Multiple decrement life tables of *Cephus cinctus* Norton (Hymenoptera: Cephidae) across a set of barley cultivars: The importance of plant defense versus cannibalism

Buddhi B. Achhami\(^1\)*, Robert K. D. Peterson\(^1\), Jamie D. Sherman\(^2\), Gadi V. P. Reddy\(^3\), David K. Weaver\(^1\)

\(^1\) Department of Land Resources and Environmental Sciences, Montana State University, Bozeman, MT, United States of America, \(^2\) Department of Plant Sciences and Plant Pathology, Montana State University, Bozeman, MT, United States of America, \(^3\) USDA ARS-Southern Insect Management Research Unit, Stoneville, MS, United States of America

* buddhiachhami@gmail.com

Abstract

Accurately estimating cause-specific mortality for immature insect herbivores is usually difficult. The insects are exposed to abiotic and biotic mortality factors, causing cadavers to simply disappear before cause of mortality can be recorded. Also, insect herbivores are often highly mobile on hosts, making it difficult to follow patterns for individuals through time. In contrast, the wheat stem sawfly, *Cephus cinctus* Norton, spends its entire egg, larval, and pupal period inside a host stem. Therefore, with periodic sampling stage-specific causes of mortality can be ascertained. Consequently, we examined *C. cinctus* mortality in eight barley, *Hordeum vulgare* L., cultivars in two locations in Montana from 2016 to 2018 by collecting stem samples from stem elongation to crop maturity at weekly intervals, and collecting overwintered barley stubs the following spring and summer from the same plots. If larvae were present, we examined larval status—dead or alive—and categorized dead individuals into one of 5 mortality categories: plant defense, cannibalism, parasitism, pathogens, and unknown factors. We used multiple decrement life tables to estimate cause-specific mortality and irreplaceable mortality (the proportion of mortality from a given cause that cannot be replaced by other causes of mortality). Plant defense (antibiosis) caused 85.7 ± 3.6%, cannibalism (governed by antixenosis) caused 70.1 ± 7.6%, parasitism caused 13.8 ± 5.9%, unknown factors caused 38.5 ± 7.6%, and pathogens caused 14.7 ± 8.5% mortality in the presence of all causes of mortality. Similarly, irreplaceable mortality due to plant defense was 22.3 ± 6.4%, cannibalism was 29.1 ± 4.2%, unknown factors was 6.2 ± 1.8%, pathogens was 0.9 ± 0.5%, and parasitism was 1.5 ± 0.6%. Antibiosis traits primarily killed newly emerged larvae, while other traits supported more favorable oviposition decisions by females, increasing mortality by obligate cannibalism. Our results suggest that breeding barley for resistance to *C. cinctus* targeting both categories of traits (antibiosis and antixenosis) is a highly valuable tactic for management of this important pest.
Introduction

Wheat stem sawfly, *Cephus cinctus* Norton (Hymenoptera: Cephidae), successfully adapted from its ancestral wild grass hosts to domesticated spring and winter wheat cultivars [1]. In addition, this species is not only widening its range southward to Colorado and Kansas [2], but it also is increasing infestation and larval survival in barley in Montana [3,4]. To mitigate economic loss, an integrated pest management (IPM) approach has been recommended for wheat across the northern Great Plains [5]. However, to further formulate and implement IPM tools against this pest, it is important to estimate the mortality of *C. cinctus* across life stages with other crop species in multiple environments.

*Cephus cinctus* is a univoltine insect. In spring, when plant hosts are at the stem-elongation stage, females lay eggs in suitable host stems. Approximately one week after oviposition, a larva emerges. The larva feeds primarily on the parenchymous tissues of the stem until the host plant reaches maturity. Once the host plant ripens and desiccates, the larva moves to the base of the stem, makes a v-shaped groove by chewing a uniform gouge around the stem interior, and plugs the stem below this groove with frass and plant materials to make a hibernaculum. The v-shaped groove contributes to stem lodging when gravity and wind impact the ripened stem. The remaining part of the lodged stem that is intact in the soil and contains an overwintering larva is called a "stub." Within this stub, the larva moves close to the root crown where it hibernates and spends 8–9 months as a prepupa. The following spring, and after sufficient warming, the hibernated prepupa fully metamorphoses before chewing through the frass plug in the stub and emerging as an adult [6,7].

The probability of death of an individual at each developmental stage due to cause-specific mortality can be estimated using a multiple decrement life table [8]. The multiple decrement life table mathematically accounts for a number of mutually exclusive causes of mortality. In addition, the multiple decrement life table can be used to estimate the contribution of each cause of mortality in the presence and absence of other mortality causes.

Parasitism, host plant resistance, and cannibalism are three major causes of mortality of *C. cinctus* in wheat [9]. However, the mortality due to parasitism varies by locations and by host types [10,11]. These three major causes of mortality act on different stages of *C. cinctus* larvae. For instance, host plant defense kills mostly neonates, at an early stage of development, while cannibalism and parasitism occur in instars larger and later than neonates. The larvae that survive plant defense mechanisms are still vulnerable to parasitism and are more vulnerable to cannibalism if multiple live larvae inhabit in a stem [12–14]. Thus, cannibalism and plant defense are important factors influencing *C. cinctus* fitness at both the individual and population levels. However, the role and importance of these three causes of mortality—plant defense, cannibalism, and parasitism—for *C. cinctus* developing in a barley (*Hordeum vulgare* L.) hosts is yet to be determined.

Although barley was historically categorized as resistant to *C. cinctus*, a decade-long study revealed an increasing infestation rate and larval survival rate of *C. cinctus* in barley in Montana [4]. Montana ranked second among barley producing states based on the amount of production in the US in 2019 [15].

Growing solid-stem—filled with pith that reduces the larval survival rate [16–18]—cultivars and swathing are two tools that can mitigate economic loss incurred by *C. cinctus* in Montana wheat crops [19]. However, both management tactics are not implementable for barley because all currently cultivated barley cultivars have hollow stems and swathing can substantially reduce grain quality, especially in malt barley. In Montana, malt barley was 61.8% of total barley production by hectare in 2019 [20].

To reduce the increasing economic losses caused by *C. cinctus* in Montana, estimating age- and cause-specific mortality in populations is crucial. Cannibalism and host plant resistance
are two major causes of mortality in wheat [9]. Both causes of mortality are mediated by host plant traits. For instance, cannibalism ultimately occurs because the host plants release attractive compounds like (Z)-3-hexenyl acetate [21,22] and (E)- and (Z)-β-ocimene to gravid females [23]. These, along with other physical characters, stimulate the abundant gravid females to lay numerous eggs in a stem at high population densities [22,23]. In wheat, stem solidness obstructs larval movement and most of the larvae eventually die [18,24]. The pith also plays a negative role in maternal oviposition decisions in stems with varying solidness due to physically encountered factors [25] and because the hollow stem wheat cultivar releases greater amount of (Z)-3-hexenyl acetate compared to solid stem wheat [26]. Therefore, estimating causes of mortality by barley cultivars will provide a foundation to estimate cultivar-specific mortality and may allow for the development of cultivars that cause consistently greater levels of larval mortality. Consequently, in this study, we estimated age- and cause-specific mortality of *C. cinctus* in eight barley cultivars (Celebration, Champion, Craft, Haxby, Haybet, Hockett, Lavina, and Tradition) at two locations in Montana.

**Materials and methods**

**Field site, field preparation, and seeding**

To estimate *C. cinctus* mortality in barley, we conducted a field study in Montana near Amsterdam in 2016 (45°45’27.3” N, 111°24’00.9” W) and 2017 (45°45’33.2” N, 111°23’50.0” W), and near Big Sandy in 2017 (48°15’42.1” N, 110°22’19.1” W). In each site-year, we planted eight barley cultivars in the spring. The cultivars were ‘Haxby’ and ‘Champion’ (feed barley); ‘Haybet’ and ‘Lavina’ (forage or hay barley); and ‘Celebration,’ ‘Craft,’ ‘Hockett,’ and ‘Tradition’ (malt barley). All cultivars were two-rowed seeded heads except ‘Celebration and ‘Tradition’, which had six-rowed heads.

Each study used a randomized complete block design. The study field was divided into three blocks based on field variability due to slope. Each block was divided into eight plots and each plot was 1.8 m × 3.6 m with a 0.3-m spacing between each plot. The seeding rate was 9 g/m². We performed several manual weeding operations to maintain a better crop stand.

**Summer samples**

We began stem sampling 59 days after seeding because more than 50% of plants were at stem elongation [27]. The stem elongation stage of the host plant is when *C. cinctus* females can first lay eggs because they do not deposit eggs on foliage—or on the later developing seed heads, for that matter. The sampling continued at weekly interval to crop maturity; in total, we collected for 9 consecutive weeks. The procedure for stem sampling was conducted as described previously [3]. For each sampling week, we randomly measured a 0.3-m row length in each plot, uprooted all the plants within the measured length, and wrapped the uprooted plants in a uniquely labelled paper bag. We repeated this uprooting procedure 3 times within a plot; so that each plot had 3 bags of samples per sampling week. Subsequently, we dissected the collected samples lengthwise and, if infested, recorded the status of *C. cinctus* by counting eggs and dead or live larvae. We selected only 35 primary stems from each bag of the samples (the sample that we collected until the 8th week) because all samples could not be dissected on the day they were collected. Unprocessed samples were stored in a 4°C storage room and we dissected them as rapidly as possible on the following day to prevent wilting and to maintain larval status as when collected. In total, we dissected 105 stems per plot per sampling week. However, we dissected all ripened stems from the harvest samples collected at crop maturity (9th week of sampling).
Spring and summer stub samples

We collected stub samples from each plot where we collected the stem samples during the previous summer. We collected stubs twice, at both before and after the flight period of adults. Once again, these stubs are the remaining lower part of a cut stem where the hibernating larva is located. 

The first collection of stubs was conducted in the first week of the following April or May (Amsterdam 2016: 4 April 2017; Amsterdam 2017: 28 April 2018; Big Sandy 2017: 3 May 2018). In mid to late May through early July, female adults emerge, mate, and begin laying eggs. Therefore, we named these collected samples “pre-emergence” or “pre-flight period.” For stub collection, we divided each plot into two equal sections. From the first half of each remaining plot area, we uprooted all the stubs and wrapped these inside a uniquely labelled paper bag. Thereafter, we randomly selected at least 25 stubs that contained hibernaculae with diapaused larvae from each plot and recorded the status, dead or alive, of the overwintered larvae after dissection. We selected 25 stubs to maintain uniform sample size across all cultivars and sites. Additionally, we categorized the dead larvae by mortality causes, including parasitism, pathogens, or unknown factors.

Next, in July (Amsterdam 2016: 10 July 2017; Amsterdam 2017: 12 July 2018; Big Sandy 2017: 20 July 2018), we collected the remaining stubs from each plot which we considered “post-emergence” samples. Then, we again randomly separated 25 stubs per plot, dissected them and recorded the status as either emerged or dead. For assessing potential emergence, we additionally considered whether the top part of the stub, the frass plug, had only a single emergence hole and did not have any indicators of larval or pupal mortality, such as a cadaver or more correctly, fragments of a cadaver, or a parasitoid cocoon. Otherwise, we recorded the life stage as dead by the appropriate cause of mortality.

Causes of mortality

As described, we collected samples for 3 stages of *C. cinctus* development that occur within a stem: summer samples (pre-diapause period), overwintered (pre-flight period), and post emergence (at the end of the flight period). After determining the status, we categorized the death of an individual by one of the following mortality causes: cannibalism, plant defense, parasitism, pathogens, or unknown factors.

Within obligate cannibalism events, we categorized two groups. The first was egg cannibalism when we found a stem with multiple eggs or a larva and an egg(s). The second was larval cannibalism if a stem contained more than one larva. In both cases, a single larva eventually consumes all others, both eggs and larvae, in that stem. Only a single larva per infested stem proceeds to cut the stem and hibernate if the larva survives other causes of mortality [14,24,28,29].

We categorized plant defense as a cause of death if we found a stem contains dead neonates or the stem lining had only couple of millimeters of feeding scars at the larval emergence site as recently described by Buteler et al [9]. Additionally, if we saw only feeding scars, tiny amounts of frass or a small fragment of a dead neonate in the stem, then we also categorized the cause of death as plant defense.

We combined the mortality due to parasitoids and predators under the category of parasitism because we only found two stems with larvae of the clerid predator, *Phyllobaenus dubius* Wolcott (Coleoptera: Cleridae), described in Morrill et al. [30], at the site near Amsterdam in 2016. Thus, otherwise for this paper, “parasitism” represents a stem containing either a parasitoid larva, a parasitoid cocoon or a diagnostic emergence hole in the stem wall. For the rate of parasitism, we categorized the larvae into two groups based on their relative size: early growth...
stages or stages vulnerable to parasitism. The early larval growth stages are not vulnerable to parasitism. Therefore, an early stage before parasitism was possible occurred from the 2\textsuperscript{nd} to 4\textsuperscript{th} week of sampling and the records for the period with stages vulnerable to parasitism occurred from the 5\textsuperscript{th} to 9\textsuperscript{th} week of sampling, as well as being evident in stub samples taken for the pre- and post-flight periods.

If the larval cadaver was covered with white or pink hyphae or was characteristically pink [31], we assumed the cause of death was entomopathogenic fungi and we categorized it as “pathogens.” If the larva mined through multiple internodes but was found dead from other than the above-mentioned mortality causes, it was classified as having died due to “unknown factors.”

**Construction of multiple decrement life tables**

To construct multiple decrement life tables, we used abridged life tables to calculate mortality causes and relevant mortality proportions by those causes in the presence of other mortality causes [32]. We produced the multiple decrement life tables by cultivar and study site using the spreadsheet program M-DEC [33]. After accounting for mortality causes from summer samples, as well as pre-emergence and post emergence stubs, we calculated the percentage of mortality for each cause of mortality in the presence of other mortality causes, and subsequently the irreplaceable mortality. Irreplaceable mortality is mathematical estimates of the degree of impact of a specific mortality factor if it was removed from the mix of mortality factors operating on the insect at the time the study was conducted. For each specific cause, the irreplaceable mortality was determined using the methods described by [10,32–34]. In this technique, the variables are defined as: \(x\) = life stage index, \(l_x\) = the number of individuals alive at each \(x\), \(d_x\) = the total number of deaths in each stage, \(a_{lx}\) = the proportion of the cohort living at the beginning of the stage (starting at 1.0 for the first state and calculated by \(a_{lx} = a_{lx-1} - a_{dx-1}\)), \(a_{dx}\) = proportion of death attributable to one cause, \(a_d\) = proportions of deaths from all causes (\(a_{dx} + a_{dx} + \ldots + a_{dx}\)), \(a_{qx}\) = stage specific probability of dying from all listed causes \((d_x/l_x)\).

Similarly, the probability for cause of death in the absence of other causes can be estimated using a quadratic solution. Elimination-of-cause analysis relies on the probability of surviving each source of mortality \((px)\) and its complement \((1 - qx)\) where \((1 - q_1) \times \ldots \times (1 - q_n)\) is the chance of jointly surviving a set of mortality causes and its complement, \(1 - [(1 - q_1) \times \ldots \times (1 - q_n)]\) is the chance of jointly dying from a set of mortality causes. To estimate mortality in the absence of one or more causes, two simultaneous equations with two unknowns are used. For example, by expression \(q_1\) (e.g. cannibalism) in terms of \(q_2\) (e.g. all other mortality causes), \(D_1\) and \(D_2\) (the proportion of all individuals observed that died of cause 1 and cause 2), this yields the quadratic equation:

\[
a_{q_1}^2 + b_{q_1} + c = 0, \quad \text{where} \quad a = D_1, \quad b = -(D_1 + D_2), \quad \text{and} \quad c = D_2(D_1 + D_2).
\]

The value of \(q_2\) can be calculated by substituting \(a, b,\) and \(c\) in the quadratic formula. Similarly, to calculate irreplaceable mortality for any cause of mortality, for instance irreplaceable mortality due to cannibalism, we subtracted total mortality caused by all estimated mortality causes (cannibalism, parasitism, plant defense, pathogens, and unknown) from the total mortality (parasitism, plant defense, pathogens, and unknown) except for the mortality from cannibalism [10,33].

**Data analyses**

We compared the percentage mortality of each mortality cause in the presence of other mortality causes by cultivars and by sites. In addition, we compared percentage of irreplaceable
mortality caused by each mortality cause by cultivars and by sites. We used linear and nonlin-
ear mixed effects model (nlme) [35] with percentage mortality attributed to each cause as the
response variable, while cultivars, sites and their interaction were fixed effects. Year was con-
considered as a random factor for the analysis. All the percentage data were square-root trans-
formed to meet the assumption of homogeneity of variance, but untransformed means and the
associated standard errors are presented. First, we included a site x cultivar interaction for the
analysis and later dropped it if $P > 0.05$ for interaction. Post-hoc analysis was done whenever
$P \leq 0.05$ by using Tukey HSD (multcomp) [36]. Tukey HSD was used to reduce error due to
unequal sample sizes across cultivars. We used the package ‘scales’ [37] to prepare a heat map
to visualize mortality percentage by mortality causes, by cultivars, and by site x year for the C.
cinctus stages on visualized life tables. Additionally, using the packages ggpubr [38] and
ggcorrplot [39], we prepared correlation plots, by cultivars, to show relationship between mor-
tality percentage caused by plant defense and cannibalism. All data were analyzed in Compre-
hensive R Archive Network (CRAN) version 3.5.3 [40] and visualization by using ggplot2 [41].

Results

A total of 64,076 stems and 3,200 stubs were split to assess C. cinctus mortality among eight
cultivars over 3 site x years. Multiple decrement life tables were produced for each study site
and each cultivar (S2 Table and Fig 1). Cannibalism, plant defense, and unknown factors were
the three main causes of mortality for eggs through pre-diapause larvae, whereas unknown fac-
tors, pathogens, and parasitism were main causes of mortality recorded for large larvae before
and after overwintering (Table 1, Fig 1, S1 and S2 Tables).

The probability of death from a specified cause in the presence of combinations of other
causes by all stages is shown in S1 Table. For instance, cannibalism alone would kill 71% of
population (S1 Table). The combination of three mortality causes—cannibalism, unknown

Fig 1. Heat map displaying the percentage of mortality by each cause of mortality at each developmental stage for Cephus cinctus in barley cultivars that were
grown in Montana. Larva I: Pre-parasitism period; Larva III: Parasitism period; Larva IV: Overwintered larva (pre-flight period; post-parasitism); Larva V:
Overwintered larva (post-flight period; post-parasitism).

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factors, and plant defense—would kill 92% of the population, while the combination of all 5 listed causes of mortality would kill 93% of the population. Thus, the effect of mortality due to parasitism or pathogens for the population in the presence of the other 3 causes was negligible (Table 1, S1 Table).

There was an interaction between site × years and cultivars for the percentage of larval mortality caused by plant defense (\( F_{14,35} = 3.09, P = 0.003 \)). However, the cultivar 'Hockett' had the lowest larval mortality due to plant defense compared to the other cultivars at all sites (Fig 2).

Mortality from cannibalism was significantly different for site × cultivar (\( F_{14,35} = 4.12, P = 0.004 \)). The variation in mortality due to cannibalism was greatest near Amsterdam in 2016 when compared with near Amsterdam in 2017 and near Big Sandy in 2017 (Fig 3).

Parasitism and pathogens caused less than 20% mortality individually, while unknown factors caused approximately 30% (Fig 4). All sites had a similar pattern of mortality within each cause of mortality in all cultivars (Parasitism: \( F_{14,35} = 1.833, P = 0.073 \); Pathogens: \( F_{14,35} = 0.637, P = 0.815 \); Unknown factors: \( F_{14,35} = 1.314, P = 0.248 \)).

There was no site × year effect on irreplaceable mortality by cultivars within the specified cause of mortality: cannibalism (\( F_{14,35} = 0.97, P = 0.493 \)), pathogens (\( F_{14,35} = 0.97, P = 0.494 \)), plant defense (\( F_{14,35} = 0.74, P = 0.715 \)), and unknown factors (\( F_{14,35} = 0.96, P = 0.502 \)). The irreplaceable mortality due to cannibalism in 'Hockett' was greater than for the other cultivars (Table 2). Similarly, irreplaceable mortality due to pathogens was the lowest compared to other causes of mortality (Table 2).

Irreplaceable mortality due to parasitism was less than 4% but was significantly different by cultivar and by site × year (\( F_{14,35} = 2.19, P = 0.029 \)) (Fig 5). The lowest irreplaceable mortality due to parasitism was found near Big Sandy in 2017.

Discussion

Our findings indicate that plant defense, cannibalism, and unknown factors were the three major causes of mortality for \( C. \) cinctus developing in barley (Table 1). Although irreplaceable mortality due to plant defense and cannibalism was similar to previous reports for wheat [9,10], the mortality because of parasitism and pathogens was quite low in our study compared with previous observations made in wheat. Both cannibalism and plant defense are mediated

Table 1. Multiple decrement life tables of wheat stem sawfly, \( Cephus \) cinctus, in barley.

| Category | Alive \( a_{lx} \) | Proportion of dying \( q_{lx} \) | Proportion of living \( a_{lx} \) | Proportion of total dying \( a_{dx} \) | Parasitism \( a_{q1x} \) | Plant defense \( a_{q2x} \) | Cannibalism \( a_{q3x} \) | Unknown \( a_{q4x} \) | Pathogens \( a_{q5x} \) |
|----------|------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Egg      | 8968             | 0.619           | 1               | 0.619           | 0               | 0               | 0.619           | 0               | 0               |
| Larva I  | 18447            | 0.478           | 0.381           | 0.182           | 0               | 0.102           | 0.072           | 0.008           | 0               |
| Larva III| 17254            | 0.554           | 0.199           | 0.110           | 0.010           | 0.063           | 0.009           | 0.027           | 0               |
| Larva IV | 1546             | 0.169           | 0.089           | 0.015           | 0.002           | 0               | 0               | 0.012           | 0.0007          |
| Larva V  | 1499             | 0.1368          | 0.074           | 0.010           | 0.002           | 0               | 0               | 0.006           | 0.0013          |
| Adult    | 1291             | 0               | 0.064           | 0.064           | 0               | 0               | 0               | 0.064           | 0               |
| Total    |                 |                 | 0.015           | 0.165           | 0.701           | 0.054           | 0.002           |                 |                 |

Larva I: Pre-parasitism period; Larva III: Parasitism period; Larva IV: Overwintered larva (pre-flight period; post-parasitism); Larva V: Overwintered larva (post-flight period; post parasitism).

\( a_{lx} \) = proportion of death caused by all the given mortality causes in stage \( x \) given that the individual is alive at the beginning of stage \( x \); \( a_{lx} \) = proportion of survivors at stage \( x \) out of original cohort of all individuals; \( a_{dx} \) = proportion of deaths in stage \( x \) from all the mortality causes; \( a_{qix} \) = proportion of death from cause \( i \) in stage \( x \) in the presence of all other mortality causes given that the individual is live at the beginning of state \( x \).

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by host plant traits, either relative preference by gravid females [26] or antibiosis that limits larval survival rate. Thus, capitalizing on cultivar traits that cause both a greater proportion of antibiotic neonate mortality and those traits that result in obligate cannibalism when attracting more females to oviposit [23,42] are critical considerations to reduce economic losses caused by this species.

Our results indicate that cannibalism caused the greatest mortality in the presence of other mortality causes (Fig 3) and high levels of irreplaceable mortality (Table 2). The mean irreplaceable mortality due to cannibalism was 5.59 ± 1.29% in 'Craft' and 29.15 ± 4.19% in 'Hockett' (Table 2). Buteler et al. [9] observed similar variation in irreplaceable mortality due to cannibalism when comparing susceptible (hollow stem) and resistant (solid stem) wheat cultivars.
Cannibalism is obligate in *C. cinctus* when a stem contains multiple eggs [9,24,28]. Why a cohort of *C. cinctus* females deposit multiple eggs in a stem over a flight period of several weeks is not known, but the following could be potential reasons. First, females prefer to lay eggs in hosts that release greater amounts of attractive volatiles such as β-ocimene, (Z)-3-hexenyl acetate, and (Z)-3-hexenol acetate [23,26] and therefore may indiscriminately lay eggs in a host even though conspecific females have already laid eggs [22,43]. Second, the females are short-lived, typically surviving about 7 days, which may provide an overarching imperative to lay eggs in any available host instead of dying with a greater complement of eggs while searching for an optimal host. For instance, in the absence of primary hosts for oviposition, *C. cinctus* can lay eggs in flax stems [44] and in wheat straw [45]. The behavior of accepting a lower
quality host for oviposition in the absence of higher quality hosts can be accompanied by higher levels of cannibalism on less suitable hosts in phytophagous insect species [46]. However, in our study more cannibalism occurred in cultivars with less evidence of plant defense which also received more eggs; these cultivars are arguably more, rather than less suitable as hosts. Regardless of which of these reasons cause females to lay multiple eggs in a stem, our results strongly suggest cannibalism plays a considerable role in influencing the population dynamics of *C. cinctus* in barley compared to other mortality causes.

Multiple eggs in a stem are not laid by a single female [22,25], but rather by more than one female because of failure to recognize a previously infested stem [22]. This non-discriminatory behavior suggests neither self-provisioning nor opportunistic oviposition facilitates the occurrence of cannibalism in larval *C. cinctus*, as might be expected from studies on species that

![Fig 4. Mean mortality percentages by parasitism, pathogens, and unknown factors by site and cultivar. The percentage of mortality was calculated in the presence of other causes of mortality. Bars without letter within the mortality causes are not different (α = 0.05). Panel A: Parasitism (Cultivar: F\(_7, 35\) = 1.40, \(P = 0.233\)); Panel B: Pathogens (Cultivar: F\(_7, 35\) = 1.47, \(P = 0.208\)); Panel C: Unknown factors (Cultivar: F\(_7, 35\) = 1.31, \(P = 0.248\)). SE is standard error of the mean.](https://doi.org/10.1371/journal.pone.0238527.g004)
oviposit in clutches, some of which may have evolved parental guarding [47–49]. For example, ‘Craft,’ ‘Champion,’ and ‘Celebration’ had fewer stems with multiple eggs so that they had a lower percentage of mortality due to cannibalism compared to ‘Hockett.’ Fewer eggs per stem were likely because these cultivars were less attractive to foraging gravid females. This host-mediated trait of fewer eggs in a stem is somewhat similar to those observed for solid-stem wheat [25], which inhibits the deposition of eggs when probing with the ovipositor.

Individual females do not sacrifice their own eggs by laying multiple eggs in a stem to avoid cannibalism. Cannibalism poses risks to survivors of being infected by species-specific pathogens from their kin and potential mortality due to fighting [50]. We hypothesize that having multiple eggs from different females in a stem is mediated primarily by host plant volatiles and not by cues released from already oviposited eggs [22], yet this strategy favors increased cannibalism.

Cannibalism is a major cause of mortality. From the perspective of population ecology, cannibalism seems to be an important cause of mortality when considering recruitment and it possibly has fitness implications in the host that contains defense compounds or lacks essential nutrients [51]. Thus, fitness can potentially be improved by gaining additional nutritional resources or by reducing the rate of parasitism to larvae [14,52]. Further, cannibalism in other species is a well-known phenomenon to gain additional nutritional resources. For instance, *Harmonia axyridis* Pallas shortens development time and increases survivorship by consuming conspecific larvae than by consuming prey aphid species that sequester plant compounds as a defense mechanism [53]. By consuming conspecific larvae, *H. axyridis* avoids the potential negative effect of plant toxins [53]. Several previous reports [4,54] and the current study have indicated that a large proportion of *C. cinctus* neonate mortality indicates that either the host stem tissues contain antibiotic toxins, or possibly the tissues lack some essential nutrients required for neonate growth and development (categorized as plant defense: Table 2, Fig 2). But by consuming conspecific larvae and eggs, first, each cannibal avoids both tissue toxins or diet deficiency and acquires additional nutrient resources. Second, due to extra nutritional resources, cannibalizing individuals would be more vigorous and could also consume vulnerable parasitized hosts, effectively eliminating the larval ectoparasitoids [13,14]. Third, cannibalism reduces larval host density available for parasitism and can ultimately reduce the rate of parasitism [52,55]. Cannibalism also reduces injury to stem tissue by eliminating other feeding individuals, thus limiting tritrophic communication.

### Table 2. Irreplaceable mortality by cause of mortality and cultivar.

| Cultivar | Irreplaceable mortality percent (mean ± SE) | Cannibalism | Unknown factors | Pathogens | Parasitism |
|----------|---------------------------------------------|--------------|----------------|-----------|------------|
| Plant defense | | | | | |
| Celebration | 18.74 ± 2.69 | 8.96 ± 1.78 | 5.81 ± 1.79 | 0.11 ± 0.07 | 1.03 ± 0.47 |
| Champion | 17.02 ± 3.33 | 9.41 ± 2.17 | 2.24 ± 0.25 | 0.45 ± 0.23 | 0.40 ± 0.16 |
| Craft | 21.65 ± 7.05 | 5.59 ± 1.92 | 1.72 ± 0.46 | 0.04 ± 0.03 | 0.26 ± 0.12 |
| Haxby | 17.27 ± 6.08 | 16.10 ± 3.87 | 2.75 ± 0.96 | 0.51 ± 0.37 | 0.27 ± 0.08 |
| Haybet | 12.26 ± 4.35 | 18.38 ± 2.56 | 5.32 ± 1.40 | 0.10 ± 0.07 | 0.77 ± 0.28 |
| Hockett | 6.42 ± 2.60 | 29.15 ± 4.19 | 4.92 ± 1.06 | 0.40 ± 0.19 | 1.41 ± 0.41 |
| Lavina | 15.61 ± 3.72 | 12.80 ± 2.96 | 6.44 ± 1.22 | 0.28 ± 0.18 | 1.10 ± 0.39 |
| Tradition | 22.29 ± 6.35 | 11.13 ± 3.82 | 6.22 ± 1.80 | 0.39 ± 0.25 | 1.03 ± 0.10 |
| DF | 7.49 | 7.49 | 7.49 | 7.49 | 7.49 |
| F-value | 2.44 | 11.35 | 2.69 | 1.16 | 4.77 |
| P-value | 0.037 | <0.001 | 0.024 | 0.347 | <0.001 |

*SE is standard error of the mean.

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to parasitoids that could immediately favor the infested stem [56]. Thus, cannibalism can increase the fitness of survivors and populations in this system [57].

Our results show that mortality due to plant defense caused the second-highest percentage of mortality (Table 1, S1 Table), while irreplaceable mortality due to plant defense was greatest (Table 2). For example, in the cultivar ‘Craft,’ the percentage of irreplaceable mortality due to plant defense was 4-fold higher than the irreplaceable mortality due to cannibalism, 12-fold higher than irreplaceable mortality due to unknown factors, and almost 20-fold higher than irreplaceable mortality due to parasitism or pathogens (Table 2 and Fig 5). Thus, ‘Craft’ has stronger antibiosis traits compared to the other cultivars.

In wheat, a higher proportion of larval mortality occurs in pith-filled solid stems than in hollow stems [18,58]. Even though all barley cultivars in this study have hollow stems, our
results reveal the percentage of mortality due to plant defense was higher than the mortality reported in any solid-stem wheat cultivar. Therefore, we suggest that larval mortality in barley is likely caused by traits other than those associated with stem architecture.

Barley plants synthesize the defensive alkaloids hordenine (N,N-dimethyltriamine) and gramine (N,N-dimethylindolemethyl-amine), which are feeding deterrents to *Heliothis* spp. [59] and aphids [60,61]. Further, barley plants synthesize protease inhibitors in the form of trypsin inhibitors [62] that decrease the access to essential amino acids and impair protein functions that eventually increase the mortality rate in the spider mite, *Tetranychus urticae* Koch (Arachnida: Tetranychidae) [63]. However, it was beyond the scope of our study to identify associated physical strength measurements or biochemical mechanisms for stem tissue, which can collectively be considered antibiosis, although these both may play important roles in resistance to *C. cinctus*.

Although both cannibalism and plant defense were mortality causes mediated through plant traits, they were negatively correlated (S1–S4 Figs). Plant defense kills neonates at the site of emergence, and it is density independent. Cannibalism occurs after larvae gain a certain size and it is density dependent. Thus, cannibalism allows for survival of one individual in a cohort of multiple immatures within a single stem. As a result, the surviving mature larvae cut the stems in preparation for overwintering, which is agronomically undesirable [19]. Further, due to cannibalism, cannibals consume less stem tissue, which could be a bridge for this species when adapting from wild grass to domesticated wheat and currently, to barley. However, both causes of mortality need to be considered to best manipulate cultivars traits against *C. cinctus*.

Irreplaceable mortality due to pathogens was lowest across all cultivars at < 1% (Table 2). Similarly, Buteler et al. [9] observed that pathogens contributed a small fraction (< 5%) to the mortality of *C. cinctus* in wheat. We did not assess which pathogen species killed larvae. Studies show that entomopathogenic fungi [64,65] and certain *Fusarium* spp. [31] can colonize plant tissue without developing highly apparent symptoms in the plants, but these fungi can kill diapausing larvae [31,65]. Thus, our results suggest that barley cultivars may also harbor entomopathogenic fungi, but the level of mortality was negligible.

The overall parasitism rate was < 20% in the presence of other causes of mortality across all cultivars and sites (Table 1). Similarly, irreplaceable mortality due to parasitism was < 4% (Fig 5, Table 2). Conversely, in wheat, several other studies have reported that *C. cinctus* mortality due to parasitism was as much as 60% [9,10,54,66] and irreplaceable mortality due to parasitism was 7–11% [9,10]. In our study, the irreplaceable mortality due to parasitism was lower compared to wheat; as a result, the mortality from parasitism would most likely have been largely replaced by other mortality causes if parasitism was not present. Further, the discrepancies in the rate of parasitism are associated with cultivar traits. For instance, Buteler et al. [9] found a greater rate of parasitism in *C. cinctus* in a susceptible hollow stem wheat cultivar compared to a resistant solid stem wheat cultivar. The cultivars included in our study had a wide range of resistance levels against *C. cinctus* and that could explain the varying rates of parasitism by cultivars. This is not because plant resistance directly acted on the parasitoid females, but rather because it reduced the number of larval hosts available to parasitoid females. Thus, cultivars with greater levels of plant defense killed neonates, which reduced the number of larval hosts available to parasitoid females, ultimately causing a lower rate of parasitism in those cultivars.

Further, foraging parasitoids generally exploit visual and olfactory cues as well as optimized innate memories to detect their hosts [67]. Additionally, odor cues present on parasitoid pupal cocoons are important when developing host-seeking behaviors [68]. Thus, parasitoid adults might not be able to fully use cues from host *C. cinctus* larvae in infested barley. This is because common use of barley as a host by *C. cinctus* is relatively new, when compared to wheat at our study locations [4] and thus these specialist idiobiont parasitoids, with *C. cinctus* as their only known host, might have a reduced probability of locating large larvae in barley stems.
We conducted our research at two disparate locations. These locations have different agroclimatic conditions (e.g. the duration of crop growth was 115 days near Amsterdam in 2016 and only 99 days near Big Sandy in 2017). Therefore, we observed relative differences in larval mortality, as well as the expression of host-plant performance related to larval mortality, within cultivars across our study sites. However, the overall pattern we observed across the cultivars remains clear, where we see that there is a major tradeoff between plant defense and cannibalism as causes of irreplaceable mortality (Table 2, Fig 5, S1–S4 Figs).

Finally, our overall results suggest that cannibalism and plant defense were the two major causes of mortality for pre-diapause larvae across study sites. Further, irreplaceable mortality due to these two factors has a significant impact that reduces the size of pre-diapause larval populations of *C. cinctus*, which may drive the population growth the following year. Both of these causes of mortality are mediated by host plant traits, either relative preference [26] or antibiosis (Fig 2). In the future, transcriptomic and metabolomic studies could reveal the biochemical pathways that cause a greater proportion of neonate mortality (antibiosis) and those that result in obligate cannibalism when attracting more ovipositing females [23,42]. Thus, cultivars with both of these traits are critical to reducing economic losses caused by this species.

**Supporting information**

S1 Fig. Correlation coefficients of mortality percentages for given causes of mortality in the presence of other causes of mortality, combined over site × years and cultivars. (TIF)

S2 Fig. Correlation coefficients of irreplaceable mortality percentages of given causes of mortality combined over site × years and cultivars. (TIF)

S3 Fig. Correlation of mortality percentage caused by plant defense and cannibalism in the presence of other causes of mortality by cultivar, combined over site × years. Each solid circle represents a replication of each site × years. R indicates the correlation coefficient and shaded area around the blue line indicates standard errors of the regression line for each cultivar. (TIF)

S4 Fig. Correlation between irreplaceable mortality percentages of cannibalism and plant defense by cultivar, combined over site × years. Each solid circle represents a replication of each site × years. R indicates the correlation coefficient and shaded area around the blue line indicates standard errors of regression line for each cultivar. (TIF)

S1 Table. Estimated total mortality in *C. cinctus* for different cause-specific combinations. (DOCX)

S2 Table. Multiple decrement life tables of wheat stem sawfly, *Cephus cinctus*, in barley cultivars that were grown at Amsterdam 2016 and 2017, and Big Sandy 2017 in Montana. (DOCX)

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Author Contributions
Conceptualization: Buddhi B. Achhami, Robert K. D. Peterson, David K. Weaver.
Data curation: Buddhi B. Achhami.
Formal analysis: Buddhi B. Achhami, David K. Weaver.
Funding acquisition: David K. Weaver.
Investigation: David K. Weaver.
Resources: David K. Weaver.
Visualization: Buddhi B. Achhami.
Writing – original draft: Buddhi B. Achhami, Robert K. D. Peterson, David K. Weaver.
Writing – review & editing: Buddhi B. Achhami, Robert K. D. Peterson, Jamie D. Sherman, Gadi V. P. Reddy, David K. Weaver.

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