Multimodal Integration of Carbon Dioxide and Other Sensory Cues Drives Mosquito Attraction to Humans

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SUMMARY

Multiple sensory cues emanating from humans are thought to guide blood-feeding female mosquitoes to a host. To determine the relative contribution of carbon dioxide (CO₂) detection to mosquito hostseeking behavior, we mutated the AaegGr3 gene, a subunit of the heteromeric CO₂ receptor in Aedes aegypti mosquitoes. Gr3 mutants lack electrophysiological and behavioral responses to CO₂. These mutants also fail to show CO2-evoked responses to heat and lactic acid, a human-derived attractant, suggesting that CO2 can gate responses to other sensory stimuli. Whereas attraction of Gr3 mutants to live humans in a large semi-field environment was only slightly impaired, responses to an animal host were greatly reduced in a spatial-scale-dependent manner. Synergistic integration of heat and odor cues likely drive host-seeking behavior in the absence of CO₂ detection. We reveal a networked series of interactions by which multimodal integration of CO₂, human odor, and heat orchestrates mosquito attraction to humans.

INTRODUCTION

Anthropophilic female mosquitoes possess a strong innate drive to find and blood-feed on humans, whose blood provides a protein source needed for egg development. As a consequence of such host-seeking behavior, many mosquito species including the yellow fever mosquito $Aedes\ aegypti$ and the African malaria mosquito $Anopheles\ gambiae$ efficiently spread blood-borne diseases such as dengue and malaria between humans. To orient toward hosts, female mosquitoes detect a variety of chemical and physical cues including body odor, CO_2 , moisture, heat, and visual contrast (reviewed in Gibson and Torr, 1999). Of these sensory cues, the $\sim 4\%\ CO_2$ exhaled in breath is a potent behavioral activator and attractant for mosquitoes and is often considered to be the most important sensory cue used by these disease vectors to find humans (Gillies, 1980).

In a broad range of mosquito species, CO₂ elicits a suite of context-dependent effects (reviewed in Gillies, 1980) that are thought to promote host-seeking behavior. Exposure to CO₂ alone strongly activates mosquito flight, increasing the probability of flight take-off and locomotor activity in the absence of directional air currents (Eiras and Jepson, 1991). Filamentous intermittent plumes of CO₂ (Dekker et al., 2005; Geier et al., 1999; Healy and Copland, 1995) elicit stereotyped and sustained patterns of upwind flight toward the CO₂ source (Dekker and Cardé, 2011).

In Ae. aegypti, CO₂ increases both the kinetics and fidelity of flight responses to host odor plumes, thereby enhancing source finding (Dekker and Cardé, 2011; Dekker et al., 2005). Several studies have also shown that CO₂ enhances mosquito attraction to warmth (Burgess, 1959; Kröber et al., 2010; Maekawa et al., 2011). Despite earlier attempts to quantify the overall effect of exhaled CO₂ from breath on attraction to live humans (Snow, 1970), the scale over which CO₂ acts and its relative importance compared to all the other sensory cues encountered during mosquito orientation toward a live host remain unclear.

Ae. aegypti maxillary palps house a specialized class of olfactory sensory neurons (OSNs) that are exquisitely sensitive to changes in CO2 concentration (Grant et al., 1995). In the vinegar fly Drosophila melanogaster, two gustatory receptors (DmelGr21a and DmelGr63a) function together to mediate CO₂ detection (Jones et al., 2007; Kwon et al., 2007). Direct orthologs of DmelGr21a and DmelGr63a have been identified in a wide range of insect species including Ae. aegypti, where they are named Gr1 and Gr3, respectively (Robertson and Kent, 2009). In most non-Drosophila insect species examined, a Gr1 paralog named Gr2 is present (Figure 1A). Functional studies of An. gambiae Gr1, Gr2, and Gr3 orthologs using heterologous expression in Drosophila (Lu et al., 2007) and transient knockdown studies in Ae. aegypti (Erdelyan et al., 2012) have suggested a conserved role for these genes as candidate CO2 receptors, but the functional requirement of one or more subunits for mosquito CO₂ detection is unknown.

Here we used genome editing to engineer *Ae. aegypti* unable to sense CO₂, allowing us to uncover the relative contribution of CO₂ detection to mosquito host-seeking behavior. We find that attraction to heat and lactic acid are absent in *Gr3* mutants, suggesting that CO₂ detection gates responses to these other



sensory cues. Despite these dramatic deficits, Gr3 mutants show only mild impairment in their attraction to live human subjects in semi-field conditions. Using live animals, we demonstrate that effects of CO_2 detection on mosquito host-seeking behavior are dependent on spatial scale. We further provide compelling evidence that mosquito attraction to humans and blood-feeding behavior are driven by multimodal integration of CO_2 and other host sensory cues.

RESULTS

Generation of Targeted Mutations in Gr3

Of the three Ae. aegypti CO₂ receptors (Figure 1A), we chose to mutate Gr3 because mutations in the fly ortholog, DmelGr63a, were sufficient to disrupt CO₂ reception (Jones et al., 2007). We generated null mutations in this gene by injecting a zinc-finger nuclease (ZFN) pair targeting exon 3 of Gr3 (Figure 1B) into preblastoderm stage mosquito embryos. Two independent Gr3 alleles were isolated and characterized, Gr3⁴ and Gr3^{ECFP}. Gr3⁴ harbors a 4 bp deletion in exon 3 generated by nonhomologous end-joining (Figure 1C), leading to a frameshift and multiple nonsense mutations. We also generated a homologous recombination allele Gr3^{ECFP} (Figure 1D) by coinjecting a donor plasmid for homology-directed repair along with the ZFN mRNAs. In this allele, 33 bp of exon 3 were deleted, and a 2.45 kb cassette was inserted via ZFN-mediated homologous recombination. This insertion marked mutants visually with enhanced cyan fluorescent protein (ECFP) fluorescence (Figure 1E) and could be detected in the Gr3 locus by PCR (Figure 1F). After outcrossing each isolated allele to wild-type mosquitoes for at least five generations, we established heterozygous ($Gr3^{4/+}$ and $Gr3^{ECFP/+}$), homozygous ($Gr3^{4/4}$ and $Gr3^{ECFP/ECFP}$), and heteroallelic $(Gr3^{EC\dot{F}P/4})$ mutant lines for behavioral characterization.

${\it Gr3}$ Mutants Cannot Sense or Respond Behaviorally to ${\it CO}_2$

CO2 is detected by OSNs of large-spiking amplitude that are housed within capitate peg sensilla on the Ae. aegypti maxillary palp, along with a small-spiking neuron responsive to 1-octen-3-ol and a third neuron of unknown ligand specificity (Figure 1G) (Grant et al., 1995; McIver, 1972). Previous studies in An. gambiae demonstrated that Gr1, Gr2, and Gr3 are coexpressed in the CO₂-responsive OSN, and Or8/orco are coexpressed in the 1-octen-3-ol-responsive OSN (Jones et al., 2007; Lu et al., 2007) (Figure 1G). The organization and function of these OSNs are believed to be conserved in Ae. aegypti (Bohbot et al., 2007). To determine whether Gr3 mutants could sense CO₂, we performed extracellular recordings of CO₂-evoked activity from female capitate peg sensilla using single sensillum electrophysiology. Wild-type and heterozygous female mosquitoes responded robustly to a 1 s pulse of 0.5% CO2, whereas homozygous and heteroallelic mutants were totally unresponsive (Figure 1H). In the mutants, this lack of a CO₂-evoked response was accompanied by the complete absence of spontaneous activity in this neuron. In contrast, the 1-octen-3-olsensitive cell of all genotypes responded similarly to a 1 s pulse of 10⁻⁶ (R)-1-octen-3-ol (Figure 1I), indicating that our extracellular recordings were targeted correctly to capitate peg sensilla.

Summary data showing spontaneous and stimulus-evoked activity are in Figure 1J.

To test whether CO₂-evoked behavior was affected in these mutants, we tracked the flying activity of groups of female mosquitoes using 3D multi-insect tracking in a large assay chamber. Following a period of baseline tracking, we introduced a pulse of CO₂ into the airstream, which caused large bursts of flight activity in wild-type and heterozygous female mosquitoes (Figures 2A–2C; top three rows). In contrast, homozygous and heteroallelic mutant mosquitoes remained indifferent to this stimulus (Figures 2A–2C; bottom three rows).

To rule out nonspecific locomotor defects in Gr3 mutants, we tracked the activity of wild-type mosquitoes and a representative homozygous mutant genotype, $Gr3^{ECFP/ECFP}$, over a 23 hr period. No significant differences in either flying distance tracked per mosquito or flight velocity were observed between genotypes, and their daily rhythms of activity were qualitatively similar (Figures 2D and 2E). We conclude that Gr3 mutants are selectively impaired in their response to CO_2 .

CO₂ Detection Can Gate Mosquito Heat-Seeking Behavior

Warm-blooded hosts provide mosquitoes with thermal contrast that facilitates the localization of a suitable blood meal. To investigate the potential for an interaction between CO₂ detection and heat perception, we developed a heat-seeking assay that scored the ability of mosquitoes to land on a 37°C target (Figures 3A–3C). Trials with wild-type females revealed that mosquitoes only landed on the heated target when a pulse of CO₂ was presented in the assay (Figures 3D–3F). Wild-type males were not attracted using identical assay conditions (Figure 3G), showing that heat seeking stimulated by CO₂ is sexually dimorphic in *Ae. aegypti*.

To further verify that CO₂ detection was required for heatseeking behavior in female *Ae. aegypti*, we tested the heterozygous, homozygous, and heteroallelic *Gr3* mutants in the same assay. Heterozygous genotypes landed on the heated target in a CO₂-dependent manner (Figures 3H and 3I), but homozygous and heteroallelic *Gr3* mutants did not land (Figures 3J–3L), indicating that CO₂ detection is required for this behavior. Heatseeking data are summarized in Figures 3M and 3N.

CO₂ Detection Enhances Mosquito Attraction to Lactic Acid and Human Scent

In addition to exhaled CO_2 , humans emit a blend of hundreds of volatile organic compounds in breath and skin odor (Gallagher et al., 2008). To evaluate whether CO_2 detection influences the ability of $Ae.\ aegypti$ to orient toward human odorants, we used a two-port olfactometer to assay attraction of wild-type, heterozygous, and homozygous Gr3 mutant female mosquitoes to various odor stimuli in the presence or absence of supplemental CO_2 (Figure 4A). After characterizing background levels of attraction using blank trials that contained no stimuli (Figure 4B), we evaluated responses to CO_2 . We observed moderate numbers of wild-type and heterozygous Gr3 females trapped in response to stimulation with CO_2 alone. However, attraction appeared to be nondirectional with respect to the CO_2 source in this assay (Figure 4C, $Gr3^{+/+}$ and $Gr3^{ECFP/+}$). This general

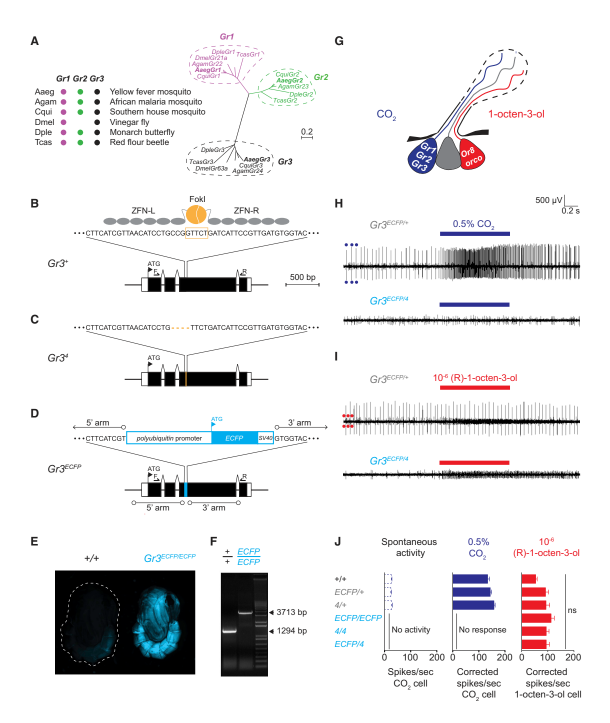


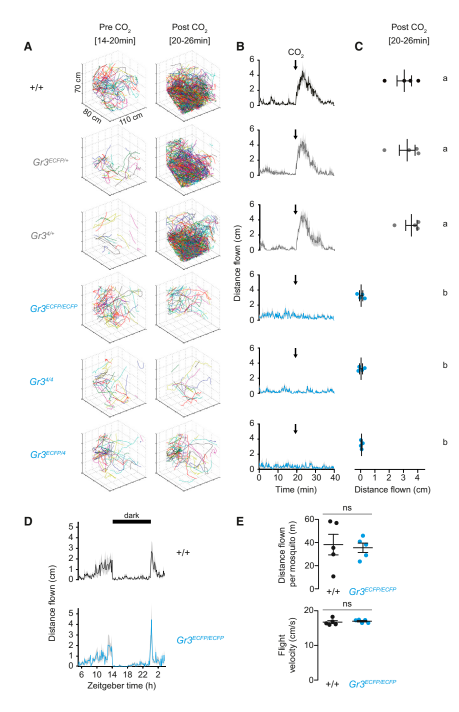
Figure 1. Electrophysiological Characterization of Gr3 Mutants Generated by ZFN Gene Editing

(A) Phylogenetic comparison of insect CO2 receptor genes. Aaeg: Aedes aegypti, Agam: Anopheles gambiae, Cqui: Culex quinquefasciatus, Dmel: Drosophila melanogaster, Dple: Danaus plexippus, Tcas: Tribolium castaneum. Branch lengths are proportional to sequence divergence. Scale bar, 0.2 amino acid replacements per site.

- (B) The Ae. aegypti Gr3 genomic locus, with a schematic of the Gr3 ZFN pair binding to exon 3 DNA and the Fok1 cleavage site indicated in orange.
- (C) Diagram of 4 bp deletion in the mutant *Gr3*⁴ genomic locus.
- (D) Diagram of the mutant *Gr3^{ECFP}* genomic locus.
- (E) Representative fluorescent images of female wild-type and Gr3^{ECFP/ECFP} mutant pupae.
- (F) PCR products generated across the ECFP insertion site using the F and R primers indicated by small arrows in (B) and (D).
- (G) Diagram of receptor expression and ligand specificity of maxillary palp capitate peg sensillum OSNs.

 (H) Representative spike traces during extracellular recordings from a *Gr3*^{ECFP/+} heterozygous (top) and *Gr3*^{ECFP/4} mutant (bottom) capitate peg sensillum evoked by 0.5% CO₂. Dark blue circles mark large amplitude spikes from the CO₂-sensitive OSN. Stimulus bar (blue): 1 s.

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increase in capture rate in both ports may reflect random exploratory behavior of females in the two-port assay chamber upon activation with CO2 in the absence of other olfactory cues. This contrasts with the known attraction of mosquitoes to CO2 in a

Figure 2. Ae. aegypti Gr3 Mutants Are Not Activated by CO2 and Retain Normal Locomotor Behavior

(A-C) CO₂-evoked flight activity of groups of female Ae. aegypti wild-type, heterozygous, and mutant mosquitoes tracked for 20 min before and after a 40 s pulse of CO₂ (n = 4 replicates per genotype; n = 20 mosquitoes/trial).

(A) Representative 3D tracks of the indicated genotypes in the 6 min pre- (left) and post-application (right) of a 40 s pulse of CO2.

(B) Distance flown (cm/s/animal) in response to stimulation with CO₂ (black arrow). Data are shown as mean (solid line) ± SEM (grav shading).

(C) Mean distance flown per second for mosquitoes post-application of CO₂ (t = 20-26min). Each replicate is indicated by a dot, and mean \pm SEM as bars. Variation among genotypes was significant (oneway ANOVA, p < 0.0001). Genotypes marked with different letters are significantly different by post hoc Tukey's HSD test (p < 0.001).

(D) Locomotor activity of wild-type and Gr3^{ECFP/ECFP} mutant mosquitoes measured as cm/min/animal over a 23 hr period (dark period indicated by black bar) without supplemental CO2 (n = 5 replicates per genotype; 20 females per replicate). Data are shown as mean (solid line) ± SEM (gray shading).

(E) Total distance flown per mosquito and average flight velocity of data in (D). Each replicate is indicated by a dot, and mean ± SEM as bars (ns, not significant; t test, p > 0.05).

wind tunnel (Dekker and Cardé, 2011), likely because CO2 is not presented as a filament in our two-port assay. Responses to CO₂ alone were abolished in Gr3^{ECFP/} ECFP mutants, consistent with their inability to sense this stimulus (Figure 4C).

We next evaluated responses to L-(+)-lactic acid, a volatile organic compound present in human breath and skin odor, which has previously been shown to synergize behaviorally with CO₂ to enhance mosquito attraction (Acree et al., 1968; Eiras and Jepson, 1991). L-(+)-lactic acid was not attractive to female mosquitoes when presented alone (Figure 4D). However, when L-(+)-lactic

acid was presented with CO2, many wild-type and heterozygous females were attracted. Gr3^{ECFP/ECFP} mutants did not respond in these same trials, indicating that CO2 detection is required for synergistic responses to lactic acid (Figure 4E).

⁽I) Representative spike traces from heterozygous and mutant basiconic sensilla evoked by 10⁻⁶ (R)-1-octen-3-ol. Red circles mark small amplitude spikes from the 1-octen-3-ol-sensitive OSN. Stimulus bar (red): 1 s.

⁽J) Summary of spontaneous activity and CO2- and (R)-1-octen-3-ol-evoked responses in the indicated genotypes. Data are presented as mean ± SEM. Genotypes did not differ in their response to (R)-1-octen-3-ol (one-way ANOVA, p = 0.07; ns, not significant; n = 8-16 sensilla recorded from at least three mosquitoes per genotype).

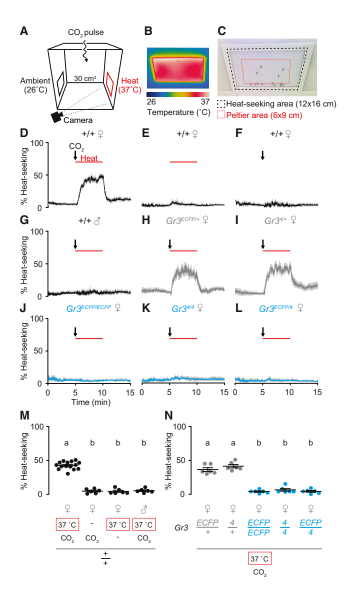


Figure 3. ${\rm CO_2}$ Detection Gates Heat-Seeking Behavior in Female Ae. ${\it aegypti}$

- (A) Schematic of the mosquito heat-seeking assay.
- (B) Thermal image of target Peltier heated to 37°C.
- (C) Image indicating Peltier location (red outline) and area scored for heat seeking (black dashed line).
- (D–L) Mosquitoes of the indicated sex and genotype seeking an ambient or heated target, with or without a pulse of CO_2 . The 5 min 37°C stimulus is indicated by the red bar. The 20 s pulse of CO_2 is indicated by the black arrow. Data are shown as mean (solid line) \pm SEM (gray shading) (n = 6 trials, except D, where n = 15; 20–25 mosquitoes/trial).

(M and N) Quantification of heat seeking during minutes 7–10 of assay. Each replicate is indicated by a dot, and mean \pm SEM as bars. Variation among both stimulus regimes and mosquito genotypes was significant (one-way ANOVA, p < 0.0001 for both M and N). Data labeled with different letters are significantly different using Tukey's HSD test comparing all pairs of means (p < 0.001).

To evaluate whether CO₂ detection modulates perception of human skin odor, we examined mosquito responses to human odor collected on nylon sleeves. Human odor in the absence

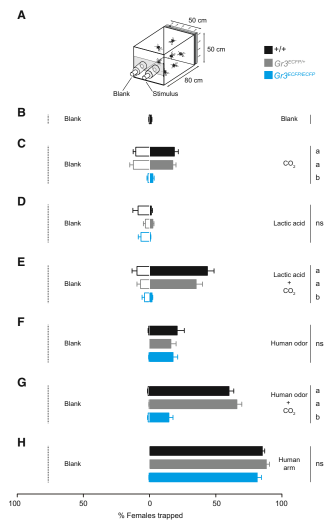


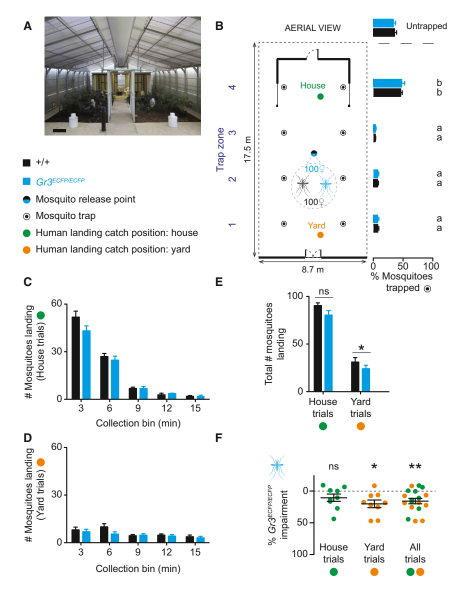
Figure 4. CO₂ Synergizes with Host Odor to Drive Mosquito Attraction

(A) Diagram of the two-port olfactometer.

(B–H) Response of the indicated genotypes to a choice of the blank port compared to the stimulus indicated at the right. Data are plotted as mean \pm SEM (n = 8–11 trials per condition; n = 50 mosquitoes per trial). Significance was assessed with one-way ANOVA. Genotypes marked with different letters are significantly different by Tukey's HSD test (p < 0.0001; ns, not significant).

of CO_2 elicited moderate attraction that was not significantly different among genotypes (Figure 4F). However, supplementation of CO_2 to human odor enhanced levels of attraction observed for wild-type and heterozygous females, but not homozygous mutants (Figure 4G).

To assess a more complex multisensory stimulus, we tested mosquito attraction to a live human arm inserted into the stimulus port. Very high levels of attraction were observed for all mosquito genotypes to a live human arm in this assay (Figure 4H). We speculate that additional stimuli presented by the human arm, including odorants not trapped on nylon sleeves, moisture, heat, and visual cues, overcome the behavioral defect seen with simpler stimulus combinations in the mutants.



Mosquito Attraction to Humans in a Semi-Field Environment Is Diminished but Not Abolished in Gr3 Mutants

To obtain a more biologically relevant estimate of the contribution that CO₂ detection makes to mosquito host attraction, we tested the ability of Gr3 mutants and wild-type females to locate and land on live human volunteers in the James Cook University semi-field cage in Northern Queensland, Australia (Figure 5A). This semi-field cage contained the underside of an elevated Queenslander style house, including furniture used as resting sites of Ae. aegypti, and a lush tropical backyard that mimicked the native habitat of Ae. aegypti in this region (Ritchie et al., 2011). Equal numbers of wild-type and mutant females were released into the semi-field cage during each trial for simultaneous comparisons of mosquito host-seeking behavior.

To examine the distribution of the released mosquito population over the semi-field cage without humans present, we used

Figure 5. Gr3 Mutants Have Diminished Responses to Live Humans in a Semi-Field Environment

(A) Interior view of the semi-field cage. Scale bar. 0.5 m.

(B) Schematic of the semi-field cage showing the average distribution of an equal mix of wild-type and $Gr3^{ECFP/ECFP}$ mutant female mosquitoes (n = 100 per genotype) in the cage in the absence of humans (n = 5 trials). Distribution varied significantly according to trap zone but not genotype (two-way ANOVA, p < 0.0001 for trap zone and p = 0.773 for genotype). Different letters indicate significantly different means by post hoc Tukev's HSD test.

(C and D) Number of mosquitoes of the indicated genotype landing on human subjects in 3 min bins during 15 min at house (C) or yard (D) position (n = 9 trials per position).

(E) Total number of mosquitoes landing in humanlanding catch trials with human volunteers at house or vard position. Paired t tests were used to assess statistical significance in comparisons between genotypes (ns, not significant; p = 0.09; *p = 0.018). (F) Relative impairment in host-seeking of mutants compared to wild-type mosquitoes (one-sample t test relative to zero; ns, not significant; p = 0.105; *p = 0.011; **p = 0.002). In (B)–(F), all data are plotted as mean \pm SEM.

mosquito traps to capture females in four designated trap zones (Figure 5B, left). The majority of the mosquitoes of both genotypes were trapped near the house, and there were no differences between the numbers of each mosquito genotype trapped in each zone (Figure 5B, right). Therefore, under semi-field conditions, both wild-type and Gr3 mutant mosquitoes retained an innate preference for human habitation. The mutants showed similar dispersal across the semi-field cage relative to wild-type mosquitoes,

consistent with their performance in flight-tracking assays (Figures 2D and 2E).

To test mosquito attraction to humans, we counted the number of wild-type and mutant mosquitoes landing on a seated volunteer positioned within either the house or yard of the semi-field cage (Figure 5B, green and orange circles, respectively). The mosquito traps were inaccessible during humanlanding catch trials. Slightly fewer Gr3 mutants landed on humans compared to wild-type mosquitoes during the initial two collection bins of house and yard trials (Figures 5C and 5D). These small cumulative differences were maintained over the duration of each assay, leading to fewer Gr3 mutants being collected in total on average compared to wild-type mosquitoes at either human-landing catch position (Figure 5E). Overall, landing varied significantly according to genotype (p = 0.043) and position in the cage (p < 0.0001), but a two way-ANOVA showed that there was no interaction between these two variables (p = 0.713). Variation in mosquito landings between

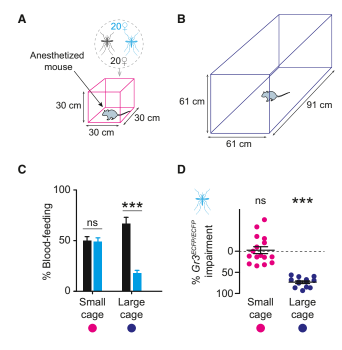


Figure 6. Spatial Scale-Dependent Responses of *Gr3* Mutants to Live Animal Hosts

(A and B) Diagrams of small (A) and large (B) cage assays. Mice not to scale. (C and D) Percent blood feeding of the indicated genotype (C) and impairment of Gr3^{ECFP/ECFP} mutants relative to wild-type (D).

Significance was assessed in (C) with paired t tests and in (D) with a one-sample t test relative to zero (ns, not significant; ***p < 0.0001).

All data are plotted as mean \pm SEM (small cage: n = 16 trials; large cage: n = 13 trials; n = 20 mosquitoes/genotype/trial).

human volunteers was not significant (data not shown). Overall, $Gr3^{ECFP/ECFP}$ mosquitoes were impaired by approximately 15% relative to wild-type mosquitoes in the semi-field cage (Figure 5F). Thus, a loss of CO_2 detection diminishes but does not abolish the ability of mosquitoes to find humans in the semi-field cage.

Effects of CO₂ Sensation on Mosquito Host-Seeking Behavior Are Dependent on Spatial Scale

Host sensory cues such as heat, CO₂, human odor, moisture, and visual contrast are most potent when a mosquito is close to the human host and decline at a distance. We therefore reasoned that the host-seeking defects in *Gr3* mutants would depend on spatial scale, such that a more dramatic defect would be apparent if the mutants were tested in a larger semi-field cage. Because to our knowledge no larger semi-field facilities currently exist, we modeled host-seeking behavior at two different spatial scales with live animal hosts in laboratory cages of different sizes (Figures 6A and 6B).

In these assays, mixed populations of equal numbers of wild-type and $Gr3^{ECFP/ECFP}$ mosquitoes were released into each cage and assayed for their ability to blood-feed on an anesthetized mouse. Similar numbers of wild-type and $Gr3^{ECFP/ECFP}$ females blood-fed during trials in the small (0.027 m³) cage, but significantly fewer Gr3 mutant females blood-fed relative to wild-type

mosquitoes in the larger (0.339 m³) cage (Figure 6C). On average, blood-feeding rates in $Gr3^{ECFP/ECFP}$ females were unimpaired in the small cage consistent with semi-field cage results but impaired by approximately 70% relative to wild-type mosquitoes in the larger cage (Figure 6D). Two-way ANOVA on blood-feeding rates revealed a highly significant interaction (p < 0.0001) between mosquito genotype and cage size, suggesting that the effects of CO_2 sensation on host-seeking behavior are dependent on spatial scale. We note that presence of wild-type mosquitoes did not rescue the major deficits in host seeking seen in $Gr3^{ECFP/ECFP}$ mutants in the large cage (Figure 6C). This rules out the possibility that motor, auditory, or pheromonal cues from wild-type mosquitoes influence the behavior of mutant mosquitoes in this assay.

Sensory Stimuli Synergize in Multiple Binary Combinations to Drive Mosquito Blood-Feeding Behavior

Given that CO_2 is a major host-seeking cue and strongly influences mosquito perception of heat and odorants, we were intrigued by the nearly normal attraction of homozygous $\mathit{Gr3}$ mutants to live humans in a semi-field environment. To ask how the phenotypic deficits of $\mathit{Gr3}$ mutants may be compensated during attraction to live hosts, we developed an artificial membrane-feeding assay (Figure 7A) to test the contribution of individual and combined sensory cues including CO_2 , heat, and human odor to mosquito blood-feeding behavior.

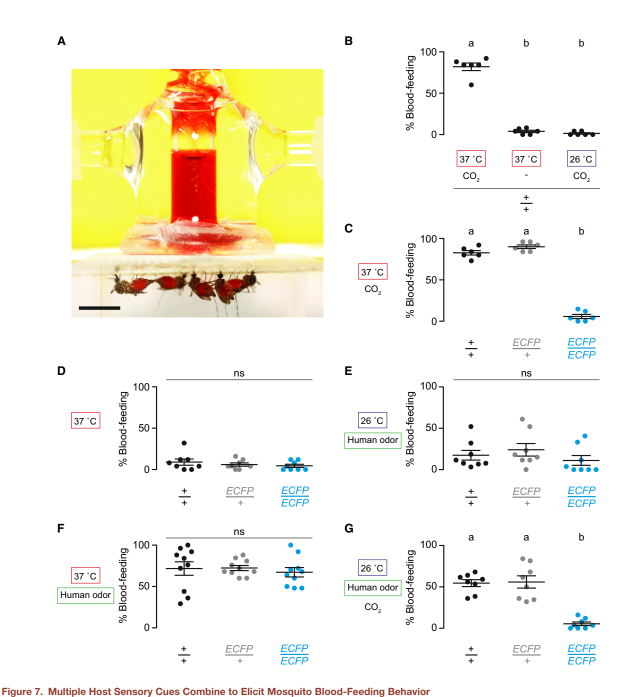
To test whether heat and CO_2 are integrated to drive blood-feeding behavior, we manipulated the temperature of blood inside the membrane feeder to reflect human body (37°C) or ambient (26°C) temperatures in the presence or absence of CO_2 . Wild-type and heterozygous mosquitoes blood-fed from membrane feeders only when the blood was heated to 37°C and CO_2 was added to the assay (Figure 7B). Homozygous $\mathrm{Gr3}$ mutants rarely blood-fed under these conditions (Figure 7C).

We then tested whether human odor presented with heat can rescue blood feeding in *Gr3* mutants. Very few females of any genotype blood-fed from a heated membrane feeder in the absence of other cues (Figure 7D). Human odor when applied alone to a membrane feeder filled with ambient-temperature blood also elicited low levels of blood feeding (Figure 7E). However, when we applied human odor to the membrane of heated feeders, we observed high rates of blood feeding in all genotypes (Figure 7F).

To determine whether CO₂ and odor cues can also synergize to drive blood feeding in the absence of heat, we applied human odor to a membrane feeder maintained at ambient conditions and added CO₂. High numbers of wild-type and heterozygous mosquitoes, but very few homozygous *Gr3* mutants, blood-fed on this room-temperature blood (Figure 7G). Taken together, these results indicate that CO₂, odor, and heat synergize in all possible binary combinations to drive *Ae. aegypti* blood-feeding behavior.

DISCUSSION

Here we used ZFN-mediated targeted mutagenesis and homologous recombination to target the Ae. aegypti Gr3 gene to



(A) Image of female Ae. aegypti mosquitoes blood feeding from a glass membrane feeder. Scale bar, 0.5 cm. (B) Blood-feeding responses of wild-type mosquitoes with different blood temperatures and ± CO2. (C-G) Blood-feeding responses of the indicated genotypes with different blood temperatures \pm CO₂ \pm human odor. In (B)–(G), significance was assessed with one-way ANOVA. Data labeled with different letters are significantly different using Tukey's HSD test (p < 0.001; ns, not significant). Replicate data points and mean ± SEM are shown (n = 6-10 trials/genotype and condition; n = 25 mosquitoes per trial).

generate mosquitoes unable to detect or behaviorally respond to volatile CO₂. Interestingly, mutating only Gr3 while sparing Gr1 and Gr2 was sufficient to disrupt CO₂ responses. This suggests that this subunit plays an essential role not compensated for by the two remaining CO2 receptor subunits. The stoichiometry of the CO₂ receptors in Aedes, Anopheles, and Drosophila currently remains unknown. In Drosophila, it has been shown that the two CO₂ receptor subunits together reconstitute a functional receptor (Jones et al., 2007; Kwon et al., 2007). Ectopic expression of An. gambiae CO₂ receptor subunits in Drosophila OSNs suggests that Gr1 and Gr3, as well as the combination of Gr1, Gr2, and Gr3, form a functional CO₂ receptor (Lu et al., 2007). In Ae. aegypti, transient RNAi knockdown experiments that assessed mosquito orientation toward a CO₂ source suggested that Gr1 and Gr3 but not Gr2 play a role in CO_2 reception (Erdelyan et al., 2012). The respective functional contribution of Gr1 and Gr2 to CO_2 detection therefore remains unclear and awaits further genetic and biochemical characterization. It will be interesting to determine whether all three subunits are functionally required to detect CO_2 or whether Gr1 and Gr2 are functionally redundant. Further, it is not known whether one CO_2 receptor subunit functions as a non-ligand-selective coreceptor whereas others interact with the ligand, as is the case with insect odorant receptors and ionotropic receptors (Benton et al., 2009; Larsson et al., 2004).

Several volatile odorants have recently been shown to antagonize, activate, or hyperactivate the CO₂ neuron in a variety of mosquito species, in some instances leading to behavioral disorientation (Lu et al., 2007; Tauxe et al., 2013; Turner et al., 2011). The availability of Gr3 mutants will allow us to ask whether these compounds act through the CO2 receptor complex or through other uncharacterized receptors or signaling machinery associated with this neuron. Further, multiple odorants have been shown to antagonize responses from the CO₂ neuron and to reduce mosquito attraction when applied at high concentration (10⁻¹) to human skin (Tauxe et al., 2013). We suggest that these chemicals are unlikely to mask attraction to humans via a loss of activity in the CO₂ receptor complex alone because CO₂insensitive Gr3 mutants remain highly attracted to live humans in our semi-field assays. Whether these odorants also activate additional mosquito chemosensory receptors on the antenna, palp, or proboscis at high concentration to elicit mosquito repulsion remains to be determined.

Our behavioral data support the conclusion that multimodal integration of CO₂, heat, and odor drives host-seeking and blood-feeding behavior in *Ae. aegypti*. We show that CO₂ gates mosquito perception of both heat and the monomolecular attractant lactic acid. This binary synergism extended to other combinations of sensory stimuli, including human skin odor with heat or CO₂. This synergism of multiple sensory cues may explain why attraction to live humans was diminished but not abolished in *Gr3* mutants in semi-field and two-port olfactometer trials. Interestingly, we also observed that synergy between CO₂ and host odor was sufficient to induce wild-type female mosquitoes to blood-feed on unheated blood. Consistent with this notion, *Ae. aegypti* have been observed to blood-feed on cold-blooded animals such as lizards when warm-blooded vertebrates were unavailable (Woke, 1937).

The absence of a major deficit in *Gr3* mutant attraction to live humans in the semi-field cage tested here does not preclude the possibility that more pronounced effects may have been observed if the human host was presented in a larger semi-field cage. Indeed, we used tests with animal hosts presented in small or large cages to determine that the effects of CO₂ detection on host-seeking behavior appear to be dependent on spatial scale. Synergism between multiple remaining sensory cues including visual contrast, moisture, heat, and volatile odorants from the skin and breath may explain why attraction to live humans was diminished but not abolished in *Gr3* mutants in our assays.

When combined, heat and human odors have been found to influence cross-wind flight behavior and increase source-finding and landing responses in female *An. gambiae* (Spitzen et al.,

2013). Although it is not known over what distance mosquitoes perceive heat, it is generally assumed that thermal convection currents emanating from human skin rapidly dissipate to reach equilibrium with ambient temperature conditions within ≤ 1.5 m of the host (Lewis et al., 1969; Spitzen et al., 2013). Synergistic interactions between human odor or CO₂ with heat may potentially act over short distances to guide landing behavior as mosquitoes approach hosts.

Previous studies suggested that *Ae. aegypti* is attracted to both stationary and moving visual contrast, and that the activating effects of human scent combined with visual feedback allow mosquitoes to navigate upwind toward hosts (Kennedy, 1940). In addition, it is possible that diurnal patterns of flight periodicity in *Ae. aegypti* (Trpis et al., 1973) may randomly bring females into the effective range of host cues where multimodal integration may then occur. The role of mosquito vision in host-seeking and its interactions with odors warrants further investigation.

Multisensory integration allows organisms to form salient percepts from combined sensory stimuli to drive context-dependent behaviors. An example of such integration in humans is the fusion of smell and taste of food or drink into the single percept of flavor (reviewed in Small, 2012). Neither chemosensory input in isolation fully encodes the experience of the food or beverage, which may explain why humans with olfactory dysfunction report difficulties with enjoyment of food and wine (Keller and Malaspina, 2013). Another benefit of multisensory integration is improvement in the reliability of signal detection. In noisy auditory environments, verbal comprehension can be improved dramatically by the practice of lip reading, thus combining auditory and visual perception to enhance the accuracy of signal detection (reviewed in Campbell, 2008). For the female mosquito, such multisensory integration may be crucial for efficient host seeking in a cluttered sensory environment. In the case of female mosquitoes living near human settlements, there are likely to be many sensory distractors in the environment that emit heat, human scent, or CO₂ in isolation. By requiring the simultaneous presentation of at least two such cues before host-seeking behavior is triggered, the mosquito increases her chances of high-fidelity host localization. Furthermore, body odor is itself a complex blend of multiple odorants that acts to shape both mosquito host preference (DeGennaro et al., 2013) and the tendency of Ae. aegypti and An. gambiae to bite certain regions of the body including the lower legs, ankles, and feet (reviewed in Smallegange et al., 2011).

What are the neural mechanisms employed by the female mosquito to integrate sensory cues to trigger attraction to humans? One can envision peripheral mechanisms by which CO_2 or heat directly sensitize peripheral odor-detecting OSNs, but no such mechanism has yet been described in insects. Conversely, odors have been shown to affect responses of the CO_2 OSN (Lu et al., 2007; Tauxe et al., 2013; Turner et al., 2011), but this modulation required very high odor concentrations unlikely to be attained in human skin odor. A more plausible mechanism would be central integration of these cues, as is known to occur for human taste and smell integration in the anterior ventral insula (reviewed in Small, 2012). One can imagine odor and CO_2 synergy occurring early in sensory

processing pathways because both send afferents to the antennal lobe (Anton et al., 2003). Much less is known about what brain areas process heat cues, so it seems plausible that higher brain centers are responsible for the synergistic interactions reported here. It is intriguing that male mosquitoes do not show synergy between CO2 and heat sensation. We presume that this is not due to defects in primary detection of the cues but more likely due to differences in wiring the male brain or its susceptibility to modulation by multiple sensory inputs. The availability of genetically encoded calcium sensors and novel circuit-tracing methods (Ruta et al., 2010) opens up the possibility of carrying out functional imaging in the mosquito brain to reveal the anatomical sites and mechanisms of multisensory integration.

The ability of Ae. aegypti to improve host source localization via multimodal integration of sensory cues could act to increase the probability that the female mosquito obtains a blood meal, a behavior inextricably linked to mosquito reproduction. Because integration between any pairing of CO₂, heat, and host odor is sufficient to drive mosquito blood feeding, chemicals targeting a single sensory modality (Tauxe et al., 2013; Turner et al., 2011; Jones et al., 2011) may be less effective in perturbing mosquito behavior than their combined use to disrupt multiple sensory pathways. Such synergistic approaches to manipulate mosquito sensory perception may ultimately yield a powerful strategy to deter mosquitoes from finding humans.

EXPERIMENTAL PROCEDURES

Mosquito Rearing and Maintenance

Ae. aegypti wild-type (Orlando) and Gr3 mutant strains were maintained and reared at 25°C-28°C, 70%-80% relative humidity, with a photoperiod of 14 hr light:10 hr dark (lights on 8 a.m.) as previously described (DeGennaro et al., 2013). Adult mosquitoes were provided constant access to 10% sucrose solution for feeding. Females were provided with a blood source for egg production. All laboratory blood-feeding procedures with mice and humans and use of humans in behavioral assays were approved and monitored by The Rockefeller University Institutional Animal Care and Use Committee and Institutional Review Board, protocols 11487 and LVO-0652, respectively. Semi-field cage experiments performed with human subjects at James Cook University were approved and monitored by the JCU Institutional Biosafety Committee and Human Research Ethics Committee (Ethics approval H4450). All human subjects gave their informed consent to participate in laboratory and semi-field experiments. Female mosquitoes used for electrophysiological and behavioral assays were 5-10 days old and mated but not blood-fed. Before behavioral assays, mosquitoes were sexed and sorted under hypothermia (4°C) or aspiration and fasted for 14-24 hr in the presence of a water source.

Phylogenetic Analysis

Amino acid alignment and phylogenetic analysis of insect CO₂ receptor genes were performed with MegAlign version 8.0.2 (DNASTAR) using the ClustalW and neighbor-joining methods. The phylogenetic tree was visualized as a radial phylogram using Dendroscope version 3.2.8 (Huson and Scornavacca, 2012).

ZFN-Targeted Mutagenesis and Homologous Recombination

ZFNs targeting Ae. aegypti Gr3 were produced by the CompoZr Custom ZFN Service (Sigma-Aldrich Life Science). Details of the design of the ZFNs and isolation of deletion and homologous recombination mutants are in the Extended Experimental Procedures.

Single Sensillum Electrophysiology

Extracellular recordings of CO2 (Matheson Tri-Gas) and (R)-1-octen-3-ol (C.A.S. 3687-48-7, Bedoukian) responses from female maxillary palp capitate peg sensilla of the indicated genotype were carried out as previously described (DeGennaro et al., 2013; Pellegrino et al., 2010). All capitate peg sensilla sampled from wild-type and Gr3 heterozygous mosquitoes responded to both (R)-1-octen-3-ol and CO2. Corrected spike increases were calculated by subtracting the spontaneous activity for 1 s before stimulus application from the activity evoked during 1 s of stimulus. Spontaneous activity was determined by counting spikes for 1 s prior to the application of paraffin oil (Sigma-Aldrich) to the same sensillum.

Three-Dimensional Multi-Insect Tracking System

A multi-insect 3D tracking system was custom designed and built in collaboration with SciTrackS GmbH. Details of the fabrication and operation of the system are in the Extended Experimental Procedures.

Heat-Seeking Assay

For each 15 min trial, 20–25 mosquitoes were introduced into a custom-made Plexiglass box (30 \times 30 \times 30 cm) and allowed to acclimate for 5 min. Two Peltier elements (6 x 9 cm surface area; Tellurex) located on opposite walls of the enclosure, and covered with white paper and white plastic mesh, were used to present heat stimuli. During acclimation, both Peltiers were kept at ambient temperature (26°C) via controllers (Oven Industries) commanded by a custom MATLAB script. Throughout the trial, carbon-filtered air was pumped into the box via a diffusion pad (59-144, Flystuff.com) installed on the ceiling of the enclosure. After acclimation, 10% CO2 was added to the air stream for 20 s via a solenoid valve (Parker-Hannifin) to increase CO2 concentration 1,000 ppm above background levels. At this time, one Peltier was warmed to 37°C for 5 min and subsequently cooled to ambient for the remaining 5 min of the trial. The side of the heated Peltier was randomized across trials. Mosquito landings at the Peltier were monitored by fixed cameras (FFMV-03M2M-CS, Point Grey Research) acquiring images at 1 Hz. We noted that mosquitoes were attracted to the Peltier surface as well as to adjacent surfaces, presumably due to heat dissipation and convective currents. Therefore, images were analyzed using custom MATLAB scripts to count heat-seeking mosquitoes within a fixed target region comprising a 12 cm × 16 cm area around the center of the Peltier. Peltier surface temperatures were measured using a thermal imaging camera (E60, FLIR).

Two-Port Olfactometer Assays

Two-port olfactometer assays were performed as previously described (DeGennaro et al., 2013). Experimental details are in the Extended Experimental Procedures.

Semi-Field Cage and Human-Landing Catch Assays

Semi-field cage experiments were conducted at the James Cook University Mosquito Research Facility Semi-Field System (Ritchie et al., 2011) in Cairns, Queensland, Australia from January 11, 2013-February 5, 2013. Mated, nulliparous 5- to 10-day females were used for these assays. Prior to use, mosquitoes were sexed via aspiration and fasted for 16 hr in holding containers with access to water only. To populate the semi-field cage for human-landing catch trials, at 10 a.m. each day, an equal mix of 100 each wild-type Orlando and $\mbox{\it Gr3}^{\mbox{\it ECFP/ECFP}}$ mutant female mosquitoes was released from a central release point in the cage (Figures 5A and 5B). Mosquitoes were allowed to disperse across the cage and acclimate to semi-field conditions for 5 hr. At 3 p.m. each day, a human volunteer entered the cage and sat down for a period of 15 min at one of two predesignated human-landing catch positions: either within the house structure or 11 m away in the yard of the cage (Figure 5B). During each trial, volunteers wore full-length white lab coats to standardize visual cues and used a hand-held mechanical aspirator (model 2809C, BioQuip Products) to collect mosquitoes that landed on their exposed legs and ankles. Mosquitoes were aspirated during five consecutive 3 min collection windows across each 15 min trial period using separate aspirator cartridges (2809V, BioQuip Products). The number of wild-type and Gr3^{ECFP/ECFP} mutant mosquitoes landing on each volunteer was then scored post hoc by assaying tissue homogenates from each collected mosquito for ECFP fluorescence using a POLARstar Omega microplate reader (BMG Labtech) equipped with a 440/485 nm excitation-emission filter set. To prepare samples for analysis, whole mosquitoes were homogenized at 4°C in 2 ml microcentrifuge tubes containing 100 μI PBS (pH 7.2) (GIBCO) with 2 mm borosilicate beads (Sigma-Aldrich) using a QIAGEN TissueLyser II (QIAGEN) set to maximum speed for 5 min. Samples were centrifuged at 12,100 \times g at 4°C for 5 min to pellet debris and obtain a soluble protein fraction that was assayed in triplicate for ECFP fluorescence on black 384-well small volume HiBase polystyrene microplates (784076, Greiner Bio-One). Three adult male volunteers (ages 27–46) participated in human-landing catch trials. Each volunteer was tested three times in each of the two house and yard positions (n = 9 trials total per position). A single experimental replicate was performed each day.

In separate experiments, the distribution of wild-type and $Gr3^{ECFP/ECFP}$ mosquitoes in the cage in the absence of humans was determined using eight Biogents Sentinel (BGS) Mosquito traps (Biogents GmbH) devoid of chemical lures and symmetrically placed, two in each of four trap zones, across the semi-field cage (Figures 5A and 5B). During BGS trials (n = 5), each mixed population of 100 females per strain was released at 9 a.m., and mosquitoes were allowed to distribute across the cage for 1 hr. After this acclimation period, traps were uncovered and turned on until 5 p.m. the same day. The BGS trap socks containing catches were then collected and emptied, and mosquitoes genotyped as described above.

Between experimental replicates, the semi-field cage was further depleted of all remaining mosquitoes by a mass-trapping procedure using 1 hr human-landing catches, mechanical aspiration, and continuous BGS trap runs. Cage depletion between replicates was verified in the morning prior to initiating the next experimental replicate by the prerequisite condition that no mosquito landings were observed during two mock 15 min human-landing catch periods at both human-landing catch positions in the semi-field cage.

Host-Seeking Assays with Mice

Assays were performed in small (30 cm \times 30 cm \times 30 cm; DP1000, MegaView) or large (91 cm \times 61 cm \times 61 cm; BioQuip Products) screened mosquito cages. For each trial, an equal mix of 20 each wild-type and $Gr3^{ECFP/ECFP}$ females was released and acclimated for 5 min. After this period, a single female BALB/c mouse (Harlan Laboratories) anaesthetized with ketamine:xylazine was introduced into the center of the cage, and mosquitoes were given the opportunity to blood-feed for 10 min. The number of mosquitoes of each genotype that blood-fed on the mouse was scored post hoc by *ECFP* fluorescence with a Nikon SMZ1500 dissecting microscope fitted with a C-HGFI Intensilight Illuminator (Nikon Instruments Inc.). A separate mouse was used in each experimental replicate. Humans were not present during acclimation or feeding to eliminate human host cues.

Membrane-Feeding Assay

A membrane-feeding assay was assembled using six custom 20 mm flanked glass jacketed membrane feeders (JHU-0804-081MS, Chemglass Life Sciences) connected in series via silicone tubing to a digital 12 I water bath (VWR International) set to 37°C. Technical details of the construction and operation of the assay are in the Extended Experimental Procedures.

Statistical Analysis

Statistical analysis was performed using GraphPad Prism Software version 5.0b (GraphPad Software, Inc.).

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2013. 12.044.

AUTHOR CONTRIBUTIONS

C.J.M. designed and carried out all the experiments in the paper except those in Figures 2 and 3, which were conducted by B.J.M. and R.A.C., respectively, with assistance from C.J.M. S.A.R. coordinated the semi-field cage trials in

Figure 5. C.J.M., R.A.C., B.J.M., and L.B.V. composed the figures. C.J.M. and L.B.V. together wrote the paper.

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Supplemental Information

EXTENDED EXPERIMENTAL PROCEDURES

ZFN-Mediated Targeted Mutagenesis of *AaegGr3*

Prior to ZFN design, the coding sequence and exon content of AaegGr3 (Vectorbase Accession number AAEL010058) was verified in the recipient Orlando strain with 3'RACE on maxillary palp cDNA. The following primers were used 5'-ATGAATCTCAACCAA GACCC-3' and 5'-GCGAGCTCCGCGCCGTTTTTTTTTTTTT-3' with AccuPrime HF Tag polymerase (Invitrogen). For genomic PCR across the Gr3 locus the following primers were used: (1) 5'-AGGGCATTGTTAGCGGGCATATGAA-3' and 5'-GGTTTGT GCATCCGAGTTCATCCAA-3' and (2) 5'-TTGGATTTCCGCCAATCTCTTCGAG-3' and 5'- ATCGTTGCCAAAAGCTGGAATGGAA-3' with Novagen KOD polymerase (EMD Millipore). The resulting PCR products were cloned using pCR4-TOPO (Invitrogen) and Sanger sequenced (Genewiz) to develop a consensus Gr3 sequence. Of 16 candidate ZFN pairs predicted in silico to bind and cleave within conserved regions of nucleotide sequence across exons 1, 2, and 3 of Gr3, one ZFN pair (Gr3 PZFN1/PZFN2) targeting exon 3 was chosen for use. This pair exhibited the highest levels of in vitro activity (106.7%) relative to the CCR5 control ZFN pair in a yeast-based MEL1 validation screen (Doyon et al., 2008) of all candidates. The nucleotide sequence of the ZFN-binding sites and nonspecific 5 nucleotide cut site for wild-type heterodimeric Fok1 endonuclease for Gr3 PZFN1/PZFN2 are denoted in upper case and lower case letters, respectively: 5'-ATCGTTAACATCCTGCCGgttctGATCATTCCGTTGATGTG-3'.

Nonhomologous End-Joining Deletion Mutants

To generate deletion alleles, Gr3 PZFN1/PZFN2 mRNAs (400 ng/ul) were microinjected into 550 pre-blastoderm stage Orlando Ae. aegypti embryos by Genetic Services Inc. Post-injection, 54 G0 larvae (9.8%) hatched, yielding 8 female and 8 male fertile G0 adults (from 31 adults total) that were outcrossed individually to wild-type Orlando mosquitoes to generate 16 G1 lines. PCR-based barcoding of amplicons crossing the target ZFN site coupled with Illumina sequencing was used to identify G1 pools containing mutant progeny as previously described (DeGennaro et al., 2013) with modifications. For this selection strategy, G1 progeny within each G0 progenitor line were self-crossed, blood-fed, and eggs collected en masse, before all adults from an individual G1 family were sacrificed and DNA extracted as a pool using the DNeasy protocol (QIAGEN). 16 G1 DNA pools were derived from 981 G1 adults total. Subsequently, a 388 bp gene-specific amplicon spanning the Gr3 ZFN cut site was amplified from each DNA pool by AccuPrime HF Taq polymerase (Invitrogen) using Gr3-specific primers (bold) with Illumina adapters and 6 nucleotide barcodes (nnnnnn): 5'-AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCTnnnnnnTTCGAAGAGGCCG TCATCGCTTA-3' and 5'-CAAGCAGAAGACGGCATACGAGCTCTTCCGATCTTCAAAGCCCTTTGGAAATCCTCC-3' (total amplicon size = 486 bp). Gr3 amplicons from each G1 pool were barcoded twice in separate PCR reactions using sets of barcodes: (PCR reaction 1) cgtgat, acatcg, gcctaa, tggtca (PCR reaction 2) cactgt, attggc, gatctg, tcaagt; such that mutant sequences could be assigned to a specific pool by the binary combination of two unique barcodes in a 4 × 4 matrix of 16 G1 families total. Barcoded PCR amplicons from each G1 pool were combined in equimolar ratios and sequenced using 100 bp single reads on an Illumina HiSeq 2000 sequencer (Rockefeller University Genomics Resource Center), and data filtered and analyzed as previously described (DeGennaro et al., 2013). In this case, 7 out of the 16 G0 lines (43.8%) passed deletion alleles to their G1 progeny

Mutant alleles identified during Illumina sequencing were isolated using size-based genotyping of single-pair crosses between progeny derived from candidate G1 pools containing the mutations and wild-type mosquitoes. Gr3 amplicons for size-based genotyping were amplified using fluorescently labeled primers 5'-6FAM-TTCGAAGAGGCCGTCATCGCTTA-3' and 5'-TCAAAG CCCTTTGGAAATCCTCC-3' with AccuPrime HF Taq polymerase (Invitrogen), and analyzed by capillary gel electrophoresis using an Applied Biosystems 3730 DNA analyzer (Genewiz). PCR amplicons containing the deletions were cloned and Sanger sequenced to confirm mutations, and the alleles named for the number of deleted bases. After isolation, the $Gr3^4$ allele was outcrossed to wildtype mosquitoes for five successive generations, using size-based genotyping to select for mutant alleles at each generation before homozygotes were generated. The 4 bp frameshift in $Gr3^4$ alters an amino acid residue at position 150 in the 445 amino acid wild-type peptide sequence from P to F, and results in a premature stop directly after this residue. To verify that the genomic deletion led to the expected mutant transcript, mRNA was extracted from head tissue of wild-type and Gr34 mutant mosquitoes using RNeasy and Oligotex mRNA mini kits (QIAGEN), reverse transcribed using Superscript III Reverse Transcriptase (Invitrogen). A 555 bp amplicon (551 bp in mutants) spanning Gr3 Exon 2 and 3 was amplified with AccuPrime HF Taq polymerase (Invitrogen) using the primers 5'-GATATAGTTTACGGCACCACGA-3' and 5'-ACTGAGAGTTTCGCAGGTCATA-3'. This amplicon was subcloned into pCR4-TOPO (Invitrogen) and then Sanger sequenced (Genewiz). The sequence confirmed the proper splicing of exon 2 to 3, the presence of the 4 bp deletion, and the creation of premature stop codons caused by this frameshift.

Homologous Recombination Mutants

To facilitate ZFN-mediated homologous recombination in Ae. aegypti, a donor plasmid pSL1180-HR-PUbECFP (Addgene plasmid #47917) was made containing a fluorescent cassette flanked on either side by multiple restriction sites for inserting homologous arms. The fluorescent cassette consisted of the Ae. aegypti polyubiquitin (PUb) promoter (Anderson et al., 2010) sourced from pMos-3xP3DsRed-PUb-EGFP-attP (a kind gift of Zach Adelman) driving a single nuclear copy of ECFP with a SV40 polyadenylation signal. To target Gr3 using ZFN-mediated homologous recombination, homologous arms 800 bp in length flanking the Gr3 PZFN1/ PZFN2 cut site were amplified with Novagen KOD polymerase (EMD Millipore) from two genomic DNA clones tiling the Gr3 locus using the primers: 5'-atatatgcggccgcGACGATGATGCGTTAAATATTCATCGGA-3' and 5'-atatatgatatcCGATGAAGAGATAAGCG ATGACGGCCTC-3' for the left arm; and 5'-atatatggatccTGTGGTACGAAACTCGTAAGGTCTCCAG-3' and 5'-atatatggatccAT TATTTGAAAACAGGATTATTCAATCT-3' for the right arm; with underlined bases indicating restriction enzyme recognition sites added for cloning. *Gr3* PZFN1/PZFN2-binding sites were partially represented in each arm: 5'-ATCGTTAACATCCTGCCGgttctGAT CATTCCGTTGATGTG-3', with underlined bases indicating the start of each arm. Using this design, 33 bp of the *Gr3* PZFN1/PZFN1 site in exon 3 were deleted and a 2452 bp *PUb-ECFPnls-SV40* cassette was inserted during targeting. Left and right arms were cloned directionally into the Not1/EcoRV and BamH1 sites of *pSL1180-HR-PUbECFP*, respectively, using standard techniques to generate *pSL1180-HR-PUbECFP-GR3*.

To generate homologous recombination alleles, Gr3 PZFN1/PZFN2 mRNAs (200 ng/μl) and pSL1180-HR-PUbECFP-GR3 (750 ng/μl) were microinjected into 1000 pre-blastoderm stage embryos. From these injections, 295 G0 larvae (29.5%) hatched, yielding 86 female and 119 male fertile G0 adults (from 247 adults total) that were outcrossed individually to wild-type Orlando mosquitoes to generate 205 independent G1 lines. To select for homologous recombinants, progeny of each G1 line were screened as larvae for ECFP fluorescence using a Nikon SMZ1500 dissecting microscope excited by a Nikon C-HGFI Intensilight Illuminator. We detected two G1 lines out of 205 screened in which a proportion of larvae in each family (11/83 and 1/187 G1 larvae, respectively in lines 1 and 2) constitutively expressed ECFP. Positive G1 adults derived from these larvae (6 G1 females and 3 G1 males in line 1; 1 G1 female in line 2) were individually outcrossed to wild-type mosquitoes, and eggs collected before adults were sacrificed for DNA extraction. To confirm directed insertion of the PUb-ECFPnls-SV40 cassette into Gr3 Exon 3 in these mosquitoes, two diagnostic PCR products were amplified with Novagen KOD polymerase (EMD Millipore) using ECFP (5'-ACAAGCAGAAGAACGGCAT CAAG-3') and SV40 (5'-CCTCCCCTGAACCTGAAACATA-3') forward primers anchored within the inserted cassette and a reverse primer in Gr3 Exon 4 (5'-TCGTCTGGTATGCTGATTTGGAAC-3') outside the boundary of the right arm; and PCR products cloned and Sanger sequenced. All 9 individuals tested from line 1 contained directed insertions into Gr3 Exon 3, whereas no evidence for directed insertion was obtained from the single female in Line 2, which was discarded. A homologous recombination rate of 0.49% was calculated and defined as the percentage of fertile G0 individuals giving rise to directed homologous recombination events in F1 progeny, or 1/205 G0 lines.

A single mutant allele (*Gr3*^{ECFP}) from a G1 female in line 1 was chosen for further characterization, and outcrossed for eight generations to wild-type mosquitoes before homozygotes were generated. *Gr3*^{ECFP} mutants were genotyped with Novagen KOD polymerase (EMD Millipore) using the primers 5′-CTTCGAGGAGAGTGTTCGGTTTG-3′ and 5′-ACCAAATAGGTCACCATCG TTGC-3′ anchored in *Gr3* exons 1 and 4 (labeled F and R, respectively, in Figures 1B and 1D). This PCR reaction produced a 3,713 bp amplicon in *Gr3*^{ECFP} homozygous mutants, and a 1,294 bp amplicon in wild-type mosquitoes (Figure 1F).

SciTrackS 3D Multi-Insect Tracking System

The system consists of a flight chamber (1.25 m L \times 0.75 m W \times 1.00 m H) with clear acrylic sides and front. Carbon-filtered air was continuously provided to this chamber by a pump (Quiet Pressure Pump, 79610-81, Cole-Parmer). CO_2 was added to this airstream when required through the use of a computer-controlled solenoid valve and a valve controller (Model 360D1X75R, NResearch Inc.) operated by a MATLAB (Mathworks) script through an analog/digital interface (model USB-1208FS, Measurement Computing Corporation). Tracking was accomplished by imaging the chamber with two offset cameras (model piA640-210 gm; Basler AG) with 1/2 4-12 mm F/1.2 IR aspherical objectives (Tamron). Lighting was provided by two banks of IR LEDs (model ISO-23-IRN-24-AL2, Metaphase Technologies). 3D position information of each visible mosquito was calculated in real-time at 200 Hz using custom-built software based on previously published algorithms (Fry et al., 2000). Tracks were further processed, filtered, and analyzed using MATLAB.

 CO_2 activation assay: For each trial, 20 female mosquitoes were added to the flight chamber with a mechanical aspirator. After acclimation and tracking of baseline activity for 30 min, a 40 s pulse of 100% CO_2 was added to the airstream at a flow rate of 1,360 ml/min. This raised the concentration of CO_2 by 1,000 ppm above background levels, as measured in the middle of the flight chamber with a Carbocap Hand-Held CO_2 Meter (model GM70, Vaisala Inc.). Tracking continued for another 20 min to measure activity in response to this pulse of CO_2 . Female mosquitoes had access to cotton balls soaked in 10% sucrose placed in a dish in the flight chamber. Data are presented as the population distance tracked in 10 s bins (Figure 2B), or as cumulative distance tracked per mosquito in the 6 min immediately following CO_2 addition (Figure 2C). Experiments and analysis were performed blind to genotype.

Daily locomotor activity assay: For each trial, 20 female mosquitoes were placed into the apparatus and tracked for 23 hr. White LED lights provided overhead illumination during the "daylight" phase of the mosquito rearing cycle (14 hr lights on: 10 hr lights off). A cotton wick soaked in 10% sucrose solution was provided in the flight chamber. Data are presented as the distance tracked per mosquito in 10 min time bins (Figure 2D), cumulative tracked distance per mosquito over the 23 hr experiment, and the mean velocity of all individual tracks (Figure 2E). Experiments and analysis were performed blind to genotype.

Two-Port Olfactometer Assays

During each trial, 50 female mosquitoes were introduced into the central chamber of the olfactometer and allowed to acclimate to assay conditions for 10 min. Treatment odors \pm CO₂ were then introduced into the stimulus port at the beginning of the assay, after which time mosquitoes were allowed to fly upwind into the stimulus or blank port for 8 min. The number of mosquitoes attracted to stimulus and blank trapping chambers at the cessation of this time period was then scored.

In trials where supplemental CO₂ was added, a 10% CO₂: 90% custom air mixture (GTS-Welco) was introduced into the system via Nalgene tubing placed inside the stimulus port for a final concentration of 2,000–3,000 ppm CO₂, as measured using a CARBOCAP Hand-Held Carbon Dioxide Meter (GM70, Vaisala Inc.). To test mosquito responses to lactic acid, we placed 4 ml L(+)-lactic acid solution (C.A.S. 79-33-4, catalog number 27714; 88%–92%; Sigma-Aldrich) into the open base of a 60 mm diameter × 15 mm height Petri dish (0875713A, Thermo Fisher Scientific) positioned at the center of the stimulus port. To test mosquito responses to human odor or live humans we placed a nylon sleeves infused with human-odor or a live human arm respectively into the stimulus port as described previously (DeGennaro et al., 2013). Supplemental CO₂ was not added in assays with live human arms due to pilot observations of a ceiling-like effect with high levels of attraction that was independent of whether CO2 was added or not. For all treatment conditions, stimulus and blank ports were alternated between experimental replicates to control for side bias. Background responses were evaluated using blank trials with wild-type mosquitoes where no stimulus was presented in either port.

Membrane-Feeding Assay

Two 137016 extra-large fountain pumps (Sunterra) were used to circulate water through the outer jacket of the feeders, with three feeders connected in each series to each pump. Membrane feeders from these two linked series were placed inside the central assay chamber which consisted of a translucent polypropylene storage box 36 cm L × 31 cm W × 32 cm H (The Container Store) with a removable lid. Four 1.5 cm holes made on opposite walls of the chamber allowed the exit and entry of silicone tubing connecting the feeders to pumps and water bath. A CO₂ diffusion pad (8.9 cm × 12.7 cm; Tritech Research) was affixed to the inner center of the lid to allow delivery of purified air and CO_2 to condition the chamber atmosphere when required. A plastic platform (31 cm \times 26 cm) with 4 cm × 3 cm legs was inserted into the inner chamber to create an elevated false floor that containing six evenly spaced circular holes (8 cm diameter) designed to hold 6 × 16 oz paper cups (KH16A-J8000, Solo Cup Company) covered in white 0.8 mm Polyester Mosquito Netting (American Home & Habit Inc.) to house mosquitoes. Feeding positions were randomized according to genotype during assays.

Membrane feeders were prepared for use by stretching a double-layer of Parafilm M laboratory film (Bemis Company, Inc.) over the feeder reservoir, and filling them with 400 µl of heparinized whole sheep blood (Hemostat Laboratories). In experiments where human odor was required, 3.5 cm × 3.5 cm squares of sheer knee high nylon stockings (Duane Reade) previously worn continuously for 33 hr by human volunteers were affixed to membrane using a rubber band. Clean stocking was affixed to the membrane in all other experiments. During each trial, 6 cups of mosquitoes (n = 25 per cup) were introduced into their feeding positions in the central chamber, the apparatus closed, and mosquitoes allowed to acclimate to assay conditions. Charcoal-purified air was provided continuously throughout each assay via a quiet pressure pump (Part No. 2415B-01, Welch-Gardner Denver). After 5 min acclimation, fountain pumps were switched on to circulate warm water to the membrane feeders for a period of 15 min. In trials involving supplemental CO₂, a custom 10% CO₂: 90% air mixture (GTS-Welco) was added continuously to the assay atmosphere during heating at a rate of 1,592 ml/min using a Single Tube Rotameter Flow meter (P16A1-BA0A-023-92-ST, Aalborg) to increase CO₂ concentration to 15,000 ppm above background levels. After 15 min, cups were removed from the apparatus and mosquitoes cold anaesthetized. Blood feeding was visually scored and any mosquitoes that did not appear to have blood-fed were crushed between absorbent papers. The animal was scored as blood-fed if there was any evidence of blood on the paper.

SUPPLEMENTAL REFERENCES

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