Automated Quantification and Topographical Distribution of the Whole Population of S- and L-Cones in Adult Albino and Pigmented Rats

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PURPOSE. To quantify the whole population of S- and L-cones in the albino (Sprague-Dawley; SD) and pigmented (Piebald Virol Glaxo, PVG) rats and to study their topographical distribution within the retina.

METHODS. Retinal radial sections and whole-mounted retinas were double immunodetected with antibodies against UV-sensitive and L-opsins to detect the S- and L-cones, respectively. Two automated routines were developed to quantify the whole population of S- and L-cones. Detailed isodensity maps of each cone type were generated. In both strains, the presence of dual cones was detected, these were semiautomatically quantified and their distribution determined. The matching distribution of retinal ganglion cells (RGCs) and L-cones was attained by double immunodetection of Brn3a and L-opsin, respectively.

RESULTS. The mean number ± SEM of L- or S-cones in SD and PVG retinas was 231,736 ± 14,517 and 239,939 ± 6,494 or 41,028 ± 5,074, and 27,316 ± 2,235, respectively. There was an increasing gradient of S-cone density along the inferonasal quadrant, although the highest densities were found in the retinal rims. The distribution of L-cones seemed to be complementary to the S-cones. The highest densities were observed in the superior nasotemporal axis, paralleling the distribution of Brn3a-positive RGCs.

CONCLUSIONS. These data establish, for the first time, the total number and the topographical distribution of S- and L-cones in two rat strains and demonstrate the correlation of L-cones and RGC spatial distribution. (Invest Ophthalmol Vis Sci. 2010;51:3171–3183) DOI:10.1167/iovs.09-4861

I n the mammalian retina, cone photoreceptors are responsible for daylight (photopic) vision and color discrimination. Color discrimination in nonprimate mammals is achieved by two types of cones: S and M/L. These cones are distinguished mainly by the portion of light spectrum to which each is more sensitive: short wave lengths stimulate S-cones and medium to long wave lengths stimulate M/L-cones. This spectral sensitivity is conferred by the protein part of the visual pigments, the opsins. Thus, opsins are slightly different, depending on their sensitivity and color preference. Each cone type carries a specific opsin. In rodents, S-cones express the ultraviolet-sensitive or SWS1 opsin,1,2 and M/L-cones express the LWS opsin,3 which detects only green light4 and is known as L-opsin. Therefore, the strict nomenclature of these photoreceptors is L-cones. In some species, there are cones that coexpress both opsins; these are known as dual cones.4–6

The rat retina has been the subject of intensive research for several decades, because it is an easily accessible model for the study of the response of central nervous system neurons to a variety of insults and associated therapies.7–12 It is, as well, a model for the study of experimentally induced photoreceptor degeneration, such as phototoxic insult,13 or that caused by inherited retinal diseases, such as retinitis pigmentosa14 or retinal dystrophy.15,16 Although there is a body of literature reporting the number and distribution of cones in several species of mammals,17,18 including pigs,19 primates,5,20–23 and several rodents,1,24–31 there is little information with respect to the common rat.3,32–35 Using immunohistochemical techniques, Szél and Rohlich34 in 1992 reported the presence of S- and L-cones in the albino rat retina and estimated their proportions. Among cones, 6.7% were S-cones, giving a ratio of S- to L-cones of 1:15. According to their article, the greatest percentage of cones was found in the inferior and nasal retina, followed by the superior and temporal areas, although in the central retina few cones were detected. Last year, it was reported that the mean density of L-cones in the albino rat (Sprague-Dawley; SD) is approximately 2000 cells/mm²; however, no total number or spatial distribution was provided.32 Of importance, it has been demonstrated that, in the pigmented Long-Evans rat strain, these two types of cones allow dichromatic color vision.5

We have developed an automated method to quantify the total population of retinal ganglion cells (RGCs) that were identified by either retrograde tracing or immunohistofluorescence techniques.36–37 This approach allows the objective quantification of all the RGGs present in naive or injured retinas and thus enables the assessment of RGC loss or the
degenerative course after different insults.\textsuperscript{36-38,39} Moreover, the quantitative data aid in the construction of detailed isodensity maps illustrating the distribution of RGCs, as well as their density peaks. Based on our previous experience and taking advantage of the availability of commercial antibodies raised against the UV-sensitive opsin (S- and cone) or the opsin red/green (L-cones), we proposed to double detect and quantify by an automated method, the whole population of S- and L-cones and to study their topographical distribution in flat-mounted retinas from two rat strains commonly used in the laboratory: one albino (Sprague-Dawley; SD) and the other pigmented (Piebald Virol Glaxo; PVG). The data gathered establish, for the first time, the normal values of both types of cones in adult rats and lay the foundations for further studies to analyze the effect of degenerative diseases, experimental insults, or ageing on each cone population.

**MATERIAL AND METHODS**

**Animal Handling**

All animals were treated in accordance with our institutional rules, European Union regulations, and the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. Adult rats of the SD and PVG strains were obtained from Harlan Laboratories, Inc. (Milan, Italy, [SD] and Longborough, UK [PVG]). The SD and PVG rats weighed 50 to 560 g (age, 3–3.5 months) and 80 to 160 g (age, 3–3.5 months), respectively.

**Experimental Design**

To study the S- and L-cone population, we used six retinas from each strain for radial sections and 12 (PVG) or 14 (SD) for analysis on whole, flat-mounted retinas. To quantify the total number of cones, we analyzed six whole, flat-mounted retinas from each strain. To study the correlation of RGCs and L-cone spatial distribution, we analyzed six additional retinas from each strain.

**Double Immunohistofluorescence**

All animals were deeply anesthetized with an overdose of sodium pentobarbital and perfused transcardially with 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer after a saline rinse. After death, the superior pole of the eyes was sutured for posterior orientation, as described elsewhere.\textsuperscript{38} The eyes were then enucleated and postfixied for 2 hours in 4% PFA at 4°C.

**Oriented Radial Sections**

The superior pole was labeled with yellow ink (Bradley Products, Inc., Bloomington, IL). The optic cups were dissected and cryoprotected in 15% sucrose (Sigma-Aldrich, Alcobendas, Madrid, Spain) before they were embedded, with the superior pole in known position, in optimal cutting temperature (OCT) compound (Sakura Finetec, Torrance, CA) for cryostat sectioning. Sections (15 μm thick) were blocked in 5% normal donkey serum (NDS; Jackson Immunoresearch Inc., Cambridge, UK) in phosphate buffered saline (PBS) with 0.1% Triton X-100. S- and L-opsins were double detected by an overnight incubation at 4°C with a mixture of the primary antibodies goat anti-OPN1SW (N-20), which detects the UV-sensitive opsin (i.e., S-cones; Santa Cruz Biotechnology, Heidelberg, Germany), and rabbit anti-red opsin, which detects the L-opsin in rodents (i.e., L-cones; Chemicon-Millipore Iberica, Madrid, Spain) diluted in blocking buffer (1:1100 for anti-OPN1SW and 1:1200 for anti-opsin red/green). Secondary detection was performed by incubating the sections, during 1 hour at room temperature, with Alexa Fluor-568 donkey anti-rabbit IgG (H+L) and Alexa Fluor-488 donkey anti-goat IgG (H+L; Molecular Probes–Invitrogen, Barcelona, Spain) each diluted 1:500 in PBS 0.1% Triton X-100. Finally, sections were thoroughly washed and mounted with anti-fade mounting medium containing DAPI, to counterstain the nuclei (Vectashield mounting medium with DAPI; Vector Laboratories, Alicante, Spain).

**Flat-Mounted Retinas**

Retinas from both eyes were dissected as flattened whole-mounts, as previously reported.\textsuperscript{36-38} They were permeabized in PBS 2% Triton by freezing them for 15 minutes at −70°C. After they were thawed at room temperature, the retinas were rinsed in new PBS-0.5% Triton. Primary and secondary detection was performed as for radial sections, with the exception that 2% Triton was used. In those retinas used to quantify the total number of cones, both opsins were detected with the same fluorophore, and thus, in these experiments, the secondary antibodies were Alexa Fluor-568 donkey anti-rabbit and Alexa Fluor-568 donkey anti-goat (Molecular Probes–Invitrogen), each diluted 1:500 in PBS 2% Triton. Finally, all retinas were thoroughly washed in PBS and mounted vitreal side down on subbed slides and covered with anti-fade mounting medium (Vectashield; Vector Laboratories).

Double immunodetection of L-cones and RGCs was performed as described, with the exception that the incubation with the primary antibodies was a mixture of goat anti-Brn3a (1:1000, Santa Cruz Biotechnologies) and of rabbit-anti opsin red/green (1:1200). In these immunodetections, Brn3a was detected in green (Alexa Fluor-488 donkey anti-goat) and opsin red/green in red (Alexa Fluor-568 donkey anti-rabbit). The retinas were mounted in anti-fade medium, vitreal side down, and the L-cone signal was acquired. Afterward, the retinas were mounted again, vitreal side up, to acquire the Brn3a signal.

**Retinal Analysis**

Whole-mounted retinas (left and right eyes) were analyzed for OPN1SW or Brn3a (detected with Alexa Fluor-488, i.e., green signal) and L-opsin (detected with Alexa Fluor-568, i.e., red signal). In the experiment performed to quantify the total number of cones, both opsins were detected in red. For the radial sections, S- and L-cone DAPI signals were acquired. All images were taken with an epifluorescence microscope (Axioskop 2 Plus; Zeiss Mikroskopie, Jena, Germany) equipped with a computer-driven, motorized stage (ProScan H128 Series; Prior Scientific Instruments, Cambridge, UK), controlled by image-analysis software (Image Pro Plus software; IPP 5.1 for Windows; Media Cybernetics, Silver Spring, MD), as previously described.\textsuperscript{13,36-37} Briefly, to make reconstructions of retinal whole-mounts, we photographed retinal multiframe acquisitions for each fluorophore, in a raster scan pattern in which the frames were captured, with a ×10 objective, contiguously, side-by-side, with no gap or overlap between them (Plan-Neofluar, 10×0.30; Zeiss Mikroskopie). Single frames were focused manually before the capture of the digitized images, which were then fed into the image-analysis program. Depending on the retina’s size and orientation on the slide, a scan area is defined to cover the entire retina. This scan area consisted of a matrix of m-frames in columns and n-frames in rows, where the total number of frames in the scan area was indicated by frames in columns times frames in rows (m × n). The frame size was 840 × 623 pixels, and 140 images usually were taken for each retina. The capture calibration was 1 pixel, which equals 0.0011 mm. The images taken for each retina were saved in a folder as a set of 24-bit color images. Later, these images were combined automatically into a single, tiled, high-resolution composite image of the whole retina (IPP 5.1 for Windows; Media Cybernetics). In this study, photomontages of radial sections were performed the same way as those of whole-mounted retinas, with the exception that the objective was ×20, and, usually, 114 images were taken per section. Individual images, in flat-mounted or radial sections, were taken with an ×20 or ×40 objective, acquiring the red signal first; then, without moving the microscope stage, the green signal; and finally the blue (DAPI) signal, when needed. All images were captured at a resolution of 300 dpi. Reconstructed images were further processed with image-editing computer software (Adobe Photoshop CS; ver. 8.0.1; Adobe Systems, Inc., San Jose, CA), when correct orientation of the retina or image coupling was needed.
Image Processing: Automated Quantification of the Total Population of S-cones and L-cones in Whole-Mounted Retinas

To quantify the total number of S- and L-cones present in flat whole-mounted retinas, we developed an automated counting routine, using the macro language of the software (IPP for Windows; Media Cybernetics) for each type of cone. These routines counted positive cells in each of the 180 frames spanning a flat-mounted retina.

**S-cones.** In the first step, images were converted to 16-bit gray scale to discard color information, followed by the application of the flatten filter to eliminate luminosity variations in the image. The resulting image was then filtered with a top-hat filter, which enhances the brightness of objects over that of background. The S-cones were counted within predetermined parameters to exclude objects that were too large or too small to be outer segments. Finally, data of each count were displayed and exported by dynamic data exchange to a spreadsheet (Microsoft Office Excel 2003; Microsoft Corp.), and the data were filed and saved for further analysis.

**L-cones.** In a first step, the images were converted to 16-bit gray scale to discard color information. The flatten filter was applied on the gray-scale image to eliminate luminosity variations in the image. Then, a best fit was performed to enhance the positive objects, followed by use of an enhancement filter (HiGauss, IPP software; Media Cybernetics), which highlights the image by using a Gaussian filtering method. Finally, data of each count were displayed and exported by dynamic data exchange to a spreadsheet (Microsoft Office Excel 2003; Microsoft Corp.), and the data were filed and saved for further analysis. This routine was used, as well, to quantify the total number of cones (S+L+dual) detected with the same fluorophore.

**Brn3a.** Automated quantification of Brn3a-positive RGCs was performed according to a published method.56

Validation of the S- and L-cones Automated Counting Methods

To validate both automated counting methods, three different experienced investigators counted manually, in a masked fashion, a total of 50,695 L-cones and 8,077 S-cones present in 22 frames. These frames represented different density regions and were randomly selected from eight whole-mounted retinas. The automated quantification of these images resulted in 51,928 and 8,130 L- and S-cones, respectively. Finally, each automated method was statistically compared (SigmaStat for Windows ver. 3.11; Systat Software, Inc.) to its manual counterpart, and a correlation coefficient of 0.99 (Pearson correlation test $R^2 = 0.986$) was obtained for the L-cones automated routine and of 0.98 (Pearson correlation test $R^2 = 0.977$) for the S-cones automated routine. Thus, both automated counting methods were reliable and valid.

Manual Quantification and Topographical Distribution of Dual Cones

The presence of dual cones, that is to say cones that express both opsins, was observed in both rat strains. These cones were manually quantified, in the same six representative retinas per strain for which isodensity maps were generated, as follows: At first the retinal whole-mount images for the S (green signal) and L (red signal) cones were edited and coupled (Adobe Photoshop). Then, the whole retina was examined at high magnification, frame by frame. Every dual cone, detected because the co-localization of both opsins was yellow, was dotted in a new layer. Afterward, a counting routine was developed (Image Pro Plus macro language; Media Cybernetics) and launched, to quantify the number of dots (dual cones), as described elsewhere.49 The layer with the dots was extracted and resized, to illustrate their topographical distribution.

Retinal Area Measurement

The area of each whole-mounted retina was measured on the high-resolution photomontage image of the complete retina, with the software (IPP; Media Cybernetics) calibrated off the stage movement.

Isodensity Maps

Detailed topographical distribution of S- and L-cones was demonstrated with isodensity maps in six retinas from each strain: three right and three left. These maps were constructed as previously described for RGCs,36,37 with the exception that each original frame was divided into 25 sampling areas instead of 64. These maps are filled contour plots generated by assigning to each one of the 25 subdivisions of each frame a color code according to its cone density, within a 28-step, color-scale range from 0 (purple) to 6500 or higher (red) L-cones/mm² in both rat strains. There is a significantly higher number of S-cones in the albino than in the pigmented strain; therefore, to appreciate the spatial distribution in both strains, a different density scale was given to each one. Thus, in the albino strain, the scale was from 0 (purple) to 1300 or higher (red) S-cones/mm², whereas in the pigmented strain, red was set at values of 900 or higher S-cones/mm². Isodensity maps of Brn3a-positive RGCs were generated, as previously described.36

Statistical Analysis

The analysis of the differences between groups of retinas or groups of animals was performed with the Mann-Whitney or t-tests (SigmaStat for Windows, ver. 3.11; Systat Software, Inc.). Data are shown as the mean ± SEM, and differences are considered significant at $P < 0.05$.

RESULTS

Identification of S- and L-cones in Oriented Retinal Radial Sections

The first objective was to analyze the expression pattern of both opsins in oriented radial sections. Figure 1 shows the double detection of the UV-sensitive opsin (green signal) and L-opsin (red signal) in the retina of the albino rat, and Figure 2 shows these findings in the retina of the pigmented strain. The images in both Figures 1 and 2 show the expression patterns of both opsins in different areas of the retina: dorsal (B, C, F, G, J, K), ventral (D, E, H, I, L, M), peripheral (B, E, F, I, J, M), and central (C, D, G, H, K, L). There was a mutually exclusive labeling of cone subpopulations, demonstrating that each antibody recognized a different opsin. In both strains, opsin expression was confined to the external segments of the S (UV-sensitive opsin) and L (L-opsin) cones, although at lower antibody dilutions, the cone somata and the synaptic terminals were also detected (not shown). At this low dilution, the L-opsin antibody faintly recognized, in both strains, cells located in the ganglion layer, which have been described and are known as atypical cones.41

We observed as well that both types of cones were distributed throughout the retinal section and that the L-cones were more abundant than the S-cones. A closer inspection of the coupled images disclosed that some cones expressed both opsins, which means that, in adult SD and PVG rats (yellow signal, arrows in Figs. 1J, 2J), there were dual cones (addressed in detail later). In this analysis, no differences were found in the two strains in the expression pattern of opsins and distribution of S- and L-cones.

Identification of S- and L-cones in Flat-Mounted Retinas

Next, we proposed to detect both types of cones in flat-mounted retinas. The results of this experiment are illustrated in Figure 3. These images correspond to eight areas of the
retina (Fig. 3A–H from SD and 3I–P from PVG rats) as follows: two images, one central and one peripheral, were acquired for each one of the four retinal quadrants (superior, inferior, nasal, and temporal). Central images were taken at 1 mm from the optic nerve and peripheral ones at the retinal border. In both rat strains, we observed that the S-cones were more abundant in the peripheral than in the central retina in all retinal quadrants (compare green signal in Figs. 3A, 3B, 3E, 3F, 3I, 3J, 3M, 3N with the green signal in 3C, 3D, 3G, 3H, 3K, 3L, 3O, 3P). In fact, S-cone density seemed higher in the far retinal periphery and the retinal rim (see gradient from top to bottom in Figs. 3A, 3B, 3E, 3F, 3I, 3J, 3M, 3N). Besides, when comparing the quantity of S-cones detected in each strain, it seemed that these were more abundant in the SD than in the PVG rats (compare Figs. 3C, 3D, 3G, 3H with 3K, 3L, 3O, 3P). The L-cones were more abundant in the central than in the peripheral retina (compare red signal in Figs. 3C, 3D, 3G, 3H, 3K, 3L, 3O, 3P to red signal in Figs. 3A, 3B, 3E, 3F, 3I, 3J, 3M, 3N), and no clear differences are observed between the strains.

In agreement with the radial section results, L-cones were more abundant than S-cones (compare green to red signal) and, although most of the cones in both strains were genuine S- or L-cones—that is, cones that expressed only one opsin (higher magnification in Fig. 3Q)—some cones were dual (i.e., expressed both opsins; Figs. 3R–T, arrows).

**Automated Quantification of the Whole Population of S- and L-cones**

To quantify the total number of S- and L-cones, we developed two automated counting routines using the macro language of commercial software (IPP for Windows; for details and validation of each routine, see the Material and Methods section). Figure 4 shows a representative left whole-mounted retina of each rat strain (SD, Figs. 4A, 4B; PVG, Figs. 4C, 4D) where the S (Figs. 4A, 4C) and the L (Figs. 4B, 4D) cones were doubly detected. The number of S- and L-cones counted in each retina is also shown. It is worth highlighting that, because double immunodetection was performed, both types of cones were counted in the same retinas. In total, S- and L-cones from 14 SD and 12 PVG retinas were quantified (Table 1A, Fig. 4). The mean number ± SEM of S-cones counted in the SD and PVG strains was 41,028 ± 5,074 and 27,316 ± 2,235, respectively. The difference in the total number of S-cones in both strains is statistically significant (Mann-Whitney test; P < 0.001). The mean ± SEM number of L-cones was 231,736 ± 14,517 and 239,939 ± 6,494 in the SD and PVG rats, respectively; the difference was not significant (Mann-Whitney test; P = 0.227). Therefore, the ratio of S- to L-cones was 1:5.7 ± 0.6 and 1:8.7 ± 2.9 in the SD and PVG strains, respectively.

**FIGURE 1.** Immunodetection of S- and L-cones in oriented retinal radial sections from SD rats. (A) Photomontage of an oriented radial section from an SD rat retina spanning the optic nerve. The superior pole is located at 12 o’clock. These sections were immunoreacted for UV-sensitive opsin, which is expressed in S-cones (left, green signal) and L-opsin which is expressed in L-cones (middle, red signal). These images were coupled with the nuclear stain DAPI (blue) in the right image. (B-I) Magnifications corresponding, respectively, to the squares (b–i) shown in (A). (B–E) Immunodetection of S-cones; (F–I) immunodetection of L-cones; (J, K) coupled images showing S- and L-cones and DAPI counter-staining. Arrows: dual cones. Bar, 100 μm.
Because it was also possible to determine the area of each retina, the mean density of both cones was calculated, and so in the SD strain, there was a mean ± SEM of 689 ± 84 S-cones/mm² and of 3906 ± 358 L-cones/mm², whereas in the pigmented strain their density was 442 ± 43 S-cones/mm² and 3878 ± 167 L-cones/mm². In conclusion, in both rat strains there were more L- than S-cones; furthermore, the albino strain had a significantly higher number of S-cones than did the pigmented one.

Topographical Distribution of the Whole Population of S-cones

The next step was to analyze the spatial distribution of each cone type in both rat strains. For this analysis, we generated isodensity maps based on the quantification data (for explanations see the Material and Methods section). Figure 5 shows the topography of S-cones in six retinas, three left and three right, from each strain (SD, Figs. 5A–F; PVG, Figs. 5G–L). According to these maps, S-cones were similarly distributed in both rat strains. Their highest densities (red, 1300 or higher S-cones/mm² in the SD strain, and 900 or higher S-cones/mm² for the pigmented rat) were found on the far retinal periphery and retinal rims (100–50 µm from the retinal end), both superior and inferior, although this distribution was more evident in the superior retina. Their lowest densities (blue) were found in the superior retina in the area from the optic nerve to the retinal edge. In the inferior retina, the S-cones were of intermediate density (green, yellow). Finally, an increase in the density of these cones was noted toward the inferonasal quadrant (orange).

Topographical Distribution of the Whole Population of L-cones

Figure 6 shows the topography of L-cones in the three left and three right retinas from the same SD (Figs. 6A–F) and PVG (Figs. 6G–L) rats as are shown in Figure 5. As observed for the S-cones, there were no differences between the strains in the pattern of L-cone distribution. L-cones were distributed in a fashion that seemed to be complementary of the topography of S-cones; that is, their lowest densities were found in the retinal borders (blue); the medium densities (green, yellow) were located in the medial, superior, and inferior retina; and the highest densities (orange, red) are observed above the optic nerve, along the nasotemporal axis. This distribution was easily observed when looking at the patterns of both types of cones in the same retina (for example, Figs. 5A and 6A, and so on for each panel in each figure). Of interest, the highest densities (red) were found within the central region, above the optic nerve along the nasotemporal axis, mimicking the spatial distribution of retinal ganglion cells described by our group.
which are also densest in this retinal area.\textsuperscript{36,37} This phenomenon is addressed in detail in the next section.

Quantification and Topographical Distribution of the Whole Population of Dual Cones

Because dual cones were detected, it was of interest to know how many of them there were and where in the retina they were located. In the same 12 retinas (n = 6 per strain) from which the S and L isodensity maps were obtained, dual cones were manually dotted on the retinal photomontage, and these dots were quantified by our automated system. The mean number of dual cones was 8607 ± 3325 and 7758 ± 1182 in the SD and PVG rats, respectively (Table 1A). Of note, the mean percentage of dual cones, with respect to the total number of cones, was close to 3% in both rat strains (Figs. 7A, 7B). This approach may underestimate the number of dual cones in this species, since only yellow cones in the coupled images (red signal) were indicated by the dots, which means that only those cones with a similar expression of both opsins were counted. Thus, it is possible that there were dual cones that coexpressed both opsins, with the expression of one of them being below the detection sensitivity of the protocol. For the assessment of this possibility, confocal analysis would probably render better results; however, this method is not feasible since, to capture the whole retina, at least 180 individual images are needed. Nevertheless, quantification of the total number of cones (see below) showed that these numbers are quite accurate.

![Figure 3](http://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/933456/ on 06/26/2017)

**Figure 3.** Immunodetection of S- and L-cones in flat-mounted retinas from SD and PVG rats. Immunodetection of S (green signal) and L (red signal) cones in flat-mounted retinas from SD (A–H, Q–R) and PVG (I–P) rats. Images in (A–I, Q–I) correspond to the periphery (retinal rim) of each one of the four retinal quadrants (superotemporal: A, I; inferotemporal: E, M; superonasal: B, J; and inferonasal: F, N). All images are oriented from the retinal border (top) to the medial retina (bottom). Images in (C, D, G, H, K, L, O, P) correspond to the central retina (1 mm from the optic nerve) of the same quadrants (superotemporal: C, K; inferotemporal: G, O; superonasal: D, L; and inferonasal: H, P). Most of the cones in both rat strains, express one opsin, either S (green signal) or L (red signal), these were genuine S- or L-cones, respectively (magnification in Q). However, dual cones were also identified (arrows and magnifications in R–T), these cones express L-opsin (R) and S-opsin (S). In (T) is shown the coupled image (R+S). Bar: (A–P) 100 μm; (Q–T) 50 μm.

![Figure 4](http://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/933456/ on 06/26/2017)

**Figure 4.** Quantification of the total population of S- and L-cones in SD and PVG whole-mounted retinas. Whole-mounts of representative left SD (A, B) and PVG (C, D) retinas where the S-cones (A, C) and the L-cones (B, D) were double immunodetected. Bottom right of each map: the number of S- or L-cones counted in that retina. S, superior; I, inferior; T, temporal; N, nasal. Bar, 1 mm.
As observed in Figures 8A–L, dual cones were similarly distributed in both rat strains. These images indicate that the dual cones were more abundant in the retinal far periphery, close to the retinal rim, the areas where the S-cones were denser. It is worth highlighting that, even though most of the dual cones were found in the far retinal periphery, they did not reach the retinal rim, where most of the cones were genuine S-cones (Fig. 8M). In the rest of the retina, however, they were homogenously distributed, with no resemblance to the S (superior void and inferonasal gradient) or L (nasal) cones.

TABLE 1. Total Number of Dual, S-, L-, Genuine S-, and Genuine L-cones and Total Cone Counts in the SD and PVG Rats

A. The Total Number of S- and L-cones Quantified in the Same Retina by an Automated Routine

<table>
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<th>Cone Type</th>
<th>Retinal Area (mm²)</th>
<th>Densities (mm²)</th>
<th>Estimated Total S-cones</th>
<th>Estimated Total L-cones</th>
<th>Estimated Total Dual Cones</th>
<th>Estimated Total Genuine S-cones (S-dual)</th>
<th>Estimated Total Genuine L-cones (L-dual)</th>
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For this experiment S-cones were detected in green and L-cones in red. The number of dual ones was assessed with a semiautomated method. The number of genuine S- and genuine L-cones was calculated by subtracting the number of S- and L-cones from the number of dual cones. The estimated number of total cones is the sum of dual, genuine S- and genuine L-cones. Each retinal area was automatically measured, which allowed calculating the densities of each cone type. At the bottom of each column is shown the mean ± SEM.

B. Total Number of Cones (S+L+dual)

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<td>PVG</td>
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For this quantification, S, L, and dual cones were detected using the same fluorophore; hence, all cones were labeled in red, whether they express one or both opsins. These cones were automatically quantified. Mean and SEM values are shown at the right of the table.
temporal axis) cone topography shown in Figures 5 and 6, respectively.

**Numbers and Ratios of Genuine S- and L-cones and of Total Cones**

Because, as mentioned, the number of dual cones may have been underestimated, in six retinas from each strain, both opsins were detected with the same fluorophore. In this manner, all cones were detected in red, irrespective of their opsin expression. Results from the automated quantification of cones thus detected (genuine S\(^{/H11001}\)genuine L\(^{/H11001}\)dual) are shown in Table 1B. These data show that the total number of cones did not differ from the estimated population of cones calculated by adding up the number of genuine S, genuine L, and dual cones (Table 1A). These results indicate that most, if not all, of the dual cones were identified in the double-immunodetection experiment (Table 1B).

If the total number of cones is taken into account, there were no significant differences (t-test; \(P = 0.461\)) between the strains, as the SD strain had a mean number of cones of 263,375\(6\)/4,508 and the PVG strain of 257,737\(12\)/223. Thus, the percentage of L- and S-cones with respect to the total number of cones was, respectively, 87.5% and 15.5% in the albino strain and 92.4% and 10.5% in the pigmented one (Fig. 7).

Knowing the number of dual cones present in both rat strains enabled calculating the number of genuine S- and L-cones (Table 1) by mere subtraction of the number of dual cones from the total number of S- or L-cones. Thus, these counts were 36,165\(4,475\) and 219,863\(17,708\) in the albino rat and 20,758\(2,350\) and 231,272\(7,108\) in the pigmented strain, respectively. Statistically, the number of genuine S-cones in the albino strain is significantly higher than in the pigmented rat (Mann-Whitney test; \(P < 0.001\)), and the number of genuine L-cones present in the PVG rat is significantly higher (\(P = 0.042\)) than in the SD strain. When only the number of genuine cones is considered, the ratio of S- to L-cones is 1:5.2 and 1:8.5 in the albino and pigmented strain, respectively and their percentage with respect to the total number of cones is, respectively, 83% and 14% in the albino strain and 89% and 9% in the pigmented rat—ratios and percentages close to those calculated when all the S- and L-cones are taken into account (Fig. 7).

**Parallel Distribution of RGCs and L-cones**

The spatial distribution of L-cones in both rat strains mimics the spatial distribution of RGCs described in detail by our group.\(^{36,37,42}\) To further analyze this possible co-distribution, double immunolabeling of RGCs (Figs. 9E, 9F) and L-cones was carried out in the same retinas, both types of cells quantified, and isodensity maps generated (Figs. 9A–D). The isodensity maps showed, first, that RGCs (Fig. 9A, SD strain; and 9C, PVG strain) were distributed thorough the whole retinal surface.
and, in agreement with our previous work, that their topography was not homogeneous, as they were densest along the superior nasotemporal axis, an area that has been proposed to be the visual streak in rodents,36,37,42 and, second, that L-cone distribution (SD, Fig. 9B; PVG, 9D) matched that of RGCs, as their density peak was found in the same area (Fig. 9, compare isodensities A with B and C with D). Furthermore, we observed that RGC and L-cone co-distribution expanded across the entire retinal surface, as high and low densities of each cell type followed the same pattern in both topographies, indicating that, in those areas where there were more L-cones, there are also more RGCs.

Quantitatively, the total number of RGCs in the SD retina was 82,158 and in the PVG retina was 85,117 (Figs. 9A, 9C, respectively), in accordance with our previous work, were a mean of 83,449 Brn3a+/H11001 RGCs in the SD rats36 and 85,341 in the PVG rats (Nadal-Nicolás et al., unpublished data, 2009, n = 7 retinas) was reported. In these retinas, the number of L-cones...
was 215,739 in the SD strain and 239,502 in the PVG one (Figs. 9C and 9D, respectively). Considering the mean number of RGCs$^{36,37}$ and of L-cones shown in this work, there was a ratio of RGCs to L-cones of 1:2.8 in both strains, or 1 RGC for 3 L-cones.

**DISCUSSION**

The retina of the common rat, *Rattus norvegicus*, is widely used in studies aimed at increasing the understanding of induced$^{13,43}$ or inherited$^{14,16,44}$ retinal degenerations that affect photoreceptors. However, there is little information regarding the number and distribution of cone photoreceptors in this species.$^{1,3,34,35}$ Our work determined, for the first time, the total cone counts and total number and the detailed topographical distribution of S- and L-cones present in adult animals from two inbred laboratory rat strains: one albino and one pigmented. In addition, we demonstrated the presence of dual cones in adult rats, their number, and their retinal distribution. Finally, we provided direct evidence that the topography of L-cones parallels the spatial distribution of RGCs. These data and approaches may be used as baseline and hallmark to reliably quantify, in rats, the degeneration of each type of cone associated with different insults$^{13}$ or mutations$^{15}$ as well as to assess the effects of neuroprotective therapies on the survival of each cone type.

To our knowledge, there is one report$^{3}$ in which an approximate density of cones in the Long-Evans pigmented strain is given (from 2000 to 7000 [S+L]/mm$^2$). They also report that S-cones were, in this pigmented strain, approximately 10% of the cone population (ratio S:L-cones 1:10). These data are in agreement with the results presented herein, albeit in a different pigmented strain. With respect to the albino rat, there is a recent article$^{52}$ reporting a density of L-cones in SD rats of $\sim$2000 cells/mm$^2$. This density is lower than that reported herein, but they sample the retina while we quantify the whole L-cone population. Our quantification data in the albino strain cannot be compared with the classical work by Szel and Röhlich,$^{34}$ as they show the proportion of S- and L-cones and rods related to the 5000 photoreceptors counted, rather than...
density or total numbers. In this study, the percentage of S-cones accounted for 6% to 7% of the total cone population, which gives a ratio of S-L-cones of 1:15, a ratio higher than ours. Regarding cone distribution, Szel et al. 34 have observed that in most species, including the rat, the ratios shown in the albino and pigmented rat, respectively, to S-cone ratio goes up to 15:1 and 27:1 in the albino and pigmented rat, respectively. If we consider the cone densities in the dorsomedial retina, where the L-cones peak and there is an absence of S-cones, this picture changes and the L-to S-cone ratio would decrease to 0.7:1 and 1:1 in the albino and pigmented strains, respectively. We quantified the total population of both types of cones and assessed their topographical distribution based on these data, rather than sampling the retina.

The different S-to-L ratio between both strains is explained by the only quantitative difference observed in them: The albino strain has a significantly higher number of S-cones than does the pigmented one and, in turn, the latter has a higher number of L-cones. The biological meaning of these differences is unknown, but they may be related to the differences in photopic visual function observed between albino and pigmented rats. 53, 54

There were no differences in the total number of cones between both strains or in the spatial distribution of each type of cone. In fact, in both strains, the S pigment dominates the peripheral retina, whereas the L pigment is preferentially expressed in the medial and central retina. This rough distribution is widespread in most of the mammalian species with separate cone populations, including primates and humans. 18 55 S-cones are densest in the far retinal periphery and retinal rims, both superior and inferior. Although cone-rich rims have been described in humans 23, 46 and recently in rats, 46 this is the first time that the existence of an S-cone rim in the four retinal quadrants has been fully documented, probably because the whole retina was analyzed. The function of the cone-rich rim remains a matter for discussion, 46 and indeed it is not known whether these cones are functional at all. However, Vugler et al. 46 have reported that melanopsin-expressing RGCs form a well-organized plexus at the retinal border that mirrors the S-cone rich rim, an arrangement that has also been observed in the cone-rich fovea of primates. 57 This finding leads to the hypothesis that the S-cone-rich rim in rats may be related to the function of melanopsin-expressing RGCs, which play a role in driving the circadian and the pupillomotor systems. 57 The medium S-cones densities are found in the inferior retina, more so in the nasal quadrant. The lowest S-cone density is found in the dorsal-central retina, where the L-cones and RGCs are densest. This finding is very interesting, because this area has been proposed to be the visual streak of rats. 57 The visual streak is a region of the retina specialized to provide the best vision at some point in the visual space. In primates, some birds, and reptiles, this rounded area is known as the area centralis. The main characteristic of the visual streak is that it contains a high concentration of RGCs, L-cones, and bipolar cells, but is mostly devoid in adult primates including humans, of S-cones and rods. 5, 48 Although the densities of bipolar cells and rods are not accounted for herein, we showed that in this dorsal area above the optic nerve, both L-cones and RGCs reach their maximum densities, whereas the S-cones reach their lowest, thus further supporting the role of this retinal region as the visual streak in rats. Moreover, our data demonstrate, for the first time, that in both rat strains, L-cones parallel the distribution of RGCs, as has been observed in primates.
(reviewed in Ref. 49). Furthermore, the quantification of the total population of both types of neurons gives a ratio of 1 RGC to 3 L-cones. This ratio was fairly conserved throughout the retinal surface, since, as shown in Figure 9, the density peaks and valleys of both cell types are found in the same retinal regions.

The presence of dual cones has been reported for several species of mammals, (reviewed in Ref. 5 including mice4,5,50). In mice, most of the cones are dual, but this opsin co-expression does not impair dichromatic color discrimination.50 In contrast with Szel et al.,55 we found dual cones to be present in the adult rat retina. These cones account for 5% of the total cones in both rat strains and are found highly packaged in the retinal far periphery, superior and inferior retina, but homogeneously distributed in the medial and central retina. This pattern does not fit with any of the five patterns of dual cone distribution reviewed by Lukats et al.13 It has, however, characteristic of human retinas, where dual cones are uniformly distributed throughout the entire whole retina.

In conclusion, in this work, we established for the first time the total cone counts and the total number of S- and L-cones as well as of dual cones, together with their detailed topography, in two rat strains: pigmented and albino. Furthermore, we demonstrated that RGCs and L-cone spatial distributions match, with both cell types densest in the visual streak of the rat. Altogether, these data provide the normal values of both types of cones in adult rats and are the pillar that will support further studies analyzing the effects of degenerative diseases, experimental insults, or ageing on each cone population.

Acknowledgments

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References


