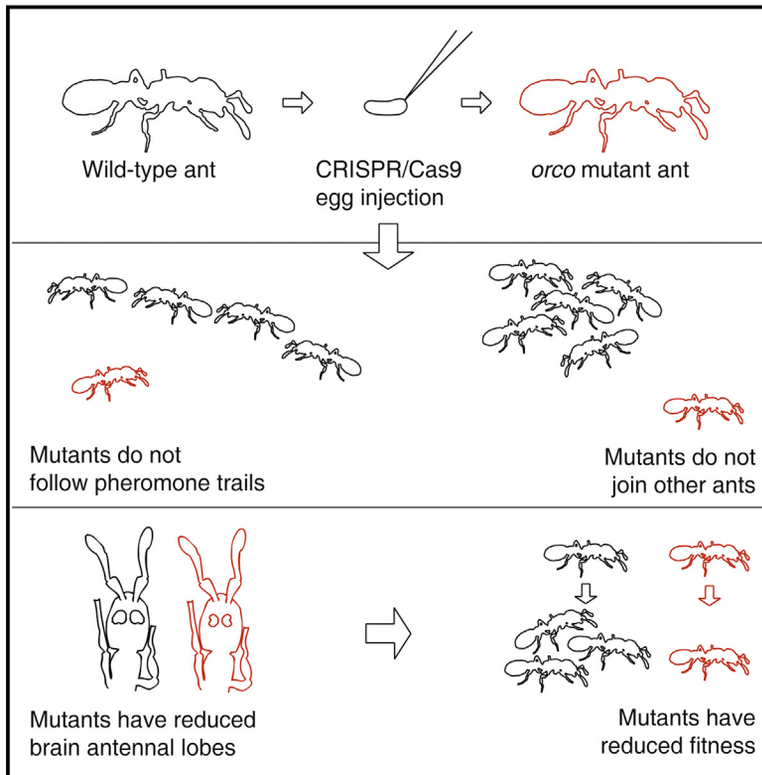


orco Mutagenesis Causes Loss of Antennal Lobe Glomeruli and Impaired Social Behavior in Ants

Graphical Abstract



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In Brief

Development of the first line of mutant ants, using CRISPR/Cas technology, reveals what happens inside of a colony when ants lose the ability to recognize odors.

Highlights

- We generated odorant receptor co-receptor (*orco*) mutants in an ant
- *orco* mutants did not follow pheromone trails or cluster with other ants
- Mutant ants also had greatly reduced antennal lobes in the brain
- Odorant receptors are essential for the complex organization of ant societies



orco Mutagenesis Causes Loss of Antennal Lobe Glomeruli and Impaired Social Behavior in Ants

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SUMMARY

Life inside ant colonies is orchestrated with diverse pheromones, but it is not clear how ants perceive these social signals. It has been proposed that pheromone perception in ants evolved via expansions in the numbers of odorant receptors (ORs) and antennal lobe glomeruli. Here, we generate the first mutant lines in the clonal raider ant, *Ooceraea biroi*, by disrupting *orco*, a gene required for the function of all ORs. We find that *orco* mutants exhibit severe deficiencies in social behavior and fitness, suggesting they are unable to perceive pheromones. Surprisingly, unlike in *Drosophila melanogaster*, *orco* mutant ants also lack most of the ~500 antennal lobe glomeruli found in wild-type ants. These results illustrate that ORs are essential for ant social organization and raise the possibility that, similar to mammals, receptor function is required for the development and/or maintenance of the highly complex olfactory processing areas in the ant brain.

INTRODUCTION

Ants live in complex societies and display sophisticated social behavior, including reproductive division of labor between queens and workers, behavioral division of labor between nurses and foragers, the formation of adaptive foraging networks, nestmate versus non-nestmate discrimination, and collective nest construction (David Morgan, 2009; Grüter and Keller, 2016; Leonhardt et al., 2016; Richard and Hunt, 2013). All of these behaviors are largely mediated via chemical communication using a wide range of pheromones. In *Drosophila*, pheromone receptors have been identified that belong to multiple insect chemosensory receptor families, including odorant receptors (ORs), gustatory receptors (GRs), ionotropic receptors (IRs), and pickpocket channels (PPKs) (Kohl et al., 2015). Ants have numbers of GRs, IRs, and PPKs that are typical for insects, while their OR repertoire is highly expanded (McKenzie et al., 2016;

Oxley et al., 2014; Smith et al., 2011a, 2011b; Zhou et al., 2012) (Figure 1A; Table S1). This raises the possibility that the expansion of ORs specifically, rather than chemoreceptors in general, may underlie the evolution of complex chemical communication in ants. Ants also have exceedingly large numbers of glomeruli in their antennal lobes, which likely mirror their expanded OR gene repertoire (McKenzie et al., 2016; Zube et al., 2008) (Table S1). Insect ORs function as chemosensory receptors by dimerizing with the highly-conserved co-receptor protein Orco to form ligand-gated ion channels (Jones et al., 2005; Larsson et al., 2004; Sato et al., 2008). *orco* null mutants in fruit flies, locusts, mosquitoes, and moths therefore lose OR function and show impaired responses to odors such as food volatiles and sex pheromones (Asahina et al., 2008; DeGennaro et al., 2013; Koutroumpa et al., 2016; Li et al., 2016; Yang et al., 2016). Thus, *orco* constitutes a prime candidate to test the hypothesis that the expanded OR repertoire of ants is required for chemical communication.

The receptor families involved in pheromone perception in ants have not been functionally characterized, in part because the complex life cycle of ants has hindered the development of functional genetic tools (Grüter and Keller, 2016; Kohno et al., 2016; Reid and O'Brochta, 2016; Schulte et al., 2014). Indeed, to this point, no mutant lines in ants have been generated (e.g., using transgenesis, random mutagenesis, or targeted mutagenesis). Most ant species reproduce sexually, so generating homozygous mutant lines requires multiple generations of crosses, which is challenging given that the generation times of ants tend to be many months (Yan et al., 2014). Furthermore, performing such crosses raises methodological concerns, as inbreeding individuals to generate homozygous mutants also has the potential to homozygote off-target mutations (Fu et al., 2013). In other model organisms, this issue can be addressed by using multiple generations of outcrossing, but this is not feasible in species with long generation times such as ants (Fu et al., 2013; Kistler et al., 2015). Thus, most ant species pose major challenges to the generation of mutant lines that can be used to generate reliable scientific results.

In this study, we take advantage of the unique biology of the clonal raider ant, *Ooceraea biroi*, to generate the first mutant lines in ants. This species is a promising genetic model system

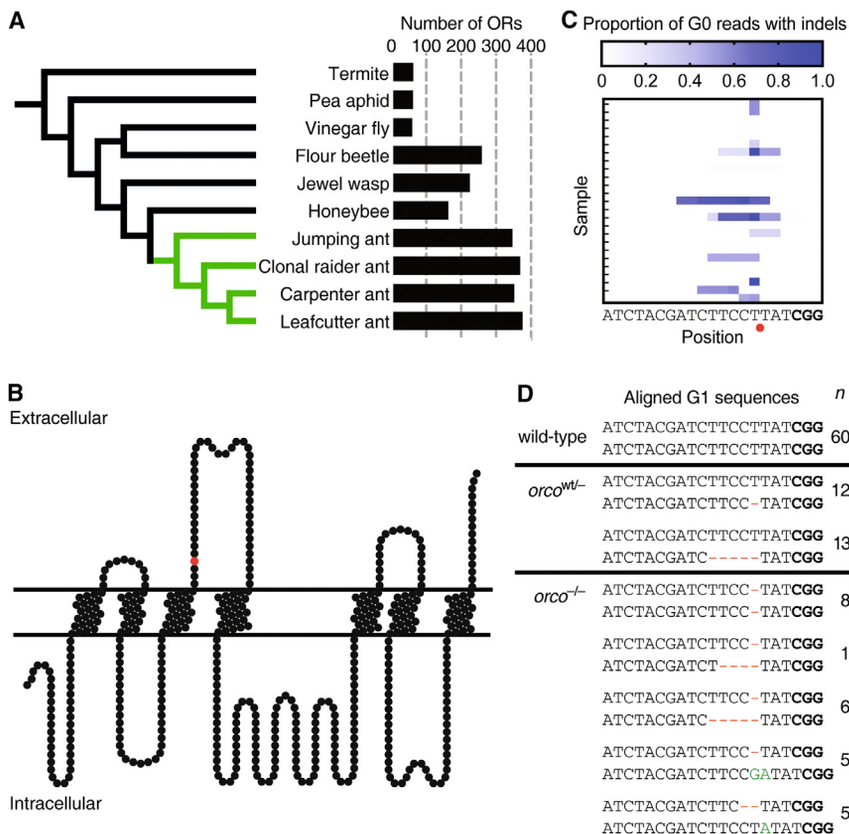


Figure 1. Number of OR Genes and *orco* Mutagenesis

(A) Phylogeny with numbers of ORs for ants (green) and other insects (black), showing ant OR expansion (Table S1).

(B) Position of predicted CRISPR/Cas9 cut site in Orco protein model (red circle). Frameshift mutations at this position truncate the wild-type protein between the third and fourth transmembrane domains, and the resultant mutant protein is unlikely to form functional ion channels.

(C) Proportion of Illumina sequencing reads of *orco* amplicons with insertions or deletions (indels) relative to gRNA sequence in G0s, showing mutation rates of at least 97% in some individuals. Red circle indicates predicted CRISPR/Cas9 cut site. Protospacer adjacent motif (PAM) in bold.

(D) Wild-type *orco* sequence compared to sequences for the two *orco^{wt/-}* and the five *orco^{-/-}* mutant lines. Deletions are shown in red and insertions in green. *orco^{-/-}* ants have two frameshift alleles and are therefore expected to be complete loss-of-function *orco* mutants. Each of these lines arose independently; *n* indicates the number of ants of each line (G1s and subsequent generations) used across all experiments in this study (see Table S3 for a description of the lines used in each specific experiment). PAM in bold. See also Figure S1 and Tables S1 and S2.

because, unlike most other ant species, *O. biroi* reproduces via parthenogenesis, so stable germ-line modifications are obtained from the clonal progeny of injected individuals without laboratory crosses (Oxley et al., 2014). On the other hand, this unusual reproductive system implies that it will be challenging to conduct laboratory crosses for other purposes. *O. biroi* also has a generation time of ~2 months, which is among the shortest known for any ant species (Oxley et al., 2014). Here, we develop a CRISPR/Cas9 protocol in *O. biroi* and use this technique to create *orco* null mutants, allowing us to conclusively test the role of ORs in chemical communication in ants. To reduce the potential for off-target mutagenesis, we designed a guide RNA (gRNA) that has a unique sequence relative to the rest of the genome (see STAR Methods) and characterized multiple independently generated lines of heterozygous and homozygous *orco* mutants. We find that *orco* mutant ants, as expected, lack Orco antibody staining and possess deficiencies in general olfactory behaviors. These ants also have deficiencies in social behavior, including an inability to nest with other ants or follow pheromone trails, and exhibit reduced fitness. In contrast with previously studied insects, we also find that *orco* mutant ants possess striking neuro-anatomical phenotypes, including a loss of olfactory sensory neurons (OSNs) in the antennae and a striking reduction in the number of glomeruli in the antennal lobes. Collectively, these results illustrate the crucial significance of *orco* and ORs in ant biology.

RESULTS

Generation of *orco* Mutant Ants

We confirmed the identity of *orco* in the *O. biroi* genome (Figure S1; Table S2) and designed and synthesized a gRNA to target *orco* (Figure 1B). To produce *orco* mutants, we injected Cas9 protein and gRNA into 3,291 eggs <5 hr of age and produced 42 G0 adults, some of which displayed mutations in at least 97% of PCR amplicons of the *orco* target site (Figure 1C; STAR Methods). G0 mutations in the germline can be inherited by G1s to produce stable modifications to the genome (Reid and O'Brochta, 2016). Given that *O. biroi* reproduces through parthenogenesis (Oxley et al., 2014), stable mutant lines can be clonally propagated from individual mutant G1s and subsequent generations. *orco* loss-of-function mutant lines are thus derived from G1 eggs with independent frameshift mutations in both *orco* alleles. We recovered a diverse set of *orco* mutant lines, including two *orco^{wt/-}* lines with one frameshift allele and five *orco^{-/-}* lines with two frameshift alleles (Figure 1D). The phenotypes reported below were consistent across the two *orco^{wt/-}* lines and across the five *orco^{-/-}* lines, respectively. Descriptions of the specific lines used in each experiment along with the associated phenotypes for each individual are given in Table S3.

Antennal Morphology

To test whether *orco* frameshift mutations cause a loss of the full-length Orco protein and/or change the distribution of olfactory

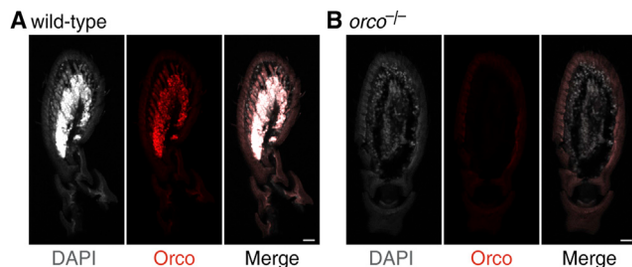


Figure 2. Loss of Orco Expression and OSNs in *orco*^{-/-} Ants

(A) Antennal section of wild-type *O. biroi* showing DAPI counterstain (gray), Orco immunostain (red), and the merged image (black). Wild-type ants possess a dense region of Orco-positive OSNs in the center of the antenna. (B) Antennal section of *orco*^{-/-} ant. *orco*^{-/-} ants lack Orco staining, indicating that the full-length Orco protein is absent. *orco*^{-/-} ants also lack the dense region of cells in the center of the antenna, indicating that most or all of the OSNs that would express *orco* in wild-types are absent in *orco*^{-/-} ants. Scale bars, 20 μ m.

sensory neurons (OSNs), we immunostained *O. biroi* wild-type and *orco* mutant antennal sections with an Orco antibody and counterstained the cell nuclei with DAPI. Wild-type and *orco*^{wt/-} *O. biroi* antennae contained a dense region of Orco-positive OSNs in the center of the antennal club, as indicated by DAPI and Orco staining (Figures 2A, S2A, and S2B). As predicted, antennae of *orco*^{-/-} ants lack Orco staining, indicating that the full-length Orco protein is absent (Figures 2B and S2C). Surprisingly, the majority of OSNs in the *O. biroi* antenna are also absent in *orco* mutants, as indicated by a reduction in DAPI signal (Figures 2B and S2C). This result suggests that the *orco*-positive OSNs in the *O. biroi* antenna are absent in *orco* mutants, but the smaller number of other types of sensory neurons may remain present. This contrasts with *D. melanogaster*, in which the antennal OSNs that express *orco* in wild-types are still present in *orco* mutant adults (Chiang et al., 2009).

Antennal Lobe Morphology

In *D. melanogaster*, all antennal lobe glomeruli that have been examined remain present in *orco* mutants, implying that OR function is not required for gross antennal lobe morphology (Chiang et al., 2009; Larsson et al., 2004). These results contrast strongly with mice, where olfactory receptor function and neuronal activity are essential for the formation and maintenance of the analogous brain region, the olfactory bulb (Yu et al., 2004). However, *D. melanogaster* has only 60 ORs and a similar number of glomeruli, while mice possess over 1,000 olfactory receptors and glomeruli. This striking difference suggests that complex olfactory systems must rely on receptor function for their development and/or maintenance. Ants have highly expanded numbers of ORs and antennal lobe glomeruli (McKenzie et al., 2016; Oxley et al., 2014; Zhou et al., 2012; Zube et al., 2008) (Table S1), raising the possibility that the development and/or maintenance of ant antennal lobes may require additional mechanisms to exceed the complexity found in other insects.

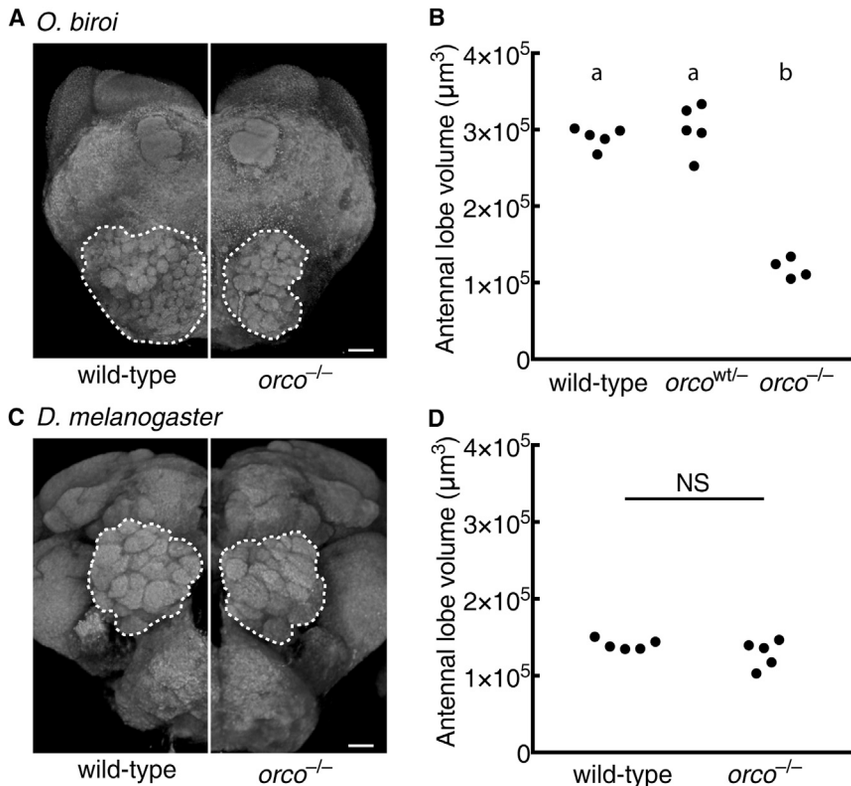
To address whether OR function might be required for the structure of the ant antennal lobe, we imaged brains using confocal microscopy, measured antennal lobe volumes and the number of glomeruli, and reconstructed antennal lobes in

wild-type and *orco* mutant adults in *O. biroi* and *D. melanogaster*. We found that the antennal lobes of *orco*^{-/-} ants measured only one-third of the volume of wild-type and *orco*^{wt/-} antennal lobes, and ~82% of the glomeruli were lost (Figures 3A and 3B; Movie S1). However, all six glomeruli in the T7 cluster of the ant antennal lobe, which is believed to be innervated by olfactory sensory neurons that do not express ORs (Nakanishi et al., 2010), were still present in *orco*^{-/-} individuals. No differences were observed in the volume of the protocerebrum, mushroom bodies, or central complex relative to wild-type ants ($p = 0.45, 0.17, \text{ and } 0.20$, respectively; t test). In contrast, we detected no significant difference in antennal lobe volumes and only minor potential differences in glomerulus numbers between wild-type and *orco*^{-/-} flies (Figures 3C and 3D; Movie S1; STAR Methods). To test whether antennal lobes degenerate as a function of age in *orco*^{-/-} flies, we imaged antennal lobes of additional wild-type and *orco*^{-/-} flies aged for 30 days, showing that antennal lobe glomeruli were not reduced (Figure 3C). These results demonstrate that development and/or maintenance of antennal lobes in ants, but not flies, are indeed dependent on Orco function.

Behavioral Phenotypes

Based on the general observation that ants are often repelled by the smell of permanent markers, we developed a simple assay to test whether *orco* mutants have compromised chemosensory abilities. We found that wild-type and *orco*^{wt/-} *O. biroi* are indeed strongly repelled by lines drawn with Sharpie permanent marker and rarely contact or cross Sharpie lines (Figures 4A–4C). Given that *O. biroi* is blind, and the ants are often repelled before touching the Sharpie lines (Movie S2), it is clear that this behavior is mediated via olfaction, rather than visual or tactile cues. However, *orco*^{-/-} *O. biroi* are significantly less repelled by Sharpie (Figures 4A–4C; Movie S2), implying that *orco* is required to perceive the odorants that cause Sharpie lines to be repulsive. These results suggest that *orco* mutant ants possess general olfactory deficiencies, similar to *orco* mutants in other types of insects (Asahina et al., 2008; DeGennaro et al., 2013; Koutroumpa et al., 2016; Li et al., 2016; Yang et al., 2016).

Pheromone trails are a major feature of chemical communication in many ants and are important for coordinating collective behaviors (David Morgan, 2009; Zube et al., 2008). In many doryline ants, including *O. biroi*, disturbance of colonies leads to robust trail formation, likely for the purpose of nest relocation (Gotwald, 1995; Hölldobler, 1982; Schneirla, 1971). To test whether *orco* influences the ability of *O. biroi* to follow pheromone trails, we set up five experimental colonies composed of 12–14 identically reared G1s with wild-type, *orco*^{wt/-}, and *orco*^{-/-} genotypes and individually tagged each ant with color dots (Figure 4D). We recorded videos of each colony and used a novel custom-built automated behavioral tracking system employing painted color tags (rather than paper barcodes) (Mersch et al., 2013) to individually identify the ants and quantify their behavior (Figures 4D and S3; Movies S3 and S4). We disturbed each colony at the beginning of each video, causing the colonies to form conspicuous trails. During the ensuing period of high activity, we created a 2D histogram, or density map, of movement for each ant and measured the Pearson



(D) Antennal lobe volumes for wild-type ($n = 5$) and *orco*^{-/-} ($n = 5$) flies. Volumes of wild-type and *orco*^{-/-} antennal lobes are not significantly different ($p = 0.20$, t test). Scale bars, 20 μm . NS, not significant. Genotypic classes marked by different letters are significantly different ($p < 0.05$) after ANOVA followed by Tukey's test (B).

See also Tables S1 and S3.

correlation coefficient of this density map with the density map of the other ants in the colony, reasoning that density maps would be more highly correlated when ants were following pheromone trails (Figures 4E and 4F). To provide a null expectation, we also compared the density maps of individual ants to a randomized density map of other ants in the colony (Figures 4E and 4F). We found that the density maps of individual ants had significantly higher correlations with the density maps of the rest of the ants in the colony than with the randomized density maps in wild-type, but not *orco*^{-/-}, ants (Figure 4F; average correlation coefficients were 0.31 versus 0.04 in wild-type and 0.07 versus 0.02 in *orco*^{-/-}, respectively). These findings imply that trail following behavior is reduced or absent in *orco*^{-/-} ants, likely because they are unable to perceive chemical pheromone trails.

Nesting behavior, and the formation of aggregations more generally, is a ubiquitous feature of social insect biology (Depickere et al., 2004). Immediately upon eclosion, we noticed that some G1s did not nest with other ants, but instead showed a wandering phenotype (Figure 4G). In a set of 16 G1 colonies, we used this wandering phenotype to identify colonies containing *orco*^{-/-} ants with 100% accuracy ($p < 0.001$, Fisher exact test, see STAR Methods). To more precisely measure nesting behavior in *orco* mutants, we recorded and analyzed 24 hr videos of each experimental colony. We found that wild-type and *orco*^{wt/-} ants aggregated into tight clusters and exhibited

little movement outside the cluster, while *orco*^{-/-} ants frequently exited the cluster and wandered around the dish (Figure 4H; Movie S3). Overall, *orco*^{-/-} ants spent a significantly larger fraction of time without contact with other ants when compared to wild-type and *orco*^{wt/-} ants (Figure 4I). These findings demonstrate that typical nesting behavior is compromised in *orco*^{-/-} ants. This observation is consistent with the idea that *orco* mutants are unable to perceive odorants, such as aggregation pheromones (Bell et al., 1972; Depickere et al., 2004; Li et al., 2016), that might be involved in nesting behavior.

Fitness of *orco* Mutants

Finally, we investigated whether *orco* mutations influence fitness by measuring egg-laying and survival rates of wild-type and *orco* mutant ants. We found that *orco*^{-/-} ants laid significantly fewer eggs than wild-type and *orco*^{wt/-} ants over a 2-week period (Figure 5A), and *orco*^{-/-} ants exhibited significantly higher mortality than wild-types over a 34-day period (Figure 5B). These results contrast with *orco* mutant *Drosophila* and mosquitoes, in which no fitness effects are observed under typical laboratory rearing conditions (Asahina et al., 2008; DeGennaro et al., 2013). This suggests that the *orco* mutant phenotype has serious consequences for ant fitness (for a discussion of the potential role of off-target effects, see the STAR Methods). It is possible that these fitness effects result because *orco*^{-/-} ants are unable to

Figure 3. Reduced Antennal Lobes in *orco*^{-/-} Ants

(A) *O. biroi*: dorsal (n-ventral) 3D projections of *orco*^{-/-} and wild-type ants. Antennal lobes indicated by dashed lines. *orco*^{-/-} antennal lobe is highly reduced relative to wild-type. Two *orco*^{-/-} ants had 90 and 91 glomeruli relative to 493 and 509 glomeruli for two wild-type ants (one of the wild-type reconstructions has been published previously [McKenzie et al., 2016]); small differences between replicates within treatments might reflect reconstruction errors or actual biological variation.

(B) Antennal lobe volumes for wild-type ($n = 5$), *orco*^{wt/-} ($n = 5$), and *orco*^{-/-} ($n = 4$) ants (*orco*^{wt/-} and *orco*^{-/-} were age-matched at ~ 4 months old). *orco*^{-/-} ants, but not *orco*^{wt/-} ants, have significantly smaller antennal lobes than wild-type.

(C) *D. melanogaster*: anterior (n-ventral) 3D projections for wild-type and *orco*^{-/-} brains from 1-month-old flies. Antennal lobes indicated by dashed lines. *orco*^{-/-} antennal lobe is similar to wild-type. Two *orco*^{-/-} flies had 43 and 44 glomeruli, and two wild-type flies each had 46 glomeruli. These glomerulus numbers were higher than has been previously reported, which is likely due to differences in sample preparation and imaging techniques. Slight differences in glomerulus numbers between replicates may be due to reconstruction errors, or may reflect modest antennal lobe phenotypes in *orco* mutant flies (STAR Methods).

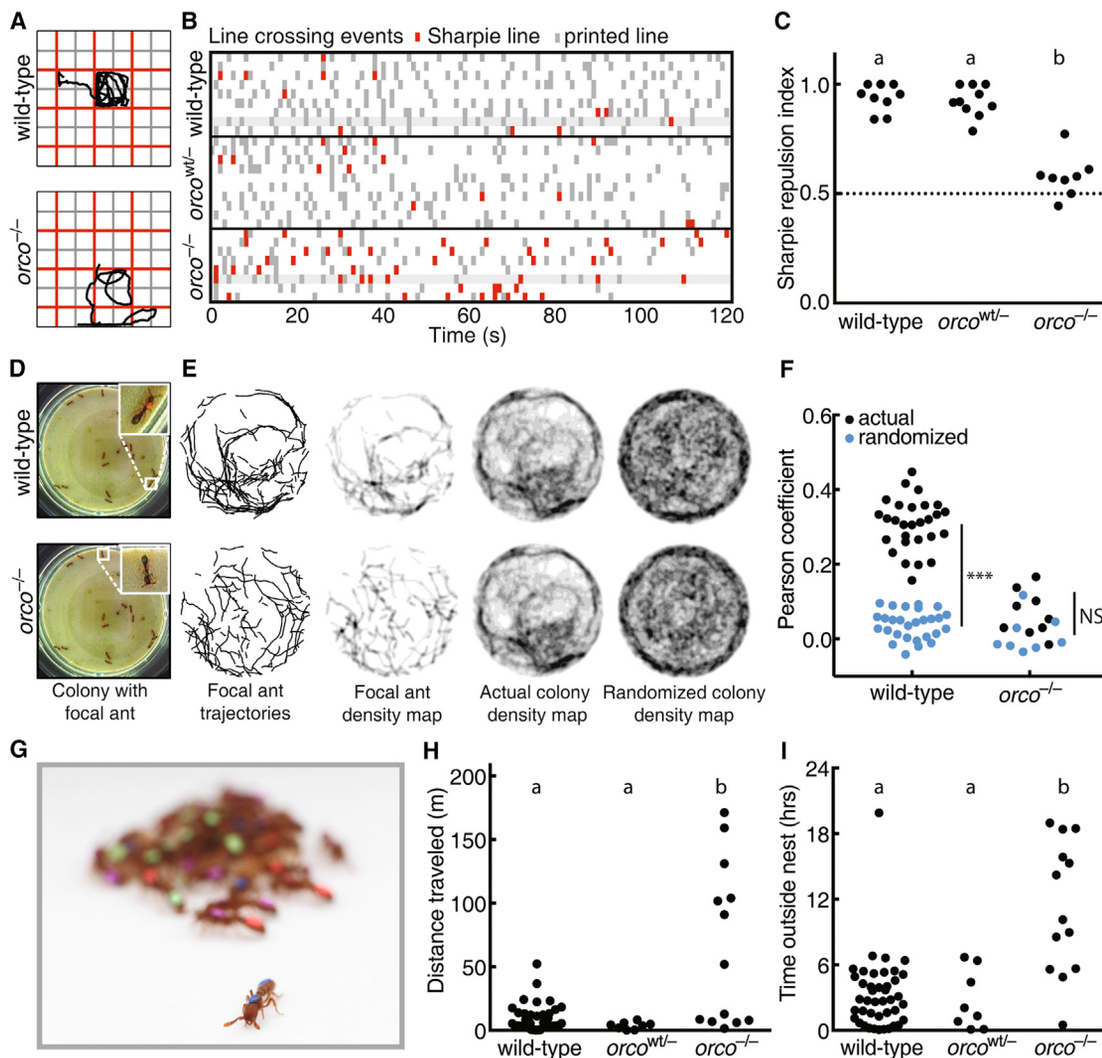


Figure 4. Deficient Olfactory and Social Behavior in *orco*^{-/-} Ants

(A) Example trajectories of wild-type and *orco*^{-/-} ants in Sharpie assay.

(B) Line crossing events for wild-type ($n = 9$), *orco*^{wt/-} ($n = 10$), and *orco*^{-/-} ($n = 8$) ants in Sharpie assays, with ants from (A) highlighted. Wild-type and *orco*^{wt/-} ants cross Sharpie lines (red) less frequently than printed lines (gray), but *orco*^{-/-} ants cross both lines at approximately equal frequencies.

(C) Repulsion indices for ants in Sharpie assays. Repulsion index is calculated as proportion of printed line crosses. *orco*^{-/-} ants, but not *orco*^{wt/-} ants, are significantly less repelled than wild-types.

(D) Example colony used for trail pheromone analysis. The same colony, containing a mixture of wild-type, *orco*^{wt/-}, and *orco*^{-/-} ants, is shown twice, with a wild-type or *orco*^{-/-} focal ant highlighted.

(E) Example trail pheromone analysis. Trajectories, during which ants were moving and edges were excluded, were used to create 2D histograms, or density maps, for each ant in the colony. These density maps were compared to the actual and randomized density maps for all other ants in the colony. The wild-type density map is more strongly correlated with the actual colony density map than with the randomized colony density map, while the *orco*^{-/-} density map is poorly correlated with both colony density maps.

(F) Pearson correlation coefficients for individual ant density maps with the actual or randomized density map of the other ants in the colony. Pearson correlation coefficients for wild-type ants ($n = 28$), but not for *orco*^{-/-} ants ($n = 9$), were significantly higher in actual than randomized density maps (sample sizes were too small to test *orco*^{wt/-}).

(G) Example colony showing an individual outside of the nest.

(H) Distances traveled in 24 hr videos by ants in experimental colonies. *orco*^{-/-} ants ($n = 10$), but not wild-type ($n = 40$) or *orco*^{wt/-} ($n = 8$) ants, exhibit a wandering phenotype.

(I) Time without contacting other ants in 24 hr videos. *orco*^{-/-} ants spend more time without contact than wild-type or *orco*^{wt/-} ants. *** $p < 0.001$; NS, not significant. Genotypic classes marked by different letters are significantly different ($p < 0.05$) after ANOVA followed by Tukey's test (C), or from log-likelihood ratio tests on generalized linear mixed models followed by Tukey's tests with colony as a random variable and actual/randomized maps (F) or genotypic class (H and I) as fixed variables.

See also Figure S3 and Table S3.

than any other social insect model system (Yan et al., 2014). Thus, the clonal raider ant provides a unique opportunity to rapidly generate and phenotype mutant lines in a social insect. While all data in the present study were collected from germ-line mutants (G1s and subsequent generations), we also found very high somatic cut-rates among the G0 ants (Figure 1C). Multiple G0s displayed nearly 100% mutation rates and exhibited wandering phenotypes similar to germ-line mutants in subsequent generations (Figure 1C; STAR Methods). Thus, our results show that CRISPR/Cas9 can have very high efficiency in social insects and may be useful for conducting experiments in G0s even in the large number of species where it is not feasible to maintain stable mutant lines.

Major transitions in evolution require the coordinated action of individuals to operate as a functional, higher-level unit (Maynard Smith and Szathmari, 1997). During the transition from solitary to eusocial living in ants, this coordination was largely achieved via pheromones, and ants accordingly possess highly expanded antennal lobes and OR repertoires. Our results illustrate the functional significance of these striking changes and imply that the expansion of ORs may have been an important component of the evolution of eusocial behavior.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
- METHOD DETAILS
 - CRISPR Methods
 - Treatment of Eggs and Larvae
 - Tagging, Genotyping, and Sequencing
 - Neuro-anatomy
 - Behavior and Fitness
- QUANTIFICATION AND STATISTICAL ANALYSIS
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes three figures, three tables, and four movies and can be found with this article online at <http://dx.doi.org/10.1016/j.cell.2017.07.001>.

A video abstract is available at <http://dx.doi.org/10.1016/j.cell.2017.07.001#mmc7>.

Supporting Citations

The following references appear in the Supplemental Information: Dippel et al. (2016); Engsontia et al. (2008); Engsontia et al. (2015); Flanagan and Mercer (1989); Kelber et al. (2009); Kollmann et al. (2011); Robertson et al. (2003); Robertson et al. (2010); Smadja et al. (2009); Terrapon et al. (2014).

AUTHOR CONTRIBUTIONS

W.T. and D.J.C.K. conceived the project with input from B.M. W.T., B.M., and L.O.-C. conducted preliminary experiments to develop CRISPR methods in *O. biroi*. W.T., N.-C.C., and L.O.-C. performed injections and ant rearing. W.T., N.-C.C., L.O.-C., and P.R.O. performed sequencing and analysis with input from B.M. L.O.-C. performed brain dissections and antibody assays.

S.K.M. performed remaining neuroanatomical experiments and analyses. W.T., L.O.-C., N.-C.C., and J.S. performed behavioral experiments. J.S. developed automated tracking techniques and performed behavioral analyses involving automated tracking. W.T., N.-C.C., and L.O.-C. performed fitness experiments. D.J.C.K. supervised the project. W.T. and D.J.C.K. wrote the paper. All authors discussed the results and the manuscript.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-Orco antibody	Vanessa Ruta	N/A
3C11 Anti-SynORF-1	Developmental Studies Hybridoma Bank	3C11
NC82	Developmental Studies Hybridoma Bank	NC82
Alexa Fluor 647 donkey anti-mouse	Millipore Sigma	AP192SA6
Alexa Fluor 594 donkey anti-mouse	Invitrogen	A21203; RRID: AB_141633
Critical Commercial Assays		
QIAamp DNA Micro Kit	QIAGEN	56304
MEGAscript T7 Transcription Kit	ThermoFisher Scientific	AM1334
MEGAclear Transcription Clean-Up Kit	ThermoFisher Scientific	AM1908
Agencourt AMPure XP	Beckman Coulter Life Sciences	A63882
Agilent RNA 6000 Pico Kit	Agilent Technologies	5067
Nextera XT DNA Library Preparation Kit	Illumina	FC-131-1096
MiSeq Reagent Kit v3 (150 cycle)	Illumina	MS-102-3001
KAPA Library Quant kit (Illumina)	KAPA	KK4824
Universal qPCR Mix		
Mwol with CutSmart Buffer	New England Biolabs	R0573
Chemicals, Peptides, and Recombinant Proteins		
Cas9	PNABio	CP01
Oligonucleotides		
O4_F: 5'- TCGTCGGCAGCGTCAGATGTGTATA AGAGACAGTCCAACCTTGCTGTAAATTTGGAT-3'	IDT	N/A
O4_R R:5'-GTCTCGTGGGCTCGGAGATGTG TATAAGAGACAGCTCTTCTTGGTCGGCGGTA-3'	IDT	N/A
LCO: 5'- GGTCACAAATCATAAAGATATTGG-3'	IDT	N/A
HCO: 5'-TAAACTTCAGGGTGACCAAAAAATCA-3'	IDT	N/A
O4_gRNA_primer: 5'- GAAATTAATACGACTCAC TATAGGTCTTCATGACCCTTATGTGTTTTAGAG CTAGAAATAGC-3'	IDT	N/A
Software and Algorithms		
CRISPRseek	Zhu et al., 2014	http://bioconductor.org/packages/release/bioc/html/CRISPRseek.html
genomepy	This paper	https://github.com/oxpeter/genomepy
phmmer	Eddy, 1998	http://hmmer.org/download.html
MAFFT	Katoh and Standley, 2013	http://mafft.cbrc.jp/alignment/software/
RAxML	Stamatakis, 2006	https://github.com/stamatak/standard-RAxML
MEGA	Kumar et al., 2016	http://www.megasoftware.net/mega4/
Mutation Surveyor	Softgenetics	http://www.softgenetics.com/mutationSurveyor.php
MATLAB	Version R2016a, The MathWorks, Inc.	https://www.mathworks.com/products/matlab.html
R	R Core Team	http://www.R-project.org/
lme4	Bates et al., 2014	https://github.com/lme4/lme4/
Prism 7	GraphPad Software	https://www.graphpad.com/scientific-software/prism/

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Daniel J.C. Kronauer (dkronauer@rockefeller.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Ooceraea biroi colonies were maintained at 25°C in circular Petri dishes (50 mm diameter, 9 mm height) with a plaster of Paris floor ca. 4 mm thick. Colonies were fed 3 times weekly with fire ant (*Solenopsis invicta*) brood and cleaned and watered at least once per week. Two *O. biroi* clonal lines, which are genetically distinguishable at the mitochondrial *cytochrome oxidase subunit 1* (CO1) gene (Kronauer et al., 2012), were used in this study. All experimental ants belonged to Line B, while Line A ants were only used as chaperones to raise experimental Line B individuals. Experimental ants were reared by placing Line B larvae (G0s) or eggs (G1s and subsequent generations) in colonies of 20 Line A chaperones, and chaperones were removed once the callows had eclosed. This rearing method results in a small fraction of Line A offspring of chaperones in colonies with the G0s and subsequent generations. For this reason, all individuals were genotyped following experiments, and Line A individuals were removed from all analyses. All experimental colonies in this study had eggs removed twice weekly so that adults were maintained without larvae or pupae. All individuals, including those that died during experiments, were genotyped (see below) to determine their clonal line, and *orco* amplicons from Line B individuals were sequenced to determine *orco* genotype.

METHOD DETAILS

CRISPR Methods

Confirmation of the *O. biroi orco* identity

Candidate *orco* orthologs for eight insect species were detected as reciprocal best hits using phmmer (Eddy, 1998) with *D. melanogaster orco* (flybase id: FBgn0037324) as the initial query sequence (Table S2). To confirm orthology, homologs \pm 50% the length of *orco* were aligned with MAFFT (Katoh and Standley, 2013) using default parameters. This alignment was then used to construct a bootstrapped phylogeny with RAxML (Stamatakis, 2006), providing unambiguous support for a single copy ortholog of *orco* in *O. biroi* (Figure S1).

Design of *orco* gRNA

Identification of cut sites and assessment of off-target sites was performed using the script *cris.py*, part of the *genomepy* package (commit #94cc628), available at <https://github.com/oxpeter/genomepy>. The genomic sequence for *O. biroi orco* was searched on both strands for the CRISPR guide RNA (gRNA) recognition sequence 5' - N₂₀NGG - 3' using BLASTN (Altschul et al., 1990) and checked for off-target hits using CRISPRseek (Zhu et al., 2014). We detected no off-target sites with 2 or less mismatches and only one site with 3 mismatches for our *orco* gRNA, which is expected to lead to low or no off-target cutting (Fu et al., 2013).

Potential off-target effects

It has been shown that in some cases CRISPR/Cas9 injections can lead to off-target mutagenesis that in turn can give rise to non-specific phenotypes (Fu et al., 2013). In the present study we used multiple measures of precautions, making it highly unlikely that the phenotypes we report arise due to off-target effects. First, we used a high-quality reference genome to design the gRNA in this study to have no additional target sites in the genome that are likely to lead to off-target cutting (Fu et al., 2013; Oxley et al., 2014). Second, mutations induced by Cas9 are stochastically generated (Fu et al., 2013), such that any off-target effects would likely be present in some G1 lines but not others. The phenotypes we report are consistent across five independently generated *orco*^{-/-} lines, and we do not observe the same phenotypes across two independently generated *orco*^{wt/-} lines (Table S3). Third, the striking reduction of OSNs and antennal lobes in *orco* mutant ants are phenotypes specific to the ant chemosensory system that are unlikely to arise from random off-target effects. These phenotypes provide a direct functional link between the *orco*^{-/-} genotype and the chemosensory deficiencies described in this study. Importantly, the antennal lobe phenotype was entirely discrete: every *orco*^{-/-} brain had substantially smaller antennal lobes than any *orco*^{wt/-} or wild-type brain, even though this phenotype was measured across multiple independently derived *orco*^{-/-} and *orco*^{wt/-} lines (Figure 2B; Table S3). Therefore, while we cannot exclude the possibility that our injections gave rise to some level of off-target mutations, it is unlikely that the specific phenotypes reported in this study arise from off-target effects.

CRISPR reagent preparation

Recombinant Cas9 protein was purchased from PNAbio, and gRNA was synthesized via in-vitro transcription from PCR amplicons (Kistler et al., 2015). Activity of Cas9 and gRNA was validated using in-vitro digestion of PCR amplicons containing the gRNA recognition site with Cas9 and buffer from New England Biolabs. Immediately prior to injection, Cas9 and gRNA were mixed in water to produce a solution with 100 ng/ μ L Cas9 and 10 ng/ μ L gRNA.

Treatment of Eggs and Larvae

Egg collection

Eggs for injections were collected from egg-laying units consisting of 70 *O. biroi* adults without larvae or pupae. Eggs were collected and placed on double-sided tape with the ventral side up on a glass slide, and injected into the anterior end (Oxley et al., 2014). Slides were prepared with up to 80 eggs for injection and ~25 control eggs to validate incubation conditions.

We observed that the presence of eggs in *O. biroi* colonies inhibits the production of new eggs and employed this observation for efficient egg collection. Egg-laying units were left with eggs for 7 days to inhibit worker egg-laying. On day 0 eggs were removed to release inhibition, and on day 2 eggs were removed again to further prevent inhibition. This led workers to synchronously activate their ovaries, and on days 5, 6, and 7 eggs were collected for injection. Following day 7, eggs were not collected from these colonies for 7 days, causing workers to become inhibited, and the protocol was then repeated.

Eggs were collected from colonies under a stereoscope using insect pinning needles. On a typical injection day, eggs were removed from colonies from 10-11 am, and those eggs were used as uninjected incubation controls or fostered into rearing units. Eggs were collected for injections from 2-3 pm and 6-7 pm and injected from 3-4 pm and 7-8 pm, respectively. Therefore, all injected eggs were less than 5 hr old, when *O. biroi* eggs are in a syncytial stage of development with <100 nuclei (Oxley et al., 2014). Typical egg-laying units produced 2-5 eggs per day, and we collected from up to 60 egg-laying units, injecting 100-300 eggs per day.

Egg injection

Injection needles were prepared as in previous studies (Lobo et al., 2006). Injections were performed using an Eppendorf Femtojet with a Narishige micromanipulator. The Femtojet was typically set to Pi 1800 hPa and Pc 500 hPa. Needles were broken by gently touching the needle against a capillary submerged in halocarbon oil. Alternatively, sharper needles were generated by setting the Femtojet to maximum pressure (6000 hPa) and lightly touching the capillary against fibers on the tape. Data in this manuscript result from a combination of both methods.

The anterior end of *O. biroi* eggs is slightly wider than the posterior end, and the ventral surface is concave while the dorsal surface is convex. To inject, eggs were placed on double-sided 3M tape (Model S-10079 from Uline) on a glass slide, with the anterior end forward and the ventral side upward. Eggs were injected into the anterior end, where nuclei are located in early *O. biroi* embryos (Oxley et al., 2014). The ventral side was placed upward, so that larvae hatched with the mouth facing away from the tape, which facilitated successfully recovering larvae from the tape. To inject, eggs were individually submerged in 1-2 μ L drops of water. Eggs were gently pierced with the needle and injected for 1-2 s. During successful injections, little or no cytoplasm is discharged from the egg when the needle is removed. Preliminary trials showed that injection under liquid was necessary to remove the needle without rupturing the chorion, and that water led to higher survival than Ringer, PBS, or halocarbon oil. A video was recorded of every injection session, allowing us to verify that hatching larvae had been successfully injected.

Preliminary injections were conducted using multiple batches of reagents and variable Cas9 and gRNA concentrations. These trials suggested that hatch rates varied inversely with cutrates. Batch effects in hatch rates were observed across different days of injection within the same experimental treatment, requiring multiple injection days and large numbers of eggs (~400) to accurately estimate hatch rates of any given experimental treatment. In the final injection round, eggs were injected with either low (1800 hPa) or high (6000 hPa) pressure, with sharper needles and lower injection volumes used in injections with high pressure. 46 of 2535 eggs (1.8%) hatched after injections with low pressure, and 58 of 756 eggs (7.6%) hatched after injections with high pressure. 25 of the 42 G0s were Illumina sequenced (see *orco* sequencing), and we observed an average of at least 27% cutrates at the predicted cut site resulted from low pressure injections ($n = 17$) relative to 22% from high pressure injections ($n = 8$). *orco*^{wt/-} and *orco*^{-/-} G1s were recovered from G0s injected with each method.

Egg incubation

Following injections, slides with eggs were incubated in air-tight plastic boxes (0.9L SpaceCube boxes from ClickClack). Incubation boxes were prepared with a plaster of Paris floor (85 g plaster of Paris mixed with 50 mL distilled water). The plaster was dried completely after casting, and water was then added until the plaster became saturated with moisture to determine the saturation volume. The plaster was then dried completely once again, after which 20% of the saturation volume of distilled water was added. This procedure produced suitable incubation conditions for 2 weeks, after which the plaster was discarded. Incubating eggs were checked daily, and any water that had condensed on the eggs was removed with Kimwipes™ tissue. Fungus frequently grew on injection slides. Growth was controlled by spacing the eggs at least ~2 mm apart and mechanically breaking up fungal hyphae and overgrown eggs in 100% ethanol using insect pinning needles. This egg-incubation protocol yielded ~60% hatch rates of uninjected control eggs, which is similar to hatch rates of eggs in laboratory colonies.

To synchronize hatching of larvae from injected eggs, eggs injected on days 5, 6, and 7 were incubated at different temperatures. Preliminary trials showed that eggs incubated at 25°C hatch after 9-10 days, while eggs incubated at 30°C hatch after 7-8 days. We therefore incubated eggs injected on days 5 and 7 at 25°C and 30°C, respectively, while eggs injected on day 6 were incubated at 25°C for the first 5 days and then at 30°C until hatching. This protocol resulted in most larvae hatching on days 14 and 15. Once hatching had commenced, larvae were manually removed from the egg membrane with an insect pinning needle, taking care to prevent them from becoming stuck to the double-sided tape. Eggs that were expected to hatch overnight were surrounded with a sheet of Parafilm® (stretched to be as thin as possible) to prevent larvae from falling onto the tape.

Larval rearing

To rear G0 larvae, uninjected control eggs slightly older than the eggs injected on day 5 were placed with ~20 adult Line A workers in a Petri dish with a plaster of Paris floor and maintained at 25°C. These eggs hatched slightly earlier than the injected eggs, priming the workers to rear larvae derived from injected eggs. When the larvae hatched from injected eggs, control larvae were replaced with experimental (injected) larvae. Preliminary trials showed that higher survival was obtained by fostering a minimum of 7 larvae at a time, so control larvae were added to experimental larvae if insufficient experimental larvae were available. The G0 adults reported in this study therefore include an unknown fraction of adults derived from control larvae. Survival of larvae under these conditions was approximately 50%.

Tagging, Genotyping, and Sequencing

Tagging

All ants in behavioral and fitness experiments (Figures 4 and 5) were tagged with two color dots, one on the thorax and one on the gaster, using uni-Paint markers (models PX-20 or PX-21) such that each individual could be identified within the colony (Figures 4D and 4G). For automated behavioral tracking, four colors were used (blue, green, orange and pink) for a total of 16 unique combinations. Ants were tagged with a randomly assigned color pair at least 10 days prior to any behavioral experiments. Tagged ants had a leg removed for genotyping and sequencing either before (Figures 3A, 3B, and 4A–4C) or after (Figures 4D–4I and 5) experiments.

Genotyping

To distinguish Line A and Line B, eggs and adults were genotyped using PCR of mitochondrial *CO1* with standard DNA barcode primers (HCO and LCO, Folmer et al., 1994) followed by a restriction digest with *MwoI* from New England Biolabs. This enzyme cuts the PCR product derived from Line B, but not from Line A.

orco sequencing

To screen for *orco* mutations, we designed PCR primers that flanked the *orco* cut site and sequenced the resulting PCR products using Sanger and Illumina sequencing. Primer sequences were:

F:5'TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGTCCAACTTGCTGTAATTTGGAT3'
R:5'GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGCTCTTCTTGGTCGGCGGTA3'

Illumina methods followed a previously described protocol (Kistler et al., 2015). Primers included tails at the 5' end (underlined) that were used as adapters to add indices to individual samples for Illumina sequencing (Kistler et al., 2015). Sequences were aligned to the *orco* genomic sequence and reads at each base pair that aligned with an insertion or deletion were counted with the script `crisp-align.py` from the `genomepy` package, available at <https://github.com/oxpeter/genomepy>. *orco* amplicons from 25 of 42 recovered G0s were subjected to Illumina sequencing (Figure 1C). Three of these individuals, all of which were found to have nearly 100% mutation rates, displayed a wandering phenotype similar to the wandering phenotype observed in G1s, indicating that somatic CRISPR in G0s may be useful for functional genetic studies even in the many social insect species where it is not logistically possible to generate or maintain stable mutant lines (Schulte et al., 2014; Yan et al., 2014).

Identification of mutant sequences

Mutant lines were identified via Sanger sequencing of eggs and adults of G1s and subsequent generations. All Sanger sequencing traces were scored manually. For the G1 dataset (below), manual identifications were verified by checking for misalignment against a reference sequence using MEGA (Kumar et al., 2016) and by using the program Mutation Surveyor (Softgenetics) for automated allele identification. Mutant lines were defined as groups of ants that possess identical *orco* genotypes, and *orco* amplicons from representatives of each mutant line were Illumina sequenced and individual reads were manually inspected to ensure both alleles were properly identified.

Neuro-anatomy

Immunohistochemistry

Immunofluorescence was performed as described previously (Dobritsa et al., 2003; Pitts et al., 2004). Antennae were removed and prefixed in a solution of 4% paraformaldehyde in PBS with 0.1% Triton (PBSTx) for 30–60 min at room temperature then rinsed three times with PBST. Due to the scarcity of material, antennae were collected from wild-type and *orco*^{-/-} ants that had died less than 24 hr previously. Antennae were stored overnight in a solution of 25% sucrose in PBST at 4°C, then embedded in Tissue-Tek OCT (Sakura Finetek, Torrance, CA) and frozen. Antennae were sectioned at 12 μm at –20°C and allowed to dry at room temperature for 3 hr. Slides were then fixed in 4% paraformaldehyde in PBSTx at room temperature for 30 min, rinsed, then blocked with PBSTx plus 5% normal goat serum (NGS, Vector Laboratories) at room temperature for 60 min. An anti-Orco polyclonal mouse antibody (gift from Vanessa Ruta) was diluted 1:1000 in PBSTx plus 5% NGS and incubated on slides at room temperature overnight. Slides were rinsed and incubated with secondary Alexa594 donkey anti-mouse (Invitrogen) at a 1:500 dilution in PBSTx plus 5% NGS at room temperature for 2 hr then rinsed. Nuclei were labeled using DAPI diluted to 1 μg/mL in PBS at room temperature for 20 min followed by a brief wash in water. Slides were mounted with Dako Fluorescent mounting medium. Images were captured using a confocal microscope (Zeiss LSM 780).

Glomerulus counts and antennal lobe volumes

One of the wild-type *O. biroi* antennal lobe reconstructions was based on published data (McKenzie et al., 2016). For the remaining data, *D. melanogaster* and *O. biroi* brains were dissected in PBS and immediately transferred to a fixative solution of either 1% glutaraldehyde or 2% PFA and 2.5% glutaraldehyde in PBS, and fixed at room temperature on a shaker for 1–30 days. To dehydrate, brains were rinsed in PBS and then suspended for 5 min each in an ascending series of 50%, 70%, 90%, 95%, 100%, 100%, and 100% ethanol. Brains were cleared and mounted in methyl salicylate. Glutaraldehyde-enhanced autofluorescence was imaged using a confocal laser scanning microscope (Zeiss LSM 8800) with excitation by a 488 nm laser. Three-dimensional projections were created from confocal image stacks using Fluorender (Wan et al., 2012). Three-dimensional reconstructions of glomeruli and antennal lobes were produced by manually segmenting confocal image stacks using the Segmentation Editor plugin in the Fiji distribution of ImageJ (Schindelin et al., 2012). Antennal lobe volumes were calculated using the Object Counter3D ImageJ plugin (Bolte and Cordelières, 2006), blindly with respect to genotype.

Fly and ant brains depicted in Figure 3 were immunostained with NC82 and SYNORF1, respectively (both deposited to the Developmental Studies Hybridoma Bank by Buchner, E. (DSHB Hybridoma Products 3C11 (anti SYNORF1) and NC82)). For these stains, brains were fixed at room temperature in 4% paraformaldehyde (with 0.5% triton-x for NC82) for 30min (NC82) or 4hrs (SYNORF1), washed 6x10min in PBS (with 0.5% triton-x for NC82), blocked for 1–2hrs in PBS with 0.5% triton-x and 5% normal goat serum, and incubated in the blocking solution with 1% sodium azide and primary antibody (1:20 in both cases) for three days at room temperature. Brains were then washed 6x10min in PBS and then incubated in blocking solution with 1% sodium azide and secondary antibody (goat anti-mouse Alexafluor647, 1:100) for two days at room temperature. Brains were then washed 3x10min in PBS. Ant brains were subsequently dehydrated in an ascending series of 50%, 70%, 90%, 95%, 100%, 100%, and 100% isopropanol (30 s per solution) and cleared and mounted in methyl salicylate. Fly brains were mounted in Dako fluorescent mounting medium (Agilent Technologies). Brains were imaged as above. Three-dimensional projections were created from confocal image stacks using ZEN software (Zeiss). Flies in this experiment were aged for one month to test whether glomeruli become visibly reduced in old flies.

D. melanogaster glomerulus counts

Our reconstructions of *D. melanogaster* antennal lobe glomeruli yielded different glomerulus numbers than what has been published previously (Table S1). Our reconstructions also showed small differences in glomerulus numbers between wild-type and *orco*^{-/-} flies (Figure 3). To address this possibility, we imaged and reconstructed two additional *D. melanogaster* antennal lobes, one from an *orco*^{-/-} ant and one from a wild-type individual. These reconstructions were not performed strictly de novo, as the *O. biroi* and *D. melanogaster* reconstructions reported in the main text, but by referring to the published map of the *D. melanogaster* antennal lobe (Laissue and Vosshall, 2008; Laissue et al., 1999). Due to differences in sample preparation and imaging methods, it was not possible to unambiguously match each glomerulus to the published map. However, we identified structures in our newly reconstructed wild-type and *orco*^{-/-} antennal lobes that corresponded to all published wild-type glomeruli. These results suggest that *orco*^{-/-} flies have no systematic reduction in the number of antennal lobe glomeruli compared to wild-type flies, although it is possible that some neighboring glomeruli in *orco*^{-/-} flies have divisions that appear less distinct or may even be fused relative to wild-types (Figure 3).

Behavior and Fitness

Sharpie assay

Preliminary trials showed that wild-type ants are strongly repelled by lines drawn with Sharpie permanent markers, frequently without even touching the Sharpie lines (Movie S2). Since *O. biroi* is blind, these results indicate that this is likely an olfaction-mediated behavior. Sharpie assays were conducted with tagged ants (Figures 4A–4C) on printer paper in a 5.25x5.25 in open-air arena bounded by a clear acrylic barrier. Six horizontal and 6 vertical black lines were printed on the paper using an HP LaserJet printer (Figures 4A and 4B). Immediately before each assay, 3 alternating horizontal and vertical black lines were traced with red Fine Point Sharpie Permanent Marker (item 30002). Then the ant was placed in the center of the grid. A 2 min video was recorded and the number of times the ant crossed black and Sharpie lines was manually counted (Movie S2). A Sharpie repulsion index was calculated as the ratio of black line crosses to total line crosses. Once the experiment had concluded, we determined that low numbers of line crosses caused the repulsion indices to be unreliable, and we therefore excluded four assays that had less than 10 line crosses total. As a positive control, a wild-type worker was assayed after each assay with a low repulsion index to ensure the Sharpie lines retained a repulsive effect. All positive controls had high repulsion indices and as a population were statistically indistinguishable from the other wild-type workers assayed ($p = 0.42$, t test).

G1 preparation for behavior and fitness experiments

G1 rearing resulted in a set of 34 colonies containing a mixture of G1 ants and Line A progeny of chaperones. These colonies were used to identify mutants for egg-laying, automated behavioral tracking, and survival experiments (Figures 4D–4I and 5; Table S3). Once each colony started producing eggs, we collected all eggs 5 times over a 14–16 day period. CO1 amplicons from all eggs were genotyped to identify Line A and Line B eggs, and *orco* amplicons from Line B eggs were Sanger sequenced to identify *orco* mutants. We sequenced 2,184 eggs from the 533 Line B ants in these colonies, corresponding to ca. 4 eggs per ant. During the period of egg collection and one week after egg collection had concluded, we subjectively determined whether any individuals in any given colony displayed a wandering phenotype. Colonies in which *orco* mutant eggs were detected or in which wandering phenotypes were observed were selected for the egg-laying dataset.

Egg-laying dataset

We included 16 colonies in the egg-laying dataset. A subset of ants in these colonies were later also used for behavioral and survival experiments (Table S3, see below). After the experiments had been concluded, all ants had a leg removed, from which *orco* amplicons were sequenced. For each *orco*^{wt/-} and *orco*^{-/-} adult we identified, we counted the number of eggs of its genotype in its colony. If several adults of the same genotype were identified in a colony, for each individual we calculated the number of eggs of that genotype divided by the number of adults of that genotype. We used a two-tailed Wilcoxon test to test whether *orco*^{wt/-} or *orco*^{-/-} G1s produced different numbers of eggs than the average of wild-type G1s in this experiment (Figure 5A).

Behavioral and survival dataset

Before removing legs from ants for genotyping, workers from colonies that produced a high frequency of *orco* mutant eggs or contained individuals with wandering phenotypes were pooled to create 5 experimental colonies with a mixture of 12–14 G1 wild-type and *orco* mutant ants. Before pooling, all workers in these colonies were individually tagged with two color dots. These 5 colonies were recorded in 24 hr videos.

Experimental *O. biroi* colonies initially contained a total of 68 G1 ants, with 42 wild-type, 8 *orco*^{wt/-}, and 14 *orco*^{-/-} individuals (Figure 1D). These colonies also contained 4 *orco* mutant individuals with in-frame mutations, which were not included in the current analyses because sample sizes were too small. G1s in experimental colonies varied in *orco* genotype but were otherwise identical in rearing methods, genetic background, and did not differ systematically in age. Before the start of each 24 hr video, plaster nest boxes were cleaned and the plaster was moistened. For four weeks after the video was recorded, we also recorded survival of all ants in the 5 experimental colonies.

Video recording and automated behavioral tracking

Automated behavioral tracking was performed in custom-made tracking setups under constant illumination. Temperature was maintained at 25°C. Videos were acquired using C910 Logitech USB webcams controlled with custom MATLAB (version R2016a, The MathWorks, Inc.) software at 10 fps at 960x720 pixel resolution (13 pixels/mm).

Tracking was performed blind with respect to genotype. Videos were processed and analyzed using custom MATLAB software. In each frame, ants were segmented from the background of the dish using a fixed threshold. Resulting components, or blobs, were linked into trajectories using the optical flow computed between consecutive frames (Horn–Schunck method (Barron et al., 1994)). Trajectories ended and new ones were initiated whenever blobs split or merged between two consecutive frames. Trajectories stored the following data, collected from the respective blob in each frame: centroid (position of center of mass), orientation (angle between the major axis of the best-fitting ellipse and the horizontal axis) and area (in mm²).

We used a threshold size to select trajectories that corresponded to a single ant and lasted longer than two seconds. Each trajectory was then assigned a combination of color tags using a custom classification algorithm. For each experiment, at least 500 frames per ant were manually identified to create a training set, 70% of which was used to train, and 30% of which was used to validate an automated identity classifier.

For each trajectory, a naive Bayes color classifier (Fletcher et al., 2011) was used to compute the pixel color probabilities for each pixel in the blob of each frame for six color classes: the four tag colors, the ant cuticle color and the color of the plaster of Paris (Figure S3). Predicted probabilities for all four tag colors were used to determine whether both tags were visible and, if so, the orientation of the ant in the frame was deduced from the relative position of the tags with respect with the cuticle color. If both procedures were successful, the pixel color probabilities were fed into another naive Bayes classifier to assign an identity to the ant.

For each trajectory, frames were tested in a random order until 20 frames were successfully identified or no more frames were available. If at least one frame was identified, the trajectory, and thus all its underlying positions, was assigned the most frequently predicted identity. The identity classifier had an empirical error rate of 20% on single frames. However, the error rate decreased with the number of frames tested within a trajectory. Overall, we estimate that less than 10% of the total identified positions were misclassified, a performance equivalent to that reported previously for a functionally similar tracking setup for ants (Mersch et al., 2013).

Analysis of automated behavioral tracking data

Cleaning and watering the nests at the beginning of each experiment caused the ants to establish pheromone trails and actively move around the dish. We used this initial period of high activity to measure trail following behavior, reasoning that trail pheromones may cause two or more ants to move along the same path within the dish. *O. biroi* is a blind species and experiments from our laboratory indicate that their behavior is not influenced even by very bright light (A. Gal, personal communication), so it is unlikely that light or landscape cues cause correlation in the movement of the ants. We have developed assays causing *O. biroi* to develop trails both in response to food and disturbance (A. Gal, personal communication), but we chose to use the disturbance-induced trails for this particular analysis because they are more robust.

For each ant, we measured the correlation between its own movement and the movement of the remaining ants in the colony during the first hour of the tracking experiment. As *O. biroi* have a tendency to walk along the edges of the dish, we discarded segments of trajectories close to the edge of the dish and included only segments where the ants moved faster than 1 mm/s continuously for at least 1 s and without contact with any other ant. We then computed a 2-D histogram, or density map, for each ant by counting the number of times one of the remaining positions in the 970x720 pixel original image fell into each bin of a 120x90 bin grid. We then computed the Pearson correlation coefficient between the density map of each ant and a density map constructed from the trajectories of all ants in the colony but the focal ant (Figure 4E). For each ant, the actual correlation coefficient provided an estimate of the

correlation of movement of that ant with the actual movement of other ants in the colony, which presumably results from following pheromone trails.

As a baseline comparison, the density map of each ant was correlated with a randomized density map constructed by rotating the trajectories of all ants but the focal ant in the colony by a random angle around the center of the dish (Figure 4E). This randomized correlation coefficient provided an estimate of the correlation of the movement of that ant with the randomized movement of other ants in the colony. This residual correlation reflects the portion of the correlation that is due to non-local effects such as turning frequencies, linear and angular velocity dynamics, and radial preference for certain regions of the petri dish. After examining the data, two experimental colonies were excluded from the trail following analysis because they did not form clear trails during the videos, resulting in Pearson correlation coefficients of approximately zero for all ants in the colony.

To measure wandering phenotypes, in each experiment we calculated the total distance traveled by each ant over the 24 hr video by computing the distances between all pairs of successive positions in meters in all identified trajectories. Time without contact was calculated as the total time each ant was identified in each experiment (since ants were only identifiable when they were separated from other ants). This provides a minimum estimate of the time without contact for each ant, given that it was also possible for ants to be spatially separated from other ants, yet unidentifiable, for example if their posture did not allow the detection of both tags.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical details of all experiments can be found in the figure legends. Behavioral tracking and antennal lobe volume measurements were performed blindly with respect to genotype. Other analyses were not performed blindly with respect to genotype. Mixed model statistics were performed in R v 3.3.1 using the *lmer* function in the lme4 library as described previously (Bates et al., 2014; Ulrich and Schmid-Hempel, 2012, 2015). All other statistics were performed using GraphPad Prism 7. Normality was determined by D'Agostino-Pearson normality tests. Datasets used for ANOVA analyses had equal variance across treatments. Single groups were compared against a predicted mean using two-tailed Wilcoxon tests. Proportional data were compared between treatments using a Fisher exact test. Unpaired two-tailed Student's t tests or paired Wilcoxon tests were used to compare two groups, when appropriate, and two-way ANOVAs followed by Tukey's multiple comparisons test were used to compare more than two groups.

DATA AND SOFTWARE AVAILABILITY

Raw data for all experiments are available in Table S3. Raw illumina sequencing reads are available through the National Center for Biotechnology Information Short Reads Archive. gRNA design and sequencing analyses were performed using the script *crispr-align.py* from the *genomepy* package, available at <https://github.com/oxpeter/genomepy>.

Supplemental Figures

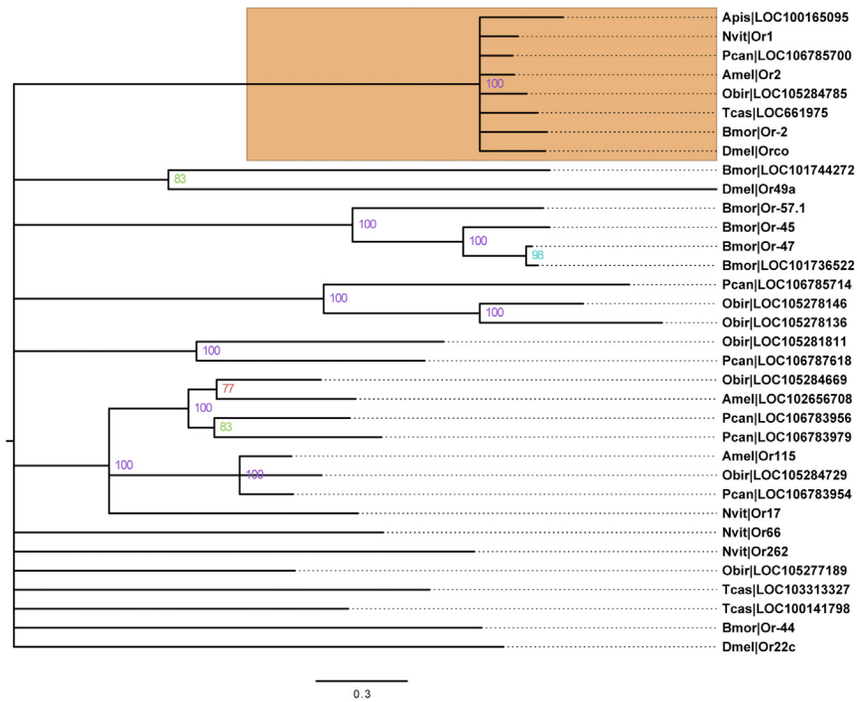
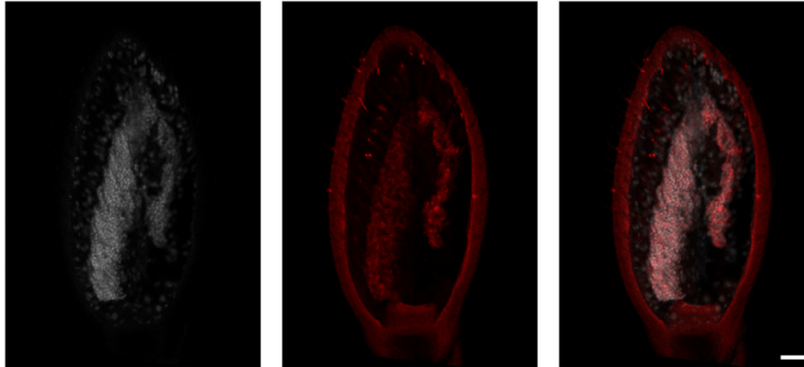
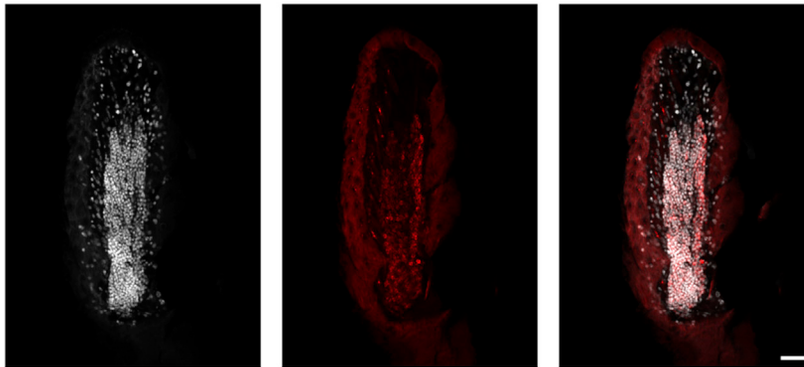
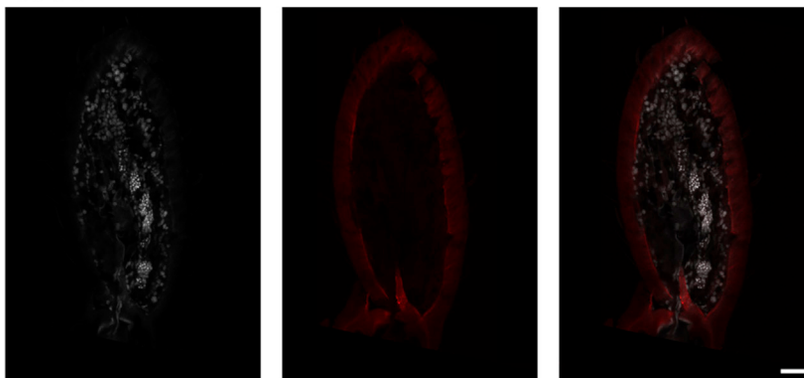


Figure S1. Phylogeny of *orco* and Other Odorant Receptor Genes in Insects, Related to Figure 1

orco orthologs in red. *O. biroi orco* is included in a clade with the *orco* genes from all other studied insect species (100% bootstrap support). All nodes with less than 75% bootstrap support have been collapsed for clarity. All other bootstrap values are shown at the respective node. Species are indicated by a four letter code: Apis – *Acyrtosiphon pisum*; Nvit – *Nasonia vitripennis*; Pcan – *Papilio canadensis*; Amel – *Apis mellifera*; Obir – *Ooceraea biroi*; Tcas – *Tribolium castaneum*; Bmor – *Bombyx mori*; Dmel – *Drosophila melanogaster*. NCBI gene symbols are shown on the right. The scale bar indicates an average of 0.3 substitutions per site.

A wild-type**B** *orco*^{wt/-}**C** *orco*^{-/-}

DAPI

Orco

Merge

Figure S2. Immunohistochemistry of Antennal Sections from Additional Wild-Type and *orco* Mutant Ants, Related to Figure 2

(A) Antennal section of wild-type ant showing DAPI counterstain (gray), Orco immunostain (red), and the merged image (black).

(B) Antennal section of *orco*^{wt/-} ant, showing similar patterns of DAPI and Orco staining as wild-type.

(C) Antennal section of *orco*^{-/-} ant, showing reduction in DAPI staining and loss of Orco staining, indicating that Orco-positive OSNs are absent in *orco*^{-/-}, but not wild-type or *orco*^{wt/-}, *O. biroi*. Scale bars are 20 μ m.

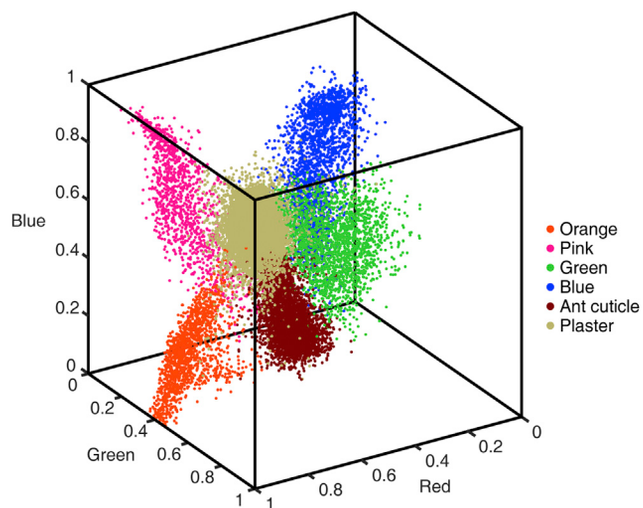


Figure S3. Color Definition for Automated Behavioral Tracking in RGB Space, Related to Figure 4

Measurements were collected manually from the four different tag colors (orange, pink, green and blue), the plaster (background) and the ant cuticle. These colors are clearly distinguishable in RGB space, allowing automated behavioral tracking based on identifying the color tags of each ant.