Freeze–thaw *Caenorhabditis elegans* freeze–thaw stress response is regulated by the insulin/IGF-1 receptor *daf-2*

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**Abstract**

**Background:** Adaption to cold temperatures, especially those below freezing, is essential for animal survival in cold environments. Freezing is also used for many medical, scientific, and industrial purposes. Natural freezing survival in animals has been extensively studied. However, the underlying mechanisms remain unclear. Previous studies demonstrated that animals survive in extremely cold weather by avoiding freezing or controlling the rate of ice-crystal formation in their bodies, which indicates that freezing survival is a passive thermodynamic process.

**Results:** Here, we showed that genetic programming actively promotes freezing survival in *Caenorhabditis elegans*. We found that *daf-2*, an insulin/IGF-1 receptor homologue, and loss-of-function enhanced survival during freeze–thaw stress, which required the transcription factor *daf-16/FOXO* and age-independent target genes. In particular, the freeze–thaw resistance of *daf-2(rf)* is highly allele-specific and has no correlation with lifespan, dauer formation, or hypoxia stress resistance.

**Conclusions:** Our results reveal a new function for *daf-2* signaling, and, most importantly, demonstrate that genetic programming contributes to freezing survival.

**Keywords:** *C. elegans*, Freeze–thaw stress response, Insulin/IGF-1 receptor *daf-2*, Transcription factor *daf-16/FOXO*

**Background**

Cold temperature is a critical environment stimulus to animals. Subzero temperatures especially may adversely affect animals by direct lethal effects and damage caused by ice formation [1]. The ability of animals to sense and respond to cold temperatures, even below freezing, is essential for survival in cold environments. Freezing is also widely used for many medical, scientific, and industrial purposes, such as strain preservation and organ preservation. Understanding how to enhance survival and maintain normal physiological functions in the presence of freeze stress is critical for animals in nature and human research.

Natural freezing survival in animals has been extensively studied. Previous studies demonstrated that animals survive in extremely cold weather by avoiding freezing or controlling the rate of ice-crystal formation in their bodies [2], which indicates that freezing survival is a passive thermodynamic process. However, response to freeze–thaw stress and other biological phenomena, such as longevity and hypoxia resistance, may be genetically programmed. Because it is a powerful model in molecular genetics, the nematode *Caenorhabditis elegans* is suitable for studying genetic response to freeze–thaw stress.

Insulin/insulin-like growth factor 1 (IGF-1)-like signaling is the best-characterized pathway that regulates the lifespan and other stress-resistance traits of *C. elegans*. The insulin/IGF-1 receptor homologue *daf-2* activates a conserved phosphatidylinositol-3-OH kinase (PI(3)K)/3-phosphoinositide-dependent kinase-1 (PDK1)/Akt signal transduction pathway, which prevents FOXO transcription factor *daf-16* entry into the nucleus [3–5]. *daf-2* reduction/loss-of-function (rf) produces a longer lifespan [6]. Conversely, the PTEN phosphatase homologue, *daf-18*(rf), suppresses life-span extension induced by *daf-2(rf)* [7, 8].
Insulin/IGF-1 signaling is also involved in formation of dauer larvae, which have an alternative, developmentally arrested third larval stage (L3) [9]. In unfavorable environments, such as crowding or food shortage, insulin/IGF-1 signaling or transforming growth factor-beta (TGF-β) is suppressed; unliganded nuclear receptor DAF-12 regulates dauer diapause [10]. Insulin/IGF-1 signaling is also involved in other stress tolerances, including oxidative stress, ultraviolet light, heat shock, or hypoxia stress. The longevity and stress tolerance produced by insulin/IGF-1 signaling mutants require nuclear translocation or nuclear activity of daf-16 [11]. Savory et al. [12] showed that daf-16 is important for delta-9 desaturase gene expression, which is important for survival at low temperatures. Ohta [13] showed that the insulin-signaling pathway in the intestines and neurons is essential for temperature experience-dependent cold tolerance in animals. Animal survival in freezing conditions is a more complicated phenomenon than in cold temperatures without ice formation. Organs can be injured during freezing by physical factors, such as ice-crystal formation, dehydration, and cold [14]. Moreover, animals can also suffer biochemical damage, such as oxidative stress or hypoxia stress. [15]

To date, few studies have addressed the genes that regulate freezing tolerance or survival. We investigated the roles of daf-2(rf) in freeze–thaw stress-response regulation. We exposed daf-2(e1370) and other rf strains to freeze–thaw stress to identify daf-2(rf)-improved freeze-induced mortality and cell damage. Then, we tested other daf-2(rf) alleles with various phenotypic severities and performed molecular analysis to determine which signaling is involved. We performed the experiments through both genetic and morphological analyses with different C. elegans mutants.

**Results**

Insulin/IGF-1 receptor homologue daf-2(rf) regulates freeze–thaw stress survival

To investigate the role of daf-2(rf) in freezing tolerance, we evaluated the survival rate of daf-2(e1370) and wild-type (N2) strains exposed to freeze–thaw stress. Two-day-old adults were exposed to -80 °C for 8 min and then thawed in a water bath at 30 °C; results showed that the daf-2(e1370) strain had a significantly increased survival rate compared with the N2 strain (p < 0.01; Fig. 1a). To confirm the results of enhanced freezing survival produced by the reduction of daf-2/insulin-like signaling, we evaluated the survival rate of N2 animals with daf-2 RNAi interference or IGF,R inhibitor treatment. Animals with both daf-2 RNAi inactivation and IGF,R inhibitor treatment had enhanced survival rates after freeze–thaw treatment (p < 0.01; Fig. 1a).

At high freezing rates, intracellular freezing occurs, which can lead to cell damage, mainly by ice-crystal formation [15, 16]. To reduce ice-crystal formation and cell damage, cells and organisms can be cryopreserved by slowly lowering the temperature until deep freezing temperature. daf-2(e1370) and N2 animals were gradually cooled at a rate of about −1 °C/min (Additional file 1: Figure S1). Moreover, we also found that daf-2(e1370) animals had higher survival than N2 at any growth stage. This result indicates that survival of daf-2(rf) animals following freeze–thaw damage is not stage-specific (Fig. 1, Additional file 1: Figure S1A–C). We also found that the survival rate of L3/L4-stage worms was dramatically decreased compared with L1/L2-stage worms (Fig. 1b, Additional file 1: Figure S1A–B). This result indicates that freezing tolerance changed with development and age.

**Freezing-induced behavioral defects and cell defects blocked by daf-2(rf)**

Maintaining physiological functions in freezing temperatures is challenging; therefore, evaluation of response to different freezing conditions is important. Under freeze–thaw conditions, animals frozen at 0 °C and thawed in 30 °C water baths did not differ in survival rate (Additional file 2: Figure S2), which indicates that the animals’ physiological function loss is mainly caused by freezing rather than thawing. daf-2(e1370) animals survived and fully recovered locomotion after freeze–thaw stress. N2 animals displayed significant locomotion defects after recovery from some stress (Fig. 2).

To investigate cell defects from freezing and protection by daf-2(rf), we examined the muscle cell morphology of animals exposed to freeze–thaw stress. Freezing caused striking nuclear fragmentation in muscle myocytes. RNAi inactivation of daf-2 and downstream genes age, pdk-1 maintained intact nuclei and protected myocytes from both nuclear fragmentation and death [Fig. 3].

**daf-2 allelic specification for freeze–thaw stress survival is not a consequence of lifespan, dauer formation, or other stress resistance mechanisms**

daf-2(rf) alleles tend to promote a prolonged lifespan and form dauer larvae. Consequently, daf-2(rf) mutants are always resistant to harsh environments [17]. To analyze whether freeze–thaw survival is correlated with lifespan, we tested freezing survival rates associated with 11 daf-2 alleles’ (Fig. 4). The freeze–thaw stress survival phenotypes were not well correlated with lifespan (r = 0.538; p = 0.088).

We found that daf-2(e1370) worms, which did not have the longest lifespan, had the highest freeze–thaw stress survival, followed by e1391, e1371, e979, and m579. Five alleles that were weak or produced no increased freezing survival had significantly increased lifespans as long as or longer than e1370 [17]. Similarly,
one allele that did not increase survival (m41) and two weaker alleles (e1391 and e979) produced stronger Daf-c phenotypes than e1370, and one allele that did not increase survival (e1368) produced the same Daf-c phenotype as e1370 [18, 19]. With regard to stress resistance, four daf-2 alleles (e1391, e979, e579, and m596) were significantly resistant to thermal stress [18] but had weaker or no increased freeze–thaw survival. daf-2 allele e579 was significantly resistant to hypoxia stress but had weaker freeze–thaw survival. Three daf-2 alleles (e1371, e1391, and e979) were weak or non-resistant to hypoxia stress but had higher freeze–thaw survival. daf-2(e1370) freeze–thaw stress survival was highly allele-specific and did not appear to be a consequence of mechanisms that regulate lifespan, dauer formation, or other stress-resistance traits (Additional file 3: Figure S3).

The insulin-signaling pathway, but not dauer signaling, longevity genes, or trp-1, is essential for freeze–thaw stress survival

To determine the molecules downstream of the insulin receptor involved freeze–thaw stress survival, we tested various mutants defective in the known insulin-signaling pathway (Fig. 5). Phenotypic analysis showed that mutants defective in the daf-2/insulin receptor or its downstream molecules had abnormal enhancement or reduction of freeze–thaw stress survival (Fig. 5). Abnormal increments of freeze–thaw stress survival in daf-2 and downstream molecules mutants were suppressed by mutation or RNAi in the daf-16/FOXO-type transcriptional factor (Fig. 6).

Because the insulin-signaling pathway and other signaling molecules are essential for dauer larva formation, we tested other molecular components, including TGF-β and steroid hormonal signaling (Fig. 5) [19, 20]. However, we did not observe a considerable increase in these mutants (Fig. 5). The results indicate that the daf-2/insulin-signaling pathway, but not other dauer larva formation signaling, is essential for freezing survival.
daf-2 is the best-characterized longevity gene in *C. elegans*. We tested other longevity genes mutants, including *eat-2*, *clk-1*, and *isp-1* mutants, and found that these animals did not have enhanced freezing survival compared with the N2 strain. Recently, it was reported that the cold receptor transient receptor potential (TRP) channel, which is encoded by *trpa-1*, plays a central role in ageing and stress response to cold temperatures [21]. However, no reduced freeze–thaw survival was observed in *trpa-1* loss-of-function mutants. These results indicate that longevity genes and *trpa-1* are not essential for freeze–thaw stress survival.

**daf-16 nuclear translocation is not responsible for daf-2(rf)-enhanced freeze–thaw survival**

To elucidate how *daf-16* affects *daf-2(rf)* enhanced freeze–thaw stress survival, cellular distributions of a *daf-16*:GFP fusion protein were first categorized on a scale of 1 (unlocalized) to 3 (fully nuclear localized) (Fig. 7). Distributions were scored in *daf-2* RNAi animals and compared with controls maintained at 20 °C. As a positive control, *daf-2* inactivation resulted in a marked translocation of *daf-16* to the nucleus (Fig. 7a–d, left histogram). With freeze–thaw stress, *daf-16* nuclear translocation significantly increased in both *daf-2* RNAi and control animals (Fig. 7e–h). However, the difference
in daf-16 nuclear translocation between daf-2(rf) and control animals disappeared, because there was no significant discrepancy in the subcellular localization of daf-16::GFP (Fig. 7, right histogram). These observations indicate that altering the subcellular localization of daf-16 does not explain why daf-2(rf) has higher freeze–thaw stress survival than wild-type animals. We found that other mechanisms responsible for daf-2(rf) increased freezing survival.

To gather additional evidence to elucidate the nuclear activity of daf-16 responsible for daf-2(rf) increased freezing survival, we assayed daf-16 target genes. If the insulin pathway promotes daf-16 nuclear activity, this pathway should regulate the expression level of daf-16 target genes. We first examined the expression level of 21 direct transcriptional target genes of daf-16 that are also involved in ageing, larval arrest, or fat formation. We found that mRNA levels of C36A4.9 (acs-19, acetyl-CoA synthetase) and C46A10.7 (srh-99, class H chemoreceptor/olfactory receptor) were markedly upregulated in daf-2(e1370) worms (Additional file 4: Figure 4). Mice deficient in the homologous AceCS2 (acetyl-CoA synthetase 2) gene cannot maintain normal body temperatures when starved or fed a LC/HF diet [22]. In Drosophila, central and peripheral elements of the olfactory receptor system are responsible for temperature adaptation [23].

We infer that acs-19 or srh-99 may be required for freeze–thaw adaptation in C. elegans. For further elucidation, we assayed freeze–thaw survival rates with acs-19 and srh-99 RNAi in daf-2(e1370) and wild-type animals. We found that acs-19 and srh-99 RNAi significantly reduced daf-2(e1370) freeze–thaw survival, although there was still higher survival than in the daf-16;daf-2 double mutant (Fig. 8). This result indicates that the daf-16 target genes acs-19 and srh-99 are involved in the observed daf-2(rf) increased freeze–thaw stress survival. While, there is no evidence indicating other daf-16 target genes sup-37 or lig-1 involved in daf-2(rf) increased freezing survival (Fig. 8).

Discussion
In this study, we revealed genetic regulation of freeze–thaw stress responses in C. elegans. Phenotypic analysis of genetic deletion strains revealed that insulin/IGF-1 receptor daf-2 controls both survival and behavior during freeze–thaw stress. daf-2 reduction improved freeze–thaw stress survival, locomotion, and muscle cell protection. daf-2(rf) freeze–thaw response is highly allele-specific and not a consequence of lifespan, dauer formation, or other
stress-resistance trait regulation. We also revealed that insulin signaling, but not TGF-β, are related to freeze–thaw survival. Steroid hormone signaling participated in \( \text{daf-2(rf)} \) enhanced freeze–thaw stress regulation. \( \text{daf-16:GFP} \) cellular localization analysis and \( \text{daf-16} \) target gene screening revealed that \( \text{daf-16} \) regulated the target genes \( \text{acs-19} \) and \( \text{srh-99} \) but not \( \text{daf-16} \) nuclear translocation or \( \text{daf-2(rf)} \) enhanced freeze–thaw stress survival.

In the past few decades, there has been rapid progress in our understanding of how physiological mechanisms can protect freeze injury [24]. By contrast, very little is known about how genetic regulation promotes freeze resistance. At least in \( \text{C. elegans} \), freeze resistance is not purely a passive thermodynamic process. In this study, we characterized insulin/IGF-1 receptor \( \text{daf-2} \), which regulates freeze–thaw stress survival improvement in \( \text{C. elegans} \). Our results indicate that genetic programming actively contribute to enhanced survival and physiological function recovery from freezing conditions.

Previous work revealed that insulin signaling is required for lifespan regulation, dauer formation, and stress tolerance. Recent studies demonstrated that the insulin-signaling pathway or \( \text{daf-16} \) is required for temperature experience-dependent cold tolerance of animals [12, 13]. To ascertain whether insulin signaling plays roles in regulating freeze resistance in very stressful conditions, we performed survival experiments with freeze–thaw stress. In these experiments, we confirmed that insulin signaling via the \( \text{daf-16/FOXO} \) pathway is essential for freeze–thaw survival. Reduction-of-function of insulin/IGF-R \( \text{daf-2} \) protected muscle cell damage and promoted physiological activity recovery from freeze–thaw stress.

We found that freeze–thaw survival was \( \text{daf-2(rf)} \) allele-specific and is not a consequence of ageing, larval arrest, or other stress-resistance traits. Insulin signaling is also involved in dauer formation. To determine whether other dauer signaling participated in the freeze–thaw stress resistance, we assayed freeze–thaw survival of mutants. We found that dauer formation pathways, including TGF-β and steroid receptor signaling, were not essential for freeze–thaw stress survival. In addition, \( \text{daf-2} \) is the best-characterized longevity-regulated gene. Therefore, we also tested other freeze survival longevity gene mutants, including \( \text{eat-2} \) [25], \( \text{clk-1} \) [26], and \( \text{isp-1(rf)} \) [27]. We found that these other longevity genes were not required for freeze-stress resistance. Low temperatures led to increased longevity, in which the cold receptor TRP channel encoded by \( \text{trpa-1} \) is essential [21, 28]. Therefore, we tested the function of \( \text{trpa-1} \) and found that loss-of-function mutant \( \text{trpa-1} \) produced a normal phenotype compared with the wild-type strain. These results indicate that freeze–thaw survival is independent on cold receptor TRPA-1 signaling. These results are consistent with the \( \text{daf-2(rf)} \) allele relationship analysis between freeze–thaw survival and lifespan, larval arrest, and other stress-resistance traits. Freezing resistance may have an independent biological mechanism that differs from mechanisms that control ageing, dauer formation, and response to other stresses.

Finally, our results indicate that the \( \text{daf-16} \) target genes \( \text{C36A4.9 (acs-19)} \) and \( \text{C46A10.7 (srh-99)} \) are required for freezing survival of \( \text{daf-2(e1370)} \). Previous studies demonstrated that \( \text{acs-19} \) maintained core body temperatures of mice in fasting conditions [22], and olfactory receptor be acclimated to the environmental temperature of \( \text{D. melanogaster} \) [23]. Inactivation by RNAi of \( \text{acs-19} \) or \( \text{srh-99} \) contributed to both decreased fat storage and enhanced dauer formation of \( \text{C. elegans} \) [29]. It is possible that an unrecognized underlying mechanism contributes to both fat storage and dauer formation phenomena, which are related to freezing tolerance and survival. Nevertheless, our analyses did not exclude the contribution of additional mechanisms to freezing survival. \( \text{acs-19} \) and \( \text{srh-99} \) RNAi reduced freezing survival of \( \text{daf-2(e1370)} \); however, \( \text{daf-2(e1370) acs-19 RNAi and daf-2(e1370) srh-99 RNAi mutants exhibited significantly higher survival compared with the daf-16 mutant, which indicates that additional daf-16 target genes or mechanisms must be involved. For example,
genes involved in the synthesis of trehalose [30–34], glycerol [30], heat-shock proteins [32, 35, 36], antioxidant enzymes [37], and Δ9 desaturase enzymes [12] could participate in freeze–thaw stress survival by enhancing cold tolerance of *C. elegans* and other species. Future studies are needed to identify other unknown genetic programming and additional components of the *daf-2*(rf)/ *daf-16* pathway that ultimately may lead to a thorough understanding of how *daf-2*(rf) promotes freezing survival in *C. elegans*.

**Conclusion**

In conclusion, our mutant survival assay revealed that insulin/IGF-1 receptor *daf-2* played important roles in freeze–thaw stress responses in *C. elegans*. Freezing resistance is *daf-2* allele-specific and not a consequence of ageing, dauer formation, and other stress-regulation traits. Reduction-of-function of *daf-2* enhanced freeze–thaw survival, because it is dependent on insulin signaling pathway. The *daf-16*/FOXO-regulating target genes *acs-19* are *srh-99* essential for *daf-2*(rf) enhanced freezing resistance. Considering that the insulin/IGF-1 receptor showed striking conservation across phylogeny [38, 39], our work indicates that a similar phenomenon may also occur in other organisms.

**Methods**

**Strain selection and maintenance**

*Caenorhabditis elegans* were maintained on nematode growth medium (NGM) agar plates seeded with OP50, which is a slow-growing *Escherichia coli* mutant. The strains used included *N2*, *eat-2(ad465)*, *isp-1(qm150)*, *clk-1(qm30)*, *daf-1(m40)*, *daf-7(e1372)*, *daf-12(rh286)*, *daf-2(e1370)*, *daf-2(e979)*, *daf-2(e1391)*, *daf-2(e1368)*, *daf-2(e1371)*, *daf-2(m596)*, *daf-2(m577)*, *daf-2(m579)*, *daf-2(Sa193)*, *daf-2(m41)*, [daf-16(mgDf47); daf-2(e1370)], *TJ356(daf-16::GFP)*, and *PD4251(pmyo-3::GFP)*. All strains were provided by the Caenorhabditis Genetics Center funded by the National Institutes of Health National Center for Research Resources. Unless otherwise stated, all strains were cultured at 15 °C to the L4 stage and then transferred to animals at 20 °C for 2 d.
Freeze–thaw stress conditions

Adult animals (2 d after L4) on NGM plates were washed with M9 buffer and transferred into FACS tube with 1 ml M9 buffer. *Caenorhabditis elegans* does not survive freezing very well, so to determine optimal freezing length for analyses, we placed animals from room temperature into −80 °C for different amounts of times (4 min, 5 min, 6 min, 8 min, 12 min, or 16 min). At 4 min, there was no survival difference between N2 and daf-2(e1370). Ice began to form at 5 min, and all water froze in the tube at 6–7 min; however, all animals, including N2 and daf-2(e1370), died at 16 min. At 8 min, there were significant freezing survival differences among mutants. Therefore, animals were treated with freezing stress at −80 °C for 8 min for subsequent analyses. This experiment revealed that freezing stress damaged animals; it is possible that in the body, some cells did not completely freeze, but the freezing stress still injured animals.

Animals were exposed to freezing stress and then thawed at different temperatures (0 °C and 30 °C) for different lengths of time (10 min for 0 °C and approximately 1 min for 30 °C). We removed the tubes at the end of thawing process (when the ice was completely thawed), when the temperature in the tube is still 0 °C. All treated animals were placed on dry NGM plates for 6 h before determining mortality score with an optical microscope. L1–adult animals were used for the daf-2(e1370) survival test for different stages.

**Programmed freezing conditions**

L1–adult animals on NGM plates were gently washed with M9 buffer. For the freezing procedure, a Cryo 1 °C freezing container was used with a gradient cooling rate of −1 °C/min. Animals were transferred into freezing tubes with 0.9 ml buffer and 0.9 ml 30 % glycerin. At different time points (every 20 or 40 min), animals were thawed in a 30 °C water bath, and the survival rate was then assayed.

**Feeding RNAi**

dsRNA-expressing *E. coli* were streaked onto LB agar plates that contained ampicillin (50 μg ml⁻¹) and tetracycline (12.5 μg ml⁻¹), and then incubated at 37 °C overnight. Bacteria were inoculated in 3 ml LB liquid medium that contained only ampicillin (100 μg ml⁻¹) and then incubated at 37 °C overnight. All 3 ml of culture was spun down, the supernatant poured off until 150 μl was left (20× concentrated culture), and pellets were resuspended. Then, 50 μl of cells were resuspended to the center of RNAi plate (NGM/IPTG/ampicillin), allowed to dry (wrapped in aluminum foil), and induced overnight at room temperature (RNAi-seeded plates can be stored at room temperature for 2–3 d before use). Synchronized L1 worms were placed on each plate and incubated at 15 or 20 °C until they reached the desired stage for further experiments.

**Scoring mortality**

Worms were scored on NGM plates after 6 h of recovery from the thawing procedure. Worms were prodded with a pick at least three times over approximately 10 s; any that failed to move were counted as dead and removed from the plate.

**Morphologic cell defects**

The *PD4251* reporter gene *pmyo-3::gfp*, which is located in body wall muscle nuclei, was assayed with a fluorescence microscope. *pmyo-3::gfp* expression of strain *PD4251* by RNAi treatment with freezing and thawing was assayed.

**Quantitation of locomotion rate by counting body bends**

Ten animals were selected from fresh plates with fairly thin lawns, and one worm each was placed new plates. Assays started 24 h later (±1 h). A 3-min timer was used to count number of body bends. Every time the part of the worm just behind the pharynx reached a maximum bend in the opposite direction from the bend last counted was considered one body bend.
**daf-16 nucleus translocation**

*TJ356* nuclear protein *daf-16::GFP* expression was assayed with a fluorescence microscope. *daf-16::GFP* expression of [daf-2[RNAi]; *TJ356*] treated with freezing and thawing was assayed compared with that of *TJ356*.

**RNA isolation for qRT-PCR**

Total mRNA was extracted by TRIzol reagent (Invitrogen, Carlsbad, CA) from certain *C. elegans* and treated with RNase-free DNase (Promega, Madison, WI). Then, reverse transcription (RT) was performed with a TaKaRa RNA PCR kit (Takara, Dalian, China) following the manufacturer’s instructions. The primers used are described in Additional file 5: Table S1 and sequenced in the DNA Sequencing Department of Biosune Systems Biology (Shanghai, China).

**mRNA reverse transcription and qRT-PCR**

One nanogram in vitro transcribed RNA was added to the RNA sample (500–1000 ng). DEPC H2O was added to the RNA sample to 29.5 μl. Then, 0.5 μl each of 1 μg/μl random hexamer and 1 μg/μl poly dT were added. The mixture was incubated at 65 °C for 10 min, immediately put on ice for 5 min, and let stand at room temperature for 10 min. Then, 18.5 μl pre-mixture, which contained 2.5 μl 10 mM dNTP mix, 10 μl 5× first-strand buffer, 5 μl 0.1 M DTT, and 1 μl RNase OUT, was added. The mixture was then mixed and spun, and placed at room temperature for 2 min. Then, 1 μl Superscript II RT was added and gently mixed. The mixture was then spun down and let stand at room temperature for 10 min, incubated at 42 °C for 50 min, and heat inactivated at 70 °C for 15 min. Then, 1 μl RNase H was added to the solution, which was gently mixed, spun down, and blocked at 37 °C for 30 min.

SYBR Green qRT-PCR was performed on the LightCycler® 480 II System (Roche, Pleasanton, CA) using 5 μl 2× SYBR Green master mix (Roche), 2 μl RNase free water, 1 μl of forward, 1 μl of reverse primer (10 μM), and 1 μl cDNA per reaction. Primer efficiency was assessed by a dilution series, and a dissociation curve was used to assess primer specificity. qRT-PCR data analysis was performed using genEx software (MultiD).

**Statistical analyses**

Percent survival was reported as mean ± SEM per trial. Every test was repeated at least three times under the same conditions. Survival rate was analyzed by the non-parametric Mann–Whitney test. Correlation analysis between *daf-2(rf)* allele freeze–thaw survival and lifespan/other stress traits were conducted by the nonparametric Spearman correlation test.

**Additional files**

Additional file 1: Figure S1. *daf-2(e1370)* enhanced freezing survival under programmed freezing conditions. Reduction-of-function mutant *daf-2(e1370)* (squares) exhibited significantly increased survival compared with wild-type (N2) animals under the programmed cooling conditions (1 °C/min) at different stages. (JPG 259 kb)

Additional file 2: Figure S2. Survival rates of wild-type (N2) and *daf-2(e1370rf)* under different thawing processes. With different thawing treatments (30 °C water bath for 1 min or ice for 30 min), wild-type (N2) and *daf-2(e1370rf)* animals had unchanged survival rates after freeze–thaw stress. (JPG 94 kb)

Additional file 3: Figure S3. Correlated analysis of *daf-2(rf)* freeze–thaw survival and lifespan, larval arrest, and other stress-resistance traits. Freeze–thaw stress survival phenotypes were not correlated with lifespan, larval arrest, or hypoxia resistance, but was moderately correlated with heat shock survival. (JPG 41 kb)

Additional file 4: Figure S4. Screening for daf-16 target genes required for freezing survival by QF-PCR. daf-16 target genes C01B7.1 (ac-19), C29A12.3 (dgf-7) and C6A10.7 (str-99) have higher expression in *daf-2(e1370)* compared with wild-type (N2) animals. (JPG 92 kb)

Additional file 5: Table S1. *daf-2(rf)* allelic variation influence on freeze–thaw stress survival. (JPG 169 kb)

**Competing interests**

Authors have no financial and nonfinancial competing interests.

**Authors’ contributions**

Hu JP assayed the freezing survival and drafted the manuscript. Xu XY performed the target gene analysis and feeding RNAi. Huang LY measured the *daf-16::GFP* and muscle GFP analysis. Wang LS and Fang NY contributed to design the experiment. Wang LS analyzed the data and Fang NY revised the manuscript. All authors read and approved the final manuscript.

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