Influence of dietary nutritional composition on caterpillar salivary enzyme activity

Branislav Babica, Alexandre Poissona, Shireef Darwisha, Jean Lacassea, Magali Merkx-Jacquesa, Emma Desplandb, Jacqueline C. Bedea,*

aDepartment of Plant Science, McGill University, 21,111 Lakeshore, Ste-Anne-de-Bellevue, Quebec, Canada H9X 3V9
bBiology Department, Concordia University, 7141 Sherbrooke St. W., Montreal, Quebec, Canada H4B 1R6

Received 26 June 2007; received in revised form 19 September 2007; accepted 28 September 2007

Abstract

Caterpillars are faced with nutritional challenges when feeding on plants. In addition to harmful secondary metabolites and protein- and water-limitations, tissues may be carbohydrate-rich which may attenuate optimal caterpillar performance. Therefore, caterpillars have multiple strategies to cope with surplus carbohydrates. In this study, we raise the possibility of a pre-ingestive mechanism to metabolically deal with excess dietary sugars. Many Noctuid caterpillars secrete the labial salivary enzyme glucose oxidase (GOX), which oxidizes glucose to hydrogen peroxide and gluconate, a nutritionally unavailable carbohydrate to the insect. Beet armyworm, Spodoptera exigua, larvae were restricted to diets varying in protein to digestible carbohydrate (P:C) ratio (42p:21c; 33p:30c; 21p:42c) and total nutrient concentration (42% and 63%). High mortality and longer developmental time were observed when caterpillars were reared on the C-biased, P-poor diet (21p:42c). As the carbohydrate content of the diet increased, caterpillars egested excess glucose and a diet-dependent difference in assimilated carbohydrates and pupal biomass was not observed, even though caterpillars restricted to the C-biased diet (21p:42c) accumulated greater pupal lipid reserves. Larval labial salivary GOX activity was also diet-dependent and gluconate, the product of GOX activity, was detected in the frass. Unexpectedly, GOX activity was strongly and positively correlated with dietary protein content.

Keywords: Larval performance; Salivary enzymes; Nutritional quality; Spodoptera exigua

1. Introduction

Invertebrate herbivores, particularly generalists, face dietary nutritional variation in plants, in terms of species, developmental stage and tissues and responses to the environment (Bae and Sicher, 2004; Cross et al., 2006; Mattson, 1980; Roberts and Paul, 2006; Roslin et al., 2006; Yeoh et al., 1992). The proper feeding decision is critical and can greatly impact herbivore growth, development and reproductive success (Auger, 1995; Awmack and Leather, 2002; Barton-Browne, 1993). Therefore, caterpillars must be able to monitor their nutrient intake and often have behavioral and/or biochemical mechanisms to cope with non-ideal foods (Bede et al., 2007; Behmer et al., 2002; Raubenheimer and Simpson, 2003; Simpson and Simpson, 1990; Waldbauer and Friedman, 1991). One such mechanism may be to switch food sources (Behmer et al., 2002; Bernays et al., 2004; Hägele and Rowell-Rahier, 1999; Moreau et al., 2003; Wright et al., 2003). Although caterpillars are relatively mobile (McAuslane and Alborn, 2000; Smits et al., 1987) and may, within one or amongst closely neighboring plants, find dietary variation between protein-rich reproductive tissues and carbohydrate-rich leaves (Cross et al., 2006; Mattson, 1980; Roulston et al., 2000), caterpillars are often restricted to nutritionally unbalanced diets and, presumably, have mechanisms to cope with nutrient variation.

Dietary proteins (P) and digestible carbohydrates (C) and, in particular, the ratio of these two nutrients (P:C) are critical for proper insect growth and development (Simpson and Raubenheimer, 1993). Generalist caterpillars often
select and perform better on a slightly P-biased diet (Lee et al., 2006, 2002; Merkx-Jacques et al., in press; Waldbauer et al., 1984). However, plants, particularly forbs, are rich in digestible mono- and disaccharide carbohydrates and are relatively nitrogen-limited (Mattson, 1980; National Research Council, 2001; Scriber and Slansky, 1981), which may negatively impact caterpillar growth and development (Felton, 1996; Karowe and Martin, 1989; Lee et al., 2002, 2003; Merkx-Jacques et al., in press; Raubenheimer et al., 2005). Not only are plants often P-limited, but in response to herbivory, they often have strategies to obstruct protein digestion by the caterpillar such as inducible proteinase inhibitors and amino acid deaminases that specifically digest proteins by the caterpillar such as inducible proteinase inhibitors and amino acid deaminases that specifically digest proteins (Chen et al., 2005; Kang et al., 2006; Ryan, 1990). On these P-poor diets, many caterpillar species engage in compensatory feeding (Raubenheimer et al., 2005; Simpson et al., 2004; Warbrick-Smith et al., 2006), exposing them to even greater C excesses and more noxious plant secondary metabolites. For many insect species, the necessity to achieve the optimal balance of dietary P:C results in tradeoffs between growth (protein) and accumulating sufficient lipid stores (carbohydrate) for adults, and respiratory metabolism (Simpson and Simpson, 1990; Telang et al., 2003; Zanotto et al., 1997). Adaptation to excess dietary C is so critical that generalist caterpillars of Plutella xylostella show generational plasticity, which may come at a fitness cost (Warbrick-Smith et al., 2006).

Another strategy to cope with nutritional imbalances is to metabolize nutrients before ingestion by secreted salivary enzymes. Differential salivary digestion has been observed for the desert spider, Stegodyphus lineatus, where nutrients were differentially extracted from their prey (Mayntz et al., 2005). During the feeding process, caterpillars secrete saliva onto the plant (Ribeiro, 1995) to digest and transport chewed plant tissue into their buccal cavity. In addition to aiding digestion, salivary enzymes may detoxify noxious plant secondary metabolites or, as is proposed in this paper, help to regulate nutrient levels in the plant before ingestion. Many Noctuid caterpillars, including Spodoptera exigua, Helicoverpa armigera, Helicoverpa assulta and Helicoverpa zea possess the salivary enzyme glucose oxidase (GOX) (Eichenseer et al., 1999; Merkx-Jacques and Bede, 2005; Zong and Wang, 2004), which catalyzes the oxidation of glucose to hydrogen peroxide and gluconate, which to the insect is a dietary unavailable form of carbohydrate. A number of roles for caterpillar salivary GOX have been proposed (Eichenseer et al., 1999; Musser et al., 2002, 2005a, b). The hydrogen peroxide produced by this enzyme may act as a signaling molecule and interfere with induced plant defensive pathways that are induced in response to caterpillar herbivory (Bede et al., 2006; Musser et al., 2002, 2005a) or an antimicrobial agent (Musser et al., 2005b). The consumption of oxygen by the reaction may help maintain an anaerobic midgut which would lower the reactivity of plant compounds, such as phenolics and tannins in the ingested leaves (Eichenseer et al., 1999). We propose that, in addition, salivary GOX may be a pre-ingestive way for the caterpillar to cope with dietary C excesses since by metabolizing glucose to gluconate, caterpillars may dispose of excess dietary sugars in nutritionally imbalanced diets.

Caterpillar salivary GOX activity is diet-dependent in S. exigua caterpillars (Merkx-Jacques and Bede, 2005). When caterpillars are transferred from the legume, Medicago truncatula, to a wheat germ-based artificial diet, salivary GOX activity significantly increases (Merkx-Jacques and Bede, 2005). Peiffer and Felton (2005) also noted that caterpillar salivary GOX is affected by plant diet; enzyme activity decreases as plant nutritional quality increases.

In this study, we wished to determine if caterpillar salivary GOX could be a pre-ingestive mechanism to cope with dietary C excesses which allows these caterpillars to achieve optimal performance on P-rich diets. Previous studies have shown that when given choices, S. exigua caterpillars select a diet with an intake target of 22p:20c (Merkx-Jacques et al., in press). As noted in other Lepidopteran species, when S. exigua caterpillars are restricted to a C-rich, P-poor diet, high mortality and delayed development are observed (Lee et al., 2002, 2003, 2004a). However, in contrast to related Spodoptera species, higher pupal lipid reserves and biomass was not observed in S. exigua caterpillars reared on P-rich diets (Karowe and Martin, 1989; Lee et al., 2002, 2004a). Therefore, S. exigua caterpillars exhibit flexible and efficient pre-ingestive nutrient intake regulation and post-ingestive utilization and we wished to determine if salivary GOX is a possible mechanism involved in this nutrient balancing.

To answer this, S. exigua caterpillars were restricted to chemically defined diets ranging in their P:C ratio over their development from 3rd larval instars to pupation. These diets only contained glucose as their digestible carbohydrate source. Larval stadium duration, mortality and pupal biomass, indicators of caterpillar performance, were assessed. Carbohydrates egested in the frass (including glucose and gluconate) as well as those converted to lipid stores in the pupa were measured to determine possible mechanisms of carbohydrate metabolism. Salivary GOX activity in actively feeding 4th instar caterpillars was also examined to see if diet affected enzyme activity.

2. Materials and methods

2.1. Caterpillar colony

Eggs of the beet armyworm, S. exigua (Hübner) (Lepidoptera: Noctuidae) were obtained from AgriPest...
Inc. (Zebulon, NC). The laboratory colony was reared for multiple generations on an artificial wheat germ-based diet (Bio-Serv, Frenchtown, NJ) in a growth chamber (16:8 light:dark hours; 28–40% relative humidity; 28.5 °C). Adult moths were allowed to mate and collected eggs were used to maintain colony.

2.2. Defined synthetic diet

Chemically defined artificial diets were prepared according to Simpson and Abisgold (1985). All ingredients were obtained from Bio-Serv unless otherwise indicated. Four diets were prepared containing different ratios of protein:-

2.7. Catalogue of artificial diets

Glucose (Sigma) and gluconate (Megazyme) levels were determined in the caterpillar frass according to manufacturer instructions and modified for a 96-well plate format. μ-Glucose levels were specifically measured spectrophotometrically using a GOX/peroxidase-o-dianisidine coupled reaction (Sigma). A standard curve was prepared from a NIST standard glucose solution. All frass samples were assayed in triplicate. An experimental control without frass (blank) was included. μ-Gluconate/δ-glucono-δ-lactone levels were specifically measured by a 6-phosphogluconate dehydrogenase/gluconate kinase assay (Megazyme). Changes in NADPH levels were measured spectrophotometrically. Gluconate standard curve and all frass samples were assayed in triplicate and an experimental control without frass (blank) was also included.

Calculations of carbohydrate assimilation were estimated from the amount of glucose ingested (see Section 2.3) minus the glucose and gluconate detected in the frass.

2.7.1. Experiment 1

Since the hypothesis being tested was that caterpillar salivary GOX is a pre-ingestive mechanism to deal with excess dietary carbohydrates, caterpillars were reared from eggs on a slightly P-biased diet, 22p:20c, which had low carbohydrate levels to ensure that they had minimal background GOX activity. Third instar caterpillars were individually restricted to chemically defined foods (21p:42c, 22p:20c, 33p:30c, 42p:21c). In this first experiment, cater-
pills were left on these diets for as long as possible (approximately 72 h) to ensure that they had time to adjust their salivary GOX activity to the diet. Labial salivary glands were dissected from mid-5th instar caterpillars and assayed for GOX activity. A diet-dependent difference in mass of the caterpillars used in this experiment was not observed (average mass = 0.47 ± 0.02 g; \( F_{(2,135)} = 0.53, p = 0.67 \)). Cold-anesthetized caterpillars were placed ventral side up and rinsed with cold, sterile Nathanson’s saline (150 mM NaCl, 3 mM KCl, 3 mM CaCl\(_2\), 10 mM TES, 20 mM MgCl\(_2\), pH 6.9; Christensen et al., 1991) to remove any debris or frass. The head was turned to expose the salivary glands and the salivary glands were placed in a microfuge tube containing saline and a broad-spectrum proteinase inhibitor (Sigma). Pairs of labial salivary glands from four caterpillars were pooled and homogenized in Nathanson’s saline with proteinase inhibitor. The homogenate was centrifuged and the supernatant filtered using 0.22 \( \mu \)m low protein binding membrane filters (Millipore). The GOX activity of the soluble homogenate was assayed using the o-dianisidine-horseradish peroxidase assay (Bergmeyer, 1974). A reaction cocktail containing 10% \( \beta \)-D-glucose, 0.0066% o-dianisidine in 0.1 M potassium phosphate buffer, pH 7.0 was brought to 35 °C. The salivary gland homogenate and horseradish peroxidase (final concentration 3 U; Sigma) were added to the reaction cocktail and GOX activity was measured in duplicate spectrophotometrically at 460 nm. Controls included all reaction compounds except for the salivary gland extract (blank) and boiled homogenate (negative control). Fungal GOX (Sigma) was used as a positive control. Enzyme activity (blank) and boiled homogenate (negative control). Fungal GOX activities and assayed for GOX activity as outlined above (Section 2.7.1). A diet-dependent effect on caterpillar biomass was not seen (average mass = 0.079 ± 0.006; \( F_{(2,135)} = 0.056, p = 0.95 \)). To determine if P level affected the level of salivary proteins, soluble labial salivary protein levels were measured by the Bradford assay (Bradford, 1976) as described above. Six to eight biological replicates of pooled pairs of labial salivary glands were analyzed for each diet.

### 2.8. Statistical analyses

Statistical analyses were performed using SPSS version 14 (SPSS Inc., Chicago, IL). Diet consumption and performance of caterpillars surviving until pupation were analyzed using general linear models, in particular, analysis of variance (ANOVA), analysis of covariance (ANCOVA) and multiple analysis of covariance (MANCOVA). Statistical differences were determined by Tukey post-hoc tests. Significant differences (\( p \leq 0.05 \)) are indicated in graphs and tables by different alphabetical letters. Caterpillar survivorship was analyzed by Fisher’s exact test.

### 3. Results

#### 3.1. Nutrient intake and performance


Previous studies have shown that *S. exigua* caterpillars select an intake target of 22p:20c (Merkx-Jacques et al., in press). On P-biased and C-biased diets, caterpillar face nutritional excesses (Fig. 1) (Lee et al., 2002, 2006; Raubenheimer and Simpson, 1999; Waldbauer et al., 1984).

A survival of greater than 86% was observed, irrespective of diet (Fisher’s exact test, \( p = 0.39 \)) (Fig. 2A). Caterpillars took longer to reach pupation on the C-biased, P-poor diets (two-way ANOVA, \( F_{(3,108)} = 17.417, p < 0.001 \)) (Fig. 2B). Supernumerary 6th instar caterpillars were occasionally observed on all diets. The pupal biomass attained by these caterpillars restricted to the different foods was not statistically different (two-way ANOVA, \( F_{(3,108)} = 0.683, p = 0.565 \)). Caterpillar gender did not significantly affect the nutritional or performance criteria which were assessed in this study (total food consumption: \( F_{(1,108)} = 0, p = 0.99 \); consumption rate: \( F_{(1,108)} = 0.20, p = 0.89 \); development time: \( F_{(1,105)} = 0.17, p = 0.69 \); pupal biomass: \( F_{(1,108)} = 0.17, p = 0.69 \).

![Fig. 1. Nutrient consumption rate of Spodoptera exigua caterpillars](image-url)
and insects restricted to the C-biased diets have pre- and/or post-ingestive mechanisms for disposing of excess carbohydrates.

Insect gender did not effect the amount of carbohydrates eaten or retained or the accumulation of pupal lipid stores (C ingestion: $F_{(1,95)} = 0.11$, $p = 0.74$; C retention: $F_{(1,39)} = 1.85$, $p = 0.18$; pupal lipids: $F_{(1,95)} = 0.57$, $p = 0.45$).

### 3.3. Caterpillar labial salivary glucose oxidase activity

#### 3.3.1. Experiment 1

The activity of caterpillar labial salivary GOX was significantly affected by diet ($U/LSG: F_{(3,23)} = 14.017$, $p < 0.001$; U/mg x pair LSG: $F_{(3,23)} = 14.319$, $p < 0.001$) (Fig. 4); this appears to reflect the P level of the food. In terms of total activity ($U/$pair LSG) or activity per milligram soluble protein ($U/$mg x pair LSG), a significant difference in GOX activity was not observed between caterpillars on diets with comparable protein levels (22p:20c vs. 21p:42c). Instead, the highest salivary GOX activity was found in caterpillars transferred to 42p:21c.

#### 3.3.2. Experiment 2

In the previous experiment, dietary protein appeared to affect labial salivary GOX activity, but this was difficult to conclusively determine since carbohydrate levels were also changing. In this experiment, 3rd instar caterpillars fed for a shorter time on a chemically defined diet which had the same glucose composition (30%) but differed in protein levels (5%, 15%, or 33%). Diet significantly affected the activity of caterpillar labial salivary GOX ($U/$pair LSG: $F_{(2,17)} = 43.46$, $p < 0.001$) (Fig. 5A), with the highest activity associated with caterpillars feeding on the P-rich diet (33p:30c). Soluble protein levels in the labial salivary gland of these caterpillars were also significantly higher on the P-rich diet (mg/LSG: $F_{(2,17)} = 24.21$, $p < 0.001$) (Fig. 5B).

### 3.4. Frass carbohydrate composition

Caterpillar frass was analyzed for glucose and gluconate: two possible glucose-associated excretion products. Gender did not effect frass glucose or gluconate levels (glucose: $F_{(1,46)} = 0.32$, $p = 0.58$; gluconate: $F_{(1,53)} = 0.63$, $p = 0.43$). Frass glucose levels increased with the digestible C content of the diet (total glucose (mg): $F_{(3,46)} = 53.349$, $p < 0.001$; Glucose/frass (mg/mg): $F_{(3,46)} = 77.900$, $p < 0.001$) (Fig. 6A). In comparison, frass gluconate was only present at low levels and varied slightly between treatments (two-way ANOVA, $F_{(3,53)} = 2.944$, $p = 0.043$) (Fig. 6B). Cellulose, gluconate and glucose account for the dry weight of the frass from caterpillars reared on the C-biased diet (Fig. 6C). In comparison, caterpillars reared on equal-ratio and P-biased diets also egest a small percentage of unknown components, which presumably contains protein and/or amino acid metabolites.
Discussion

Previous studies have shown that the self-selected nutrient intake ratio for *S. exigua* caterpillars is a slightly P-biased diet (22p:20c, 42%) (Merkx-Jacques et al., in press). In this study, we wanted to evaluate caterpillar coping strategies when restricted to diets containing nutritional excesses, specifically to determine if salivary GOX may be a mechanism for the caterpillar to cope with excess dietary carbohydrates. Therefore, *S. exigua* caterpillar performance was assessed on a range of diets which had higher nutritional concentrations (63%) were compared to this self-selected diet. This is consistent with plant nutrient content which commonly ranges from 40% to 66% with typical P:C values for the foliage of herbaceous dicots in the range between 10–16% P and 40–45% C (National Research Council, 2001; Yeoh et al., 1992).

It was unexpected that *S. exigua* caterpillars did show evidence of compensatory feeding when restricted to the dilute diet (22p:20c vs. 30p:30c) as has been observed for other closely related *Spodoptera* caterpillar species (Lee et al., 2004a, b; Wheeler and Slansky, 1991). We speculate that the reason for this is that even though 22p:20c has roughly 66% of the nutritive value of 30p:30c, it still has satisfactory dietary value, compared to the extremely dilute samples used in previous studies (i.e. 8.4p:8.4c) (Lee et al., 2004b).

On C-biased, P-poor diets *S. exigua* caterpillars must still attain sufficient protein resources for growth and development. Therefore, in this and other studies, longer developmental stadia are observed on these diets (Despland and Noseworthy, 2006; Lee et al., 2002, 2003, 2004a, 2006; Manuwoto and Scriber, 1985; Petersen et al., 2000; Raubenheimer et al., 2005; Telang et al., 2001; Thompson et al., 2005; Waldbauer et al., 1984; Whitford et al., 1992). Supernumerary 6th instar caterpillars are observed on all diets; however, more caterpillars required an additional instar to attain proper resources on the P-poor diet.

In *S. exigua*, pupal biomass, which strongly reflects lipid reserves, is positively and closely correlated with adult female egg production (Tisdale and Sappington, 2001). In a previous study, a diet-dependent difference in *S. exigua* pupal biomass and lipid reserves was not observed (Merkx-Jacques et al., in press). This suggested that.

---

**Fig. 3.** Pupal lipid content and lipid conversion efficiency. 2nd instar *Spodoptera exigua* caterpillars were restricted to a single food (nutrient concentration 42%: 22p:20c; nutrient concentration 66%: 42p:21c, 33p:30c, 21p:42c) until pupation. (A) Conversion of total ingested carbohydrates into pupal lipids: scatterplot illustrates the relationship between pupal lipid accumulation and total digestible carbohydrate ingested by caterpillars restricted to different diets. (B) Digestible carbohydrates assimilated: scatterplot illustrates the relationship between the amount of digestible carbohydrates eaten and retained by caterpillars restricted to different diets. (C) Carbohydrate retention: scatterplot illustrates the relationship between the amount of glucose ingested and the carbohydrates egested by caterpillars restricted to different diets. (D) Conversion of assimilated carbohydrates into pupal lipids: scatterplot illustrates the relationship between the amount of digestible carbohydrates retained by caterpillars restricted to different diets and their pupal lipid reserves.
caterpillars on the C-poor diet had efficient mechanisms to process carbon skeletons from excess amino acids. In this experiment where caterpillars were reared on diets with a higher nutritional concentration, a food-dependent difference in pupal biomass was not observed but higher pupal lipid reserves were found in caterpillars reared on the C-rich diets (Fig. 3A). Similar findings were observed in studies with related Spodoptera species where pupal lipid stores increase when caterpillars are reared on a C-rich diet compared with those on the P-rich, C-poor diets (Lee et al., 2002, 2003, 2004a, b). Nonetheless, the amount of lipid stored relative to carbohydrate consumed was higher on P-biased foods, confirming that the efficiency of carbohydrate metabolism increases with P-bias of the diet (Fig. 3A).

Caterpillars on the C-biased diets egested more glucose (Fig. 6C), leading to similar values for carbohydrate assimilation, despite differences in consumption on different diets (Fig. 3B and C). These findings suggest a pre-absorptive mechanism for C regulation, where excess glucose is voided directly in the frass. This contrasts with previous observations with hemimetabolous locusts, where higher carbohydrate assimilation was observed on C-limited diets and locusts voided excess carbohydrates post-ingestively by increasing their respiration (Zanotto et al., 1993). In this study, we were interested in determining if a pre-ingestive mechanism of C regulation, such as the use of salivary enzymes, may occur in addition to mechanisms such as the elevation of respiration rate.

Labial salivary GOX is a digestive enzyme produced and secreted through the spinneret onto leaf material by many Noctuid caterpillar species (Eichenseer et al., 1999; Merkx-Jacques and Bede, 2005; Zong and Wang, 2004). Recently, there has been speculation as to the biological role of this enzyme, particularly since its activity appears to be regulated by diet (Merkx-Jacques and Bede, 2005; Peiffer and Felton, 2005). GOX catalyzes the oxidation of glucose to gluconate, making carbohydrates inaccessible to the caterpillar. Gluconate was identified in caterpillar frass (Fig. 6B), albeit at low levels, indicating that GOX is involved in the conversion of glucose to gluconate which is egested. These levels probably do not represent actually enzyme conversion since, though non-accessible to insects, gluconate is an important carbon source for some bacteria, including many gut-associated Enterobacteria, such Escherichia coli (Chang et al., 2004; Peekhaus and Conway, 1998; Portais and Delort, 2002).

Therefore, secretion of GOX may be a strategic adaptation of generalist caterpillars to oxidize excess...
dietary carbohydrates that they may encounter. Based on this hypothesis, we expected labial salivary GOX to reflect the digestible carbohydrate content of the diet and be highest on sugar-rich sources. Instead, there was a strong relationship between dietary protein content and GOX activity. Labial salivary GOX activity \( \text{U/mg} \times \text{pair LSG} \) was over three-fold higher in caterpillars transferred to the protein-rich (42p:21c) than the carbohydrate-rich (21p:42c) diet (Fig. 4). When the C-content of the diet was held constant, GOX activity was associated with high P content and an overall increase in soluble salivary proteins were observed (Fig. 5A and B). This implies that on protein-limited diets, *S. exigua* caterpillars may be “unwilling” to synthesize excess digestive enzymes which are secreted into the environment, resulting in a loss of nitrogen which is already limiting.

Findings from this study seem to imply that GOX is not a metabolic strategy to cope with excess carbohydrates. On the C-limited, P-rich diet, C utilization efficiency was increased (Fig. 3A) and on the C-rich diets, caterpillars coped with glucose excesses by increasing C egestion. However, nutrient utilization by caterpillars reared on a P-biased diet (42p:21c) with those reared on their intake target (22p:20c) is worth focusing attention on. On a daily basis, caterpillars reared on these diets ingested similar amounts of digestible carbohydrates (Fig. 1) and reached similar performance in terms of mortality, development, pupal lipid stores and pupal biomass (Fig. 2A and B). Yet, caterpillars on the P-biased diet, which had higher salivary GOX activity, had less glucose egested in the frass (Fig. 6A); this undoubtedly reflects the increased C utilization efficiency of these caterpillars, but, perhaps, also supports the notion the caterpillar salivary GOX does indeed metabolize excess glucose, but much of the gluconate produced is metabolized by gut-associated bacteria.

### 4.1. Ecological implications

Understanding how caterpillars mediate feeding decisions and nutrient utilization and, hence, the regulation of GOX activity, is not only important from both ecological and pest management viewpoints, but also because salivary GOX has also been implicated in subverting plant defense responses (Musser *et al.*, 2002, 2005). The hydrogen peroxide produced by this enzyme activity is believed to act as a signaling molecule and acts to attenuate octadecanoid- or ethylene-mediated plant defense responses (Beckers and Spoel, 2006; Shoji *et al.*, 2000; Winz and Baldwin, 2001).

This study indicates that caterpillar labial salivary GOX activity is closely correlated with dietary protein intake. Martin (1991) proposed that for the insect, it is not advantageous to produce enzymes involved in complex carbohydrate digestion, such as cellulases which degrade the cellulose polymers to glucose sugars, since the plant provides sufficient carbohydrate resources and protein is

---

Fig. 6. Frass levels of glucose and its metabolite gluconate. *Spodoptera exigua* caterpillars were reared until pupation on chemically defined diets (nutrient concentration 42%: 22p:20c; nutrient concentration 66%: 42p:21c, 33p:30c, 21p:42c) containing glucose as the sole source of digestible carbohydrates. The digestible carbohydrate source in the food was glucose which can be metabolized by labial salivary glucose oxidase to gluconate. Total frass collected from these caterpillars were analyzed for (A) glucose and its metabolite (B) gluconate: bars represent the mean metabolite level (mg) ± S.E. for 9–21 individuals. Significant differences are indicated by different alphabetical lettering (Tukey post-hoc tests, \( p < 0.05 \)). (C) Frass metabolite levels: bars represent the mean frass egested ± S.E. for 15 caterpillars. Frass contains cellulose, glucose and gluconate and unaccounted for metabolites, which are presumably proteins and catabolic products. Significant differences in frass egested are indicated by different alphabetical lettering (Tukey post-hoc tests, \( p < 0.05 \)). Percentages represent the proportion of the frass composed of unaccounted, presumably nitrogen-based metabolites.
clearly limiting to phytophagous insects. Similarly, when feeding on most plant tissues, generalist caterpillars are faced with a C-biased diet and must weigh the detrimental effects of dietary C-excess with P-limitations (Merkx-Jacques et al., in press; Raubenheimer et al., 2005). When the caterpillar has ingested sufficient P, then salivary GOX might be produced and secreted to act either as a strategy to attenuate plant induced defenses or a pre-ingestive mechanism to cope with excess dietary C (Bede et al., 2007; Musser et al., 2002).

In many studies, caterpillars show better performance on heavily fertilized plants (Al-Zubaidi and Capinera, 1983; Goverde and Erhardt, 2003; Manuwoto and Scriber, 1985; Wheeler et al., 1998). Even though nitrogen-based plant defense might be strengthened on nitrogen-fertilized crops (Fritz et al., 2006), the increased quantity and quality of foliar protein levels often result in shorter developmental times, higher pupal weight and egg production (Throop et al., 2004). P-biased tissues, such as anthers or young photosynthetic leaves, or heavily nitrogen-fertilized plants, benefit the caterpillar herbivore by contributing the building blocks for growth, development and reproduction. As well, dietary P may influence salivary enzyme activity, producing GOX, which may function to regulate excess C and/or as a strategy to circumvent induced plant defenses (Musser et al., 2002, 2005).

Acknowledgments

We thank Jeremy McNeil and Steve Simpson for insightful comments on an earlier version of this manuscript. This research was funded through an NSERC operating grant and CFI infrastructure grant to J.C.B.

References


