits a bimodal spectral profile \([L + M] - S\). A ‘red-green’ channel combines L-cones in opposition to M-cone inputs \([L - M]\), which also produces a bimodal spectral profile. Under certain conditions, this second channel may receive S-cone inputs leading to a ‘cyan-red’ (S-cones combined to M-cones) or a ‘magenta-green’ opponency (S-cones combined to L-cones). Although cells in the lateral geniculate nucleus (LGN) of primates respond preferentially to yellow-blue and red-green modulations (Derrington et al. 1984), cells responding to cyan-red or magenta-green modulations might be found in the cortex and were recently described in area V1 of the primate cortex (Conway, 2000). In Ingling and Tsou’s model (1977), the two versions of the red-green channel are expressed as a linear combination of two \([L - M]\) or three \([L + S] - M\) type of photoreceptors, depending on the light adaptation conditions. The two channels show a di- or triphasic spectral profile, respectively (Figure 4B).

Due to their opponent process, the post-receptoral channels show band-pass filter characteristics in the comb frequency domain, with a high frequency cutoff equivalent to that imposed by the receptors. Moreover, they attenuate low-frequency components. Depending on the type of cone interactions, the channels are tuned to different ranges of comb frequencies \([L + M] - S\) channel peaks at 1c; \([L - M]\) channel at 1.2 c and the \([L + S] - M\) channel at 1.5 c / T). The yellow-blue channel is thus involved in the extraction of lower comb frequencies compared to the red-green channel, but the difference in tuning is even more marked when compared to the magenta-green channel, which shows a further shift to higher comb-frequencies in its response (Fig. 5a) In the absence of one type of retinal receptor, as is the case in inherited human dichromacy (Nathans et al. 1986) the best that can be achieved is a single channel with a simple cone opponency \([L - S]\), for deuteranopy or \([M - S]\) for protanopy. The net consequence is a reduction in the width of the tuning curve in the comb frequency domain (Figure 5B).

Measurements of human sensitivity to comb-filtered spectra were performed for the first time by Barlow et al (1983) using a Michelson interferometer. Sensitivity thresholds plotted as a function of comb frequencies exhibit curves with jagged profiles, indicating the high degree of dependency of the color system on phase (i.e. in a Michelson interferometer the phase of the comb-filtered spectra varies with the comb frequency). In 1991, we developed a stimulator that allows us independent control of the modulation parameters (Bonnardel and Varela, 1991). In its latest version (Bonnardel, 1998), the stimulator uses the same optical arrangement as in Newton’s set-up. The main difference lies in the use of a linear interferential filter to
The response of cone fundamentals and opponent-color channels are computed for comb frequencies varying from 0.1 to 6 c/T (in intervals of 0.1 c/T) and for phase, varying from 0 to 330° (in intervals of 30°). For each phase the value for the optimal phase only is plotted.

Figure 5
Response of the human chromatic system to comb frequencies at various levels of description:
C. Mean spectral modulation sensitivity function of normal trichromat and deuteranopic observers adapted from Bonnardel et al. (1997).

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obtain a linear spectrum, and the use of a Liquid Crystal Display to replace the comb, thus providing the advantage of displaying electronic masks of any desired spectral profile under computer program control (Figure 1C). The type of measurements obtained were defined as the Spectral Modulation Sensitivity Function (SMSF) (Bonnardel et al. 1996), which corresponds to sensitivity to the optimal phase plotted only as a function of the comb frequency. The SMSF thus provides an estimate of the modulation transfer characteristics of the chromatic system. Several studies have since followed (Bonnardel et al. 1997; Romero et al. 1997) in which, despite differences in technique and methods, all SMSFs show band-pass profile with a maximum sensitivity between 1 and 1.6 c / T and a high comb frequency cutoff estimated between 4.5-6 c / T.

In Figure 5C, the SMSF determined by trichromatic observers is well fitted in the low comb frequencies by the response function of the yellow-blue channel and in the high comb frequencies by the magenta - green (triphasic) channel. However, the red-green (diphasic) channel fails to fit the second shoulder situated at 1.5 c / T. The SMSF of deuteranope observers shows a corner frequency of 1.1 c / T with a reduction in sensitivity in the comb frequency domain which is consistent with the lack of red-green opponency. The curve’s profile is reasonably well described by the response of the red-blue channel.

Color dimensionality

The analysis of biochrome colors and the filtering properties of the color system allow us to draw specific conclusions regarding the relationship between color dimensionality and color constancy for human color vision. A more general conclusion with respect to coherent band-pass theory that applies to color vision in other species is proposed.

1- The frequency cut-off of biochrome colors for the criterion of 99% coincides with the frequency cut-off of the photoreceptors. This coincidence might not be fortuitous. The molecular-interaction hypothesis, as proposed by Maloney (1986), offers an explanation of the observed low-pass constraints based on the physics and chemistry occurring at the molecular level for biochrome colors, where colors are produced by selective absorption of light by pigments. This explanation should also apply to visual pigments in which the combination of a protein (opsin) with an aldehyde of vitamin A (retinal) should yield a similar cut-off frequency.

2- As already pointed out by Maloney (1986), the number of types of receptors is not dictated by the signal approximation requirement; if it were, a perfect color constancy would be expected, and this is obviously not the case for human color vision. However, although the system largely undersamples the signals, Maloney (1986) showed that the low pass filtering properties of photoreceptors enhance their approximation. At the second stage of color processing, in his spatio-temporo-spectral optimal filter theory, van Hateren (1993) demonstrated that, by attenuating the lower frequencies, opponent-color mechanisms exhibit an elementary form of color constancy. This filtering process primarily concerns illuminants that, as mentioned earlier, correspond to the lowest comb frequency components in the color signals. The windowing imposed at the color opponent stage can thus be regarded as the second step of signal filtering to optimize the signal approximation that can be performed at later stages.

3- Buchsbaum and Gottschalk (1983) showed that the design of the two opponent-color channels corresponds to the optimum transformation from the three overlapping types of receptors, to an optimization of information transmission by decorrelating receptor output and thus reducing redundancy. As discussed previously, Buchsbaum and Gottschalk’s opponent-color functions shared the same characteristics as the basis functions, which, as noted by Hurlbert (1998) is non-accidental: both analyses undergo a
decomposition into principal components. However, the opponent-color channels postulated by a psychophysical model such as Ingling and Tsou (1977) bear a closer resemblance to the functions resulting from Buchsbaum and Gottschalk’s optimum transformation than to basis functions. In the comb frequency domain, the opponent-color channels outperform the basis functions, extending sensitivity to high frequencies Figure 6. This result can be attributed to the fact that the basis functions so far computed are derived from a limited set of reflectances and, apart from the study of Chiao et al. (2000), computations made by excluding the contribution of the illuminant did not consider the color signal per se.

If color constancy performance is dependent upon the number of receptors and the subsequent transformation of their response, this raises the question of maintaining color constancy in the case of innate dichromats, whose color system is missing the red-green opponent mechanism. Although this mechanism has a crucial ecological role in the detection and discrimination of fruits (Regan et al. 1998) or young leaves (Dominy and Lucas, 2001) against mature foliage (and is indeed believed to have evolved to fulfill this function in frugivore and folivorous monkeys), the performance of daltonians does not seem to be very impaired when tested for color constancy under laboratory conditions (Ruttiger et al. 2001). The lack of obvious color constancy impairment could be partially explained by the more drastic filtering process imposed by the yellow-blue mechanism, which helps to retrieve color descriptors with only two basis functions. However, if dichromats’ failure in color constancy seems less obvious than their color confusions under laboratory testing conditions, observed results are different in natural settings. It is from a lack of constancy in the color appearance of a pelargonium when moving from candlelight to daylight that John Dalton gained his first acquaintance with his visual condition (Collins, 1925). Indeed, color signals under candlelight compared to daylight illuminants might have a slightly smaller percentage of the total variance recovered at 1 c / T (limit of recovery for a 2-dimensional linear model). In addition, more basis functions would be required to recover surface descriptors seen under this illuminant (see Table I, for the percentage of total energy recovered for a biochrome color under a tungsten bulb and daylight illuminant). In the natural environment, certain conditions might be more challenging to dichromats, thus revealing the more obvious failure of color constancy. It would complement our argument if, in relation to the geographical distribution of human color deficiencies as given in Cruz-Coke’s (1964) survey, specific changes in light and/or surface were found to be more critical for color deficient color constancy.

In light of what has been presented, we can now reconsider the question behind Barlow’s interest in comb-filtered spectra: Why trichromacy? It is clear that there are a number of factors that must enter into an understanding of color dimensionality: (1) the band limitation in natural color signals; (2) the frequency characteristics of photoreceptors; (3) the constraints to achieving color constancy beginning with color signals and photoreceptor responses and the establishment of color-opponent stages. These different factors are, to varying degrees, relatively independent
from one another, and include: sources of illumination, structure of surface materials, receptor photochemistry derived from ancestral biochemical pathways (Nathans et al. 1986), neural developments, and the behavioral and ecological requirements of color constancy (Thompson et al. 1992; Thompson, 1995). The result achieved through the biological evolution of all of these relatively independent natural factors can be assumed to originate from a coherent relationship. No single aspect involved in color vision can in itself account for trichromacy; it is the mutual constraints among them that lead to an evolutionarily satisfactory color dimensionality in the primate group. In other words, there are plenty of metamerisms with 3 types of receptors (and even more with two), but the signal windowing achieved at the post-receptoral level allows a ‘good-enough’ signal approximation for the purpose of color constancy with three or two basis functions. This ‘sufficing’ argument provides a well-grounded rationale for the fact that humans are trichromats rather than deca-chromats, if we follow the strict limit of the Sampling Theorem. Trichromacy appears to be an ecological compromise between accuracy and simplicity in the use of resources; we may call this multi-factor account of trichromacy in primates a coherent band-pass theory of color dimensionality.

**Implications for the color systems of other species**

Considering the central role of color constancy in object recognition, it is a reasonable conjecture that this property has been subject to a selective pressure that presumably makes it inseparable from the evolution of color vision itself in the entire animal kingdom (Thompson et al., 1992). Indeed, in addition to primates, insects, fishes and birds exhibit color constancy when tested behaviorally (Neumeyer, 1998). The coherent band-pass theory of color dimensionality can thus be applied to tetrachromatic systems.

Tetrachromacy is a common feature among insects, fishes, reptiles and birds (Goldsmith, 1991). In these species, the addition of a fourth type of receptors in the context of the comb-frequency analysis could serve the purpose of extending the sensitivity of the chromatic system in the wavelength or in the comb-frequency domain. Diurnal mid-water fish that show what is probably the largest visible window (from 350 to 800 nm) encounter the first advantage among vertebrates. The absorption curves of their photoreceptors, based on porphyropsin, are slightly broader than those based on cone opsin. Thus, the comb frequency cut-off is expected to be slightly inferior to that of the human system. In this case tetrachromacy is of no advantage in the comb frequency domain (Bowmaker, 1983).

<table>
<thead>
<tr>
<th>Proportion of spectral energy below the frequency-limit (c / T)</th>
<th>Visible window : 400-710 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orange x Daylight</td>
<td>0.8845 0.9958 0.9979 0.9983 0.9985 0.9986</td>
</tr>
<tr>
<td>Orange x Tungsten</td>
<td>0.8572 0.9962 0.9988 0.9995 0.9995 0.9996</td>
</tr>
<tr>
<td>Blue feather x Daylight</td>
<td>0.7562 0.9505 <strong>0.9910</strong> 0.9967 0.9975 0.9981</td>
</tr>
<tr>
<td>Blue feather x Tungsten</td>
<td>0.8455 0.9583 0.9882 <strong>0.9957</strong> 0.9972 0.9981</td>
</tr>
</tbody>
</table>

**TABLE I**

At 1 c / T the percentage of total energy recovered from an orange is lower under the tungsten bulb than under daylight illuminant. In this example, the structural color signal displays higher cut-off frequencies than the biochrome color signal (shaded cells) (see text for explanation).
On the other hand, in birds (Maier and Bowmaker, 1993), reptiles (Lipetz, 1984) and crustaceans (Cronin and Marshall, 1994), the spectral sensitivity curve of the pigment is narrowed by means of colored oil droplets that act as cut-off filters in the wavelength domain. The Fourier transform of a 569 nm photopigment with a red oil droplet for chicken retinas extends to comb frequencies nearly twice as high as that of human cones (Barlow, 1982; Bowmaker, 1983) (figure 7). The combination of oil droplet-receptor as a functional unit with a narrow spectral sensitivity curve suggests that color signals with higher band-limits exist in the environment of these species.

Under daylight illuminants, such color signals may be found among a second type of biological coloration called 'schemochrome' or structural colors. Structural colors result from the selective reflection of light by scattering, interference or diffraction produced by the interaction of the light with the microstructure of the biological material. In particular, interference colors, which are rare in plants, are widespread in the animal kingdom (wings of insects, fish scales, eye, outer layer of the skin of some snakes and feathers of birds). A representative example is given by the iridescent (coherent scattering) green reflection spectra of Indian peafowl with two sharp reflectance peaks in the green region (500 nm) and the blue region (400 nm) or shifted in the ultra violet range (350 nm). The conditions are the same for the feathers of the quetzal bird (Finger and Burkhardt, 1994). A recent analysis performed by a team of ornithologists on the non-iridescent blue color (which, until now was incorrectly attributed to the Rayleigh scattering) has identified the process by which its coloration occurs. Interestingly, the predicted spectrum of this structural non-iridescent blue color also shows sharp peaks (Prum et al. 1999).

The Molecular Interaction Hypothesis does not apply to schemochrome colors and no PC or Fourier analysis is available to test the Low-pass hypothesis. However, considering the profile of their spectra, it is likely that structural colors encompass higher comb frequencies and that a higher number of basis functions would be required for their approximation by a linear model. As an illustration, Figure 8 presents the profile of color signals resulting from a biochrome (orange) and a schemochrome (blue feather) color under daylight or tungsten illumination. A comparison of their relative Fourier powers indicates a substantial power beyond the second Fourier component in the case of the schemochrome color, whereas it is only residual in the case of the biochrome color under both illuminants. The criteria of 99% of total energy is reached for frequencies below 2 c / T for color signals resulting from the orange under the two illuminants, and at frequencies below 3 c and 4 c / T, for the blue feather illuminated by daylight and tungsten bulb, respectively (see Table I for values).

If the combination oil droplet-retinal receptor in some tetrachromatic species allows higher comb frequencies to pass, this introduces the possibility that the band-pass response of their chromatic systems would consequently be widened to higher comb frequencies. In this case, a new coherence would be obtained by increasing the number of color channels, narrowing the retinal

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**Figure 7**

Compared to the Fourier transform of the three types of human photoreceptors (continuous and dashed lines), the Fourier transform of the spectral sensitivity of a chicken cone screened by an oil droplet (dotted line) indicates an extension to higher comb frequencies. It should be noted that the scale is in c / Tera Hz (each division corresponds to 1.6 c / T), for which the responses of the three type of human receptors become similar.
photopigments by oil droplets, and adjusting the neural mechanisms in a tetravalent manner to handle color constancy for schemochrome colors. The recent findings of Ventura, Zana, de Souza and DeVoe (Ventura et al 2001) support this hypothesis. For the first time, the authors reported a tetraphasic opponent color ganglion cell [(UV+M) – (S+L)] in the turtle retina (Pseudemys Scripta Elegans), giving rise to the possibility that higher comb-frequency components can be processed beyond the photoreceptoral stage. However, the determination of the transfer characteristics of tetrachromatic systems, as well as the test of the Low-pass hypothesis for surface reflectances from the corresponding natural environment, are key steps towards validating the present hypothesis.

REFERENCES


Figure 8

Four color signals and their relative Fourier Power (RFP) resulting from the product of two surface reflectances (SS) (biochrome and schemochrome colors, top row) and two illuminants (natural daylight and tungsten bulb, left column). The spectral power distribution (SPD) of the daylight illuminant is taken from Romero et al. (1997) measurements, the reflectance function of the biochrome colors from Chittka (1996) and that of the schemocrome color was reconstituted from Prum et al. (1999).
A real-valued function cannot be localized in the variable (say within an interval $T$) and in the frequency domain (say in an interval $B$) at the same time. The best that can be achieved is a compromise such as that of Gaussian profiles of intermediate spread. This ‘limitation’ result for band-limited functions is well known and takes different forms, the most famous being the Heisenberg uncertainty principle. It is also the basis of what is known as the ‘folk’ version of the so-called Sampling Theorem. Roughly stated, the Sampling Theorem asserts that one can recover (up to negligible error) a band limited signal from $2BT$ sampling or basis functions. In this form the folk Sampling Theorem is usually attributed to Whittaker and Shannon, who first used it in the 1940’s in an empirical fashion. A rigorous justification was not available until Landau and Pollak (1962), motivated by the use of time and band limited functions in communication theory, achieved their result through a very extensive study of time and band limited functions. Their results apply to the ensemble of square-integrable real functions, which are said to be approximately variable-limited to $|x| \leq T/2$, that is:

$$\frac{\int_{|x| \leq T/2} |f(x)|^2 dx}{\int_{-\infty}^{\infty} |f(x)|^2 dx} = 1 - \varepsilon^2_T$$

where one assumes that $\varepsilon^2_T$ is small and, under normalization of the unit area, measures the degree to which $f(x)$ fails to be concentrated in the interval $|x| \leq T/2$. Similarly, one assumes that the Fourier transform $f(\omega)$ of a function $f(x)$ nearly vanishes for $|\omega| \leq 2\pi B$. Under these conditions, Landau and Pollak (1962) show there is a linear model based on $2BT$ independent functions that approximate any function with a frequency limit given by $B$, with an error comparable to $\varepsilon^2_T$. This linear model is defined by wavelets described by the so-called ‘prolate spheroidal functions,’ which are the optimal base for this type of functions. Any other orthonormal set is not as efficient, in that it can be shown that the number of required functions grows beyond $2BT$ quite rapidly. We call this a strict relationship between band limitation and sampling basis.