Human blood type influences the host-seeking behavior and fecundity of the Asian malaria vector *Anopheles stephensi*.

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The nutritional requirements of mosquitoes include both sugar (generally derived from the nectar of flowers) and blood (humans or animals). Mosquitoes express different degrees of preferences towards hosts depending on behavioral, ecological, and physiological factors. These preferences have implications for mosquito-borne disease risk. The present study is directed to reveal the effect of the human blood groups on the fecundity and fertility of the malaria vector *Anopheles stephensi*.

In laboratory tests, mosquitoes were fed on ABO blood groups via artificial membrane feeders, and the level of attraction against different blood groups was tested by the electroantennogram and wind tunnel bioassay under control conditions. Results indicate that the female mosquitoes had a strong preference towards the blood group B, while in the case of females fed on O blood group had the highest digestibility rate. Overall, the human blood type had a significant impact on the fecundity and fertility of female *An. stephensi*. The highest numbers of eggs are laid, in the case of blood group B, (mean (± SD)) 216.3 (8.81) followed by the AB, 104.06 (7.67), and O, 98.01 (7.04). In the case of blood group B, females attain the highest fertility of about 92.1 (9.98). This study provides novel insight into the ABO blood type host choice of the mosquitoes that are still partially unknown and suggests encouraging personal protection for relevant individuals within communities at risk, which is a useful tool for preventing malaria where the *An. stephensi* is present as a dominant vector.

Mosquitoes pose a significant threat to human health due to the close association with the human habitat of many species, their propensity to bite people for blood, and their role in outbreaks of mosquito-borne diseases1–3. *Anopheles stephensi* Liston (Diptera: Culicidae) is among the critical vectors of malaria parasites in Pakistan, Iraq, Afghanistan, and India4. Mosquitoes of the genus *Anopheles* are important vectors of the human malaria parasites *Plasmodium falciparum* and *P. vivax*5. As most vector mosquitoes are anautogenous, females require a blood meal for the oogenesis6, and understanding blood-feeding preferences is critical for assessing mosquito-borne disease risk. The volume and quality of blood meal can play an important role in egg production and subsequently influence potential population dynamics and vector competence7. *Aedes aegypti* Linnaeus, a globally important vector of dengue viruses, when fed on animal blood, laid fewer eggs than after feeding on the blood of a human8. When *Culex quinquefasciatus*, a globally important mosquito of pets and public health importance, fed on bovine blood, it hurt fertility and fecundity compared to when provided chicken blood9.

Similarly, the source of blood provided to *Anopheles* mosquitoes can impact fecundity with human, cow, and chicken blood, having differing effects over the egg production rate of *An. gambiae*10. Beyond the source of blood, whether it is from animals or humans, the constituents of blood may play an important role, with some studies
indicating the importance of isoleucine, and amino acid, in the blood, influencing the rate of egg production in some species of mosquitoes\textsuperscript{11–13}.

Understanding the attraction of many vector mosquitoes to human hosts could assist in determining the role of individual mosquito species in outbreaks of mosquito-borne diseases and provide critical information to inform mosquito control and surveillance programs\textsuperscript{14}. The preferences of some mosquitoes to feed on humans are well documented, especially \textit{Ae. aegypti}\textsuperscript{15}. While many \textit{Anopheles} mosquitoes display a similar anthropophilic feeding behavior, which is a critical factor in determining their role in malaria outbreaks, there is also often a demonstrated difference in host-feeding preferences of \textit{Anopheles} mosquitoes. For example, \textit{An. gambiae} has been shown to have strong anthropophilic feeding preferences while \textit{An. stephensi} has zoophilic preferences\textsuperscript{16}. Understanding how these mosquitoes, considered primarily zoophilic, may opportunistically, or at times preferentially, respond to humans in host-seeking behavior.

Female mosquitoes use different cues to locate hosts; physical, visual, and chemical cues\textsuperscript{17}. Organic molecules released from the host body are considered the most important with regard to mosquito attraction. Sweat excreted from the human body has various components that influence attraction, and even slight variations in released organic chemicals influence host attraction by mosquitoes\textsuperscript{18}. Further, variations in attraction to individual humans have been of great interest. Hematophagous female mosquitoes show different levels of attraction towards different human beings\textsuperscript{19}. It is assumed that the host-seeking response to humans is directly linked with body odor\textsuperscript{19,20}. For example, \textit{Aedes} and \textit{Anopheles} mosquitoes are attracted more towards pregnant women, possibly because of their increased secretion of estrogen in urine\textsuperscript{21,22}. The perception that some individuals are more likely to be bitten by a mosquito than others is frequently debated, and differences in human odor profiles and the response of mosquitoes have been a focus of research\textsuperscript{23,24}. While an individual's odor profile may be complex and vary due to various factors, blood type is often proposed as a determinant of propensity to be bitten by mosquitoes\textsuperscript{25}.

In this study, the effect of human ABO blood groups on female mosquito oogenesis and fertility was investigated under laboratory conditions using methods of electroantennography and olfactometry bioassays. The investigation aimed to determine that a specific blood group represents the following qualities (a) more attractive than all other blood groups (phagostimulant); (b) fully engorged when feeding; (c) initiated the vitellogenin; (d) must have a positive effect on oogenesis; and (e) positive effects on health and fitness of offspring. There is a paucity of published data on the effect of human blood types on the fertility and fecundity of mosquitoes. We tested the hypothesis that human blood type groups significantly influence specific components of the life cycle of \textit{An. stephensi} under laboratory conditions.

**Results**

**Fecundity and fertility.** Female \textit{An. stephensi} that were fed on blood group B laid an average of highest numbers of eggs 216.3 (8.81), whereas those fed on blood groups O and AB laid an average of (mean (± SD)) 98.01 (7.04) and 104.06 (7.67) eggs per female. Mosquitoes fed on blood group A laid the lowest mean numbers of eggs 65 (3); this is in line with the results from scanning electron microscope with the lowest development in the oogenesis was recorded (Fig. 1). But in the case of percent fertility, the females fed on blood group B acted as the best nutrition for the female mosquitoes because the females had the highest percent fertility 92.1 (9.98). On the other hand, the females fed on blood group A had the lowest percent fertility 59.7 (4.68). The data suggest that fecundity, fertility, and oogenesis are directly linked with the blood groups and the availability of the blood types.
Digestibility. A difference was observed in the rate of digestion of different blood groups after 12 h with visible differences in specimens of engorged mosquitoes according to the human blood group types provided. Approximately 6 h post-feeding, those fed on blood groups A partially changed into black while those fed on B and O completely changed into black. However, the abdomen of females fed on the blood groups, AB, still maintained a reddish color. After 12 h of the female fed on O blood groups in their stomachs showed two-third of the abdomen distended. After 24 h, the O blood group was half-digested in the abdomen while the AB batch has changed the color of the abdomen red to black, and the females engorged on blood group B had abdomen slightly more than three quarter still filled with blood. After 48 h, the O blood group color completely disappeared from the abdomens of all the females. In contrast, the blood groups A and B disappeared after about 60 h of post-fed. The AB blood group was the last one, which was disappeared after 72 h from all blood groups.

The percent rate of digestion of different Blood groups after feeding by female An. stephensi under control condition was shown in (Fig. 2). The blood group O has the highest rate of digestion, and it is significantly different from the other blood groups (P<0.0001).

The use of chemical tests, such as precipitin and bendizidine (determining the presence of iron porphyrins) tests have been used in mosquito blood feeding experiments and here, a positive reaction was observed for the both tests up to 100 h in the case of the AB blood group after the engorgement. Defaecated matter collected from the cages of each blood group also gave positive results.

Effect on oogenesis. Using a scanning electron microscope (SEM), two clusters of ovarioles were identified in the ovaries of the female fed over blood group B, 36 h post engorgement. While the development of ovaries commenced after 48 h in the case of blood groups A and AB. The result indicated that after the single engorgement of blood group O, the development of ovaries had not started. But here in this study, the females were selected after 36 h and were presented to SEM to test the ovary development stages. The ovarioles were situated at the middle of the spongy fat bodies like structure (Fig. 3). After 36 h, the ovaries development in female mosquitoes fed on blood group B looks mature. While the females fed over blood groups A, AB and O are still under immature conditions 36 h post-feeding (Fig. 3).

The total number of ovarioles in all females mosquitoes fed on different blood groups ranged from 100 to 600, strictly linked to the blood groups26. The highest numbers of ovarioles were observed in females fed on blood group B; all the three females have, on average of 550 ovarioles. The lowest numbers of ovarioles were counted in the female fed on blood group O about 150 ovarioles (Fig. 3).

Electroantennography. The electroantennography (EAG) experiments were conducted to test the response of different olfactory receptors on the antennae. First of all, the electroantennogram responses of An. stephensi were recorded against the known olfactory stimuli, including lactic acid, 1-octen-3-01, and isovaleric acid. After recording the results of control stimuli, the treatment stimuli were applied, and the response was recorded. The 5–7 days old unfed female An. stephensi mosquitoes gave a high blood group linked EAG response. Air current from blood group B indicated a significantly stronger response over the other blood groups. The amplitude response for the other blood group O is lower than the B. While the A blood group is the least attractive one blood group because the female showed the lowest level of response amplitude (Fig. 4A). The electroantennogram response is directly related to the time of air current in seconds; as the time of air current increases, the olfactory response of antennae was also increased. All the blood groups led to a significant response against the control of air currents (Fig. 4B).
Wind tunnel bioassays. In wind tunnel bioassays, *An. stephensi* exhibited a clear preference for human blood type B, as evident by the significantly higher number of mosquitoes attracted to that human blood type (Fig. 5). The second and third most attractive blood groups were the O and AB, while there was no significant ($P = 0.1454$) difference between the number of mosquitoes attracted to blood group A compared with control.

Similarly, *An. stephensi*, when exposed to olfactory cues derived from volunteers of differing blood group types, displayed a strong preference for blood group B compared to other blood groups (Fig. 5). The second most important stimulus was the O blood group, while the A and AB were the least attractive blood groups.

Mosquito fitness. The females fed on blood groups B and O showed the highest level of mosquito fitness, as evident by the lowest mortality rates for eggs, larvae, pupae, and adults (Fig. 6). When compared with the females fed on blood, groups A and AB have a significantly high larval mortality, egg infertility, and adult death rate.

Feeding rate and egg production. About 91% of the females fed on blood group B were fully engorged compared with the 70%, 54%, and 45% in the case of blood groups O, AB, and A (Fig. 7). On the other hand, the percent numbers of eggs were also higher in the case of females fed on blood group B (Fig. 8A). The rate of egg production and engorgement with blood group B increased more than 12% relative to blood group O and 55% in the case of A blood type (Fig. 8B), suggesting that the blood group B can be highly beneficial and attractive to female mosquitoes.
Discussion

This is the first time the host-seeking and reproductive response of *An. stephensi* to human blood group types has been undertaken. The findings of these laboratory studies suggest a strong preference in host-seeking of individuals with type B blood group with concomitant increased fecundity of mosquitoes feeding on that blood. Our results reflect those of other researchers who have found that the source of blood-fed on by mosquitoes of pets and public health importance influences fecundity and other aspects of their life cycle.

The amount and the quality of blood meal is a vital thing to female mosquitoes for their gonotrophic cycles and rate of egg production. However, the variation in the egg production rate and the number of females moving towards their first gonotrophic cycles is strictly linked with the amino acids and the protein contents in each blood meal. Djamila et al. evaluated the effects of two blood meals, the chicken and cattle blood, on the egg development rate of *An. maculipennis* and found fecundity were significantly lower with the chicken blood as compared with the cattle blood, while fertility is not associated with blood meals. The fertility, fecundity, hatching rate, developmental time of the larvae of *Cx. theileri* is strongly associated with the source of blood.

**Figure 4.** Electroantennogram response of *Anopheles stephensi*. (A) The electroantennogram response of female *An. stephensi* antennae to all ABO blood groups along with the scented air puffs. (B) The graph is showing the observed changes in numbers of spikes of antennae to the different blood group stimuli applied during the gray box. The value shown is the number of spikes per second, per female mosquito (*An. stephensi*). A value of 1 (dashed line) indicates no change in spike rate. The control stimuli applied to the antenna: Puffs of unscented air in black color demonstrate the lack of response of the antenna.

**Figure 5.** The response of female mosquitoes in wind tunnel bioassay. (A) The response of female mosquitoes against different blood groups. There is no significant difference observed between the unscented air and the blood group A ($P = 0.0172$), and there is also no significant difference was also present between the blood groups A and AB ($P < 0.0053$). The arm having the blood group B have significantly higher numbers of mosquitoes at the opposite end ($P < 0.0001$). (B) The response of female mosquitoes against filth collected from the hands and the armpit of the person having ABO blood groups.
The fecundity of females fed on chicken blood was significantly higher than the females fed on human and cow blood. We have analyzed the rate of digestion of different blood types by An. stephensi female mosquitoes by several methods; observational studies, precipitin test, and benzidine test. The visual and the tests indicated that the blood group O is the blood that females could easily digest in all three replications. This may be because of the chemical structure of the O blood group; it is simpler than the other blood groups in case of antigen absence.

Several methods are used to study the egg development mechanism or oogenesis in female mosquitoes. But the most important and widely used of them are a light microscope, scanning electron microscope, and fluorescent confocal microscopy. This study demonstrated that SEM was an effective method to study the ultrastructure of ovaries of female mosquitoes fed on different blood groups.

Experiments on females fed on ABO blood groups and control reveal the different levels of changes in the structure of reproductive organs that lead to the series of changes in cell structure. Blood type B elucidated the significantly higher changes in the cell structure and ovaries development. One of the protuberant changes in the cell structure of females fed on blood group B is a large number of un-oriented microvilli in the area of oocytes. The same type of microvilli was also observed in the case of females fed on blood groups O and AB but few in numbers and small in size. Results indicated that these microvilli are not uniform in length, and these are not regularly present in all females fed on different blood groups. This type of result is in line with the Brandt microvilli are not uniform in length.

The results are also in line with the scanning electron microscopy results obtained by Clements. The total numbers of ovarioles ranged from 120 to 650 in blood group B and 60 to 400 in the case of O and AB blood.

Figure 6. The percent fertility in all the rearing boxes and in 10 replications. All the blood groups are significantly different from each other, and the blood group “B” has the highest percentage fertility in the next generation.

Figure 7. The percent fed and unfed female mosquitoes. The female Anopheles stephensi mosquitoes were reared under different control conditions on different blood groups.
groups. Only 20–30 ovarioles were identified in the case of females fed on blood group A. The results are also in agreement with Roth and Porter46, who described the oogenesis in Ae. aegypti.

Most of the autogenous mosquitoes detect the volatile semiochemicals by the use of olfactory neurons present on antennae. Earlier studies indicated the EAG response of An. gambiae towards the volatile compounds of human sweat and the carboxylic acid from the cheese40,41. About eight volatile compounds were identified from the chicken feces that provoked the electroantennogram response from the Cx. quinquefasciatus42. These volatile also derived the behavior of female Cx. quinquefasciatus under control conditions. The female Ae. albopictus showed the inverse dose-dependent EAG response against certain fatty acids and alcohols derived from the human skin emanations. In the case of the present study, substantial electroantennogram responses were displayed by the antennae of selected mosquitoes against known stimulants l-lactic acid, 1-octen-3-ol, and isovaleric acid. After confirming the sensitivity and response of antennae and the EAG towards the human-specific known stimulants, treatment stimuli ABO blood groups were applied and demonstrated a positive response of An. stephensi to the blood group B. The results from the electroantennogram response clearly stated that the blood groups have significant effects on the behavior of female An. stephensi; the blood group A is the least attractive.

Artificial blood-feeders with collagen membranes are an effective method to evaluate attraction responses of blood and chemical compounds released by the human skin and the sweat. The chemical volatiles permeated from the collagen membrane attracted the female mosquitoes and indicated the landing behavior in choice assays. In most of the previous studies, the collagen membrane was used in artificial membrane feeders to study the hematophagous behavior of female mosquitoes44. The blood emitted chemical volatiles that attracted the hungry female mosquitoes. Female Cx. quinquefasciatus and Ae. aegypti showed different levels of attraction towards the avian and bovine44. Blood composition and concentration are the key drivers of this type of behavior in female mosquitoes, mainly in a wide choice.

In the present study, both tests conducted in the olfactometer indicated that blood groups have a substantial impact on the behavior, fertility, and fecundity of the female An. stephensi mosquitoes. In both, cases blood group B acted as a mosquito magnet so that the human-specific blood groups may act as an important part for the identification of autogenous female mosquitoes such as An. stephensi.

An. stephensi is one of the most important vectors of malaria on the Indian subcontinent46 and is also distributed across the South Asia region and the Middle Eastern countries such as Afghanistan, Bahrain, Bangladesh, China, Egypt, Iran, India, Iraq, Pakistan, Oman, Saudi Arabia and Thailand46. The percentage of the population in these countries under threat of malaria due to this vector are Bahrain (23%), Bangladesh (34%), China (29%), Egypt (13%), Iran (25%), India (34%), Iraq (28%), Pakistan (33%), Saudi Arabia (18%) and Thailand (37%). The countries with the 2nd highest percentage of B blood group are Burkina Faso (29%), China (29%), Ghana (19%), Indonesia (29%), Iraq (28%), Ivory Coast (23%), Laos (35%), Mongolia (30%), Myanmar (33%), North Korea (30%), Philippines (25%), Singapore (25%), Sri Lanka (27%), Thailand (37%) and Vietnam (31%)46. According to the present study An. stephensi prefer to feed on blood group B. So the countries with a higher percentage of blood group B need more protection against this malarial vector An. stephensi.

In conclusion, the present study supports the hypothesis that blood groups have effects on the fertility, fecundity, and behavior of female An. stephensi, just like An. albimanus44. An. stephensi is a well-known vector involved in the transmission of malarial parasites (P. falciparum and P. vivax), and given blood from different vertebrates affects the behavior, fertility, and fecundity of mosquitoes; our results suggest this may be due to differences in
the chemical composition and concentration of blood of vertebrate\textsuperscript{29,44}. Significant differences were observed in both the host-seeking behavior in response to differing ABO blood group types and concomitant differences in fecundity following blood-feeding by \textit{An. stephensi} in a laboratory setting. The results of this study represent an important breakthrough in the field of parasitology and malaria control. It provides clear insights about the behavior of female mosquitoes that not only may have implications for determining in whom the community most at risk of exposure to malaria parasites, and consequently possibly prioritized for anti-malarial medication. It opens potential opportunities for the development and adaptation of novel mosquito control and surveillance strategies that exploit the host-seeking behaviours demonstrated here.

Methods

Rearing. All mosquitoes used in these experiments were derived from a laboratory-reared colony of \textit{An. stephensi} is initially established (six generations) in University College Agriculture, University of Sargodha, Pakistan. Uninfected mosquitoes were maintained in the laboratory; in gauze-covered boxes (30 cm wide $\times$ 30 cm deep) under control conditions of temperature $27 \pm 2\, ^\circ\text{C}$ and relative humidity 75–80\%. A 12:12 h light:dark cycle was set. Feeding success was determined by calculating the percentage of engorged female adult mosquitoes. The engorged female adult mosquitoes were culled from the breeding box at 8 h intervals, rubbed over the filter papers (25 cm wide $\times$ 25 cm deep) to break the scotophase (dark) period in the control conditions of the laboratory. Mosquitoes that were killed at 8 h intervals, rubbed over the filter papers (25 cm wide $\times$ 25 cm deep) were used in experiments. Mosquitoes were provided two days following the second blood meal. The larvae were reared under laboratory conditions described above and provided a certified Laboratory Rodent Diet (LRD) Lab Diet 5001\textsuperscript{51}.

Fecundity and fertility. To determine differences in fecundity (number of eggs) and fertility (percentage of fertile eggs) in \textit{An. stephensi} mosquitoes, cages of mosquitoes were provided ABO blood groups and control (distilled water) via artificial membrane feeders (as described above for mosquito rearing). The blood was obtained from the blood bank of DHQ (one batch of each blood group was used for each replication). After each replication along with the new batch of each blood group, a new strain of mosquitoes was used just to reduce the learning behavior of mosquitoes. Feeding success was determined by calculating the percentage of fed mosquitoes, and also, the numbers of fully engorged female mosquitoes were recorded.

To determine fecundity, females from each blood group were removed, killed, and dissected under a microscope, and the numbers of eggs per female were counted 60 h post-blood-feeding. Additionally, to determine oviposition and larval development, 10 fully fed female mosquitoes were caged in one of three replicate glass cages with gauze (25 cm wide $\times$ 25 cm deep) and provided wet filter papers were placed for egg-laying. The total number of eggs was counted twice every 12 h from 48 h until 96 h post-blood-feeding under a microscope. The total numbers of eggs/40 females/box for each human blood group for 10 replicates were calculated.

For fertility estimation, an additional 40 gravid \textit{An. stephensi} from each treatment and replication, including the control group, the females were gently transferred to the cages with triangular Whatman filter paper No. 1 using the mouth aspirator. The egg laid in each experimental and control group was reared in plastic trays filled with distilled water. The numbers of hatched larvae were recorded for a fertility test. While the eggs that could be hatched in 2 days were recorded as fertile. The number of fertile and infertile eggs was recorded in all experimental and control cages. The collected eggs from each experimental box were placed into the plastic trays (24 cm wide $\times$ 12 cm deep) with water, and the development of mosquitoes was observed until adult mosquitoes had emerged from all pupae for each blood group. The water in these plastic trays was maintained at a constant level throughout immature mosquito rearing. The larvae were fed a certified Laboratory Rodent Diet (LRD) Lab Diet 5001\textsuperscript{51}.

The rearing was done according to the standard mass rearing of \textit{Anopheles} techniques\textsuperscript{52}. Pupae were counted and removed from the tray and placed in cages according to each human blood group type fed to allow emergence, and the percent of male and female mosquitoes was recorded. Adult mosquitoes were maintained on a 10% fructose solution supplemented with 0.05% para-minobenzoic acid (PABA) but were not provided with a blood meal. The mortality of adult mosquitoes was recorded daily until total mortality reached 100%.

Digestibility tests. The precipitin and benzidine tests were used to test the effect of human blood groups ABO (on the rate of digestion in mosquitoes). The experiment was conducted in controlled laboratory conditions where the temperature, humidity, and day and night periods were maintained as described above. Mosquitoes (10 mosquitoes were used for each blood group and the same experiment was repeated 10 times) that had not been fed previously on either a sugar solution or blood were used in experiments. Mosquitoes were provided one of four different human blood group types, as previously described. After feeding the female were kept in the same boxes without any further food and water, and boxes were placed in an incubator where the temperature and the relative humidity was at a constant level (28 $\pm$ 2 $^\circ\text{C}$ and 80 $\pm$ 5\%). The engorged female adult mosquitoes were killed at 8 h intervals, rubbed over the filter paper\textsuperscript{49}, and the filter papers were placed inside the refrigerator until the test could be conducted. Approximately 48 female mosquitoes were used in each boxed marked for each blood group. The rate of blood digestion in the engorged blood was classified according to the Sella scale, following Detinova et al.\textsuperscript{53}.

To perform the precipitin test, the physiological saline and the filter paper swabs were extracted in a small capillary tube. The specific antiserum was also extracted in the same capillary tube at the end; the change in
color, clumping, and cloudiness of the solution indicates the presence of human blood in the tissue smears. The collected material was heated in a steam oven for 10–12 min to apply the benzidine test at 108–110 °C. The test was used to check the traces of iron porphyrins in the abdomen of mosquitoes.

**Effect of blood groups on oogenesis.** To test the blood-specific effects on the development of the ovaries of *An. stephensi*, the ovaries of fully fed female mosquitoes were collected separately from the box of each blood group 36 h post engorgement. For scanning electron microscopy (SEM), the whole female mosquitoes were selected from each box of every blood group separately (10 females for each blood group ABO). In preparing specimens for the scanning electron microscope, the process is divided into two fixations, the primary and the secondary fixation. For the primary fixing process, the 2.5% glutaraldehyde in 0.1 M cacodylate buffer was used for the period of 2 h, followed by the three consecutive washing with the same buffer for the 30 min. While for the secondary fixing process, 1% osmium tetroxide was used for the 2 h. Then the samples were rinsed for the final time with the 0.1 M cacodylate buffer three times for 30 min.

The ovaries were dehydrated by using the graded series of acetone (50%, 70%, 80%, 90%, and 100%). The dehydrated ovaries were then transferred to the critical point drying apparatus. The recommended quantity of acetone solution was also poured into the drying chamber to avoid over-drying. Liquid nitrogen was also added into the drying critical point drying chamber. The CO2 and acetone were allowed to be mixed freely; the same process was repeated eight times to confirm the drying of the specimen. The dried mosquito specimens were mounted over the stubs, and the specimens tubes were coated with a thin layer of silver. Gold-spotted SCD005 was used, and then the samples were photographed with SEM.

**Electroantennography (EAG).** To measure the response of mosquitoes to each human blood group type, EAG recordings from one antenna of *An. stephensi* female mosquito were made. Unfed female mosquitoes were anesthetized by the use of CO2 and were permanently fixed on the reference electrode by the use of spectra 360 electrode gel. It was made sure that the mosquito was completely immobile except for the antennae. The tips of the antennae were pressed into the small drop of electrode gel on the recording electrode. Both of the electrodes are silver wires coated with silver chloride with a diameter of 0.2 mm. The experimental preparations were done in continuous airflow (600 mL/min, 1.5 m/s) by the Helon tube of 0.7–0.8 cm diameter, containing about 100 mL/min dry air and the 600 mL/min moist air passed through the charcoal filter. At this stage, little modification was done in the structure of the electroantennogram, and an artificial blood feeder of mosquitoes with the membrane was attached to the same.

The blood feeder was packed in a glass jar through which continuous airflow was passed, and this air flow ends as stimulus near the mounted mosquito. The diameter of the glass tube was 0.5 mm, and the flow of air was controlled by using an ON/OFF switch; these bursts of 0.5 s of air from the blood jar were provided as a stimulus to host-seeking mosquitoes. The blood groups were tested systematically together with the control group. The amplifier amplified the generated signals while the well-known software decoded the recordings (EAG 2000, Syntech, Hilversum, and the Netherlands). All of the test blood groups were also dissolved separately in tetyl-butyl-methyl ether (MTBE), and about 30 uL of this test solution was applied onto a piece of filter paper (1.5–2 cm). About 30 min was given to the TMME solution to evaporate from the filter paper leaving behind the blood; then, this piece of filter paper was placed into the Pasteur pipette. In the case of the control treatment, distilled water was used, and the same treatment was applied with the distilled water for the test compounds. The stimulus controller CS-04/b, Syntech, was used to inject the odor cues originating from the treated filter paper in the Pasteur pipette into the humidified and filtered air stream directed towards the antennae of immobilized mosquitoes. All olfactory stimuli were tested randomly against different mosquito specimens with a total of five specimens exposed to each of the human blood type groups and control.

To minimize the chances of error and to test the electrophysiological activity of the stuck female mosquito, lactic acid, 1-octen-3-01, and isovaleric acid were used as known stimulators. After that, each blood group was replicated three times to record the activity of the olfactory neurons of antennae. All the treatments of blood groups were tested randomly, and a regular interruption of control stimulus (0.1% lactic acid) was done. The regular interruption of the control stimulus was used to control the activity of the antennae. All the stimulants were expressed as a mean percent response to the control treatment. The response of different female mosquitoes to human blood groups was indicated as a mean percent response. The results were analyzed by the use of the Student’s t-test.

**Wind tunnel bioassays.** Wind tunnel bioassays were used to determine the response of *An. stephensi* to four human blood groups (A, B, AB, and O) and control stimuli (distilled water). Wind tunnel bioassays have been used to evaluate the response of *Ae. aegypti*, *Cx. quinquefasciatus* and *Cx. nigripalpus* towards the blood volatiles. A dual choice wind tunnel was converted into a “five-choice” tunnel with all five glass tubes having glass jars at their end with openings to accommodate an artificial blood feeder. A continuous flow of warm water ensured the blood remained in liquid and produced its specific smell.

A batch of approximately 100 female mosquitoes was released at the downwind end of the tunnel in the air stream coming from the five upwind end chambers. After 30 min, the numbers of mosquitoes in each of the five glass jars were counted. The mosquitoes were then sent back towards the downwind end of the wind tunnel, and the positions of odor cues were changed, including the control. Before the second time release, the fresh air was passed through the tunnel. Again the mosquitoes were released from the releasing box, and the response of the mosquitoes towards the new cues and the number of mosquitoes in each chamber at the upwind end was counted after 30 min; the same process was repeated, and for the third time with randomization. The same process was repeated 10 times with each blood group and with a new batch of mosquitoes each time.
To test the response of female mosquitoes towards human-emitted olfactory cues, an olfactometer was used in previous studies. The olfactometer test was conducted in the control room; the temperature was 27 ± 2 °C with 70–80% relative humidity. The optimum activity of the An. stephensi was observed late at night, so the experiment was conducted at 2–6 AM.

Steel balls rubbed in the hands of persons (ten volunteers per blood group) of having ABO blood groups along with the few drops of blood group-specific sweat were placed in the glass jars at the upwind end of the olfactometer. Approximately 100 female mosquitoes were released at the downwind end of the olfactometer from the releasing cage. After 30 min, the total number of mosquitoes in each box at the upwind end was counted, including the control. After that, the mosquitoes were returned to the releasing cage; then, the positions of steel balls at the upwind end of the glass jar of the olfactometer were changed randomly to decrease the biases from the data. To remove the smell of a sweat from the olfactory tube after cleaning, the fresh air was passed for about 10 min continuously. Mosquitoes were then again allowed to enter into the olfactometer, and after 30 min, the total number of mosquitoes was counted. The same process was repeated for the third time. The same experiment was repeated 10 times with different persons and mosquitoes to decrease the chances of error. A new batch of mosquitoes was selected for each replication.

**Statistical analysis.** The mean number of eggs of An. stephensi were evaluated with the help of a linear model (ANOVA) and Tukey’s test on Minitab® software (12.2, version, Minitab). Before performing the ANOVA, with the help of angular transformation (arcsine √x), egg viability was also transformed for fertility. Data obtained were analyzed using R 3.2.2 software. The Shapiro–Wilk normality test was carried out, which showed that the data were not normally distributed. Hence, the Kruskal–Wallis Chi-square test was used to compare the averages of the responses of An. stephensi in relation to blood-group treatments.

**Ethics statement.** All applicable international, national, and/or institutional guidelines for the care and use of animals were followed.

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The authors declare no competing interests.

Additional information

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