Abstract

It is well known that the peripheral visual system declines with age: the yellowing of the lens causes a selective reduction of short-wavelength light and sensitivity losses occur in the cone receptor mechanisms. At the same time, our subjective experience of colour does not change with age. The main purpose of this large-scale study (n = 185) covering a wide age range of colour-normal observers (18–75 years of age) was to assess the extent to which the human visual system is able to compensate for the changes in the optical media and at which level of processing this compensation is likely to occur. We report two main results: (1) Supra-threshold parafoveal colour perception remains largely unaffected by the age-related changes in the optical media (yellowing of the lens) whereas our ability to discriminate between small colour differences is compromised with an increase in age. (2) Significant changes in colour appearance are only found for unique green settings under daylight viewing condition which is consistent with the idea that the yellow-blue mechanism is most affected by an increase in age due to selective attenuation of short-wavelength light. The data on the invariance of hue perception, in conjunction with the age-related decline in chromatic sensitivity, provides evidence for compensatory mechanisms that enable colour-normal human observers a large degree of colour constancy across the life span. These compensatory mechanisms are likely to originate at cortical sites.

Introduction

The purpose of this study was to assess in a large sample of adult colour-normal observers whether higher-order colour appearance mechanisms are affected by ageing. Human colour vision starts in the retinal photoreceptors, where the lights is absorbed by the long-, medium- and short-wavelength-sensitive (L-, M-, or S-) cones [1]. The cone outputs are recombined in post-receptoral cone-opponent retinal channels and their spectral tuning is inherited by the Lateral Geniculate Nucleus (LGN), a sub-cortical visual relay station [2]. These early cone-opponent mechanisms feed into cortical colour mechanisms with different spectral properties that have not been fully characterised [3–5]. The cortical colour mechanisms that mediate the unique hues have been characterised using behavioural methods [6–11], but the neural basis of the unique hues is still an unresolved issue [12,13]. In our study we use unique hue judgements as a way to gauge higher-order colour appearance mechanisms since unique hue judgments allow us to obtain absolute appearance judgements without the need for a reference.

Previous studies suggest that hue perception is fairly constant across the life span [14,15]. At the same time age-related changes in the optical media (yellowing of the lens; [16]) occur; the number of retinal ganglion cells is likely to decrease with age [17], at the cortical level losses in dendritic spines and changes in the morphology of myelin sheaths and axons lead to a decrease in neural efficiency [18]. A possible explanation for hue constancy with an increase in age is that the higher-order colour mechanisms that mediate hue perception take the difference between the cone signals and are therefore not greatly affected by a signal loss in the peripheral system [14]. For example, an L-M mechanism will stay roughly constant with age, if the decline in the L and M cone signal is parallel. Here we test this hypothesis directly by assessing hue perception in a large-scale study (n = 185) for observers spanning a wide age range (18–75 years of age) and then compare the hue settings with the prediction derived from the changes in the optical media. Our second aim was to evaluate whether adaptation mechanisms that ensure constancy across different ambient viewing conditions, are affected by age [17]. We find that, to a large extent, hue perception is invariant with age; the direction but not the magnitude of the small observed age-related hue changes are predicted by the yellowing of the lens. Our results are consistent with compensatory mechanisms that operate on higher-order colour vision but not on early colour discrimination mechanisms.

Methods

We report a series of three experiments conducted with the same set of colour-normal observers (n = 185) covering a wide age range (18–75 y.o.a). All observers made settings on a different occasion: under dark viewing conditions, under a day-light simulator (close to D65), or under typical cool white office lighting (CWF).
Stimuli and procedure

To obtain settings of the unique hues we used a modified hue selection task [6]. Patches of slightly different hues were arranged along an annulus at constant eccentricity (Figure 1a) and the task of the observer was to select a patch that contains neither yellow nor blue (to obtain unique red and green). Unique yellow (blue) was obtained by asking observers to select a patch that contains neither red nor green. Each patch had a diameter of 2° of visual angle and was presented at an eccentricity of 4°. Each unique hue was determined at nine combinations of different saturation and brightness levels (ranging from 0.5 cd/m² to 60 cd/m²). The exact luminance values varied slightly between the different hues so as to maximise the available monitor gamut (Figure 1b). Each of these nine settings was repeated three times. In total, 360 test colours (4 unique hues×9 combinations of different saturation-brightness levels×10 colour patches per test) were displayed. On a particular trial, the patches were of the same saturation and brightness, and only differed in hue (for an example, see Figure 1a), in order to facilitate the task for the observer.

To obtain achromatic settings (neutral grey), the annulus consisted of greyish patches of a higher luminance (60 cd/m²) than the background and the observer was asked to select that patch that contained neither red nor green and, at the same, contained neither yellow nor blue. To obtain the achromatic settings, each trial consisted of two stages; first the greyish patches in the annulus contained small random modulations (around a theoretical achromatic point set to D65) along a reddish-greenish axis and the observer had to select that patch that contained neither red nor green. In the 2nd stage, the red-green coordinates were set to the selection obtained in the 1st stage and random modulations were along a yellow-bluish axis were added. In the 2nd stage the observer selected that greyish patch that appeared neither yellow nor blue. Each trial was repeated three times.

Figure 1. The modified hue selection task. (a) 2-deg colour patches (test stimuli) are presented at an eccentricity of 4 degrees. To obtain unique red (or green) the observer is asked to click on that patch that contains neither yellow nor blue. Similarly, to obtain unique yellow (or blue), an annulus consisting of yellowish (or bluish) patches is displayed and the observer is asked to click on that patch that is neither red nor green. In a previous experiment, we confirmed that the hue settings are not affected by the arrangement of the hue patches; randomised and ordered hue patches yield the same mean results. (b) The chromatic properties of the test stimuli are shown in a 3D solid in CIELUV space. The outermost white line indicates the gamut of our monitor. On a particular trial, hues of similar saturation and brightness were displayed in an annulus (see a). E.g. To obtain unique red, reddish hues lying on an arc in the CIELUV space, are displayed and the observer is asked to select that reddish hue that is neither yellow nor blue.

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Chromatic specifications

To calculate the cone absorptions in the long-, medium-, and short-wavelength-sensitive cones, we used the 2-deg cone fundamentals derived by Stockman and Sharpe [20]. The cone fundamentals were then scaled in terms of luminance using the new \( V^*(\lambda) \) (available at www.cvrl.org) such that luminance equals the sum of the L and M cone absorptions (Figure S1b). Then the appropriate scaling factor \( (k_m = 683) \) was applied to \( V^*(\lambda) \) to obtain candela units [21]. The phosphor spectra were measured at peak output (Figure S1a), upsampled to 1 nm spacing and multiplied with the scaled cone fundamentals to obtain the transformation from linearized RGB space to LMS space. All calculation performed in LMS cone space; to visualize the unique hue settings in a chromaticity diagram, we transform the LMS coordinates to the Boynton-McLeod (BML) chromaticity coordinates [2,22]. In the BML space, the x-axis corresponds to the difference between the L and M cone signals, normalized to luminance; the y-axis shows the modulation of the S cones, again normalized to luminance (Figure S1d).

Estimating the cone weightings

Colour-opponent mechanisms were first mentioned by Hering [23] who proposed that any hue can be described in terms of its redness or greenness and its yellowness or blueness. Red and green are opposite hues because they cannot—and under normal viewing conditions without retinal stabilisation—be elicited simultaneously by a single colour stimulus; the same is true for blue and yellow. Hering therefore postulated the existence of two colour-opponent channels coding red-green and yellow-blue sensations. Quantitative estimates of these two colour-opponent channels were first obtained by Jameson and Hurvich [24–26] using a hue cancellation technique. For example, when a subject adjusts the red and green component of a yellowish stimulus such that it contains neither red nor green, the red-green opponent channel (RG) is at equilibrium; consequently a red-green opponent mechanism produces a zero response for this yellowish stimulus (for details see [6]):

\[
w_L * L + w_M * M + w_S * S = RG_Y = 0 \tag{1}
\]

A yellowish stimulus with these differential L,M,S cone weights \( (w_L,w_M,w_S) \) is void of any red and green, since it silences the putative red-green mechanism (i.e. \( RG_Y = 0 \)); the stimulus is therefore referred to as ‘unique yellow’. Equation 1 defines a plane in three-dimensional LMS cone space. The vector \( (w_L,w_M,w_S) \) is orthogonal to this plane and is called the normal vector for this plane; it characterises the chromatic properties of the red-green mechanism \( RG_Y \). By definition, this mechanism is silenced by all yellowish colours on this plane and the plane is therefore the null plane for the \( RG_Y \) mechanism. We can derive an analogous equation for unique blue. A bluish stimulus that contains neither red nor green, produces a zero response in a red-green opponent channel:

\[
w_L * L + w_M * M + w_S * S = RG_B = 0 \tag{2}
\]

Unique red (Eq 3) and unique green (Eq. 4) are defined as colours that produce zero output in a yellow-blue opponent channel:

\[
w_L * L + w_M * M + w_S * S = YB_R = 0 \tag{3}
\]

\[
w_L * L + w_M * M + w_S * S = YB_G = 0 \tag{4}
\]

Altogether, we therefore estimate 12 coefficients, three for each unique hue, by a principal component analysis (PCA); the last eigenvector (explaining the least variance) is the normal vector. The length of the normal vector is normalised to unity (for details see [6]).

The Lens model by Pokorny et al

We used the lens model by Pokorny et al [16] to predict the changes in lens transmission as a function of age. To calculate the age-related relative loss in the three cone classes (L,M,S), the tabulated 2-deg cone fundamentals derived by Stockman and Sharpe [20] were multiplied with the linear lens transmission (up to 650 nm), for each year of age (Figure S1c). The resulting loss in the L,M,S cone signals a function of age is normalised to unity at 32 years of age (Figure 2).

Results

1. Effect of age on colour appearance

To evaluate whether hue perception changes with age and whether these changes are predicted by the age-related changes in the optical media, we proceed as follows.

(1) From the unique hue settings, we first estimate the cone weightings of the mechanisms that mediate the four unique hues by a principal component analysis (PCA); the last eigenvector (explaining the least variance) is the normal vector. The length of the normal vector is normalised to unity (for details see [6]).

(2) We then predict the cone weightings using the lens model by Pokorny et al [16]. By comparing the observed with the predicted cone weightings as a function of age, we can ascertain whether...
changes in hue perception are explained by the yellowing of the lens.

Figure 3 shows the observed cone weightings under the dark viewing conditions. The cone weightings (in columns: L,M,S) are plotted as a function of age, for the chromatic mechanisms associated with the four unique hues (Figure 3: in rows: YBR, YBG, RGY, RGb). Observed cone weightings are shown by open circles; the best fitting regression lines for the observed cone weightings are indicated by dashed lines. For example, the first row shows the cone weightings of the chromatic mechanism that is silenced by unique red, denoted YBR (cf Eq. 3); the leftmost graph shows the L cone weighting (wL) for this chromatic YB mechanism; the 2nd column indicates the M cone weightings (wM) and the 3rd column the S cone weightings (wS) of the mechanism that is silenced by unique red (YBR). Row 2 shows the cone weightings for the chromatic mechanism that is at equilibrium for unique green (YBc; cf. Eq 4); unique green is nulling a mechanism that takes the difference between the L and S cone signals, with very little M cone input. The mechanism that is silenced by unique red (YBc; cf Eq. 3) has a very different chromatic tuning; it has an L cone coefficient close to zero; indicating that this yellow-blue mechanism is parallel to the L cone plane and is primarily driven by the difference between the M cone and the S cone inputs. The observation that not a single linear mechanism is silenced by unique red and green confirms previous findings [6,26].

Cone weightings for the mechanisms that are silenced by unique yellow and unique blue are shown in rows 3 and 4, respectively (cf. Eq. 1 and 2). As expected, the cone weightings for these two hues are very similar, consistent with the hypothesis that unique yellow and blue are generated by silencing a single, approximately linear mechanism [25,26], that combines the difference between the L and S cone signals, with very little M cone input. The mechanism that is silenced by unique red (YBc; cf Eq. 3) has a very different chromatic tuning; it has an L cone coefficient close to zero; indicating that this yellow-blue mechanism is parallel to the L cone plane and is primarily driven by the difference between the M cone and the S cone inputs. Identical scales on the y-axes allow for a comparison of the spread of the cone weightings (cf row 3 and 4); for all three cone classes, the spread for unique blue is smaller than for unique yellow indicated that observers are able to make more precise settings for unique blue in comparison to unique yellow. Most importantly, cone weightings are invariant with age for all hues and all cone classes: the slope of the regression lines (dotted lines in Figure 3) do not differ significantly (p<0.05) from zero for any hue and for any cone class.

The solid lines in Figure 3 indicate the predicted cone weightings under the lens model (see Figure 2; Experimental Procedures). To arrive at these predictions, we adjusted the L,M,S cone co-ordinates of the hue settings to incorporate the age-dependent lens transmission. We then calculated the normal vectors for each hue (see Methods: Estimating the Cone Weightings) which reflects the chromatic properties of the mechanism that is being nulled by a particular hue. For example, for the chromatic mechanism that is silenced by unique red (denoted YBr in Figure 3, row 1) the lens model predicts a greater S cone weighting with an increase in age, to stay at equilibrium. This is explained by the significant decrease in short-wavelength light passing through the lens when age increases (Figure 2, and Figure S1c). To compensate for the weaker S cone signal, this mechanism would need to increase the weighting of the S cone input. The discrepancy between the predicted and the observed regression lines reflect the amount of compensation of the hue mechanisms. All predicted regression lines differ significantly from zero, at p<0.001, whereas none of the observed slopes differs from zero. Slopes, coefficients of determination (R^2), and the corresponding p-values are provided in Table 1 for all hues and all cone classes, for the observed cone weightings (OBS) and the predicted cone weightings (PRED). The first part of Table 1 describes the statistical results when stimuli were viewed under dark conditions.

Cones weightings obtained under the dark viewing conditions are shown as a function of age, for the chromatic mechanisms (YBR, YBG, RGY, RGb) that are silenced by unique red (row 1), green (row 2), yellow (row 3), and blue (row 4) are plotted as open symbols (see Eq 1–4 for details). Dashed lines (OBS) indicate the best fitting straight line to the data; solid lines (PRED) are the predictions based on the lens model (see Experimental Procedures). None of the observed slopes differ significantly from zero (p<0.05), indicating that the unique hue settings do not change with age. All slopes predicted from the lens model, however, differ from zero (p<0.001).

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Table 1. Coefficients of determination (R^2) and p-values for all hues and all cone classes. Predicted and observed cone weightings (OBS) under the dark viewing conditions are plotted as a function of age (columns: L,M,S) for the four chromatic mechanisms (YBR, YBG, RGY, RGb) that are silenced by unique red (row 1), green (row 2), yellow (row 3), and blue (row 4). The cone weightings (in columns: L,M,S) are plotted as a function of age, for the chromatic mechanisms associated with the four unique hues (Figure 3: in rows: YBR, YBG, RGY, RGb). Observed cone weightings are shown by open circles; the best fitting regression lines for the observed cone weightings are indicated by dashed lines. For example, the first row shows the cone weightings of the chromatic mechanism that is silenced by unique red, denoted YBR (cf Eq. 3); the leftmost graph shows the L cone weighting (wL) for this chromatic YB mechanism; the 2nd column indicates the M cone weightings (wM) and the 3rd column the S cone weightings (wS) of the mechanism that is silenced by unique red (YBR). Row 2 shows the cone weightings for the chromatic mechanism that is at equilibrium for unique green (YBc; cf. Eq 4); unique green is nulling a mechanism that takes the difference between the L and S cone signals, with very little M cone input. The mechanism that is silenced by unique red (YBc; cf Eq. 3) has a very different chromatic tuning; it has an L cone coefficient close to zero; indicating that this yellow-blue mechanism is parallel to the L cone plane and is primarily driven by the difference between the M cone and the S cone inputs. The observation that not a single linear mechanism is silenced by unique red and green confirms previous findings [6,26].

Figure 3. Cone weightings as a function of age. The cone weightings obtained under the dark viewing conditions are shown as a function of age. The LMS cone weightings (columns: L,M,S) of the four chromatic mechanisms (YBR, YBG, RGY, RGb) that are silenced by unique red (row 1), green (row 2), yellow (row 3), and blue (row 4) are plotted as open symbols (see Eq 1–4 for details). Dashed lines (OBS) indicate the best fitting straight line to the data; solid lines (PRED) are the predictions based on the lens model (see Experimental Procedures). None of the observed slopes differ significantly from zero (p<0.05), indicating that the unique hue settings do not change with age. All slopes predicted from the lens model, however, differ from zero (p<0.001).
settings since the optical density of the lens is not changing the transmission of short-wavelength light dramatically (Figure S1c). The older age group (>60 y.o.a., Figure 4b), however, shows a significant age shift in the absence of compensatory mechanisms. E.g. blue hues should shift downwards in the BML diagram towards yellow-green hues; green is predicted to shift towards yellow. Achromatic settings (indicated by open grey symbols) are predicted to appear yellowish (indicated by a solid grey triangle in Figure 4b).

While we find no evidence for age-related hue changes, the chromatic sensitivity, as assessed with the Cambridge Trivector Test [19] declines significantly (p<0.001) with age along the protan, the deutan, and the tritan line (Figure 5). The most robust age-related sensitivity changes are observed along the tritanopic confusion line (rightmost panel in Figure 5), as expected from the differential loss of short-wavelength light in the ageing lens (see also Figure S1c).

### Table 1. Dependency of cone weightings on Observer age.

| ILLUM       | HUE  | L COEFF SLOPE | R²   | p   | M COEFF SLOPE | R²   | p   | S COEFF SLOPE | R²   | p   |
|-------------|------|---------------|------|-----|---------------|------|-----|---------------|------|-----|
| DARK        | YBr  | -1.57E-05     | 0.002| 0.539| 4.00E-05      | 0.003| 0.426| 2.73E-06      | 0.003| 0.466|
|             | OBS  |               |      |      |               |      |      |               |      |      |
|             | PRED | -1.14E-04     | 0.118| 0.000| 7.17E-04      | 0.459| 0.000| 5.20E-05      | 0.501| 0.000|
| YG          | OBS  | -8.34E-05     | 0.005| 0.343| 1.21E-04      | 0.033| 0.435| -9.60E-06     | 0.001| 0.726|
|             | PRED | -1.06E-03     | 0.461| 0.000| 1.17E-03      | 0.256| 0.000| -2.16E-04     | 0.267| 0.000|
| RG          | OBS  | -1.98E-04     | 0.012| 0.138| 4.17E-04      | 0.014| 0.103| 1.72E-04      | 0.011| 0.152|
|             | PRED | -1.67E-03     | 0.475| 0.000| 3.24E-03      | 0.481| 0.000| 1.36E-03      | 0.410| 0.000|
| RGa         | OBS  | -5.55E-07     | 0.000| 0.993| 3.40E-05      | 0.001| 0.743| 2.66E-06      | 0.000| 0.938|
|             | PRED | -1.09E-03     | 0.658| 0.000| 2.30E-03      | 0.728| 0.000| 6.76E-04      | 0.694| 0.000|
| DAYLIGHT    | YBr  | -1.15E-05     | 0.002| 0.564| 2.84E-05      | 0.003| 0.447| 1.07E-07      | 0.000| 0.973|
|             | OBS  |               |      |      |               |      |      |               |      |      |
|             | PRED | -1.10E-04     | 0.156| 0.000| 7.15E-04      | 0.573| 0.000| 5.18E-05      | 0.594| 0.000|
| YG          | OBS  | -2.46E-04     | 0.031| 0.006| 4.03E-04      | 0.026| 0.028*| -6.25E-05     | 0.023| 0.040*|
|             | PRED | -1.16E-03     | 0.422| 0.000| 1.36E-03      | 0.238| 0.000| -2.37E-04     | 0.261| 0.000|
| RG          | OBS  | -1.38E-04     | 0.004| 0.379| 2.99E-04      | 0.006| 0.313| 1.25E-04      | 0.003| 0.430|
|             | PRED | -1.65E-03     | 0.388| 0.000| 3.17E-03      | 0.398| 0.000| 1.37E-03      | 0.299| 0.000|
| RGa         | OBS  | 4.76E-05      | 0.003| 0.424| -5.32E-05     | 0.001| 0.610| 4.27E-06      | 0.000| 0.895|
|             | PRED | -1.05E-03     | 0.636| 0.000| 2.21E-03      | 0.710| 0.000| 6.47E-04      | 0.665| 0.000|
| COOL WHITE  | YBr  | -1.94E-05     | 0.004| 0.367| 2.78E-05      | 0.002| 0.577| 2.10E-06      | 0.002| 0.541|
| FLOURESCENT | OBS  |               |      |      |               |      |      |               |      |      |
|             | PRED | -1.17E-04     | 0.148| 0.000| 7.34E-04      | 0.540| 0.000| 5.30E-05      | 0.556| 0.000|
| YG          | OBS  | -1.73E-04     | 0.018| 0.069| 2.52E-04      | 0.012| 0.138| -3.24E-05     | 0.006| 0.290|
|             | PRED | -1.11E-03     | 0.448| 0.000| 1.25E-03      | 0.243| 0.000| -2.32E-04     | 0.251| 0.000|
| RG          | OBS  | -1.42E-04     | 0.006| 0.299| 2.95E-04      | 0.007| 0.251| 1.06E-04      | 0.004| 0.397|
|             | PRED | -1.60E-03     | 0.443| 0.000| 3.10E-03      | 0.456| 0.000| 1.26E-03      | 0.377| 0.000|
| RGa         | OBS  | -4.54E-06     | 0.000| 0.925| 3.28E-05      | 0.001| 0.699| 5.64E-06      | 0.000| 0.839|
|             | PRED | -1.04E-03     | 0.700| 0.000| 2.20E-03      | 0.758| 0.000| 6.34E-04      | 0.729| 0.000|

Slopes, coefficients of determination (R²) and the p-values for observed and predicted cone weightings (L COEFF, M COEFF, S COEFF) for all hues and all ambient illumination conditions (DARK, DAYLIGHT, COOL WHITE FLOURESCENT). Under the null hypothesis all slopes are zero, indicating no age dependency. All predicted slopes differ from zero (p<0.001; rows labelled with PRED). Observed slopes (rows labelled with OBS) are significantly different from zero (p<0.05) only for the unique green settings under D65. For all other hues and ambient illumination settings we find no age-related changes in perceived hue.

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2. Effect of ambient illumination on unique hue settings for different age groups

Our second aim was to assess whether the adaptation mechanisms that underlie the hue constancy across age operate in a similar manner for different ambient viewing conditions. Figures 6 and 7 show the cone weightings obtained under ambient illumination condition close to D65 (daylight) and under cool white fluorescent lighting (CWF), respectively. In contrast to the dark viewing condition, green settings change slightly with age under adaptation to D65. Slopes, coefficients of determination and p-values are listed in Table 1 (Daylight (D65), row for YBr); observed slopes that differ significantly from zero are indicated with a star (*) in Table 1 and in Figure 6 (row 2: p<0.05 for L, M and S cone weightings). The direction of the age-related changes in the observed L,M,S coefficients (dashed lines) is predicted by the lens model (solid line) but not the magnitude. The lens model predicts greater changes in the coefficients than is observed in the data.

Under adaptation to CWF, the results mirror those for dark adaptation (Figure 7; Table 1, lower part): cone coefficients are...
independent of age (p > 0.05 for all hue settings) whereas the lens model predicts highly significant age-related changes. For all ambient illumination conditions the lens model predicts the direction of the observed changes but not the magnitude. Table 1 shows that the observed (OBS) and predicted slopes (PRED) as a function of age; in all cases the sign of the slope is the same; but only for the green settings under adaptation to daylight do these age-related hue changes reach statistical significance.

Discussion

Our main finding is that colour appearance mechanisms are to large extent unaffected by the known age-related changes in the optical media (yellowing of the lens) whereas the ability to discriminate between small colour differences is compromised with an increase in age.

Figure 4. Comparison between the lower (<30 y.o.a.) and the upper age group (>60 y.o.a). (a) Observed unique hue settings for the dark viewing conditions are shown as circles in Boynton-MacLeod chromaticity diagram for the younger age group (<30 y.o.a). Symbol size is proportional to the number of observations per data point. Dotted lines indicate the observed unique hue lines (summarised by the 1st principal component); the solid lines denote the predicted unique hue lines assuming the lens model by Pokorny et al. The grey triangle is the average prediction for neutral grey settings, assuming the same lens model. For the younger age group, only very small hue shifts are expected assuming the lens model. All predictions are made with respect to a 32 old observer. (b). For the older age group (>60 y.o.a) large hue shifts are predicted under the lens model. The difference between the observed and predicted hue lines indicates the amount of compensation present in these hue judgements. doi:10.1371/journal.pone.0063921.g004

Werner and colleagues [14] suggested that the approximate hue constancy across the life span could be explained by a concurrent parallel decline in cone signals. A mechanism that takes the difference between the L and M cone signals, for example, will not be greatly affected by a decline in both the L and M cone outputs. We tested this hypothesis directly by calculating the cone weightings for each of the four unique hue mechanisms (Eq. 1–4) and then used a lens model to predict how perceived hues should change given the changes in the lens transmission for different wavelengths. The predictions for the hue changes expected under the lens model (Figure 2) are shown by solid lines in Figures 3, 6, and 7 (cf. Table 1 for statistical regression results). While the directions of the predictions are consistent with the data, e.g. for unique green (row 2 in figures 3, 6 and 7, the lens model predicts a decrease in the L and the S cone weightings and an increase in the M cone weightings, the magnitude of the observed

Figure 5. Chromatic sensitivity as a function of age. Thresholds along the protan (left most), deutan (middle) and tritan line (right most) is plotted as function of age for all 185 observers. A linear regression analysis revealed that the slope for all three sets of thresholds differs significantly from zero: Protan: slope = 0.004 R² = 0.119 p < 0.001; Deutan: slope = 0.003 R² = 0.139 p < 0.001; Tritan: slope = 0.254 R² = 0.258 p < 0.001. doi:10.1371/journal.pone.0063921.g005
hue changes is significantly smaller than the observed age-related cone weightings. We therefore conclude, in line with Webster et al. [27], that hue constancy across the life span is not a simple consequence of the differential cone combinations of the higher-order chromatic mechanisms.

The second possibility is that the human visual system can adjust the cone weightings of the chromatic mechanisms over the lifespan and thereby compensate for a decline in peripheral cone signals. Our data are consistent with this hypothesis. The decline in sensitivity along the protan, deutan and tritan line we documented in the same set of observers (Figure 5) suggests that these compensatory mechanisms are likely to arise at a late, probably cortical site and that these higher-order appearance mechanisms operate in parallel to the discrimination mechanisms revealed in the chromatic sensitivity task. The decline in chromatic discrimination sensitivity is consistent with the deutan and tritan decline reported for the 50+ age groups [28–30].

This dissociation between discrimination and appearance mechanisms is supported by Neitz and colleagues [31,32] who showed that shifts in unique yellow induced by long-term changes in the chromatic environment are not due to receptor or subcortical changes, but must be of cortical origin, probably after chromatic information from both eyes has been integrated. The question remains how these higher order colour mechanisms receive feedback on the strength of their cone inputs. Neitz et al’s unique yellow data are consistent with the hypothesis that the gains of the L and M cones are adjusted such that the red-green opponent mechanism is at equilibrium for the average daylight

Figure 6. Cone weightings obtained under D65. Details are as in Figure 3. Under D65, the cone coefficients associated with unique green are age-dependent (p<0.05; for L,M,S cone coefficients). Observed slopes that differ significantly from zero are indicated with a star (*). All predicted slopes are different from zero (p<0.001). The direction of the observed age-related changes (dashed lines; row 2) are predicted by the lens model (solid lines), but not the magnitude (cf Table 1).

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Figure 7. Cone weightings obtained under CWF. Details are as in Figure 1. As for the dark viewing condition, the observed coefficients do not depend on age (p>0.05), whereas the lens model predicts significant age-related changes (p<0.001).

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[33], but this recalibration is by no means complete [34]. Our results provide further evidence that the brain uses information about the statistical properties of our chromatic environment to adjust the weighting of the receptor signals to achieve hue constancy across the life span.

Our second aim was to assess whether there is a differential age-related effect of ambient illumination on hue constancy. Comparing the cone weightings under the dark viewing condition (Figure 3) with those under adaptation to D65 (Figure 6) and CWF (Figure 7) reveals that the mean unique hue settings across the entire sample are not affected by the illumination conditions (see also Figure S2 and S3). A closer look at the age dependency of the observed cone weightings (Table 1) shows that there is a differential effect of adaptation on hue constancy: only under daylight (D65) adaptation do the age-related changes in the green settings reach statistical significance (Table 1; Figure 6, row 2); what appears uniquely green for young observers appears more yellowish for older observers. Older observers require more S cone input to achieve unique green when the settings are obtained under simulated daylight, but still much less than predicted by the lens model. This is consistent with the idea that the yellow-blue mechanism (YBG) which is silenced by the unique green settings is most affected by the yellowing of the lens (Eq. 2; [35]). Similar conclusions were reached by Okajima & Takase [36] who assessed age-related colour changes using Munsell Colour chips instead of self-luminous lights.

If the visual system were able to fully compensate for the changes in the optical media, the observed cone weightings should not vary with age. We find that, under most of the tested conditions, this is the case; only under adaptation to daylight (D65), green hues change slightly with age.

In summary, we find evidence for compensatory mechanisms operating on higher-order colour functions and thereby ensuring that hue remains approximately constant despite the known age-related changes in the lens. The concurrent age-related decline in the chromatic discrimination sensitivity suggests that the neural site of these compensatory mechanisms is probably cortical; the underlying mechanism is still poorly understood, but is consistent with the idea that it is based on invariant sources in our visual environment.

Supporting Information

Figure S1 Spectra, cone fundamentals and lens model.
(a) The Spectral radiation distribution of the CRT phosphors b) The Stockman-Sharpe cone fundamentals used to obtain the lens predictions c) Lens transmission as a function of wavelength, based on the Pokorny et al. lens model. Each line indicates the transmission of a particular age group, from 20 to 80 y.o.a. From top to bottom. (d) The Boynton-MacLeod Coordinates derived from the Stockmann-Sharpe cone fundamentals.

Figure S2 Hue settings for the lower and the upper age group under D65. Details as in Figure 4 (a) Observed unique hue settings for D65 are shown as circles in Boynton-MacLeod chromaticity diagram for the younger age group (<30 y.o.a). Symbol size is proportional to the number of observations per data point. Dotted lines indicate the observed unique hue lines (summarised by the 1st principal component); the solid lines denote the predicted unique hue lines assuming the lens model by Pokorny et al. (1987). The grey triangle is the average prediction for neutral grey settings, assuming the same lens model. For the younger age group, only very small hue shifts are expected assuming the lens model. All predictions are made with respect to a 32 old observer. (b) For the older age group (>60 y.o.a) large hue shifts are predicted under the lens model.

Figure S3 Hue settings for the lower and the upper age group under CWF. Details as in Figure 4. (a) Observed unique hue settings under CWF are shown as circles in Boynton-MacLeod chromaticity diagram for the younger age group (<30 y.o.a). Symbol size is proportional to the number of observations per data point. Dotted lines indicate the observed unique hue lines (summarised by the 1st principal component); the solid lines denote the predicted unique hue lines assuming the lens model by Pokorny et al. (1987). The grey triangle is the average prediction for neutral grey settings, assuming the same lens model. For the younger age group, only very small hue shifts are expected assuming the lens model. All predictions are made with respect to a 32-year old observer. (b) For the older age group (>60 y.o.a) large hue shifts are predicted under the lens model.

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Author Contributions

Conceived and designed the experiments: SW. Performed the experiments: SW. Analyzed the data: SW. Wrote the paper: SW.

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