Cold shock induces a terminal investment reproductive response in C. elegans

Leah Gulyas1,2 & Jennifer R. Powell1*

Challenges from environmental stressors have a profound impact on many life-history traits of an organism, including reproductive strategy. Examples across multiple taxa have demonstrated that maternal reproductive investment resulting from stress can improve offspring survival; a form of matricidal provisioning when death appears imminent is known as terminal investment. Here we report a reproductive response in the nematode Caenorhabditis elegans upon exposure to acute cold shock at 2 °C, whereby vitellogenic lipid movement from the soma to the germline appears to be massively upregulated at the expense of parental survival. This response is dependent on functional TAX-2; TAX-4 cGMP-gated channels that are part of canonical thermosensory mechanisms in worms and can be prevented in the presence of activated SKN-1/Nrf2, the master stress regulator. Increased maternal provisioning promotes improved embryonic cold shock survival, which is notably suppressed in animals with impaired vitellogenesis. These findings suggest that cold shock in C. elegans triggers terminal investment to promote progeny fitness at the expense of parental survival and may serve as a tractable model for future studies of stress-induced progeny plasticity.

Environmental stressors can severely jeopardize the ability of an organism and its progeny to survive and reproduce, endangering organismal fitness. Consequently, organisms must launch effective stress response mechanisms to mitigate damage, promote recovery, and/or ensure progeny survival. A traditional view of life-history trade-offs assumes that challenge from a stressor should invoke a response in which resources are shuttled away from reproduction to support organismal defense and repair1. While this approach temporarily decreases reproductive fitness, improved long-term survival may allow for greater reproduction post-recovery. However, in a stressful environment where survival is perceived to be unlikely, an alternate strategy may instead be to funnel resources into reproduction, ensuring maximal output prior to parental death and subsequent survival of the F1 generation1.

The benefit of this strategy has been demonstrated in a number of organisms in different environmental conditions2. Both biotic (i.e. pathogen exposure) and abiotic (thermal, osmotic) stress conditions may potentiate molecular changes to reproduction. Alterations ranging from epigenetic modifications to simple maternal cytosolic investments of macromolecules and nucleic acids have been associated with increased offspring resilience to ensuing threats2,3. Yet despite myriad examples, the precise mechanisms by which stressors induce these responses are still widely under exploration.

The nematode Caenorhabditis elegans recently arose as an ideal model for investigating stress-induced parental progeny investment. Several studies have detailed molecular tactics by which worms modify and provision gametes and embryos in response to dietary restriction, infection by pathogenic pseudomonads and microsporidia, and osmotic stress4–7. Among an array of environmentally-relevant stressors, though, the reproductive impacts of thermal stress (specifically cold stress) on C. elegans have not been closely assessed. Both flies and mice employ reproductive strategies to respond to cold, with flies inducing a more general stress response of reproductive dormancy and mice depositing epigenetic modifications in sperm to promote retention of brown adipose tissue favoring increased metabolic rates in offspring8,9. Given the likelihood of C. elegans regularly encountering diurnal and seasonal temperature fluctuations, it seems probable that worms may also have evolved rigorous reproductive programs to deal with cold exposure.

Several aspects of cold response in C. elegans have been well-characterized in the literature, including thermotaxis and habituation10–12. Neurons and other tissues rely on a multitude of signaling mechanisms for these physiological responses to occur, which are in turn supported by a bevy of channels and other proteins. These processes are described in detail in a recent review from Takeishi et al.12. Only lately, though, have links between...
cold and reproduction began to emerge. Sonoda et al.\textsuperscript{13} demonstrate that the presence and integrity of sperm is required for normal cold tolerance by exerting an effect through multiple tissues, including AS\textsuperscript{1} neuronal activity. At the level of population dynamics, it has also been reported that \textit{C. elegans} warming from a 2-h cold exposure enters a state of programmed death which is thought to result in kin selection for the survival of younger (and presumably more productively fit) animals\textsuperscript{14}. However, the extent to which acute cold stress directly impacts reproductive capacity in these animals remains unknown. As alternating periods of cold and moderate temperatures preceding a winter season may comprise a substantial portion of the short reproductive window in \textit{C. elegans}, this necessitates further study.

We have previously shown that acute cold shock at 2 °C causes loss of intestinal pigmentation, immobility, and reproductive disruption in wild-type hermaphrodite worms. In many cases, this phenotypic program results in lethality\textsuperscript{15}. Here we extend this work by characterizing the induction of these phenotypes via the neuronal TAX-2/TAX-4 thermostransduction channel, mediated by the canonical stress response regulator SKN-1/Nrf2. We further describe a role for the normal vitellogenesis machinery in realocating pigmentated intestinal lipid supplies to the germline, which appears to come at the expense of parental survival following cold exposure. Overall, our findings suggest that acute cold shock provokes a terminal investment reproductive response in \textit{C. elegans}, making this discovery an exciting opportunity to explore the phenomenon in a genetically tractable system.

**Results**

**Acute cold shock causes drastic phenotypic alterations.** The duration of cold exposure for young adult hermaphrodite \textit{C. elegans} at 2 °C is negatively correlated to post-shock survival rates\textsuperscript{15}. Wild-type hermaphrodite worms exposed to a 4-h cold shock (CS) do not initially display high mortality rates (Fig. 1a); this allows observation of a range of phenotypic transitions as they recover from the limited-duration cold stress at their preferred temperature of 20 °C. One of the most striking phenotypes exhibited in post-cold shock (post-CS) animals during the recovery period is a dramatic decrease in pigmentation in the normally highly pigmented intestine, so that the body becomes almost entirely clear (Fig. 1b, c)\textsuperscript{15}. This is often accompanied by motor and reproductive disruptions such as mobility loss, withering of the gonad arms, decreased number of internal embryos, and the eventual death of about 30% of the population (Fig. 1a–d)\textsuperscript{15}. It should be noted that these phenotypic responses do not appear to be due to any relative heat shock following the transition from 2 to 20 °C as the expression of GFP-tagged HSP-4 (heat shock protein) is not induced following cold shock (Fig. 1e).

Neither is the reduced pigmentation following cold shock due to a period of starvation presumably experienced by the worms while they are at 2 °C. At this extreme cold temperature, the worms enter a “chill coma” in which pharyngeal pumping and virtually all other movement ceases\textsuperscript{15,16}; however, a total absence of food for a similar time period does not induce a comparable clearing phenotype (Supplemental Fig. S1). Interestingly, some CS wild-type animals regain pigmentation after clearing; these worms do not die and display a general reversal of the other negative impacts of cold shock (Fig. 1b)\textsuperscript{15}. We sought to better understand the factors regulating the post-CS recovery program in wild-type worms, focusing particularly on the functional role of pigmentation loss and the genetic components involved in producing it.

**Cold shock induces lipid reallocation from somatic tissues to the germline.** Since pigmentation in the \textit{C. elegans} intestine is due in part to the presence of lipid-storing fat droplets\textsuperscript{17}, we wondered whether the decrease in pigmentation in cold-shocked worms corresponds to a depletion of intestinal lipid supplies, potentially as a result of increased metabolism meant to fuel post-cold shock recovery. Using Nile Red, which accumulates and fluoresces in hydrophobic environments\textsuperscript{18}, as an indicator of lipid content, we therefore analyzed the fat stores of worms 12 following CS (note that all cold shocks were performed for 4 h at 2 °C). We indeed observed a significant decrease ($P < 0.0001$) in the average lipid content per worm (Fig. 2a); visually, this presents as a significant reduction in average fluorescence that is most striking in the intestine (Fig. 2c). However, we unexpectedly noted that the lipid content of the gonad appears to concomitantly increase, marked by the presence of fluorescent (and therefore lipid-rich) internal embryos in the gonads of many cold-shocked worms that were mostly absent in their non-cold shocked counterparts (Fig. 2b, c). These embryos are somewhat sporadic, usually accounting only for a proportion of all embryos in the germline and are interspersed with non-fluorescent embryos. Importantly, the percentage of fluorescent internal embryos per worm was found to be elevated two-fold in CS worms relative to that of non-CS controls (Fig. 2b). These observations suggest that rather than just metabolically depleting lipid supplies, \textit{C. elegans} may also reallocate lipids from the intestine to the germline. To further analyze this process spatiotemporally, we performed a time course of Nile Red staining following cold shock (Fig. 2c). After exit from cold shock and the corresponding chill coma\textsuperscript{15,16}, most worms gradually resume movement over the first 30 min of recovery at 20 °C. At this point, the distribution of Nile Red-staining lipids is indistinguishable from non-cold shocked controls, indicating that lipid reallocation occurs after, rather than during, the cold shock. By 4 h post-cold shock, we observed visually decreased but rarely absent fluorescence in the intestine; at this time, many worms have brightly fluorescing proximal oocytes, suggesting that lipids are beginning to relocate from the intestine into the germline. This process of reallocation appears to continue over the next 14 h, with many 12 h-recovered worms containing fluorescent embryos and oocytes but only slight intestinal fluorescence, and most 18 h-recovered worms fluorescing almost exclusively in the embryos. Though it is unclear what other factors (e.g. metabolism) may contribute to the overall pigmentation loss, these observations suggest that the dramatic decrease in intestinal pigmentation results in large part from the reallocation of lipids from somatic tissues to the germline during the recovery phase following a severe cold shock.
Thermosensation is required to initiate the cold shock response. We next asked how wild-type *C. elegans* perceive and interpret cold shock as a signal for lipid reallocation. The cGMP-gated channel subunits TAX-2 and TAX-4 are important for thermosensation and acquired cold tolerance in worms, acting specifically in the ASJ neurons\(^{11,19,20}\). We reasoned that worms might also rely on a neuronal mechanism of sensation that acts through the TAX-2/TAX-4 channel to induce the cold shock response in the absence of prior cold exposure. *tax-2(p671); tax-4(p678)* double loss-of-function mutants were therefore tested for sensitivity to cold shock. Not only do these mutants show 100% survival following cold shock (Fig. 3a), but, unlike wild-type worms, they also do not display any significant difference (\(P > 0.9999\)) in overall lipid content 12 h after CS, nor the striking lipid reallocation phenotype characterized by large numbers of fluorescent embryos (Fig. 3b–d). Interestingly, despite this resilience to shock, *tax-2(p671); tax-4(p678)* mutants still show decreased internal embryo quantities, though not to the same degree as N2 worms (Fig. 3e), suggesting that cold exposure has some additional
impact on fertility independent of this thermosensation pathway. Taken together, though, these data support the hypothesis that fat reallocation as a stress response following CS may depend on a neuronal mechanism for induction. Since TAX-2/TAX-4 plays a role in temperature sensation, we speculate that canonical cold thermosensation is required for lipid reallocation to take place.

Figure 2. Cold shock recovery is associated with lipid localization shifts from the intestine to the germline. Young adult N2 hermaphrodites were exposed to 2 °C cold shock or control nonCS conditions (20 °C) for 4 h and recovered before Nile Red staining and imaging for lipid content. (a) Average fluorescence per worm (n ≥ 29 per condition; Mann–Whitney U test: U = 48; two-tailed $P < 0.0001$) and (b) percent of internal fluorescent embryos per worm (n ≥ 111; Mann–Whitney U test: U = 1885; two-tailed $P < 0.0001$) were quantified 12 h post-CS. Error bars are mean ± s.e.m. (c) Representative images of Nile Red-stained worms at indicated time points post-CS show progression of intestinal lipid loss and relocalization to the embryos.
Figure 3. TAX-2; TAX-4-mediated thermosensation is required for lipid relocalization following cold shock. (a) *tax-2(p671); tax-4(p678)* double loss-of-function mutants were cold shocked and recovered while monitoring survival rates (n ≥ 120 worms per condition). (b) At 12 h post-CS or nonCS control, *tax-2(p671); tax-4(p678)* were Nile Red lipid-stained and the average fluorescence per worm was quantified (Kruskal–Wallis test: $H = 83.83$, $P < 0.0001$; Dunn’s Multiple Comparison test: ****$P < 0.0001$). (c) Representative images of Nile Red staining show retention of fluorescence in the intestines of *tax-2; tax-4* mutants. (d) Number of embryos (Kruskal–Wallis test: $H = 171.1$, $P < 0.0001$; Dunn’s Multiple Comparison test: *$P = 0.0484$, ****$P < 0.0001$) and (e) percent fluorescent internal embryos (Kruskal–Wallis test: $H = 100.1$, $P < 0.0001$; Dunn’s Multiple Comparison test: ****$P < 0.0001$) were quantified per worm from Nile Red images (n ≥ 24 worms per condition for b-e; error bars are mean ± s.e.m.).
**SKN-1 promotes cold stress resistance.** After determining a requirement for neuronal signaling in cold-shock-induced lipid reallocation, we next wondered what intermediate factors were needed to translate the cold stimulus into a signal to mobilize resources. The master regulatory transcription factor SKN-1/Nrf2 coordinates a return to homeostasis following a variety of stresses, including oxidative stress, pathogen infection, and osmotic stress. We therefore predicted that skn-1 would perform a similar function during cold stress recovery. Indeed, skn-1(lax188) gain-of-function mutants are highly resistant to cold shock, with nearly 100% survival rates 96 h post-CS (Fig. 4a). Consistent with a retention of overall lipid stores by Nile Red Staining (Fig. 4b.d), these mutants did not contain significantly greater numbers of fluorescing internal embryos following cold shock (P > 0.9999), though they did have a marginal reduction in the number of internal embryos (Fig. 4d–f). Conversely, skn-1(mg570) loss-of-function mutants displayed a wild-type cold shock response, with significant reallocation (P < 0.00001) of lipids to the germline and loss of somatic fats (Fig. 4c–f). Because mg570 eliminates only the skn-1a mRNA isofrom, but skn-1(lax188gf) activates both skn-1a and skn-1c, we additionally tested the CS response of skn-1(z15) loss-of-function mutants, which are reported to have a 76% reduction in both skn-1a and skn-1c mRNA levels. We observed a similar clear phenotype (Supplemental Fig. S2); however, these animals appeared slow-growing and less robust, which may have contributed to more substantial CS-induced lethality. Taken together, these data suggest that the skn-1a and perhaps also the skn-1c isofroms are not required for lipid mobilization in response to cold shock. However, SKN-1a/c activation protects cold-shocked worms from recovery phase lethality and prevents lipid reallocation.

**Lipid reallocation results from upregulated vitellogenesis following cold shock.** The process governing the normal movement of lipids from the soma to the germline is vitellogenesis, requiring the vitellogenin proteins coded for by vit-1–6 genes. Once coupled to somatic lipid supplies, the resulting lipoprotein complexes shuttle to the germline. We hypothesized that the lipid mobilization induced by cold shock is a result of the normal vitellogenesis machinery being commandeered as part of a stress response.

One of the major vitellogenins is VIT-2, which generates a 170 kD yolk protein product known as YP170B. We predicted that loss of vit-2 would reduce lipid reallocation following CS. Since preventing reallocation in our previous mutants also increased post-CS survival, we expected that vit-2 loss of function would enhance protection from CS. Consistent with these hypotheses, impairment of vitellogenesis in vit-2(ok3211) loss-of-function mutants produced phenotypes similar to tax-2(p671); tax-4(p678) animals. In addition to maintaining high survival rates, CS vit-2(ok3211) mutants did not contain a substantial loss of intestinal lipid supplies, nor did they exhibit a marked increase in internal embryo fluorescence (despite a modest decrease in internal embryo counts) (Fig. 5a–e).

Since other vitellogenin transcripts produce different lipoprotein forms for movement to the germline, we decided to additionally test whether a member of the YP170A-generating class of vitellogenins would recapitulate our results with vit-2. To this end, we analyzed the CS phenotypes of vit-5(ok3239) loss-of-function animals. Indeed, we found that as in vit-2 mutants, survival was rescued in these animals following CS, corresponding with a retention of somatic lipids and inhibition of embryonic lipid reallocation (Fig. 6a–c.e). As with other mutants that prevent lipid reallocation, there was still an effect of cold in reducing overall embryo output 12 h post-CS, hinting that temperature impacts fertility independent of the other reproductive alterations exhibited by wild-type worms (Fig. 6d).

**Cold shock response is a form of a terminal investment.** Our data thus far suggest that upon recovery from acute CS, wild-type *C. elegans* induce a reproductive phenotype whereby somatic lipid supplies are massively relocalized to the germline using the normal vitellogenesis machinery. This appears to come at the expense of the parent’s own mortality, as retaining somatic lipids by preventing thermosensation or vitellogenesis is sufficient to rescue survival. Such a trade-off between survival and reproduction is replete with the terminal investment hypothesis (reviewed in Gulyas and Powell, 2019), which predicts that in some instances of acute stress where the likelihood of death is high, organisms can preferentially funnel resources to reproduction to maximize reproductive fitness at the cost of survival.

To determine whether cold stress-associated phenotypes in *C. elegans* are an example of such a process, we eliminated all potential for reproductive investment by assaying sterile glp-1(e22141) and glp-1(q231) worms to ask whether these worms still exhibited lipid loss or death upon cold shock. Strikingly, the absence of a germline in these animals completely prevented both intestinal clearing and death, again confirming that lipid movement from the soma to the germline is associated with parental lethality (Fig. 7a). If resource reallocation to the progeny is indeed meant to increase reproductive fitness in inclement conditions, there should be some benefit to the progeny of CS worms that offspring of nonCS parents do not receive. We speculated that in the case of environmental CS, a sudden, seasonal, cold to warm cycle might signal likelihood of future cold conditions that would impede the ability of embryos to hatch and survive to reproductive age. We therefore devised an assay to test for the relative fitness of embryos in cold conditions depending on whether they came from nonCS or CS parents and were thus more likely to have received lipid provisioning. To do this, embryos were collected from nonCS and CS parents within a 2 h time window when post-CS lipid reallocation seems to peak and subsequently exposed them to a cold stress of 24 h. We then assessed hatching rates 24 h following the embryonic cold shock. Excitingly, embryos coming from parents that had experienced cold shock prior to reproduction exhibited a small but significant increase in hatching rates relative to their counterparts from nonCS hermaphrodite parents. Furthermore, preventing lipid reallocation by impairment of vitellogenesis in either vit-2(0) or vit-5(0) was able to substantially ablate this effect, suggesting that the improved survival is attributable specifically to lipid investment in the F1 generation (Fig. 7b, c). Altogether, it appears that while preventing embryonic lipid endowment may promote adult survival post-CS, offspring that go on to experience
Figure 4. Activated SKN-1 prevents lipid relocalization following cold shock (a) *skn-1(lax188)* gain-of-function mutants and *skn-1(mg570)* loss-of-function mutants were cold shocked and recovered while monitoring survival rates (*n* ≥ 120 worms per condition). (b–c) At 12 h post-CS or control nonCS, (b) *skn-1(lax188)* and (c) *skn-1(mg570)* were Nile Red lipid-stained and average fluorescence per worm quantified (*N2* v. *skn-1(lax188)* - Kruskal–Wallis test: *H* = 47.46, *P* < 0.0001; Dunn's Multiple Comparison test: **** *P* < 0.0001; *N2* v. *skn-1(mg570)* - Kruskal–Wallis test: *H* = 64.55, *P* < 0.0001; Dunn's Multiple Comparison test: ** *P* = 0.0064, **** *P* < 0.0001). (d) Relative to wild-type *N2s*, representative images of *skn-1(lax188)* mutants show maintenance of intestinal fluorescence contrary to *skn-1(mg570)*, which show lipid relocalization. (e) Number of embryos (Kruskal–Wallis test: *H* = 223.1, *P* < 0.0001; Dunn's Multiple Comparison test: ** *P* = 0.0020, ** *N2* CS vs. *skn-1(gf)* nonCS = 0.0014, **** *P* < 0.0001) and (f) percent of fluorescent internal embryos (Kruskal–Wallis test: *H* = 132.0, *P* < 0.0001; Dunn's Multiple Comparison test: *** *P* = 0.0002, **** *P* < 0.0001) were quantified per worm from Nile Red images (*n* ≥ 26 worms per condition for b–e; error bars are mean ± s.e.m.).
Figure 5. VIT-2-regulated vitellogenesis during recovery promotes lipid relocalization and impairs survival. (a) vit-2(ok3211) loss-of-function mutants were cold shocked and recovered while monitoring survival rates (n ≥ 130 worms per condition). (b) At 12 h post-CS or nonCS, vit-2(ok3211) were Nile Red lipid-stained and average fluorescence per worm quantified (n ≥ 31 worms per condition; Kruskal–Wallis test: H = 75.18, \( P < 0.0001 \); Dunn's Multiple Comparison test: ****\( P < 0.0001 \)). Error bars are mean ± s.e.m. (c) Images of Nile Red Staining show intestinal lipid retention in vit-2 mutants. (d) Number of embryos (Kruskal–Wallis test: H = 169.9, \( P < 0.0001 \); Dunn's Multiple Comparison test: *\( P = 0.0149 \), ****\( P < 0.0001 \)) and (e) percent of fluorescent internal embryos (Kruskal–Wallis test: H = 101.5, \( P < 0.0001 \); Dunn's Multiple Comparison test: ****\( P < 0.0001 \)) were quantified per worm from Nile Red images (n ≥ 31 for b–e; error bars are mean ± s.e.m.).
future inclement conditions suffer diminished survival rates, underscoring the evolutionary advantage of terminal investment as a reproductive strategy.

Figure 6. VIT-5 vitellogenin family also promotes embryo lipid reallocation during cold shock recovery. 
(a) vit-5(ok3239) loss-of-function mutants were cold shocked and recovered while monitoring survival rates (n ≥ 170 worms per condition). At 12 h post-CS or control nonCS, vit-5(ok3239) were Nile Red lipid-stained and average fluorescence per worm quantified (n ≥ 67 worms per condition; Kruskal–Wallis test: H = 172.0, P < 0.0001; Dunn’s Multiple Comparison test: ****P < 0.0001. Error bars are mean ± s.e.m. (c) Representative Nile Red staining shows intestinal fluorescence in vit-5(ok3239) following cold shock. (d) Number of embryos (Kruskal–Wallis test: H = 137.2, P < 0.0001; Dunn’s Multiple Comparison test: ****P < 0.0001) and (e) percent of fluorescent internal embryos (Kruskal–Wallis test: H = 37.58, P < 0.0001; Dunn’s Multiple Comparison test: H = 37.58, ****P < 0.0001) were quantified per worm from Nile Red images (n ≥ 67 worms per condition; error bars are mean ± s.e.m.).
Cold shock response overlaps with other stress response phenotypes. We have provided here the first evidence that *C. elegans*, upon acute cold shock (CS), is able to improve the success of progeny survival during future cold exposure by mobilizing lipids to the germline as reproductive provisioning. This process seems to depend on neuronal sensation of the cold temperature, which permits a shift of lipid localization from the soma to the germline. Lipid reallocation is mediated by vitellogenins, and despite an overall decrease in embryo output, the embryos in these worms are more resilient to future cold exposure and display more robust hatching. The switch to a sudden last reproductive event and resulting parental mortality is consistent with the adoption of a more semelparous lifestyle\(^29\). In this instance, potentiation by a severe environmental stress to induce a higher quality (rather than quantity) final reproductive investment suggests that *C. elegans* has evolved a terminal investment response to deal with acute cold exposure.

**Figure 7.** Cold shock response is a form of terminal investment. (a) *glp-1(e2141)* and *glp-1(q231)* were grown at the restrictive temperature (25 °C) to induce sterility and then habituated at 20 °C for 4 h. Worms were then cold shocked or mock shocked (nonCS) and recovered at 20 °C for 24 h and phenotypes were scored (n ≥ 197 worms for all conditions; Chi-Squared Test for Homogeneity, \(P < 0.0001\) for N2 CS vs. e2141 CS or q231 CS). (b) Young adult hermaphrodite N2 and *vit-2(ok3211)* (P0) were cold shocked or mock shocked (nonCS) for 4 h and recovered for 16 h. Embryos from P0 treatments were collected from 16 to 18 h and cold shocked for 24 h. Embryos were allowed to recover 24 h at 20 °C and then the number of hatched embryos was quantified. Data points represent rates as percent hatched for a plate containing 50–100 embryos (n ≥ 10 plates per condition; 2-way ANOVA (F(3,32) = 20.87, \(P < 0.0001\)) with Tukey multiple comparison test (****\(P < 0.0001\)). Bars are mean ± s.e.m. (c) Performed as in b with N2 and *vit-5* (ok3239) worms (n = 10 plates per condition; 2-way ANOVA (F(3,27) = 35.28, \(P < 0.0001\)) and Tukey multiple comparison test (****\(P < 0.0001\)).
Though, to our knowledge, no previous study has linked terminal investment to *C. elegans*, there are other documented phenomena characterized by a similar trade-off. For example, the “age-dependent somatic depletion of fat (ASDF)” phenotype is characterized by age-dependent changes in lipid homeostasis that are induced during both starvation and oxidative stress and are highly reminiscent of post-cold-shock reallocation of lipids from the intestine to the germline. As is also the case with cold shock-induced terminal investment, reduction of vitellogenic transcripts (including vit-2) suppresses lipid mobilization. Lipid reallocation during ASDF does appear to negatively impact parental survival, but it is unclear whether this phenotype also holds functional significance for offspring survival. Surprisingly, while the process governing the ASDF phenotype and cold shock recovery are both regulated by the master stress response transcription factor SKN-1/Nrf2, activating SKN-1 has opposite effects. When measuring ASDF, skn-1(gf) stimulates the shift of lipids from the intestine to the germline; in contrast, skn-1(gf) prevents lipid mobilization and promotes intestinal lipid sequestration following cold shock. It is possible that the role of SKN-1 varies with age since ASDF occurs in much older adults than those analyzed following cold stress. This potential will be analyzed in future work, in addition to elaborating the roles of specific skn-1 alleles during CS. The major outstanding question of how stress-altered metabolic state impacts lipid governance and the nature of the lipids involved also merits further exploration. Although regulation of lipid homeostasis may be via distinct pathways, the existence of mechanistically similar responses to starvation, oxidative stress, and cold stress suggests that terminal investment may be a more generalized response to severe stressors in *C. elegans*.

Despite the fact that terminal investment requires extremely high parental resource investment, there are other similar and less severe forms of maternal provisioning in *C. elegans*. For example, osmotic stress shifts embryonic contents to include less glycerol and more glycerol, a cryoprotectant. It is particularly relevant to our study that mild nutrient deprivation in hermaphrodites leads to intergenerational plasticity in which embryos tend to be slightly larger and offspring are somewhat protected from the effect of larval starvation. Remarkably, this corresponds to decreased maternal insulin signaling and upregulated vitellogenin provisioning in the germline, much as we report during cold exposure. It will be interesting in future experiments to dissect how thermosensory signaling may hyperactivate vitellogenesis upon acute cold stress, especially as it relates to different vitellogenin yolk protein products. Additionally, analyzing how various stressors impact and modify embryonic composition will be important for understanding the extent to which maternal investments are conserved. Taken together, it is clear that stressors in the parental generation can have remarkable consequences on reproductive and survival trade-offs as well as offspring physiology and plasticity.

**Terminal investment in response to cold may have evolved in regularly fluctuating environmental conditions.** Since the preeminent goal of survival is reproduction, terminal investment is sensible from a fitness standpoint. But why would cold shock elicit this phenotype? Experimental evolution in *C. elegans* reveals that deterministic maternal effects (those that attempt to successfully invest offspring for survival in a particular environment) evolve in response to predictably fluctuating environments. Thus, we conjecture that predictably encountered cold environments, such as those during seasonal freeze–thaw cycles in temperate regions, are likely to have given rise to the reallocation phenotype. This could happen by one of the two possible mechanisms. First, cold shock-induced damage may signal to the worm that death is imminent and reproductive timespan is limited. Worms may then funnel all their resources into embryos which are then better capable of surviving cold stress. However, this seems unlikely since these data suggest that *C. elegans* that avoid reallocation (or are genetically prevented from doing so) survive CS extremely well, which is inconsistent with CS causing large amounts of damage. It is possible that these worms are simply more resistant to CS-induced damage or are better able to repair damage in the absence of reallocation, though this would need to be tested more rigorously. A second, more plausible scenario is that seasonal cold cycles signal a coming winter season with a low chance of offspring survival. At specific larval stages, worms can enter a highly stress-resistant state (called the dauer stage) which may enable them to survive the winter season and resume development when conditions are more favorable. If the better-provisioned embryos of worms that have recently experienced a cold cycle survive successive cold shock, this could favor a larger percentage of offspring that are able to successfully enter the dauer state. Long-term, this would translate to a higher probability that offspring survive to reproduce when winter conditions are alleviated, continuing the parent’s lineage. This model accounts for the selection of the reallocation trait in evolutionary history. It further suggests that rather than being induced as a result of damage that signals impending death (and thus a final chance to reproduce), reallocation may instead be prompted by a more general prediction of future damage to both the parent and the offspring. Further experimentation is needed to assess this model and to determine how it would fit into the known framework for terminal investment strategies.

Although we have largely considered simple maternal effects involving cytosolic investments in this work, there is also the potential for epigenetic effects at the level of gene expression to play a role in intergenerational and transgenerational offspring survival. Indeed, the progeny of *C. elegans* submitted to various stressors demonstrate improvements in stress survival to the same and other types of stress. Recently, a study by Burton et al. examined multigenerational responses to a number of stressors including nutrient deprivation and pathogenic stress and found that multiple *Caenorhabditis* species regulate gene expression in the F1 generation in a stress-specific manner to benefit survival upon exposure to the same. These studies highlight the evolutionary advantage of provisioning offspring for what is perceived to be a likely future. It would be interesting to extend analyses of multigenerational gene regulation to see if this is also at play in the context of CS.

**Terminal investment has consequences for population-level dynamics.** Terminal investment occurs across a broad phylogenetic range in response to many stressors, and the potential ecological and evolutionary implications are serious in today’s global climate. Pathogen exposure commonly elicits terminal invest-
Cold shock survival. For adult cold shock, approximately 20–70 young adult worms (not yet gravid) were picked to OP50-seeded 3.5 cm NGM and placed at 2 °C for 4 h. After 4 h cold shock, plates were transferred to 20 °C for worm recovery. Worms were scored at 1, 4, 24, 48, 72, and 96 h post-cold shock for survival and qualitative phenotypic assessments. A worm was considered dead when nose tap did not elicit any movement. Clear worms were determined by a lack of almost all intestinal pigmentation, and immobile worms by a nose tap that elicited only movement in the head region.

For embryonic cold shock, parents were cold shocked as above, allowed to recover for 15 h, then transferred to fresh plates to lay embryos for 2 h. This time window is the peak of lipid reallocation following cold shock. Approximately 50–100 embryos were transferred to very lightly seeded 3.5 cm NGM plates and cold shocked at 2 °C for 24 h. Following a 24-h recovery at 20 °C, the number of hatched and unhatched embryos were counted. Embryos were considered hatched if the entirety of the L1 larva was visible in a non-curled state, and the larva was not dead.

For adult starvation, approximately 50–150 young adult worms were washed with M9 to remove excess OP50 and pipetted onto seeded or unseeded 3.5 cm NGM plates. Worms on unseeded plates were starved for 4 h at 20 °C, then recovered by the addition of concentrated OP50. Worms on seeded plates were cold shocked at 2 °C for 4 h, then recovered by transfer to 20 °C. Worms were scored for pigmentation and viability after 24 h of recovery.

During all cold shock experiments, plates were placed directly on the incubator shelf in a monolayer rather than a stack both during cold shock and the early stages of recovery to facilitate uniform temperature changes among plates.

C. elegans fluorescence imaging and quantification. zcIs4[hs:4::GFP] worms were either heat shocked at 35 °C for 2 h, cold shocked according to standard protocol, or non-shocked. After 12 h recovery at 20 °C, worms were picked to a drop of M9 containing 5 mM sodium azide. Worms were imaged on a Nikon Eclipse 90i microscope at exposure times of 2.9 ms for DIC and 900 ms for GFP.

The fixed Nile Red staining protocol was modified from Pino et al.35 and Escorcia et al. 201936. Briefly, at 12 h post-cold shock, 50–75 worms were washed once with M9 and fixed for three minutes at room temperature in 25 μl M9 + 150 μl 40% isopropanol. Worms were stained for 2 h in the dark with gentle rocking in 175 μl 40% isopropanol containing 0.6% Nile Red stock (0.5 mg/ml in acetone). Samples were washed with M9 for 30 min in the dark with gentle rocking, then mounted on a 2% agarose pad for imaging. Worms were imaged on a Nikon Eclipse 90i microscope at exposure times of 3 ms for DIC and 400 ms for GFP (Figs. 1, 3, 4, 5) or on a Zeiss Axio Imager Z1 microscope with exposure times of 3 ms for DIC and 500 ms for GFP (Figs. 2, 6).

Scale bars (100 μm) were added and all images were rotated and cropped using ImageJ. No other image manipulations were performed.

Images of fluorescent worms taken on the Nikon Eclipse 90i microscope (Figs. 1, 3, 4, 5) were analyzed using NIS Elements Data Analysis software; images taken on the Zeiss Axio Imager Z1 microscope (Figs. 2, 6) were analyzed using ImageJ. In either case, total fluorescence was determined by outlining the entire worm as a Region...
of Interest (ROI) and calculating the average fluorescence for that ROI. The number of internal embryos and the number of fluorescent embryos were determined by visual counts.

**Statistical analysis.** All data represent at least three independent replicates. Statistical analysis of continuous data was performed in GraphPad Prism 9.2.0 (www.graphpad.com) with an alpha value of P < 0.05. For samples with 2 conditions, nonparametric Mann–Whitney U tests were performed and reported with two-tailed P values. In cases with greater than 2 conditions, Kruskal–Wallis nonparametric ANOVA were performed with Dunn’s Multiple Comparison test. For hatching analyses, a Grubb’s test was conducted through GraphPad to identify outliers in the data, with one point identified as an outlier (Z = 3.4005, two-tailed P < 0.01) and removed. d’Agostino & Pearson normality tests were then applied to the remaining values and normal distribution was confirmed by a P > 0.05 for each condition. After validating data normality, 2-way ANOVAs were used to compare the mean of each condition with a Tukey’s Multiple Comparison test. Categorical data were analyzed using a Chi-Squared Test for Homogeneity in Microsoft Excel, with an alpha value of P < 0.05.

Received: 11 September 2021; Accepted: 8 December 2021
Published online: 25 January 2022

**References**

1. Duffield, K. R., Bowers, K. E., Sakaluk, S. K. & Sadd, B. M. A dynamic threshold model for terminal investment. *Behav. Ecol. Sociobiol.* **71**, (2017).
2. Gulyas, L. & Powell, J. R. Predicting the future: Parental progeny investment in response to environmental stress cues. *Front. Cell Dev. Biol.* **7** (2019).
3. Herman, J. J. & Sultan, S. E. Adaptive transgenerational plasticity in plants: Case studies, mechanisms, and implications for natural populations. *Front. Plant Sci.* **0**, 102 (2011).
4. Hirschman, J. D., Hung, A. & Baugh, L. . R. Maternal diet and insulin-like signaling control intergenerational plasticity of progeny size and starvation resistance. *PLoS Genet.* **12** (2016).
5. Frazier, H. N. & Roth, M. B. Adaptive sugar provisioning controls survival of *C. elegans* embryos in adverse environments. *Curr. Biol.* **19**, 859–863 (2009).
6. Burton, N. O. et al. Cysteine synthases CYSL-1 and CYSL-2 mediate *C. elegans* heritable adaptation to *V. parzenovis* infection. *Nat. Commun.* **11** (2020).
7. Burton, N. O. et al. Intergenerational adaptations are evolutionarily conserved, stress-specific, and have deleterious trade-offs. *bioRxiv* 2021.05.07.431118 (2021). https://doi.org/10.1101/2021.05.07.431118.
8. Lirakis, M., Dolezal, M. & Schütter, C. Redefining reproductive dormancy in *Drosophila* as a general stress response to cold temperatures. *J. Insect Physiol.* **107**, 175–185 (2018).
9. Sun, W. et al. Cold-induced epigenetic programming of the sperm enhances brown adipose tissue activity in the offspring. *Nat. Med.* **2018** 249, 1372–1383 (2018).
10. Garrity, P. A., Goodman, M. B., Samuel, A. D. & Sengupta, P. Running hot and cold: Behavioral strategies, neural circuits, and the molecular machinery for thermotaxis in *C. elegans* and *Drosophila*. *Genes Dev.* **24**, 2365–2382 (2010).
11. Ohta, A., Ujisawa, T., Sonoda, S. & Kuhara, A. Light and pheromone-sensing neurons regulates cold habituation through insulin signalling in *Caenorhabditis elegans*. *Nat. Commun.* **5** (2014).
12. Takeishi, A., Takagaki, N. & Kuhara, A. Temperature signaling underlying thermotaxis and cold tolerance in *Caenorhabditis elegans*. https://doi.org/10.18608/1677063.2020.1734001.
13. Sonoda, S., Ohta, A., Maruo, A., Ujisawa, T. & Kuhara, A. Sperm affects head sensory neuron in temperature tolerance of *Caenorhabditis elegans*. *Cell Rep.* **16**, 56–65 (2016).
14. Jiang, W. et al. A genetic program mediates cold-warming response and promotes stress-induced senescence in *C. elegans*. *Elife* **7** (2018).
15. Robinson, J. D. & Powell, J. R. Long-term recovery from acute cold shock in *Caenorhabditis elegans*. *BMC Cell Biol.* **17**, 1–11 (2016).
16. Collinet, H., Lee, S. F. & Hoffmann, A. Temporal expression of heat shock genes during cold stress and recovery from chill coma in adult *Drosophila melanogaster*. *FEBS J.* **277**, 174–185 (2010).
17. McGhee, J. D. The *C. elegans* intestine. *WormBook* 1–36 (2007). https://doi.org/10.1895/WORMBOOK.1.133.1.
18. Greenspan, P., Mayer, E. P. & Fowler, S. D. Nile red: A selective fluorescent stain for intracellular lipid droplets. *J. Cell Biol.* **100**, 965–973 (1985).
19. Coburn, C. M. & Bargmann, C. I. A putative cyclic nucleotide–Gated channel is required for sensory development and function in *C. elegans*. *Neuron* **17**, 695–706 (1996).
20. Komatsu, H., Mori, I., Rhee, J. S., Akaike, N. & Oshima, Y. Mutations in a cyclic nucleotide–gated channel lead to abnormal thermosensation and chemosensation in *C. elegans*. *Neuron* **17**, 707–718 (1996).
21. Hoeven, R. van der, McCallum, K. C., Cruz, M. R. & Garsin, D. A. Ce-Duox1/BLI-3 generated reactive oxygen species trigger protective SKN-1 activity via p38 MAPK signaling during infection in *C. elegans*. *PLoS Pathog.* **7**, e1002453 (2011).
22. Papp, D., Csermely, P. & Soti, C. A role for SKN-1/Nrf in pathogen resistance and immunosenescence in *Caenorhabditis elegans*. *PLoS Pathog.* **8**, e1002673 (2012).
23. Dodd, W. et al. A damage sensor associated with the cuticle coordinates three core environmental stress responses in *Caenorhabditis elegans*. *Genetics* **208**, 1467–1482 (2018).
24. Lynn, D. A. et al. Omegas-3 and -6 fatty acids allocate somatic and germline lipids to ensure fitness during nutrient and oxidative stress in *Caenorhabditis elegans*. *Proc. Natl. Acad. Sci. U. S. A.* **112**, 15378–15383 (2015).
25. Lehrbach, N. J. & Ruvkun, G. Proteasome dysfunction triggers activation of SKN-1/Nrf1 by the aspartic protease DDI-1. *Elife* **5** (2016).
26. Tong, L., Dodd, W. & Choe, K. Isolation of a hypomorphic skn-1 allele that does not require a balancer for maintenance. *G3 Genomes Genet.* **6**, 551–558 (2016).
27. Perez, M. F. & Lehner, B. Vitellogenins—Yolk gene function and regulation in *Caenorhabditis elegans*. *Front. Physiol.* **0**, 1067 (2019).
28. Spiehl, T. & Blumenthal, T. *The Caenorhabditis elegans vitellogenin gene family includes a gene encoding a distantly related protein*. *Mol. Cell. Biol.* **5**, 2495–2501 (1985).
29. Hughes, P. W. Between semelparity and iteroparity: Empirical evidence for a continuum of modes of parity. *Ecol. Evol.* **7**, 8232–8261 (2017).
30. Jordan, J. M. et al. Insulin/IGF signaling and vitellogenin provisioning mediate intergenerational adaptation to nutrient stress. *Curr. Biol.* 29, 2380-2388.e5 (2019).
31. Proulx, S. R. & Teotónio, H. What kind of maternal effects can be selected for in fluctuating environments? *Am. Nat.* 189, E118–E137 (2017).
32. Brannelly, L. A., Webb, R., Skerratt, L. F. & Berger, I. Amphibians with infectious disease increase their reproductive effort: Evidence for the terminal investment hypothesis. *Open Biol.* 6 (2016).
33. Hargrove, J. W., Muzari, M. O. & English, S. How maternal investment varies with environmental factors and the age and physiological state of wild tsetse Glossina pallidipes and Glossina morsitans morsitans. *R. Soc. open Sci.* 5 (2018).
34. Brenner, S. The genetics of *Caenorhabditis elegans*. *Genetics* 77, 71–94 (1974).
35. Pino, E. C., Webster, C. M., Carr, C. E. & Soukas, A. A. Biochemical and high throughput microscopic assessment of fat mass in *Caenorhabditis elegans*. *J. Vis. Exp.* https://doi.org/10.3791/50180 (2013).
36. Escorcia, W., Buter, D. L., Nhan, J. & Curran, S. P. Quantification of Lipid Abundance and Evaluation of Lipid Distribution in *Caenorhabditis elegans* by Nile Red and Oil Red O Staining. *J. Vis. Exp.* 2018 (2018).

**Acknowledgements**

The authors thank Sean Curran, Ralph Baumeister, and Thomas Heimbucher for reagents and useful discussions. Some strains were provided by the CGC, which is funded by NIH Office of Research Infrastructure Programs (P40 OD010440). Funding was provided by Gettysburg College.

**Author contributions**

L.G. and J.R.P. conceived and performed experiments, analyzed and interpreted data, and prepared the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1038/s41598-022-05340-6.

**Correspondence** and requests for materials should be addressed to J.R.P.

**Reprints and permissions information** is available at www.nature.com/reprints.

**Publisher’s note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2022