How and When Do Insects Rely on Endogenous Protein and Lipid Resources during Lethal Bouts of Starvation? A New Application for $^{13}$C-Breath testing

Marshall D. McCue¹ *, R. Marena Guzman¹, Celeste A. Passement¹, Goggy Davidowitz²

¹ St. Mary’s University, Department of Biological Sciences, San Antonio, Texas, United States of America, ² University of Arizona, Department of Entomology, Tucson, Arizona, United States of America

* mmccue1@stmarytx.edu

Abstract

Most of our understanding about the physiology of fasting and starvation comes from studies of vertebrates; however, for ethical reasons, studies that monitor vertebrates through the lethal endpoint are scant. Insects are convenient models to characterize the comparative strategies used to cope with starvation because they have diverse life histories and have evolved under the omnipresent challenge of food limitation. Moreover, we can study the physiology of starvation through its natural endpoint. In this study we raised populations of five species of insects (adult grasshoppers, crickets, cockroaches, and larval beetles and moths) on diets labeled with either $^{13}$C-palmitic acid or $^{13}$C-leucine to isotopically enrich the lipids or the proteins in their bodies, respectively. The insects were allowed to become postabsorptive and then starved. We periodically measured the $\delta^{13}$C of the exhaled breath to characterize how each species adjusted their reliance on endogenous lipids and proteins as energy sources. We found that starving insects employ a wide range of strategies for rationing these limited resources during starvation. Although lipids and proteins are critical metabolic fuels for both vertebrates and insects, insects apparently exhibit a much wider range of strategies for rationing these limited resources during starvation.

Introduction

All animals face the possibility of food limitation during which they must rely solely on endogenous nutrients to fuel their continued metabolic demands. Most of our understanding about
how animals respond to starvation comes from studies of vertebrates (see reviews by [1–6]). While vertebrates demonstrate a number of physiological strategies for surviving starvation these strategies may not necessarily be generalizable to invertebrates such as insects that have also evolved under the omnipresent challenges of food limitation (e.g., [7–11]).

Insects outnumber vertebrates in number, species, and biomass [12]. Because they play central roles in most terrestrial and aquatic food webs [13] it is important to understand how these animals cope with fluctuating food resources. Although the diversity of starvation tolerance among insects may be no less impressive than that documented among vertebrate animals, we know quite little about how they physiologically respond to starvation [14–18]; in fact, recent reviews have explicitly cited the need for additional comparative studies of starvation among insects and other invertebrates [19–21].

Characterizing the progression of starvation

Can researchers use the same physiological toolbox developed for vertebrate animals to study starvation in insects? Starvation-induced changes in blood metabolites used routinely in vertebrates (e.g., glucose, ketone bodies, and nitrogen metabolites) are rarely reported for insects in part because of logistical issues related to body size and peculiarities of hemolymph metabolites (e.g., trehalose and proline [18,22]), and perhaps also in response to the growing awareness that circulating metabolites offer limited mechanistic insight into systemic nutrient fluxes [3,23,24]. Indeed, studies have shown that starving insects mobilize glycogen and triglycerides stored in their fat bodies [17,18,22,25,26], but destructive sampling of body composition precludes continual measurements of fuel oxidation over long periods.

Changes in respiratory exchange ratios (RERs) of starving insects are often difficult to quantitatively interpret [15,19,27] and therefore may preclude accurate assessments of changes in metabolic fuel mixtures [28–30]. This limitation is confounded with the fact that VCO₂ can usually be measured in insects with greater accuracy than VO₂ [31]. Other non-invasive approaches like NMR microscopy can be used to quantify changes in the fat and water content in small insects over time and therefore may be used to estimate lipid oxidation [32]; but the method is not suitable for tracking changes in protein use or for making measurements in large numbers of individuals. Quantitative magnetic resonance (QMR) has been shown to be suitable for accurate measures of lean mass in large insects [33], but it is not suitable for quantifying the lipid content in their bodies.

Recently developed approaches where different nutrient pools in the body (e.g., carbohydrates, lipids, and proteins) are selectively enriched with a stable isotope (e.g., ¹³C) have been coupled with ¹³C-breath testing to characterize the starvation-induced changes in metabolic fuels among birds and mammals [34,35], but they have not yet been used to study starvation in insects. Here we examine whether insects of different species and age classes exhibit strategies of rationing oxidative fuels that are generally similar to those seen among most vertebrates. Based on the traditional three-phase paradigm about fuel switching (reviewed in [1,36–38]), it is likely that carbohydrates in the hemolymph provide a readily available fuel source and therefore lipid and protein oxidation are minimal at the onset of starvation. As starvation progresses, lipids are expected to become the predominant source of metabolic energy. Protein oxidation is expected to increase dramatically, but only during the lattermost phases of starvation when most, or all, of the lipid reserves have been depleted. Because of the ethical concerns of starving vertebrate animals to death, this pre-mortem increase in protein oxidation, often used to delimit the transition from phase II to phase III, is rarely documented in vertebrates. However, insects are convenient models in which to test the prediction that death from starvation is invariably preceded by a dramatic, unsustainable increase in protein oxidation (sensu [39]).
Species selection

In contrast to vertebrates in which much of the work on fuel use during starvation has been done, insects have a greater diversity of life history strategies for growth and development. Insects of some orders are hemimetabolous, whereas others are holometabolous. In general, juveniles of hemimetabolous insects eat the same food as do the adults, whereas juveniles of holometabolous insects typically live in very different habitats and eat substantially different foods than do the adults. Some insects are capital breeders in which (nearly) all of the resources used for reproduction are acquired during the juvenile (larval) phase of growth \[40,41\]. In contrast, income breeders accumulate resources used in reproduction during the adult stage. Insects vary dramatically in size and insect growth is exponential \[42\] so that nearly all of growth occurs in the last larval instar \[43\]. Last, longevity of the juvenile stage relative to the adult stage and longevity of lifespan in general is extremely variable among insect species.

To account for this complexity of life histories we chose five species of insects that span a range of life history strategies with the restriction that we could rear sufficient numbers to accommodate the study. The Madagascar hissing cockroach, *Gromphadorhina portentosa* is hemimetabolous, large sized and relatively longed lived. The eastern lubber grasshopper, *Romalea microptera* is also hemimetabolous and large sized, but has a shorter lifespan. The house cricket, *Acheta domesticus*, is hemimetabolous but smaller with a shorter lifespan than the grasshoppers. The darkling beetle, *Zophobis morio* is a small holometabolous income breeder. The hawk moth, *Manduca sexta*, is a large holometabolous capital breeder. It was not our intention to exhaustively sample all insect life histories, rather, the insects used in this study represent a limited subset of insect life history strategies that represent a significant diversity of strategies to provide a picture of possible strategies of fuel use during starvation. As mentioned above, there are no studies of starvation strategies in insects. With this perspective as a starting point, subsequent studies can test specific hypotheses of starvation strategies across taxa, life history stages, and environmental conditions.

Methods

Animals and experimental diets

The five phylogenetically diverse species of insects were raised in the laboratory on one of two $^{13}$C-labeled tracers ($^{13}$C-1-palmitic acid or $^{13}$C-1-L-leucine; Cambridge Isotope Laboratories, Tewksbury, MA) with the aim of isotopically enriching either their body lipids or proteins, respectively. They were then starved while we measured the δ$^{13}$C in their exhaled breath to track how starvation affected their reliance on endogenous lipid and protein oxidation.

Madagascar hissing cockroaches (*G. portentosa*) nymphs (n = 300; age 1–14 days; 10–20mg) were selected from a larger colony maintained for several generations in our laboratory and randomly assigned to one of two diet treatment groups (Table 1). They were raised to adulthood (4–5 months of age) on a base diet consisting of ground chick starter food (Nutrena; Naturewise) supplemented with one of the two $^{13}$C-labeled tracers. Lots of n = 20 adult cockroaches from each population were placed into five metabolic chambers (1.5 L) where they were starved. The bottom of each metabolic chamber consisted of a false floor of 1cm×1cm wire mesh that allowed feces to pass through thereby preventing coprophagy. Segments of PVC tubing were also placed inside the metabolic chambers to provide a hiding place where they naturally reside during the day. The cockroaches were given 24 hours to become postabsorptive before beginning the breath collection.

House crickets (*A. domesticus*) nymphs (n = 500; age 2 weeks) were obtained from a commercial vendor (Fluker Farms; Port Allen, LA) and were randomly divided into one of two diet
treatment groups (Table 1). They were raised to adulthood (6-weeks of age) on a base diet of ground tilapia pellets mixed with one of the two 13C-labeled tracers. Lots of n = 40 adult crickets from each population were relocated into five metabolic chambers (1.0 L) where they were starved. The metabolic chambers were lined with 1cm×1cm plastic mesh to provide three-dimensional contour. The crickets were given 12 hours to become postabsorptive before beginning the breath collection.

Eastern lubber grasshoppers (R. microptera) nymphs (n = 80, male, 2–3cm) were collected by Prof. John Hatle with permission of a private property owner in Jacksonville, Florida; no collecting permit was required. They were randomly assigned to one of two diet treatment groups and fed a base diet of romaine lettuce leaves lightly dusted with one of the two 13C-labeled tracers (Table 1). They were raised to adulthood over the following 6 weeks. Individual grasshoppers were placed inside 100ml plastic syringes that served as metabolic chambers. The grasshoppers were given 24 hours to become postabsorptive before beginning the breath collection.

Darkling beetles (Z. morio) larvae (mealworms; n = 100, <1cm) were selected from a colony maintained for several generations in our laboratory and randomly assigned to one of two diet treatment groups (Table 1). They were raised to their penultimate larval instar (~300–500mg) on a diet of unlabeled wheat bran and ground tilapia pellets mixed with one of the two 13C-labeled tracers (Table 1). Individual larvae were placed inside 20ml syringes that served as metabolic chambers. The beetle larvae were given 12 hours to become postabsorptive before beginning the breath collection.

Hawk moth (M. sexta) larvae (tobacco hornworms; n = 60) were hatched from eggs of adults from a colony maintained for several generations in our laboratory. They were raised to their penultimate larval instar (~300–500mg) on a prepared diet [44] supplemented with one of the two 13C-labeled tracers (Table 1). Individual larvae were placed inside 60ml syringes that served as metabolic chambers. The larvae were given 12 hours to become postabsorptive before beginning the breath collection.

For all species, ambient temperature was 28°C and photoperiod was 14:10D during the rearing and starvation trials. Breath samples were collected once or twice a day depending on the species. For group-housed animals (i.e., crickets and cockroaches) dead individuals or those who lost normal locomotory ability were removed from cages twice each day to prevent cannibalism. Determination of the post absorptive period was based on size with the three smaller species (cricket, beetle, caterpillar) for a shorter period of 12 h and the two larger species (grasshopper and roach) a period of 24 h.

**Breath sampling and tissue analyses**

During starvation, all animals had access to hydrated gelatin polymer water crystals to maintain humidity within the metabolic chambers and provide them with drinking water so that

| Species                               | Base diet       | $^{13}$C-1-Leucine | $^{13}$C-1-palmitic acid |
|---------------------------------------|-----------------|--------------------|-------------------------|
| Gromphadorhina portentosa             | Chick food      | 1 g kg$^{-1}$      | 1 g kg$^{-1}$           |
| Acheta domesticus                     | Tilapia chow    | 1.667 g kg$^{-1}$  | 1 g kg$^{-1}$           |
| Romalea microptera                    | Romaine lettuce | variable           | variable                |
| Manduca sexta                         | Prepared diet   | 8 g kg$^{-1}$      | 8 g kg$^{-1}$           |
| Zophobis morio                        | Bran+Tilapia chow| 1 g kg$^{-1}$     | 1 g kg$^{-1}$           |

Bran was mixed in with the 13C-labeled tilapia chow for the larval beetles.

Bran was mixed in with the 13C-labeled tilapia chow for the larval beetles.

Table 1. Summary of the five insect species used and their experimental diets used to isotopically enrich their tissues.

doi:10.1371/journal.pone.0140053.t001
Starvation stress was not confounded with dehydration stress [45]. When hydrated this material is >99.9% water and does not provide a significant energy source during starvation. The metabolic chambers were ported to allow fresh air to passively circulate, but these ports were closed prior to gas sampling to ensure CO₂ concentrations were between 2 to 4% at the time of breath collection. Gas samples were collected using gas tight syringes and injected into evacuated 12ml Exetainer™ vials (Labco Limited; Ceredigion, UK). The metabolic cages were cleaned every other day to minimize microbial growth. The starvation trials were stopped once half of the experimental animals succumbed to starvation (i.e., lethal time; LT₅₀). Because the LT₅₀ includes total time food was withheld, it includes the period during which the insects were becoming postabsorptive and thus overestimates the actual ‘starvation’ period by 12 (cricket, and beetle and moth larvae) or 24 (grasshopper, cockroach) hours as described above.

The δ¹³C in each gas sample was analyzed using a HeliFAN Plus nondispersive infrared spectrometer (Fischer, ANalyesen Instrumente GmbH; Germany) interfaced with a FANas autosampler. The ¹³C-analyzer was internally and externally calibrated at the start of each batch of samples. Vials containing a standard gas (2.5% CO₂; Mesa Specialty Gases) were analyzed after every five unknown breath samples to identify and correct for analytical drift.

Postabsorptive insects were selected from each population and killed at the start of the starvation trials to measure their initial tissue δ¹³C. The carcasses were dried to a constant mass at 70°C in a convection oven, ground with a mortar and pestle, and further homogenized using a dental amalgamator. The lipid and lean fractions of the carcasses were chemically isolated using a modified Folch extraction method [46]. The δ¹³C of each fraction was analyzed at the University of Arizona using a Picarro (Sunnyvale, CA) G2121-i Cavity Ring Down Spectroscopy (CRDS) δ¹³C stable isotope analyzer with the A0502 ambient CO₂ interface, an A0201 Combustion Module, and an A0301 gas interface (CM-CRDS). All ¹³C concentrations are expressed in δ¹³CVPDB [47,48].

Calculations and statistics

Because the amount of ¹³C in the exhaled breath is primarily a function of the absolute concentration of the ¹³C in the diets, reporting of the actual δ¹³C of the breath provides only limited insight [49]. It is more informative to document how the δ¹³C of the breath changes as the insects gradually transition from the nourished state to starvation. We therefore used the δ¹³C from the first, postabsorptive breath samples as a starting reference point for each species. All of the subsequent δ¹³C breath values during starvation are expressed in terms of that reference point. For example, positive values mean that the breath of the starving animal contained a higher concentration of ¹³C than the recently postabsorptive animals, whereas negative values mean that the breath contained a lower concentration of ¹³C. We used a one sample t-test against a mean (SigmaPlot 12, San Jose, CA) at each time point to determine whether the δ¹³C in the breath significantly differed from the prestarvation values. Evaluations were made using Because we were asking whether a specific time point along a specific sequence was different from the initial value (and not more generally which values were different) there was no need for a Bonferroni-type correction and therefore α = 0.05.

¹³C-breath testing relies on the assumption that the oxidation and subsequent excretion of ¹³CO₂ from the ¹³C-palmitic acid and ¹³C-leucine tracers is proportional to the rates of lipid and protein oxidation in the whole body. This assumption is not unrealistic because these two monomers comprise such a large component of their parent nutrient pools. For example, palmitic acid accounts for approximately 30% (21.8–46.5%) of the fatty acids in insects used for human consumption [50] and leucine accounts for approximately 7% (1–9%) of all of the amino acid residues in body proteins [50]. It is worth noting that studies of vertebrates have
shown that the relative amount of leucine, an essential amino acid, in the body proteins remained constant despite substantial protein losses over a six-month starvation period [51] supporting our contention that leucine content is an accurate proxy for total protein levels.

The $\delta^{13}$C of the exhaled breath of starving animals, either reported in absolute terms or in terms of the difference between the freshly postabsorptive and the starved states as described above, follows highly complex time-dependent functions. The complexities of these functions are not unlike those seen during the specific dynamic action (SDA) response in postprandial animals. We therefore borrowed and modified some of the descriptive variables routinely used to characterize SDA responses among animals (see [52,53]) to describe the changes in lipid and protein oxidation during starvation. Specifically, we defined several metrics including:

- $LT_{50}$: The time (in days) after which food was removed required for 50% of the individuals of a species to succumb to starvation.
- $lipid_{peak}$: The time (in days) at which peak lipid oxidation occurred denoted by the maximal $\delta^{13}$C in the breath of the palmitic acid treatment groups.
- $lipid_{peak\%LT_{50}}$: The relative timing of the lipid$_{peak}$ expressed in terms of a percent of $LT_{50}$.
- $protein_{minimum}$: The time (in days) at which minimum protein oxidation occurred denoted by the minimum $\delta^{13}$C in the breath of the leucine treatment groups.
- $protein_{minimum\%LT_{50}}$: The relative timing of the protein$_{minimum}$ expressed in terms of a percent of $LT_{50}$.
- $sparing_{duration}$: The time (in days) during which protein sparing was occurring. Protein sparing was defined as the period during which the mean $\delta^{13}$C in the breath of the leucine treatment was lower (negative) during starvation than in freshly postabsorptive insects.
- $sparing_{duration\%LT_{50}}$: The relative length of sparing$_{duration}$ expressed in terms of a percent of $LT_{50}$.
- $protein_{peak}$: The time (in days) at which peak protein oxidation occurred denoted by the maximal $\delta^{13}$C in the breath of the leucine treatment groups.
- $protein_{peak\%LT_{50}}$: The relative timing of protein$_{peak}$ expressed in terms of a percent of $LT_{50}$.

Strategies for maximal longevity

We addressed the question of what the best strategies of resource utilization are to maximize longevity, by plotting the relationships between $LT_{50}$ versus $lipid_{peak}$, $protein_{peak}$, and sparing$_{duration}$. We fit either a linear or a non-linear exponential regression whichever had the higher coefficient of determination ($R^2$).

Results

Tissue enrichment

The tissues of the growing insects became enriched in $^{13}$C roughly in the proportions that their respective diets were isotopically enriched (Table 2). For example, the highest $\delta^{13}$C values were seen in tissues of the moth larvae raised on 8g kg$^{-1}$ (dry mass) of tracers in their prepared diet and the lowest enrichments were seen in the tissues of the beetle larvae raised on unlabeled bran supplemented with tilapia food spiked with 1g kg$^{-1}$ of $^{13}$C-tracer. In general the $^{13}$C from...
the leucine tracer remained in the nonlipid pool and the $^{13}$C from the palmitic acid tracer remained in the lipid pool (Table 2). See the Discussion for more details.

### Oxidation of endogenous lipids

All of the insects increased their rate of lipid oxidation at the onset of starvation, but the responses thereafter were species-specific. The starving grasshoppers quickly increased lipid oxidation and maintained a high reliance on lipid oxidation that peaked at 11 days (lipidpeak), 69% of the LT50 (Table 3). Immediately preceding death their reliance on lipids remained significantly higher than prestarvation values (Fig 1A). The cockroaches, crickets, and larval moths exhibited their lipidpeak during the first third of the starvation period (Table 3), but near the point at which they succumbed to starvation their reliance on lipids was varied. Immediately preceding death the cockroaches exhibited rates of lipid oxidation that were significantly higher than prestarvation levels (Fig 1B). In contrast, after reaching peak lipid oxidation early in the starvation period, the moth larvae gradually reduced their reliance on lipid oxidation.
In the moth larvae these values were not significantly different from the prestarvation values (Fig 2A). The beetle larvae were unique in that they did not rapidly increase lipid oxidation at the onset of starvation and only reached peak lipid oxidation ($\text{lipid}_{\text{peak}}$) near the point of death (Fig 2B). At no time point did any of the starving insects exhibit rates of lipid oxidation (Fig 2C).
Fig 2. Starvation-induced changes in the δ\(^{13}\)C of the exhaled breath of insects. Adult crickets (A) and larval beetles (B) and moths (C) raised on diets supplemented with \(^{13}\)C-palmitic acid tracers (grey) or \(^{13}\)C-leucine tracers (black). The dashed line represents the δ\(^{13}\)C of the breath in postabsorptive insects at the start of fasting. Error bars represent standard deviations. The bold lines indicate time points at which fasting values were statistically different from the prefasting values according to two-way, one-sample t-tests.

doi:10.1371/journal.pone.0140053.g002
oxidation that were significantly lower than the prestarvation states; however, this was not the case with protein oxidation (see below).

**Oxidation of endogenous proteins**

The responses with regard to protein oxidation were more complex than those described above for lipid oxidation. Four of the five insect species exhibited some tendency for protein sparing whereby their reliance on protein oxidation was lower than it was in the prestarvation state, but this response was only significant in the grasshoppers, cockroaches, and moth larvae. Protein minimum occurred during the first quarter of the experiment in cockroaches, crickets, and grasshoppers. These three species were apparently able to spare proteins for 30% to 88% of the starvation period (sparing duration; Table 3). Although the moth larvae were also able to spare proteins for most of the experiment, they exhibited minimum rates of protein oxidation only at the end of the experiment (Fig 2C). The beetle larvae were apparently unable to reduce protein oxidation below prestarvation values at any time point (Fig 2B). Notably, the beetle larvae exhibited peak protein oxidation 18% into the experiment (protein peak; Table 2)—a point at which most other species were exhibiting protein sparing.

At the time of death the reliance on protein oxidation was as varied among the five species as it was for lipid oxidation. For example, the cockroaches and crickets were oxidizing proteins at or near their peak rates that were also significantly higher than starvation rates. In contrast, the grasshoppers and beetle larvae relied on protein oxidation at the time of death to an extent similar to their prestarvation levels. The moth larvae were unique in that they exhibited their minimal reliance on protein oxidation immediately preceding death at rates that were significantly lower than prestarvation rates.

**Strategies of resource use**

Protein peak showed a linear relationship with LT50 ($y = 0.9475x + 2.1826$, $R^2 = 0.99$, Fig 3A). Lipid peak and sparing duration showed non-linear relationships with LT50 (sparing duration: $y = 3.7105e^{0.1175x}$, $R^2 = 0.91$, Fig 3B; lipid peak: $y = 2.7102e^{0.2085x}$, $R^2 = 0.86$, Fig 3C). There were no significant relationships between LT50 and any of the other metrics defined above suggesting these were not as important in determining starvation tolerance.

**Discussion**

**13C Tracer integration into the body**

The $^{13}$C-leucine and $^{13}$C-palmitic acid tracers added to the insect diets were effective at enriching the lean and lipid fractions in the bodies, respectively (Table 2). Experimentally controlling the dose of $^{13}$C-tracers was more effective for species that consumed a homogenized, prepared diet (Table 1) than those that specialized on bran or fresh lettuce (mealworms and grasshoppers, respectively), however it may be possible to individually force feed a fixed amount of tracer to larger species (e.g., grasshoppers; J.D. Hatle, unpublished observation).

Interestingly, in most cases the $\delta^{13}$C of the body lipids of the palmitic acid groups was higher than the $\delta^{13}$C of the lean tissue of the conspecific leucine groups (Table 2). The higher tissue $^{13}$C enrichment within the lipids was initially surprising given that the molecular mass of palmitic acid is nearly twice that of leucine and thus the insects in the leucine treatments ultimately consumed nearly twice the number of tracer-derived $^{13}$C-atoms than those in the palmitic acid treatment. We did not quantify the rates of exogenous (i.e., dietary) tracer oxidation in the insects during growth, but previous studies on birds [54,55], rodents [56], and bats [57], have shown that exogenous leucine is oxidized at a rate nearly an order of magnitude greater.
Fig 3. Correlations between fuel use and longevity. Relationships between starvation tolerance (LT50) and physiological metrics related to fuel oxidation in five species of insects (data from Table 2). The dashed lines illustrate isometry.

doi:10.1371/journal.pone.0140053.g003
than palmitic acid during the postprandial phase. Comparisons of the oxidation of $^{13}$C-labeled leucine and oleic acid (another common fatty acid) in postprandial grasshoppers shows that 1.6% and 0.64% of the exogenous $^{13}$C was recovered, respectively, in the breath by 11 hours [58]. Previous studies of crickets injected with $^{14}$C-radiolabeled tracers into their hemocoel showed that the amino acid, glycine was oxidized about three-fold more rapidly than palmitic acid [59,60]. Consequently, the observed differences in $\delta^{13}$C between the lipid and lean pools in the nourished insects may be explained by the fact that less exogenous palmitic acid was oxidized thereby leaving more to be allocated to the growing tissues [49].

A negligible amount of the $^{13}$C from the leucine tracer was bioconverted into the lipid pool. The mean $\delta^{13}$C of the lipids in control populations (populations that were not exposed to any $^{13}$C-tracers) of cockroaches and crickets (-22.4‰) was not statistically different from the mean $\delta^{13}$C of the lipids in cockroaches and crickets in the $^{13}$C-leucine treatment (-22.0‰; t-test, df = 14, p = 0.107). We did not raise parallel, control populations of the other species, but we assume the basic biochemistry of these insects is generally similar. The minimal 'leakage' (sensu [35]) of the $^{13}$C from the leucine tracer was similar to that reported for mice raised for seven weeks on $^{13}$C-leucine tracers.

A small amount of the $^{13}$C from the palmitic acid tracer was recovered in the lean tissues of the insects. The mean $\delta^{13}$C of the lean tissue in control populations of cockroaches and crickets (-18.5‰) was statistically different from the mean $\delta^{13}$C of the lean tissue in cockroaches and crickets in the $^{13}$C-palmitic acid treatment (-12.9‰; t-test, df = 14, p < 0.001). This extent of $^{13}$C leakage from the lipid pool into the lean pool was greater than previously reported in mice raised on $^{13}$C-palmitic acid [35], yet it remains unclear precisely how the $^{13}$C from the palmitic acid tracer was partitioned between the carbohydrate and protein components of the lean tissue fraction because we did not analyze these fractions separately. Compound specific stable isotope analyses would be useful to determine how the palmitic acid-derived $^{13}$C atoms become distributed among non-lipid components [61–63]. Nevertheless, we estimate that >95% of the $^{13}$C atoms from the palmitic acid tracer remained within the lipid pool of the body, and conclude that the $\delta^{13}$C in the exhaled breath of the palmitic acid treatment groups was an effective proxy of endogenous lipid oxidation during starvation.

What strategies contribute to improved starvation tolerance?

The LT$_{50}$ ranged from 3 days in moth larvae to 52 days in adult cockroaches. Comparisons of starvation tolerance across studies are difficult given the dearth of values reported in literature for these species and life stages. American cockroaches apparently begin to succumb to starvation within two weeks [17,64], and all died by 42 days [65]. Many of the cockroaches in this study lived beyond 60 days. Studies of Gryllus crickets report that they are capable of surviving starvation for two days longer than LT$_{50}$ of the Acheta species in the present study [15,26]. While we did observe a few crickets tolerating at least seven days of starvation we do not consider these to be representative of the general population.

Some crickets and cockroaches are known to reduce their locomotor activity during starvation [26,65]. The crickets and the cockroaches in this study were maintained in groups, and not individually, which may have increased activity and affected their apparent starvation tolerance. We eventually improved our sampling protocols so that we could measure the breath on individuals of the other three focal species. The three-day LT$_{50}$ for the penultimate instar moth larvae is not surprising as other work has shown that these moths enter their last larval instar with almost no fat reserves (Helm and Davidowitz, unpublished data).

A superficial perspective reveals that starvation tolerance in this study was positively correlated with body mass, but we consider this to be a spurious correlation and an artifact of our
selection of two large species and three species (or life stages) that were an order of magnitude smaller (see Methods). Positive correlations have been made between starvation tolerance and body mass in *Drosophila* [21], ant lion larvae [8], and backswimmers [14], but bumble bees show the opposite relationship [66], and we have no evidence that body size in insects is optimized to improve starvation tolerance as it may in mammals [67,68]. Measures of starvation tolerance among other insect species will be useful to confirm this.

Researchers recently developed a dynamic energy budget model to describe different physiological responses to starvation in insects [14]. The model includes three strategies that insects might employ including 1) reducing somatic maintenance costs, 2) maximizing the mobilization of endogenous nutrients, and 3) regulating energy mobilization to only pay for maintenance costs. It was not an objective of this study to document the extent to which these insects engaged in starvation-induced hypometabolism as has been done for many vertebrate species (reviewed in [3,69]). One study on starving *Gryllus* crickets found that they did not reduce metabolic rates during prolonged starvation [15], but future studies designed to document changes in standard metabolic rates and activity levels will be useful to document energy saving strategies during starvation in other species. Nevertheless our results do provide insight into the latter two, mutually exclusive, starvation strategies. If the starving insects were maximizing the rates of nutrient oxidation, we would expect to see evidence of increased protein and increased lipid oxidation at the onset of starvation. While the larval beetles exhibited a response weakly resembling this strategy, all of the other species exhibited inverse changes in endogenous protein and lipid oxidation that are suggestive of physiological regulation and not a state where all possible fuels were maximally oxidized.

The insects in this study exhibited varying strategies with regard to the regulation of fuel oxidation. Are these differences related to their ability to withstand starvation? We found several correlations between survival time (LT50) and particular performance metrics (Fig 3A). The strongest of these was a linear relationship between LT50 and proteinpeak. The relationship between LT50 and proteinpeak was isometric supporting the long-standing idea that starvation tolerance is generally limited by an unregulated increase in protein oxidation. But the strategies occurring during the earlier phases of starvation might play a role in delaying this reliance on protein. If we assume that the five species used in this study are representative of insects in general (an assumption that needs to be tested in future studies), then correlations between the starvation tolerance and duration of protein sparing and the timing of maximal lipid oxidation (lipidpeak) could offer insight into variation in starvation tolerance. In particular, all of the species besides the cockroaches exhibited roughly isometric relationships between sparingduration and lipidpeak versus LT50 (Fig 3B and 3C). Interestingly, the values for the cockroaches did not follow these linear relationships, suggesting possible mechanisms responsible for the comparatively high starvation tolerance in the cockroaches. While each of the aforementioned relationships are correlative, they provide a basis for formulating hypotheses and predictions about which particular physiological strategies are responsible for differences in the starvation tolerance among insects. Future comparisons among the starvation strategies of other insects that exhibit high and low tolerances to starvation could be useful to examine this possibility. It would also be informative to correlate the timing of these physiological events with variation in starvation tolerance within conspecifics at the same life stage that may differ in their adiposity (*sensu* [70,71]); unfortunately, this would involve destructive sampling that would not be compatible with breath testing.

Starvation strategies of insects, like other life history traits, appear to be shaped in part by the life stage [60]. The adult cockroaches, grasshoppers, and crickets tended to exhibit inverse patterns of oxidation of their lipids and proteins. These species are income breeders and would normally allocate ingested food among the demands associated with maintenance, locomotion,
and reproductive activity. However, many other insect species are capital breeders and may normally fast (sensu [72]) during adulthood [73]. It would be useful to characterize how those species partition their lipids and proteins to meet energy demands during adulthood. Growth is an imperative component of the life history of larval insects and the two larval species examined in this study exhibited comparatively low reliance on protein oxidation during starvation, suggesting that larval insects maximize protein sparing to support larval growth. Future studies comparing the starvation strategies among the different life stages of both holometabolous and hemimetabolous species will be useful to characterize how starvation strategies may change ontogenetically.

**Conclusion**

The findings of this study underscore the growing awareness among comparative physiologists that starving animals do not necessarily follow the classic paradigm that canalizes the physiological progression into three discrete phases [74,75]. In fact, only two of the five species exhibited any evidence of sharply increased rates of protein oxidation immediately preceding death as predicted by Lusk (1928) and since retained as part of the paradigm in nutritional physiology [39]. Although there is a need for additional comparative studies of starvation physiology among many key groups of vertebrates and insects [76,77], the diverse responses we describe here raise the possibility that insects employ a broader range of strategies for regulating lipid and protein use than expected. Furthermore, the ability to regulate critical transitions in lipid and protein mobilization may help explain the differences in starvation tolerance, especially the physiological responses that immediately precede death. In addition to characterizing starvation responses in other insect species it will be important to explore the evolutionary underpinnings of this remarkable physiological diversity in the context of phylogenetic relationships and ecological factors.

**Acknowledgments**

We thank Joey Sandoval, Evelyn Oliva, and Crystal Lafleur for helping rear the insect populations and collect breath samples and Autumn Moore for the carbon isotope measures of the solid samples. We are grateful to John Hatle for providing us with grasshopper nymphs.

**Author Contributions**

Conceived and designed the experiments: MDM GD. Performed the experiments: MDM RMG CAP GD. Analyzed the data: MDM GD. Contributed reagents/materials/analysis tools: MDM GD. Wrote the paper: MDM RMG CAP GD.

**References**

1. Castellini MA, Rea LD (1992) The biochemistry of natural fasting at its limits. Experientia 48: 575–582. PMID: 1612138
2. Wang T, Hung CCY, Randall DJ (2006) The comparative physiology of food deprivation: from feast to famine. Annual Review of Physiology 68: 223–251. PMID: 16460272
3. McCue MD (2010) Starvation physiology: reviewing the different strategies animals use to survive a common challenge Comparative Biochemistry and Physiology 156A: 1–18.
4. Kalm LM, Semba RD (2005) They starved so that others be better fed: remembering Ancel Keys and the Minnesota experiment. Journal of Nutrition 135: 1347–1352. PMID: 15930436
5. Navarro I, Gutierrez J (1995) Fasting and starvation. In: Hochachka PW, Mommsen TP, editors. Biochemistry and molecular biology of fishes. New York: Elsevier. pp. 393–434.
6. Lignot J-H, LeMaho Y (2012) A history of modern research into fasting, starvation, and inanition. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 7–24.

7. Abbott KC, Dwyer G (2007) Food limitation and insect outbreaks: complex dynamics in plant-herbivore models. Journal of Animal Ecology 76: 1004–1014. PMID: 17714279

8. Arnett AE, Gotelli NJ (2003) Bergmann’s rule in larval ant lions: testing the starvation resistance hypothesis. Ecological Entomology 28: 645–650.

9. Irwin JT, Lee RE (2003) Cold winter microenvironments conserve energy and improve overwintering survival and potential fecundity of the goldenrod gall fly, Eurosta solidaginis. Oikos 100: 71–78.

10. Renault D, Hervant F, Vernon P (2003) Effect of food shortage and temperature on oxygen consumption in the lesser mealworm, Alphitobius diaperinus (Panzer) (Coleoptera: tenebrionidae). Physiological Entomology 28: 261–267.

11. Wu L, Culver DA (1994) Daphnia population dynamics in western Lake Erie: regulation by food limitation and yellow perch predation. Journal of the Great Lakes 20: 537–545.

12. Gullan PJ, Cranston PS (2010) The Insects: An Outline of Entomology. Oxford: Wiley-Blackwell.

13. Schowalter TD (2006) Insect Ecology: An Ecosystem Approach. Amsterdam: Academic Press.

14. Gerds A, Jager T (2014) Body size-mediated starvation resistance in an insect predator. J Anim Ecol 83: 758–768. doi: 10.1111/1365-2656.12195 PMID: 24417336

15. Sinclair BJ, Bretman A, Tregenza T, Tomkins JL, Hosken DJ (2011) Metabolic rate does not decrease with starvation in Gryllus bimaculatus when changing fuel use is taken into account. Physiological Entomology 36: 84–89.

16. Marron MT, Markow TA, Kain KJ, Gibbs AG (2003) Effects of starvation and desiccation on energy metabolism in desert and mesic Drosophila. J Insect Physiol 49: 261–270. PMID: 12770001

17. Park MS, Park P, Takeda M (2013) Roles of fat body trophocytes, mycetocytes and urocyctes in the American cockroach, Periplanta americana under starvation conditions: an ultrastructural study. Arthropod Structure and Development 42: 287–295. doi: 10.1016/j.asd.2013.03.004 PMID: 23567491

18. Bede JC, McNeil JN, Tobe SS (2007) The role of neuropeptides in caterpillar nutritional ecology. Peptides 28: 185–196. PMID: 17161504

19. Overgaard J, Wang T (2012) Metabolic transitions during feast and famine in spiders. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 53–68.

20. Kirk KL (2012) Starvation in rotifers: physiology in an ecological context. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 25–36.

21. Gibbs AG, Reynolds LA (2012) Drosophila as a model for starvation: evolution, physiology, and genetics. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 37–52.

22. Arrese EL, Soulages JL (2010) Insect fat body: energy, metabolism, and regulation. Ann Rev Entomol 55: 207–225.

23. Jenni-Eiermann S, Jenni L (1998) What can plasma metabolites tell us about the metabolism, physiological state and condition of individual birds? an overview. Biol Cons Fauna 102: 312–319.

24. Price ER, Valencak TG (2012) Changes in fatty acid composition during starvation in vertebrates: mechanisms and questions. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 237–256.

25. Boutton TW, Smith BN, Harrison AT (1980) Carbon isotope ratios and crop analyses of Arphia (Orthoptera: Acrididae) species in southeastern Wyoming grassland. Oecologia 45: 299–306.

26. Konuma T, Morooka N, Nagasawa H, Nagata S (2012) Knockdown of the adipokinetic hormone receptor increases feeding frequency in the two-spotted cricket Gryllus bimaculatus. Endocrinology 153: 3111–3122. doi: 10.1210/en.2011-1533 PMID: 22619358

27. Jensen K, Mayntz D, Wang T, Simpson SJ, Overgaard J (2010) Metabolic consequences of feeding and fasting on nutritionally different diets in the wolf spider Pardosa prativaga. Journal of Insect Physiology 56: 1095–1100. doi: 10.1016/j.jinsphys.2010.03.001 PMID: 20227417

28. Gerson AR, Guglielmo CG (2011) House sparrows (Passer domesticus) increase protein catabolism in response to water restriction. American Journal of Physiology 300: R925–R930. doi: 10.1152/ajpregu.00701.2010 PMID: 21248307

29. Kleiber M (1975) The Fire of Life. Huntington: Krieger. 453 p.
30. Sonko BJ, Fennessy PV, Donnelly JE, Bessesen D, Sharp TA, Jacobson DJ, et al. (2005) Ingested fat oxidation contributes 8% of 24-h total energy expenditure in moderately obese subjects. J Nutr 135: 2159–2165. PMID: 16140892

31. Lighton JRB (2008) Measuring Metabolic Rates: a Manual for Scientists. New York: Oxford University Press.

32. Schilling F, Dworschak K, Schopf R, Kuhn R, Glaser SJ, Haase A (2012) Non-invasive lipid measurement in living insects using NMR microscopy. J Exp Biol 215: 3137–3141. doi: 10.1242/jeb.071209 PMID: 22660788

33. O'Regan SM, Guglielmo CG, Taylor GM (2012) Measurement of arthropod body composition using quantitative magnetic resonance. Invertebrate Biology 131: 215–223.

34. McCue MD, Amaya JA, Yang AS, Erhardt EB, Wolf BO, Hanson DT (2013) Targeted $^{13}$C enrichment of lipid and protein pools in the body reveals circadian changes in oxidative fuel mixture during prolonged fasting: a case study using Japanese quail. Comp Biochem Physiol 166A: 546–554.

35. McCue MD, Pollock ED (2013) Measurements of substrate oxidation using $^{13}$CO$_2$ breath testing reveals shifts in fuel mix during prolonged fasting. Journal of Comparative Physiology 183B: 1039–1052.

36. Belkhou R, Cherel Y, Heitz A, Robin J-P, Le Maho Y (1991) Energy contribution of proteins and lipids during prolonged fasting in the rat. Nutrition Research 11: 365–375.

37. Hervant F, Meathieu J, Durand J (2001) Behavioural, physiological and metabolic responses to long-term starvation and refeeding in a blind cave-dwelling (Proteus anguinus) and a surface-dwelling (Euproctus asper) salamander. Journal of Experimental Biology 204: 269–281. PMID: 11136613

38. Le Maho Y, Van Kha HV, Koubi H, Dewasmes G, Girard J, Ferre P, et al. (1981) Body composition, energy expenditure, and plasma metabolites in long-term fasting geese. American Journal of Physiology 241: E342–E354. PMID: 7304738

39. Lusk G (1928) The Elements of the Science of Nutrition. Philadelphia: W.B. Saunders Co. 844 p.

40. Jonsson KI (1997) Capital and income breeding as alternative tactics of resource use in reproduction. Oikos 78: 57–66.

41. Boggs CL (1997) Reproductive allocation from reserves and income in butterfly species with differing adult diets. Ecology 78: 181–191.

42. Nijhout HF, Davidowitz G, Roff DA (2006) A quantitative analysis of the mechanism that controls body size in Manduca sexta. Journal of Biology 5: 1–15.

43. Davidowitz G, D’Amico LJ, Nijhout HF (2004) The effects of environmental variation on a mechanism that controls insect body size. Evolutionary Ecology Research 6: 49–62.

44. Davidowitz G, D’Amico LJ, Nijhout HF (2003) Critical weight in the development of insect body size. Evolution and Development 5: 188–197. PMID: 12622736

45. Schimpf NG, Matthews DE, White CE (2011) Cockroaches that exchange respiratory gases discontinuously survive food and water restriction. Evolution 66: 597–604. doi: 10.1111/j.1558-5646.2011.01456.x PMID: 22276551

46. McCue MD, Amitai O, Khozin-Goldberg I, McWilliams SR, Pinshow B (2009) Effect of dietary fatty acid composition on fatty acid profiles of polar and neutral lipid tissue fractions in zebra finches, Taeniopygia guttata. Comparative Biochemistry and Physiology 154A: 165–172.

47. Werner RA, Brand WA (2001) Referencing strategies and techniques in stable isotope ratio analysis. Rapid Communications in Mass Spectrometry 15: 501–519. PMID: 11268135

48. Slater C, Preston T, Weaver LT (2001) Stable isotopes and the international system of units. Rapid Communications in Mass Spectrometry 15: 1270–1273. PMID: 11466782

49. McCue MD (2011) Tracking the oxidative and non-oxidative fates of isotopically labeled nutrients in animals. BioScience 61: 217–230.

50. Bukkens SGF (1997) The nutritional value of edible insects. Ecology of Food and Nutrition 36: 287–319.

51. McCue MD (2007) Western diamondback rattlesnakes demonstrate physiological and biochemical strategies for tolerating prolonged starvation. Physiological and Biochemical Zoology 80: 25–34. PMID: 17160877

52. McCue MD (2006) Specific dynamic action: a century of investigation. Comparative Biochemistry and Physiology 144A: 381–394.

53. Secor S (2009) Specific dynamic action: a review of the postprandial metabolic response. Journal of Comparative Physiology 179: 1–56. doi: 10.1007/s00360-009-0283-7 PMID: 18597096
54. McCue MD, McWilliams SR, Pinshow B (2011) Ontogeny and nutritional status influence oxidative kinetics of exogenous nutrients and whole-animal bioenergetics in zebra finches, *Taeniopygia guttata*. Physiological and Biochemical Zoology 84: 32–42. doi: 10.1086/657285 PMID: 21043807

55. McCue MD, Sivan O, McWilliams SR, Pinshow B (2010) Tracking the oxidative kinetics of carbohydrates, amino acids, and fatty acids in the house sparrow using exhaled 13C2O. Journal of Experimental Biology 213: 782–789. doi: 10.1242/jeb.039842 PMID: 20154194

56. McCue MD, Voigt CC, Jefimow M, Wojciechowski M (2014) Thermal acclimation and nutritional history affect the oxidation of different classes of exogenous nutrients in Siberian hamsters, *Phodopus sungorus*. J Exp Zool 321: 503–514.

57. Voigt CC, Sorgel K, Suba J, Keiss O, Petersons G (2012) The insectivorous bat *Pipistrellus nathusii* uses a mixed-fuel strategy to power autumn migration. Proc Roy Soc B 279: 3772–3778.

58. Nicholas J, Awan A, McCue MD, Williams CM, Hahn DA, Hatle JD (2015) Life-extending ovariectomy and dietary restriction each alter leucine metabolism in grasshoppers, but in different ways. Integrative and Comparative Biology 55: E308.

59. Zera AJ, Zhao Z (2006) Intermediary metabolism and life-history trade-offs: differential metabolism of amino acids underlies the dispersal-reproduction trade-off in a wing-polymorphic cricket. American Naturalist 167: 889–900. doi: 10.1086/503578 PMID: 16609924

60. Zhao Z, Zera AJ (2006) Biochemical basis of specialization for dispersal vs. reproduction in a wing-polymorphic cricket: morph-specific metabolism of amino acids. J Insect Physiol 52: 646–658. PMID: 16643945

61. Newsome SD, Wolf N, Peters J, Fogel ML (2014) Amino acids δ13C analysis shows flexibility in the routing of dietary protein and lipids to the tissue of an omnivore. Integrative and Comparative Biology 54: 890–902. doi: 10.1093/icb/icu106 PMID: 25104856

62. Bos C, Metges CC, Gaudichon C, Petzke KJ, Pueyo ME, Morens C, et al. (2003) Postprandial kinetics of dietary amino acids are the main determinant of their metabolism after soy or milk protein ingestion in humans. Journal of Nutrition 133: 1308–1315. PMID: 12730415

63. Mc Clelland JW, Montoya JP (2002) Trophic responses and the nitrogen isotopic composition of amino acids in plankton. Ecology 83: 2173–2180.

64. Park MS, Takeda M (2014) Cloning of PaAtg8 and roles of autophagy in adaptation to starvation with respect to the fat body and midgut of the American cockroach, *Periplanta americana*. Cell Tissue Res 356: 405–416. doi: 10.1007/s00441-014-1802-3 PMID: 24696316

65. Park MS, Takeda M (2008) Starvation supresses cell proliferation that rebounds after refeeding in the midgut of the American cockroach, *Periplanta americana*. J Insect Physiol 54: 386–392. PMID: 18067918

66. Couvillon MJ, Dornhaus A (2010) Small worker bumble bees (*Bombus impatiens*) are harder against starvation than their larger sisters. Insect Soc 57: 193–197.

67. Lindstedt SL, Boyce SJ (1985) Seasonality, fasting endurance, and body size in mammals. American Naturalist 125: 873–878.

68. McNab BK (1999) On the comparative ecological and evolutionary significance of total and mass-specific rates of metabolism. Physiological and Biochemical Zoology 72: 642–644. PMID: 10521332

69. McCue MD, Lillywhite HB, Beaupre SJ (2012) Physiological responses to starvation in snakes: low energy specialists. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 103–132.

70. Beauplet G, Guinet C, Arnould JPY (2003) Body composition changes, metabolic fuel use, and energy expenditure during extended fasting in subantarctic fur seal (*Arctocephalus tropicalis*) pups at Amsterdam island. Physiological and Biochemical Zoology 76: 262–270. PMID: 12794680

71. Goodman MN, McElaney MA, Ruderman NB (1981) Adaptation to prolonged starvation in the rat: catabolism of skeletal muscle proteolysis. American Journal of Physiology 241: E321–E327. PMID: 7315958

72. Mrosovsky N, Sherry DF (1980) Animal anorexias. Science 207: 837–842. PMID: 6928327

73. Boggs CL, Niltepoidk (2014) Insights from stable isotope tracers on reproductive allocation under stress. Integrative and Comparative Biology 54: 880–889. doi: 10.1093/icb/icu074 PMID: 24920750

74. McKenzie DJ, Vergnet A, Chatain B, Desmarais E, Steffensen JF, et al. (2014) Physiological mechanisms underlying individual variation in tolerance of food deprivation in juvenile European sea bass, *Dicentrarchus labrax*. J Exp Biol 217: 3283–3292. doi: 10.1242/jeb.101657 PMID: 25232198

75. Khalilieh A, McCue MD, Pinshow B (2012) Physiological responses to food deprivation in the house sparrow, a species not adapted to prolonged fasting. American Journal of Physiology 303: R551–R561. doi: 10.1152/ajpregu.00076.2012 PMID: 22785424
76. McCue MD (2012) Horizons in starvation research. In: McCue MD, editor. Comparative Physiology of Fasting, Starvation, and Food Limitation. New York: Springer-Verlag. pp. 409–420.

77. Welch KC, Perronet F, Voigt CC, Hatch K, McCue MD (2015) Combining respirometry with stable isotopes to investigate fuel use in animals. Annals of the New York Academy of Sciences in press: