Retinoic acid inhibitors mitigate vision loss in a mouse model of retinal degeneration

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Rod and cone photoreceptors degenerate in retinitis pigmentosa (RP). While downstream neurons survive, they undergo physiological changes, including accelerated spontaneous firing in retinal ganglion cells (RGCs). Retinoic acid (RA) is the molecular trigger of RGC hyperactivity, but whether this interferes with visual perception is unknown. Here, we show that inhibiting RA synthesis with disulfiram, a deterrent of human alcohol abuse, improves behavioral image detection in vision-impaired mice. In vivo Ca²⁺ imaging shows that disulfiram sharpens orientation tuning of visual cortical neurons and strengthens fidelity of responses to natural scenes. An RA receptor inhibitor also reduces RGC hyperactivity, sharpens cortical representations, and improves image detection. These findings suggest that photoreceptor degeneration is not the only cause of vision loss in RP. RA-induced corruption of retinal information processing also degrades vision, pointing to RA synthesis and signaling inhibitors as potential therapeutic tools for improving sight in RP and other retinal degenerative disorders.

INTRODUCTION

Age-related macular degeneration (AMD) and retinitis pigmentosa (RP) are the most prevalent photoreceptor degenerative disorders, impairing vision or causing blindness in hundreds of millions of people worldwide (1, 2). Both AMD and RP progress gradually, with retinal light responses and visual perception declining over years to decades. Despite the slow deterioration of rods and cones, most retinal ganglion cells (RGCs) survive and maintain synaptic connectivity with the brain until late stages in disease (3–5). This has made RGCs a potential substrate for artificial vision restoration by optoelectronics (6, 7), optogenetics (8, 9), and opthalmopharmacology (10, 11). Because vision restoration technologies supplant light responses initiated by residual rods and cones, treatments based on these technologies are being deployed only in a small fraction of people with late-stage retinal degeneration. For patients whose vision is severely impaired but not yet eliminated, treatments for improving sight remain an unmet medical need.

Studies on mouse, rat, and rabbit models of RP show that, while downstream retinal neurons do survive, they nonetheless undergo morphological and physiological remodeling. Months after photoreceptors are lost, new dendritic branches begin to sprout from downstream neurons and their cell body positions start to change, mirroring remodeling events that occur over years to decades in advanced human RP (12–15). Much earlier, within days of photoreceptor loss, some RGCs become hyperactive, firing spontaneously at up to eight times their normal rates in healthy retinas (16, 17). RGC hyperactivity results partly from changes in presynaptic neurons (18, 19) and partly from changes intrinsic to RGCs, including up-regulation of voltage-gated ion channels, which increases membrane excitability (11), and up-regulation of ligand-gated channels, which increases resting membrane permeability (20). Recent studies indicate that retinoic acid (RA) is the signal that triggers morphological (21) and physiological (17, 22) remodeling. Adding exogenous RA to healthy retinas mimics remodeling, adding RA receptor (RAR) inhibitors to degenerated retinas suppresses remodeling, and increased RA-induced gene expression can be detected in degenerated retinas (17). Local atrophy of photoreceptors, induced by subretinal implantation of a metallic chip, also caused elevation of RA-induced gene expression in corresponding RGCs, leading to local hyperactivity (23). This suggests that RA-induced hyperactivity is a common sequel to photoreceptor loss, whether the underlying cause is hereditary, as in RP, or environmental, as in many cases of AMD.

The functional consequences of remodeling on information processing and visual perception have remained uncertain (24). Heightened spontaneous RGC firing could mask responses triggered by residual photoreceptors, as has been observed in the rd10 mouse model of RP (17). However, degeneration-induced hyperactivity (18) and hyperpermeability (20) apply only to Off-RGCs. Accelerated spontaneous firing in Off-RGCs is predicted to compress the dynamic range to light decrements that normally provoke firing and expand the dynamic range to light increments that normally suppress firing. How these seemingly opposite effects might affect higher-order information processing in the brain is unknown. The effects of hyperactivity on visual perception are also unclear. Genetically inhibiting RAR increased light avoidance behavior in vision-impaired rd10 mice (17), but whether this reflects improved visual acuity or enhanced photophobia is unknown.

There is circumstantial evidence that retinal hyperactivity also applies to humans with degenerative blindness (14, 24). Patients with RP report persistent photopsia (25, 26) and they have a heightened threshold for perceiving phosphenes induced by electrical stimulation of the eye, consistent with interference from heightened spontaneous retinal activity (27). However, without tools to suppress hyperactivity, it is impossible to differentiate the direct effects of photoreceptor signal loss from the indirect effects of increased RGC background firing (i.e., “noise”).

Here, we have used a pharmacological approach to block RA signaling, giving us a way to discriminate the effects of decreased signal...
from increased noise. By investigating visual perception with behavioral experiments and higher-order neural processing with functional brain imaging, we show that RA-induced retinal hyperactivity is a major contributor to vision impairment in the rd10 mouse. Moreover, by inhibiting RA, we reveal a new therapeutic strategy for mitigating vision loss that may be applicable across a wide range of photoreceptor degenerative disorders, regardless of the underlying etiology.

**RESULTS**

**Inhibitors of RA synthesis or RA signal transduction reduce degeneration-induced gene expression in the retina of rd10 mice**

We intervened at two different steps in the RA signaling pathway in an attempt to suppress or reverse degeneration-induced remodeling of RGCs in the rd10 mouse model of RP (Fig. 1A). RA is a retinoid, derived from dietary retinol (vitamin A). Retinol is converted into

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**Fig. 1. Disulfiram and BMS 493 reduce signaling through the RA pathway in the degenerated retina.** (A) RA signaling pathway, including sites of intervention by disulfiram and BMS 493. RDH, retinol dehydrogenase. (B) Design of RAR reporter. CMV, cytomegalovirus promoter; RFP, red fluorescent protein; P2A, 2A self-cleaving peptide; RARE, RA response element; SV40, simian virus 40 promoter; GFP, green fluorescent protein. (C) Design of disulfiram experiment. AAV2 encoding the RAR reporter was injected into both eyes of rd10 mice at P30. Starting at P60, mice received food with disulfiram or no drug for 20 to 30 days, and retinas were obtained at P80 to P90 for imaging. (D) Flattened Z-stacks of the ganglion cell layer showing cells expressing RFP and GFP (spinning disk confocal microscopy). (E) Quantification of RA-induced expression. The value of GFP/RFP was measured for each labeled cell. The violin plot depicts the whole dataset. The bar graph shows the mean values for each experimental group ± SEM. Control: n = 733 cells from 15 retinal samples from six eyes in three mice; disulfiram: n = 535 cells from 15 retinal samples from six eyes in three mice. F(1,1266) = 12.51, ***P = 4.20 × 10^-4; two-tailed F test. (F) Design of the BMS 493 experiment. AAV2 encoding the RAR reporter was injected into both eyes at P30. At 45 to 60 days later, one eye was injected with 1 μl of vehicle [phosphate-buffered saline (PBS) ×1] and the contralateral eye with 1 μl of BMS 493 (5 μM). Retinas were obtained 4 to 6 days after injection for imaging. (G) Representative images for (F). (H) Quantification of RA-induced gene expression similar to (E). Control: n = 385 cells from 15 retinal samples from eight eyes from four mice; BMS 493: n = 448 cells from 15 retinal samples from eight eyes from four mice. F(1,831) = 66.44, ***P = 1.32 × 10^-15; two-tailed F test.
retinaldehyde by retinol dehydrogenase, which is expressed in retinal pigment epithelium (RPE) cells and Müller glial cells (28–30). 11-Cis retinaldehyde, the opsin chromophore for rod and cone phototransduction, can be regenerated after photoisomerization by enzymes of the visual cycle, which are also expressed in RPE and Müller glial cells. However, retinaldehyde can also be converted into RA by the enzyme retinaldehyde dehydrogenase (RALDH), which is expressed in choroid, RPE, and retina (31, 32). RA crosses cell membranes and binds to RAR, a nuclear protein that heterodimerizes with the retinoid orphan receptor (RXR). The complex, in conjunction with other coactivator proteins, then binds to specific DNA sequences to enhance transcription of genes (33–35). RAR and RXR are expressed in many cells, but because physiological changes intrinsic to RGCs can account for much of degeneration-induced hyperactivity (17), we focused specifically on RGCs.

To assess pharmacological inhibition of degeneration-induced gene expression, we used a reporter gene construct delivered to RGCs with an adeno-associated virus (AAV) vector (Fig. 1B). The construct encodes red fluorescent protein (RFP), which is expressed constitutively in all virally transduced cells, and green fluorescent protein (GFP), which is expressed only upon RAR activation by RA (17, 23). Because GFP expression reflects the level of RAR activation and RFP expression reflects the level of viral transduction that can vary between retinas, the GFP/RFP ratio can be used as a metric of degeneration-induced RA signaling (17).

Our first target for intervention was RALDH. RALDH is a member of a large family of enzymes known as aldehyde dehydrogenases (ALDHs). Disulfiram (Antabuse) is a Food and Drug Administration (FDA)–approved irreversible inhibitor of ALDHs (36), including all members of the RALDH subfamily (37). Disulfiram is usually prescribed for chronic alcoholism and is taken orally. Ingested ethanol is converted into acetaldehyde, which is broken down by ALDHs. By preventing its breakdown, disulfiram allows buildup of acetaldehyde, discouraging alcohol consumption by causing severe hangover symptoms, known collectively as the disulfiram ethanol reaction (38). In the absence of alcohol use, treatment with disulfiram causes infrequent adverse reactions that resolve completely after stopping treatment (39).

To assess whether disulfiram can inhibit degeneration-dependent activation of the RA pathway, we injected the eyes of rd10 mice with the RAR reporter virus early in degeneration (Fig. 1F). In each mouse, we delivered BMS 493 in one eye and vehicle in the contralateral eye by intravitreal injections. Last, we imaged the retinas isolated from these mice (Fig. 1G). RGCs in BMS 493–injected eyes had a mean GFP/RFP value of 0.12 ± 0.01, significantly lower than RGCs in vehicle-injected eyes with a ratio of 0.38 ± 0.07 [F(1,831) = 66.44, P = 1.32 × 10−15; gLMEM: fixed effect for treatment, random effects for mouse and retina grouped by mouse; Fig. 1H]. These findings establish that delivery of BMS 493 through intravitreal injection suppresses RA-induced gene expression even more effectively than orally administered disulfiram.

### Disulfiram and BMS 493 reduce RA-induced RGC hyperactivity without affecting residual photoreceptor function

We next asked whether inhibiting the RA pathway could suppress RGC hyperactivity in the rd10 retina. We used mice midway through the photoreceptor degeneration process (P40 to P45) when RGCs are already hyperactive (17, 41). Oral disulfiram must cross several barriers to inhibit RA synthesis and subsequent downstream effects in the retina, so we assessed its effects on hyperactivity after 40 days of continuous treatment (Fig. 2A), sufficient time to inhibit RA-induced gene expression (Fig. 1E). Control littermates were given food without disulfiram for the same period.

After treatment, we isolated the retina and prepared flat-mounted samples for multielectrode array (MEA) recordings of spontaneous RGC activity. Recordings showed less spontaneous firing in darkness in samples from disulfiram-treated mice than from control mice (Fig. 2B). Overall, mean firing frequency across all units was 5.34 ± 0.54 Hz with disulfiram, significantly less than 7.90 ± 0.78 Hz for control [F(1,972) = 4.36, P = 0.037; gLMEM: fixed effect for treatment, random effects for mouse and retina grouped by mouse; Fig. 2C]. Cumulative probability analysis shows that disulfiram scales down firing in highly active units (P = 1.75 × 10−5, Kolmogorov-Smirnov test; Fig. 2D). In disulfiram–treated retinas, we detected a mean value of 44.3 spontaneously active units per retinal sample, as compared to 51.6 active units in control retinas, a reduction of 14.2% (table S1).

Next, we assessed BMS 493 (Fig. 2, E to H). One eye of each mouse was injected with BMS 493 and the contralateral eye with vehicle, allowing intra-animal comparisons (similar to Fig. 1F). We again used mid-degeneration rd10 mice (P40 to P45), but because the drug was injected directly into the eye and administered only once, inhibition of RA signaling begins much more quickly and is more transient. Therefore, recordings were obtained much earlier in the life of the mice, at 4 to 6 days after injection (P45 to P50) (Fig. 2E). This is long enough to substantially inhibit RA-induced gene expression (Fig. 1H), but at a younger total age than in the disulfiram experiments, accounting for the lower-baseline firing rate (compare Fig. 2, B versus F).

Retinas obtained from eyes injected with BMS 493 showed less spontaneous activity than retinas from contralateral eyes injected with vehicle (Fig. 2F). The mean firing rate of BMS 493–treated retinas was 1.71 ± 0.34 Hz, significantly lower than in vehicle-injected retinas of 3.31 ± 0.35 Hz [F(1,404) = 20.87, P = 6.53 × 10−6; gLMEM: fixed effect for treatment, random effects for mouse and retina grouped by mouse; Fig. 2G]. Cumulative probability analysis of all units shows that similar to disulfiram, BMS 493 decreased the probability of encountering high–firing frequency RGCs (P = 0.001, Kolmogorov-Smirnov test; Fig. 2H). In BMS 493–treated retinas,
the mean number of spontaneously active units per retinal sample was 22.5, while the same value was 38.7 in the vehicle-injected counterpart eye of the same mouse, a decrease of 41.9% (table S1).

RAR expression is abundant in the outer retina, particularly in cone photoreceptors (42). Hence, it seemed possible that disulfiram and BMS 493 might affect events in the outer retina, independent of their effect on RGCs. To address this possibility, we recorded the photopic electroretinogram (ERG) in rd10 mice (43). We focused on the b-wave, which reflects light-modulated synaptic transmission primarily from cones to bipolar cells, because the rods are largely degenerated at this age (P60 to P80). We found that neither disulfiram (fig. S1, A and B) nor BMS 493 (fig. S1, C to E) caused a significant change in the amplitude of the b-wave across a range of photopic intensities (see also table S2). These results suggest that inhibiting RA does not affect events in the outer retina, leaving events in the inner retina as the primary mechanism of RA-induced hyperactivity.

Disulfiram and BMS 493 enhance behavioral detection of images in rd10 mice

Genetically interfering with RA signaling augments light avoidance behavior in rd10 mice (17), but whether inhibiting RA signaling can improve image-forming vision has remained unknown. To investigate this, we used a contrast sensitivity test involving operant conditioning, inspired by a previous paradigm for measuring visual threshold (44). We used a behavioral cage outfitted with a unidirectional running wheel, orienting the mouse toward a computer display (Fig. 3A). The cage also has a nose poke sensor, which, when engaged, triggers delivery of a water droplet reward (10% sucrose). Running on the wheel digitally gates the onset of a full-contrast grating displayed on the screen, while simultaneously unlocking the nose poke. Mice are water- and food-deprived before each session, providing motivation for finding the association between the visual stimulus and availability of reward. The period of reward availability was shortened during three successive training sessions to increase...
**Fig. 3. Measuring behavioral image detection in mice.**

(A) Apparatus for operant-conditioned image detection. Wheel running gates the display of a conditioned stimulus image associated with the availability of reward (a 10% sucrose water droplet). Reward is obtained by engaging the nose poke within the allotted reward interval.

(B) Training to 100% contrast images occurs over three sessions (each 12 hours) with decreasing reward intervals, shaping response latency (table S3).

(C) Contrast test was performed under the same conditions as session 3, but the visual stimulus presented was randomized between discrete contrasts from 0 to 100%. See table S3 and fig. S2.

(D to F) Responses of an individual wild-type (WT) mouse during training sessions 2 (D) and 3 (E) and to three different contrasts (0, 8, and 100%) during contrast sensitivity test (F). Trials of equal contrasts were grouped for quantifying latency, although they were randomized during the test. Red lines in responses to 100% contrast show the upper and lower limits of the value range for a single trial to be considered successful (means ± 2 SD).

(G) Grouped data showing full contrast sensitivity curve. The response speed (latency in seconds) plotted against the contrast of each visual stimulus (n = 5 WT mice, P60).

(H) Full contrast sensitivity curve showing success rate as a function of contrast in the same group of mice as in (G).

(I) Comparison of response speed and success rate in the same group of WT mice at P60 versus P100, with no training session carried out between the two time points. Individual data for each mouse are shown in gray; mean ± SEM values are shown in green (latency) and red (success rate), respectively. P > 0.05, Wilcoxon rank sum test.
response stringency (Fig. 3B and table S3). Only those mice achieving a criterion level of success during training (>80% of trials with a response latency of <4 s) with 100% contrast were carried forward for subsequent testing of contrast sensitivity. This involved randomly varying the image contrast over nine distinct values over the course of a testing session (Fig. 3C). All images were equiluminant irrespective of contrast, and monitors across different cages were calibrated for uniform brightness (see Materials and Methods).

Figure 3 (D to F) exemplifies the behavioral responses of an individual wild-type (WT) mouse during training and testing. During training, the mouse learns to associate the full-contrast image with the availability of reward (Fig. 3D). At the beginning of training session 2, response latencies were prolonged and inconsistent, but by the end, responses occurred within 2 to 3 s of stimulus presentation. The response latency remained at 2 to 3 s during training session 3 when the reward availability period was reduced (Fig. 3E), further reinforcing the association.

We tested contrast sensitivity on a mouse that had achieved the criterion level of success during training. We presented stimulus images at different contrasts, randomized from trial to trial (Fig. 3F). Low-contrast images elicited responses with longer and more variable latencies than high-contrast images (Fig. 3F). The mean latency of five WT mice subjected to the contrast sensitivity test is gradually reduced from ~8 s at 1% contrast to ~2 s at 100% contrast (Fig. 3G). The mean latency at 100% contrast was used to set the range of responses considered as successful ones for all other contrasts and subsequent tests (Fig. 3F). A nose poke was considered successful if it happened at a time range of −2 SD to +2 SD of the mean (a mouse with a response SD ≥ 40% of the mean was excluded from analysis; see Materials and Methods and tables S4 and S5). Success rate was then used to generate a full contrast sensitivity curve (Fig. 3H). To test the possibility that mice might fail the test because of behavioral extinction, we tested the same five mice 40 days apart with no “refresher” training session in between (Fig. 3I). We found that their response latency and success rate did not differ in the second test from the first, demonstrating that, once they learn, mice are able to retain this visually guided operant conditioning for long periods of time.

Next, we carried the same procedure to quantify the effects of disulfiram on contrast sensitivity. Mice younger than P35 run on the wheel infrequently and are poor learners of this task. Therefore, we started training at P35 to P40, when some photoreceptor degeneration had already begun (Fig. 4A) but early enough to ensure consistent short-latency responses to the 100% contrast image (Fig. 4B and C). All mice were first trained and tested in the absence of any pharmacological manipulation and were randomly assigned to treatment or control groups ahead of the second test.

Figure 4B shows the behavior of a trained rd10 mouse early (test #1) and late (test #2) in degeneration. Without drug treatment, short-latency responses disappeared nearly completely because of vision loss. However, in a mouse receiving disulfiram, short-latency responses were retained (Fig. 4C). Group data from early degeneration rd10 mice show the already impaired contrast sensitivity caused by partial death of photoreceptors (compare Fig. 4D to Fig. 3H). After test #1 was complete, mice were randomly assigned to the control or disulfiram group, but retrospective analysis of their performance in test #1 showed similar contrast sensitivity (P > 0.05, Mann-Whitney test; table S4). However, when tested for the second time 40 days later, mice fed with disulfiram-containing food (2 mg/kg) performed significantly better than control when reacting to high-contrast images (64 and 100% contrast; P = 0.005 and P = 0.002, respectively, Mann-Whitney test; table S4). As expected, experiments on WT
Inhibiting the RA pathway sharpens the tuning of cortical neurons to complex visual stimuli

Behavioral evaluation of image-forming vision in mice requires days of training and testing, but physiological evaluation of neuronal responses can be characterized rapidly, without behavioral training, through functional Ca$^2+$ imaging. Calcium imaging has the additional advantage of providing responses from many neurons simultaneously. Responses to a wide range of visual stimuli, from bars of specific orientation to movies with complex naturalistic scenes, can be elicited in a single session. Cortical imaging can provide a more thorough description of how photoreceptor degeneration degrades visual function and how the RA pathway inhibitors disulfiram and BMS 493 might alleviate vision impairment.

To test the effect of disulfiram, rd10 mice were given a diet with or without the drug for 28 days, beginning early in the process of degeneration and ending late, after visual function was severely impaired (Fig. 6A). Midway through the treatment period, we performed a surgical procedure (a hemicraniectomy) under anesthesia to facilitate intracortical injection of an AAV encoding the Ca$^2+$ indicator GCaMP6s, which included the Ca$^{2+}$/calmodulin-dependent protein kinase II (CaMKII) promoter to target expression to excitatory neurons (see Materials and Methods). Following injection, we inserted a glass cranial window to provide optical access for subsequent imaging of GCaMP6s expressing neurons at the end of the treatment period. Imaging data were collected blind to treatment conditions.

Many excitatory neurons in the primary visual cortex respond selectively to stimuli of a particular orientation (45). To determine whether disulfiram treatment influences orientation tuning, we measured responses to drifting periodic gratings presented to the eye contralateral to the cranial window (Fig. 6B and C). We found that a lower percentage of neurons were orientation-tuned in rd10 mice (16.06%) than typically observed in WT mice using a similar approach (~30% in (46)), but the fraction tuned was significantly increased with disulfiram treatment (22.69%). In addition, disulfiram mice showed no significant effect of disulfiram on behavioral contrast sensitivity (fig. S3A), consistent with disulfiram acting specifically by inhibiting RA synthesis.

A similar series of tests were conducted to assess the efficacy of BMS 493 in boosting contrast sensitivity in degenerative rd10 mice (Fig. 5A). Similar to experiments with disulfiram (Fig. 4), rd10 mice were trained and tested and then randomly assigned for binocular intravitreal injections of BMS 493 or vehicle. An individual rd10 mouse receiving intravitreal BMS 493 showed more short-latency responses than an age-matched rd10 mouse injected with vehicle (Fig. 5, B and C). Contrast sensitivity curves show that BMS 493 treatment preserved short-latency responses to 100% contrast stimuli (Fig. 5, D and E), while almost none of these responses were seen in vehicle-treated mice ($P = 0.041$, Mann-Whitney test; table S5). Control experiments on WT mice showed no augmentation of vision after BMS 493 treatment (fig. S3B). This is consistent with our previous studies showing little or no ongoing RA-induced gene expression in healthy retinas of WT mice (17), supporting the conclusion that BMS 493 improves vision specifically by blocking RAR. Together, these findings indicate that behavioral vision loss can be alleviated by inhibiting the RA pathway, consistent with RA-induced remodeling, making a major contribution to corrupting vision in retinal degenerative disorders.

**Fig. 5. BMS 493, an RAR inhibitor, improves visual contrast sensitivity in rd10 mice.** (A) Experimental design. Rd10 mice were reared in the dark until P55. At P55 to P60, they were trained and tested for contrast sensitivity and placed under 14:10 light/dark conditions to induce degeneration. They received binocular intravitreal injections containing 1 μl of BMS 493 (5 μM) or vehicle (PBS ×1) and tested for a second time 4 to 6 days later, at P80 to P85. (B and C) Responses of individual mice in the vehicle (B) and BMS 493 (C) groups. The response speed (latency) to the visual stimulus at 100% contrast is shown for each trial, at early degeneration and before intervention (test #1, left) and at late degeneration and after intervention (test #2, right). (D) Results of 12 mice before administration of drug during test #1. Success rate is plotted as a function of contrast; values are shown as means ± SEM. Retrospective analysis of mice for test #1, according to condition in test #2, found no significant differences ($P > 0.05$ for all contrasts, Mann-Whitney test; table S5). (E) Contrast sensitivity curve obtained later in degeneration and following treatment with vehicle or BMS 493. *$P = 0.041$, Mann-Whitney test (see also table S5).
treatment caused significantly stronger orientation tuning as compared to control \( F(1,1127) = 4.77, P = 0.029; \) gLMEM: fixed effect for treatment, random effect for mouse; see Materials and Methods; Fig. 6, D and E]. This was quantified with a cross-validated orientation selectivity index (OSI) measure that compares the magnitude of response to preferred versus nonpreferred orientations, where the preferred orientation is determined using a separate subset of trials to eliminate spurious orientation tuning due to bursts of activity (see Materials and Methods).

We next asked whether inhibiting the RA pathway can improve the detection of responses to complex naturalistic stimuli. To investigate this, we presented a 30-s natural scene (http://observatory.brain-map.org/visualcoding/stimulus/natural_movies) that contains a rich diversity of visual information. Others have found that individual
neurons in the mouse visual cortex respond preferentially at different time points in the movie, reflecting their tuning to different combinations of higher-order features (47). By presenting the same clip to the mouse many times, we could measure the response reliability across trials while remaining unbiased to any specific visual feature (Fig. 7A). A highly reliable response suggests that a neuron is coding information about that particular frame with high fidelity.

We found high variability in the response properties of cortical neurons in rd10 mice, with many exhibiting low reliability. However, the distribution of reliability values was shifted significantly by disulfiram treatment, with an increase in the proportion of neurons showing high reliability \( F(1,1610) = 4.79, P = 0.029; \) gLME: fixed effect for treatment, random effect for mouse; Fig. 7B]. On average, reliability across the population was increased by \( >1.7 \) -fold, from 0.1022 to 0.1751 by disulfiram, consistent with higher-fidelity detection of high-order visual features. Because of the high variability in response properties, we questioned whether disulfiram affected all cells equally (an additive shift) or unequally (a multiplicative shift). To answer this, we first calculated deciles of the reliability for each population of neurons and then separated and compared reliability within each matched decile group under control and disulfiram conditions (Fig. 7C). We found a significant interaction between treatment condition and decile group, suggesting that the effects of disulfiram treatment differentially affect neurons, depending on their response fidelity [treatment and decile group interaction: \( F(1,1610) = 223.61, P = 2.13 \times 10^{-37}; \) gLME: fixed effect for treatment and decile, random effect for mouse].

We next reasoned that sharpened feature selectivity would improve the population-level representation of the naturalistic movie stimulus. To test this, we used an ideal observer analysis to decode the current frame of the movie based on the neuronal population responses (Fig. 7D; see Materials and Methods). We carried out this analysis over many iterations, using different sized pools of randomly selected neurons from both disulfiram and control mice. To quantify the performance of the decoder, we calculated the percentage of iterations in which the decoded frame was within 10 frames (1 s) of the actual frame (Fig. 7E). We found that decoding in disulfiram-treated mice consistently outperformed that of control mice across neuronal pool sizes [treatment group effect: \( F(1,79) = 16.20, P = 1.30 \times 10^{-4}; \) gLME for treatment and pool size]. These experiments indicate that disulfiram treatment leads to more robust encoding of complex features found in naturalistic scenes, consistent with improved visual perception.

We next repeated these analyses to ask whether inhibiting RAR with BMS 493 could also improve visual processing of complex stimuli (Fig. 8A). For these experiments, each mouse was anesthetized and

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**Fig. 7. Disulfiram treatment improves coding of naturalistic scenes in V1.** (A) Top: Example frames from the movie clip presented to rd10 mice. Bottom: Example responses from three different neurons to repeated presentations of the movie. Each row is the response to a single presentation. Different neurons show varying amounts of response reliability to repeated presentations. (B) Comparison of the response reliability of neurons from disulfiram-treated versus control-treated mice (experimental design similar to Fig. 6A). Neurons in disulfiram mice show significantly greater reliability compared to neurons from control mice. (C) Comparison of reliability segmented by deciles. Reliability is similar in lower deciles but becomes significantly greater for disulfiram-treated mice in higher deciles. (D) Performance of a population-level decoder across control (left) versus disulfiram-treated (right) mice. (E) Across all population sizes used for decoding, disulfiram treatment shows significantly improved performance (predicted frame within 10 frames of actual frame) compared to control. (B, C, and E) Values shown are means ± SEM. Control: \( n = 1006 \) neurons across seven mice; disulfiram: \( n = 2075 \) neurons across nine mice. Two-tailed \( t \) test in (B), \( F(1,2066) = 5.27, *P = 0.022; \) in (C), \( F(1,2064) = 58.08, ***P = 3.82 \times 10^{-14}; \) in (E), \( F(1,79) = 16.20, ***P = 1.30 \times 10^{-4}. \)
BMS 493 was injected intravitreally into one eye while the contralateral eye received vehicle. At 4 to 6 days after injection, we compared responses of cortical neurons to visual stimuli displayed individually to each eye. We recorded Ca$^{2+}$ responses from the monocular zone of the primary visual cortex, which selectively receives information from the contralateral eye. This procedure allowed a within-mouse comparison of retinal degeneration between individual mice.

Comparison across treated and untreated neuronal populations showed significantly increased OSIs in neurons receiving input from the BMS 493–treated eye than from the vehicle-treated eye [$F(1,1076) = 10.67, P = 0.001$; two-tailed $F$ test]. These neurons also exhibited markedly increased response reliability to the naturalistic movies [$F(1,1959) = 191.58, ***P = 1.23 \times 10^{-41}$; two-tailed $F$ test]. As with disulfiram, comparing reliabilities across deciles also suggested a multiplicative effect of BMS 493 [$F(1,1959) = 191.58, ***P = 1.23 \times 10^{-41}$; two-tailed $F$ test]. Last, ideal observer analyses...
showed much greater decoding accuracy in V1 neurons contralateral to the BMS 493–treated eye compared to neurons contralateral to the vehicle-treated eye \( F(1,144) = 14.49, P = 2.08 \times 10^{-3} \); gLMEM for treatment and pool size; Fig. 8, F and G). Together, these experiments indicate that BMS 493 can rescue responses to complex visual stimuli in downstream cortical neurons during retinal degeneration.

**DISCUSSION**

**Degeneration-induced retinal hyperactivity corrupts vision in rd10 mice**

As photoreceptors degenerate, the retina undergoes morphological and physiological plasticity, some of which is thought to be adaptive, but much of which is maladaptive. In certain respects, downstream circuit function is resilient. Selective loss of rods leads to compensatory changes in the rod synaptic pathway to restore synaptic strength \((48, 49)\), and loss of cones leads to similar changes in the cone pathway \((50)\). However, our studies have shown that RA-induced hyperactivity is maladaptive, adding background noise that obscures signals that are already attenuated by the loss of photoreceptors \((17)\). Here, we find that RA-induced hyperactivity impairs not only simple light detection but also higher-order visual capabilities, such as reliable detection of specific visual scenes.

Visual information is ordinarily transmitted from the retina to the brain in the form of an RGC spike frequency code. Hypothetically, the code might be replicable with prosthetic devices or optogenetic tools that artificially stimulate RGCs in response to light \((51)\). However, accurate replication of the code is limited by the background firing rate of RGCs. If artificially stimulated firing is superimposed on hyperactive spontaneous firing, then the proper encoding of input stimuli will be altered, corrupting the accurate representation of visual features. Corrupted neural processing masks visual system function at multiple levels, from RGC responses \((17)\) to orientation tuning and representation of scenes by the visual cortex (Figs. 6 to 8) to behavioral image recognition (Figs. 4 and 5). The loss of photoreceptors is the ultimate cause of vision loss, but our findings indicate that physiological remodeling in downstream neurons contributes significantly to visual decline. Moreover, inhibiting RA-induced changes can mitigate impairment of simple or complex visual functions, even in mice with late-stage photoreceptor degeneration. These findings suggest RA synthesis or signaling inhibitors as tools for therapeutic improvement of low vision in humans.

**Relevance for human vision disorders**

Indirect evidence strongly suggests that RA-induced retinal hyperactivity contributes to human vision impairment \((24)\). Direct evidence, however, will be difficult to obtain because methods for detecting hyperactivity are invasive and therefore inappropriate for humans. In animal models of RP, hyperactivity can be detected directly with electrophysiological recordings from isolated retina, but RP is a rare disease, greatly restricting the availability of postmortem human retinal samples for possible electrophysiological recording. Noninvasive methods appropriate for the intact human visual system include field potential recordings from retina (ERG) or brain (electroencephalogram) and radiological methods such as functional magnetic resonance imaging or positron emission tomography scans. However, at least present, these methods are designed to report stimulus-evoked responses, not background activity.

Likewise, directly detecting heightened RA in the human eye is fraught with difficulties. RAR binds to all-trans RA with a dissociation constant in the subnanomolar range \((K_d = 0.4 \text{ nM} (33))\), suggesting that the ambient concentration of RA is very low. In addition, RA is labile owing to rapid enzymatic degradation by enzymes of cytochrome P450 family 26 (Cyp26) \((52)\). The combination of low concentration and short half-life makes direct measurement of RA practically impossible, particularly in subregions of the retina (e.g., in regions of geographic atrophy). An alternative is indirect detection, for example, with a RAR reporter gene incorporating the RA response element (RARE) sequence to drive fluorescent protein expression. However, delivery of a nontherapeutic reporter gene is considered too invasive and risky for implementation in humans, at least at present. Unfortunately, nonhuman primates with RP have been found only very recently (two individual macaques) \((53)\), limiting direct validation of heightened RA signaling to other mammalian models.

These issues make our results with disulfiram particularly exciting and compelling. Because disulfiram is FDA-approved with safety that has been established over decades \((36)\), it should face low regulatory hurdles for trials in humans with RP or other similar disorders. Our results on vision-impaired mice show that disulfiram improves behavioral contrast sensitivity, sharpens cortical neuron representations of spatial orientation, and increases the fidelity of responses to naturalistic scenes, all consistent with improved visual perception. Whether disulfiram will improve vision in humans remains to be seen, but the barriers to answering this question seem relatively low. If disulfiram shows efficacy, then it could be administered orally, but local ocular delivery involving a new drug formulation might ultimately be more appropriate for avoiding the undesired systemic consequences associated with alcohol consumption. Disulfiram nonselectively inhibits all ALDH isozymes, but new drug candidates selectively targeting RALDH isoforms in the retina \((54)\) might also help remove these concerns.

Our results show that inhibiting RAR also improves vision. We used BMS 493, a pan–RAR inverse agonist that acts on RAR co-repressor protein \((33, 40)\). BMS 493 is potent and effective, but there is a rich pharmacopeia of other small-molecule agents that also interact with the RA binding site on RAR, the co-repressor or the co-activator, acting as agonists, antagonists, or inverse agonists. There are also agents that inhibit RXR, a protein that forms an obligate dimer with RAR before binding to DNA \((55)\). Determining which agent will have the most appropriate pharmacological properties for therapeutic vision rescue requires further studies of efficacy, toxicity, pharmacokinetics, and pharmacodynamics.

RAR expression or activity can also be inhibited by gene therapy. We have shown that a dominant-negative mutant of the RARα isoform \((RARP\alpha)\), delivered to RGCs with a targeted AAV vector, suppresses hyperactivity and augments light responses in rd10 mice \((17)\). AAV-mediated gene delivery has the advantage of allowing cell type targeting, which may benefit from the discovery of gene regulatory elements that may be expressed preferentially in retinal neurons that are most severely affected by RA-induced remodeling. However, in nonhuman primates, AAV-mediated gene delivery is limited by uneven penetration of the virus across the inner limiting membrane, restricting viral transduction to a ring of RGCs surrounding the fovea \((56)\). This scenario will presumably apply to humans as well. To avoid viral delivery limitations, other genetic tools may be appropriate for suppressing RAR expression, including
RNA-interfering molecules such as antisense oligonucleotides (ASOs) and small interfering RNAs (siRNAs). These molecules are much smaller than viruses, and after lipid modification, they can cross the inner limiting membrane and inhibit gene expression across the lateral extent of the retina (57, 58). ASOs and siRNAs are in clinical trials for other eye disorders, and advances in their design may generally benefit this strategy for suppressing RA signaling.

In principle, light responses could be unmasked and vision could be improved by directly inhibiting the ion channels that underlie hyperactivity. Gap junction proteins (59), P2X receptors (20), and hyperpolarization-activated cyclic nucleotide-gated (HCN) channels (11) have all been implicated in remodeling, making them potential drug targets for vision improvement. However, these channels are all normally expressed in the healthy retina, and it might be impossible to block them without affecting normal function. It is possible that RA is the common signal for up-regulating these channels, in which case, a single RA inhibitor might reduce their expression in a coordinated manner. RAR is largely inactive in adult RGCs (17), so an RAR inhibitor might reverse hyperactivity in degenerated regions of the retina without affecting normal function in regions with healthy photoreceptors.

Outer segments of rods and cones completely disappear in no-light perception patients with end-stage RP. However, at least some patients retain cone cell bodies that continue to express cone-specific genes, and these cells remain synaptically connected to bipolar cells (60). The hyperactivity of downstream RGCs may obscure small light responses produced by these remnant cones, raising the possibility that elevated RA might obscure perception. Thus, there is the possibility that RALDH or RAR inhibitors might awaken light perception in patients otherwise considered completely blind.

Because RAR is relatively inactive in healthy RGCs, RAR inhibitors should act preferentially on regions of the retina where degeneration has taken place, bypassing healthy retinal regions. RA-induced gene expression is not limited to inherited retinal degeneration; it also occurs in response to photoreceptor degeneration caused by photodamage (21) and local physical detachment from the RPE (23). This suggests that corruption of retinal processing may apply to many disorders that can cause loss of rods and cones, including retinal detachment, photodamage, and perhaps AMD.

Rescue of visual function by RAR may extend useful vision for months or years, but there is no evidence that it will slow photoreceptor degeneration itself. The strategy of rescuing vision with RA inhibitors is distinct from the strategy of restoring vision with retinal prosthetics, optogenetic and optopharmacological tools, or cell-based therapies. Vision restoration therapies are aimed, at least for now, at the small fraction of patients with end-stage photoreceptor degeneration, but treatments targeting the RA pathway may be relevant to the much larger patient population with low vision. Moreover, reducing RGC hyperactivity with RA inhibitors might be beneficial even after all the photoreceptors have degenerated and light perception is absent. Responses evoked by optoelectronic (61, 62), optogenetic (8, 63), or optopharmacological (64, 65) stimulation of the degenerated retina are superimposed on the heightened background activity of RGCs, curtailing the encoding of visual images. The combination of a light-sensitive actuator with an RAR inhibitor could have a synergistic effect, boosting neural signals to more effectively restore visual function to patients with no light perception.

Materials and Methods

Animals

C57/Black (WT) and rd10 (Pde6β−/− mutant) mice were purchased from the Jackson Laboratory (strain #00664 and #004297, respectively). All mice were kept in a 12:12 light/dark cycle room, except for experiments including dark rearing (see below). All animal procedures were approved by the Institutional Animal Care and User Committee at University of California (UC) Berkeley and UC Santa Barbara.

Chemicals and solutions

Dissections and ex vivo retinal assays (MEA and imaging) were performed in artificial cerebrospinal fluid (ACSF) containing 119 mM NaCl, 2.5 mM KCl, 1 mM KH₂PO₄, 1.3 mM MgCl₂, 2.5 mM CaCl₂, 26.2 mM NaHCO₃, and 20 mM d-glucose. Disulfiram (tetraethylthiuram disulfide, C₁₀H₂₀N₂S₄) was purchased from Sigma-Aldrich (#86720) and formulated in mouse chow by Dyets Inc. (PA, USA) at a final concentration of 2 mg/kg. Food was refreshed every 4 to 5 days during treatment. BSM 493 was purchased from Tocris (#3509) and stored at −20°C in a final concentration of 5 μM in phosphate-buffered saline (PBS).

Viruses

RAR reporter

To detect RAR-induced transcription, we used an AAV vector (Vigenes Biosciences, Maryland, USA) that included a cytomegalovirus (CMV) promoter upstream to the coding sequence for the RFP (“mStrawberry”), followed by a polyadenylate tail and a stop sequence (17, 23). A fragment containing three repetitions of the RARE sequence followed by the weak promoter SV40 was subcloned from pGL3-RARE-luciferase (Addgene, plasmid #13458), a gift of the Underhill Laboratory (66). Last, a GFP sequence was subcloned downstream to SV40 for a final construct of AAV2-CMV-RFP-stop-RARE(x3)-SV40-GFP. The presence of inverted terminal repeat sequences was confirmed by enzymatic digestion.

Genetically encoded calcium indicator

To measure activity-dependent increases in intracellular Ca²⁺ resulting from spike activity, we used a commercially available AAV vector encoding the Ca²⁺ indicator GCaMP6s under the control of the CaMKIIα promoter (Addgene, #107790, AAV9-CamKII-GCaMP6s-WPRE-SV40), which is expressed in excitatory neurons.

Intravitreal injections

Adult WT and rd10 mice (>P21) were intravitreally injected with drugs or virus. Before injection, animals were anesthetized with isoflurane (2%), and their pupils were dilated with tropicamide (1%) and phenylephrine (2.5%). Proparacaine (0.5%) was used as a topical analgesic. GenTeal was applied under a glass coverslip to keep the cornea lubricated. An incision was made through the sclera below the ora serrata with a 30-gauge needle. Solutions were injected into the vitreous with a blunt-ended 33-gauge Hamilton syringe or a 25- to 30-μm tip diameter glass pipette beveled at a 45° angle. After injection, the antibiotic tobramycin (0.3%) was applied to the eye. Injections of RAR reporter AAV2 were binocular, at a maximal volume of 1.5 μl, and using a titer of >10¹⁴ particles/μl. Injections of vehicle (PBS x1) and BMS 493 (5 μM) were binocular or monocular (depending on the experiment), using a final volume of 1.0 μl. Final BMS 493 concentration was 500 nM, assuming a 1:10 dilution in the vitreous volume of the mouse (~10 μl).
Retinal dissections
All dissections were performed in ACSF continuously gassed with 95% O₂/5% CO₂. Mice were euthanized using saturating levels of isoflurane gas followed by cervical dislocation. Mice were then enucleated, the cornea was excised from the eye, and the lens was removed. For live retinal imaging and MEA recordings, retinas were detached from the RPE and cut in quarters. Each retinal piece was mounted on nitrocellulose paper, RGCs facing up (imaging), or placed on top of the recording electrodes, RGCs facing down (MEA). Retinal pieces were kept at 34° to 35°C.

Cortical surgery
All surgeries were conducted with the mouse under isoflurane anesthesia (3.5% induction, 1.5 to 2.5% maintenance). Before incision, the scalp was infiltrated with lidocaine (5 mg/kg, subcutaneously) for analgesia. Meloxicam (2 mg/kg, subcutaneously) was administered preoperatively to reduce inflammation. Once anesthetized, the scalp overlying the dorsal skull was sanitized and removed. The periosteum was removed with a scalpel, and the skull was abraded with a drill burr to improve adhesion of dental acrylic.

For mice in the disulfiram experiments, a single 4-mm craniotomy was made over each bilateral visual cortex (centered at 4.0 mm posterior, 2.5 mm lateral to bregma on each side), leaving the dura intact. We used a motorized micro-injector (Stoelting, 53311) to deliver 0.75 µl of virus AAV9-CamKII-β injector (Stoelting, 53311) to deliver 0.75 µl of virus AAV9-CamKII-GaMP6s (titer = 1 × 10¹³ to 3 × 10¹³) to three to four injection sites within area V1 of the cerebral cortex. A cranial window was implanted over the craniotomy and sealed, first with a silicon elastomer (Kwik-Sil, World Precision Instruments) and then with a dental acrylic (C&B-Metabond, Parkell) mixed with black ink to reduce light transmission. The cranial windows were made of dental acrylic (0.5%) was used as a topical analgesic. To create and maintain contact between the lens containing the recording electrode and the cornea, a small drop of hypropromellose was placed on the eye, and the lens was gently pressed against it. After recording, GenTeal was applied to both eyes, and mice were allowed to recover from anesthesia.

Electroretinogram
Photopic ERG responses were recorded in adult mice with the HMxERG LAB System (OcuScience) under isoflurane anesthesia (3.5% induction, 1.5 to 2.5% maintenance) at a stable rectal temperature of >35.8°C. The ground electrode was placed on the back above the tail, while reference electrodes were placed in the chick under the cheek (all subdermal). Mouse pupils were dilated with tropicamide (1%) and phenylephrine (2.5%). Proparacaine (0.5%) was used as a topical analgesic. To create and maintain contact between the lens containing the recording electrode and the cornea, a small drop of hypropromellose was placed on the eye, and the lens was gently pressed against it. After recording, GenTeal was applied to both eyes, and mice were allowed to recover from anesthesia.

Visual behavioral assay
Cage design and manipulation
The automated behavioral cages were based on a design developed by E. Pugh (44) and custom-built by Lafayette Instrument (IN, USA). The cage includes three main components: (i) a unidirectional running wheel, orienting the mouse toward a computer display and a revolutions counter; (ii) a nose poke area with an infrared sensor, connected to a peristaltic pump triggering delivery of a water droplet reward; and (iii) a Raspberry Pi minicomputer and 1366-pixel by 768-pixel liquid crystal display (LCD) screens. Screens were calibrated to have the same brightness. Custom software linked image display with wheel running and reward availability. Cages were used in a dark room, and experiments were conducted throughout the night. The involvement of the animal handlers was minimal (introduction and retrieval), while protocol implementation and data acquisition were fully automated.
Behavioral experiment design and data analysis

Adult (>P35), male and female WT and rd10 mice, in littermate tandems, were used for these experiments. The cohort of rd10 mice used in disulfiram experiments (Fig. 4) was reared under 14:10 light/dark conditions. The cohort of rd10 mice used in BMS 493 experiments (Fig. 5) were dark-reared from P10 to P55 to P65 (depending on training and testing) and under 14:10 light/dark conditions afterward. Mice were water-deprived for >9 hours ahead of each experiment. Mice learned the task through operant conditioning. Training and testing sessions were conducted for 12 hours (6 p.m. to 6 a.m.). During the habituation protocol, the visual stimulus (a full-contrast grating with a 0.51 cycle per degree frequency) associated with reward availability is continuous, and the mouse can poke and receive 10% sucrose water ad libitum. During training, running on the wheel gates the visual stimulus, displayed on the screen. This is paired with unlocking of the nose poke, enabling the mouse to obtain the reward. The reward availability period and the duration of pump activation are shortened during training sessions to increase response stringency. During testing, stimuli have the same total luminance, but the contrast is randomly chosen from nine different possible steps, including 0, 1, 2, 4, 8, 16, 32, 64, and 100%.

In all sessions, latency was defined as the elapsed time between onset of the visual stimulus to nose poke. The mean latency to the 100% contrast stimulus during the first contrast sensitivity test was used as a reference to assess performance in response to all other contrasts and for all subsequent tests. A successful response was defined as one with mean latency falling within the ±2 SD of the latency during the reference test. Exclusion criterion during training is as follows: if during session 3 success rate was <80% and/or mean latency was >4 s. Exclusion criterion during testing is as follows: if SD > 40% of the mean (SD/mean > 0.4; see tables S5 and S6). Initial analysis was performed with custom software to extract the data from text files and arrange responses by contrast steps in datasheets. Analysis of success rate was performed manually.

Cortical imaging

Visual stimuli

All visual stimuli were generated with a Windows PC using MATLAB and the Psychophysics toolbox (67). Visual stimuli were presented on two LCD monitors that can display visual images to either eye independently. Each monitor (17.5 cm by 13 cm, 800 pixels by 600 pixels, 60-Hz refresh rate) was positioned symmetrically 5 cm from each eye at a 30° angle right of the midline, spanning 120° (azimuth) by 100° (elevation) of visual space. The monitors were located 3 cm above 0° elevation and tilted 20° downward. A non-reflective drape was placed over the inactive monitor to reduce reflections from the active monitor.

Orientation tuning was measured with drifting sine wave gratings (spatial frequency, 0.05 cycles/deg; temporal frequency, 2 Hz) presented in 1 of 12 directions, spanning from 0° to 330° in 30° increments. For a single repeat, each grating was presented once for 2 s with a luminance-matched 4-s intertrial gray screen between presentations. This was repeated for eight repetitions per session.

Natural scenes stimuli consisted of 900 frames from Touch of Evil (Orson Wells, Universal Pictures, 1958) presented at 30 frames/s, leading to a 30-s presentation time per repeat with a 5-s intertrial gray screen. The clip consists of a single continuous scene with no cuts, as has been previously described and is commonly used as a visual stimulus (https://observatory.brain-map.org/visualcoding/ stimulus/natural_movies). Each presentation was repeated 30 times, with a 5-s intertrial gray screen.

Two-photon imaging

After >2 weeks of recovery from surgery, GCaMP6s fluorescence was imaged using a Prairie Investigator two-photon microscopy system with a resonant galvo-scanning module (Bruker). For fluorescence excitation, we used a Ti:sapphire laser (Mai-Tai eHP, Newport) with dispersion compensation (DeepSee, Newport) tuned to λ = 920 nm. For collection, we used GaAsP photomultiplier tubes (Hamamatsu). To achieve a wide field of view, we used a 16×/0.8 numerical aperture microscope objective (Nikon) at ×1 (850 μm by 850 μm) or ×2 (425 μm by 425 μm) magnification. Laser power ranged from 40 to 75 mW at the sample depending on GCaMP6s expression levels. Photobleaching was minimal (<1%/min) for all laser powers used. A custom stainless steel light blocker (eMachineShop.com) was mounted to the head plate and interlocked with a tube around the objective to prevent light from the visual stimulus monitor from reaching the photomultiplier tubes. During imaging experiments, the polypropylene tube supporting the mouse was suspended from the behavior platform with high-tension springs (Small Parts) to reduce movement artifacts.

Two-photon post-processing

Images were acquired using the Prairie View acquisition software and converted into TIF files. All subsequent analyses were performed in MATLAB (MathWorks) using a custom code (https://labs.mcdb.ucsb.edu/goard/michael/content/resources). First, images were corrected for X-Y movement by registration to a reference image (the pixel-wise mean of all frames) using 2D cross-correlation.

To identify responsive neural somata, a pixelwise activity map was calculated using a modified kurtois measure. Neuron cell bodies were identified using local adaptive threshold and iterative segmentation. Automatically defined ROIs were then manually checked for proper segmentation in a graphical user interface (allowing comparison to raw fluorescence and activity map images). To ensure that the response of individual neurons was not due to local neuropil contamination of somatic signals, a corrected fluorescence measure was estimated according to

\[
F_{\text{corrected}}(n) = F_{\text{soma}}(n) - \alpha (F_{\text{neuropil}}(n) - \bar{F}_{\text{neuropil}})
\]

where \(F_{\text{neuropil}}\) was defined as the fluorescence in the region <30 μm from the ROI border (excluding other ROIs) for frame \(n\), and \(\alpha\) was chosen from [0 1] to minimize the Pearson’s correlation coefficient between \(F_{\text{corrected}}\) and \(F_{\text{neuropil}}\). The ∆F/Φ for each neuron was then calculated as

\[
\Delta F/Φ(n) = \frac{F_n - F_0}{F_0}
\]

where \(F_n\) is the corrected fluorescence (\(F_{\text{corrected}}\)) for frame \(n\) and \(F_0\) defined as the first mode of the corrected fluorescence density distribution across the entire time series.

Analysis of two-photon imaging data

Blinding to experimental condition

For disulfiram experiments, disulfiram-containing chow or control chow of the same composition (Dyets Inc.) was given neutral codes by an investigator not involved in the study and administered to the mice with the experimenter blind to experimental condition. For BMS 493 experiments, vials of drug and vehicle were given neutral...
codes, and each solution was used for one eye. In both cases, the experimental condition was revealed only after the primary analysis was complete.

**Orientation tuning**

Neural responses to the orientation tuning stimulus were first separated into trials, each containing the response of the neuron across all tested queried orientations. For each neuron, we averaged the baseline-subtracted responses to each orientation, creating an orientation tuning curve for each trial. To calculate the OSI in a cross-validated manner, we first separated the orientation tuning curves into even and odd trials. We then aligned the even trials using the maximal response of the averaged odd trial for each neuron and vice versa, resulting in aligned responses. We then calculated the OSI from the averaged tuning curves for each neuron using the following equation

$$\text{OSI} = \frac{R_{\text{pref}} - R_{\text{pref}+\pi}}{R_{\text{pref}} + R_{\text{pref}+\pi}}$$

where $R_{\text{pref}}$ is the neuron’s average response at its preferred orientation, defined by cross-validation on a different set of trials using the above procedure. Aligning the orientation tuning curves of the neurons using cross-validation provides a more accurate measurement of the orientation tuning of the neuron, as it prevents nonselective neurons from having high OSI values due to spurious neural activity.

**Naturalistic movies reliability**

Neural responses to the naturalistic movies were first separated into trials, with each trial containing the full response of the neuron to the entire presented movie. The reliability of each neuron to the naturalistic movie was calculated as follows

$$R_c = \frac{\sum_{t=1}^{T} \text{CC}(r_{c,t}, \bar{t}_{c,[1:T]}|\pi)}{T}$$

where $R_c$ is the reliability for cell $c$, $t$ is the trial number from [1, $T$], CC is the Pearson correlation coefficient, $r_{c,t}$ is the response of cell $c$ on trial $t$, and $\bar{t}_{c,[1:T]}|\pi$ is the average response of cell $c$ on all trials excluding trial $t$. To separate neurons by reliability deciles, we independently calculated the decile cutoffs for each condition and then binned neurons into their respective deciles.

**Naturalistic movies decoding analyses**

To decode naturalistic movie responses from the population data, we first randomly selected neurons of a given pool size (pool sizes: 2, 4, 8, 16, 32, 64, 128, 256, or 512 neurons) from across all recordings. The neural responses to natural movies within that pool were then divided into even and odd trials. The average population activity across even trials was used to calculate a “template” population vector for each frame of the movie. We then estimated the movie frame ($F_{\text{Decoded}}$) from the population activity of each actual frame ($F_{\text{Actual}}$). To accomplish this, we calculated the population vector from the odd trials during $F_{\text{Actual}}$ and compared to the template population vectors (even trials) for all of the frames. The frame with the smallest Euclidean distance between population vectors was chosen as the decoded frame ($F_{\text{Decoded}}$). This process was repeated for each frame ($F_{\text{Actual}}$) of the movie. For each pool size of neurons used, the entire procedure was iterated 1000 times, picking new neurons for each iteration. This resulted in a confusion matrix that describes the similarity of neural activity patterns for each frame between nonoverlapping trials. To assess decoder performance, we measured the percentage of decoded frames that fell within 10 frames of the actual frame (chance level = 7%).

**Statistical analysis**

For data shown in Figs. 1, 2, and 6 to 8, we used gLMEMs to account for the individual differences between mice in our statistical analyses. The formula for these models is as follows

$$y = X\beta + Zb + \epsilon$$

where $y$ is the response vector, $X$ is the fixed-effects design matrix (denoting treatment condition), $\beta$ is the fixed effects vector, $Z$ is the random-effects design matrix (denoting different mice), $b$ is the random effects vector, and $\epsilon$ is the observation error vector. After fitting the models for each experiment, we performed $F$ tests on the appropriate contrasts to determine significance. For data presented in Fig. 2 (D and H), significance between cumulative probabilities was tested using the Kolmogorov-Smirnov test. For data presented in Figs. 3 to 5 and figs. S1 and S3, comparison between groups used nonparametric tests (Mann-Whitney $U$ test for independent samples or Wilcoxon sign test for paired data), unless the data passed the normality test (Shapiro-Wilk) and could be analyzed with parametric tests (two-tailed $t$ test).

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.sciencemag.org/content/11/10.T2/SI/siadv.abm4643

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