Recovery Of Visual Function In Pacific Halibut (Hippoglossus Stenolepis) After Exposure To Bright Light

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Abstract—Commercial fishing exposes Pacific halibut (Hippoglossus stenolepis) to a myriad of stressors during capture, processing, and discarding, including exposure to direct sunlight that causes diminished retinal sensitivity. It is unknown, however, whether recovery occurs. We therefore employed both electroretinography and a behavioral assay to measure recovery of retinal sensitivity and visual function in halibut exposed to 15 min of simulated sunlight. We used electroretinography to measure changes in retinal light sensitivity after recovery periods of 2, 4, 6 and 10 weeks and a behavioral assay to measure responsiveness to simulated prey (i.e., in behavioral trials) to measure visual function after recovery periods of 2 to 6 d. Exposure to simulated sunlight significantly reduced retinal sensitivity to light with no apparent recovery after 10 weeks. Although retinal sensitivity was reduced, fish exposed to direct sunlight displayed no demonstrable deficits in visual function during behavioral trials.

Recovery of visual function in Pacific halibut (Hippoglossus stenolepis) after exposure to bright light

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One-quarter of the catch of worldwide fisheries comprises nontarget species (i.e., as bycatch or incidental catch) that are often discarded (Alverson et al., 1994). Fish may be dead when discarded, or may subsequently expire as a consequence of physical injury and stress incurred during capture and release. Mortality rates for discarded fish are, however, rarely known and represent a large source of uncertainty in fisheries models (Davis, 2002). In some instances, compromised fish succumb to predation hours or days after being discarded (Davis, 2002). For example, juvenile walleye pollok (Gadus chalcogrammus) and sablefish (Anoplopoma fimbria) subjected to stressors simulating escape through trawl codend meshes have been shown to be more vulnerable than control fish to predation in staged predator encounters (Ryer, 2002, 2004). In other instances, fish may recover but experience lower fitness as a consequence of injuries or stress. Atlantic cod (Gadus morhua) stressed through simulated trawl avoidance produced poor quality eggs and larvae (Morgan et al., 1999); and sockeye salmon (Oncorhynchus nerka) that escaped gill nets incurred physical injuries and physiological impairments that reduced spawning success by 50% (Baker and Schindler, 2009). Reduced growth and body size may also impact reproduction. Using a bioenergetics model, Meka and Margraf (2007) estimated that catch-and-release can reduce growth of rainbow trout (Oncorhynchus mykiss) up to 15% when there is no physical injury, and up to 164% where debilitating hook injuries are incurred. Although these studies have documented outcomes of bycatch stress, they rarely address the mechanisms that cause the stress. In particular, scant information exists on how capture and release may impair sensory systems such as vision, which fish rely on to locate food and avoid predation.

Pacific halibut (Hippoglossus stenolepis) are captured in trawl and longline fisheries targeting groundfishes along the contiguous United States and Canada (Davis and Olla,
Trawl fisheries are, however, required to discard all Pacific halibut, thus subjecting a significant portion of the Bering Sea and Gulf of Alaska population to capture stress (Williams and Wilderbuer). Methods to determine health and condition of Pacific halibut destined for discard are based on the physical condition of the fish and variables related to the actual fishing process (Kaimmer and Trumble, 1998). Information on fish condition, stress, and variables related to the fishing process are collected by fisheries observers, but these data can vary greatly owing to subjective differences in assessment of fish condition and trawl tow characteristics (e.g., catch weight, depth of tow, tow speed) (Pikitch et al.). Therefore, the amount of time on deck may be a better indicator of condition at release than the means of capture (i.e., trawl or longline) (Davis and Schreck, 2005).

Recent studies indicate that Pacific halibut biomass remains relatively stable, although recruitment remains weak (Stewart and Hicks), and bycatch mortality is approximately 20% within directed groundfish fisheries (Benaka et al., 2014). Also, bycatch has been slowly decreasing, although rates fluctuate depending on the location of the fishery itself (Dykstra). Continued reductions in bycatch mortality could be facilitated by a better understanding of both the physiological and behavioral mechanisms that are compromised at the time of release of bycatch and affect survival.

Pacific halibut are visual predators (Hurst et al., 2007) and frequently live in turbid coastal waters at depths ranging from 90 to 900 m (i.e., on the continental shelf) (IPHC) and therefore under low ambient light levels. After capture in trawl fisheries, individual fish are often left on deck for tens of minutes before they are discarded (Trumble et al., 1995; Davis and Olla, 2001). During this time, they can be exposed to direct sunlight (i.e., at light levels orders of magnitude above ambient levels on the seafloor) that potentially causes impaired visual function (Loew, 1976; Meyer-Rochow, 1994; Wu et al., 2006). Previous research has documented a reduction in retinal sensitivity to light in Pacific halibut after 15 min of exposure to simulated sunlight (Brill et al., 2008). This reduction in sight could have consequences for foraging success after release by diminishing the ability of a fish to perceive and capture prey. It is unknown, however, whether this deficit is permanent or whether it reduces the ability of Pacific halibut to detect and capture prey. Our objective was to extend previous research (Brill et al., 2008) and to assess specifically whether retinal sensitivity and overall visual function can recover after exposure to simulated sunlight. We addressed these objectives by using both electroretinography (ERG) and behavioral methods. ERG measures the summed potential of electrical signals within the retina, providing a technique for rapidly and quantitatively assessing retinal function (Brown, 1968). An evaluation of the behavior of Pacific halibut subjected to bright light, namely an evaluation of their ability to accomplish essential tasks, such as perceiving and capturing prey, will help determine the effects of bycatch on somatic growth, fecundity, and survival.

### Materials and methods

All fish capture, maintenance, handling, and experimental procedures followed accepted protocols and were in compliance with all relevant laws and regulation. Age-0 Pacific halibut (40–70 mm in total length (TL)) were acquired by trawl net in Chiniak Bay, Kodiak Island, Alaska (57°40′N, 152°30′W) and delivered to the Hatfield Marine Science Center, Newport, Oregon. Pacific halibut were kept in 3.1-m diameter fiberglass tanks (at a 1-m depth) with flowing seawater at 8–10°C degrees for 2 or 3 years before use in the experiments. The tanks were maintained under low-illumination fluorescent lighting (photon flux density of 0.01 µmol·m⁻²·s⁻¹) and day time and night time were set on a 12-h photoperiod. Fish were fed 3 times per week during the first year and twice per week during the second year with a gel food consisting of gelatin, vitamins, amino acid supplements, krill (*Euphausia superba*), pelleted food, Pacific herring (*Clupea pallasii*), and squid.

### Exposure to bright light

Individual 2-year-old Pacific halibut (13–17 cm TL) fish were lifted by dip net from their holding tank, lightly anesthetized with a tricaine methanesulfonate (TricaineS™ [MS-222], Western Chemical, Inc., Ferndale, WA) solution of ~5 mg/L to reduce movement and stress, and held in a shallow seawater bath (12°C). They were

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6 Mention of trade names or commercial companies is for identification purposes only and does not imply endorsement by the National Marine Fisheries Service, NOAA.
then exposed to simulated sunlight for 15 min by using a light source and a fiber optic guide aimed at the right eye of a fish. The left eye was covered with a light-blocking cloth. The 15-min simulated sunlight exposure was chosen to correspond with the time fish are left on deck during commercial trawl sorting operations (Davis and Olla, 2001; Davis and Schreck, 2005). Control fish were treated in kind, except that the light source was not turned on. Fish were subsequently returned to their holding tanks and separated with a barrier to allow both control and light exposed fish to be held under identical conditions.

Sunlight was simulated by using a high-intensity xenon lamp (Spectral Products, Putnam, CT) and its spectral range was ~320–700 nm, which approximates the visible (400–700 nm) and the UV range of sunlight (Lalli and Parsons, 1997). Light intensity exiting the fiber optic light guide was ~2000 μmol·m⁻²·s⁻¹ (measured over 400–700 nm of spectral range) and simulated sunlight (2010 μmol·m⁻²·s⁻¹) and measured at Newport, Oregon, under ideal clear conditions at 1200 noon PST on 5 October 2007 and by using a IL 1700 Research Radiometer (International Light Technologies, Inc., Peabody, MA) equipped with a photosynthetically active radiation-filtered waterproof sensor.

**Evaluation of visual function with the use of an ERG**

To evaluate visual function by using ERG, fish were moved into a dark room in a light-proof container. Individuals were then lightly anesthetized with a buffered MS-222 solution (~5-mg/L) and the neuromuscular blocking drug gallamine triethiodide (Flaxedil, Sigma Chemical Co., St. Louis, MO, dose ~20 mg/kg) injected into the caudal vein to reduce movement. Fish were then placed on a sling and enclosed in a light-blocking container placed in an acrylic box. The body of the fish was submerged in a manner such that only a small portion of the head and the eye would remain above water to receive the light stimulus. The container was supplied with flow-through seawater (12°C) and the gills of the fish remained aerated by means of a small submersible pump for water circulation. Fish were adapted to darkness for a minimum of 1 h before physiological measurements were taken.

Teflon-coated silver wire electrodes with a silver chloride electroplated coating, were used to record the ERG responses. The recording electrode was placed lightly on the corneal surface and the reference electrode was placed on the skin over the head of the fish. The recording chamber was illuminated with a dim red light (peak wavelength 660 nm) produced by light-emitting diodes (LEDs); these remained on while the electrodes were positioned. The recording system was grounded by using a stainless-steel plate within the experimental apparatus. ERG signals were amplified (10,000x gain) with 1-Hz high pass and 1-kHz low-pass filter settings on a DAM50 amplifier (World Precision Instruments, Inc., Sarasota, FL). The signal was also filtered with a HumBug active electronic noise eliminator (Quest Scientific Instruments, Inc., North Vancouver, Canada) that removed 60-Hz noise and was digitized at a 1-kHz sampling frequency with a multifunction data acquisition card (DAQCard-6024E, National Instruments Corp., Austin, TX). Light stimuli and all data were controlled by a custom program developed by Eric Warrant (University of Lund, Lund, Sweden) for use in the LabVIEW graphical programming system for measurement and automation (National Instruments Corp.).

A circular (3.8-cm diameter) light source (SL2420 spot light, Advanced Illumination, Inc., Rochester, VT) was used to produce a white LED light stimulus, and a thin diffuser and collimating lens were used to produce an even field of illumination (~10%). An intensity controller (CS410, Advanced Illumination, Inc.) was used to control light output. The intensity controller was connected and controlled by the analog output of the data acquisition card. To extend the range of available light levels, a series of neutral density filters (Kodak Optical Products, Eastman Kodak Co., Rochester, NY) were used to dim the light stimulus.

As in previous studies (e.g., Brill et al., 2008), we examined changes in retinal sensitivity to light resulting from exposure to simulated sunlight by recording the summed potential of electrical signal in response (in volts [V]) to a range of light intensities (I) and subsequently used the data to construct voltage in relation to log light intensity response curves (V-log I). Light intensities were increased by 0.2 log-unit steps from a level with no measurable response, to a level that produced a max response. A light stimulus consisted of a train of five 200-ms light flashes delivered 200 ms apart. This stimulus was presented every 5 s and repeated 5 times at each light intensity. The ERG responses to the final flash of each train were recorded and averaged. At the conclusion of an experiment, fish were euthanized with either a massive overdose (>300 mg/kg) of sodium pentobarbital (Beuthanasia-D, Merck Animal Health, Madison, NJ) injected into the caudal vein, or by immersion in a bath of clove oil where the clove oil solution was circulated over the gills by a small submersible pump.

Initially, we compared ERG data for the left and right eyes of control fish (n=4) that had not been exposed to simulated sunlight. Preliminary analysis indicated that right eyes produced a consistently stronger voltage signal than left eyes. Our original intention had been to use unexposed left eyes as ‘within-fish’ controls for the exposed right eyes in the exposure recovery experiment. However, because of the difference in signal strength between left and right eyes, we abandoned this strategy and relied instead upon a comparison of right eyes between control fish and sunlight exposed fish after various periods of recovery. Fish exposed to simulated sunlight were divided in groups with recovery times of 2, 4, 6, and 10 weeks. Each group consisted of 8–10 individuals.

In addition to voltage response data we also calculated voltage percent maximum (p-max) data; for each
fish, namely the percentage of maximal response at each tested light intensity. Finally, the data from each individual ERG curve was fitted by using a second-order polynomial equation with SYSTAT software, vers. 13 (Systat Software, Inc., San Jose, CA) or Microsoft Office 2013 (Microsoft Corp., Redmond, WA), because the ERG response curves generally were of a sigmoid shape. To provide a summary measure of visual impairment, we calculated log-scale illumination required to produce a 50% p-max response from each fish. In the left and right eye, and exposure recovery experiments, ERG responses presented as voltages and pmx responses were examined with repeated measures analysis of variance (ANOVA) (Sokal and Rohlf, 1969). For examination of the light level required to produce a 50% p-max response, we compared treatment groups, using one-way ANOVA (Sokal and Rohlf, 1969). Where appropriate, we employed a Tukey’s honestly significant difference (HSD) test (Sokal and Rohlf, 1969) to examine differences in treatment means. Tests were considered significant at the P<0.05 level.

Behavioral evaluation of fish in relation to visual function

Individual 3-year-old Pacific halibut (21–27 cm TL) were anesthetized with MS-222 as described above, but in this case both eyes were subjected to a 15-min exposure to simulated sunlight before behavioral experiments. After light exposure, pairs of fish were moved into 1.9-m diameter × 80-cm deep circular tanks to recover. The tanks were located within a light-controlled laboratory and supplied with constantly flowing seawater at ~9°C.

Experiments were conducted with 8–10 pairs of fish at six light intensities simulating environmental conditions typical for Pacific halibut (~90–900 m): 1×10⁻³, 1×10⁻⁴, 1×10⁻⁵, and 1×10⁻⁶ µmol-m⁻²-s⁻¹, and complete darkness (<0.01×10⁻⁷). Light levels were measured on the bottom of the experimental tank with a IL1700 Research Radiometer equipped with a photosynthetically active radiation-filtered waterproof sensor. To reduce shadows, all lighting was attached to an overhead ring suspended 1.8 m above the tank bottom and approximately 0.7 m outside the tank circumference. Four cone lamps with green LED (~555-nm) clusters were mounted on the ring. The LED clusters were linked to a rheostat that was used to vary light intensity. The lights were placed directed perpendicular to the tanks to avoid glare and hot spots.

We recorded fish movements with an overhead video camera (Ikegami Electronics, Inc., Mahwah, NJ) and under infrared illumination. Infrared illumination ranged from 760–880 nm, which is a range undetectable by Pacific halibut (John, 1964; Higgs and Fuiman, 1996; Brill et al., 2008). Infrared lights were placed below the bottom of the tank and provided a silhouette of the fish; these lights were left on for all experimental trials, regardless of the light treatment being used. Each experimental tank had a clear Plexiglas tube placed in the middle that held a white fishing jig that was attached to the ceiling with a counter-weighted line and to the bottom of the tank with an elastic band. The bottom 20 cm of the Plexiglas tubes were covered with black tape, such that the jig would not be visible to the fish when not in use.

Fish were allowed to recover for at least 48 h after exposure to simulated sunlight before use in further trials. Each pair of fish was tested at all 6 levels of illumination: 2 illumination levels on each of the first 2 days, and a single illumination level on the last day. The illumination level set with the rheostat and fish were allowed to acclimate for 2 h before the trial began, 2 h were allowed between trials, and the order of testing with respect to illumination level was randomized. A trial at each illumination level consisted of two 5-min periods before and after presentation of the visual stimulus (white jig). After the first 5-min period, the jig was moved up and down rapidly (within the Plexiglas tube) for 60 s and then allowed to sink back below the masked bottom of the Plexiglas tubes, where it was out of sight. Each minute was split into 10-s intervals and scored as to whether the pair of fish reacted to the visual stimuli. A reaction was considered positive if the fish either 1) moved one body length, 2) made oral contact with the column while attempting to bite at the jig, or 3) re-oriented itself such that the long axis of the fish was pointing toward the jig (~10°).

Scoring behavior of fish

Scores were recorded as either 0 (no reaction by either fish), 1 (reaction by one fish), or 2 (reaction by both fish). For each 1-min trial, the 10-s scores were summed to arrive at an activity index. We compared activity indexes of fish exposed to simulated sunlight and control fish over time at each light level by using repeated measures ANOVA (n=6-9). Where ANOVA results indicated significant differences, a Tukey’s HSD was used to determine differences between group means. During the scoring process and in preliminary analysis it became apparent there was no difference between the lowest light levels (1×10⁻⁵, 1×10⁻⁶, and 1×10⁻⁷ µmol-m⁻²-s⁻¹ and complete darkness). Hence, we decided to show only the highest 4 light intensities (1×10⁻³, 1×10⁻⁴, 1×10⁻⁵, and 1×10⁻⁶ µmol-m⁻²-s⁻¹).

Results

Electroretinography experiment

At the same light intensities, voltages measured on the corneal surface of the right eyes of control fish were significantly higher than those measured on the corneal surface of left eyes. This finding was manifest by a significant interaction between eye (left vs. right) and light intensity in our ANOVA (F[16, 32]=4.18, P<0.0001). The difference in the responses of right and left eyes increases with increasing light intensities (Fig. 1). When voltage data for each fish were converted to p-
Figure 1

Comparison of responses to increasing illumination or light intensities (I, measured in log candela/m² by using electroretinography) between right and left eyes of previously unexposed Pacific halibut (*Hippoglossus stenolepis*) (n=4). To construct voltage in relation to log light intensity (V-log I) response curves, light intensities were increased in 0.2 log units from levels that produced no measurable responses to those that produced maximal responses. The data are reported either as voltage or as log-normalized by expressing the average response to an intensity step as a percentage of the maximum observed average response (p-max). V-log I response curves were created with both (A) voltage data and (B) log-normalized data expressed by the average response to an intensity step as a percentage of the maximum observed average response. Data points represent mean values, and error bars indicate standard errors of the means.

max, a significant difference was no longer present between left and right eyes (F[1, 2]=0.00, P=0.963), nor was there a significant interaction between eye and light intensity (F[16, 32]=0.90, P=0.575). P-max continued to increase with increasing test light level (F[16, 32]=17.68, P<0.001).

Exposure to simulated sunlight for 15 min resulted in a visual deficit that did not improve during the 10 weeks of recovery. Voltages measured from the right eyes of control fish (i.e., no exposure to simulated sunlight) were generally greater than those of the right eyes of fish that were exposed to simulated sunlight and allowed to recover for 2–10 weeks. This was particularly evident at lower test light levels, as evidenced by a significant interaction between treatment and light intensity (Fig. 2A; F[16, 32]=1.55, P=0.009). Conversion of voltages to p-max did not appreciably change this relationship (Fig. 2B). Again, there was a significant interaction between treatment and light intensity (F[16, 32]=2.04, P<0.001).

There were significant differences in light intensities required to produce a response 50% of maximum (F[4, 21]=11.1, P<0.001) between treatments (control, and 2, 4, 6 and 10-weeks recovery) (Fig. 3). The light intensity required to produce a response 50% of maximum was significantly lower for control fish, than for fish in any of the recovery treatments (Tukey’s HSD: P<0.05). Among the recovery treatments, the light intensity required to produce a response 50% of maximum increased over the 10-week recovery period and was lower at week 2 than at week 10 (Tukey’s HSD: P<0.05). The light intensity required to produce a response 50% of maximum at week 2 did not differ from those at either weeks 4 or 6, and similarly, the response at week 10 did not differ from responses at weeks 4 or 6 (Tukey’s HSD: P<0.05). In context, it took approximately 17 times the photons to produce a response of 50% of maximum in fish exposed to simulated sunlight after 10 weeks than it did for control fish.

**Behavioral experiment**

There was no effect of exposure to simulated sunlight on the behavioral response of Pacific halibut to the visual cues associated with a simulated prey (F[1, 2]=0.40, P=0.539). This lack of difference between control and treated fish was consistent throughout the trials, as well as across ambient light levels, because ANOVA showed no significant interactions between treatment (control vs fish exposed to simulated sunlight) and any of the other factors (e.g., time, ambient light level). Pacific halibut were generally active and responded strongly to the appearance of prey (presented at the beginning of minute 6) at the highest ambient illumination (3×10⁻³ µmol·m⁻²·s⁻¹), but responsiveness progressively declined at lower ambient light levels (Fig. 4). This finding is supported by a significant interaction between time and ambient light level in our ANOVA for Pacific halibut activity (F[1, 2]=4.16, P<0.001). At the 2 highest ambient light levels, fish would orient themselves toward the simulated prey when it appeared, swim toward it, and repeatedly strike at the sides of the Plexiglas tube containing the simulated prey. This behavior was characterized by a sharp in-
Comparison of responses to increasing illumination or light intensities ($I$, measured in log candela/m²) by using electroretinography between a control group of Pacific halibut (*Hippoglossus stenolepis*) and another group of Pacific halibut after 2, 4, 6, and 10 weeks of recovery ($n=8–10$). To construct voltage in relation to log light intensity ($V$-log $I$) response curves, light intensities were increased in 0.2 log-unit steps from levels that produced no measurable responses to those that produced maximal responses. The data were log normalized by expressing the average response to an intensity step as a fraction of the maximum observed average response. Each curve was then fitted with a second-order polynomial equation because the ERG response curves generally indicated a sigmoid response to light intensities. Light intensities required to produce a response 50% of the maximum response were taken from the predicted values produced from the quadratic equation for each model. All data points were those recorded from the right eye. Data points represent means, and error bars indicate standard errors of the means.

Discussion

Prior research (Brill et al., 2008) has shown that exposure to simulated sunlight (i.e., imitating the situation experienced on the deck of a vessel) impairs the retinal function of Pacific halibut. The authors speculated that exposure to simulated sunlight resulted in damage and apoptosis of photoreceptor cells containing the longer wavelength (520–540-nm) absorbing visual pigments. A predominance of receptors with maximal sensitivity in the green wavelengths is characteristic of coastal and continental shelf species (Levine and MacNichol, 1979; Bowmaker, 1990). If permanent, a deficit in these retinal receptors could have negative consequences for post release foraging success, somatic growth, reproductive success, and ultimately survival.
Results of the behavioral experiment quantifying responses of pairs of Pacific halibut (*Hippoglossus stenolepis*) to a visual stimulus (i.e., a white jig that simulated prey) at 4 ambient light levels (photon flux density): (A) $1 \times 10^{-3}$, (B) $1 \times 10^{-4}$, (C) $1 \times 10^{-5}$, and (D) $1 \times 10^{-6}$ µmol·m$^{-2}$·s$^{-1}$. For each experiment, ambient light level was set with a rheostat and fish were allowed to acclimate for 2 h before the next trial began. Each trial consisted of a 5-min period before and a 5-min period after presentation of the visual stimulus. After the initial 5-min period, the jig was rapidly moved up and down within a Plexiglas column for 60 s and then allowed to sink back to the level at which it was out of sight of the fish (i.e., to the masked bottom of the column). A reaction was considered positive if the fish 1) moved one body length, 2) made oral contact with the column as it attempted to bite at the jig, or 3) reoriented itself such that its long axis was directly pointing toward the jig. Scores were recorded at 10-s intervals as either 0 (no reaction by either fish in the pair), 1 (reaction by one fish), and 2 (reaction by both fish). For each minute, the scores were summed to arrive at an activity index.

Using both ERG and a behavioral assay, we tested the hypothesis that Pacific halibut recover from retinal damage and visual function resulting from exposure to direct sunlight. Our ERG data indicated damage to the Pacific halibut visual system and no significant recovery during the 10 weeks after exposure. Even after 10 weeks, it took approximately 17 times the light intensity to elicit a response 50% of maximum than with control fish. This result equates to an approximate 94% reduction in retinal sensitivity. In contrast, our behavior assay (which occurred 2–6 d after exposure to simulated sunlight) could not reveal impairment of the ability of Pacific halibut to detect visual cues associated with simulated prey across a broad range of ambient light levels.

Electroretinography is a procedure in which the summed electrical responses from the retinal photoreceptors are recorded by placing electrodes on the corneal surface and skin adjacent to the eye. In our study, we exposed fish to 15 min of simulated sunlight, an intensity equivalent to ambient sunlight under clear skies at noon (Newport, Oregon, 5 October 2007; the same exposure used by Brill et al., 2008). Light-exposed fish required approximately 5 times the amount of light to generate an ERG response equal to control fish. This was manifest as a depression in both voltage and $p$-max voltage plotted against log illumination. These curves remained depressed over a 10-week post exposure period, compared with controls that indicated no recovery of retinal sensitivity. Brill et al. (2008) speculated that the mechanism of damage was disruption of photoreceptor cells and predicted that the process would be progressive and permanent. Our ERG data support this contention. The illumination required to stimulate a 50% maximum response, shows that vision deteriorated from 2 weeks to 10 weeks.
after exposure, indicating a progressive worsening of Pacific halibut retinal sensitivity over time. In an environmental context, a sunlight exposed Pacific halibut would have to move to water that is 18 m shallower to have the same visual acuity as that of an unexposed fish, assuming a light extinction coefficient of 0.15 (e.g., simulating typical conditions in the Gulf of Alaska). This level of illumination would potentially result in a shoaling effect among fish discarded as bycatch. Our data further indicate that the visual deficit associated with sunlight exposure was most pronounced at the low end of the Pacific halibut visual range. As a consequence, fish captured in and subsequently returned to relatively shallow well-lit waters may be less affected than fish captured from and then returned to deeper water, where impaired fish may be at the limit of their range of visual sensitivity. Whether or not discarded Pacific halibut move to shallower water to mitigate visual impairments could be tested in future research with mark-recapture techniques. It should be noted that Pacific halibut size generally increases with depth. The fish used in our ERG were 2 year olds and therefore were smaller than most fish encountered in commercial fisheries. Although we have no reason to believe that the visual systems of our fish differed from those of larger Pacific halibut, future work in this area would benefit from an examination of a wider range of fish sizes.

The impairment of retinal sensitivity revealed by ERG contrasts with the results from our behavioral assay that produced no statistical evidence of significant visual impairment associated with exposure to simulated sunlight. The simulated prey bobbed up and down within a clear Plexiglass tube that minimized cues associated with water movements and the possibility that Pacific halibut would respond to nonvisual cues. The fact that the responsiveness of fish, as measured by activity, decreased with decreasing ambient light levels clearly indicates that Pacific halibut use vision to detect prey. Yet, across the range of ambient light levels there were no consistent statistical differences between control fish and those exposed to simulated sunlight, with the possible exception of a slight reduction in behavioral activity (i.e. movement, bait strike, etc) among the latter at an ambient light level of 1x10^{-4} µmol·m^{-2}·s^{-1} (Fig. 4B). Pacific halibut are visual predators and at light levels of 1x10^{-4} µmol·m^{-2}·s^{-1} primarily use visual cues to locate and attack prey, shifting to tactile and olfactory cues as light levels fall below 1x10^{-5} µmol·m^{-2}·s^{-1} (Hurst et al., 2007). For immobile baits, Pacific halibut feeding performance is likewise facilitated by vision (Stoner, 2003). We initially reasoned that the threshold ambient light level for visual foraging would be that at which a deficit would be most pronounced. It is possible that we performed tests over too wide a range of ambient light levels. For example, we might have seen a difference between sunlight-exposed and control fish by testing over finer gradations of ambient light levels between 1x10^{-5} to 1x10^{-4} µmol·m^{-2}·s^{-1}). Additionally, conditions in this behavior-
sensitivity in Pacific halibut across a broad range of illuminations, and the ~94% reduction in light sensitivity does not recover for during 10 weeks. Visual sensitivity appeared to be most affected at low ambient light levels. If this impairment is permanent, we speculate that fish may either make the best of a bad situation if they are released into deep waters, or attempt to move to shallower water to compensate for their visual deficit. However, these conclusions from our ERG data conflict with our behavioral data and observations, where no clear impairment in simulated prey detection was observed. We suspect that our behavioral assay may not have been ideally designed to show differences in visual sensitivity. We are not aware of any other studies that have attempted to link visual function, as measured by methods such as ERG, with behavioral performance, which ultimately determines the fitness of a species with visual deficits. This is an area of research that will be needed to assess the consequences of damage to the visual system resulting from conditions onboard vessels before discard of bycatch (Pacific halibut and other fish species), and to assess the implications of such damage for fisheries management.

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