Reflection-based imaging of macular pigment distributions in infants and children

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Abstract. We have developed a reflection-based capability of the RetCam® platform, an FDA-cleared pediatric retinal-imaging instrument, for the purpose of measuring macular pigment levels as well as their spatial distributions in infants and children. Our modifications include narrow-band blue-wavelength excitation of the macular pigment absorption in combination with spectrally selective blue-wavelength readout of the reflection signals received by the instrument’s CCD detector array. Furthermore, an algorithm is developed that allows the computation of optical density maps for the macular pigment relative to peripheral retinal areas. This made it possible for the first time to directly measure macular pigment levels and their spatial features in the developing human retina. In contrast to adults, infants with measurable pigment levels had almost exclusively a narrow, circularly symmetric, pigment distribution. The described methodology holds promise for future investigations into the role of macular pigment in the developing human retina and the effect of dietary interventions in diseases resulting from a lack of normal carotenoid levels.

Keywords: macular pigment; carotenoids; reflection spectroscopy; retinal imaging; newborns.

Paper 130639R received Sep. 4, 2013; revised manuscript received Oct. 8, 2013; accepted for publication Oct. 9, 2013; published online Nov. 6, 2013.

1 Introduction

The healthy adult human retina contains high concentrations of blue-light-absorbing carotenoid compounds in its macular region, which is the ∼1-mm-diameter central tissue location concentrated with the cone photoreceptors that are responsible for high-acuity color vision. Known as macular pigment, MP, they are constituted of three carotenoid compounds: lutein, zeaxanthin, and meso-zeaxanthin, are located anterior to the photoreceptor outer segments and the retinal pigment epithelium (RPE),1,2 and are therefore thought to shield these vulnerable tissues from light-induced damage by effectively absorbing the photoionizing short wavelengths of ambient light otherwise reaching the deeper retinal layers. The MP carotenoids are also thought to directly protect the tissue cells in their immediate vicinity through their well-known function as antioxidants. Much research carried out over the last two decades has investigated the role of the macular carotenoids in the prevention of age-related macular degeneration (AMD),3-5 and more recently also their role in the improvement of visual acuity via reduction of deleterious glare effects is investigated.6

MP is very likely to play an important role also in early phases of foveal development, in the enhancement of infant visual acuity, and in the protection against light-induced oxidative damage,7 and it may be essential also for the development and/or maintenance of a normal distribution of RPE cells.8 Furthermore, the development of AMD in the elderly is potentially linked to phototoxic light exposure and oxidative stress accumulated over the lifetime, including the childhood years,9 in which oxidation-induced, potentially harmful, photosensitive compounds such as lipofuscin can already be generated in high concentrations.10 Without knowledge of normal MP levels and their spatial distributions in infants and children, further progress on their potentially beneficial roles cannot proceed. Motivated by this challenge, we have developed a novel reflection-based capability of the RetCam® platform, an FDA-cleared pediatric retinal-imaging instrument (Clarity Medical Systems, Pleasanton, California) widely used for the collection of fundus images of the infant retina for general tissue health diagnostics purposes, new for the purpose of measuring macular pigment levels as well as their spatial distributions in infants and children.

Several methodologies for optical MP assessment exist already for adults including the widely used psychophysical method of heterochromatic flicker photometry,11 and three quantitative imaging methods based on direct detection of the MP carotenoids via resonance Raman spectroscopy (RRS),12,13 indirect detection via lipofuscin fluorescence excitation spectroscopy, also known as “autofluorescence” spectroscopy,14,15 and detection via fundus reflection spectroscopy (FRS).16-18 A reflection-based imaging approach with blue-light excitation is implemented in a commercially available fundus camera (Visucam, Zeiss Meditec, Jena Germany).19 However, none of the described methodologies is applicable to infants, since besides the lack of FDA clearance for any of the corresponding platforms, infants are either not able to participate in the measurement or they would be exposed to risky high-laser light intensities in view of the anticipated low MP levels (RRS and laser-based FRS).

2 Reflection-Based Imaging with the RetCam®

The RetCam® is a fully integrated digital imaging system permitting the capture of wide-field, high-resolution, video streams...
of the infant retina, which facilitates optimum visualization of topographic retinal features. Importantly, it also offers the possibility to capture and store high-resolution still images. A usable RetCam® fundus image can be obtained within a few minutes in the anesthetized infant eye. The instrument is fiber-coupled to hand-held light delivery and collection modules that are brought in direct contact with the cornea of the infant. The contact module routes the excitation light in a peripheral, ring-shaped, illumination geometry to a detachable front lens and detects the reflected light with a centrally positioned CCD camera. An ophthalmic coupling gel is used to fill any remaining spatial gaps between the contact module’s front lens and the subject’s cornea. Front lenses with varying focal lengths exist to achieve, respectively, 30, 80, 120, or 130 deg fields-of-view of the retina. The CCD camera consists of a dispersive 3-chip configuration, “RGB triplet,” with each spatially separated 640 × 480 pixel array having enhanced sensitivity in either the red (“R-chip”), green (“G-chip”), or blue (“B-chip”) spectral range.

Two optional light sources exist for excitation, a tungsten-halogen (“white-light source”) and a blue-filtered xenon arc lamps (“blue-light source”). The first one is typically used for general white-light fundus reflection imaging, and the latter, in combination with a blue-light suppressing barrier filter, for fluorescein angiography. The relatively high power levels for the continuous excitation light sources result in temporary bleaching of the vast majority of photoreceptor pigments.

The emission spectra of the two light sources, measured with a high-resolution spectrograph, are shown in Fig. 1, along with the absorption spectrum of MP in excised retinal tissue. The emission spectrum of the blue-filtered continuous xenon arc lamp, peaking near 485 nm, has a high-spectral overlap with the MP absorption, which reaches maximum strength at ~460 nm but still has strong vibronic sidebands near 430 and 490 nm. Since the emission spectrum of the blue-light source is much narrower than the MP absorption band, it is well suited for highly wavelength-selective blue-light reflection imaging of MP in a detection scheme where the instrument’s fluorescence angiography blue barrier filter is simply removed.

The MP detection concept is illustrated in Fig. 2. As customary in a simplified FRS scheme, we assume that the excitation light, \( I_{\text{exc}}(\lambda) \), traverses all anterior ocular media and the retinal layer system toward the sclera, where it is reflected, and that it subsequently traverses all layers again in a 180-deg detection geometry. After traversing the MP-containing central region, the detected light intensity is, in general, given by

\[
P_{\text{Mac}}(\lambda) = I_{\text{exc}}(\lambda) \cdot T_{\text{OM}}^2(\lambda) \cdot T_{\text{MP}}^2(\lambda) \cdot T_{\text{PRL}}^2(\lambda) \cdot T_{\text{RPE}}^2(\lambda) \\
\cdot T_{\text{CHR}}^2(\lambda) \cdot R(\lambda) + I_{\text{OM}}(\lambda_{\text{OM}}) + I_{\text{RPE}}(\lambda_{\text{RPE}}),
\]

where the \( T(\lambda) \)s are, respectively, the transmission of anterior optical media (OM), the MP-containing layer, the photoreceptor layer (PRL), the RPE, and the choroid (CHR). All the transmissions are squared due to the double-path light propagation. \( R(\lambda) \) is the reflection at the sclera and is assumed to be constant across the sclera including macular and peripheral sclera regions. \( I_{\text{OM}}(\lambda_{\text{OM}}) \) is the fluorescence intensity originating from the ocular media (mostly from the crystalline lens) at fluorescence wavelengths, \( \lambda_{\text{OM}} \), that are longer than the excitation wavelength, and \( I_{\text{RPE}}(\lambda_{\text{RPE}}) \) the fluorescence intensities of chromophores in the RPE layers such as lipofuscin.

Several simplifications of Eq. (1) are possible. First, we can assume that the white-light exposure of the retina, occurring during the fundus video-capturing mode of the measurement, which precedes the MP reflection imaging, will bleach the vast majority of photoreceptors, i.e., \( T_{\text{PRL}} = 100\% \). We also know that the ocular media of infants and children are typically highly transparent, i.e., \( T_{\text{OM}} = 100\% \), and that the inner limiting membrane (ILM) and nerve fiber layer (NFL) are also highly transparent. Furthermore, from an inspection of all RGB triplet pixel intensities for wavelengths longer than the excitation wavelength, achieved with the help of a long-pass wavelength filter, we saw that the fluorescence of the ocular media and lipofuscin in the RPE was negligible in infants and children. With these assumptions, Eq. (1) can be reduced to

\[
P_{\text{Mac}}(\lambda) = I_{\text{exc}}(\lambda) \cdot T_{\text{MP}}^2(\lambda) \cdot R(\lambda).
\]

The excitation intensity, \( I_{\text{exc}}(\lambda) \), and reflection coefficient, \( R(\lambda) \), cancel out by referencing the reflection detected from the macular region to the reflection detected from a peripheral retinal region. This yields a ratio

![Absorption spectrum of macular pigment (MP) and emission spectra of light sources provided in RetCam® platform. (a) Absorption spectrum of MP in excised retinal tissue, featuring characteristic maximum near 460 nm and vibronic sidebands near 430 and 490 nm; (b) unfiltered emission spectrum of halogen lamp “white-light source;” and (c) emission spectrum of blue-filtered continuous xenon arc lamp, peaking near 485 nm.](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics/116001-2)

![Schematic representation of the excitation and reflection light paths through the retinal layer system involved in our in vivo reflection-based measurements of MP in the infant retina. OM, ocular media; ILM, inner limiting membrane; NFL, nerve fiber layer; MP, macular pigment containing area; PRL, photoreceptor layer; RPE, retinal pigment epithelium; CHR, choroid; SC, sclera; \( I_{\text{exc}} \), excitation light from blue-filtered xenon lamp; \( I_{\text{Mac}} \), light intensity reflected by sclera and traversing MP-containing retinal area (macula); \( I_{\text{PRL}} \), light intensity reflected by cornea and traversing the region outside the macula (peripheral retina).](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics/116001-2)
In this ratioing procedure, we have to assume that chromophores existing and absorbing in the deeper retinal layers are evenly distributed across its posterior pole. Mainly, this includes melanin in the RPE and blood in the CHR. Furthermore, we neglect reflections at internal retinal layer interfaces, which can be detected with high-sensitivity optical coherence tomography detection schemes, but are roughly one order of magnitude lower than the reflection at the sclera, and since as Fresnel reflections, they vary only weakly with wavelength over the visible/NIR region. For the processing of the raw reflection images, we followed basically the same procedure used previously in our autofluorescence-based imaging method for the measurements of MP in adults. Briefly, this involves selecting several small peripheral pixel areas in the raw images that exhibit uniform illumination, then calculating an average pixel intensity for these areas, and finally using this average as a reference for the intensity of a small macular “test” pixel area.

The transmission of the excitation light at wavelength $\lambda$ can be expressed through the transmission at the peak of the MP absorption at 460 nm via

$$T_{\text{MP}}(\lambda) = T_{\text{MP}}(460 \text{ nm})^{K_{\text{MP}}(\lambda)},$$

where $K_{\text{MP}}(\lambda)$ is the extinction coefficient. With this expression, the right side of Eq. (3) reduces to $[T_{\text{MP}}(460 \text{ nm}) / T_{\text{MP}}(460 \text{ nm})]^{2[K_{\text{MP}}(\lambda)]}$. With the optical density (OD) equaling $\log(1/T)$, the MP optical density (MPOD) difference between macula and periphery can be written as

$$\text{MPOD} = \text{OD}_{\text{Mac}}(460 \text{ nm}) - \text{OD}_{\text{Per}}(460 \text{ nm})$$

$$= \frac{1}{2 \Delta K} \left\{ \log \left( \frac{T_{\text{refl}}(\lambda)}{T_{\text{refl}}^{\text{Mac}}(\lambda)} \right) \right\},$$

i.e., this difference can be approximated as the log ratio between foveal and peripheral reflection intensities when recorded with the blue subset of the CCD camera pixel array. Since the blue-excitation light of the RetCam® peaks at 485 nm, it overlaps the MP absorption spectrum at only a slightly reduced extinction relative to the maximum absorption at 460 nm (see Fig. 2), resulting in a $\frac{1}{2K}$ value of 1.15 for all MPOD calculations.

The imaging of an infant’s retina started with the recording of several fundus images under white-light exposure to find the proper alignment of the macula relative to the camera’s light delivery and collection modules. Subsequently, the blue-light source was quickly fiber-coupled into the contact module, and several still images were recorded. The instrument’s 80-deg front lens was found to be optimal for our purposes, since due to its relatively high field-of-view and resulting high contrast of retinal features, it facilitated “finding” the macula, which in the infant eye generally has only weakly distinguished topological features compared with the peripheral retina. The blue-light excitation disk size resulting with this lens had a diameter of ~1 cm at the cornea. The light power was ~10 mW in these measurements. Typical total measurement times were several minutes including excitation with white light for about 1 to 2 min in order to “find” the macula, and subsequent excitation with blue light for a fraction of a minute to generate the reflection image of interest. These light exposures are well below the 5-min safety limit for the RetCam’s blue-light source. To develop the reflection method for clinical use and to survey the main MP properties in infants and children, we enrolled 51 subjects ranging in age from prematurity up to age 6. Parental consent was obtained in all cases under an IRB-approved protocol that complied with the Declaration of Helsinki. Subjects were imaged with pupil dilation according to the standard of care for their age. Preterm infants were imaged at the same time they were scheduled for retinopathy of prematurity screening, all others during examinations scheduled for other medical reasons. At the time of enrollments, all demographic information was collected including age, sex, race, and ethnicity.

In Figs. 3 and 4, we compare fundus images for a 4-month-old infant obtained, respectively, with white-light and blue-light excitations and differing color readouts of the CCD arrays. As can be seen from the white-light color fundus image of Fig. 3, recorded with full RGB readout of the three CCD arrays, the macular region is only vaguely discernible, since distinguishing topological features such as a foveal depression do not yet exist, and color changes due to yellow MP, expected in the macular region, are largely washed out. Basically, only blood vessels appear with high contrast in this operation mode, as desired for the intended routine fundus-imaging modes of the RetCam®. To enhance MP features, we derived 8-bit grayscale reflection and corresponding absorption images via selective B, G, and R-chip readouts, respectively. For better visual appearance of the images, illumination nonuniformities were removed from the raw images via fast Fourier transform (FFT) filtering (ImageJ software, NIH). All processed image results are shown in Figs. 3 and 4, clearly revealing enhanced visualization of the macular region.

Calculating apparent MPOD levels along the horizontal meridian through the macula, the highest level of ~0.1 is obtainable from the B-chip absorption image and 0.03 from the G-chip absorption image, roughly corresponding in relative strengths to the absorption behavior of macular pigment, which is stronger in the blue versus green. The R-chip image reveals absorption, too, most likely due to melanin. In principle, this would be a confounding factor in the measurement, but fortunately the OD is very low (smaller than about 0.01). Assuming that the melanin absorption is about a factor of 2 higher in the blue relative to the red wavelength region, the percentage melanin contribution to the MPOD would be at most 0.02/0.1 = 20% for this subject. When using blue-light imaging in combination with selective B-chip readout only, as shown in Fig. 4, a dim blue image with central dark MP disk results. The absence of blood vessel features demonstrates the selective sensitivity of the B-chip image to the yellow carotenoid chromophores. After carrying out identical image processing as described above, the MP-containing macular region reveals an MPOD density of ~0.2 in the macular region.
Fig. 3 Fundus imaging of 4-month-old infant with white-light excitation. Left: raw image obtained with combined RGB-chip readouts of CCD detector array. First column: reflection images derived from raw image via selective B, G, and R-chip readouts, respectively. Second column: B, G, and R absorption images derived from reflection images. For better visual appearance, all B, G, and R images were FFT filtered (intensity variations removed in small structures up to three pixels, and in large structures averaged over 40 pixel increments). Right column: MP optical density (MPOD) line profiles obtained from raw image B, G, and R chip intensities, respectively, along meridians running horizontally through the center of the macula.

Fig. 4 Fundus imaging of infant of Fig. 3 with blue-light excitation. Left: raw image obtained with combined RGB-chip readouts of CCD detector array. First column: reflection images derived from raw image via selective B- and G-chip readouts, respectively, and subsequent FFT filtering (same settings as for Fig. 3). Second column: B and G absorption images derived from reflection images. Right column: MPOD line profiles obtained along meridians running horizontally through the center of the macula. Note that the MPOD level of ~0.2 in the center of the MP distribution is about twice as high as the level obtained with white-light excitation (shown in Fig. 3).
B-chip image, which is about double the OD level obtainable with the white-light imaging of Fig. 3. The G- and R-chip images revealed no absorption, in agreement with the absence of excitation intensity in those wavelength regions.

Measurements of several dozen subjects showed that the MPOD imaging improved significantly with this approach with the percentage increases of MPOD levels relative to white-light imaging ranging from about 20% to 100%. Interestingly, the exact increase in MPOD levels is not a constant factor, but rather depends on the alignment of the contact module during the initial white-light imaging phase of the measurement. This effect is shown in Fig. 5, where two B-chip images and the corresponding line plots along the horizontal meridian through the center of the macula are compared for the same subject for white-light excitation. The apparent peak MPOD levels are \( \sim 0.18 \) in one case and \( \sim 0.08 \) in the other, thus differing by more than a factor of 2 and indicating a strong directional dependence of the contact module alignment on the apparent MPOD results. This directional sensitivity also manifests itself as a color change in the RGB chip images under white-light imaging, where the fundus images change from a normal orange/red color at optimum alignment (i.e., macula exactly centered in the image) to a yellowish/orange color for off-center locations. The described alignment effects are noticeable even if the macula is not too far off-center.

In most infants, all white- and blue-light reflection images also revealed a narrow, ring-shaped, high-intensity reflection feature surrounding the macula at a larger distance. Probably, this feature originates from spatial irregularities of the ILM or deeper retinal layers and is typically seen also in images of adult subjects.17 In the processing of the images for MPOD calculations, these highly reflecting areas were excluded.

Visualization of the spatial MP features is further facilitated by line plots along multiple meridians and by pseudocolor displays of the pixel intensities, as illustrated in Fig. 6, showing a peak MPOD level of \( \sim 0.4 \) in a particular subject with a nearly circularly symmetric MP distribution. Iterative measurements of MPOD levels for two subjects demonstrated a satisfactory high agreement.
repeatability of our methodology, as shown in Fig. 7, for the peak MPOD levels of the MP distributions in these two subjects. The standard deviation for each series was ∼5%.

The main MP image features for the measured population are illustrated in Fig. 8, displaying representative grayscaled and processed images for three infants aged 3, 0.5, and 0.6 years, respectively. Centered on the macula in each case, two images show nearly circularly symmetric MP distributions with different central levels, and one shows a nonsymmetrical distribution characterized by a strong central peak surrounded by a concentric ring of lower MP levels. Peak MP levels varied between 0 and 0.4 OD units in the measured subject population, with the highest levels reaching MP levels in adults. Out of 39 subjects with measurable MP levels, 37 had a relatively narrow, centrally peaked distribution, and only two subjects had a ring-shaped distribution. This behavior is different from about five distinguishable MP distribution patterns observable in healthy adults and AMD patients.

In all nine prematurely born infants, MP was found to be absent. In all these cases, a distinguishable macular region could not be detected, as seen from a representative B-chip image in Fig. 9. Instead of any MP-resembling feature, usually
only one, and sometimes two small, gray-shaded, retinal areas were seen for each subject, at roughly constant distances from the optical disk, but in locations with eccentricities too large for the normal macular location of an infant (too large by at least a factor of 2). Supporting the conclusion that these spots are not associated with yellow MP chromophores is our finding that these spots remain visible also when imaging with white-light excitation and selectively reading out only the G- and R-chip arrays. Such a spectral behavior would be consistent with a broadly absorbing chromophore, such as melanin, but its nature remains unclear at this time.

For all subjects with measurable MPOD levels in both eyes, the values for right and left eyes correlated strongly ($R = 0.97$; $P < 0.0001$) over a wide range of levels, as shown in Fig. 10, for the peak MPOD levels of the distribution in each case. The behavior is similar to the respective correlation in adult eyes.13,15,23 While at any particular age, the peak MPOD levels can differ by about one order of magnitude between subjects, there is on average a steady linear increase of MPOD levels over the first 6 years of life, as shown in Fig. 11. We did not find any significant correlation between the MP-containing macula, measured in square pixels, versus the age of the infant, as seen from Fig. 12, i.e., the width of the MP distribution appears constant on average over this age range. A more comprehensive description of the clinical MP results along with their correlations to skin carotenoid levels, which can be measured via RRS techniques, such as our previously described autofluorescence-imaging (AFI) method.15 This method makes use of the spectral properties of lipofuscin, which is located posteriorly to the macular pigment, and features a strong fluorescence over a broadband in the blue/green spectral region, essentially in the same region as the macular pigment, and features a strong fluorescence over a broadband peaking at 620 nm.30 MPOD levels and their spatial distributions are obtained in this method by exciting the lipofuscin chromophore in its blue absorption region and comparing the unattenuated lipofuscin fluorescence intensity in the peripheral region with the MP attenuated fluorescence intensity in the macular region.

Comparing reflection and AFI images obtained for the same adult, the AFI image features a much higher image quality relative to the reflection image, as shown in Fig. 13. This effect is consistent with the different light–tissue interaction scenarios in the reflection versus the AFI method. Compared with the double-path blue-light propagation through all anterior ocular media and deeper retinal layers is strongly confounded by scattering and absorption. This observation is consistent with macular pigment reflection imaging of adults with a standard fundus camera, where images with acceptable quality could only be obtained in relatively young subjects ranging in age from 18 to 24.29

It is interesting here to compare the merits of reflection-based imaging of MP in adults with other applicable imaging methodologies, such as our previously described autofluorescence-imaging (AFI) method.15 This method makes use of the spectral properties of lipofuscin, which is located posteriorly to the macular pigment in the RPE.30 Lipofuscin absorbs over a broadband in the blue/green spectral region, essentially in the same region as the macular pigment, and features a strong fluorescence over a broadband peaking at 620 nm.30 MPOD levels and their spatial distributions are obtained in this method by exciting the lipofuscin chromophore in its blue absorption region and comparing the unattenuated lipofuscin fluorescence intensity in the peripheral region with the MP attenuated fluorescence intensity in the macular region.

Comparing reflection and AFI images obtained for the same adult, the AFI image features a much higher image quality relative to the reflection image, as shown in Fig. 13. This effect is consistent with the different light–tissue interaction scenarios in the reflection versus the AFI method. Compared with the double-path blue-light propagation through all ocular layers in the reflection method, which includes the strongly absorbing CHR, the AFI method uses blue-light propagation only in a single path and only up to the retinal pigment epithelial layer, thus avoiding
the CHR. Furthermore, the fluorescence signals of the return path occur at far-red/near-infrared wavelengths, which are much less severely affected by scattering relative to blue-light wavelengths.

4 Conclusion
We developed a novel capability of the RetCam® for the rapid quantitative measurement of MP levels and their spatial distributions in infants and children and surveyed for the first time the main MP features in the developing human retina. Comparing white-light with blue-light imaging, the latter clearly results in improved MPOD imaging with higher image contrast in the macular region, as evidenced by significantly higher MPOD levels. It is conceivable that white-light excitation generates stronger scattering contributions from shorter excitation wavelengths into peripheral areas relative to narrow-band blue-light excitation or that the convolution of the white-light excitation spectrum with the spectral sensitivity of the blue camera channel peaks at a longer wavelength relative to the blue excitation, thus degrading the image contrast between macula and periphery or decreasing the MP absorption, respectively, and making it less accurate.

Measuring MPOD levels in a clinical population of 51 infants ranging in age between preterm to age 6, we find that MP levels are absent in all measured preterm infants and that they increased on average with age in all other subjects. In the oldest measured infants of about age 6, the peak MP-approached MP levels seen in adults. In contrast to adults, where widely varying spatial distribution patterns are seen, infants with measureable MP had almost exclusively only circularly symmetric MP distributions. The described methodology holds promise for future investigations into the role of MP in the development of the human retina and in combination with intravenous feeding and/or dietary interventions could potentially help improve the health of infants lacking normal carotenoid levels.

Acknowledgments
The authors acknowledge the clinical measurements of Robert A. Hoffman and financial support from the National Eye Institute Grant EY-11600, Research to Prevent Blindness, Abbott Nutrition (Columbus, Ohio), and Image Technologies (Salt Lake City, Utah).

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