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# The Repeatability of *In Vivo* Parafoveal Cone Density and Spacing Measurements

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## **Abstract**

**Purpose:** To assess the repeatability and measurement error associated with cone density and nearest neighbor distance (NND) estimates in images of the parafoveal cone mosaic obtained with an adaptive optics scanning light ophthalmoscope (AOSLO).

**Methods:** Twenty-one participants with no known ocular pathology were recruited. Four retinal locations, approximately  $0.65^\circ$  eccentricity from the center of fixation were imaged 10 times in randomized order with an AOSLO. Cone coordinates in each image were identified using an automated algorithm (with or without manual correction), from which cone density and NND were calculated. Owing to naturally occurring fixational instability, the 10 images recorded from a given location did not overlap entirely. We thus analyzed each image set both before and after alignment.

**Results:** Automated estimates of cone density on the unaligned image sets showed a coefficient of repeatability of 11,769 cones/mm<sup>2</sup> (17.1%). The primary reason for this variability appears to be fixational instability, as aligning the 10 images to include the exact same retinal area, results in an improved repeatability of 4,358 cones/mm<sup>2</sup> (6.4%) using completely automated cone identification software. Repeatability improved further by manually identifying cones missed by the automated algorithm, with a coefficient of repeatability of 1,967 cones/mm<sup>2</sup> (2.7%). NND showed improved repeatability, and was generally insensitive to the undersampling by the automated algorithm.

**Conclusions:** As our data were collected in a young, healthy population, this likely represents a best-case estimate for corresponding measurements in patients with retinal disease. Similar studies need to be carried out on other imaging systems (including those using different imaging modalities,

wavefront correction technology, and/or cone identification software), as repeatability would be expected to be highly sensitive to initial image quality and the performance of cone identification algorithms. Separate studies addressing inter-session repeatability and inter-observer reliability are also needed.

**Keywords:** retina, cones, adaptive optics, repeatability, photoreceptors

The use of ophthalmoscopes equipped with adaptive optics (AO) enables direct visualization of individual cone and rod photoreceptors in the living human retina.<sup>1, 2</sup> The higher transverse resolution provided by AO makes it possible to examine features of the photoreceptor mosaic such as the spatial arrangement of the different spectral types of cone within the mosaic,<sup>3, 4</sup> temporal reflectance changes of individual cones and rods,<sup>5-9</sup> and even the orientation tuning of individual cones.<sup>10</sup> However the most exciting applications of this imaging technology are perhaps the clinical ones, as AO imaging tools offer the promise of a more sensitive means with which to characterize and track retinal degeneration than is currently possible with conventional clinical tools. This capability is especially pertinent to those conditions for which treatments are available or will soon become available.

Central to the realization of the clinical potential of AO imaging is the development of robust techniques with which to analyze such high-resolution images. The ability to use retinal images to make a determination about whether the photoreceptor mosaic of a particular individual has changed over time, or whether it differs from normal depends, among other things, on the reliability and repeatability of the metric being used. Metrics currently used include cell density,<sup>11</sup> mosaic geometry,<sup>12, 13</sup> and cell spacing,<sup>14, 15</sup> though there remains inconsistency in how these are derived. While numerous studies have examined photoreceptor density and spacing in the normal<sup>16-18</sup> and diseased<sup>14, 19-23</sup> retina, there have been only a few reports examining the repeatability of such measurements, outlined below.

A recent study by Talcott *et al.* performed a repeated-measures analysis of cone spacing in three normal eyes and found no significant change in cone spacing over time periods ranging from 16 to 53 months.<sup>24</sup> They provide an estimate of error in cone density

measurements of 6.3%, which takes into account cone selection/misidentification, spectacle magnification errors, distortion in cone images from eye motion, and the selection of the region of interest for analysis. In a single patient with a red-green color vision defect, Rha *et al.* observed a 3.9% change in cone density over a period of six years.<sup>25</sup> Boretzky *et al.* reported a standard deviation of less than 1,000 cones/mm<sup>2</sup> for repeated measures of the same retinal location, though the identification of cone cells was reported to be highly dependent on the confocal pinhole diameter (which would affect the contrast of individual cells) and no additional repeatability statistics were reported.<sup>26</sup> Song *et al.* imaged a single retinal location in one subject at two time points separated by six months and observed cone density estimates from the two sessions within 2%.<sup>18</sup> Despite these isolated reports, there remains a pressing need to rigorously define repeatability statistics for cone density measurements in a larger population, in order to facilitate their application to larger clinical studies. In other words, it is difficult to determine whether a significant change has occurred without an estimate of the repeatability of any one measurement. As such, the purpose of the present study was to assess the intrasession repeatability of *in vivo* cone density measurements based on automated and semi-automated cone identification, and to quantify the measurement error. In addition, we investigated the intrasession repeatability of a metric of cone spacing, mean nearest neighbor distance (NND), also using automated and semi-automated cone identification. For both metrics, we also assessed the effect of the size of the retinal area sampled, as different sampling strategies are often used by different investigators. These results provide a valuable starting point in the discussion of repeatability, and similar systematic approaches will be required for different systems and cone identification software.

## Methods

### *Subjects*

All research followed the tenets of the Declaration of Helsinki and study protocols were approved by the Institutional Review Boards at the Medical College of Wisconsin and Marquette University. Subjects provided informed consent after the nature and possible consequences

of the study were explained. Axial length measurements were obtained on all of the subjects using an IOL Master (Carl Zeiss Meditec, Dublin, CA) to calculate the scale of the retinal images. Twenty-one subjects (13 males and 8 females, aged  $25.9 \pm 6.5$  years) were recruited for the study (Table 1). No subjects had any vision limiting pathology, though one subject (JC\_0002) was found to have an inherited color vision deficiency (deuteranopia). While some individuals with color vision defects have been shown to have disrupted cone mosaics,<sup>20, 27</sup> this subject was previously shown to have a contiguous cone mosaic of normal density and did not harbor any genetic mutation known to affect cone structure in red-green color vision defects, and was thus included in the present study.

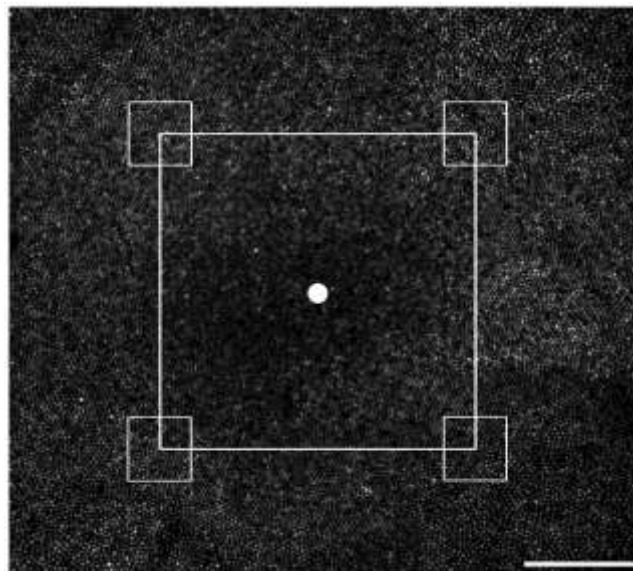
**TABLE 1.**  
Subject Demographics

Subject	Age (yr)	Gender	Axial length (mm)
JC_0002	28	M	24.72
JC_0007	36	M	27.43
JC_0138	27	F	22.67
JC_0200	24	M	24.72
JC_0364	21	M	23.41
JC_0395	23	M	23.75
JC_0571	26	M	24.08
JC_0616	24	M	24.35
JC_0617	27	M	23.77
JC_0645	21	M	23.76
JC_0654	25	F	23.57
JC_0655	23	F	22.40
JC_0656	23	F	25.95
JC_0659	21	F	24.08
JC_0660	21	M	24.31
JC_0667	22	F	23.78
JC_0668	22	F	24.31
JC_0669	23	M	23.08
JC_0677	22	F	24.03
JC_0692	40	M	24.54
JC_0820	45	M	24.27

### *Imaging the Photoreceptor Mosaic*

Each subject's head was stabilized using a chin and forehead rest similar to those found on standard clinical imaging instruments. There was no pupil dilation or control of accommodation using eye drops. A previously described AOSLO was used to image the parafoveal cone mosaic of the right eye.<sup>28, 29</sup> The wavelength of the

super luminescent diode used for retinal imaging was 775nm, subtending a field of view of  $0.96^\circ \times 0.96^\circ$ . The system's pupil used for imaging was 7.75mm, however the eye's pupil was undilated and certainly less than this. We thus calculated that the  $30\mu\text{m}$  confocal pinhole of our system was one Airy disk diameter or less. Separate image sequences of 150 frames each were acquired at four parafoveal locations, each approximately  $0.65^\circ$  from the center of fixation (Figure 1). The four parafoveal locations were imaged in a random order, with the subject staying positioned on the chin/forehead rest for each set of four image sequences. Randomization of the imaging order had two potential benefits. First, the image quality may be best at the first location imaged when the tear film might be more evenly distributed across the cornea (though subjects were instructed to blink normally during each imaging set). Second, the randomized order would mitigate any effect in decreased fixational stability over the course of the imaging session, which might result from fatigue. This procedure was repeated 10 times for each subject with a short break after each set of four locations. The image acquisition software had an "active blink removal" algorithm, which discarded frames that had a mean intensity below a specified threshold. This process improved the percentage of frames in the recorded image sequence (always 150 frames) that contained useable retinal image data.



**Figure 1** Parafoveal imaging locations used in this study. Shown is a foveal montage from subject JC\_0645. Montages were not created for each subject, this one is presented simply to assist with understanding the relationship between the size of



the scanning raster and that of the sampled areas for density analysis as well as the relationship between the foveal center and the location of the parafoveal sampling locations. The large white box represents the extent of the AOSLO scanning raster ( $0.96^\circ \times 0.96^\circ$ ), with the approximate location of the foveal center (fixation) marked with a white circle at the center of the box. The subject was asked to fixate at each of the four corners of the scanning square, and the central portion of each of these images was cropped for density analysis, indicated by the smaller white squares. In this illustration, the small white squares are  $55\mu\text{m} \times 55\mu\text{m}$  in size. Scale bar is  $100\mu\text{m}$ .

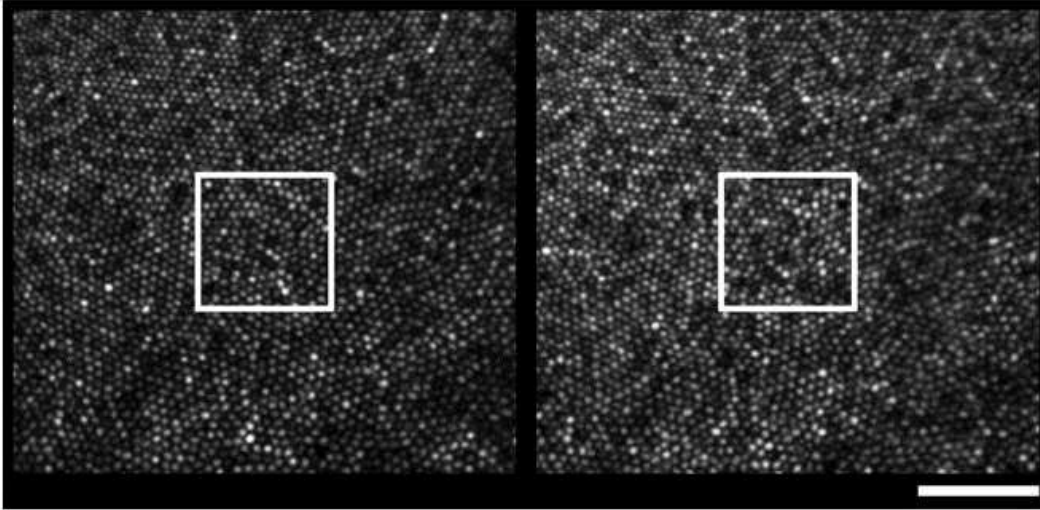
To correct for intraframe distortions within the frames of the raw image sequence due to the sinusoidal motion of the resonant optical scanner, we estimated the distortion from stable images of a Ronchi ruling, and then re-sampled each frame of the raw image sequence over a grid of equally spaced pixels. After desinusoiding, a reference frame was manually selected from within each image sequence, for subsequent registration using custom software. Registration of frames within a given image sequence was performed using a "strip" registration method, in which the frames were registered by dividing the frame of interest into strips, aligning each strip to the location in the reference frame that maximizes the normalized cross correlation between them.<sup>30</sup> Once all the frames were registered, the 40 frames with the highest normalized cross correlation to the reference frame were averaged, in order to generate a final registered image with an increased signal to noise ratio for subsequent analysis.

### *Analyzing the Cone Mosaic*

A total of 840 registered images (21 subjects, four locations each, 10 images at each location) were analyzed. The same retinal area ( $55\mu\text{m} \times 55\mu\text{m}$ ) within the central portion of each image was cropped and used for subsequent analysis of cone density at each location (Figure 1). The cropped images were analyzed three different ways. First, a completely automated algorithm implemented in Matlab (Mathworks, Natick, MA) was used to identify the cones in each cropped image. This is a modified version of the previously described algorithm of Li & Roorda (2007).<sup>12</sup> This algorithm first applies a finite-impulse-response low-pass filter to the retinal image. The original version of the algorithm required manual setting of cutoff frequency of this filter, which dramatically affects the performance of the algorithm. In our study, the filter applied to the image was objectively and

automatically determined based on the image itself (by first automatically estimating the modal cone frequency in the image being analyzed). Local maxima were then identified in the filtered image, and complete details of the method for applying the filter and identifying local maxima have been previously published,<sup>12</sup> which were applied similarly here. The number of cones in each cropped image was simply divided by the retinal area ( $0.003025\text{mm}^2$ ) to derive an estimate of cone density for a given cropped image. The  $(x,y)$  coordinates of the cones were stored in a text array and the Delauney triangulation of the coordinates was obtained. From this triangulation, the built in `dsearch` function in Matlab was used to find the distance of the closest cone in the array for each of the cones (NND). This is identical to the newer function, `nearest Neighbor`.

We then repeated the analysis, except in the second analysis, the 10 averaged images from a given location were first aligned to one another (using the same strip registration as described above) before cropping the central portion (see Figure 2 and Supplementary Digital Content 1 available at **[LWW insert link]**). This ensures that cone density and NND estimates were derived from exactly the same retinal area. The third analysis incorporated manual identification of cones missed by the automated algorithm, using the same aligned image sets utilized in the second analysis. All manual additions for the 840 aligned and cropped images were performed by the same observer (author JC). The identity of the images was not known to the observer and were presented in random order. During the manual addition step, the brightness and contrast of the image was adjusted by the observer to assist in determining whether a cone was present or not. While the opportunity to remove cones was also available to the observer, no such removals were necessary in our image set.



**Figure 2** Effect of fixation instability on the retinal area sampled across the 10 images for a given retinal location. Shown are unaligned (left) and aligned (right) image sequences of the 10 images acquired using the temporal-inferior fixation location for JC\_0616. The white box depicts a  $55\mu\text{m} \times 55\mu\text{m}$  sampling window, demonstrating how different photoreceptors are sampled in each of the 10 images in the unaligned condition, while in the aligned image sequence, the exact same photoreceptors are analyzed in each of the 10 images. See Supplemental Digital Content 1 (available at [LWW insert link]) for the full video sequences. Scale bar is  $50\mu\text{m}$ .

These three analyses were then applied to two additional cropped image sets utilizing smaller sampling windows. As we were interested in the effect of the sampling window size, we simply selectively truncated the  $(x,y)$  cone coordinate list to leave just those cones falling within  $40\mu\text{m}$  or  $25\mu\text{m}$  of the center. This resulted in  $40\mu\text{m} \times 40\mu\text{m}$  and  $25\mu\text{m} \times 25\mu\text{m}$  cropped image sets, respectively.

### *Calculating Measurement Error*

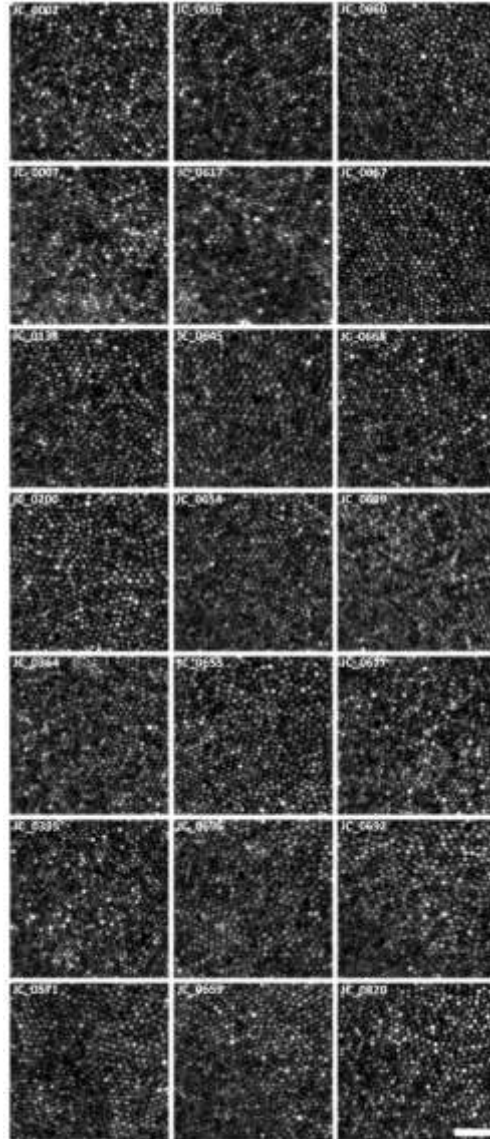
The repeatability for each of the analysis conditions described above was calculated based on the within-subject standard deviation ( $S_w$ ) as outlined by Bland & Altman (1996).<sup>31</sup> To estimate  $S_w$ , we first calculated the standard deviation of the repeated measures for each subject, and then squared this to get variance for each subject. The square root of the average variance for the 21 subjects gives  $S_w$ , and repeatability is defined as  $S_w$  times 2.77.<sup>31</sup> The 95% confidence interval (CI) for repeatability is  $1.96 (S_w/\sqrt{2n(m-1)})$ , where  $n$  is the number of subjects and  $m$  is the number of observations for each

subject. Repeatability is reported both in terms of the measurement unit as well as a percentage of the mean. The measurement error is defined as  $S_w$  times 1.96, and the difference between a subject's measurement and the true value would be expected to be less than the measurement error for 95% of observations.

## Results

### *Repeatability of Cone Density & NND Measurements Based on Automated Cone Identification*

Figure 3 shows representative images of the parafoveal cone mosaic ( $\sim 0.65^\circ$  eccentricity) for all 21 subjects, acquired at the temporal-superior fixation location. As can be seen in the figure, contiguous images of the cone mosaic were obtained in all subjects. In assessing the repeatability of cone density measurements using the completely automated algorithm, we find an average repeatability of 11,769 cones/mm<sup>2</sup>, or 17.1%. This means that the difference between two measurements for the same subject would be less than this value for 95% of pairs of observations. The measurement error in this case was 8,328 cones/mm<sup>2</sup>, which represents the expected difference between a single measurement and the true value for 95% of observations. Compared to cone density, NND showed enhanced repeatability of 0.29  $\mu\text{m}$  (8.4%), with a measurement error of 0.20  $\mu\text{m}$ . A summary of the repeatability statistics is provided in Table 2 and Table 3. In examining the left panel of Figure 2, we see that despite instructing the subject to fixate at a given location 10 times, a slightly different patch of cones was imaged each time. Thus, the relatively poor repeatability here is due to the fact that fixation is unstable even in "normal" subjects and the density/spacing of the underlying mosaic is changing rapidly near the fovea. As a result, even small deviations in fixation would result in differences in cone density or NND between successive images.



**Figure 3** Cone photoreceptor images for all 21 subjects, acquired using the temporal-superior fixation location. So as not to bias the reader, the representative image for each subject was chosen randomly from the 10 images from this location. Scale bar is 25 $\mu$ m.

**TABLE 2.**  
Intraseession repeatability of parafoveal cone density measurements (55  $\mu\text{m} \times 55 \mu\text{m}$  sampling window)

Fixation location	Mean density (cones/mm <sup>2</sup> )	Measurement error (cones/mm <sup>2</sup> )	Repeatability (cones/mm <sup>2</sup> )	95% CI for repeatability (cones/mm <sup>2</sup> )	Repeatability (%)	95% CI for repeatability (%)
Unaligned, automated cone identification						
Bottom left	66,786	7,867	11,119	10,714–11,524	16.6	16.0–17.3
Bottom right	65,466	8,364	11,820	11,390–12,250	18.1	17.4–18.7
Top left	72,967	8,751	12,368	11,918–12,818	17.0	16.3–17.6
Top right	69,987	8,329	11,771	11,343–12,199	16.8	16.2–17.4
Average	68,802	8,328	11,769	11,341–12,198	17.1	16.5–17.8
Aligned, automated cone identification						
Bottom left	67,207	2,557	3,614	3,482–3,746	5.4	5.2–5.6
Bottom right	66,812	3,013	4,258	4,103–4,413	6.4	6.1–6.6
Top left	70,928	3,277	4,631	4,462–4,800	6.5	6.3–6.8
Top right	69,192	3,487	4,928	4,749–5,107	7.1	6.9–7.4
Average	68,535	3,084	4,358	4,199–4,517	6.4	6.1–6.6
Aligned, automated cone identification with manual addition						
Bottom left	70,575	1,305	1,844	1,777–1,911	2.6	2.5–2.7
Bottom right	70,204	1,387	1,960	1,889–2,031	2.8	2.7–2.9
Top left	75,418	1,373	1,940	1,869–2,011	2.6	2.5–2.7
Top right	73,914	1,503	2,124	2,047–2,201	2.9	2.8–3.0
Average	72,528	1,392	1,967	1,896–2,039	2.7	2.6–2.8

**TABLE 3.**  
Intraseession repeatability of parafoveal NND measurements (55  $\mu\text{m} \times 55 \mu\text{m}$  sampling window)

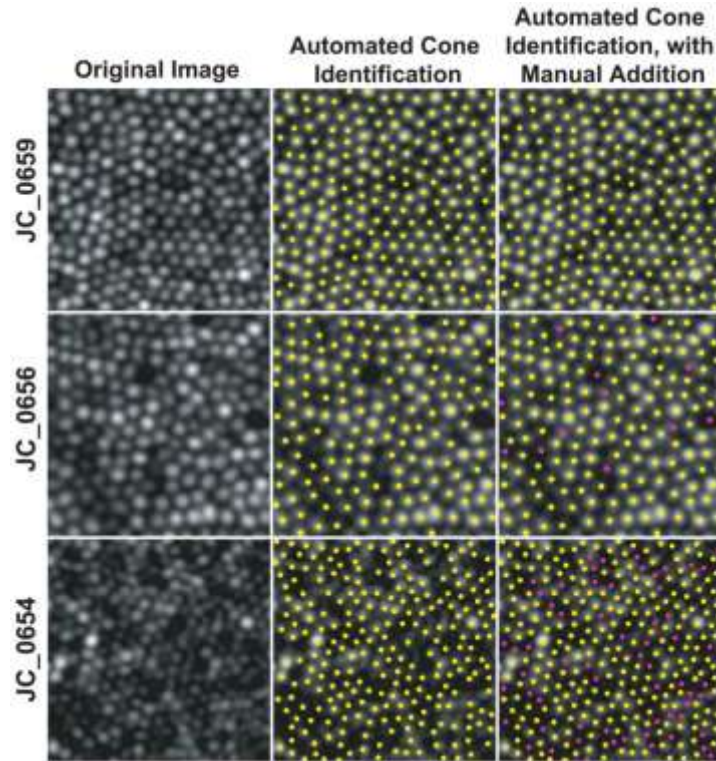
Fixation location	Mean spacing ( $\mu\text{m}$ )	Measurement error ( $\mu\text{m}$ )	Repeatability ( $\mu\text{m}$ )	95% CI for repeatability ( $\mu\text{m}$ )	Repeatability (%)	95% CI for repeatability (%)
Unaligned, automated cone identification						
Bottom left	3.52	0.177	0.251	0.242–0.260	7.1	6.8–7.4
Bottom right	3.54	0.211	0.298	0.287–0.309	8.4	8.1–8.7
Top left	3.34	0.212	0.300	0.289–0.311	9.0	8.7–9.3
Top right	3.40	0.214	0.303	0.292–0.314	8.9	8.6–9.2
Average	3.45	0.204	0.288	0.278–0.299	8.4	8.1–8.7
Aligned, automated cone identification						
Bottom left	3.50	0.049	0.069	0.066–0.072	2.0	1.9–2.1
Bottom right	3.50	0.057	0.081	0.078–0.084	2.3	2.2–2.4
Top left	3.37	0.058	0.082	0.079–0.085	2.4	2.3–2.5
Top right	3.39	0.057	0.081	0.078–0.084	2.4	2.2–2.5
Average	3.44	0.055	0.078	0.075–0.081	2.3	2.2–2.4
Aligned, automated cone identification with manual addition						
Bottom left	3.45	0.060	0.085	0.082–0.088	2.5	2.4–2.6
Bottom right	3.44	0.068	0.095	0.092–0.098	2.8	2.7–2.9
Top left	3.31	0.072	0.102	0.098–0.106	3.1	3.0–3.2
Top right	3.32	0.056	0.080	0.077–0.083	2.4	2.3–2.5
Average	3.38	0.064	0.090	0.087–0.094	2.7	2.6–2.8

To account for fixational instability, the 10 images from a given fixation location were first aligned to each another before cropping out the central 55 $\mu\text{m} \times 55\mu\text{m}$  for analysis. As shown in the right panel of Figure 2, this results in a situation where exactly the same cones are included in the analysis. As summarized in Table 2, this results in an improved average repeatability of 4,358 cones/mm<sup>2</sup>, or 6.4% for the aligned images. In this case, the measurement error was 3,084

cones/mm<sup>2</sup>, which again represents the expected difference between a single measurement and the true value for 95% of observations. For the 55µm × 55µm cropped images an average of 207 cones were identified by the automated algorithm, so our repeatability indicates that the number of cones missed between two measurements for the same subject would be fewer than 13 for 95% of pairs of observations. The average repeatability for the NND measurements improved to 0.078µm (2.3%), with a measurement error of 0.055µm (Table 3).

### *Effect of Manual Addition of Cones on the Repeatability of Cone Density & NND Measurements*

The third analysis allowed the manual addition of cones that were missed by the automated algorithm. Despite good image contrast and resolution, the performance of the automated cone identification algorithm was highly variable, and this can be seen in Figure 4. An average of 12 cones were manually added across the 840 images analyzed (range=0–62 cones added), resulting in an average of 219 total cones in the 55µm × 55µm cropped images. The top row of Figure 4 shows an example of an image where the user added no cones. In other words, by the judgment of the user, no cones were missed by the automated algorithm. The middle row of Figure 4 shows an example of an image where the user identified 12 cones missed by the automated algorithm, and the bottom row shows an example of an image where the user identified 62 cones missed by the automated algorithm. The manual addition step further improves the repeatability of cone density measurements, with an average repeatability of 1,967 cones/mm<sup>2</sup>, or 2.7% (Table 2). For our data, this is equivalent to about 6 cones, indicating that the number of cones missed between two measurements for the same subject would be fewer than 6 for 95% of pairs of observations. The associated measurement error improves to 1,392 cones/mm<sup>2</sup> and the average standard deviation for the 10 repeated measures across the 21 subjects was 710 cones/mm<sup>2</sup>.



**Figure 4** Variable performance of the automated cone identification algorithm. Shown are images from three subjects, JC\_0659, JC\_0656, and JC\_0654. These images illustrate the variable performance of the automated algorithm across all 840 images analyzed in the aligned case. In the image from JC\_0659 the algorithm missed no cones, while in the image from JC\_0654, the user added 62 cones. The average number of cones added manually across all images was 12 (5.5%), which is the number missed by the automated algorithm in the image from JC\_0656. Yellow circles represent cones identified by the automatic algorithm; pink cones indicate those added by the user during the manual addition step. All images are  $55\ \mu\text{m} \times 55\ \mu\text{m}$  in size.

In contrast to cone density, the NND measurements showed no improvement over those obtained using the completely automated algorithm, highlighting the insensitivity of this metric to small amounts of undersampling. The average repeatability for the NND measurements was  $0.090\ \mu\text{m}$  (2.7%), with a measurement error of  $0.064\ \mu\text{m}$ . (Table 3)

### *Effect of Sampling Window Size*

We repeated all of the above analyses on our image sets using two smaller sampling windows,  $40\ \mu\text{m} \times 40\ \mu\text{m}$  and  $25\ \mu\text{m} \times 25\ \mu\text{m}$ .



These were chosen based on those reported previously by other groups.<sup>16, 18</sup> Interestingly, as the sampling window size decreased, we observed a decrease in the repeatability and an increase in the measurement error for both cone density and NND, though there was some variability in the effect. Complete statistical summaries for cone density for the 40 $\mu\text{m}$   $\times$  40 $\mu\text{m}$  sampling window are given in Table 4, while those for the 25 $\mu\text{m}$   $\times$  25 $\mu\text{m}$  sampling window are given in Table 5. Table 6 and Table 7 provide similar summaries of the NND measurements. These data illustrate the importance of specifying the size of the sampling window used to derive density estimates in order to facilitate comparison of different studies.

**TABLE 4.**  
Intraseession repeatability of parafoveal cone density measurements (40  $\mu\text{m}$   $\times$  40  $\mu\text{m}$  sampling window)

Fixation location	Mean density (cones/mm <sup>2</sup> )	Measurement error (cones/mm <sup>2</sup> )	Repeatability (cones/mm <sup>2</sup> )	95% CI for repeatability (cones/mm <sup>2</sup> )	Repeatability (%)	95% CI for repeatability (%)
Unaligned, automated cone identification						
Bottom left	67,131	8,680	12,267	11,821–12,713	18.3	17.6–18.9
Bottom right	65,804	9,125	12,896	12,427–13,365	19.6	18.9–20.3
Top left	73,565	9,508	13,437	12,948–13,926	18.3	17.6–18.9
Top right	70,405	9,219	13,029	12,555–13,503	18.5	17.8–19.2
Average	69,226	9,133	12,907	12,438–13,377	18.7	18.0–19.3
Aligned, automated cone identification						
Bottom left	67,518	2,937	4,151	4,000–4,302	6.1	5.9–6.4
Bottom right	67,071	3,525	4,982	4,801–5,163	7.4	7.2–7.7
Top left	71,414	3,763	5,318	5,124–5,512	7.4	7.2–7.7
Top right	69,949	3,664	5,178	4,990–5,366	7.4	7.1–7.7
Average	68,988	3,472	4,907	4,729–5,086	7.1	6.9–7.4
Aligned, automated cone identification with manual addition						
Bottom left	70,741	1,826	2,581	2,487–2,675	3.6	3.5–3.8
Bottom right	70,348	2,184	3,087	2,975–3,199	4.4	4.2–4.5
Top left	75,595	2,043	2,887	2,782–2,992	3.8	3.7–4.0
Top right	74,637	2,279	3,221	3,104–3,338	4.3	4.2–4.5
Average	72,830	2,083	2,944	2,837–3,051	4.0	3.9–4.2

**TABLE 5.**  
Intrasection repeatability of parafoveal cone density measurements (25  $\mu\text{m} \times 25 \mu\text{m}$  sampling window)

Fixation location	Mean density (cones/mm <sup>2</sup> )	Measurement error (cones/mm <sup>2</sup> )	Repeatability (cones/mm <sup>2</sup> )	95% CI for repeatability (cones/mm <sup>2</sup> )	Repeatability (%)	95% CI for repeatability (%)
Unaligned, automated cone identification						
Bottom left	67,451	10,387	14,679	14,145–15,213	21.8	21.0–22.6
Bottom right	65,813	10,395	14,690	14,155–15,225	22.3	21.5–23.1
Top left	73,615	11,164	15,777	15,203–16,351	21.4	20.7–22.2
Top right	70,331	10,205	14,423	13,896–14,948	20.5	19.8–21.3
Average	69,303	10,538	14,892	14,350–15,434	21.5	20.8–22.3
Aligned, automated cone identification						
Bottom left	67,368	4,348	6,145	5,921–6,369	9.1	8.8–9.5
Bottom right	66,583	5,005	7,073	6,816–7,330	10.6	10.2–11.0
Top left	71,718	5,842	8,256	7,956–8,556	11.5	11.1–11.9
Top right	68,610	5,110	7,222	6,959–7,485	10.5	10.1–10.9
Average	68,570	5,076	7,174	6,913–7,435	10.4	10.1–10.8
Aligned, automated cone identification with manual addition						
Bottom left	70,304	3,007	4,249	4,094–4,404	6.0	5.8–6.3
Bottom right	70,019	3,348	4,732	4,560–4,904	6.8	6.5–7.0
Top left	75,674	3,519	4,973	4,792–5,154	6.6	6.3–6.8
Top right	72,701	3,213	4,451	4,376–4,706	6.2	6.0–6.5
Average	72,175	3,272	4,624	4,456–4,792	6.0	6.2–6.7

**TABLE 6.**  
Intrasection repeatability of parafoveal NND measurements (40  $\mu\text{m} \times 40 \mu\text{m}$  sampling window)

Fixation location	Mean spacing ( $\mu\text{m}$ )	Measurement error ( $\mu\text{m}$ )	Repeatability ( $\mu\text{m}$ )	95% CI for repeatability ( $\mu\text{m}$ )	Repeatability (%)	95% CI for repeatability (%)
Unaligned, automated cone identification						
Bottom left	3.54	0.206	0.291	0.280–0.302	8.2	7.9–8.5
Bottom right	3.56	0.224	0.317	0.305–0.329	8.9	8.6–9.3
Top left	3.35	0.230	0.325	0.313–0.337	9.7	9.3–10.1
Top right	3.42	0.224	0.317	0.305–0.329	9.3	8.9–9.6
Average	3.47	0.221	0.313	0.301–0.324	9.0	8.7–9.4
Aligned, automated cone identification						
Bottom left	3.53	0.066	0.094	0.091–0.097	2.7	2.6–2.8
Bottom right	3.51	0.079	0.112	0.108–0.116	3.2	3.1–3.3
Top left	3.39	0.068	0.097	0.093–0.101	2.9	2.7–3.0
Top right	3.41	0.074	0.104	0.100–0.108	3.1	2.9–3.2
Average	3.46	0.072	0.102	0.098–0.106	3.0	2.8–3.1
Aligned, automated cone identification with manual addition						
Bottom left	3.47	0.076	0.107	0.103–0.111	3.1	3.0–3.2
Bottom right	3.45	0.085	0.120	0.116–0.124	3.5	3.4–3.6
Top left	3.33	0.076	0.107	0.103–0.111	3.2	3.1–3.3
Top right	3.33	0.075	0.105	0.101–0.109	3.2	3.0–3.3
Average	3.39	0.078	0.110	0.106–0.114	3.3	3.1–3.4

**TABLE 7.**  
Intrasession repeatability of parafoveal NND measurements (25  $\mu\text{m}$   $\times$  25  $\mu\text{m}$  sampling window)

Fixation location	Mean spacing ( $\mu\text{m}$ )	Measurement error ( $\mu\text{m}$ )	Repeatability ( $\mu\text{m}$ )	95% CI for repeatability ( $\mu\text{m}$ )	Repeatability (%)	95% CI for repeatability (%)
Unaligned, automated cone identification						
Bottom left	3.56	0.247	0.349	0.336–0.362	9.8	9.4–10.2
Bottom right	3.58	0.254	0.359	0.346–0.372	10.0	9.7–10.4
Top left	3.38	0.278	0.393	0.379–0.407	11.6	11.2–12.1
Top right	3.45	0.258	0.364	0.351–0.377	10.6	10.2–10.9
Average	3.49	0.259	0.366	0.353–0.380	10.5	10.1–10.9
Aligned, automated cone identification						
Bottom left	3.55	0.115	0.162	0.156–0.168	4.6	4.4–4.7
Bottom right	3.55	0.130	0.184	0.177–0.191	5.2	5.0–5.4
Top left	3.42	0.104	0.147	0.142–0.152	4.3	4.1–4.4
Top right	3.44	0.115	0.162	0.159–0.168	4.7	4.5–4.9
Average	3.49	0.116	0.164	0.158–0.170	4.7	4.5–4.9
Aligned, automated cone identification with manual addition						
Bottom left	3.50	0.120	0.170	0.164–0.176	4.9	4.7–5.0
Bottom right	3.49	0.132	0.186	0.179–0.193	5.3	5.1–5.5
Top left	3.36	0.116	0.164	0.158–0.170	4.9	4.7–5.1
Top right	3.37	0.118	0.167	0.161–0.173	4.9	4.8–5.1
Average	3.42	0.121	0.172	0.166–0.178	5.0	4.8–5.2

## Cone Density and NND Variability Across Subjects

Accepting that the estimates of cone density and NND obtained using the aligned images with manual addition of cones are more accurate than those based on the completely automated analysis, we can examine the statistics of the normal cone mosaic. Table 8 provides the average cone density and NND for each subject using each of the three sampling window sizes. There was no significant difference in cone density across the three sampling window conditions ( $p=0.21$ , repeated measures ANOVA, GraphPad InStat, v3.1a). The average cone density for each subject ranged from 55,165 cones/mm<sup>2</sup> to 93,604 cones/mm<sup>2</sup>, with a mean ( $\pm$  SD) of the group of 72,528  $\pm$  8,539 cones/mm<sup>2</sup> (using the 55 $\mu\text{m}$   $\times$  55 $\mu\text{m}$  window). This is comparable to previous estimates at this retinal location ( $\sim 0.65^\circ$ ). For example, Li *et al.* reported a range from about 64,000 cones/mm<sup>2</sup> to 98,000 cones/mm<sup>2</sup> at a comparable eccentricity across 18 subjects.<sup>17</sup>

**TABLE 8.**  
Intersubject variability in parafoveal cone density and NND measurements<sup>a</sup>

	Cone density (cones/mm <sup>2</sup> )			NND (μm)		
	55 μm × 55 μm	40 μm × 40 μm	25 μm × 25 μm	55 μm × 55 μm	40 μm × 40 μm	25 μm × 25 μm
JC_0002	66,876	68,500	67,040	3.49	3.52	3.55
JC_0007	60,165	62,625	62,280	3.65	3.65	3.68
JC_0138	64,843	64,359	63,440	3.54	3.58	3.61
JC_0200	65,678	63,906	64,680	3.56	3.57	3.64
JC_0364	76,893	79,125	75,120	3.27	3.30	3.32
JC_0395	82,992	80,344	79,080	3.16	3.18	3.21
JC_0571	67,512	67,797	68,240	3.47	3.46	3.52
JC_0616	70,318	69,186	67,214	3.51	3.56	3.64
JC_0617	79,612	79,359	80,800	3.11	3.12	3.10
JC_0645	69,893	69,875	68,520	3.45	3.47	3.51
JC_0654	93,604	92,937	91,280	3.00	3.01	3.03
JC_0655	76,339	77,094	78,800	3.30	3.32	3.32
JC_0656	55,165	57,547	56,480	3.80	3.80	3.85
JC_0659	72,322	71,938	72,560	3.41	3.41	3.45
JC_0660	81,314	81,469	78,240	3.19	3.20	3.25
JC_0667	73,802	73,922	74,600	3.40	3.42	3.44
JC_0668	74,074	73,859	72,400	3.41	3.42	3.45
JC_0669	75,818	76,969	76,160	3.17	3.18	3.24
JC_0677	78,306	78,141	80,520	3.18	3.22	3.25
JC_0692	64,417	65,747	64,369	3.52	3.55	3.59
JC_0820	73,140	72,734	71,840	3.37	3.38	3.41
Average	72,528	72,830	72,174	3.38	3.40	3.43
SD	8,539	8,094	8,079	0.20	0.20	0.21

<sup>a</sup>All values were acquired using aligned images, with manual addition of cones missed by automated algorithm.

As seen in Table 8, there was a significant difference between the NND values across the three sampling window conditions ( $p < 0.0001$ , repeated measures ANOVA, Bonferroni corrected, GraphPad InStat, v3.1a). This presumably reflects the fact that as the sampling window decreases in size, the relative proportion of cones with undefined neighbors increases. These edge cones will serve to increase, on average, the NND – as there are only two possible scenarios with regard to the NND for that cone. Either the nearest neighbor resides within the sampling window, or it falls outside the sampling window. If it falls inside the sampling window, the NND value recorded for that cone will be equal to the true NND for that cone. If, on the other hand, it falls outside the sampling window, then the NND value for that cone will be based on the closest neighbor within the sampling window, which will always have a greater intercone distance than the true NND for that cone. While this artifact affects the overall accuracy of NND measurements, it wouldn't affect the measured repeatability, as each image within a given condition would be expected to have a similar proportion of cones at the edge of that particular sampling window size.

## Discussion

Using undilated pupils, we obtained images of the contiguous cone mosaic in 21 subjects with an AOSLO at four locations, each approximately  $0.65^\circ$  from the center of fixation. We used automated and/or manual approaches to identify the cones in each image, from which cone density and NND were calculated. These data represent an important first step in assessing the broader clinical utility of such measurements, specifically with regard to determining whether a given mosaic has changed over time or whether a given mosaic differs significantly from another or from a population mean. There are a number of important limitations and caveats to our study that we review here, with the goal of stimulating further work on this issue so as to accelerate the development of robust image analysis tools for *in vivo* images of the photoreceptor mosaic.

First, our images were acquired close to the fovea (within about  $200\ \mu\text{m}$ ). It is known from a number of studies that this is where cone density is changing most rapidly.<sup>17, 18, 32</sup> One would expect that in the periphery, where cone density is more uniform, that the repeatability would be affected less by fixational instability and that there may be less of a difference between the automated approach that does not include aligning the successive images to one another versus the automated approach that first aligns the successive images to one another.

A second issue relates to the fact that we only examined the cone mosaic. As has been shown recently, it is now possible to image the rod mosaic.<sup>2, 29, 33</sup> Unlike the cone mosaic, which appears to reach an asymptotic density beyond about 5mm, rod density changes throughout the retina; first increasing sharply moving away from the fovea and then decreasing beyond the rod-rim.<sup>32</sup> As a result, the same negative effect that small misalignments between images has on the repeatability of parafoveal cone density estimates would exist for estimates of peripheral rod density. Thus, we conclude that obtaining the highest intersession repeatability requires precise alignment of images from each session, or some other means by which one can ensure the images are from the exact same retinal location. Not doing

this severely limits the sensitivity of the corresponding photoreceptor density measurements.

Another important issue to consider relates to the use of cone density and NND as our image metrics. While our NND measurements were less sensitive to undersampling (*i.e.*, missed cones) than our estimates of cone density (Table 3), it has been shown previously that measures of cone spacing based on an exclusion radius are even less sensitive to undersampling.<sup>34, 35</sup> Such insensitivity could be viewed as either an advantage or disadvantage. From the point of view of developing image processing tools to find cones in an image, the utilization of spacing metrics relaxes the constraint that such a tool find each and every cell in the image. However from an image interpretation point of view, finding "normal" cone spacing in an image in no way ensures that the image in its entirety is "normal". Thus, these spacing measures overestimate the global health of the photoreceptor mosaic. For example, a mosaic that has sporadic loss of cones would be flagged as having normal spacing, but abnormal density. To be able to use density, one needs to be sure that they can reliably visualize every cell that remains in the mosaic. Likewise, any analysis of the geometry of the mosaic (*i.e.*, Voronoi) requires that every cell present be visualized. As suggested by Chen *et al.*,<sup>36</sup> cone spacing (and conversely, cone density) should each only be considered one aspect of image analysis. Perhaps more importantly, it will be useful to combine different mosaic metrics (both local and global) to provide a more comprehensive picture of the overall integrity of the mosaic.

In conclusion, we have defined the repeatability of parafoveal cone density measurements for our AOSLO system and accompanying semi-automated cone identification software, as well as the associated measurement error. Repeatability would be expected to differ from system to system based on image quality and individual, thus one should not generalize these results to other research or commercial AO systems, though our data provide a useful starting point for the discussion of reliability and repeatability. Our data also demonstrate the importance of specifying the size of the sampling window, as this can affect the repeatability and/or absolute values of cone density and NND. For multicenter clinical trials, it will be important to demonstrate comparable repeatability across systems, as well as establishing the

inter-session repeatability and inter-observer reliability. Equally important are the development of normative databases against which measurements of the cone mosaic in diseased retinas can be compared. There are growing databases of cone spacing<sup>14, 22, 36</sup> and cone density<sup>16-18</sup> that will need to be expanded to include information about the rod mosaic as well as define the repeatability of the measurements used to construct the databases.

## Supplementary Material

### Supplemental Digital Content 1:

Shown are unaligned (*left*) and aligned (*right*) image sequences of the 10 images acquired using the temporal-inferior fixation location for JC\_0616. The white box depicts a 55 $\mu$ m  $\times$  55 $\mu$ m sampling window, demonstrating how different photoreceptors are sampled in each of the 10 images in the unaligned condition, while in the aligned image sequence, the exact same photoreceptors are analyzed in each of the 10 images. Scale bar is 50 $\mu$ m. (.avi file).

<http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3348369/bin/NIHMS369072-supplement-1.avi> (4.8M, avi)

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## Footnotes

Supplemental digital content is available online:

Video 1: Fixation instability on the retinal area sampled across the 10 images (.avi video) <http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3348369/#SD1>

## References

1. Liang J, Williams DR, Miller DT. Supernormal vision and high-resolution retinal imaging through adaptive optics. *J Opt Soc Am (A)* 1997;14:2884–92.
2. Rossi EA, Chung M, Dubra A, Hunter JJ, Merigan WH, Williams DR. Imaging retinal mosaics in the living eye. *Eye (Lond)* 2011;25:301–8.
3. Roorda A, Williams DR. The arrangement of the three cone classes in the living human eye. *Nature*. 1999;397:520–2.
4. Hofer H, Carroll J, Neitz J, Neitz M, Williams DR. Organization of the human trichromatic cone mosaic. *J Neurosci*. 2005;25:9669–79.
5. Pallikaris A, Williams DR, Hofer H. The reflectance of single cones in the living human eye. *Invest Ophthalmol Vis Sci*. 2003;44:4580–92.
6. Jonnal RS, Rha J, Zhang Y, Cense B, Gao W, Miller DT. In vivo functional imaging of human cone photoreceptors. *Opt Express*. 2007;15:16141–60.
7. Rha J, Schroeder B, Godara P, Carroll J. Variable optical activation of human cone photoreceptors visualized using a short coherence light source. *Opt Lett*. 2009;34:3782–4.
8. Jonnal RS, Besecker JR, Derby JC, Kocaoglu OP, Cense B, Gao W, Wang Q, Miller DT. Imaging outer segment renewal in living human cone photoreceptors. *Opt Express*. 2010;18:5257–70.
9. Cooper RF, Dubis AM, Pavaskar A, Rha J, Dubra A, Carroll J. Spatial and temporal variation of rod photoreceptor reflectance in the human retina. *Biomed Opt Express*. 2011;2:2577–89.
10. Roorda A, Williams DR. Optical fiber properties of individual human cones. *J Vis*. 2002;2:404–12.
11. Chui TY, Song H, Burns SA. Adaptive-optics imaging of human cone photoreceptor distribution. *J Opt Soc Am (A)* 2008;25:3021–9.
12. Li KY, Roorda A. Automated identification of cone photoreceptors in adaptive optics retinal images. *J Opt Soc Am (A)* 2007;24:1358–63.
13. Baraas RC, Carroll J, Gunther KL, Chung M, Williams DR, Foster DH, Neitz M. Adaptive optics retinal imaging reveals S-cone dystrophy in tritan color-vision deficiency. *J Opt Soc Am (A)* 2007;24:1438–47.
14. Duncan JL, Zhang Y, Gandhi J, Nakanishi C, Othman M, Branham KE, Swaroop A, Roorda A. High-resolution imaging with adaptive optics in patients with inherited retinal degeneration. *Invest Ophthalmol Vis Sci*. 2007;48:3283–91.
15. Rossi EA, Roorda A. The relationship between visual resolution and cone spacing in the human fovea. *Nat Neurosci*. 2010;13:156–7.



16. Chui TY, Song H, Burns SA. Individual variations in human cone photoreceptor packing density: variations with refractive error. *Invest Ophthalmol Vis Sci.* 2008;49:4679–87.
17. Li KY, Tiruveedhula P, Roorda A. Intersubject variability of foveal cone photoreceptor density in relation to eye length. *Invest Ophthalmol Vis Sci.* 2010;51:6858–67.
18. Song H, Chui TY, Zhong Z, Elsner AE, Burns SA. Variation of cone photoreceptor packing density with retinal eccentricity and age. *Invest Ophthalmol Vis Sci.* 2011;52:7376–84.
19. Choi SS, Doble N, Hardy JL, Jones SM, Keltner JL, Olivier SS, Werner JS. In vivo imaging of the photoreceptor mosaic in retinal dystrophies and correlations with visual function. *Invest Ophthalmol Vis Sci.* 2006;47:2080–92.
20. Carroll J, Baraas RC, Wagner-Schuman M, Rha J, Siebe CA, Sloan C, Tait DM, Thompson S, Morgan JI, Neitz J, Williams DR, Foster DH, Neitz M. Cone photoreceptor mosaic disruption associated with Cys203Arg mutation in the M-cone opsin. *Proc Natl Acad Sci U S A.* 2009;106:20948–53.
21. Carroll J, Rossi EA, Porter J, Neitz J, Roorda A, Williams DR, Neitz M. Deletion of the X-linked opsin gene array locus control region (LCR) results in disruption of the cone mosaic. *Vision Res.* 2010;50:1989–99.
22. Duncan JL, Talcott KE, Ratnam K, Sundquist SM, Lucero AS, Day S, Zhang Y, Roorda A. Cone structure in retinal degeneration associated with mutations in the peripherin/RDS gene. *Invest Ophthalmol Vis Sci.* 2011;52:1557–66.
23. Genead MA, Fishman GA, Rha J, Dubis AM, Bonci DM, Dubra A, Stone EM, Neitz M, Carroll J. Photoreceptor structure and function in patients with congenital achromatopsia. *Invest Ophthalmol Vis Sci.* 2011;52:7298–308.
24. Talcott KE, Ratnam K, Sundquist SM, Lucero AS, Lujan BJ, Tao W, Porco TC, Roorda A, Duncan JL. Longitudinal study of cone photoreceptors during retinal degeneration and in response to ciliary neurotrophic factor treatment. *Invest Ophthalmol Vis Sci.* 2011;52:2219–26.
25. Rha J, Dubis AM, Wagner-Schuman M, Tait DM, Godara P, Schroeder B, Stepien K, Carroll J. Spectral domain optical coherence tomography and adaptive optics: imaging photoreceptor layer morphology to interpret preclinical phenotypes. *Adv Exp Med Biol.* 2010;664:309–16.
26. Boretsky A, Khan F, van Kuijk E, Motamedi M. Adaptive optics SLO imaging of macular photoreceptors: variations in automated cone density measurements based on confocal pinhole diameter. *Invest Ophthalmol Vis Sci.* 2011;52:E-Abstract 3196.

27. Carroll J, Neitz M, Hofer H, Neitz J, Williams DR. Functional photoreceptor loss revealed with adaptive optics: an alternate cause of color blindness. *Proc Natl Acad Sci U S A*. 2004;101:8461–6.
28. Dubra A, Sulai Y. Reflective afocal broadband adaptive optics scanning ophthalmoscope. *Biomed Opt Express*. 2011;2:1757–68.
29. Dubra A, Sulai Y, Norris JL, Cooper RF, Dubis AM, Williams DR, Carroll J. Noninvasive imaging of the human rod photoreceptor mosaic using a confocal adaptive optics scanning ophthalmoscope. *Biomed Opt Express*. 2011;2:1864–76.
30. Dubra A, Harvey Z. *Biomedical Image Registration*. Heidelberg: Springer Verlag; 2010. Registration of 2D Images from Fast Scanning Ophthalmic Instruments; pp. 60–71.
31. Bland JM, Altman DG. Measurement error proportional to the mean. *BMJ*. 1996;313:106.
32. Curcio CA, Sloan KR, Kalina RE, Hendrickson AE. Human photoreceptor topography. *J Comp Neurol*. 1990;292:497–523.
33. Merino D, Duncan JL, Tiruveedhula P, Roorda A. Observation of cone and rod photoreceptors in normal subjects and patients using a new generation adaptive optics scanning laser ophthalmoscope. *Biomed Opt Express*. 2011;2:2189–201.
34. Rodieck RW. The density recovery profile: a method for the analysis of points in the plane applicable to retinal studies. *Vis Neurosci*. 1991;6:95–111.
35. Cook JE. Spatial properties of retinal mosaics: an empirical evaluation of some existing measures. *Vis Neurosci*. 1996;13:15–30.
36. Chen Y, Ratnam K, Sundquist SM, Lujan B, Ayyagari R, Gudiseva VH, Roorda A, Duncan JL. Cone photoreceptor abnormalities correlate with vision loss in patients with Stargardt disease. *Invest Ophthalmol Vis Sci*. 2011;52:3281–92.