DREADDs in *Drosophila*: A Pharmacogenetic Approach for Controlling Behavior, Neuronal Signaling, and Physiology in the Fly

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SUMMARY

We have translated a powerful genetic tool, designer receptors exclusively activated by designer drugs (DREADDs), from mammalian systems to *Drosophila melanogaster* to selectively, rapidly, reversibly, and dose-dependently control behaviors and physiological processes in the fly. DREADDs are muscarinic acetylcholine G protein-coupled receptors evolved for loss of affinity to acetylcholine and for the ability to be fully activated by an otherwise biologically inert chemical, clozapine-N-oxide. We demonstrate its ability to control a variety of behaviors and processes in larvae and adults, including heart rate, sensory processing, diurnal behavior, learning and memory, and courtship. The advantages of this particular technology include the dose-responsive control of behaviors, the lack of a need for specialized equipment, and the capacity to remotely control signaling in essentially all neuronal and nonneuronal fly tissues.

INTRODUCTION

Pharmacological tools have traditionally been a primary method used to modify neural signal transduction and function in mammalian systems. Although these methods are generally quite successful, certain limitations associated with pharmacological strategies include frequent deleterious off-target effects (Keiser et al., 2009). In the model organism *Drosophila melanogaster*, approaches taking advantage of the extensive genetic toolkit to elucidate the neuronal basis of behavior have been primarily utilized instead of small-molecule-based pharmacological tools. The bipartite GAL4/UAS expression system (Brand and Perrimon, 1993) and its modifications to optimize both spatial and temporal control of gene expression have been the primary workhorse. More recent methods of control of transgene expression used in combination with the GAL4/UAS system include the use of FLP recombinase (Keller et al., 2002; Struhl and Basler, 1993) and the use of hormones or drugs to induce expression of UAS transgenes (e.g., GeneSwitch) (Osterwalder et al., 2001; Roman et al., 2001). Advantages of these newer modifications include increased control of spatial and/or temporal expression and reversibility of expression through the addition or removal of a stimulus like the hormone RU486. Although greater control of targeted expression of transgenes has allowed for an enhanced understanding of the neuronal circuits in an intact organism, disadvantages to these systems include developmental and physiological “off-target” effects associated with constitutively expressed transgenes and slow on/off rates for hormonal induction systems (e.g., several days).

Recently developed methods target expression of proteins that, when activated, inhibit or enhance neuronal activity and circuitry function. Some strategies include using constitutively active sodium channels (NaChBac) or potassium channels (Kir2.1) to excite or hyperpolarize neurons, respectively (Nita-bach et al., 2002, 2006). Conditional activation of neurons can also be achieved with dTRPA1, a temperature- and voltage-gated cation channel that regulates thermotactic behavior (Hamada et al., 2008; Parisky et al., 2008; Shang et al., 2008). The most common conditional method to silence neurons in the fly is to inhibit synaptic release using a temperature-sensitive dominant-negative allele of dynamin (shibire*). Using this system, shifting the fly to the nonpermissive temperature (>-29 °C) leads to loss of neurotransmitter vesicular recycling and a rapid inhibition of neurotransmitter release (Kitamoto, 2001). Another valuable tool for inducing activity in neuronal tissue is a light-sensitive cation channel, Channelrhodopsin-2 (ChR2) (Nagel et al., 2003). Exposure to intense blue light activates these channels and leads to depolarization and neuronal firing where expressed; therefore, activity can be precisely controlled at the temporal level (Boyd et al., 2005; Li et al., 2005; Nagel et al., 2003, 2008).
Alter the duration or intensity of the light pulse can vary the response of the channels (Zhang et al., 2007). Although ChR2 is used successfully to manipulate neural circuit activity in Drosophila embryos and adults (Schroll et al., 2006; Zhang et al., 2007), limitations of the “optogenetic” approach include poor penetration of blue light into whole organisms (Eichler et al., 2007), limitations of the “optogenetic” approach include poor penetration of blue light into whole organisms (Eichler et al., 2007). Sensitization issues, and a narrow dose-response control. A significant disadvantage of optogenetics is the need for specialized and expensive equipment like custom blue light sources with fiber optics or two-photon illumination. These limitations, taken together, contribute to the inadequacy of optogenetics for types of studies that require long-term or chronic modulation of neuronal activity in Drosophila. A significant limitation of each of these channel-based methods, as well as that of shibire<sup>12</sup> for neuronal control, is that they are essentially unidirectional switches and primarily either activate or inactivate neurons in an “all on” or “all off” manner. Additionally, such channel-based methods do not regulate G protein signaling.

Powerful new tools combining genetics and pharmacology that examine physiological functions and behaviors associated with specific neuronal circuits have recently been developed for mammalian systems that overcome many of the limitations of other methods. These systems make use of modified G protein-coupled receptors (GPCRs) that respond to synthetic ligands that the wild-type receptor does not respond to and have been termed receptors activated solely by synthetic ligands (RASSLs) and designer receptors exclusively activated by a designer drug (DREADDs) (Armbruster et al., 2007; Coward et al., 1998; Nichols and Roth, 2009). While the RASSLs represent GPCRs that have been rationally designed or found in nature, and often still respond to the native ligand, the DREADDs were created through directed evolution to respond to a synthetic ligand but to have no affinity for its native ligand. The current set of DREADDs largely consist of mammalian muscarinic acetylcholine receptors that no longer have affinity for, and therefore have no response to their natural ligand, acetylcholine, and are fully and potently activated by the synthetic ligand clozapine-N-oxide (CNO), which is inert at all known mammalian GPCRs (Armbruster et al., 2007). In mammalian systems, DREADDs have produced conditional control of neuronal function through manipulation of effector pathways that is rapid, reversible, and does not require specialized equipment (Alexander et al., 2009; Krashes et al., 2011; Ray et al., 2011; Sasaki et al., 2011). Activating ligand can be easily and conveniently administered by simply feeding or injecting the drug to the animal. Importantly, and unlike all channel-based methods, DREADDs reliably control neuronal and effector pathway function in a true dose-dependent fashion.

We have adapted and translated the DREADD technology to Drosophila and show that the mammalian DREADD receptors effectively couple to Drosophila G protein effector pathways and alter behaviors and physiological function in both larvae and adults when activated by CNO. These effects are rapid, reversible, and dose responsive. We thereby demonstrate the utility of the DREADD technology for conditional control of signaling, behaviors, and physiological processes in the fly.

## RESULTS

### CNO Is Biologically Inert in the Fly

To determine if CNO had any effects on normal flies, we grew wild-type Canton-S (CS) flies on food containing 3 mM CNO and observed no gross overt adverse developmental consequences, developmental delays, or effects on lifespan (Figure S1). To test the effects of CNO on overt fly behavior, we tested larval activity and feeding, as well as adult activity. No significant differences in either the number of mouth hook contractions (a measure of feeding behavior) or body wall contractions were observed between the control and larvae fed either 3.0 mM CNO (Figure 1). To examine overt activity levels in adult flies, we used the Drosophila activity monitoring system (DAMS) photobeam break counting system. Adult CS flies fed with food containing 10 mM CNO for 5 days did not exhibit a significant change in activity when compared to flies not fed CNO (Figure 1C).

### DREADDs Couple to Drosophila Effector Pathways and Can Alter Neuronal Properties

To verify that the mammalian DREADDs effectively couple to insect G proteins, we measured the abilities of the G<sub>A</sub>-coupled and G<sub>A</sub>-coupled DREADDs to modulate cyclic AMP (cAMP) production in Drosophila S2 cells when stimulated with CNO.
We observed that stimulation of the $G_{\alpha i}$-coupled hM4Di receptor with CNO negatively modulates cAMP levels in a dose-dependent manner and that the CNO stimulated $G_{\alpha s}$-coupled rM3BDs receptor dose-dependently increases cAMP (Figures 2A and 2B).

To examine the effects of activation of the hM1Dq receptor on neuronal properties in vivo, we coexpressed hM1Dq receptor along with a live cell fluorescent calcium sensor, GCaMP (Tian et al., 2009), in neurons of the third-instar larval brain using the pan-neuronal elavC155-GAL4 driver. The response to a depolarizing amount of KCl (75 mM) applied to the brain was measured in neurons of the ventral ganglion that were pretreated for 5 min with either ringers or ringers + CNO (6 μM). In fly brains carrying one copy of all three transgenes (elavC155-GAL4 + UAS-hM1Dq + UAS-GCaMP3), we observed that pretreatment with CNO resulted in a more rapid increase in the fluorescence intensity of neurons and in a reduced latency to the maximum peak intensity (Figures 3B and 3D). CNO had no effect on neuronal firing or maximum peak latency in control flies expressing only GCaMP by the elavc155-GAL4 driver (Figures 3A and 3B) or when added directly to the brain instead of KCl (Figure 3C). Together, these data are consistent with activation of the hM1Dq receptor facilitating neuronal depolarization, as has been previously observed in mammalian systems.

**DREADD Activation Dose-Dependently Modulates Olfactory Response in Larva**

Each DREADD receptor was expressed in larval sensory neurons under the control of the SG18.1-GAL4 (6405-GAL4) driver (Sarpal et al., 2003). Next, their response to the attractant odor, ethyl acetate, was measured. In adults and larvae, attraction to ethyl acetate has been shown to involve $G_{\alpha q}$ signaling (Kain et al., 2009). Activation of hM1Dq expressed in sensory neurons with CNO produced a dose-dependent increase in the performance index (Figure 4B). CNO activation of sensory-neuron-expressed hM4Di receptor produced a dose-dependent decrease in the preference for ethyl acetate (Figure 4C). There was no effect of CNO on any of the parental lines or for the rM3BDs × 6405-GAL4 F1 larva for the preference to ethyl acetate (Figure 4A). The maximal effective concentration in the food was approximately 1.0 mM CNO. Our previous studies with CNS active drugs at GPCRs have demonstrated robust behavioral activity in the low-millimolar range. Therefore, we chose to perform our subsequent experiments with low-millimolar CNO concentrations in the food.

**$G_{\alpha q}$ Activation in Larval Heart Disrupts Heart Rate**

To express hM4Di in the heart, the 24B-GAL4 driver was used. This driver is expressed in all larval somatic muscle, including cardiac (Schuster et al., 1996). In nearly all larvae preparations where hM4Di was expressed in heart muscle, the heart dramatically slowed immediately after exposure to 500 nM CNO (Table 1; Movie S1). Each time the heart was revitalized when the CNO was washed out with fresh HL3 saline (pH 7.1). Flies that did not express hM4Di (but containing the UAS-hM4Di element) from the same cross (control sibling progeny that lacked the 24B-GAL4 element) showed no effect upon exposure to CNO (Table 1). Although 24B-GAL4 expresses in skeletal muscles, they have no impact on heart rate in dissected preparations because the heart is directly visualized contracting and the segmental roots to skeletal muscles from the CNS have been transected.

**$G_{\alpha s}$ Activation in Cry-GAL4 Neurons Affects Diurnal Behavior in Adults**

The neurons defined by the cryptochrome (cry) GAL4 driver define many cells of the circadian clock (Im et al., 2011). An important component of the cry circuitry is the PDF receptor, which is a $G_{\alpha s}$-coupled receptor (Im and Taghert, 2010). Therefore, manipulating cAMP levels through $G_{\alpha s}$ activation will likely influence certain circadian behaviors. UAS-rM3BDs × cry-GAL4 F1 flies fed 3.0 mM CNO had decreased early night activity levels compared to the same genotype flies not fed CNO (Figure 5C). There were no observable differences in activity levels in either parental strains fed CNO (Figures 5A and 5B).
Stimulation of hM4Di in the Mushroom Body Disrupts Short-Term Learning and Memory

The mushroom body (MB) has been shown to be necessary for proper short-term memory (STM) function (Zars et al., 2000). Furthermore, cAMP levels in the MBs are crucial for olfactory learning and memory (Davis et al., 1995). Disruption of cAMP levels in the MBs with DREADDs is therefore predicted to negatively influence learning and memory. To determine if we could use the DREADD system to probe MB circuitry function, we expressed the hM4Di receptor in a subset of MB lobes using the MB247-GAL4 driver, activated them with CNO, and measured the resulting performance. The UAS-hM4Di MB247-GAL4 F1 flies fed CNO (1.0 mM) prior to training exhibited a 50% decrease in the performance index compared to the non-CNO-fed F1 flies.

Figure 3. Activation of Mammalian hM1Dq Receptor Facilitates Neuronal Depolarization

(A–D) The hM1Dq receptor was expressed in flies carrying the elav<sup>1155</sup>-GAL4 driver and the UAS-GcAMP. Brains from third-instar larvae were stimulated with KCl (75 mM), either in the presence or absence of CNO (6 μM). GCaMP fluorescence response was measured and the maximum peak latency was determined.

(A) In neurons only expressing GCaMP (elav<sup>1155</sup>-GAL4 + UAS-GCaMP), pretreatment with CNO had no effect on KCl-induced depolarization.

(B) Maximum peak latency was unaffected by pretreatment of CNO in larvae only expressing GCaMP (elav<sup>1155</sup>-GAL4 + UAS-GCaMP).

(C) In neurons expressing both GCaMP and hM1Dq (elav<sup>1155</sup>-GAL4 + UAS-GCaMP + hM1Dq), pretreatment with CNO resulted in a decreased latency to firing.

(D) Maximum peak latency was significantly reduced in CNO-pretreated brains expressing hM1Dq (elav<sup>1155</sup>-GAL4 + UAS-GCaMP + hM1Dq). *p < 0.005, Student’s t test. N = 15 brains per treatment, ten neurons per brain. Errors bars indicate SEM.
We observed no difference in the performance of any of the parental strains fed CNO (Figure 6A).

The Behavioral Effects of DREADDActivation Are Reversible
To determine if DREADD activation is reversible, we used the UAS-hM4Di MB247-GAL4 F1 flies in additional short-term learning and memory assays. The flies were removed from the CNO at T = 0, 3, and 12 hr prior to training. When the flies were immediately trained (T = 0) and tested, we observed the expected 50% decrease in the PI (Figure 6B). When flies were removed from the drug and placed on food without CNO for 3 hr prior to training, the PI was decreased only by 12% (Figure 6B). Flies removed from the CNO and placed on food without drug for 12 hr prior to training exhibited no statistically significant difference in short-term olfactory learning and memory performance compared to flies not fed CNO (Figure 6B). Based upon these results, the half-life for the reversibility of the CNO/DREADD activation, at least for this behavior, is approximately 90 min.

hM4Di Activation of 5-HT7Dro-GAL4 Neurons Modulates Behaviors
Our previous studies have indicated that the 5-HT7Dro receptor, as well as the circuitry defined by this driver, is involved in courtship and mating (Becnel et al., 2011). To further validate the role of 5-HT7Dro receptor circuitry in this behavior, we expressed the hM4Di receptor under the control of the 5-HT7Dro-GAL4 driver and assessed the effects of CNO activation on courtship. When fed CNO, the UAS-hM4Di x 5-HT7Dro-GAL4 F1 flies exhibited a decrease in the courtship index (Figure 7). These results are consistent with our previous methods utilizing receptor antagonists and RNAi methods (Becnel et al., 2011).

DISCUSSION
The major finding of this paper is that the DREADD system allows for reversible, spatiotemporal control of signaling and behavior as well as physiological processes in Drosophila melanogaster. DREADD technology overcomes many limitations of other methods, including the need for expensive specialized equipment, dedicated temperature environments, and the inability to easily titrate signaling and induced behaviors in a dose-responsive manner. As demonstrated, flies of the appropriate genotype can be simply fed CNO, either acutely or chronically, to modulate signaling and downstream behaviors. Significantly, this system not only examines the role of signal transduction effector pathways in cells, but could conceivably silence and facilitate the electrical activity of neurons in the traditional sense. For example, we observed that activation of the Gαq-coupled hM1Dq DREADD facilitates depolarization of neurons. This is possibly due to PIP2 gating of inwardly rectifying potassium channels (Alexander et al., 2009). In developing the DREADD technology, clozapine-N-oxide was chosen as the ligand to evolve the receptors around because it demonstrates no appreciable affinity to any known mammalian GPCR or enzyme (Armbruster et al., 2007). We found no overt behavioral or developmental effects of CNO in flies indicating that, as in mammals, CNO is likely biologically inert in the fly.

Another issue to address in developing this system was the coupling of mammalian GPCRs to insect effector pathways. Evidence in the literature examining this question is sparse and...
mixed with respect to heterologous coupling. A number of reports, however, demonstrate positive coupling (Han et al., 1996; Kemp and Manahan-Vaughan, 2005; Perret et al., 2003; Saudou et al., 1992; Wei et al., 2000). Mammalian muscarinic acetylcholine receptors expressed in insect cells have been previously shown to fully couple to appropriate insect heterotrimeric G proteins to activate canonical effector pathways (Knight et al., 2003). We cloned the DREADD receptors into insect expression vectors and tested them in a Drosophila S2 cell culture expression system and found that they correctly couple to adenylate cyclase with the rM3BDs increasing, and the hM4Di decreasing, cAMP production in a dose-dependent manner upon activation of G\textsubscript{i}-coupled GPCRs in mammalian heart. Dramatically, upon washout of CNO, most of the hearts began to beat again. We anticipate that other physiological processes regulated by intracellular signaling pathways in nonneuronal tissues can also be manipulated with the DREADD system.

In the adult, we examined diurnal behavior, and olfactory learning and memory. For the diurnal tests, we used a cryptochrome (cry)-GAL4 driver to drive DREADD expression. There is significant overlap with cry expression and the G\textsubscript{\alpha}i-coupled PDF receptor (PDFR) (Im et al., 2011), which is a key regulatory component of the circadian clock (Im and Taghert, 2010). Flies mutant for the PDFR primarily have defects in the evening peak, with a slight forward shift and increased general activity in the afternoon (Im and Taghert, 2010). We observed that activation of rM3BDs under expression control of a cry-GAL4 driver led to a decrease in evening peak activity, without affecting the morning peak. Because PDFR mutants demonstrate a general increase in afternoon activity, an increase in PDFR activity in cry circuitry may be predicted to decrease afternoon activity, which is consistent with our results. Our system may therefore prove useful in experiments to further elucidate the role of specific neurons and circuits in the circadian clock and the role of effectors including cAMP within these neurons in experiments utilizing circadian relevant GAL4 drivers and doses of CNO.

For adult olfactory learning and memory performance, we tested if we could negatively control the performance index by expressing and activating the hM4Di receptor under the control of the MB247-GAL4 driver. The MB247-GAL4 driver expresses in a subset of the mushroom body lobes (Aso et al., 2009), and is a commonly used GAL4 driver in the study of olfactory learning and memory. When we fed flies expressing hM4Di in the MBs CNO prior to training and testing, we significantly reduced the performance index for short-term learning and memory. Neither performance index for short-term learning and memory. Neither parental strains exhibited an effect of CNO. It is not possible at this point to determine if the effect was due to the silencing properties of receptor activation through G\textsubscript{\alpha}i-coupled opening of inwardly rectifying potassium channels and hyperpolarization of the neurons or a G\textsubscript{\alpha}i-mediated decrease in cAMP, which is a key molecule involved in learning and memory (Davis et al., 1995). We hypothesize that the inhibition is likely a combination of the two mechanisms. Regardless of the exact mechanism, we demonstrate here the ability of the hM4Di receptor to disrupt behavior in the adult. In any conditional activation system, it is crucial to determine if the effects are reversible or not, as well as the kinetics of the reversibility. We examined this

<p>| Table 1. Activation of hM4D in Larval Heart Disrupts Heart Rate |
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Mean ± SEM 77.8 ± 10.2 27.8 ± 12.9 | 80.3 ± 11.8 72.8 ± 14.4

The UAS-hM4Di was crossed to the 24B-GAL4 driver line. “24B” is heterozygous for the 24B-Gal4 transgene and homozygous for the UAS-GFP transgene. CNO (500 nM) was applied to the hearts of partially dissected live third-instar larvae and the effects directly visualized. The application of CNO rapidly and dramatically slowed the heart in nearly all larvae. Values represent heartbeats per minute from the same larvae before (saline) and after CNO application (CNO). hM4Di = 24B-GAL4+/−; UAS-mCD8:GFP +/−; UAS-hM4D +/−. Control (siblings from the same F1 cross not carrying the 24B-Gal4) = UAS-mCD8:GFP +/−; UAS-hM4D +/−.

*p < 0.01 saline versus CNO; ANOVA with Bonferroni post hoc test for multiple comparison.
utilizing the MB247-GAL4 + UAS-hM4Di fly in olfactory short-term learning and memory experiments. We observed a time-dependent recovery to a normal performance index with a predicted half-time of ~90 min after removal from food containing CNO prior to training. Although not as rapid as methods involving channels, this system is inducible on the scale of minutes and reversible on the scale of minutes to hours for modulation of behaviors. For many types of studies, these kinetics are more than adequate.

Based upon our results with these particular DREADD strains, the practical effective dose appears to be between 1 and 3 mM CNO in the food. Flies can be fed either acutely to produce immediate behaviors for observation or chronically to observe behaviors over an extended length of time. We routinely maintain adult flies on food containing CNO for 48 hr prior to a particular acute behavioral test in order to allow for steady-state accumulation of drug levels. Our results indicate a half-life of ~90 min after removal from food. This treatment regimen may result in some degree of receptor desensitization and downregulation; nevertheless, they effectively alter behaviors when activated by ligand. Flies can also be starved overnight and fed a large bolus dose and tested individually for acute behaviors. In our previous experiments, we have observed that CNS active drugs in acute administration experiments begin to have effects ~15 min, with a maximal effect ~20 min that persists at full strength until ~60 min (Nichols et al., 2002).

The elucidation of the function of unknown neural circuits and tissues defined by GAL4 drivers is a major utility we envision for this system. Although other methods employing channels are used for this purpose, they are more of a “switch” approach and maximally activate/inactivate neuronal circuits. Our system allows for a more graded analysis by gradually altering neuronal signaling through dose response experiments, which may uncover and distinguish more subtle, but distinct, behaviors mediated by the same neurons and circuits. To address this, we used our system to investigate the circuitry defined by the 5-HT7Dro-GAL4 driver, which demonstrates high expression in large field R-neurons innervating the ellipsoid body (EB) (Becnel et al., 2011). We have previously reported that the neurons defined by this driver, and the 5-HT7Dro receptor itself, are involved in courtship and mating behaviors (Becnel et al., 2011), as well as in olfactory learning and memory (Johnson et al., 2011). Previous studies by others utilizing shibire® expressed in a subset of large-field R-neurons have shown that when the neurons innervating the ellipsoid body are completely inactivated, severe locomotor and coordination deficits are produced (Krashes and Waddell, 2008), precluding an accurate analysis of the role of the EB in other behaviors dependent on normal activity and coordination. In agreement with our previous studies, activation of hM4Di expressed in 5-HT7Dro neurons dose-dependently inhibited courtship and mating, with no observable overt deficits in locomotor or coordination abilities. These results indicate that the DREADD system can indeed be used to examine more subtle behaviors otherwise masked by all-on or all-off approaches.

In summary, we have successfully translated a method for inducible and reversible remote control of behaviors and physiological processes to the fly utilizing a combination of genetics and pharmacology (e.g., “pharmacogenetics”). Perhaps the most important attribute of this system is its ability control behaviors and physiology in a dose-dependent manner, which allows for more precise and subtle examination of neuronal function and behaviors as well as physiological processes. Further, no specialized equipment is necessary; one simply feeds CNO to the fly. Due to the ubiquitous nature of GPCRs, we anticipate that this system will also be useful in the examination of the role of signal transduction pathway effectors in almost every tissue of the fly for which there is an available GAL4 driver.
The effects of DREADD activation are reversible. Flies expressing hM4Di in the mushroom bodies (MB247-GAL4 + UAS-hM4Di) were again monitored for short-term memory, this time after being removed from the CNO. Flies immediately trained and tested exhibited the expected 50% decrease in performance index (dark gray). When flies were removed from the CNO for 3 hr, the decrease in PI was only 12% (gray). Flies that were removed from the CNO for 12 hr prior to training and testing did not exhibit a decrease in PI (light gray) when compared to F1 flies that were not fed CNO (white). *p < 0.05, ANOVA with Bonferroni post hoc test for multiple comparison. N = 8 trials. Error bars indicate SEM.

EXPERIMENTAL PROCEDURES

Chemicals
General chemicals were obtained from Sigma. The odors 3-octanol and 4-methylcyclohexanol were obtained from Sigma. Ethyl acetate was from Fisher Scientific. Clozapine-N-oxide (CNO) was generously synthesized and provided by Dr. David E. Nichols at Purdue University from clozapine purchased from Enzo Life Sciences.

Drosophila Strains and Rearing
Canton-S (CS), white1118, the cry-GAL4 and the 6405-GAL4 drivers, and the UAS-GCaMP (chr3) strain were obtained from the Bloomington Stock Center. The 24B-GAL4 driver and Balancer lines (+/+; Ady1CyO;+/+) and [(+/+; +/+; Sco/CyO)] were provided by Dr. Udal Pandey (LSU Health Sciences Center, New Orleans, LA). The MB247-GAL4 strain was generously provided by Dr. Kyung-An Han (UTEP, El Paso, TX). The eya155-GAL4 strain was provided by Dr. Chunlai Wu (LSU Health Sciences Center, New Orleans, LA). The creation of the 5-HT;Dro-GAL4 transgenic line was previously described (Becnel et al., 2011). For routine maintenance, flies were reared on standard cornmeal-molasses food at 25°C under 12 hr light/dark conditions, unless otherwise stated.

Creation of UAS-DREADDs
The UAS-hm1Dq, UAS-hM4Di, and UAS-M3BDs fly strains were created, and inducibility of expression tested, as described in the Extended Experimental Procedures and as shown in Figure S2.

Generation of the pMT/V5-DREADDs
The hM4Di and rM3BDs complementary DNAs (cDNAs) were excised from the UAS-DREADD plasmids using KpnI and Xhol restriction enzymes (Promega). The pMT/V5-His vector from the DES-Inducible Kit (Invitrogen) was digested using the same enzymes. The hM4Di cDNA was ligated into the pMT-V5-His vector using the Fast Link DNA ligation kit following the manufacturer’s instructions. To generate the rM3BDs expression vector, both the excised rM3BDs fragment and the vector were blunt-ended using the End-IT DNA End Repair Kit (Epicenter) following the manufacturer’s directions and subsequently ligated together using the Fast Link DNA ligation kit following the manufacturer’s instructions. The final constructs were verified using a panel of restriction enzymes.

Culture of S2 Cells
Drosophila S2 Cells (Invitrogen) were cultured in Schneider’s Drosophila medium (Invitrogen) at 28°C in a nonhumidified, ambient-air-regulated incubator.
Larval Behavior Assays

Larvae Collection

To obtain larva, 8 oz bottles were set up with 10–15 virgin females of one parental strain and 10–15 males of the other parental line and incubated at 25°C. Flies were allowed to mate and lay eggs for 24 hr before being removed from the bottle. When the majority of the larvae in the bottle had reached the third-instar stage (day 6), the bottle was flooded with 50 ml of a 20% sucrose solution in water (w/v), causing the larvae to float to the surface. Larvae were collected, transferred to a small mesh basket, and washed twice with deionized water before further use (Nichols et al., 2012).

Chemosensory Assay

Fifty larvae were collected as described above and transferred to a small beaker containing 1:2 ml of 5% sucrose (either alone or with the appropriate concentration of CNO) for 15 min. The larvae were transferred to a Petri dish containing a 1% agarose gel with two 1 cm discs of Whatman filter paper directly opposite one another and evenly spaced from the center. One disc was spotted with 20 μl of deionized water as a control, and the other was spotted with 20 μl of ethyl acetate (1:104 dilution). The larvae were placed in the center of the plate using an overturned lid of a 35 mM dish. Once larvae were transferred to the plate, the lid was removed, the Petri dish covered, and the larvae were allowed to roam freely for 3 min, at which time the number of larvae on either side of the dish were recorded. Larvae on the half of the dish with the odorant disc subtracted by the number or larvae in the half of the dish containing the water disc, divided by the total number of larvae.

Heart Rate

The UAS-hM4Di transgenic line was crossed to a heterozygous 24B-GAL4 (24B-GAL4 +/-, UAS-mCD8:GFP +/-) transgenic line. Half of the resulting progeny will carry all three of the transgenes (24B-GAL4 +/-; UAS-mCD8:GFP +/-; UAS-hM4Di +/-). Experimental controls were siblings from the same cross that did not carry the 24B-GAL4 (UAS-mCD8:GFP +/-; UAS-hM4Di +/-). Heart rate measures in larval Drosophila are readily obtained within the whole animal or in an in situ preparation. Detailed procedures are described and illustrated in Cooper et al. (2009). In this study, we used early third-instar larvae and exposed the larval heart tube by dissecting the larvae on the ventral longitudinal axis. Dissection time takes about 3–6 min. During the dissection and for observations the preparation is bathed in HL3 saline (pH 7.1). This saline was a good media for maintaining synaptic transmission at neuromuscular junctions (Stewart et al., 1994). For heart measures, pH was carefully controlled (pH 7.1) in the HL3 saline and for the HL3 saline containing CNO. Each larva was recorded by video microscopy for the duration of an experiment. Heart rate was first observed under baseline conditions in HL3 saline alone for 5 min. Next, media was replaced with HL3 saline + 500 nM CNO and the heart observed again for 5 min. Finally, media was then replaced with HL3 saline alone (pH 7.1) (washout), and the heart observed for 5 min.

Adult Fly Behavior Assays

Courtship and Mating Assay

For the courtship assay, bottles of wild-type CS flies were cleared and newly eclosed virgin females and males were collected and matured in 10 ml glass tubes containing ~300 μl of food (10% sucrose, 1% agarose, and drug where appropriate) for 5 days. Between five and six virgin females were housed together during this process, while sexually naïve males were individually housed. During the isolation period, all flies were maintained at 25°C under a 12 hr light/dark cycle until testing. Following the isolation period, one male and one female were transferred to a single chamber of a mating wheel, and each mating pair was closely monitored for 10 min (Nichols et al., 2012). The courtship index was calculated as the amount of time a male spent performing courtship behaviors over 10 min or until copulation occurred. All testing was performed at 25°C at 70%–80% relative humidity, and between the hours of 11 am and 4 pm.

Adult Locomotor Assay and Diurnal Activity

Crosses were made between the different UAS-DREADD responder strains and the appropriate GAL4 driver strain. Then 2- to 3-day eclosed adult male flies were individually placed into 5 mm diameter glass capillary tubes with an agar plug at one end consisting of 1% agarose, 10% sucrose, and CNO (where appropriate), and then plugged with cotton at the other end. Tubes were then placed into Trikinetics activity monitor arrays, which were subsequently placed into a humidified incubator at 25°C with a 12 hr light-dark cycle (unless stated otherwise) and infrared beam breaks counted with the Trikinetics Drosophila Activity Monitor System (DAMS) at 5 min intervals. Sixteen male flies were used in each experimental trial for each treatment. Only activity data for days 3 and greater were used for analysis, omitting the first 2 days to allow for acclimation to the environment and to build up steady state drug levels.

Learning and Memory

Flies were grown in 8 oz polypropylene bottles on standard cornmeal-molasses food at 25°C under a 12 hr light/dark cycle. Both male and female adult flies were used for all conditioning procedures. For selection of flies for assays, bottles were cleared, and 48 hr later the recently emerged adult flies were transferred to large 64 oz plastic commercial juice bottle for 48 hr with the large end cut off and replaced with fine plastic mesh containing ~2 ml of food in the cap, with or without CNO where appropriate, and without anesthetization. The larger environment prevents over exposure to the food source, and allows flies to properly groom and feed, resulting in clean and dry flies for optimal performance in the conditioning apparatus.
STM was performed as previously described (Johnson et al., 2011). Essentially, ~100 flies were transferred to the training chamber containing an electroifiable grid where they received 4-methylcyclohexanol (MCH) in the presence of shock, followed by a brief period of rest in the absence of shock and odor, and finally benzaldehyde (BA) in the absence of shock. The flies were then transferred to the T-maze (choice point) for testing where they were presented with MCH and BA from either side and allowed to choose for 120 s. The PI was calculated as the number of flies avoiding the shock paired odor minus the number of flies avoiding the unpaired odor divided by the total number of flies assayed. The entire procedure was then repeated using BA as the shock-paired odor and MCH as the unpaired odor, with both calculated performance indices combined to give the overall PI for the trial. For the reversibility studies, flies were maintained on food + CNO (1 mM) for 48 hr and then either tested immediately or placed on food without CNO for 3 and 12 hours before training and testing.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, two figures, and one movie and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2013.08.003.

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REFERENCES


Supplemental Information

EXTENDED EXPERIMENTAL PROCEDURES

Creation and Expression Analysis of UAS-DREADDs

UAS-hM1Dq and UAS-hM4Di

The plasmids containing cDNA for the hM1Dq and hM4Di DREADDs (pCDNA-hM1Dq, pCDNA-hM4Di), and the Drosophila transgenic UAS plasmid, pUAST, were digested with XbaI and KpnI (Promega, Madison, WI). The digested hM1D and hM4D cDNAs were gel purified and ligated into the pUAST vector using the Fast Link DNA Ligation Kit (Epicenter, Madison, WI) following the manufactures instructions. The final products were verified by sequence analysis (Retrogen, San Diego, CA). Ten independent pUAST-hM1D and five independent pUAST-hM4D transgenic lines were generated using the services of BestGene Inc. (Chino Hills, CA).

UAS-rM3Bds

The M3D-Bar cDNA was amplified from the pCDNA-M3D-Bar plasmid by standard PCR procedures using the Apex Hot Start DNA polymerase (Genesee Scientific, San Diego, CA) with primers that contained restriction sites at the 5’ end: R (XbaI) 5’TATATCTAGACTACAAGGCTGCTT3, and F (EcoRI) 5’-AAGAATTCATGTACCCCTACGAC-3’. Both the rM3Bds cDNA PCR product and the pUAST plasmid were digested with XbaI and EcoRI (Promega, Madison, WI), followed by gel purification and ligation of the rM3Bds cDNA into the pUAST vector using the Fast Link DNA Ligation Kit following the manufactures directions. The final product was verified by sequence analysis (Retrogen) as was performed for the clones described above. Ten independent transgenic lines were generated from final cloning products using the services of BestGene Inc.

Using standard genetic mapping techniques, we mapped which chromosomes the insertions were contained in for the inducible UAS-DREADDS (see below). The UAS-hM1Dq is on the first chromosome, and the UAS-hM4Di and rM3Bds insertions are on the third chromosome.

In Vivo Expression Analysis

Each UAS-DREADD transgenic line was crossed to the ACT5C-GAL4 driver to ubiquitously express DREADD transcript. The degree of induction of expression was determined by QPCR, with the highest inducible ones being selected for subsequent use. Total RNA was isolated from adults of the parental lines and the F1 progeny (20 flies for each) using Tri-Reagent (Molecular Research Center, Cincinnati, OH) following manufacturers instructions. The final products were verified by sequence analysis (Retrogen, San Diego, CA). Ten independent transgenic UAS plasmid, pUAST, were digested with XbaI and KpnI (Promega, Madison, WI). The digested hM1Dq and hM4D cDNAs (U#55) F: 5’-TGATGCTCATCGACTACACTT-3’, R: 5’-GGAGTCCCTCAGATCGACAG-3’, rM3Bds (U#16) F: 5’-CTTCTCCTGGGCTGACAGTC-3’, R: 5’-TGTAAGCTGCGCAACTGAGTG-3’, and hM4Di (U#9) F: 5’-GAGCCTGCTGACTGTGCTG-3’, R: 5’-AGCTGCTGTGTTAGCTTGTGAT-3’. QPCR was performed with the BioRad iCycler IQ5 (BioRad, Hercules, CA) using the HotStart-IT Probe qPCR Master Mix (USB, Cleveland, OH) following the manufacturer’s instructions (25 µl reaction volume; cycle parameters: initial 95°C for 2 min, followed by 44 cycles of 95°C 15 s, 60°C 45 s) in 96 well plate format. Reactions were performed in quaduplicate for each gene and genotype. Expression of RpL32 was used as the reference control to normalize expression between genotypes. Expression levels were calculated using the ΔΔCt method (ABI: User Bulletin #2, ABI Prism 7700 Sequence Detection System, 10/2001).

Analysis of ΔF/F

A custom MATLAB program was written to analyze the effects of CNO on the brain by quantifying the change in fluorescence of a live cell reporter, GCaMP, expressed in individual neurons of the ventral ganglia by the GAL4-elayΔ155 driver in response to stimulation with KCl (75 mM) with or without CNO present. For the analysis, a real time imaging movie collected on a Leica SP2-TCS confocal microscope was loaded into our MATLAB program. Next, regions of interest were drawn around 10 neurons of the ventral ganglia, and a separate region of interest representing background fluorescence was selected. The average fluorescence intensity per neuron (minus the average background intensity) over the first 30 frames (prior to KCl administration; 1.3 s/frame) was calculated as the baseline value. This calculated baseline value was then used to normalize the fluorescence intensity for each frame over the entire time course of the assay for the fluorescence intensity of each neuron measured. The data for all ten neurons was then averaged to generate the ΔF/F data for a given brain over the 5 min of the assay. This process was repeated 14 more times for a total of 15 brains measured for each of the five treatment conditions. The software then calculated the mean ΔF/F curve ± S.E.M., and the mean fluorescence maximum frame ± SEM for each treatment group to generate the final ΔF/F data as presented in Figure 3. Although the software is capable of plotting the data, Figure 3 represents the final data plotted using GraphPad Prism 5.0.

Body Wall and Mouth Hook Contractions

Larvae were collected as described above and transferred to a small beaker containing 1.2mL of 5% sucrose (w/v) solution in water (either alone or with the appropriate concentration of CNO) for 15 min. Body wall contractions were counted as a measure of activity by placing larva singly on a Petri dish containing 1% agarose gel, letting the larvae acclimate for 1 min, and then counting the number of full posterior to anterior movements in a 60 s period. As a measure of feeding/taste, larvae exposed to CNO were placed in a dilute yeast paste solution, allowed to acclimate for 1 min, and the number of mouth hook movements were counted in a 60 s period. 20 larvae were counted for each assay (Nichols et al., 2012).
**Adult Lifespan**

Between 20-30 adult males and females were collected separately 24 hr post-eclosion and placed on 1 ml of normal food in a 15 ml culture tube either with or without 3 mM CNO. The number of dead flies were counted every 24 hr. Surviving flies were placed on fresh food every 6-7 days until the end of the experiment.
Figure S1. CNO Does Not Have Any Overt Developmental Consequences, Related to Results

(A) CNO does not delay pupation. Single pairs of mating flies were placed in vials containing either normal food, or food + CNO (3 mM), and allowed to lay eggs for five days. The time to pupation was measured for each progeny, and each progeny after eclosion was examined under the dissection microscope for gross morphological defects. There was no significant difference in the time to pupation between treatment groups (n = 3 vials/treatment; error bars indicate SEM).

(B) CNO does not effect the lifespan of the fly. Between 20-30 newly eclosed males and females were housed separately in 15 ml tubes containing 1.0 ml normal food +/- CNO (3 mM) (food changed for fresh food +/- CNO every week). The number of dead flies were counted every 24 hr to determine overall viability. CNO was found to have no effect on the lifespan of the adult fly. Data points represent the mean of three independent tubes.
Figure S2. Expression of UAS-DREADD Receptors Is Inducible, Related to Experimental Procedures
Quantitative RT-PCR demonstrates that expression of hM1Dq (left panel), hM4Di (center panel) and rM3BDs (right panel) is inducible by GAL4 drivers. RNA tested was isolated from male flies carrying either one of the UAS-DREADD transgenes alone (light gray bar), or the UAS-DREADD transgene in combination with the ACT5C-GAL4 driver (dark gray bars). (Error bars indicate SEM between quadruplicate reactions).