TRPA1 tunes mosquito thermotaxis to host temperatures

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SUMMARY

While most ectotherms thermotax only to regulate their temperature, female mosquitoes are attracted to human body heat during pursuit of a blood meal. Here we elucidate the basic rules of *Aedes aegypti* thermotaxis and test the function of candidate thermoreceptors in this important behavior. We show that host-seeking mosquitoes are maximally attracted to thermal stimuli approximating host body temperatures, seeking relative warmth while avoiding both relative cooling and stimuli exceeding host body temperature. We found that the cation channel TRPA1, in addition to playing a conserved role in thermoregulation and chemosensation, is required for this specialized host-selective thermotaxis in mosquitoes. During host-seeking, *AaegTRPA1*⁻/⁻ mutants failed to avoid stimuli exceeding host temperature, and were unable to discriminate between host-temperature and high-temperature stimuli. TRPA1-dependent tuning of thermotaxis is likely critical for mosquitoes host-seeking in a complex thermal environment in which humans are warmer than ambient air, but cooler than surrounding sun-warmed surfaces.

INTRODUCTION

Thermotaxis is a sensory-motor behavior that guides animals toward a preferred temperature. This type of sensory navigation allows animals to avoid environments of noxious cold and heat, with the goal of remaining in physiologically suitable ambient temperatures. For ectotherms such as insects, thermotaxis behavior is the primary method of thermoregulation. Many invertebrates are vulnerable to temperature extremes, facing the risk of desiccation at elevated temperatures, and rapid hypothermia at low temperatures. Therefore, mechanisms to detect environmental temperatures and trigger appropriate approach or avoidance behaviors are extremely important for their survival. For instance, adult *C. elegans* worms migrate preferentially towards a specific thermal environment determined by the temperature of their cultivation (Hedgecock and Russell, 1975; Mori and Ohshima, 1995). Adult *Drosophila melanogaster* flies prefer a narrow range of air temperatures around 24-25°C (Sayeed and Benzer, 1996; Hamada et al., 2008) and rapidly avoid air temperatures of ~31°C (Ni et al., 2013).

Interestingly, some hematophagous (blood-feeding) arthropods have evolved a specialized mode of thermotaxis to locate warm-blooded hosts. Such thermophilic behavior is seen in kissing bugs [*Triatoma infestans* (Flores and Lazzari, 1996) and *Rhodnius prolixus* (Schmitz et al., 2000)], the bedbug [*Cimex lectularius* (Rivnai, 1931)], the tick [*Ixodes ricinus* (Lees, 1948)], and many species of mosquito (Clements, 1999) including *Ae. aegypti*, a major tropical disease-vector (Bhatt et al., 2013). Female *Ae. aegypti* require a vertebrate blood meal for the production of eggs, and finding a suitable warm-blooded host is therefore an essential component of reproduction. Mosquitoes use a variety of physical and chemical senses to locate hosts in their environment (Cardé, 2015). When host-seeking, these animals become strongly attracted to warm inanimate objects, eagerly probing at them as if they were hosts (Howlett, 1910).

In nature, mosquitoes thermotax in a complex thermal landscape in which ambient air temperature, host body temperature, and surrounding surface temperatures can vary widely. For mosquitoes such as *Ae. aegypti*, host-seeking behavior can be activated by an increase in ambient carbon dioxide (CO₂) (Majeed et al., 2014). This activation elicits flight activity (Eiras and Jepson, 1991; McMeniman et al., 2014), and results in an array of behaviors including attraction to visual stimuli (van Breugel et al., 2015) and host olfactory cues (Dekker et al., 2005; McMeniman et al., 2014), and landing on warm objects (Burgess, 1959; Eiras and Jepson, 1994; Kröber et al., 2010; Maekawa et al., 2011; McMeniman et al., 2014; van Breugel et al., 2015). *Ae. aegypti* flying in a wind tunnel can detect a warmed stimulus from a distance, eliciting attraction and thermotaxis (van Breugel et al., 2015).

What are the mechanisms by which animals detect thermal stimuli, and how might these be adapted for the specialized needs of heat-seeking female mosquitoes? Thermotaxis is typically initiated by thermosensitive neurons that sample environmental temperature to inform navigational decision-making. Such neurons must be equipped with molecular thermosensors capable of detecting and transducing thermal stimuli. Diverse molecular thermoreceptors have been identified in the animal kingdom, many of which are members of the transient receptor potential (TRP) superfamily of ion channels (Barbagallo and Garry, 2015; Palkar et al., 2015). Different thermosensitive TRPs show distinct tuning spanning the thermal spectrum from noxious cold to noxious heat. Among these is TRPA1, which is a heat sensor in multiple insects, including the vinegar fly *D. melanogaster* and the malaria mosquito *Anopheles gambiae* (Hamada et al., 2008; Wang et al., 2009). Neurons in thermosensitive sensilla (Gingl et al., 2005) of *A. gambiae* female antennae express TRPA1 (Wang et al., 2009). In *D. melanogaster*, TRPA1 mutants fail to avoid high air temperature in a thermal gradient. Interestingly, animals such as snakes and vampire bats (Gracheva et al., 2010; Gracheva et al., 2011) have evolved to use thermosensitive TRP channels to locate warm-blooded prey, raising the possibility that *AaegTRPA1* may be used by mosquitoes to find hosts. Recently, a structurally distinct insect thermosensor, Gr28b, was identified in *D. melanogaster* (Ni et al., 2013). Gr28b, a gustatory re-
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RESULTS

We previously described an assay to model heat-seeking behavior in the laboratory by monitoring mosquitoes landing on a warmed Peltier element in the context of a cage supplemented with CO₂ (Figure 1A,B) (McMeniman et al., 2014). This assay has the advantages that it is simple in design, produces robust behaviors, and enables the collection of data from large numbers of animals in a short experimental timeframe. Using this system, we can examine mosquito responses to diverse thermal stimuli and measure thermotaxis in different ambient temperature environments. We first needed to determine whether heat-seeking behavior habituates over multiple thermal stimulations. In our heat-seeking assay, *Ae. aegypti* mosquitoes reliably responded to 12 serial presentations of a 40°C stimulus over the course of more than 2.5 hours, with no evidence of habituation (Figure 1—figure supplement 1).

* Ae. aegypti can feed on a variety of hosts (Clements, 1999; Tandon and Ray, 2000) with core body temperatures ranging from ~37°C (humans) to ~40-43°C (chickens) (Richards, 1971) (Figure 1C). With this in mind, we hypothesized that *Ae. aegypti* mosquitoes should be flexible in their thermotactic targeting to accommodate this wide range of host temperatures. On the other hand, one may also expect mosquito thermotaxis to be sharply tuned to optimize host-seeking by avoiding objects that are too cold or too hot. It is unknown whether there are minimal or maximal temperature thresholds constraining mosquito heat-seeking, and whether responses to thermal stimuli depend on the background ambient temperature.

To investigate these questions, we measured attraction to thermal stimuli produced by heating the Peltier to temperatures ranging from ambient (set to 26°C in these experiments) to 60°C (Figure 1D-E). We found that mosquitoes were highly sensitive to thermal contrast, and are attracted to stimuli 2.5°C above ambient (Figure 1D-F). Furthermore, mosquito occupancy on the Peltier increased with stimulus temperatures up to 40°C. However, for higher-temperature stimuli we observed a dramatic reduction in Peltier occupancy. A 50°C stimulus resulted in approximately half as many animals on the Peltier compared to a 40°C stimulus (Figure 1D-F). Stimuli of 55°C or greater resulted in occupancy rates indistinguishable from an ambient thermal stimulus (26°C) (Figure 1E-F). Spatial analysis of mosquito occupancy on or near the Peltier revealed that while mosquitoes were still attracted to high-temperature stimuli, they populated the area peripheral to the Peltier, and strongly avoided the Peltier itself for stimuli ≥ 55°C (Figure 1F). This selective thermotaxis is consistent with the natural range of thermal stimuli that mosquitoes encounter during their search for a blood meal from a live host (Figure 1C) (Richards, 1971; Cogrove and Wood, 1995; Yao et al., 2008).

Female mosquitoes searching for a warm-blooded host may be responding to the absolute temperature of a stimulus or may instead be evaluating relative warmth, defined as the differential between a
stimulus and background ambient temperature. To investigate the thermotaxis strategies constituting mosquito heat-seeking behavior, we conducted experiments at three ambient temperatures: 21, 26, and 31°C (Figure 2A-F). We found that Peltier occupancy for stimuli 21-40°C depended on the differential between the Peltier and ambient temperature (Figure 2B,C), rather than the absolute temperature of the Peltier (Figure 2A). For example, at all ambient temperatures tested, a stimulus 5°C above ambient was sufficient to elicit significant heat-seeking, and elicited approximately half as much Peltier occupancy as a stimulus 10°C above ambient. On the other hand, heat-seeking to targets 50-55°C was inhibited at all ambient temperatures tested (Figure 2D,F), despite the fact that the temperature differential varied widely in these situations (Figure 2E).

These results show that *Ae. aegypti* thermotaxis is driven by seeking relative warmth, but restricted by an absolute upper threshold of ~50-55°C. Because female mosquitoes are attracted to relative warmth, we hypothesized that they may also avoid relative cooling. This complementary behavior would serve to improve host-seeking thermotaxis. We examined mosquito responses to cooling by analyzing the rate at which animals left the Peltier when it cooled at the conclusion of a stimulus period. We found that mosquitoes left the Peltier at similar rates regardless of the absolute temperature of the stimulus (Figure 2G,H, Figure 2—figure supplement 2; based on analysis of data in Figure 1D,E), demonstrating that mosquitoes avoid relative cold during heat-seeking.

Our characterization of *Ae. aegypti* heat-seeking revealed multiple behavioral components contributing to selective thermotaxis during host-seeking: 1) the seeking of relative warmth; 2) the avoidance of relative cooling; and 3) the avoidance of thermal stimuli exceeding host temperature. Each of these sensory-motor functions may rely on the same molecular thermosensors, or may instead use distinct thermosensors. We considered the possibility that thermoreceptors ordinarily dedicated to the behavioral thermoregulation typical of most ectotherms such as *D. melanogaster*, may have evolved a function in host-seeking by mosquitoes and other hematophagous arthropods. Using this reasoning, we generated *AaegTRPA1*+/− mutants using zinc-finger nuclease-mediated genome editing (Figure 3A).

In addition to its function as a thermoreceptor, TRPA1 is a highly conserved chemosensor of electrophile irritants such as N-methylmaleimide (Macpherson et al., 2007; Kang et al., 2010). Using a modified capillary feeding (CAFE) assay (Ja et al., 2007) (Figure 3B), we found that wild-type *Ae. aegypti* mosquitoes strongly avoided consumption of N-methylmaleimide (Figure 3C), as well as the bitter compound denatonium benzoate (Figure 3D). *AaegTRPA1*+/− mutants rejected denatonium benzoate (Figure 3D), but did not avoid consumption of N-methylmaleimide (Figure 3C). We interpret this result as a loss of N-methylmaleimide detection in *AaegTRPA1*+/− mutants, leading to no preference between sucrose and sucrose containing N-methylmaleimide. We note that this simple CAFE assay could be used to discover additional mosquito anti-feedants to repel *Ae. aegypti*, beyond the two chemicals identified here.

Because TRPA1 is important in insect thermoregulation (Hamada et al., 2010), we used a modified thermal gradient assay (Sayeed and Benzer, 1996; Hamada et al., 2008) to assess thermal preference in wild-type and *AaegTRPA1*+/− mutant mosquitoes (Figure 3—figure supplement 1A). *AaegTRPA1*+/− mutants were impaired in avoidance of
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**Figure 3.** AaegTRPA1−/− mutants fail to avoid a chemical irritant and high-temperature stimuli.

A, Representative bright field (left) and fluorescence (right) images of wild-type and AaegTRPA1−/− female pupae marked by ubiquitous expression of enhanced cyan fluorescent protein (ECFP). Scale bars: 0.5 mm. B, Schematic of capillary feeding (CAFE) assay. C-D, Sucrose preference over sucrose containing the indicated concentration of N-methylmaleimide (C, n = 10-12 trials per condition) or denatonium benzoate (D, n = 7 trials per condition) for mosquitoes of the indicated genotypes (NS, not significant; *p < 0.05, **p < 0.01; one-way ANOVA with Bonferroni correction compared to wild-type). E, Percent of mosquitoes of indicated genotypes on Peltier during seconds 90-180 of stimuli of indicated temperature (mean ± s.e.m., n = 6-9 trials per genotype; ***p < 0.001; repeated measures one-way ANOVA with Bonferroni correction). F, Heat maps showing mean mosquito occupancy for the indicated genotypes on the Peltier (red square) and surrounding area, during seconds 90-180 of each stimulus period. Bold borders indicate stimuli with responses significantly different from wild-type in (E) (p < 0.05; repeated-measures ANOVA with Bonferroni correction). G, Mean percent of mosquitoes of indicated genotypes on Peltier during thermal stimuli 40-60°C and during subsequent re-presentation of 40°C. Timespans with statistically significant increases in AaegTRPA1−/− mutant Peltier occupancy compared to wild-type are indicated by purple lines (calculated from 15 second bins; p < 0.05; one-way ANOVA with Bonferroni correction).

**Figure 4.** AaegTRPA1−/− mutants fail to discriminate between host-temperature and higher-temperature targets.

A, Schematic of heat-seeking choice assay. B-C, Preference for 40°C versus 40°C (B) or 50°C versus 40°C (C) Peltiers for indicated genotypes (n = 6 trials per genotype; mean ± s.e.m., with each replicate indicated by a dot; NS, not significant; ***p < 0.001; one sample t-test versus zero preference). In (C) AaegTRPA1−/− mutants are significantly different from wild-type and heterozygous mutants (p < 0.05, one-way ANOVA with Bonferroni correction). D, Heat maps showing mean mosquito occupancy for the indicated genotypes on Peltiers (red squares) of the indicated temperatures and surrounding area, during seconds 60-240 of each stimulus period. E, Model of mosquito thermotaxis. F, Thermal image of a person (arrow) standing on a sunlit patch of grass in Central Park in New York City.
high air temperature, leading to significant mortality (Figure 3—figure supplement 1B-F). Together, these data indicate that TRPA1 has a conserved chemosensory and thermosensory function in Ae. aegypti.

We next asked if AaegTRPA1 is required for mosquito heat-seeking behavior. AaegTRPA1 +/- mutants showed normal attraction to stimuli at or below 45°C, but strikingly lacked normal avoidance of higher-temperature stimuli (50°C and 55°C) (Figure 3E,F). A detailed analysis of Peltier occupancy over time revealed that AaegTRPA1 +/- mutants persisted on the Peltier during 50, 55 and 60°C stimulus presentations, whereas control animals rapidly left these high-temperature stimuli (Figure 3G). Therefore, AaegTRPA1 is required for normal avoidance of high-temperature stimuli that exceed host body temperature. Because AaegTRPA1 +/- mutants retained attraction to warm stimuli, we used targeted mutagenesis to test a requirement for AaegGr19, the Ae. aegypti orthologue of Gr28b, in heat-seeking (Figure 3—figure supplement 2A). Although this thermosensor is required for mediating rapid avoidance of warmth in D. melanogaster (Ni et al., 2013), AaegGr19 +/- mutant mosquitoes showed no thermotaxis defects (Figure 3—figure supplement 2B).

While AaegTRPA1 +/- mutants did not show normal avoidance of high-temperature stimuli, they may still prefer host-temperature stimuli if presented with a choice. Using a heat-seeking choice assay with two independently controlled Peltiers, we examined the importance of AaegTRPA1 in guiding mosquito thermotaxis in a more complex thermal landscape (Figure 4A). In this assay, mosquitoes are simultaneously presented with two thermal stimuli. When presented with two 40°C stimuli, both wild-type and mutant mosquitoes distributed equally between the Peltiers (Figure 4B), but in a choice between a 40°C and 50°C stimulus, wild-type mosquitoes strongly preferred the 40°C Peltier (Figure 4C) and avoided the 50°C Peltier (Figure 4D). Remarkably, in this choice scenario, AaegTRPA1 +/- mutants failed to avoid the 50°C Peltier, resulting in no preference for the 40°C stimulus (Figure 4C,D, Video 1). These results demonstrate that AaegTRPA1 is required for mosquitoes to discriminate body temperatures from higher temperature stimuli, thereby tuning mosquito heat-seeking towards the temperature range of warm-blooded hosts.

DISCUSSION

We have elucidated the basic thermotaxis strategies used by host-seeking mosquitoes, and revealed an important role for TRPA1 in regulating this behavior (Figure 4E). Using a quantitative thermotaxis assay, we modelled Ae. aegypti heat-seeking behavior in the laboratory. We found that mosquitoes can search for hosts in a wide range of ambient temperatures by seeking relative warmth and avoiding relative cold. Remarkably, these animals can detect a stimulus with thermal contrast as small as 2.5°C. In an outdoor environment, however, hosts are often warmer than the surrounding air but cooler than sun-warmed soil, rocks, trees, and human-made objects (Figure 4F). For this reason, diurnal mosquitoes such as Ae. aegypti are poorly served by merely thermotaxing to the hottest object available. A more optimal strategy is to search specifically for biologically relevant stimuli, and to avoid thermal stimuli exceeding host temperature, as we have observed in our laboratory models of heat-seeking. Acquiring a blood meal is an essential component of reproduction for a female Ae. aegypti mosquito. To maximize her chances of finding a blood meal, a female mosquito should reject “distracting” stimuli that exceed host temperatures. Our results demonstrate that AaegTRPA1 is critical for this selective thermotaxis.

Mosquito heat-seeking behavior represents an excellent model system for further study of the genetics (Kang et al., 2012; Zhong et al., 2012), neuroscience (Frank et al., 2015; Liu et al., 2015), and decision-making (Lu et al., 2010) underlying thermosensation and thermotaxis. Until now, mechanistic studies of thermosensation have been restricted to traditional laboratory model organisms, such as domestic mice and Drosophila melanogaster flies, whose thermotaxis consists mainly of moving away from suboptimal thermal environments. Mosquitoes too, must undergo such behavioral thermoregulation, as we have found in our thermal gradient assay. However, their repertoire of thermotactic behaviors is expanded by the evolution of a specialized and highly tuned mode of thermotaxis to locate warm-blooded hosts. It will be interesting to investigate the neural mechanisms that regulate the divergent behavioral choices of thermoregulation and heat-seeking. We propose that these systems may be in behavioral conflict during mosquito host-seeking.

Our work identifies TRPA1 as a gene regulating mosquito avoidance of high-temperature stimuli, which we have shown to be a major behavioral component of heat-seeking. However, because both AaegTRPA1 +/- and AaegGr19 +/- mutants retain normal attraction to warmth, this aspect of heat-seeking must rely on other thermoreceptors, still to be identified. Our study shows that this attraction must be mediated either by a single thermosensor that adapts to background temperature, or multiple thermosensors each tuned to a distinct absolute threshold. Understanding the behavioral and molecular basis of thermotaxis in mosquitoes and other disease vectors (Flores and Lazzari, 1996; Schmitz et al., 2000) is of great biomedical importance. Ae. aegypti mosquitoes are potent vectors of yellow fever, chikungunya, and dengue arboviruses, resulting annually in hundreds of millions of infections (Bhatt et al., 2013). Further study of mosquito heat-seeking behavior may aid in the design of next-generation traps, repellents, and control strategies.

MATERIALS AND METHODS

Mosquito rearing and maintenance

Ae. aegypti wild-type (Orlando), AaegGr19, and AaegTRPA1 mutant strains were maintained and reared at 25-28°C, 70-80% relative humidity with a photoperiod of 14 hr light:10 hr dark (lights on at 8 a.m.) as previously described (DeGennaro et al., 2013). Adult mosquitoes were provided constant access to 10% sucrose solution for feeding, and females were provided with a blood source for egg production, either live mice or human volunteers. Blood-feeding procedures were approved and monitored by The Rockefeller University Institutional Animal Care and Use Committee and Institutional Review Board, protocols 14756 and LV-0652 respectively. Human volunteers gave their informed written consent to participate in mosquito blood-feeding procedures. Before behavioral assays, mosquitoes were sexed and sorted under cold anesthesia (4°C) and fasted for 15-24 hours in the presence of a water source.

ZFN-mediated targeted mutagenesis

Molecular Biology:

PCR was carried out using Novagen KOD polymerase (EMD Millipore, Billerica, MA), products were cloned using pCR4-TOPO (Invitrogen), and Sanger sequenced by Genewiz (South Plainfield, NJ).

ZFN Design:

ZFNs targeting AaegTRPA1 or AaegGr19 (VectorBase Accession numbers AAEL009419 and AAEL011073, respectively) were designed and produced by the CompoZr Custom ZFN Service (Sigma-Aldrich, St.
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Louis, MO). The nucleotide sequences of the ZFN-binding sites (upper case) and nonspecific cut site for wild-type heterodimeric Fok1 endonuclease (lower case) are:

| Homologous arm    | Primer sequences           |
|-------------------|----------------------------|
| AaegTRPA1 locus    | 5'-GCATGCATGGTGTA-AGAGCATGGTGTAAACTTGACAGCTCAA-3' and reverse, 5'-CGACAAGTGTTTATTGTGGTCAATC-3' |}

- **Homologous Recombination Design:**

  Donor plasmids were generated as previously described (McMeniman et al., 2014). All donor plasmids included homologous arms cloned from wild-type genomic DNA, and consisting of sequence flanking or partially overlapping the ZFN recognition sites.

  - **AaegTRPA1:** targeted using pSL1180-HP-PUbECFP-TRPA1. The left homologous arm (1608 bp, primers: forward, 5'-GATATGACATGGTGTA-ACAAAGAAGGTGT-3' and reverse, 5'-CGACAAGTGTTTATTGTGGTCAATC-3') and right homologous arm (3037 bp, primers: forward, 5'-GATATGACATGGTGTA-ACAAAGAAGGTGT-3' and reverse, 5'-CGACAAGTGTTTATTGTGGTCAATC-3') were cloned into the EcoRI and NotI sites of pSL1180-HP-PUbECFP (Addgene #47917), respectively.

  - **AaegGr19:** targeted using pSL1180-HP-PUbdsRED-Gr19. The left homologous arm (1257 bp, primers: forward, 5'-GATATGACATGGTGTA-ACAAAGAAGGTGT-3' and reverse, 5'-CGACAAGTGTTTATTGTGGTCAATC-3') and right homologous arm (1608 bp, primers: forward, 5'-GATATGACATGGTGTA-ACAAAGAAGGTGT-3' and reverse, 5'-CGACAAGTGTTTATTGTGGTCAATC-3') were cloned into the Xmal and NotI sites of pSL1180-HP-PUbdsRED (Addgene #49327), respectively.

  - **Generation of Mutant Lines:** To generate AaegTRPA1 homologous recombination mutant alleles, 2000 pre-blastoderm stage wild-type embryos were microinjected (Genetic Services Inc., Cambridge, MA) with AaegTRPA1 ZFN mRNA (200 ng/µl) and pSL1180-HP-PUbECFP-TRPA1 (750 ng/µl). To generate AaegGr19 homologous recombination mutant alleles, 1000 pre-blastoderm stage wild-type embryos were microinjected (IBBR Insect Transformation Facility, Rockville, MD) with AaegGr19 ZFN mRNA (200 ng/µl) and pSL1180-HP-PUbdsRED-Gr19 (700 ng/µl). Injected G0 animals were crossed to wild-type in multiple batches to generate G1 lines with independent ZFN mutagenesis events. G1 homologous recombination mutant individuals were recovered via fluorescence as previously described (McMeniman et al., 2014), and outcrossed separately to wild-type for 5 generations before establishing four independent homologous lines: AaegTRPA1ECFP-1, AaegTRPA1ECFP-2, AaegGr19dsRED-1, and AaegGr19dsRED-2.

  - **Capillary feeder (Cafe):** This assay was adapted for the mosquito from the original assay developed for Drosophila (Ja et al., 2007) For each trial, 5 mosquitoes were fasted with access to water for 24 hours, and placed in a polypropylene vial (#89092-742, WVR, Rednor, PA) with access to two 5 µl calibrated glass capillaries (#53432-706, WVR) containing 10 % (w : v) sucrose with 5 % (v : v) green McCormick brand food dye. Capillaries spaced ~0.5 cm apart traversed the cotton plug (#49-101, Gen-
esee Scientific, San Diego, CA) of the vial and protruded into the vial < 1 mm to provide a flush surface for mosquitoes to access the liquid while resting on the plug surface. The control capillary contained only green sucrose solution, and the experimental capillary contained green sucrose solution supplemented with 0, 1, or 10 mM of denatonium benzoate (Sigma-Aldrich) or 0, 10, 50, or 100 mM of N-methylmaleimide (Sigma-Aldrich). The experimental capillary with 0 mM chemical was identical to the control capillary, and served as a zero-choice to test for side-bias in the assay. Because a small amount of liquid evaporated during preparation of the capillaries, all choice conditions and genotypes were prepared in a time-staggered format so that any measurement error due to evaporation was spread across the conditions. The levels of remaining liquid in both capillaries were measured after 18-20 hours and were compared to the known initial liquid level. Experiments started at ZT 8-10 and ended at ZT 4-6 the following day. Consumption values were compared to control capillaries in vials without mosquitoes to account for evaporation. We note that in cases where mosquitoes did not feed from a capillary and all liquid loss was due to evaporation, consumption values could be calculated to be negative due to very small variation in evaporation rates between experimental and control capillaries. Any negative consumption values were rounded to zero. Sucrose preference was calculated by dividing the amount consumed from the control capillary (not containing denatonium benzoate or N-methylmaleimide) by the total amount consumed from both capillaries. In experiments with 0 mM denatonium benzoate or 0 mM N-methylmaleimide, one capillary was arbitrarily chosen as “sucrose only.”

Thermal gradient:
This assay was adapted from one developed for Drosophila (Sayeed and Benzer, 1996; Hamada et al., 2008). A custom-built enclosure (6 mm tall) was affixed to an aluminum thermal gradient bar (50 x 30.5 cm, TGB-5030, ThermoElectric Cooling America Corp., Chicago, IL) driven by two Peltier Elements (AHP-1200CPV, ThermoElectric Cooling America Corp.). The enclosure was separated lengthwise into 4 lanes (each 50 x 6 cm) that were visually isolated from one another. Three lanes were for testing mosquito thermal preference, while the fourth lane was dedicated to measuring air temperature via an array of 8 digital temperature sensors (DS18B20, Maxim, San Jose, CA; connected to an Arduino Uno, https://www.arduino.cc/) mounted to the top of the enclosure and distributed evenly across the length of the lane and centered in each analysis sector. An overhead camera (C910, Logitech, Lausanne, Switzerland) monitored mosquito position through the transparent lid of the enclosure and distributed evenly across the length of the lane. Images were acquired once per minute and analyzed using custom MATLAB scripts to count mosquitoes across 8 analysis sectors of the lane. The assay was conducted in a room maintained at 80-90% relative humidity and 14°C to achieve low air temperatures within the gradient enclosure. At the beginning of each 3-hour trial, the air temperature throughout the enclosure was stabilized at ~26°C, and 25-30 mosquitoes were introduced into each lane of the assay. After 90 minutes, a thermal gradient was established (air temperatures: ~19°C to ~36°C) by heating the right Peltier element and cooling the left Peltier element. Mosquitoes were monitored for an additional 90 minutes while exposed to this thermal gradient. Mosquito distributions during minutes 60-90 were monitored in both the “no thermal gradient” and “thermal gradient” conditions. Dead mosquitoes were visually identified at the conclusion of each trial. All genotypes were tested in parallel, and their lane positions were randomized across trials.

Thermal images
All thermal images were acquired with an infrared camera (E60, FLIR Systems, Wilsonville, OR).

Statistical analysis
All statistical analyses were performed using Prism 5 software (Graph-Pad Software, Inc., La Jolla, CA).

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AUTHOR CONTRIBUTIONS
R.A.C. designed and carried out all experiments in the study. R.A.C. and L.B.V. together interpreted the results, designed the figures, and wrote the paper. The authors declare no competing financial interests.

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SUPPLEMENTAL MATERIALS

1 video and 4 figure supplements

Video 1. AaegTRPA1 is required for tuning avoidance of high-temperature stimuli during heat-seeking.
Representative videos of wild-type and AaegTRPA1−/− mutants presented with a choice between two Peltiers, one at 40°C and one at 50°C. Video is sped up 10-fold (images acquired at 1 Hz and reproduced at 10 frames/sec), and shows seconds 80-240 of the stimulus period.
Available for download on bioRxiv linked to this pre-print.

Figure 1—figure supplement 1. Mosquitoes show persistent thermotaxis to repeated stimuli.
A, Peltier temperature measured by thermocouple (top trace, mean in red, s.e.m. in gray) and percent of mosquitoes on Peltier (bottom trace, mean in black, s.e.m. in gray). n = 10 trials. We note that variance in both traces is low, making s.e.m. traces difficult to see. B, Percent mosquitoes on Peltier during seconds 90-180 of each stimulus period in (A). Each replicate is indicated by a dot, and the mean by a line. There is no significant difference (p > 0.05) in Peltier occupancy between the first and last stimulus (repeated measures one-way ANOVA with Bonferroni correction).

Figure 2—figure supplement 1. Dynamics of Peltier temperature during stimulus periods.
Mean Peltier temperature measured by thermocouple during presentation of thermal stimuli (31-40°C). Dashed line indicates the end of the stimulus period.
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Figure 3 —figure supplement 1. AaegTRPA1−/− mutants fail to avoid noxious heat in a thermal gradient.
A, Schematic of the thermal gradient assay (top, side view). Representative experimental image (bottom, top view) showing mosquitoes outlined in red detected across one lane of the thermal gradient assay, with instantaneous air temperature reported for each of 8 analysis sectors. B-C, Percent of mosquitoes of indicated genotypes detected in each sector and mean sector air temperature (rounded to nearest °C) in the absence (B) or presence (C) of a thermal gradient (n = 6 trials per genotype). Data are plotted as mean ± s.e.m. (***p < 0.001; two-way ANOVA with Bonferroni correction compared to wild-type). D, Heat maps showing mean air temperature (left) and percent of mosquitoes of the indicated genotypes detected (right) in each sector over 90 minutes from the onset of a thermal gradient. E, Same data as in (D), showing total percent of mosquitoes of the indicated genotypes detected in sectors 6-8 (hot side, dotted line, mean; s.e.m., shading) during onset and maintenance of a thermal gradient. F, Percent of total mosquitoes of the indicated genotypes found dead in sectors 6-8 at the conclusion of the experiment. Each replicate is indicated by a dot, and mean ± s.e.m. by lines. Genotypes with different letters are significantly different (p < 0.01, one-way ANOVA with Bonferroni correction).

Figure 3—figure supplement 2. AaegGr19−/− mutants show normal thermotaxis.
A, Representative bright field (left) and fluorescence (right) images of wild-type and AaegGr19−/− female pupae marked with ubiquitous expression of Discosoma sp. red fluorescent protein (DsRed). Scale bars: 0.5 mm. The wild-type bright-field image is duplicated from (Figure 3A). B, Percent of mosquitoes of indicated genotypes on Peltier during seconds 90-180 of stimuli of indicated temperature (mean ± s.e.m., n = 6-9 trials per genotype). Neither AaegGr19−/− nor Aaeg-Gr19+/− were significantly different from wild-type at any stimulus temperature (repeated measures one-way ANOVA with Bonferroni correction).