Suitability of Black Soldier Fly Frass as Soil Amendment and Implication for Organic Waste Hygienization

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Abstract: Because of its nutritious properties, the black soldier fly has emerged as one of the most popular species in advancing circular economy through the re-valorization of anthropogenic organic wastes to insect biomass. Black soldier fly frass accumulates as a major by-product in artificial rearing set-ups and harbors great potential to complement or replace commercial fertilizers. We applied frass from larvae raised on different diets in nitrogen-equivalent amounts as soil amendment, comparing it to NH₄NO₃ fertilizer as a control. While the soil properties did not reveal any difference between mineral fertilizer and frass, principal component analysis showed significant differences that are mainly attributed to nitrate and dissolved nitrogen contents. We did not find significant differences in the growth of perennial ryegrass between the treatments, indicating that frass serves as a rapidly acting fertilizer comparable to NH₄NO₃. While the abundance of coliform bacteria increased during frass maturation, after application to the soil, they were outcompeted by gram-negatives. We thus conclude that frass may serve as a valuable fertilizer and does not impair the hygienic properties of soils.

Keywords: animal feedstuff; circular economy; fertilizer; greenhouse; insect larva; organic waste

1. Introduction

In recent years, the use of saprobic insect larvae from the mealworm beetle (Tenebrio molitor), the black soldier fly (Hermetia illucens; BSF), or the house fly (Musca domestica) has attracted interest in the face of rising prices of animal feedstuff and accumulating amounts of waste [1,2]. In the European Union, green waste and food waste largely contribute to an annual amount of 118 to 138 million tons of organic wastes [3]. Especially BSF larvae (BSFL) have been shown to efficiently convert organic wastes into high quality fat and protein [4]. The economic potential and meaningful re-introduction of otherwise wasted nutrients into the biosphere via a circular economy enticed researchers, investors, and the public to contribute to a more efficient recycling of organic wastes by exploiting the potential of insect larvae on a large scale [5,6]. BSFL could also play a valuable role for smaller decentralized waste management systems operated by e.g., hobbyists or farmers in areas where the fly occurs naturally [7–9]. Additionally, the exploitation of BSF and its by-products could create an affordable opportunity for revenue generation by entrepreneurs and smallholder farmers in low-income countries [9–11]. The main by-product in the bioconversion of wastes into high quality protein for animal feedstuff is summarized as ‘frass’. Frass in general describes insect excretions, but in a commercial context it often refers to a mixture of mainly insect feces, substrate residues, and shed
exoskeletons. It is an inevitable side-stream during the mass-rearing of insects that can add up to 75% of the fed substrate [12] and is often merchandised as a fertilizing product. In recent years, an increasing number of studies started focusing on meaningful applications of insect frass [9,13–15], and the first large-scale field studies provided promising perspectives for its application in agriculture, especially in terms of plant nutrient availability [10,11,16].

The substrate used to grow insects affects the properties of the frass, since undegradable residues remain unused, while the digested fraction is modified by the gut microbiota when passing through the gastrointestinal tract [17,18]. Wang et al. [19] used frass from T. molitor for subsequent rearing of BSFL to exploit leftover nutrients that T. molitor could not take up or digest. In substrates carrying a high bioburden like human feces and manure, BSFL have shown to reduce pathogenic bacteria such as Salmonella enterica [20,21] and Escherichia coli [20,22], which is attributed to their production of antimicrobial peptides [23]. In the wild, frass from various insects can help to increase the chances of survival and reproduction by either deterring [24,25] or attracting [26,27] conspecifics. Frass can act as a vector for phytopathogenic microorganisms [28,29] and as a source of probiotic yeasts [30]. Its effect on the insects’ environment can be observed in forests, where frass deposition goes hand in hand with insect canopy herbivory. It has been shown that frass has an impact on C and N dynamics, and has beneficial effects for tree growth by increasing soil total C, N, and NH₄⁺, as well as microbial soil respiration [31,32]. In industrial environments, frass pyrolyzed to biochar has been successfully tested as a biosorbent for wastewater detoxification [33]. According to recent studies, frass’ agriculturally and economically most meaningful potential could lie in its application as fertilizer [9,34].

In this study, we assessed the fertilizing potential of process residues (frass) from three generic diets degraded by BSFL. Two of the diets represent major streams of organic waste, namely grass-cuttings (GC) and fruit/vegetable (FV) mix, while the chicken feed (CF) control diet is a commonly used insect breeding substrate. We hypothesized that (I) microbial colonization increases with frass maturation and (II) frass may serve as a valuable alternative to mineral fertilizer by inducing beneficial effects on plant growth.

2. Materials and Methods

2.1. Black Soldier Fly Frass Collection

The frass was collected from a preparatory feeding experiment conducted at 27 °C, 60% relative humidity (Figures S1 and S2, Table 1). The chicken feed (CF; Grünes Legekorn Premium, Unser Lagerhaus, Klagenfurt, Austria) was processed with a Fidibus flour mill (Komo Mills, Hopfgarten, Austria) and mixed with water in a 40:60 ratio.

| Table 1. Feeding experiment termination summary. The feeding experiment was terminated after a total of 23 days when more than 90% larvae from one treatment group transitioned to prepupal stage. Different lower-case letters indicate differences between treatments (p < 0.05) according to the Tukey’s HSD test. (n = 4; average ± standard deviation; CF = Chicken feed diet, GC = Grass-cuttings diet, FV = Fruit/Vegetables diet). |
|-----------------|---------------|---------------|---------------|
|                 | CF            | GC            | FV            |
| Pupation rate [%] | 55.2 ± 5.3 b  | 98.7 ± 2.0 c  | 22.4 ± 2.0 a  |
| Prepupae fresh weight [mg] | 198 ± 10 | 167 ± 11 | 165 ± 31 |
| Prepupae dry weight [%] | 32.6 ± 1.9 | 29.6 ± 1.9 | 32.2 ± 9.3 |
| Prepupae water content [%] | 67.4 ± 3.7 | 70.4 ± 5.6 | 67.8 ± 9.9 |
| Prepupae organic content [%] | 27.6 ± 1.6 | 26.5 ± 1.8 | 31.4 ± 8.4 |
| Prepupae inorganic content [%] | 5.1 ± 0.3 c | 3.1 ± 0.2 b | 0.9 ± 0.9 a |
| Frass residues [%] | 43.7 ± 1.0 c | 46.0 ± 1.5 b | 28.5 ± 0.5 a |

The fruit/vegetable mix (FV; cucumber, tomato, apple, orange, in ratio 0.5:1:1:1) and fresh grass-cutting diet (GC) were shredded and homogenized using a Total Nutrition Center blender (Vitamix,
Olmsted Township, United States). Feeding was done in organic content-equivalents (100, 250, and 370 mg larvac⁻¹ day⁻¹ for CF, GC, and FV). After termination of the feeding experiment, the black soldier fly frass (BSFF) from each treatment was collected in plastic bags and stored at room temperature until further use. 2.2. Soil Preparation and Greenhouse Set-Up

A greenhouse trial using soil collected from an agricultural site (47°15′54″ N, 11°20′20″ E; Table 2) was set up to evaluate the fertilizing effect of the BSFF on the soil. The neutral-to-slightly basic soil (pH 7.3 ± 0.4) had an electrical conductivity of 78.0 ± 2.7 μS cm⁻¹ and a volatile solids content of 78.0 ± 26.6 g kg⁻¹. In addition to a Ptotal content of 823 ± 190 mg kg⁻¹ (Pₐvable proportion 6.88 ± 1.28 mg kg⁻¹), elemental analysis determined a C/N ratio of 24 (40 g Ctotal kg⁻¹, 1.7 g Ntotal kg⁻¹). The soil classified as a calcareous Fluvisol (IUSS Working Group WRB, 2015) was sieved (Ø < 4 mm) and homogenously mixed with a vermiculite/sand blend (1:1; v:v) at a ratio 2:1 w:w (soil:blend).

Table 2. Characterization of the soil used for the greenhouse trial. Values expressed on a dry mass basis for n = 3 (average ± standard deviation). pH (pH CaCl₂), EC (Electrical conductivity), VS (Volatile solids), Ctotal (Total carbon), Ntotal (Total nitrogen), Ptotal (Total phosphorus), Pav (Plant available P).

| Parameter | Value          |
|-----------|----------------|
| pH        | 7.3 ± 0.4      |
| EC [μS cm⁻¹] | 78.1 ± 2.7    |
| VS [g kg⁻¹] | 78.5 ± 26.6  |
| Ctotal [g kg⁻¹] | 40           |
| Ntotal [g kg⁻¹] | 1.7        |
| Ptotal [mg kg⁻¹] | 823 ± 190   |
| Pav [mg kg⁻¹] | 6.88 ± 1.28  |

The four experimental treatments were performed in 500 mL pots: soil was mixed with (1) mineral fertilizer (which served as control); (2) GC BSFF; (3) FV BSFF; (4) and CF BSFF. The mineral fertilizer (NH₄NO₃) and the different types of BSFF (Table 3) were added in an amount of 40 mg N kg⁻¹ soil, which is equivalent to 80 kg N ha⁻¹, considering the soil bulk density of 1 g cm⁻³ and a plough depth of 20 cm as described by Goberna et al. [35]. Thereby, all treatments received the same dose of total N.

Table 3. Main properties of the three different black soldier fly frass fractions (CF-F: Chicken feed frass; GC-F: Grass-cuttings frass; FV-F: Fruit/vegetables frass). Values expressed on a dry mass basis for n = 3 (average ± standard deviation). Different lower-case letters indicate differences between treatments (p ≤ 0.05) according to the Tukey’s HSD test. Different capital letters indicate significant differences between treatments (p ≤ 0.05) according to the Mann–Whitney test. EC (Electrical conductivity), VS (Volatile solids), Ctotal (Total carbon content), Ntotal (Total nitrogen content).

|        | CF-F            | GC-F           | FV-F           |
|--------|-----------------|----------------|----------------|
| pH     | 6.22 ± 0.14 C   | 5.40 ± 0.03 A  | 5.58 ± 0.01 B  |
| EC [mS cm⁻¹] | 5.67 ± 0.27 c  | 3.06 ± 0.03 b  | 2.36 ± 0.11 a  |
| Dry matter [%] | 90.9 ± 0.0     | 89.9 ± 0.0    | 90.4 ± 0.0     |
| Ctotal [g kg⁻¹] | 479 ± 8 B      | 443 ± 6 A     | 488 ± 4 B      |
| Ntotal [g kg⁻¹] | 25.9 ± 0.9 b   | 24.4 ± 0.2 b  | 18.3 ± 1.2 a   |
| C:N ratio | 18.5 ± 0.3 a   | 18.2 ± 0.4 a  | 26.6 ± 1.7 b   |
| VS [g kg⁻¹] | 910 ± 7 c       | 825 ± 9 a     | 873 ± 4 b      |

After an equilibration period of 16 h at 4 °C, pots were randomly placed in a greenhouse. Ryegrass (Lolium perenne; seed amount based on 30 kg seeds ha⁻¹) was sown and left to develop. During the incubation period of 28 days, at an average temperature of 20 °C with a light/darkness cycle of 10/14 h, the soil moisture was kept at field capacity (moisture of the soil after drainage by gravity). All treatments were applied in four replicates, resulting in a total of 16 pots in this study.
After the incubation period, plants were removed from the pots, and soil samples were sieved (Ø < 2 mm) and immediately stored at +4 °C until analyses (Table 4).

**Table 4.** Physicochemical and biological properties of the control (C-S: NH4NO3) and the frass amended soils (CF-S: Chicken feed frass + soil; GC-S: Grass-cuttings frass + soil and FV-S: Fruit/Vegetables frass + soil). Values expressed on a dry mass basis for n = 4 (average ± standard deviation). Different lower-case letters indicate differences between treatments (p ≤ 0.05) according to the Tukey’s HSD test. Different capital letters indicate significant differences between treatments (p ≤ 0.05) according to the Mann–Whitney test. EC (Electrical conductivity), VS (volatile solids), Cmic (Total carbon content), Ntot (Total nitrogen content), NH4+ (Ammonium content), NO3− (Nitrate content), DOC (Dissolved organic carbon), DC (Dissolved carbon), DN (Dissolved nitrogen), Ptot (Plant available phosphorous content), Pav (Total phosphorous content), BR (Basal respiration), qCO2: (Metabolic quotient).

|          | C-S       | CF-S       | GC-S       | FV-S       |
|----------|-----------|------------|------------|------------|
| pH CaCl₂ | 7.53 ± 0.02 a | 7.57 ± 0.01 ab | 7.58 ± 0.03 b | 7.58 ± 0.02 b |
| EC [μS cm⁻¹] | 95.5 ± 1.8 B  | 79.8 ± 6.4 A  | 77.3 ± 2.9 A  | 81.5 ± 7.8 A  |
| VS [g kg⁻¹] | 37.3 ± 1.1  | 35.4 ± 1.3  | 38.4 ± 2.0  | 37.8 ± 2.1  |
| Cmic [g kg⁻¹] | 17.9 ± 3.2  | 21.7 ± 4.7  | 22.5 ± 6.1  | 20.6 ± 5.2  |
| Ntot [g kg⁻¹] | 0.98 ± 0.35  | 0.99 ± 0.51  | 1.17 ± 0.39  | 0.98 ± 0.44  |
| C:N ratio | 20.3 ± 8.1  | 26.2 ± 11.7 | 21.0 ± 9.8  | 22.1 ± 9.0  |
| NH4+ [mg kg⁻¹] | 0.57 ± 0.13  | 0.58 ± 0.06  | 0.58 ± 0.08  | 0.61 ± 0.14  |
| NO3⁻ [mg kg⁻¹] | 45.2 ± 4.1 b | 15.4 ± 3.3 a | 17.0 ± 3.5 a | 12.1 ± 4.4 a |
| DOC [mg kg⁻¹] | 48.3 ± 3.8  | 50.2 ± 1.5  | 48.5 ± 1.9  | 51.8 ± 1.3  |
| DC [mg kg⁻¹] | 95.6 ± 1.6 a | 104.7 ± 1.4 b | 103.5 ± 4.0 b | 112.1 ± 2.0 c |
| DN [mg kg⁻¹] | 35.6 ± 3.9 C | 17.0 ± 1.5 B | 15.3 ± 1.8 AB | 15.0 ± 0.6 A |
| Ptot [mg kg⁻³] | 783 ± 46 ab  | 866 ± 35 b   | 757 ± 33 a   | 721 ± 45 a   |
| P bioavailability [%] | 67 ± 8     | 70 ± 12     | 81 ± 23     | 80 ± 12     |
| BR [μg CO₂ g⁻¹ dw h⁻¹] | 5.6 ± 0.15  | 4.6 ± 1.5   | 6.7 ± 0.7   | 5.6 ± 0.5   |
| Cmic [μg CO₂ g⁻¹ dw soil] | 416.1 ± 103.5 | 276.4 ± 38.7 | 279.0 ± 22.7 | 336.2 ± 16.1 |
| qCO₂ [μg CO₂·h⁻¹·μg⁻¹ C mic] | 14.7 ± 4.7  | 16.4 ± 4.3  | 24.5 ± 4.5  | 16.6 ± 1.4  |
| Plant biomass [mg dw] | 85 ± 7      | 80 ± 6      | 74 ± 4      | 75 ± 3      |

### 2.3. Frass and Soil Analyses

Frass and soil samples (10 g fresh weight) were placed into a glass Petri dish and oven-dried (105 °C) for 24 h to determine the content of total solids. Volatile organic solid (VS) content was determined from the weight loss following ignition in a muffle furnace (CWF 1000, Carbolite, Neuhausen, Germany) at 550 °C for 5 h. Total C and N contents were analyzed in dried samples using a CN analyzer (TruSpec CHN, LECO, St. Joseph, MI, USA). EC and pH were determined in distilled water and 0.01 M CaCl₂ extracts (1:2.5, w/v), respectively.

Soil inorganic nitrogen (NH4+ and NO3⁻) was determined in 0.0125 M CaCl₂ extracts as described by Kandeler [36,37]. Soil total P (Ptot) and plant available P (Pav) were determined as described by Illmer et al. [38]. To estimate dissolved organic carbon (DOC), dissolved carbon (DC), and dissolved nitrogen (DN), 10 g of field-moist soil were shaken in 40 mL distilled water, filtered, and immediately measured using a TOC-L analyzer (Shimadzu, Kyoto, Japan). Soil basal respiration (BR) and microbial biomass (Cmic) were measured according to Heinemeyer et al. [39]. The metabolic quotient (qCO2) was calculated from BR and Cmic according to Anderson and Domsch [40]. At the end of the trial, aboveground plant biomass was determined by cutting plant shoots at the soil surface and drying them at 60 °C for 48 h. Samples were then re-weighted to determine the dry biomass.
2.4. Preparation of Media

For the assessment of the total cultivable bacterial colony forming units (CFUs), we used standard methods agar (0.5% peptone, 0.25% yeast extract, 0.1% glucose, 1.5% agar, pH adjusted to neutral). To determine the abundance of Salmonella sp., E. coli, coliforms, and other gram-negative bacteria, XLT-4 and ChromoCult® coliform agar (Merck, Darmstadt, Germany) were prepared according to the enclosed recipe.

2.5. Pathogen Quantification/Assessment of Microbial Colonization in Frass and Soil

An amount of 2 g frass or soil sample was added to 18 mL sterile saline solution (0.95% NaCl) and placed on a rotation shaker at 200 rpm for 15 min. Samples were diluted to $10^{-2}$ and $10^{-3}$ for soil, and $10^{-5}$ and $10^{-6}$ for frass using sterile 0.95% NaCl. From each dilution, 50 µL was plated using the spread plate technique. Plates were then incubated at 37 °C for 24 h, and the CFUs were counted.

2.6. Statistical Analyses

The effect of the BSFL application on soil parameters was tested with a one-way analysis of variance (ANOVA). In case of significant F-values, a Tukey’s HSD (honestly significant difference) post hoc test ($p < 0.05$) was performed. Prior to analysis, the homogeneity of the variances was tested (Levene’s test), and data were also tested for normality. Non-normal data were subjected to non-parametric tests for several independent samples (Kruskal–Wallis test), and pairwise comparisons between treatments were performed using the Mann–Whitney U test ($p < 0.05$). Statistical analyses were performed using the SPSS v. 23.0 Software (IBM, Armonk, NY, USA). Principal component analysis was performed in R [41] using the vegan package [42]. Analysis of similarity (ANOSIM) on the physicochemical data (999 permutations) was also conducted with vegan. All graphical representations of data were created with ggplot2 [43].

3. Results and Discussion

3.1. Assessment of Microbial Load in Frass and Frass-Amended Soils

The high moisture content of substrates and air, as well as the pleasantly warm temperature common in insect breeding, favor microbial growth. While the type of diet is known to directly influence the BSFL gut microbiome [17,44], the excrements in turn may influence the microbiome in the frass. It is likely that by agitating and mixing their surrounding substrate with feces and their inherent microorganisms, the larvae have an impact on their habitat. Similar effects are known from the widely used earthworms (Eisenia fetida), which can stabilize organic wastes and introduce ammonia-oxidizing microorganisms, thereby boosting nitrification and increasing nitrate concentrations in the resulting vermicompost [45]. Other insect species inoculate the soil with excreted microorganisms and provide beneficial effects for its quality both in wild and artificial settings [46–48].

Before and after applying frass as soil amendment, the number of cultivable E. coli, coliform, and other gram-negative bacteria were assessed (Figure 1). While frass counted up to $10^9$ CFU g$^{-1}$, the count in soil was down to $10^3$–$10^5$ g$^{-1}$. With the nutrient media used in this study, untreated soil contained no cultivable E. coli or coliforms, and only low abundances of cultivable gram-negatives with $10^3$ CFUs g$^{-1}$. In particular, frass from the CF treatment acted as a reservoir for coliforms with a CFU count of $1.9 \times 10^9$, thereby exceeding CFU counts recorded on larval surfaces (Figure S2). Gram-negative bacteria predominated the cultivable microbiota in frass-amended soil with highest CFU counts of up to $10^9$ in soil treated with frass from a FV diet. High microbial load and dominance of coliforms in frass shifted to lower CFU numbers and predominantly gram-negative bacteria in the frass-soil mix, indicating that the autochthonous soil microbiota outcompeted allochthonous microorganisms introduced with frass [49–51].
3.2. Black Soldier Fly Frass Properties, Soil Quality and Plant Performance

The physicochemical properties of frass were influenced by the larval diet (Table 3). Especially CF frass was more alkaline, had a higher EC, and a higher content of VS. While total C contents were similar in all types of frass, FV frass showed a C:N-ratio of 26.6, compared to 18.5 and 18.2 in CF frass and GC frass, respectively. Similar C:N ratios as found in CF and GC frass have been reported by other studies that used brewery spent grains as larval substrate [11,16]. A C:N-ratio > 20 bears the risk of soil N immobilization, which may favor plants with a more efficient N exploitation attributed to their rhizobiome [46,52]. The addition of biochar to the larval waste conversion process might further improve the frass’ N retention, while at the same time increasing larval biomass yield [10]. Moreover, larvae pass through six instars continuously shedding their exoskeleton. Chitin, an N-acetylglucosamine-based polymer (C{sub 6}H{sub 13}O{sub 5}N){sub n}, may influence not only the C:N ratio, but its degradation product chitosan may also provide underrated benefits for plant health and pathogen resistance [53,54]. The C:N ratio is one of the major parameters to consider when it comes to deciding whether frass should be used as soil amendment or as co-substrate in anaerobic digestion or composting [55,56]. Chitin utilization by insects is often associated with chitinolytic gut symbionts [57], which still needs to be further investigated in the context of BSF larvae. Chitin-containing fertilizers have previously been found to serve as splendid nitrogen sources [58].

Frass addition to the soil before planting Lolium perenne was adjusted on a basis of N-equivalence (80 kg N ha{sup –1}; Tables 3 and 4). Soil amended with CF frass exhibited a higher P_{tot} content than the other frass-amended soils; however, P_{av} was not significantly different. Principal component analysis (Figure 2) highlighted the parameters that influenced the properties of the soil-frass mix the most, which was further confirmed by ANOSIM (R = 0.5061, p < 0.001). The three frass-amended soils clustered closely together, with P_{tot} and P_{av}, pH, DC, N_CB, C:N ratio, BR, and the qCO₂ being the most influential parameters for their similarity.
Figure 2. Principal component analysis of samples from control soil and soil mixed with the three different frass types. Data points represent replicates, and arrows show the most influential parameters for the spread of the data. \( \text{NO}_3 = \text{Nitrate}, \text{DN} = \text{Dissolved nitrogen}, \text{CN} = \text{Carbon/Nitrogen ratio}, \text{P}_{\text{bio}} = \text{Phosphorus bioavailability}, \text{BR} = \text{Soil basal respiration}, \text{qCO}_2 = \text{Metabolic quotient}, \text{C}_{\text{mic}} = \text{Microbial biomass}, \text{N}_{\text{tot}} = \text{Total nitrogen}, \text{P}_{\text{av}} = \text{Plant available phosphorus}, \text{DC} = \text{Dissolved carbon}, \text{EC} = \text{Electric conductivity}.

The \( \text{qCO}_2 \): describing the microbial soil respiration per unit \( \text{C}_{\text{mic}} \) is known to be tightly connected to the C:N ratio and increases when less N is available [59]. Higher \( \text{qCO}_2 \) can indicate stress or disturbances within the soil because, although C sources are readily available, microbial metabolism and substrate decomposition are limited by N [60]. \( \text{NO}_3 \) and \( \text{DN} \), on the other hand, were the major drivers for the deviation of the control group from the frass treatment groups, since they were both significantly higher in control soil.

In our study, the frass treatments were compared with a control that received an equivalent of 80 kg ha\(^{-1}\) nitrogen in the form of NH\(_4\)NO\(_3\). In a similar experiment, Ros et al. [61] found that such an amount of mineral N increased the maize yield by 33% compared with an unfertilized control, while N-equivalent additions of compost yielded only 15% increase. Recent observations at field-scale by Beesigamukama et al. showed that even at lower application rates of 30 kg N ha\(^{-1}\), BSFF exceeded the performance of mineral N fertilizer in terms of grain yield and nitrogen fertilizer replacement values when applied at the same rates [16]. Compared with commercial fertilizers, nitrogen recovery rates and nitrogen use efficiency of plants have been shown to be improved when amended with BSFF [11]. Additionally, the higher P concentrations in the frass could facilitate N accumulation in plants by improving N uptake, as P plays an important role in energy transfer [62,63].

Using BSFL instead of aerobic windrow composting has additionally been shown to reduce the global warming potential of treating organic wastes by 50% [64]. The addition of frass did not lead to significant differences in plant growth compared to the mineral fertilizer (Figure 3). In fact, the similar growth progress indicates that the nutrients from frass are readily available for uptake and have no
detrimental impact on plant growth. These results, however, do not support the findings of Alattar et al. [13], who reported that the development of plant height and leaves in corn (Zea mays) was inhibited by the addition of BSFL frass. In their study, they attributed the negative effects to the low porosity of larval residues that may have created anaerobic conditions. The moisture content of the frass harvested from our preliminary feeding experiment was only 10% (Table 3), thereby facilitating aeration and miscibility in soil. Insufficient oxygen supply can occur when frass has a high moisture content and is not subjected to adequate post-processing. In an environment specialized on insect rearing, a multi-step treatment of frass could increase the efficiency of degradation. With additional downstream composting or anaerobic digestion [65,66], the recovery as soil amendment represents the economically most promising option.

![Figure 3. Plant biomass yield of Lolium perenne after application of black soldier fly frass (BSFF) obtained from the degradation of various organic substrates. CF BSFF = Chicken feed frass, FV BSFF = Fruit/vegetables frass, GC BSFF = Grass-cuttings frass (n = 4).](image)

4. Conclusions

The valorization of organic wastes by insect larvae generates frass as a side-product. From our study we conclude that frass may serve as a soil nutrient source and does not impair soil hygiene. In some cases, however, frass post-processing through anaerobic digestion or composting may be advised to avoid soil nitrogen deficiencies or impairing soil gas permeability. In the light of the increasing importance of insect rearing, the agricultural utilization of frass is demanding further research, in particular, long-term studies.

Supplementary Materials: The following are available online at www.mdpi.com/2073-4395/10/10/1578/s1, Figure S1: Influence of three different diets on larval biomass increase, Figure S2: Microbial colonization of larval surfaces.

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