A systems pharmacological approach that capitalizes on the characterization of intracellular signaling networks can transform our understanding of human diseases and lead to therapy development. Here, we applied this strategy to identify pharmacological targets for the treatment of Stargardt disease, a severe juvenile form of macular degeneration. Diverse GPCRs have previously been implicated in neuronal cell survival, and crosstalk between GPCR signaling pathways represents an unexplored avenue for pharmacological intervention. We focused on this receptor family for potential therapeutic interventions in macular disease. Complete transcriptomes of mouse and human samples were analyzed to assess the expression of GPCRs in the retina. Focusing on adrenergic (AR) and serotonin (5-HT) receptors, we found that adrenoceptor α2C (Adra2c) and serotonin receptor 2a (Htr2a) were the most highly expressed. Using a mouse model of Stargardt disease, we found that pharmacological interventions that targeted both GPCR signaling pathways and adenylate cyclases (ACs) improved photoreceptor cell survival, preserved photoreceptor function, and attenuated the accumulation of pathological fluorescent deposits in the retina. These findings demonstrate a strategy for the identification of new drug candidates and FDA-approved drugs for the treatment of monogenic and complex diseases.

Conflict of interest: The authors have declared that no conflict of interest exists.

Citation for this article: J Clin Invest. 2013;123(12):5119–5134. doi:10.1172/JCI69076.
Table 1
Expression of ARs, 5-HT receptors, and ACs in the eye and retina of C57BL/6J mice, the eye of photoreceptor-degenerated Rho−/− mice, the retina of a human donor eye, and macular tissue from monkey (Macaca fasicularis)A

<table>
<thead>
<tr>
<th>Gene</th>
<th>B6 mouse eye</th>
<th>B6 mouse retina</th>
<th>Rho−/− mouse eye</th>
<th>Human retina</th>
<th>Monkey macula</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adcy1</td>
<td>17.53</td>
<td>37.50</td>
<td>9.72</td>
<td>68.60</td>
<td>81.26</td>
</tr>
<tr>
<td>Adcy2</td>
<td>14.41</td>
<td>21.70</td>
<td>22.96</td>
<td>5.78</td>
<td>11.42</td>
</tr>
<tr>
<td>Adcy3</td>
<td>5.59</td>
<td>5.09</td>
<td>1.57</td>
<td>5.70</td>
<td>21.21</td>
</tr>
<tr>
<td>Adcy4</td>
<td>1.34</td>
<td>0.54</td>
<td>2.52</td>
<td>1.18</td>
<td>0.48</td>
</tr>
<tr>
<td>Adcy5</td>
<td>7.19</td>
<td>11.15</td>
<td>11.14</td>
<td>8.00</td>
<td>12.79</td>
</tr>
<tr>
<td>Adcy6</td>
<td>17.68</td>
<td>49.98</td>
<td>11.41</td>
<td>9.99</td>
<td>7.25</td>
</tr>
<tr>
<td>Adcy7</td>
<td>4.95</td>
<td>0.75</td>
<td>0.60</td>
<td>1.14</td>
<td>2.46</td>
</tr>
<tr>
<td>Adcy8</td>
<td>2.13</td>
<td>3.80</td>
<td>2.76</td>
<td>3.73</td>
<td>8.94</td>
</tr>
<tr>
<td>Adcy9</td>
<td>3.05</td>
<td>3.99</td>
<td>3.53</td>
<td>4.64</td>
<td>ND</td>
</tr>
<tr>
<td>Adcy10</td>
<td>0.02</td>
<td>0.03</td>
<td>0.01</td>
<td>0.01</td>
<td>0.33</td>
</tr>
<tr>
<td>ARs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adra1a</td>
<td>0.45</td>
<td>0.26</td>
<td>0.63</td>
<td>0.20</td>
<td>1.35</td>
</tr>
<tr>
<td>Adra1b</td>
<td>1.60</td>
<td>1.62</td>
<td>2.35</td>
<td>1.12</td>
<td>6.61</td>
</tr>
<tr>
<td>Adra1d</td>
<td>1.72</td>
<td>3.83</td>
<td>4.26</td>
<td>0.08</td>
<td>3.97</td>
</tr>
<tr>
<td>Adra2a</td>
<td>4.60</td>
<td>9.91</td>
<td>6.14</td>
<td>0.25</td>
<td>0.67</td>
</tr>
<tr>
<td>Adra2b</td>
<td>0.20</td>
<td>0.29</td>
<td>0.20</td>
<td>0.02</td>
<td>0.06</td>
</tr>
<tr>
<td>Adra2c</td>
<td>8.96</td>
<td>7.17</td>
<td>19.28</td>
<td>2.38</td>
<td>4.30</td>
</tr>
<tr>
<td>5-HT receptors</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Htr1a</td>
<td>0.17</td>
<td>0.28</td>
<td>0.13</td>
<td>0.15</td>
<td>0.16</td>
</tr>
<tr>
<td>Htr1b</td>
<td>1.80</td>
<td>3.99</td>
<td>1.05</td>
<td>0.80</td>
<td>8.22</td>
</tr>
<tr>
<td>Htr1d</td>
<td>1.49</td>
<td>2.58</td>
<td>1.91</td>
<td>0.00</td>
<td>0.07</td>
</tr>
<tr>
<td>Htr1f</td>
<td>0.03</td>
<td>0.04</td>
<td>0.03</td>
<td>0.44</td>
<td>0.06</td>
</tr>
<tr>
<td>Htr2a</td>
<td>0.67</td>
<td>0.72</td>
<td>0.68</td>
<td>1.30</td>
<td>0.67</td>
</tr>
<tr>
<td>Htr2b</td>
<td>0.50</td>
<td>0.43</td>
<td>0.30</td>
<td>0.39</td>
<td>0.86</td>
</tr>
<tr>
<td>Htr2c</td>
<td>0.56</td>
<td>0.73</td>
<td>0.52</td>
<td>0.02</td>
<td>0.27</td>
</tr>
<tr>
<td>Htr4</td>
<td>0.02</td>
<td>0.01</td>
<td>0.01</td>
<td>0.43</td>
<td>0.05</td>
</tr>
<tr>
<td>Htr5a</td>
<td>0.43</td>
<td>0.59</td>
<td>0.48</td>
<td>0.00</td>
<td>2.17</td>
</tr>
<tr>
<td>Htr5b</td>
<td>0.19</td>
<td>0.36</td>
<td>0.29</td>
<td>0.00</td>
<td>0.29</td>
</tr>
<tr>
<td>Htr7</td>
<td>0.31</td>
<td>0.31</td>
<td>0.41</td>
<td>0.01</td>
<td>0.59</td>
</tr>
<tr>
<td>Arrestins</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sag</td>
<td>1220.54</td>
<td>1805.26</td>
<td>301.99</td>
<td>3562.22</td>
<td>2248.35</td>
</tr>
<tr>
<td>Arrb1</td>
<td>18.38</td>
<td>18.10</td>
<td>9.42</td>
<td>8.21</td>
<td>18.22</td>
</tr>
<tr>
<td>Arrb2</td>
<td>8.86</td>
<td>14.99</td>
<td>11.92</td>
<td>17.92</td>
<td>21.68</td>
</tr>
<tr>
<td>Arr3</td>
<td>50.73</td>
<td>128.48</td>
<td>32.62</td>
<td>270.16</td>
<td>680.86</td>
</tr>
<tr>
<td>GPCR kinases</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grk1</td>
<td>111.20</td>
<td>236.84</td>
<td>14.19</td>
<td>175.06</td>
<td>88.96</td>
</tr>
<tr>
<td>Adbk1</td>
<td>28.70</td>
<td>36.88</td>
<td>36.15</td>
<td>27.29</td>
<td>32.71</td>
</tr>
<tr>
<td>Adbk2</td>
<td>2.83</td>
<td>4.68</td>
<td>3.52</td>
<td>1.30</td>
<td>8.84</td>
</tr>
<tr>
<td>Grk4</td>
<td>0.86</td>
<td>0.90</td>
<td>0.42</td>
<td>5.31</td>
<td>ND</td>
</tr>
<tr>
<td>Grk5</td>
<td>3.08</td>
<td>2.97</td>
<td>2.57</td>
<td>2.99</td>
<td>5.06</td>
</tr>
<tr>
<td>Grk6</td>
<td>20.26</td>
<td>20.31</td>
<td>6.95</td>
<td>9.69</td>
<td>9.02</td>
</tr>
<tr>
<td>Grk7</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>8.94</td>
<td>20.48</td>
</tr>
<tr>
<td>Photoreceptor genes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abca4</td>
<td>59.00</td>
<td>140.14</td>
<td>15.85</td>
<td>267.71</td>
<td>129.68</td>
</tr>
<tr>
<td>Opn1sw</td>
<td>117.79</td>
<td>187.21</td>
<td>72.95</td>
<td>25.42</td>
<td>151.31</td>
</tr>
<tr>
<td>Rho</td>
<td>5853.00</td>
<td>11081.16</td>
<td>39.26</td>
<td>6386.12</td>
<td>8168.27</td>
</tr>
</tbody>
</table>

AAnalyses were done as described in Methods. Greater expression in one compartment compared with another or from one species with another indicates enrichment in that particular compartment. Values for photoreceptor genes Abca4, Opn1sw, and Rho are shown for reference. ND, not determined because no gene homolog exists in the designated species. Preliminary results have been published (47, 83, 88).

Results
Expression of GPCRs and GPCR signaling genes in human and mouse retinas. Expression analysis of retinal GPCRs by immunocytochemistry was unreliable for reasons that include poor specificity and low affinity of antibodies (data not shown) as well as low GPCR expression. Therefore, we turned to quantitative transcriptome analysis of human and mouse retinas without specific cellular localization.

Overall, we conducted gene ontology analysis in Mus musculus and found 1,766 unique gene products categorized as having GPCR activity. The same mouse transcripts were also recognized in the retinal transcriptome from Homo sapiens. Of these 1,766, we found that 165 genes displayed expression of at least 1 fragment per kilobase of exon per million mapped reads (FPKM), equivalent to 1 transcript per cell, in C57BL/6J mouse eye or retina, with 6 genes below the 1-FPKM threshold in mouse eye, but above the 1-FPKM threshold in mouse retina (Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI69076DS1). Expression of these 165 genes from human retina is also displayed as decreasing FPKM values starting from rhodopsin (RHO), the most highly expressed gene in mouse eye. The most highly expressed human GPCR was rhodopsin, followed by retinal G protein receptor (RGR) and the cone pigments OPN1SW and OPN1MW.

We performed a more detailed analysis of adenylylate cyclases (ACs), α1-adrenergic receptors (α1-ARs), α2-ARs, and serotonin receptors (5-HTRs) to detect potential pathways that could be affected by a systems pharmacological approach. Reaction-quenching molecules, such as arrestins and GPCR kinases, were also investigated (Table 1). Expression values for these genes in mouse and human retinas highlighted those pathways that should be most susceptible to pharmacological treatment as well as those targets that would best translate from mouse models to human patients. For example, among isoforms of the α1-ARs and α2-ARs, the human retina expressed α2C-AR at the highest level (Table 1), and among 5-HTRs, 5-HT2AR had the highest expression. Data from real-time PCR analyses corroborated that the expression of these genes is readily detected in the mouse retina and/or RPE (unpublished observations of the authors).

To better understand GPCR localization in the eye and even the macula, we undertook more in-depth transcriptome studies. To potentially localize such transcripts to this photoreceptor layer, we conducted transcriptome studies of rhodopsin knockout mice (40), which exhibit no rod pigment expression and fail to form rod photoreceptors. We also performed transcriptome studies with macular tissue isolated from monkeys to learn whether these GPCRs localize there and potentially mediate the high-resolution vision disrupted in macular diseases such as Stargardt disease. Our results (Table 1) showed that robust expression of Adcy1 was present in all eye tissues, but it was attenuated in the rhodopsin knockout mouse, indic-
ative of its photoreceptor localization. We also noticed that Adcy1 expression was enriched in the monkey macula.

Pharmacological compounds targeting α1-ARs: inhibition of Gq-coupled GPCRs protects Abca4−/−Rdh8−/− mouse retinas from bright light–induced retinopathy. We evaluated pharmacological compounds antagonizing α1-ARs, a class of Gq-coupled GPCRs, for their effects on bright light–induced retinopathy according to our schematized protocol (Figure 1A). Three α1-AR antagonists including doxazosin (DOX), prazosin (PRA), and tamsulosin (TAM) (Supplemental Table 2) were independently tested in 4- to 5-week-old Abca4−/−Rdh8−/− mice shown previously to display severe light-induced retinopathy (38). These mice received DOX, PRA, or TAM 30 minutes prior to their exposure to white light at 10,000 lux for 30 minutes. We then kept the mice in the dark and evaluated the effects of each treatment 7–14 days later (Figure 1A). Noninvasive OCT imaging was performed after 7 days, revealing severely disrupted photoreceptor layers demonstrated by a diminished outer nuclear layer in DMSO vehicle–treated mice (31), whereas substantial protection of photoreceptor layer morphology was observed when these mice were pretreated with DOX, PRA, or TAM (Figure 1, B and C). We also performed SLO imaging of retinal autofluorescence 8 days later to assess light-induced photoreceptor damage. We readily observed numerous autofluorescent spots in the retinas of Abca4−/−Rdh8−/− mice pretreated with DMSO control vehicle (Figure 1, D and E), a finding typical of light-induced photoreceptor damage (41). In contrast, pretreatment with DOX, PRA, or TAM significantly protected retinas from developing light-induced damage, as shown by a substantially reduced number of autofluorescent spots (Figure 1, D and E), a finding consistent with the OCT imaging findings. These combined imaging results provide experimental evidence that pharmacological inhibition of α1-ARs can protect Abca4−/−Rdh8−/− mice against bright light–induced retinopathy.

Pharmacological compounds activating Gi-coupled α2-ARs protect against bright light–induced retinopathy. We also evaluated the Gi-coupled GPCR α2-AR as a possible target for treating photoreceptor degeneration. Agonists activating α2-ARs, including lofexidine (LOF), guanabenz (GUB), and guanfacine (GUF) (Supplemental Table 2), were each administered 30 minutes prior to exposure of 4- to 5-week-old Abca4−/−Rdh8−/− mice to bright light, and OCT images were taken 7 days later. Bright light–induced retinal morphological damage was largely prevented by pretreatment with GUB, GUF, or LOF as compared with the severe photoreceptor disruption manifested by DMSO-treated mice (Figure 2, A and B). Furthermore, we observed...
a markedly decreased number of autofluorescent spots in mice pretreated with these compounds, in sharp contrast to the large number of bright autofluorescent spots seen in DMSO-pretreated mice (Figure 2, C and D). Clearly, retinal autofluorescence images correlated positively with retinal damage revealed by OCT imaging.

Pharmacological compounds inhibiting multiple Gs-coupled GPCRs protect Abca4–/–Rdh8–/– mouse retinas from bright light–induced retinopathy. To further evaluate the therapeutic potential of compounds targeting other types of GPCRs, we also examined the effect of multiple antagonists of GPCRs coupled to Gs (Supplemental Table 2). We individually tested the antagonists that block activation of the 5-HT4 receptor RS 23579-190 (RS), the 5-HT6 receptors RO 04-6790 (RO) and SGS 518 oxalate (SGS), and the 5-HT7 receptors SB 269970 (SB) and LY 215840 (LY). We administered each compound to 4- to 5-week-old Abca4–/–Rdh8–/– mice 30 minutes before white light exposure at 10,000 lux for 30 minutes and performed OCT imaging 7 days later to evaluate retinal structural changes. In contrast to the dramatically damaged photoreceptor layer displayed by DMSO-pretreated, light-exposed Abca4–/–Rdh8–/– mice, we observed a marked preservation of the photoreceptor layer in Abca4–/–Rdh8–/– mice pretreated with compounds blocking the activation of Gs-coupled GPCRs such as the 5-HT4, 5-HT6, and 5-HT7 receptors (Figure 3, A and B). The protective effects of the indicated compounds were further supported by significantly reduced formation of autofluorescent spots in Abca4–/–Rdh8–/– mice treated with antagonists of these receptors (Figure 3, C and D). These results indicate that Gs-coupled GPCRs could also be further explored as plausible therapeutic targets for light-induced degenerative photoreceptor disorders.

Inhibition of AC prevents retinas from displaying marked light-induced degeneration. The above data demonstrate that multiple antagonists of Gs-coupled GPCRs or agonists of Gi-coupled GPCRs can protect retinas of Abca4–/–Rdh8–/– mice against bright light–induced retinal degeneration (Supplemental Table 2). Given that AC is the central player mediating the intracellular function of both Gs- and Gi-coupled GPCRs, we tested the targeting of AC in our light-induced retinal degeneration mouse model. Thus, we administered the AC-selective inhibitor SQ 22536 (SQ) to Abca4–/–Rdh8–/– mice 30 minutes before bright light exposure, followed by retinal structural examination by OCT imaging 7 days later. As shown in Figure 4, A and B, SQ treatment protected retinas from light-induced degeneration in a dose-dependent manner. Furthermore, SLO evaluation also demonstrated SQ’s dose-dependent prevention of autofluorescent spot formation, indicative of light-induced photoreceptor damage (Figure 4, C and D). These data are supportive of AC involvement in the pathogenesis of bright light–induced retinal degeneration. We performed an additional examination of the possible impact of AC inhibitor on phototransduction and found that amounts of 11-cis-retinal in the bleached eyes were comparable between SQ-treated Abca4–/–Rdh8–/– mice and vehicle-treated controls, whereas mice treated with the retinoid cycle inhibitor retinylamine showed markedly lower amounts of this visual chromophore. These data indicate that SQ treatment does not significantly change the visual cycle pigment recovery rate. Likewise, ERG responses from Abca4–/–Rdh8–/– mice were not affected by SQ administration under both scotopic and photopic conditions (unpublished observations of the authors).

Pharmacological interventions targeting Gq, Gi, Gs-coupled GPCRs, and AC preserve retinal morphology after exposure of Abca4–/–Rdh8–/– mice to bright light. We performed immunohistochemical (IHC) examination to evaluate bright light–induced retinal morphological alterations in greater detail. We applied rhodopsin labeling of rod outer segments, peanut agglutinin (PNA) labeling of cone cell
matrix sheaths in the retina, and DAPI staining of cellular nuclei to retinal sections collected from Abca4–/–Rdh8–/– mice. As shown in Figure 5, bright light–exposed 4- to 5-week-old Abca4–/–Rdh8–/– mice treated with DMSO vehicle displayed severely damaged photoreceptor structures manifested by only a residual expression of rhodopsin and PNA, together with a markedly decreased thickness of the DAPI-stained photoreceptor outer nuclear layer. In contrast, photoreceptor morphology was substantially better preserved, as indicated by abundant, well-organized rhodopsin and PNA expression as well as by a well-maintained DAPI-stained outer nuclear layer in Abca4–/–Rdh8–/– mice treated with pharmacological compounds targeting various GPCRs, including antagonists of Gq-coupled α1-ARs (DOX, PRA, and TAM), agonists of Gi-coupled α2-ARs (LOF and GUB), antagonists against either multiple Gs-coupled GPCRs such as 5-HT4R (RS), 5-HT6R (RO and SGS), and 5-HT7R (LY and SB), or AC (SQ).

Contribution of Gi and Gq pathways to light-induced retinal pathogenesis in the Abca4–/–Rdh8–/– mouse model. To further elucidate the impact of Gi and Gq GPCR pathways on the pathogenesis of bright light–induced degeneration in Abca4–/–Rdh8–/– mice, we tested both the additive effects of Gi pathway activation and Gq pathway inhibition and the opposing effects of treatment with both idazoxan (IDA) (42), an α2-AR antagonist, and GUB, an α2-AR agonist. Both GUB (an activator of the Gi pathway) (Figure 6A) and DOX (an antagonist of the Gq pathway) (Figure 6B) protected the retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion. To further evaluate these positive effects, we first determined the half-maximal protective dose of GUB to be 0.3 mg/kg (Figure 6A) and that of DOX to be 0.4 mg/kg (Figure 6B). Then we found that treating mice simultaneously with half-maximal effective doses of GUB and DOX completely protected retinas of Abca4–/–Rdh8–/– mice from developing bright light–induced degeneration in a dose-dependent fashion.
of mice pretreated with IDA and then with GUB were dramatically damaged. Moreover, this damage was more evident with an increased dose of IDA (Figure 6D). This last result further confirms the positive impact of activating the Gi pathway on retinal protection from bright light–induced degeneration.

**Pharmacological interventions targeting Gq-, Gi-, Gs-coupled GPCRs, and AC preserve retinal function.** The effects of pretreatment with selected compounds affecting Gq-, Gi-, Gs-coupled GPCRs, and AC were also examined by scotopic and photopic ERG analyses performed 2 weeks after bright light exposure (Figure 7). Bright light exposure at 10,000 lux for 30 minutes nearly abolished the scotopic ERG response in Abca4−/−Rdh8−/− mice pretreated with DMSO vehicle (Figure 7, A and B). In marked contrast, we achieved substantial preservation of this response by pretreatments individually targeting multiple GPCRs. These included DOX, an antagonist of Gq-coupled α1-AR; RS, an antagonist of the Gs-coupled 5-HT4 receptor; RO, an antagonist of the Gs-coupled 5-HT6 receptor; LY, an antagonist of the Gs-coupled 5-HT7 receptor; LOF, an agonist of the Gi-coupled α2-AR; and SQ, an inhibitor of AC. These data indicate that pharmacological treatment targeting these GPCRs also protects against light-induced retinal degeneration. A complete list of therapeutics is provided in Supplemental Table 2.

**Pharmacological interventions targeting GPCRs and AC prevent light-induced degeneration in WT mice.** To investigate whether compounds that showed a protective effect against retinal degeneration in Abca4−/−Rdh8−/− mice could also prevent retinal degeneration in WT mice, DOX (α1-AR agonist), GUB (α2-AR agonist), and the AC inhibitor SQ were further tested for their effect on light-induced retinal degeneration in BALB/c mice. These drugs were administered to 4-week-old BALB/c mice by i.p. injection 30 minutes before white light exposure at 10,000 lux for 1 hour. We assessed retinal morphology 7 days after light exposure by OCT imaging and histological examination. As shown in Figure 8, severe retinal degeneration was observed in the mice treated with DMSO vehicle, whereas retinal morphology of the drug-treated mice was maintained, and no obvious signs of retinal degeneration were observed. Moreover, SLO examination revealed an infiltration of microglia and macrophages into the subretinal space, as shown by...
an increased number of autofluorescent spots in DMSO-treated and light-exposed mice, that was prevented in mice treated with DOX, GUB, or SQ. These results indicate that pharmacological interventions targeting GPCRs and AC could also prevent light-induced degeneration in WT mice.

### Figure 5

IHC examination shows protective effects on retinal morphology exerted by therapeutics targeting Gq-, Gs-, and Gi-coupled GPCRs and AC in Abca4–/–Rdh8–/– mice. Retinal morphological changes after various indicated pretreatments were further evaluated by photoreceptor IHC examination (asterisk identifies severely reduced photoreceptor outer and inner segments manifested by only residual staining of rhodopsin and PNA in DMSO-treated control mice). Rho staining (red); PNA staining of cone extracellular matrix (green); DAPI staining of nuclei (blue). Scale bars: 50 μm.

**Pharmacological interventions targeting GPCRs and AC inhibit ROS generation in mice after light exposure.** The production of ROS is closely associated with photoreceptor cell death in Abca4–/–Rdh8–/– mice (39). Thus, we also examined compounds showing the protective effect against retinal degeneration in Abca4–/–Rdh8–/– mice for their ability to modulate ROS generation in Abca4–/–Rdh8–/– mice after light exposure. Dark-adapted 4- to 5-week-old Abca4–/–Rdh8–/– mice were i.p. injected with the fluorescent ROS probe DHE, together with DOX (α1-AR agonist), GUB (α2-AR agonist), or SQ (AC inhibitor) 30 minutes before light exposure at 10,000 lux for 1 hour. Figure 9 shows that among all the mice examined, the strongest ROS signals were detected in photoreceptor nuclei in DMSO-treated mice. Pharmacological interventions targeting GPCRs and AC were able to reduce ROS generation in Abca4–/–Rdh8–/– mice after light exposure. These results indicate that ROS generation is one of the common downstream pathways potentially mediating the effects of aberrant GPCR/AC signaling in light-induced retinal degeneration in Abca4–/–Rdh8–/– mice.

**Penetration of DOX, GUB, and SQ into mouse eyes.** Though drugs administered into the systemic circulation are distributed throughout the body, they can also achieve different concentrations in various organs and tissues depending on the rates of vascular perfusion and the drugs’ molecular properties, such as their lipid solubility, pKa, and their ability to bind to carrier proteins. Moreover, drugs...
Figure 7
Therapeutics targeting Gq-, Gs-, and Gi-coupled GPCRs and AC preserve retinal function in 4- to 5-week-old Abca4−/−Rdh8−/− mice. Abca4−/−Rdh8−/− mice were exposed to 10,000 lux light for 30 minutes after pretreatment with the pharmacological agents DOX (10 mg/kg BW), LOF (2 mg/kg BW), LY (10 mg/kg BW), RO (30 mg/kg BW), RS (20 mg/kg BW), and SQ (0.5 mg/kg BW). ERGs were recorded to evaluate the effects of these agents on retinal function 2 weeks after light exposure. (A) ERG responses were compared between mice not exposed to intense light (No light), vehicle only (DMSO), and tested agents under both scotopic and photopic conditions. Amplitudes of B waves at 1.6 log cd × s/m² under scotopic and photopic conditions are shown (B). Tested compounds showed significant protective effects when compared with DMSO-treated mice, which displayed significantly impaired retinal function as indicated by decreased ERG amplitudes. *P < 0.05 compared with DMSO control. Bars indicate SDs. n = 4–6 eyes per group.

can also be cleared from target organs and tissues at different rates. To investigate whether DOX, GUB, and SQ can penetrate and persist in the eye, we quantified the amounts of these drugs present in the C57BL/6j (WT) whole-mouse eye globes within 2 hours after drug injection and compared these levels to those in serum. Liquid chromatography–mass spectrometry–based (LC-MS–based) analyses and quantification revealed the presence of tested compounds at levels in the low picomolar range in eye tissue (Table 2 and Figure 10). Given that the total volume of a mouse eye is about 0.1 ml (43), the amounts of examined compounds were comparable to levels found in 0.1 ml of serum samples for GUB and 5 times 0.1 ml (43), the amounts of examined compounds were comparable to levels found in 0.1 ml of serum samples for GUB and 5 times

Discussion
Stargardt disease is an inheritable juvenile macular dystrophy that leads to progressive vision loss and ultimate blindness. Most cases of Stargardt disease can be traced to mutations of the atRAL transporter ABCA4 (34, 45), and its pathology directly relates to the toxicity of atRAL (31). Others have postulated that condensation products of atRAL and further light exposure cause macular degeneration (46), but how atRAL actually triggers retinal degeneration is only partially understood. Toxicity induced by atRAL is mediated through a signaling cascade implicating GPCRs, PLC/IP3/Ca2+ signaling, and NADPH oxidase (39). This discovery provided several new targets for halting progression of the disease in addition to free atRAL sequestration (31). It is also likely that atRAL plays some role in AMD, just as it does in Stargardt disease. Most plausible is that AMD results from an imbalance of several signaling pathways, such that attenuation or activation of these pathways could help preserve the retina. Genetic predisposition (47), diet, cigarette smoking, bright sunlight, cardiovascular disease, and hypertension are all risk factors for AMD (48). The age-related eye disease study (AREDS) demonstrated a substantial impact of taking dietary supplements containing high-dose antioxidants and zinc on halting disease progression and preserving visual acuity (8, 9).

In the retina, a nonredundant set of genes is primarily involved in processing the light signal and regenerating the light-sensitive chromophore (35, 49, 50). But the retina, especially its rod and cone photoreceptors, is a highly labile tissue because of its high oxygen levels (51, 52), high levels of unsaturated lipids (53), high...
metabolic demands (54), high levels of excitatory neurotransmitters (55), highly reactive and oxygen-sensitive chromophore (46, 56), and its sparse neurotrophic factors (57). Moreover, this tissue is composed of postmitotic cells with a complex intercellular organization that undergo cellular renewal processes such as phagocytosis (58, 59) and autophagy (60, 61), along with direct exposure to light action (62) causing progressive degeneration that preferentially affects the macula of older individuals with a disease known as AMD (48). This and similar studies open a new rational approach to identifying suitable compounds for the treatment of Stargardt disease and possibly AMD. The large number of even more highly expressed GPCRs in the retina with as-yet undefined locations and characterization offers an even greater challenge, but also an opportunity.

GPCRs, activating Gi-coupled GPCRs, blocking Gs-coupled GPCRs, or directly inhibiting AC prevented retinal degeneration to different degrees (Figure 12). First, without establishing their precise localization in the retina, we tested the α1-AR antagonists, a group of the three highly homologous subtypes α1A-AR, α1B-AR, and α1D-AR, the latter of which is most highly expressed in mouse retina (Table 1). High doses of long-lasting DOX, a selective α1-AR blocker, effectively protected retinal tissue from light-induced damage (Figure 1 and Supplemental Table 2), supporting a previous study that identified Gq-coupled GPCR blockage as a target (ref. 39 and Figure 12). This compound, which confers positive effects on lipid profiles, is used to treat hypertensive patients and also those with urinary retention due to benign prostatic hyperplasia (63–65). Other similar antagonists, such as short-acting PRA and the antiprostatic hyperplasia drug TAM, were consistently effective, albeit with lower efficacy (66, 67). The presence of α1-ARs in rabbit retina has been reported, although their exact compartmental location remains to be established (68). Given the role of Gq-coupled GPCRs in light-induced retinopathy (39) and that α1-AR is an important member of this class of GPCRs, further efforts should be made to identify the location of α1-ARs and to clarify their physiological significance and therapeutic targeting.

The α2-ARs associated with the Gi-inhibitory heterotrimeric G protein consist of three highly homologous subtypes that include α2-A, α2B-AR, and α2C-AR. The α2C-ARs, previously localized to the soma and the inner segment of photoreceptors (69), are the most highly expressed receptors of this group in the human retina (Table 1). Activation of these receptors, accomplished by the blood pressure–lowering drugs GUB, GUF, and LOF at relatively low doses, exhibited the most consistent protective effects in all tested mice (Figure 2 and Supplemental Table 2). GUF did not induce ophthalmological deterioration, even after long-term treatment (70, 71), whereas LOF eye drops effectively increased the pupil diameter and lowered intraocular pressure (72). Interestingly, Allergan Inc. is currently conducting phase II clinical trials of the α2A-AR agonist brimonidine tartrate for the treatment of AMD (ClinicalTrials.gov Identifier: NCT00658619), but results are not yet available. Interestingly, even a low dose of a direct AC inhibitor such as SQ was highly protective against retinal degeneration in our experi-
mental setup (Figure 4 and Supplemental Table 2). Ca²⁺-stimulated AC1 is the most highly expressed AC in the retina, and an initial immunocytochemical analysis of these enzymes in the retina has been reported (73). All AC isoforms are present in the inner plexiform layer, and both Ca²⁺-sensitive isoforms of AC (AC1 and AC3) and Ca²⁺-insensitive isoforms (AC2 and AC4) are expressed in ganglion cells. Ca²⁺-sensitive isoforms are more readily detected and expressed at higher levels in the outer retina, including photoreceptors and the outer plexiform layer.

Transgenic *C. elegans* expressing rhodopsin serves as a valid model for light-induced behavioral studies (74). DOX, GUB, or SQ was also found to be able to prevent atRAL-induced neurodegeneration in this *C. elegans* model (unpublished observations of the authors), further supporting the notion that atRAL-induced toxicity is mediated by GPCR pathways affected by these compounds. Among the 5-HTRs we tested, the 5-HT2AR (Gq) antagonist nefazodone was highly protective (Supplemental Table 2), but high doses were required, and a liver damage side effect caused this drug to be withdrawn from the US market in 2004. Antagonists of Gs-coupled 5-HT4R (R5), 5-HT6R (RO and SGS), and 5-HT7R (SB and LY), which are used to induce relaxation and treat cognitive impairments such as amnesia, anxiety, and depression in patients, were effective in protecting mouse retina only at high doses (Supplemental Table 2), in which possible side effects could pose a problem. For example, transient but severe electrocardiographic changes and episodes of premature ventricular contractions were observed in patients when RS was administered at a dose that effectively blocked the 5-HT4R activity associated with periods of electrical-mechanical dissociation, leading to severe hypotension due to impaired cardiac function (75). Among these receptors, 5-HT4R is most highly expressed in the human retina (Table 1). mRNA expression from various subtypes of 5-HTRs in human retina has been documented, including 5-HT2, 5-HT3, 5-HT4, 5-HT5, and 5-HT7, with 5-HT2 expression being the most readily detected (76). In the mouse and rat, the presence of mRNAs for 5-HT1, 5-HT-2AR, 5-HT3, 5-HT6, and 5-HT7 was demonstrated (77), supporting the involvement of 5-HT1 and 5-HT2A (39) in retinal function under certain circumstances. This last finding also implies the possible therapeutic value of targeting other HTs such as 5-HT6 and 5-HT7. Alcon Inc. carried out phase II clinical trials for the treatment of atrophic/AMD with tandospirone, an eye drop formulation of a 5-HT1AR agonist used as a neuroprotective agent against excitotoxic neuronal damage, but results showed the treatment was ineffective, and

### Table 2

<table>
<thead>
<tr>
<th>Name</th>
<th>Serum (pmol/100 µl) ± SD</th>
<th>Eye (pmol/eye) ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>DOX</td>
<td>46.5 ± 12.8</td>
<td>11.0 ± 1.7</td>
</tr>
<tr>
<td>GUB</td>
<td>7.0 ± 1.8</td>
<td>13.1 ± 4.7</td>
</tr>
<tr>
<td>SQ</td>
<td>19.3 ± 5.5</td>
<td>5.6 ± 2.2</td>
</tr>
</tbody>
</table>

*a*DOX, GUB, and SQ were detected and quantified in tissues of C57BL/J WT mice by LC-MS. All three drugs were found to penetrate and persist in the eye, with GUB reaching a higher concentration in the eye than in the serum.
this study has been terminated (ClinicalTrials.gov). It also should be noted that we found this receptor to be very weakly expressed in human retina (Table 1).

Our results indicate that GPCR pathways provide distinct activation and inhibitory actions that could control photoreceptor cell survival. Thus, activation of the Gq pathway, accomplished by the α1-AR, among others, can lead to photoreceptor cell death through the phospholipase C–mediated signaling pathway (79). Photoreceptor death can also result from aberrant functioning of Gs and Gi pathways that modulate the formation of cAMP (79). Gs, affected by the action of the 5-HT4, 6, and 7 receptors, activates this pathway, whereas Gi, mediated by the α2-AR, inhibits it.

Therefore, the degenerative photoreceptor phenotypes in pathological states could be abrogated by pharmacological inhibition of either Gq or Gs pathways or by activation of the Gi pathway. It is also worth noting that rhodopsin could be a central player in light-induced retinal degeneration (80) and that reduced amounts of rhodopsin protect the retina from light-induced degeneration (81, 82). Therefore, we examined the potential impact of the AC inhibitor SQ on chromophore regeneration, given that AC is the central player in our newly identified GPCR signaling implicated in light-induced retinopathy (Figure 12). When the effects of SQ treatment on ERG responses and 11-cis-retinal levels after bleaching were evaluated, no significant changes were observed (unpublished observations of the authors), indicating that the protection against retinal degeneration conferred by SQ treatment was
Methods

Animals. Abca4\(^{−/−}\)Rdh8\(^{−/−}\) mice were generated and genotyped as previously described (39), and mixed sexes were used for the present study when they reached 4–5 weeks of age. All mice were routinely maintained in a 12-hour light (≤10 lux)/12-hour dark cyclic environment in the Animal Resource Center at the School of Medicine, Case Western Reserve University (CWRU). For bright light exposure experiments, Abca4\(^{−/−}\)Rdh8\(^{−/−}\) mouse pupils were dilated with 1% tropicamide prior to white light exposure at 10,000 lux (150 W spiral lamp; Commercial Electric) for 30 minutes. Alternatively, 4-week-old BALB/c mice (The Jackson Laboratory) were treated with the compounds indicated in Supplemental Table 2 via i.p. injection 30 minutes prior to 1 hour of white light exposure at 10,000 lux using a 150-W spiral lamp. Assessment of retinal structural and functional changes was performed 7 days after light exposure. Rhodopsin knockout mice (Rho\(^{−/−}\)) were obtained from Janis Lem (Tufts University, Medford, Massachusetts, USA) (40). Enucleated macaque (Macaca fascicularis) eyes in RNAlater (Invitrogen) from 4-year-old animals were obtained from Ricerca Biosciences. The retina was carefully dissected from an untreated eye requiring enucleation for a large ocular melanoma and immediately placed in RNAlater.

Transcriptome analyses of the eye and retina. Eyes from 4-week-old C57BL/6J mice (The Jackson Laboratory) were enucleated and immediately processed to isolate total RNA that was used to prepare cDNA libraries for sequencing with the Illumina platform of RNA sequencing instruments (47, 83). Three biological replicates were made for both whole-eye and retinal tissue to generate transcriptome data used to determine the FPKM for normalization and differential expression analyses. The retina was carefully dissected from

unlikely due to inhibition of visual pigment regeneration. Therefore, we were able to rule out the possibility of an acute effect of pharmacological compounds on phototransduction.

Together, the experimental results described here and in a previous publication (39) identify a series of intrinsically linked events, including the participation of GPCRs, PLC/IP3/Ca\(^{2+}\) signaling, and NADPH oxidase–mediated ROS production, which are collectively responsible for the pathogenesis of retinal dystrophy in Abca4\(^{−/−}\)Rdh8\(^{−/−}\) mice, a model for rod/cone degeneration resembling features of human Stargardt disease. Our findings show that arRAX toxicity in bright light–induced retinal degeneration could be mediated through a signaling cascade implicating GPCRs, PLC/IP3/Ca\(^{2+}\) signaling, and NADPH oxidase (39). Here, we report that, in addition to Gq signaling (Figure 1, new and confirmatory observations of our previous study), inhibition of the cAMP pathway also has a protective effect against retinal degeneration (Figures 2–4).

As presented here, the degenerative photoreceptor phenotypes were reversed by pharmacological inhibition of either Gq or Gs pathways or by activation of the Gi pathway (Figure 12). These proof-of-concept studies demonstrate interactions between interconnected and diverse pathways involved in the pathogenesis of retinal degeneration induced by strong light (39). Thus, systems pharmacology promises to greatly increase our knowledge of the mechanisms underlying the multiple actions of drugs and their cross-activation and inhibition of different retinal pathways involved in vision (Figure 12).

Figure 11

GPCR-targeted therapeutics prevent formation of large fluorescent granules in the RPE of 6- to 7-week-old Abca4\(^{−/−}\)Rdh8\(^{−/−}\) mice after exposure to bright light. (A) Representative TPM images of the RPE 10 days after exposure to bright light. Upper left panel, unexposed to light (No light) control; upper right panel, exposed to bright light (Bleached) and DMSO-treated control; lower left panel, pretreated with PRA; lower right panel, pretreated with GUB. Cross sections shown at the right edge and at the bottom of each en face RPE image reveal that fluorescent granules, most pronounced in the bleached DMSO-treated control, extend across the whole thickness of the RPE and into the outer retina-DMSO-treated space. Scale bars: 25 μm. (B) Emission spectra after excitation with 730 nm light (left panel) and after excitation with 850 nm light (right panel). The spectra from light-exposed, DMSO-treated control are notably red-shifted for both excitation wavelengths.
BW; SQ3, 0.25 mg/kg BW; SQ4, 0.5 mg/kg BW; and IDA, 2.5 mg/kg BW and 5 mg/kg BW. All tested compounds were dissolved in DMSO before injection except IDA, which was dissolved in 0.9% saline.

Spectral domain optical coherence tomography. Noninvasive ultra-high-resolution spectral domain optical coherence tomography (SD-OCT) (Bioptigen) was performed for in vivo imaging of mouse retinas. Mice were anesthetized with an i.p. injection of an anesthetic cocktail consisting of ketamine (6 mg/ml) and xylazine (0.44 mg/ml) diluted with 10 mM sodium phosphate, pH 7.2, and 100 mM NaCl at a dose of 20 μl/g BW. Pupils were dilated with 1% tropicamide prior to SD-OCT imaging. Five frames of OCT images were acquired in the B-scan mode and averaged for image presentation and analysis. Retinal pathology was subsequently scored according to the following criteria: grade 0, outer nuclear layer (ONL) was completely disrupted with no visible appearance observed; grade 1, extensive disruption of an ONL spanning the retina 500 μm away from the optic nerve head with an ONL thickness less than 0.01 mm; grade 2, extensive disruption of the ONL spanning the retina 500 μm away from the optic nerve head with a measured ONL thickness between 0.01 and 0.03 mm; grade 3, reduction in ONL thickness, with a measured thickness between 0.03 and 0.05 mm; grade 4, intact ONL with a measured thickness greater than 0.05 mm. For evaluation of the impact of Gi and Gq pathways, data points were plotted as the percentage of eyes that developed significant retinal degeneration with an ONL thickness less than 0.035 mm.

SLO imaging. SLO imaging (Heidelberg Engineering) was performed for in vivo whole-fundus imaging of mouse retinas (84). Mice were anesthetized with an i.p. injection of an anesthetic cocktail consisting of ketamine (6 mg/ml) and xylazine (0.44 mg/ml) diluted with 10 mM sodium phosphate, pH 7.2, and 100 mM NaCl at a dose of 20 μl/g BW. Pupils were dilated with 1% tropicamide prior to SD-OCT imaging. Five frames of OCT images were acquired in the B-scan mode and averaged for image presentation and analysis. Retinal pathology was subsequently scored according to the following criteria: grade 0, outer nuclear layer (ONL) was completely disrupted with no visible appearance observed; grade 1, extensive disruption of an ONL spanning the retina 500 μm away from the optic nerve head with an ONL thickness less than 0.01 mm; grade 2, extensive disruption of the ONL spanning the retina 500 μm away from the optic nerve head with a measured ONL thickness between 0.01 and 0.03 mm; grade 3, reduction in ONL thickness, with a measured thickness between 0.03 and 0.05 mm; grade 4, intact ONL with a measured thickness greater than 0.05 mm. For evaluation of the impact of Gi and Gq pathways, data points were plotted as the percentage of eyes that developed significant retinal degeneration with an ONL thickness less than 0.035 mm.

Chemicals. DOX was purchased from Selleckchem. LOF was obtained from Santa Cruz Biotechnology, Inc. PRA, TAM, RS, RO, SB, SGS, LY, GUB, GUF, IDA, and SQ were purchased from Tocris Bioscience. Agomelatine, nefazodone, etoproprazine, cyproheptadine, pizotifen, GR 125487, RS 39604, SB203186, SB 399885, tamsulosin, phenoxazobenzamine, phentolamine, fexofenadine, tolterodine, ABT-724, PD-168077, and yohimbine were purchased from Sigma-Aldrich. The chemical structures of all the pharmacologic agents screened are included in Supplemental Table 3.

Mouse treatments. All experimental compounds were administered to mice by i.p. injection through a 28-gauge needle 30 minutes prior to bright light exposure. Tested compounds and their doses were: DOX, 1 mg/kg body weight (BW); 2 mg/kg BW, 3 mg/kg BW, and 10 mg/kg BW, respectively; PRA, 2 mg/kg BW; TAM, 2 mg/kg BW; RS, 20 mg/kg BW; RO, 30 mg/kg BW; SB, 30 mg/kg BW; SGS, 30 mg/kg BW; LY, 10 mg/kg BW; GUB, 0.5 mg/kg BW, 1 mg/kg BW, 1.5 mg/kg, and 2 mg/kg BW, respectively; GUF, 2 mg/kg BW; LOF, 2 mg/kg BW; SQ1, 0.083 mg/kg BW; SQ2, 0.125 mg/kg BW; SQ3, 0.25 mg/kg BW; SQ4, 0.5 mg/kg BW; and IDA, 2.5 mg/kg BW and 5 mg/kg BW. All tested compounds were dissolved in DMSO before injection except IDA, which was dissolved in 0.9% saline.

Spectral domain optical coherence tomography. Noninvasive ultra-high-resolution spectral domain optical coherence tomography (SD-OCT) (Bioptigen) was performed for in vivo imaging of mouse retinas. Mice were anesthetized with an i.p. injection of an anesthetic cocktail consisting of ketamine (6 mg/ml) and xylazine (0.44 mg/ml) diluted with 10 mM sodium phosphate, pH 7.2, and 100 mM NaCl at a dose of 20 μl/g BW. Pupils were dilated with 1% tropicamide prior to SD-OCT imaging. Five frames of OCT images were acquired in the B-scan mode and averaged for image presentation and analysis. Retinal pathology was subsequently scored according to the following criteria: grade 0, outer nuclear layer (ONL) was completely disrupted with no visible appearance observed; grade 1, extensive disruption of an ONL spanning the retina 500 μm away from the optic nerve head with an ONL thickness less than 0.01 mm; grade 2, extensive disruption of the ONL spanning the retina 500 μm away from the optic nerve head with a measured ONL thickness between 0.01 and 0.03 mm; grade 3, reduction in ONL thickness, with a measured thickness between 0.03 and 0.05 mm; grade 4, intact ONL with a measured thickness greater than 0.05 mm. For evaluation of the impact of Gi and Gq pathways, data points were plotted as the percentage of eyes that developed significant retinal degeneration with an ONL thickness less than 0.035 mm.

SLO imaging. SLO imaging (Heidelberg Engineering) was performed for in vivo whole-fundus imaging of mouse retinas (84). Mice were anesthetized by i.p. injection of the anesthetic cocktail indicated above, followed...
by pupil dilation with 1% tropicamide prior to SLO imaging in the autofluorescence mode. The numbers of autofluorescent spots were counted and subjected to statistical analyses as described below.

**IH C.** Retinal IH C was performed as previously described (85). Briefly, the eyes were enucleated and, after removal of the cornea, lens, and vitreous body, the eye cups were fixed in 4% paraformaldehyde and processed for cryosectioning. Twelve-micrometer-thick cryosections were cut, collected, and examined for rhodopsin expression, and PNA staining of the cone sheath and nuclear DAPI staining were performed.

**ERGs.** ERGs were performed as previously described (85). Briefly, dark-adapted mice were examined under dim red light transmitted through a Kodak No. 1 Safelight Filter (transmittance 560 nm). Pupils were dilated with 1% tropicamide after anesthesia induced by the method described above. Contact lens electrodes were placed on the eyes, and a reference electrode and ground electrode were positioned on the ear and tail, respectively. ERGs were recorded with the UTAS E-3000 universal testing and electrophysiologic system (LKC Technologies, Inc.).

**In vivo detection of ROS.** In vivo ROS generation was evaluated as previously described (39). The ROS probe DHE, at a dose of 20 μg/kg BW in 25 μl of DMSO, was administered to Abca4−/−Rdh8−/− mice via i.p. injection 1 hour before light exposure. Eye cups obtained after removing the cornea, lens, and vitreous body from enucleated eye globes 3 hours after light illumination were fixed in 4% paraformaldehyde. Cryosections were prepared from fixed eye cups and cut at 12-μm thickness for microscopic assessment of ROS fluorescence in the retina using ImageJ software (NIH).

**TPM imaging.** Ten days after bright light exposure, TPM images were obtained as previously described (32). Briefly, we used a Leica TCS SP5 upright confocal microscope equipped with a 1.0-NA water immersion objective and tunable laser Vision S (Coherent) delivering 75-femtosecond laser pulses at an 80-MHz pulse repetition frequency. Emission spectra were obtained with a TCS SP5 (Leica) spectrally sensitive detector in the descanned configuration. Only intact, freshly enucleated mouse eyes were used for imaging. Before enucleation, mice were anesthetized by i.p. injection of the anesthetic cocktail indicated above and then euthanized.

**Quantification of selected drugs in mouse tissue.** Six-week-old C57BL/6j WT mice (The Jackson Laboratory) were treated with DOX, GUB, or SQ at a single dose of 10, 2, and 0.5 mg/kg, respectively. Each compound was dissolved in 50% DMSO in PBS (137 mM NaCl, 2.7 mM KCl, 0.67 mM Na2HPO4/KH2PO4, pH 7.4) and administrated by i.p. injection. Thirty minutes later, the mice were anesthetized. Blood samples were collected using a cardiac puncture technique and subsequently centrifuged (5 minutes at 16,000 g) for serum collection. Prior to harvesting eyes, the mice were intracardially perfused with PBS to minimize blood contamination. Eyeballs were immediately homogenized in 0.1 ml of PBS. The homogenate and serum samples were spiked with 100 pmol of internal standard (IS) (PRA - 2-[4-((2-furyl)pyrrolidin-1-yl]-6,7-dimethoxyquinazolin-4-amino or clenbuterol - RS - 1-(4-amino-3,5-dichlorophenyl)-2-(tert-butylamino) ethanol) (both from Sigma-Aldrich) for DOX, GUB, or SQ quantification, respectively (86, 87). Next, to precipitate excess proteins, 0.3 ml of methanol was added to both the eye homogenate and 0.1 ml of serum. Samples were vortexed for 30 seconds, followed by centrifugation for 15 minutes at 16,000 g. Clear supernatants were collected and used directly for liquid chromatographic mass spectrometric (LC-MS) analysis.

MS-based detection and quantification of DOX, GUB, and SQ was performed with an LXQ linear ion trap mass spectrometer (Thermo Scientific) equipped with an electrospray ionization (ESI) interface and coupled to an Agilent 1100 HPLC (Agilent Technologies). Separation of drugs and the ISs was achieved on a reverse-phase C18 Phenomenex HPLC column (250 × 4.60 mm; 5 μm) by a linear gradient of 0% to 100% acetonitrile in water within 15 minutes at a flow rate of 0.5 ml/minute. All solvents contained 0.1% formic acid (v/v). The HPLC effluent was sprayed into the mass spectrometer via an ESI probe operated in the positive ionization mode. Parameters of ionization and detection were tuned with synthetic standards for the drugs to achieve the highest possible sensitivity. DOX and PRA (IS) were detected by selected reaction monitoring (SRM) using m/z 452.2 → 344.2 and 384.2 → 247.1 transitions, whereas GUB, SQ, and their corresponding IS clenbuterol were detected by fragmentation at m/z 231.1 → 214.0, 206.1 → 136.0, and 277.1 → 259.1, respectively. The elution times for DOX, GUB, and SQ were approximately 12.8, 12.3, and 10.8 minutes, respectively. Both ISs eluted at 12.0 minutes (Figure 10). Calibration curves were calculated based on the linear relationship between ratios of the SRM ion intensity peak area corresponding to the selected drug and the IS versus molar ratios of the compounds in a range of 20 to 500 pmol (Figure 10H).

**Statistics.** Results were collected from at least 4 mice per experimental group. Data are expressed as the means ± SEM, and statistical analyses were performed using a 1-way Students t test or ANOVA. A P value of ≤0.05 was considered statistically significant.

**Study approval.** All animal procedures and experiments were approved by the IACUC of CWRU and conformed to recommendations of both the American Veterinary Medical Association Panel on Euthanasia and the Association for Research in Vision and Ophthalmology. Clinical evaluation of the human patient from whom retinal tissue was obtained was performed at the Cleveland Clinic Cole Eye Institute (Cleveland, Ohio, USA), and written permission was obtained from the patient. This research conformed to the tenets of the Declaration of Helsinki and was conducted in accordance with approved protocols of the Cleveland Clinic Cole Eye Institute.

**Acknowledgments**

We thank Leslie T. Webster Jr. and members of Palczewski’s laboratory for helpful comments on this manuscript and Mee Lee (CWRU) for technical support. We also thank Yi Yuan and John Feng for assays performed on C. elegans. We thank Hiroshi Maeno for siRNA experiments and Bhubanananda Sahu for ERG analyses. We would also like to thank Jonathan E. Sears (Cleveland Clinic Cole Eye Institute) for ophthalmic examination of the human patient and retinal tissue isolation. This work was supported by funding from the NIH (K08EY019031, to A. Maeda; K08EY019880, to T. Maeda; EY009339, to K. Palczewski; EY021126, to K. Palczewski; EY022658, to A. Maeda; AG043645, to G. Palczewska; and P30 EY11373). Support was also provided by the Research to Prevent Blindness Foundation, the Foundation Fighting Blindness, the Fight for Sight, the Ohio Lions Eye Research Foundation, and the Program for Professor of Special Appointment at Shanghai Institutions of Higher Learning (to Y. Chen). D. Mustafi was supported in part by the NIH CWRU Medical Scientist Training Program Grant (T32GM007250) and the Visual Sciences Training Grant (T32EY007157). K. Palczewski is a John H. Ford Professor of Pharmacology and CSO of Polgenix Inc.

Received for publication January 28, 2013, and accepted in revised form September 12, 2013.

**Address correspondence to: Krzysztof Palczewski, Department of Pharmacology, School of Medicine, Case Western Reserve University, 10900 Euclid Avenue, Cleveland, Ohio 44160, USA. Phone: 216.368.4631; Fax: 216.368.1300; E-mail: kxp65@case.edu.**

Yu Chen’s present address is: Yueyang Hospital and Clinical Research Institute of Integrative Medicine, Shanghai University of TCM, Shanghai, China.


