RESEARCH ARTICLE

Water-induced strong protection against acute exposure to low subzero temperature of adult Aedes albopictus

Meichun Zhang1,2,3*, Dongjing Zhang1,2,3*, Yongjun Li1,2,3, Qiang Sun1, Qin Li4, Yali Fan1,2,3, Yu Wu1,2,3, Zhiyong Xi1,5*, Xiaoying Zheng1,2,3*

1 Sun Yat-sen University—Michigan State University Joint Center of Vector Control for Tropical Diseases, Zhongshan School of Medicine, Sun Yat-sen University, Guangzhou, China, 2 Key Laboratory for Tropical Disease Control, Ministry of Education, Sun Yat-sen University, Guangzhou, China, 3 Guangdong Provincial Engineering Technology Research Center for Diseases-Vectors Control, Sun Yat-sen University, Guangzhou, China, 4 Section of Pulmonary, Critical Care and Sleep Medicine, Department of Internal Medicine, Yale School of Medicine, New Haven, CT, United States of America, 5 Department of Microbiology and Molecular Genetics, Michigan State University, East Lansing, MI, United States of America

☯ These authors contributed equally to this work.
* xizy@msu.edu (ZX); zhengxy@mail.sysu.edu.cn (XZ)

Abstract

As an important vector of dengue and Zika, Aedes albopictus has been the fastest spreading invasive mosquitoes in the world over the last 3–4 decades. Cold tolerance is important for survival and expansion of insects. Ae. albopictus adults are generally considered to be cold-intolerant that cannot survive at subzero temperature. However, we found that Ae. albopictus could survive for several hours’ exposure to -9 to -19 °C so long as it was exposed with water. Median lethal time (LT50) of Ae. albopictus exposed to -15 and -19 °C with water increased by more than 100 times compared to those exposed to the same subzero temperature without water. This phenomenon also existed in adult Aedes aegypti and Culex quinquefasciatus. Ae. albopictus female adults which exposed to low subzero temperature at -9 °C with water had similar longevity and reproductive capacity to those of females without cold exposure. Cold exposure after a blood meal also have no detrimental impact on survival capacity of female adult Ae. albopictus compared with those cold exposed without a blood meal. Moreover, our results showed that rapid cold hardening (RCH) was induced in Ae. albopictus during exposing to low subzero temperature with water. Both the RCH and the relative high subzero temperature of water immediate after cold exposure might provide this strong protection against low subzero temperature. The molecular basis of water-induced protection for Ae. albopictus might refer to the increased glycerol during cold exposure, as well as the increased glucose and hsp70 during recovery from cold exposure. Our results suggested that the water-induced strong protection against acute decrease of air temperature for adult mosquitoes might be important for the survival and rapid expansion of Ae. albopictus.
Author summary

*Aedes albopictus* is one of two most important vectors for dengue and zika. During the last 3–4 decades, this mosquito has spread from native Asian area to all continents except Antarctica, becoming the most invasive mosquitoes which imposed extensive public health threat to human beings throughout the world. Cold tolerance is important for distribution and survival of insects. During the expansion of *Ae. albopictus*, especially a spatial expansion to cooler climate areas, it needs to cope with cold temperatures. Moreover, because of such widespread distribution adult *Ae. albopictus* will certainly often encounter sudden drops in air temperature even below subzero that often happens in early spring and winter, and late autumn. Thus far, adult *Ae. albopictus* are generally considered to be cold-intolerant that can not survive at subzero temperature. In this study, we found that water can provide strong protection against low subzero temperature even below -10 °C. Cold exposure of adult female *Ae. albopictus* to low subzero temperature with water either before or after a blood meal have no detrimental impact on fitness costs of these adult mosquitoes. Considering water is common in nature, our results indicated that during the expansion of *Ae. albopictus* especially when adult mosquitoes encounter a sudden drop in air temperature water could be a good shelter for cope with such cold temperature below subzero.

Introduction

Cold tolerance or cold hardiness, the ability of an insect to survive at low temperature, is important in defining the distribution and survival of insects. There are two different cold hardening in insects at present. One is accomplished by long term (weeks or months) cold acclimatization to overwinter that occurs in an inactive or diapausing stage; the other called rapid cold hardening (RCH) which is accomplished by a brief exposure (minutes or hours) to low temperature that occurs even in feeding and reproductive stages [1–4]. As an efficient ability utilized by insects to survival in environment with rapid and unexpected changes in temperature, RCH has been found in numerous insect species belonging to different orders including Diptera [2, 4–7].

*Aedes albopictus* is an epidemiologically important vector for several arboviruses such as dengue, yellow fever, zika, and chikungunya. During the last 3–4 decades, *Ae. albopictus* has spread from native Asian area to all continents except Antarctica, becoming the most invasive mosquitoes which imposed extensive public health threat to human beings throughout the world [8–12]. During the spread of this species, a spatial expansion to cooler climate areas has also been reported and the ability to rapidly produce low temperature phenotypes has been considered as an important factor for the successful establishment in these cooler habitats [13]. Therefore, cold hardiness is a key trait for the distribution of this species and strong resistance to cold temperature provides more chances for efficient invasiveness to colder zones.

Adult *Ae. albopictus* are generally considered to be cold-intolerant that can not survive at subzero temperature and the only life stage that can cope with this low temperature is its eggs [13, 14]. Thus, the studies on the cold hardiness of this mosquito have been focused on eggs [13–19]. Moreover, the RCH has not been reported in adult and eggs of this species yet. In our current study, we found that adult *Ae. albopictus* could survive for several hours’ exposure under -10 °C when transferred from room temperature to low subzero temperature with water, which was also found in adult *Aedes aegypti* and *Culex quinquefasciatus*. Median lethal time (LT$_{50}$) of adult mosquitoes was increased by nearly 100 times compared to those
transferred directly to subzero temperature lower than -10 °C without water. This indicated that RCH was also existed in adult mosquitoes and that water in nature might provide strong protection for adult mosquitoes against sudden drop in air temperature that often happens in early spring and winter, and late autumn [2, 4, 6, 7]. The aim of this study is to compare the cold hardiness of adult Ae. albopictus exposed to low subzero temperature with water with those exposed directly, analyze the possible molecular mechanism for this cold hardiness and determine the impact of exposure to low subzero temperature with water on fitness costs of adult female Ae. albopictus.

Materials and methods

Mosquito strain and rearing
Mosquitoes used in this study including Ae. albopictus, Ae. aegypti and Cx. quinquefasciatus were all established for several years in our laboratory with Guangdong origin. All mosquitoes were reared in a climate-controlled room at 28 ± 1˚C and 80 ± 5% relative humidity with a 12:12-hour (light: dark) photoperiod. Adult mosquitoes were provided with 10% glucose solution.

Subzero temperature exposure
To evaluate the survivorship, 3 groups of 20 adult mosquitoes with 3- to 5-day-old were immobilized by CO2 and then transferred to a disposable 100ml-plastic cup with (treatment) or without (control) dechlorinated tap water (50 mL). A plastic lid was used to cover with cup for preventing escape of mosquitoes during experiment. These adults were then allowed to recover from anaesthetization at room temperature for 1 h. Subsequently, the cups of mosquitoes with water were exposed to low subzero temperatures at -9, -15 and -19 °C for 3 to 8 h, and the cups of mosquitoes without water were exposed to the same temperatures for 3 to 30 min. Twenty adults (one cup) were removed from each temperature at a 1 h interval for cups with water and 1- to 5-min interval for cups without water until 100% mortality were attained. The exposed mosquitoes were then transferred to 650ml-plastic cages with a piece of wet filter paper to keep humidity and maintained at normal climate-controlled room. The mosquito survival was recorded 24 h after cold exposure and survival was defined as the ability of righting themselves and flies [20].

Fitness determinants
To evaluate the impact of cold exposed to low subzero temperature on fitness costs of female adult Ae. albopictus, 3 groups of 30 adult mosquitoes with 3- to 5-day-old were cold exposed to -9 °C with water for 3 to 5 h as described above. Subsequently, the exposed mosquitoes were transferred to cages, maintained under normal climate-controlled room and provided with 10% glucose solution. Control groups of 90 female mosquitoes without cold exposure were also transferred to cages and maintained under normal condition. Three days after recovery from cold exposure dead mosquitoes were discarded and survived mosquitoes that have starved for 24 h were blood fed on mice for 30 min. The engorged mosquitoes were then counted and transferred to new cages, and were aspirated into individual 50 mL Corning tubes 2 days post-blood meal (PBM) with bottom lining of moist filter paper supported by water-soaked cotton [21]. Two days after oviposition, mosquitoes were aspirated out and killed by cold, eggs were removed out for maturation for 3 days and then counted. After maturation, eggs on filter paper were immersed in 50 mL water in a 100ml-plastic cup and egg hatch rates were determined by counting the number of hatched second instar larvae [22]. Adult lifespan
of female *Ae. albopictus* after cold exposure to -9 °C with water for 3 h were also monitored and compared with control group without cold exposure. Cold treatments were conducted as above and dead mosquitoes were removed at 24 h after recovery. Survived mosquitoes were reared under normal condition and dead mosquitoes were counted and removed daily for a month.

To evaluate the impact of cold exposure on blood-fed *Ae. albopictus*, engorged mosquitoes were collected and transferred to a new cage. Three groups of 30 adult mosquitoes were selected at 24 and 48 h PBM and exposed to -9 °C with water for 3 h. Mosquitoes without a blood meal were also selected and cold exposed at the meanwhile. Cold exposed mosquitoes were maintained in cages under normal condition and survivals were recorded 24 h after recovery. Two days after blood meal, mosquitoes of cold exposure at 24 h PBM were reared individually, and mosquitoes of cold exposure at 48 h PBM were reared as a pool, egg numbers and egg hatch rates were determined as above.

**Water temperature monitoring**

After transferring water from room temperature into low subzero temperature, the dynamics of water temperature was monitored by a mini-thermometer (Testo 175 H1, Lenzkirch, Germany). The temperature probe was wrapped by plastic film and immersed into 50 mL water in a disposable 100ml-plastic cup and then transferred to different low subzero temperature for 12 h. Temperature was detected and recorded at a 5-min interval.

**Glycerol and glucose analysis**

Adult mosquitoes were cold exposed as above and collected at 0, 1, 2 and 4 h after recovery from cold treatment. Six adults were pooled in a single replicate and five replicate biological assays were performed. These sampled mosquitoes were homogenized in 300 μL distilled deionized water. Two hundred microlitres of the homogenates were used for RNA extraction and another 100 μL were filtered at room temperature through a spin filter (Pall, nanosep 10k Omega, NY) at 12,000 rpm for 15 min. For glycerol analysis, filtered homogenates were diluted 1:100 v/v with distilled deionized water. Next, 10 μL of the diluted homogenates were incubated with 100 μL Master Reaction Mix (Sigma-Aldrich, MAK117, USA) for 20 min at room temperature. Absorbance was measured at 570 nm and glycerol contents were calculated from standard curve. Glucose levels were determined by using the Glucose (GO) Assay Kit (Sigma-Aldrich, GAGO-20, USA) according to the manufacturer’s protocol with minor modifications. The above filtered homogenates were diluted 1:5 v/v with distilled deionized water and 50 μL of these diluted samples were then incubated with 100 μL Assay Reagent for 30 min at 37 °C. After the incubation, 100 μL of 12N H2SO4 were added to stop the reaction. Then, the absorbance was measured at 540 nm and glucose contents were calculated from standard curve.

**Real-time PCR**

Total RNA was extracted from samples collected above using TRIZol Reagent (Invitrogen, Carlsbad, CA) and the first strand cDNA was synthesized using HiScript II Q SuperMix for qPCR (+dDNA wiper) (Vazyme, Nanjing, China) following the manufacturer’s protocol. Relative expression level of Hsp70 mRNA was performed by quantitative real-time PCR (qPCR) on the LightCycler96 Detection System (Roche, Mannheim, Germany) using TB Green Premix Ex Taq II (Tli RNaseH Plus) (TaKaRa, Otsu Shiga, Japan). The primer sequences for Hsp70 were forward (5’-TACCAACGGGCACACTCAC-3’) and reverse (5’-TTGCGGATGTCTTACCCCC-3’). Each reaction consisted 0.5 μL of cDNA, forward and reverse primer (10 μM), 10 μL of TB Green Premix Ex Taq II (2×), and 8.5 μL of distilled water.
deionized water to a final volume of 20 μL. The qPCR program was 95 °C for 30 sec, then 40 cycles of 95 °C for 5 sec and 60 °C for 30 sec followed by a melt-curve analysis. *Ae. albopictus* rpS7 was used as internal control and the relative Hsp70 expression of cold exposed samples were calibrated by samples collected at room temperature that without cold treatment. The relative expression levels of Hsp70 were determined by using the $2^{-\Delta\Delta CT}$ calculation method [23].

**Statistical analyses**

All data were analyzed by using SPSS statistics 19. Comparison of LT$_{50}$, egg numbers and egg hatch rates per female (except egg hatch rate of mosquitoes cold exposed at 48h PBM) between different groups were conducted using Student’s *t*-test. Comparison of the percent of mosquitoes imbibed a blood meal between cold exposed and non-exposed groups, of the survival of mosquitoes cold exposed at 24 h PBM with those without blood meal and of the egg hatch rate of mosquitoes cold exposed at 48 h PBM with those maintained at room temperature were performed using chi-square test. Comparisons of glycerol, glucose and hsp70 mRNA levels between groups were assessed using ANOVA followed by Tukey’s multiple comparison.

**Accession number**

The GenBank accession number of hsp70 mentioned in the text is JN132155.1.

**Results**

**Survival of adult mosquitoes experienced acute exposure to low subzero temperature with water**

We found that adult *Ae. albopictus* could survive for several hours’ exposure to subzero temperature even below -10 °C when it was transferred from room temperature to the low temperature with water. Subsequently, we analyzed the survival of adult female *Ae. albopictus* that transferred to different low subzero temperatures from -9 °C to nearly -20 °C with water and compared to the mosquitoes that directly transferred to the same temperatures without water. The results showed that when exposed adult mosquitoes to low subzero temperature with water the cold tolerance of these mosquitoes were strongly increased compared to those directly exposed without water (Fig 1). About 70 to 45% of these mosquitoes survived a 5 to 2 hours’ exposure to -9 to -19 °C when exposed with water. However, mosquitoes that directly transferred to the same low temperature without water were found to 100% mortality within 3 to 30 min. The LT$_{50}$ of these adult *Ae. albopictus* were about 292, 184 and 106 min for those exposed to -9, -15 and -19 °C with water, respectively, and these were 13.6, 145.1 and 108.7 times longer than those exposed to the same low temperature without water (Table 1). Moreover, we did the same analysis on adult male *Ae. albopictus*, female *Ae. aegypti* and *Cx. quinquefasciatus* to see if the phenomena also existed in male and other mosquitoes. The results suggested that when transferred these adult mosquitoes from room temperature to low subzero temperature (-15 °C) with water the cold tolerance were also significantly increased compared to those transferred to the same low temperature without water (Fig 2 and Table 2).

**A rapid cold-hardening response of adult *Ae. albopictus* was induced when exposed to low subzero temperature with water**

When transferring from room temperature to low subzero temperature with water adult mosquitoes will fall on the surface of water after anaesthetized by cold, we detected the change in temperature of water during this process (Fig 3). We found that when transferring from room
temperature to -9, -15 and -19 °C, the water temperature would reach subzero and iced within 55, 45 and 25 min, respectively. After that the temperature of ice kept at a relative high level of -2, -3 and -5˚C for 6, 3 and 2 h, respectively, and then rapidly decreased to the level equal to the low ambient air temperature. To evaluate whether it is the relative high temperature of ice just after dropping under subzero that protected adult mosquitoes from low temperature, we analyzed the survival of adult female *Ae. albopictus* when exposed to these relative high subzero temperature without water (Fig 4 and Table 1 in parentheses). The results indicated that when exposed to -2 to -5˚C, 100% mortality of these adult mosquitoes were reached within 60 to 100 min and that the LT$_{50}$ were about 85, 57 and 38 min for those exposed to -2, -3 and -5 ºC, respectively. Even subtracting the time needed to cool water from room temperature to the relative high subzero temperature, the LT$_{50}$ of adult female *Ae. albopictus* transferred to -9 to -19 ºC with water still were 2.7 to 2.0 times longer than those directly exposed to -2 to -5 ºC without water (Table 1). These results indicated that when exposed adult mosquitoes to low subzero temperature with water a RCH response was induced during this process.

**Fitness costs of female *Ae. albopictus* exposed to low subzero temperature with water**

Since adult *Ae. albopictus* could survive several hours’ exposure to low subzero temperature with water, it is important to know the effects of cold exposure on the bite behavior and

### Table 1. Lethal time of 50% of adult female *Aedes albopictus* after exposure to different low temperature.

| Temperature (°C) | LT$_{50}$ (min) | Fold increase (Water/Dry) | $^4$P-value | $^5$P-value |
|------------------|-----------------|--------------------------|-------------|-------------|
|                  | Dry             | Water                    |             |             |
| -9 (-2.0°)       | 21.4 ± 1.58 (85.4 ± 1.39*) | 292.2 ± 6.56 | 13.6 (2.7*) | < 5.0×10$^{-7}$ | < 1.0×10$^{-6}$ |
| -15 (-3.0)       | 1.27 ± 0.17 (57.2 ± 2.78) | 184.3 ± 2.06 | 145.1 (2.3) | < 5.0×10$^{-5}$ | < 5.0×10$^{-7}$ |
| -19°C (-5.0)     | 0.98 ± 0.15 (38.3 ± 2.03) | 106.5 ± 7.89 | 108.7 (2.0) | < 5.0×10$^{-5}$ | < 0.0005 |

* The relative high subzero temperature of water immediate after decreased below subzero when transferring to corresponding low subzero temperature.

* The LT$_{50}$ of mosquitoes after exposure to corresponding high subzero temperature.

* Fold increase in LT$_{50}$ of mosquitoes after exposure to low subzero temperature with water compared to without water.

* Fold increase in LT$_{50}$ of mosquitoes after exposure to low subzero temperature with water compared to those exposed to the corresponding high subzero temperature without water (Time needed for decrease in temperature of water after cold exposure to subzero was subtracted).

[https://doi.org/10.1371/journal.pntd.0007139.g001](https://doi.org/10.1371/journal.pntd.0007139.g001)
reproductive capacity of female adults. Our results showed that after exposure to -9˚C for 3 and 5 h with water, there still were 70.2% and 56.7% of mosquitoes that successfully had a blood meal on mice, respectively, and there was no significant difference between cold exposed and non-exposed mosquitoes (Fig 5A). The fecundity (egg numbers per female) and egg hatch rates between cold exposed and non-exposed mosquitoes also had no significant difference except for the fecundity of mosquitoes exposed to -9˚C for 5 h (Fig 5B and 5C). Although the fecundity of these mosquitoes was significantly lower than the mosquitoes without cold exposure, they still could lay about 41 eggs per female after 5 hours’ exposure to -9˚C with water. Meanwhile, we found that blood meal had no impact on cold tolerance of adult female mosquitoes compared to mosquitoes without blood meal. There were no significant difference on survival capacity between blood-fed and non-blood-fed mosquitoes after 3 hours’ exposure to -9˚C with water at both 24 and 48h PBM (Fig 6A), and these mosquitoes could still lay viable eggs (Fig 6B and 6C). Indeed, the egg hatch rate of mosquitoes cold exposed at 48h PBM was significantly higher than mosquitoes without cold exposure after blood meal (Fig 6C). Moreover, the lifespan of adult female Aedes albopictus was also compared between cold exposure and no exposure, and no significant difference was observed (S1 Fig).

Glycerol and glucose levels in adult Aedes albopictus exposed to low subzero temperature

Since the cold tolerance was significantly increased when exposed adult mosquitoes to low subzero temperature with water, we analyzed the levels of two important cryoprotectants glycerol and glucose in whole body of adult Aedes albopictus that exposed to -15˚C with water for 2.7

![Fig 2. Survival of adult mosquitoes after exposure to low subzero temperature.](https://doi.org/10.1371/journal.pntd.0007139.g002)

![Table 2. Lethal time of 50% of adult mosquitoes after exposure to -15˚C.](https://doi.org/10.1371/journal.pntd.0007139.t002)
h and compared to those exposed to -3 and -15°C directly for 1.5 h and 3 min, respectively. The duration of time were chose because at this duration mosquitoes exposed to -3 and -15°C directly were 100% mortality but mosquitoes exposed to -15°C with water were less than 30% mortality at the same duration of time as those exposed to -3°C directly after subtracting the time needed to cool water from room temperature to -3°C. The results showed that the

Fig 3. The change of water temperature after transferring from room temperature to different low subzero temperature. The temperatures of water were monitored by a mini-thermometer at 5-min interval. The mini-thermometer with ice was removed from subzero temperature at 12h after exposure and the ice was naturally melted. 
https://doi.org/10.1371/journal.pntd.0007139.g003

Fig 4. Survival of adult female *Ae. albopictus* after exposure to high subzero temperature without water. Adult female *Ae. albopictus* were transferred from room temperature to different high subzero temperatures without water. Adult mosquitoes were removed at 10-min interval from subzero temperature and recovered under normal rearing condition until 100% mortality were reached. N = 3 groups of 20 adult mosquitoes for each data point. 
https://doi.org/10.1371/journal.pntd.0007139.g004
glycerol level of mosquitoes exposed to low temperature with water was 6.2 μmol per mosquito just after recovery from cold exposure and this was significantly higher than those directly exposed without water and those maintained at room temperature (Fig 7A). Another, the glycerol level of mosquitoes exposed to low temperature with water was significantly decreased at 1 h after recovery from cold exposure. However, the glycerol levels of mosquitoes exposed to subzero temperature without water had no difference with time after recovery from cold exposure. The glucose levels of mosquitoes exposed to -15 °C without water were significantly

![Graph A](image1.png)

**Fig 5. Fitness costs of female *Ae. albopictus* exposed to low subzero temperature with water.** Adult female *Ae. albopictus* were transferred from room temperature to -9 °C with water and removed after 3 and 5 hours' exposure. Adult mosquitoes were recovered in cages under normal rearing condition. Live mosquitoes were fed on mice for about 30 min. NTC were mosquitoes maintained under normal rearing condition and fed on a blood meal. The number of mosquitoes that imbibed a blood meal (A), mean egg number (B) oviposited by and mean egg hatch rate (C) from individual female mosquitoes. Statistical differences between the numbers of blood-fed mosquitoes were determined by chi-square test and between egg numbers, egg hatch rates of individual female mosquito were determined by Student’s *t*-test. Error bars indicate SEM. "P<0.01; NS, not significant. N = 3 groups of 30 adult mosquitoes were cold exposed and 30 blood-fed mosquitoes were individually reared and laid eggs.

https://doi.org/10.1371/journal.pntd.0007139.g005

![Graph B](image2.png)

![Graph C](image3.png)

**Fig 6. The impact of exposing to low subzero temperature with water on survival, fecundity and fertility of female *Ae. albopictus* after a blood meal.** Adult female *Ae. albopictus* were transferred from room temperature to -9 °C with water at 24 and 48h PBM. The mosquitoes were recovered in cages under normal rearing condition after 3 hours’ cold exposure. Groups of mosquitoes without blood meal were also cold exposed as above. (A) The survival of mosquitoes were recorded 24h after recovery. The egg numbers (B) were counted 2 days after oviposition and egg hatch rate (C) were determined at second instar larvae. Statistical differences between the survivals of different groups, egg hatch rates between cold treated mosquitoes at 48h PBM and not treated mosquitoes (NTC) were determined by chi-square test, and between egg numbers of cold treated mosquitoes at 24h PBM and NTC were determined by Student’s *t*-test. The NTC groups that compared with cold treated mosquitoes at 24h PBM of (B) and (C) were the same as NTC groups from Fig 5 that compared with groups of -9 °C 3h. Error bars indicate SEM. """"P<0.0001; NS, not significant. N = 3 groups of 30 blood-fed and non blood-fed mosquitoes were cold exposed and 30 blood-fed mosquitoes were individually reared and laid eggs.

https://doi.org/10.1371/journal.pntd.0007139.g006
Fig 7. The change of glycerol and glucose levels and the expression of Hsp70 mRNA in adult female *Ae. albopictus* after exposure to different subzero temperature. Adult female *Ae. albopictus* were transferred from room temperature to -3 °C without water (blue) and -15 °C with (red) or without water (green). Adult mosquitoes were then removed from subzero temperature after 90 min for exposure to -3 °C, 3 min for exposure to -15 °C without water and 160 min for exposure to -15 °C with water and recovered under normal rearing condition. The contents of glycerol (A) and glucose (B) and the expression levels of hsp70 mRNA (C) were analyzed at 0 to 4h after recovery from cold exposure. The data shown are means ± SEM of five replicates with six whole mosquitoes for each. Columns that do not share the same letter are significantly different from each other within each group. The differences in glycerol and glucose contents and hsp70 mRNA levels were assessed by ANOVA followed by Tukey’s multiple comparison.

*P* < 0.05, **P** < 0.01, ***P*** < 0.001.

https://doi.org/10.1371/journal.pntd.0007139.g007
increased with time after recovery from cold exposure (Fig 7B). While those exposed to -3 \(^\circ\)C directly and to -15 \(^\circ\)C with water had no difference in glucose levels with time after recovery. However, the glucose level of mosquitoes exposed to -15 \(^\circ\)C with water was significantly higher and lower than those exposed to -3 and -15 \(^\circ\)C directly at 4 h after recovery from cold exposure, respectively. These results implied that glycerol and glucose might play important role in water-induced RCH of adult mosquitoes but at different stage of cold exposure.

**Hsp70 expression in adult Ae. albopictus exposed to low subzero temperature**

Because the upregulation of heat shock protein 70 (Hsp70) have been reported in several insect species in response to cold temperature [24–27], we wondered if this protein was also involved in mosquitoes to cope with cold temperature. We found that Hsp70 expression gradually increased with time after recovery from cold exposure of mosquitoes exposed to -3 \(^\circ\)C directly and to -15 \(^\circ\)C with water and the expression levels were the highest at 4 h after recovery (Fig 7C). The Hsp70 expression of mosquitoes exposed to -15 \(^\circ\)C without water was significantly increased at 1 h after recovery from cold exposure and maintained at these high levels till 4 h after recovery and the expression levels from 1 to 4 h after recovery of these mosquitoes were significantly higher than that exposed to -3 \(^\circ\)C directly and to -15 \(^\circ\)C with water. The results demonstrated that the degree of cold shock of mosquitoes exposed to -15 \(^\circ\)C with water was similar to those exposed to -3 \(^\circ\)C directly and both of them were significantly lesser than those exposed to -15 \(^\circ\)C without water.

**Discussion**

Adult *Ae. albopictus* has long been recognized as freeze-intolerant that can’t cope with subzero temperature. Most of the studies on cold hardiness of this species was focused mainly on eggs, which were the only life stage that can survive under subzero temperature as know so far [13–19]. However, in our current study, we found that when exposed adult *Ae. albopictus* to low subzero temperature (-9 to -19 \(^\circ\)C) with water it could survive several hours. Moreover, LT\(_{50}\) of these adult mosquitoes were increased by 13.6 to more than 100 fold changes when compared with the counterpart exposed without water. In consistent, this phenomenon also existed in *Ae. aegypti* and *Cx. quinquefasciatus*.

Cold tolerance or cold hardiness is important for the distribution and survival of insects. Over the last 3–4 decades, *Ae. albopictus* has spread from its native Asian area to all continents except Antarctica [8, 9, 12]. Such widespread distribution of *Ae. albopictus* implied the strong cold hardiness of this species and that they might have more chance to experience sharp decrease of air temperature than other local mosquitoes. Thus far, studies about the cold hardiness of *Ae. albopictus* has been focused on egg stage and previous studies indicated that they overwintered predominantly through diapause eggs [16, 17, 28–30]. A previous study showed that diapause eggs from *Ae. albopictus* could only survive for 1 hour under -12 \(^\circ\)C while non-diapause eggs could survive for 4 hours. In addition, neither *Ae. albopictus* nor *Ae. aegypti* eggs could be hatched after exposure to -15 \(^\circ\)C [19]. However, our results showed that both the adult of *Ae. albopictus* and *Ae. aegypti* could survive under -15 \(^\circ\)C for more than 3h-exposure (Figs 1 and 2). This indicated that the cold hardiness of adult mosquitoes was even stronger than eggs when exposed with water. The cold hardness of *Ae. albopictus* eggs was also highly correlated with the origin. Eggs from northern were more cold-hardy than those from southern, while eggs from tropical *Ae. albopictus* are much more susceptible to low temperature than those from temperate counterpart [15, 16, 18, 19]. Moreover, similar comparisons of *Ae. albopictus* larvae from different regions have been conducted at low temperature above zero.
So far as we knew, only adult *Culex pipiens* could tolerate for several to tens of hours’ exposure to subzero temperature (-5 °C) [20]. Interestingly, our results showed that the cold hardiness of adult *Ae. albopictus*, *Ae. aegypti* and *Cx. quinquefasciatus* could be significantly enhanced in the presence of water. When exposed to -15 °C with water these adult mosquitoes can survive for several hours more. This is the first report that adult *Aedes* mosquitoes cold also cope with low subzero temperature even below -10 °C so long as there are waters when these mosquitoes exposed to this low temperature. Meanwhile, our results suggested that water-induced enhancement of cold hardiness might be a universal phenomenon in adult mosquitoes.

This study showed that the relative high subzero temperature of water immediate after transferring to low subzero temperature just provided partial protection for adult *Ae. albopictus* against low subzero temperature (Table 1). Previous studies showed that RCH of arthropods could be induced through gradual cooling from 0.1 to 1 °C/min [4, 6]. We found that when transferred from room temperature to -9 to -19 °C, the cooling rate of water were about 0.42 to 0.95 °C/min till reaching the relatively high subzero temperature (Fig 3). These results demonstrated that RCH might be induced in these adult mosquitoes and provided partial protection during the process of cold exposure. Considering water is common in nature, our results suggested that RCH of adult mosquitoes induced by waters in nature might provide strong protection against acute decrease of air temperature to low subzero that would be lethal, which often happens in early spring and winter, and late autumn [2, 4, 7].

*Ae. albopictus* overwintered predominantly through diapause eggs, nevertheless, adult *Ae. albopictus* was also found occasionally during winter season [29, 32, 33]. This indicates these adults might experience the low subzero temperature sometimes during their lifetime like eggs do. *Ae. albopictus* has been considered to be the most invasive mosquito species worldwide and to be passively spread over long distance principally through the transportation of eggs by global shipments of used tires and other artificial containers [34–38]. In recent years, however, studies reported that, over a long distance, adult mosquitoes including *Aedes* species could be transported by aircraft [39–44], and, at a more regional level, adult *Ae. albopictus* are frequently transported by ground vehicles like cars [10, 37, 38]. These also pose a threat of experiencing low subzero temperature to the adult mosquitoes, especially those be transported to cooler climate areas. In the light of situations mentioned above, it is not known until now how the adult mosquitoes can cope with low subzero temperature in nature. In this study, we found that adult *Ae. albopictus* could cope with low subzero temperature even below -10 °C in the presence of water and that after exposure to low subzero temperature for several hours the adult mosquitoes could still bite, lay eggs, and eggs could hatch to larvae (Fig 5). In addition, after a blood meal female mosquitoes must find a micro-habitat with water to lay eggs. We found that, after cold exposure to low subzero temperature with water, blood-fed *Ae. albopictus* could still lay viable eggs (Fig 6). This implies that water in nature not only provide a micro-habitat for mosquitoes’ egg-laying but also can act as a shelter against acute decrease of air temperature. In a word, our results indicated that the water-conferred strong protection against low subzero temperature might be an important means for adult *Ae. albopictus* to survive the lethal low temperature and therefore might be important for the expansion of this species to cooler areas.

Glycerol is the most commonly produced cryoprotectant for insects to cope with freeze damage [45, 46]. High accumulation in haemolymph and tissues was important for overwintering survival of insects while lower glycerol content often resulted in higher overwinter mortality [46–49]. Furthermore, the accumulation of glycerol was also highly correlated with survival of some insects in RCH [1, 3, 50–52]. In this study, we found that the glycerol levels of the adult *Ae. albopictus* transferred from room temperature to -15 °C with water was
significantly higher than those exposed to -3 and -15 °C without water and those maintained at room temperature at 0h after recovery from cold exposure and then significantly decreased to normal level (Fig 7A). Because freeze damage might happen when adult mosquitoes anaesthetizing on -3 °C ice and underneath -15 °C air for a while, our study indicated that accumulated glycerol in adult *Ae. albopictus* during being exposed to -15 °C with water may confer strong protection to freeze damage and contribute to RCH induced by water.

Sugars are also important cryoprotectants in insects to eliminate or minimize freeze damage [45]. There were studies that the levels of glucose in some insects were increased in response to RCH or cold stress [50, 53–55]. Our results showed that the glucose level of adult *Ae. albopictus* exposed to -15 °C with water was significantly higher than those exposed to -3°C and lower than those exposed to -15 °C without water at 4 hours after recovery from cold exposure (Fig 7B). This results indicated that the accumulation of glucose in adult *Ae. albopictus* was important during recovery from cold exposure but not in the process of cold exposure. Moreover, we found that glucose levels went through significant changes with time-dependent manner during recovery from cold exposure to -15 °C without water. Heat shock protein 70 (Hsp70) also played an important role in cold hardiness of overwintering and cold stress of insects [27, 56, 57]. Our results showed that the expression of Hsp70 in adults *Ae. albopictus*, exposed to -3 or -15 °C with or without water, were all significantly up-regulated during recovery from cold exposure (Fig 7C). This is consistent with previous studies of different insects (including *Culex pipiens*) that Hsp70 expression were up-regulated during recovery from exposure to subzero temperature [20, 57–59]. Our results and others indicated that the up-regulation of Hsp70 might be required for the repair of cold injury caused by cold exposure [27]. Hsp70 protein went through obvious changes when exposing adult *Ae. albopictus* to -15 °C without water, which was similar with glucose. It increased dramatically by 6 fold from 1 to 4 h after recovery compared with those maintained under room temperature. This implied that severe acute cold shock might be happened in adult *Ae. albopictus* exposed to -15 °C without water and caused the serious disorders of glucose metabolism that eventually lead to the death of these adult mosquitoes.

In conclusion, adult mosquitoes especially *Ae. albopictus* and *Ae. aegypti*, which are the most important vectors for dengue and Zika virus, could survive at low subzero temperature even below -10 °C for several hours’ exposure in the presence of water and this cold exposure have no detriment impact on fitness costs of adult *Ae. albopictus*. Both the relative high sub-zero temperature of water immediate after cold exposure and RCH induced by gradual cooling of water provided this strong protection against low subzero temperature. The cold tolerance might be conferred by accumulation of glycerol during cold exposure stage, and contributed by both glucose accumulation and Hsp70 up-regulation during the recovery stage from cold exposure. The RCH of adult mosquitoes induced by waters in nature might provide strong protection against acute decrease of air temperature to low subzero temperature, which often happens in early spring and winter, and late autumn, and this might be important for the survival and rapid expansion of *Ae. albopictus* to cooler areas. Our subsequent studies would be performed further to identify whether water-induced protection could be eliminated by down-regulation of glycerol or Hsp70, and cold hardiness of eggs when exposed to low subzero temperature with water.

**Supporting information**

S1 Fig. The impact of cold exposure to low subzero temperature with water on the life span of adult female *Ae. albopictus*. Adult female *Ae. albopictus* were transferred from room temperature to -9 °C with water for 3 h and then transferred to a cage. Survived mosquitoes
were transferred to a new cage at 24h after recovery and reared under normal condition. Dead mosquitoes were counted and removed daily for a month for comparing the life span of adult female *Ae. albopictus* with cold exposure (blue) with those without cold exposure (red). N = 4 groups of 30 adult mosquitoes were cold exposed. (TIF)

**Acknowledgments**

The authors wish to thank Hongxin Ou for rearing the mosquito samples for this experiment.

**Author Contributions**

**Conceptualization:** Meichun Zhang, Dongjing Zhang, Yu Wu, Zhiyong Xi, Xiaoying Zheng.

**Data curation:** Meichun Zhang, Dongjing Zhang, Yongjun Li, Qiang Sun, Qin Li.

**Formal analysis:** Meichun Zhang, Dongjing Zhang, Yongjun Li, Qiang Sun, Qin Li, Zhiyong Xi, Xiaoying Zheng.

**Funding acquisition:** Meichun Zhang.

**Investigation:** Meichun Zhang, Dongjing Zhang, Yongjun Li, Qiang Sun, Yali Fan.

**Methodology:** Meichun Zhang, Dongjing Zhang, Yongjun Li, Qiang Sun, Qin Li, Yali Fan.

**Project administration:** Zhiyong Xi, Xiaoying Zheng.

**Resources:** Zhiyong Xi, Xiaoying Zheng.

**Software:** Yu Wu.

**Supervision:** Zhiyong Xi, Xiaoying Zheng.

**Validation:** Meichun Zhang, Dongjing Zhang.

**Writing – original draft:** Meichun Zhang, Zhiyong Xi, Xiaoying Zheng.

**Writing – review & editing:** Meichun Zhang, Dongjing Zhang, Qin Li, Yu Wu, Zhiyong Xi, Xiaoying Zheng.

**References**

1. Park Y, Kim K, Kim Y. Rapid cold hardening of *Thrips palmi* (Thysanoptera: Thripidae). Environ Entomol. 2014; 43(4):1076–83. https://doi.org/10.1603/EN13291 PMID: 25182622

2. Yang G, Wen J, Han Y, Hou M. Rapid Cold Hardening Confers a Transient Increase in Low Temperature Survival in Diapausing *Chilo suppressalis* Larvae. Insects. 2018; 9(2). https://doi.org/10.3390/insects9020053 PMID: 29747426

3. Lee RE Jr., Chen CP, Denlinger DL. A rapid cold-hardening process in insects. Science. 1987; 238(4832):1415–7. https://doi.org/10.1126/science.238.4832.1415 PMID: 17800568

4. Ju RT, Xiao YY, Li B. Rapid cold hardening increases cold and chilling tolerances more than acclimation in the adults of the sycamore lace bug, *Corythucha ciliata* (Say) (Hemiptera: Tingidae). J Insect Physiol. 2011; 57(11):1577–82. https://doi.org/10.1016/j.jinsphys.2011.08.012 PMID: 21872604

5. Yang S, Zhang X, Wang J, Wang S, Pan Y, Zhang J, et al. Identification and analysis of up-regulated proteins in *Lissorhoptrus oryzophilus* adults for rapid cold hardening. Gene. 2018; 642:9–15. https://doi.org/10.1016/j.gene.2017.11.002 PMID: 29104168

6. Saeidi F, Moharrampour S, Mikani A. Rapid Cold Hardening Capacity and Its Impact on Performance of Russian Wheat Aphid (Hemiptera: Aphididae). Environ Entomol. 2017; 46(4):954–9. https://doi.org/10.1093/ee/nvx087 PMID: 28541434

7. Coleman PC, Bale JS, Hayward SA. Meat Feeding Restricts Rapid Cold Hardening Response and Increases Thermal Activity Thresholds of Adult Blow Flies, *Calliphora vicina* (Diptera: Calliphoridae). PLoS One. 2015; 10(7):e0131301. https://doi.org/10.1371/journal.pone.0131301 PMID: 26196923
23. Livak KJ, Schmittgen TD. Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods. 2001; 25(4):402–8. https://doi.org/10.1006/meth.2001.1262 PMID: 11846609

24. Rinehart JP, Robich RM, Denlinger DL. Enhanced cold and desiccation tolerance in diapause adults of Culex pipiens, and a role for Hsp70 in response to cold shock but not as a component of the diapause program. J Med Entomol. 2006; 43(4):713–22. https://doi.org/10.1603/0022-2585(2006)43[713:ECADTI]2.0.CO;2 PMID: 16892629

25. Sinclair BJ, Gibbs AG, Roberts SP. Gene transcription during exposure to, and recovery from, cold and desiccation stress in Drosophila melanogaster. Insect Mol Biol. 2007; 16(4):435–43. https://doi.org/10.1111/j.1365-2936.2007.00739.x PMID: 17506850

26. Huang LH, Wang CZ, Kang L. Cloning and expression of five heat shock protein genes in relation to cold hardening and development in the leafminer, Liriomyza sativa. J Insect Physiol. 2009; 55(3):279–85. https://doi.org/10.1016/j.jinphys.2008.12.004 PMID: 19133268
27. Stetina T, Kostal V, Korbelova J. The Role of Inducible Hsp70, and Other Heat Shock Proteins, in Adaptive Complex of Cold Tolerance of the Fruit Fly (Drosophila melanogaster). PLoS One. 2015; 10(6): e0128976. https://doi.org/10.1371/journal.pone.0128976 PMID: 26034990

28. Swanson J, Lancaster M, Anderson J, Crandell M, Haramis L, Grimmst P, et al. Overwintering and establishment of Aedes albopictus (Diptera: Culicidae) in an urban La Crosse virus enzootic site in Illinois. J Med Entomol. 2000; 37(3):454–60. PMID: 15535992

29. Romi R, Severini F, Toma L. Cold acclimation and overwintering of female Aedes albopictus in Roma. J Am Mosq Control Assoc. 2006; 22(1):149–51. https://doi.org/10.2987/8756-971X(2006)22[149:CAAOOF]2.0.CO;2 PMID: 16646341

30. Pluskota B, Jost A, Augsten X, Stelzner L, Ferstl I, Becker N. Successful overwintering of Aedes albopictus in Germany. Parasitol Res. 2016; 115(8):3245–7. https://doi.org/10.1007/s00436-016-5078-2 PMID: 27112761

31. Chang LH, Hsu EL, Teng HJ, Ho CM. Differential survival of Aedes aegypti and Aedes albopictus (Diptera: Culicidae) larvae exposed to low temperatures in Taiwan. J Med Entomol. 2007; 44(2):205–10. PMID: 17427687

32. Tsunoda T, Chaves LF, Nguyen GT, Nguyen YT, Takagi M. Winter Activity and Diapause of Hypera postica (Coleoptera: Curculionidae). Environ Entomol. 2017; 46(4):960–6. https://doi.org/10.1093/ee/nvx089 PMID: 28535265

33. Hofhuis A, Reimerink J, Reusken C, Scholte EJ, Boer A, Takken W, et al. The hidden passenger of Hypera postica (Diptera: Culicidae) larvae exposed to low temperatures in Taiwan. J Med Entomol. 2007; 44(2):205–10. PMID: 17427687

34. Eritja R, Palmer JRB, Roiz D, Sanpera-Calbet I, Bartumeus F. Direct Evidence of Adult Aedes aegypti and Aedes albopictus survival during winter in the Northwestern Italy. Ann Ig. 2017; 29(1):86–90. https://doi.org/10.7416/ai.2017.2135 PMID: 28067941

35. Hawley WA, Reiter P, Copeland RS, Pumpani CB, Craig GB Jr. Water-induced protection for adult mosquitoes against low subzero temperature. J Med Entomol. 2000; 37(16):6242–7. https://doi.org/10.1093/jme/tjv122 PMID: 16606847

36. Flacio E, Engeler L, Tonolla M, Muller P. Spread and establishment of Aedes albopictus in southern Switzerland between 2003 and 2014: an analysis of oviposition data and weather conditions. Parasit Vectors. 2016; 9(1):304. https://doi.org/10.1186/s13071-016-1577-3 PMID: 27229686

37. Sukehiro N, Kida N, Umezawa M, Murakami T, Arai N, Jinnai T, et al. First report on invasion of yellow fever mosquito, Aedes aegypti, at Narita International Airport, Japan in August 2012. Jpn J Infect Dis. 2013; 66(3):189–94. PMID: 23698478

38. Brown EB, Adkin A, Fooks AR, Stephenson B, Medlock JM, Snary EL. Assessing the risks of West Nile virus-infected mosquitoes from transatlantic aircraft: implications for disease emergence in the United Kingdom. Vector Borne Zoonotic Dis. 2012; 12(4):310–20. https://doi.org/10.1089/vbz.2010.0176 PMID: 22171811

39. Saeidi M, Moharramipour S. Physiology of Cold Hardiness, Seasonal Fluctuations, and Cryoprotectant Contents in Overwintering Adults of Hypera postica (Coleoptera: Curculionidae). Environ Entomol. 2017; 46(4):960–6. https://doi.org/10.1093/ee/nvx089 PMID: 28535265
47. Izumi Y, Sonoda S, Tsumuki H. Effects of diapause and cold-acclimation on the avoidance of freezing injury in fat body tissue of the rice stem borer, Chilo suppressalis Walker. J Insect Physiol. 2007; 53 (7):685–90. https://doi.org/10.1016/j.jinsphys.2007.04.005 PMID: 17543330

48. Trudeau M, Mauffette Y, Rochefort S, Han E, Bauce E. Impact of host tree on forest tent caterpillar performance and offspring overwintering mortality. Environ Entomol. 2010; 39(2):498–504. https://doi.org/10.1603/EN09139 PMID: 20388280

49. Boychuk EC, Smiley JT, Dahlhoff EP, Bernards MA, Rank NE, Sinclair BJ. Cold tolerance of the montane Sierra leaf beetle, Chrysomela aeneicollis. J Insect Physiol. 2015; 81:157–66. https://doi.org/10.1016/j.jinsphys.2015.07.015 PMID: 26231921

50. Michaud MR, Denlinger DL. Shifts in the carbohydrate, polyol, and amino acid pools during rapid cold-hardening and diapause-associated cold-hardening in flesh flies (Sarcophaga crassipalpis): a metabolic comparison. J Comp Physiol B. 2007; 177(7):753–63. https://doi.org/10.1007/s00360-007-0172-5 PMID: 17576567

51. Park Y, Kim Y. RNA interference of glycerol biosynthesis suppresses rapid cold hardening of the beet armyworm, Spodoptera exigua. J Exp Biol. 2013; 216(Pt 22):4196–203. https://doi.org/10.1242/jeb.092031 PMID: 23948473

52. Park Y, Kim Y. A specific glycerol kinase induces rapid cold hardening of the diamondback moth, Plutella xylostella. J Insect Physiol. 2014; 67:56–63. https://doi.org/10.1016/j.jinsphys.2014.06.010 PMID: 24973793

53. Overgaard J, Malmendal A, Sorensen JG, Bundy JG, Loeschcke V, Nielsen NC, et al. Metabolomic profiling of rapid cold hardening and cold shock in Drosophila melanogaster. J Insect Physiol. 2007; 53 (12):1218–32. https://doi.org/10.1016/j.jinsphys.2007.06.012 PMID: 17662301

54. Overgaard J, Sorensen JG, Com E, Colinet H. The rapid cold hardening response of Drosophila melanogaster: complex regulation across different levels of biological organization. J Insect Physiol. 2014; 62:46–53. https://doi.org/10.1016/j.jinsphys.2014.01.009 PMID: 24508557

55. Chowanski S, Lubawy J, Spochacz M, Ewelina P, Grzegorz S, Rosinski G, et al. Cold induced changes in lipid, protein and carbohydrate levels in the tropical insect Gromphadorhina coquereliana. Comp Biochem Physiol A Mol Integr Physiol. 2015; 183:57–63. https://doi.org/10.1016/j.cbpa.2015.01.007 PMID: 25624163

56. Rinehart JP, Li A, Yocum GD, Robich RM, Hayward SA, Denlinger DL. Up-regulation of heat shock proteins is essential for cold survival during insect diapause. Proc Natl Acad Sci U S A. 2007; 104 (27):11130–7. https://doi.org/10.1073/pnas.0703538104 PMID: 17522254

57. Colinet H, Lee SF, Hoffmann A. Temporal expression of heat shock genes during cold stress and recovery from chill coma in adult Drosophila melanogaster. FEBS J. 2010; 277(1):174–85. https://doi.org/10.1111/j.1742-4658.2009.07470.x PMID: 19968716

58. Yocum GD. Differential expression of two HSP70 transcripts in response to cold shock, thermoperiod, and adult diapause in the Colorado potato beetle. J Insect Physiol. 2001; 47(10):1139–45. PMID: 12770192

59. Kostal V, Tolarova-Borovanska M. The 70 kDa heat shock protein assists during the repair of chilling injury in the insect, Pyrrhocoris apterus. PLoS One. 2009; 4(2):e4546. https://doi.org/10.1371/journal.pone.0004546 PMID: 19229329