

1.1 Voorgenomen situatie

Een melding activiteitenbesluit wordt gedaan voor de dieren aantallen zoals opgenomen in Tabel 1.

• Tabel 1: Voorgenomen situatie

Rav code	Omschrijving conform Rav	Aantal dieren	mg NH ₃ / kg/j	NH ₃ totaal
Nieuwe stal				
	Meelwormen	5 ton*	-*	0.16
	Totaal			0,16

Maximaal zijn er op enig moment 5 ton aan meelwormen aanwezig op het bedrijf. Jaarlijks bedraagt dit circa 40 ton meelwormen.

Meelwormen zijn niet opgenomen in de RAV-lijst. Daarom is voor de ammoniakemissiefactor voor meelwormen aansluiting gezocht bij wetenschappelijk onderzoek (Oonincx et al., 2010). Uit het onderzoek van Oonincx blijkt dat meelwormen maximaal 0,09 mg NH₃/kg lichaamsgewicht per dag uitstoten. Dit komt neer op een ammoniak emissie van 32,85 mg NH₃/kg lichaamsgewicht per jaar. Op het bedrijf is op enig moment maximaal 5 ton meelwormen aanwezig. In de berekening wordt er vanuit gegaan dat er het hele jaar door 5 ton meelwormen aanwezig zijn.

5.000 kg meelwormen x 32,85 mg NH₃/kg lichaamsgewicht/jaar = 164.250 mg NH₃/jaar = 0,16 kg NH₃.

1.2 Vervoersbewegingen

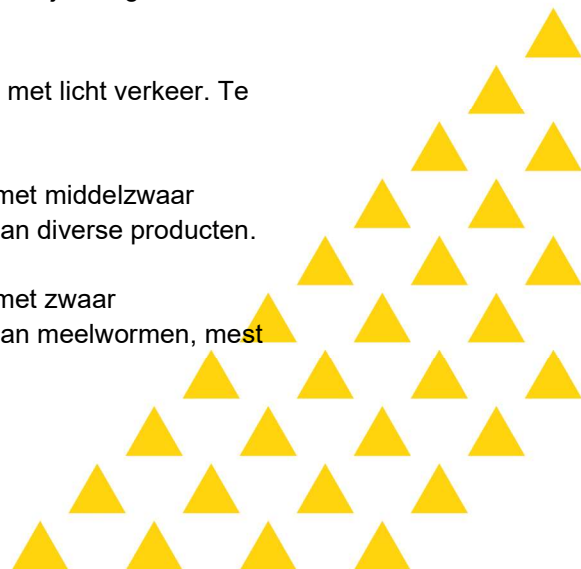
In de vergunde situatie is een akkerbouw bedrijf vergund waarbij de benodigde licht, middelzwaar en zware verkeersbewegingen komen kijken. In de beoogde situatie is een meelwormen kwekerij met een bezetting van 5 ton meelwormen vergund en de verhuur van een loods voor de opslag en het schoonmaken van bollen. De hoeveelheid vervoersbewegingen zijn afhankelijk van de gekozen bezetting bij minder tonnages meelwormen horen minder vervoersbewegingen.

Omdat de bezetting een dermate omvang heeft zal in praktijk weinig zwaar vrachtverkeer gebruikt worden voor het aan- en afvoeren van producten zoals voer, substraat en meelwormen. Onderstaande vervoersbewegingen zijn aangehouden als zijnde een worstcase benadering.

Op het bedrijf zijn gemiddeld 10 voertuigbewegingen per dag met licht verkeer. Te denken valt aan de adviseur, verkoper of overige bezoekers.

Op het bedrijf zijn gemiddeld 4 voertuigbewegingen per dag met middelzwaar vrachtverkeer. Te denken valt hierbij aan de aan- en afvoer van diverse producten.

Op het bedrijf zijn gemiddeld 4 voertuigbewegingen per dag met zwaar vrachtverkeer. Te denken valt hierbij aan de aan- en afvoer van meelwormen, mest en voer.



Projectberekening

Dit document geeft een overzicht van de invoer en rekenresultaten van een Projectberekening met AERIUS Calculator. De berekening is uitgevoerd binnen stikstofgevoelige Natura 2000-gebieden, op rekenpunten die overlappen met habitattypen en/of leefgebieden die aangewezen zijn in het kader van de Wet natuurbescherming, gekoppeld aan een aangewezen soort, of nog onbekend maar mogelijk wel relevant, en waar tevens sprake is van een overbelaste of bijna overbelaste situatie voor stikstof.



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Contactgegevens

Rechtspersoon
Inrichtingslocatie

Insectenhouderij Verkaik
IJzerpad 17,
8313 PJ Rutten

Activiteit

Omschrijving
Toelichting

Insectenhouderij Verkaik
Beoogde situatie

Berekening

AERIUS kenmerk
Datum berekening
Rekenconfiguratie

Rck1jgetFNhx
27 november 2023, 10:59
Wnb-rekengrid

Totale emissie

Situatie 1 - Beoogd

Rekenjaar	Emissie NH ₃	Emissie NO _x
2022	0,9 kg/j	23,8 kg/j

Resultaten

Situatie 1 - Beoogd
Gekarteerd oppervlak met toename (ha)
Gekarteerd oppervlak met afname (ha)
Grootste toename
Grootste afname

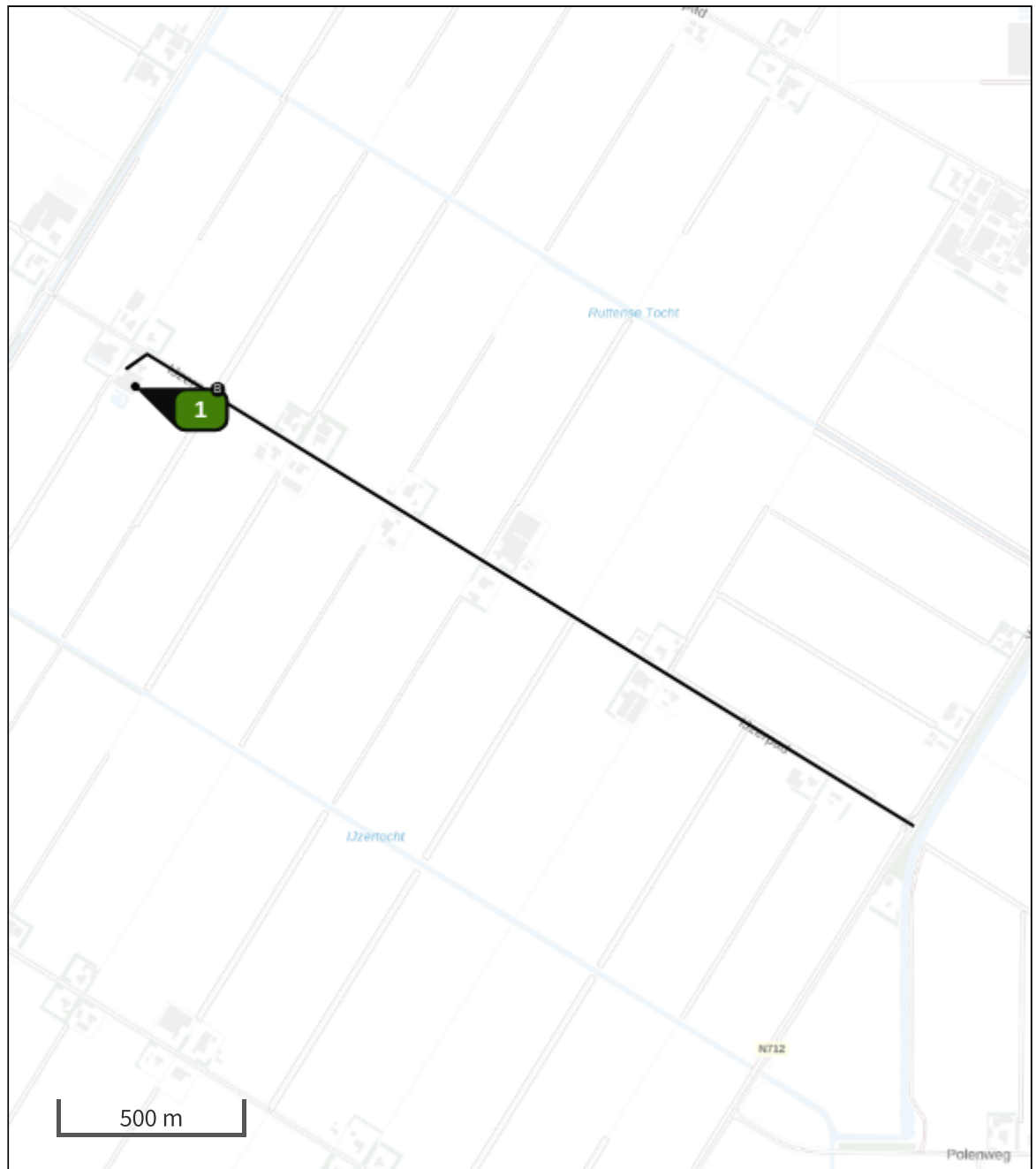
Hoogste bijdrage	Hexagon	Gebied
-		
-		
-		
-		
-		



Situatie 1 (Beoogd), rekenjaar 2022

Emissiebronnen		Emissie NH ₃	Emissie NO _x
1	Landbouw Stalemissies Insectenkwekerij	0,2 kg/j	-
	Verkeersnetwerk	0,7 kg/j	23,8 kg/j

Hoogste af- en toename op (bijna) overbelaste stikstofgevoelige Natura 2000 gebieden.



- | | |
|---|--|
|  Habitatrictlijn |  Grootste toename (projectberekening) |
|  Vogelrichtlijn |  Grootste afname (projectberekening) |
|  Vogelrichtlijn, Habitatrictlijn |  Hoogste totaal (achtergrond + projectberekening) |
|  Niet bepaald | |

De letters bij de bronlabels op de kaart geven bij welke type situaties de bronnen horen: beoogde situatie (B), referentiesituatie (R) en/of salderingssituatie (S).

Resultaten stikstofgevoelige Natura 2000 gebieden situatie "Situatie 1" (Beoogd) incl. saldering e/o referentie

	Berekend (ha gekarteed)	Hoogste totale depositie (mol N/ha/jr)	Met toename (ha gekarteed)	Grootste toename (mol N/ha/jr)	Met afname (ha gekarteed)	Grootste afname (mol N/ha/jr)
Totaal	-	-	-	-	-	-

Situatie 1, Rekenjaar 2022

1 Landbouw | Stalemissies

Naam	Insectenkwekerij	Uittreedhoogte	<u>5,0 m</u>	NH ₃	0,2 kg/j
Locatie	X:173058,36 Y:534968,02	Uittreeddiameter	0,5 m		
Wijze van ventilatie	Geforceerd	Temperatuur	<u>11,85 °C</u>		
Temporele variatie	Dierverblijven	Emissie			
		Uittreedrichting	Verticaal		
		Uittreedsnelheid	4,0 m/s		

Diersoort	RAV-code - Omschrijving	BWL-code	Aantal dieren	Stof	Emissiefactor (kg/dier/j)	Reductie	Emissie
	Meelwormen	-	1	NH ₃	0.16	-	0,2 kg/j

2 Wegverkeer | Weg

Naam	Vervoersbewegingen	Links	Rechts	NO _x	23,8 kg/j
Locatie	X:174117,18 Y:534426,43	Type scherm	-	NO ₂	5,4 kg/j
Lengte	2.548,40 m	Hoogte	-	NH ₃	0,7 kg/j
Wegtype	Buitenweg	Afstand tot de weg	-		
Rijrichting	Beide richtingen				
Tunnelfactor	1				
Type hoogteligging	Normaal				
Weghoogte t.o.v. maaiveld	0 m				

Verkeer	Max. snelheid	Aantal voertuigbewegingen	In file
Licht verkeer	Voorgeschreven factoren	10,0 /etmaal	0,0 %
Middelwaar vrachtverkeer	Voorgeschreven factoren	4,0 /etmaal	0,0 %
Zwaar vrachtverkeer	Voorgeschreven factoren	4,0 /etmaal	0,0 %
Busverkeer	Voorgeschreven factoren	0,0 /etmaal	0,0 %

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Rekenbasis

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Database versie 2023.0.1_3125d8b3c1_calculator_nl_stable

Voor meer informatie over de gebruikte methodiek en data zie:

<https://www.aerius.nl/>

An Exploration on Greenhouse Gas and Ammonia Production by Insect Species Suitable for Animal or Human Consumption

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Abstract

Background: Greenhouse gas (GHG) production, as a cause of climate change, is considered as one of the biggest problems society is currently facing. The livestock sector is one of the large contributors of anthropogenic GHG emissions. Also, large amounts of ammonia (NH₃), leading to soil nitrification and acidification, are produced by livestock. Therefore other sources of animal protein, like edible insects, are currently being considered.

Methodology/Principal Findings: An experiment was conducted to quantify production of carbon dioxide (CO₂) and average daily gain (ADG) as a measure of feed conversion efficiency, and to quantify the production of the greenhouse gases methane (CH₄) and nitrous oxide (N₂O) as well as NH₃ by five insect species of which the first three are considered edible: *Tenebrio molitor*, *Acheta domesticus*, *Locusta migratoria*, *Pachnoda marginata*, and *Blattella germanica*. Large differences were found among the species regarding their production of CO₂ and GHGs. The insects in this study had a higher relative growth rate and emitted comparable or lower amounts of GHG than described in literature for pigs and much lower amounts of GHG than cattle. The same was true for CO₂ production per kg of metabolic weight and per kg of mass gain. Furthermore, also the production of NH₃ by insects was lower than for conventional livestock.

Conclusions/Significance: This study therefore indicates that insects could serve as a more environmentally friendly alternative for the production of animal protein with respect to GHG and NH₃ emissions. The results of this study can be used as basic information to compare the production of insects with conventional livestock by means of a life cycle analysis.

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Introduction

Production of greenhouse gasses (GHG) is considered as an important cause of climate change [1,2]. The most important GHGs are carbon dioxide (CO₂), methane (CH₄) and nitrous oxide (N₂O). Since the end of the 18th century the atmospheric carbon-dioxide concentration has increased by 30% and CH₄ concentrations by 50% [3]. CH₄ and N₂O have considerably greater global warming potentials (GWPs) than CO₂. By assigning CO₂ a value of 1 GWP, the warming potentials of these other gases can be expressed on a CO₂-equivalent basis: CH₄ has a GWP of 25, and N₂O has a GWP of 298 [1]. The relative contribution of CO₂ equivalents (CO₂ eq.) of the livestock sector is large, amounting up to 18% of total anthropogenic GHG emissions [2]. Based on a Life Cycle Analysis (LCA) that takes the entire production process of animal products into account, the global contribution to GHG emissions by the animal sector are: 9% for CO₂ (fertilizer production for feed crops, on-farm energy expenditures, feed transport, animal product processing, animal

transport, and land use changes), 35–40% for CH₄ (enteric fermentation in ruminants and from farm animal manure) and 65% for N₂O (farm manure and urine) [2]. Direct CO₂ production through respiration is not relevant when determining the impact of GHGs as respiration by livestock is not considered a net source of CO₂ [2]. The respired carbon, which comes from the feed, was first taken up from CO₂ in the air and stored in an organic compound during the production of the feed. However, the ratio between body growth realised and CO₂ production is an indicator of feed conversion efficiency and thereby a relevant indicator for the environmental impact [4].

Livestock is also associated with environmental pollution due to ammonia (NH₃) emissions from manure and urine, leading to nitrification and acidification of soil [5]. Although not considered a GHG, NH₃ can indirectly contribute to N₂O emission [2], as conversion takes place by specialized soil bacteria [6]. Livestock is estimated to be responsible for 64% of all anthropogenic NH₃ emissions [2]. The main source of gaseous NH₃ is bacterial fermentation of uric acid in poultry manure [7,8] and bacterial

fermentation of urea in mammals [9]. Besides these environmental problems the livestock sector faces challenges regarding resistance to antibiotics, zoonosis and animal welfare [10].

All these problems together illustrate the need to find alternatives for conventional sources of animal protein. Mini-livestock, for instance edible insects, have been suggested as an alternative source of animal protein [11]. Production of animal protein in the form of edible insects supposedly has a lower environmental impact than conventional livestock [12,13,14]. When evaluating the total environmental impact of animal protein production, a LCA, in which all production factors are taken into account, is needed. Differences in environmental impact in a LCA can be explained mainly by three factors: enteric CH₄ emissions, feed conversion efficiencies and reproduction rates [4].

Before performing a LCA, it is necessary to know the GHG production by edible insects. This information is lacking in literature. Therefore, in this study we experimentally quantified the direct production of the GHGs CH₄ and N₂O for five insect species. CO₂ production and average daily gain (ADG) were quantified to provide an estimation of feed conversion efficiency. Additionally, NH₃ emissions were quantified. The results of this study represent a quantification of the insect physiological contribution to GHG production by insects and can in turn be used to create a LCA for insect-derived products.

Materials and Methods

2.1 Animals and housing

Five insect species were studied: fifth larval stage mealworms *Tenebrio molitor* L. (Coleoptera: Tenebrionidae), fifth and sixth nymphal stage house crickets *Acheta domesticus* (L.) (Orthoptera: Gryllidae), third and fourth stage nymphs of migratory locusts *Locusta migratoria* (L.) (Orthoptera: Acrididae), third larval stage sun beetles *Pachnoda marginata* Drury (Coleoptera: Scarabaeidae) and a mix of all stages of the Argentinean cockroach *Blaptica dubia* (Serville) (Dictyoptera: Blaberidae). Currently, *T. molitor*, *A. domesticus* and *L. migratoria* are considered edible, while *P. marginata* and *B. dubia* are not. The latter two species were included since they are a potential source of animal protein, for instance by means of protein extraction. These two species can be bred in large numbers with little time investment and are able to utilise a wide range of substrates as feed [15,16].

Per species three to six repetitions were conducted each for a period of three days. Animals were housed per species in two cages or containers per respiration chamber. These containers were placed in one of two, identical, open circuit climate respiration chambers measuring 80*50*45 cm, with a total volume of 265 L [17]. Within these climate respiration chambers, *T. molitor* and *P. marginata* were housed in two stacked plastic containers

(50*30*8.7 cm). The three other species were housed in metal wire cages (45*37.5*41 cm; mesh width 1 mm) with a glass cover plate. To increase surface area for *A. domesticus* and *B. dubia*, hollow plastic tubes (20 cm long and 3 cm in diameter), were stacked to a height of 30 cm in the wired cages, while for *L. migratoria*, two V-shaped-folded metal screens (70*15 cm) were entered per cage. Humidity, temperature, and day length were based on rearing conditions used by commercial insect rearing companies (Table 1). All animal masses reported are averages of fresh mass per cage. The starting and final animal mass per cage are provided in Table 1.

2.2 Diet

Food was provided for each species at the beginning of each repetition, except when mentioned otherwise.

Tenebrio molitor larvae were reared in 300 g mixed grain substrate (wheat, wheat bran, oats, soy, rye and corn, supplemented with beer yeast) with on top pieces of carrot ($\pm 15 \times 2$ cm) weighing a total average of 637 g per repetition.

Acheta domesticus was provided with chicken mash (501 g) with carrot pieces (784 g) on top for each repetition.

Locusta migratoria was provided with wheat bran (70 g; Arie Blok Animal Nutrition, Woerden, The Netherlands) in a metal bowl at the beginning of each repetition. Fresh Perennial ryegrass (*Lolium perenne*) was provided daily (463 g in three days). The grass was grown by Unifarm, Wageningen University and Research centre, Wageningen, The Netherlands.

P. marginata larvae were kept in a peat moss substrate (2.0 kg per respiration chamber) in which chicken mash (285 g) was mixed at the beginning of each three-day repetition. Pieces of carrot ($\pm 15 \times 2$ cm) with an average total mass of 161 g per repetition were put on top of the substrate.

B. dubia was provided with a chicken mash diet (199 g) and carrots (559 g), fresh carrot being added during the repetitions.

Peat moss, chicken mash, and carrots, offered to *A. domesticus*, *P. marginata* and *B. dubia* were provided by Kreca V.O.F, Ermelo, The Netherlands. The carrots and mixed grains substrate offered to *T. molitor* were provided by Insectra, Deurne, The Netherlands.

2.3 Gas measurements

During the experiment concentrations of CO₂ and CH₄ were measured every 9 min in the ingoing and outgoing air stream of the respiration chambers. The difference in CO₂ and CH₄ concentrations between ingoing and outgoing air thus represents the total production of CO₂ and CH₄ of insects, feed, and substrate. The exact air volumes were measured with a calibrated Schlumberger G1.6 dry gas meter and corrected for measured air temperature and pressure. CO₂ and CH₄ concentrations were

Table 1. Mean values and standard deviations of temperature, humidity, ventilation, hours of light per day and average start and final weight for five insect species.

	<i>Pachnoda. marginata</i>	<i>Tenebrio molitor</i>	<i>Blaptica dubia</i>	<i>Acheta domesticus</i>	<i>Locusta migratoria</i>
Temperature (°C)	28.0 $\pm$ 0	25.0 $\pm$ 0	28.0 $\pm$ 0	28.0 $\pm$ 0	32.0 $\pm$ 0
Humidity (%)	84.3 $\pm$ 3.3	79.8 $\pm$ 0.2	70.0 $\pm$ 0.0	69.9 $\pm$ 0.1	69.7 $\pm$ 0.2
Ventilation (L/min)	6.46 $\pm$ 2.06	6.82 $\pm$ 1.31	5.16 $\pm$ 0.05	11.18 $\pm$ 1.80	4.98 $\pm$ 0.39
Hours of light per day	0	0	12	12	12
Start weight (kg)	0.99	0.91	1.10	0.96	0.08
Final weight (kg)	1.10	1.10	1.28	1.17	0.13

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measured in dried gas. Gas was dried in a +2°C dew-point cooler. Nondispersive infrared analyzers were used to measure CO₂ (type Uras 3G, Hartmann and Braun, Frankfurt, Germany) and CH₄ (type Uras 10E, Hartmann and Braun, Frankfurt, Germany). The refreshed air volume was set so that CO₂ levels did not exceed 1%. From each climate respiration chamber, as well as from the incoming air, an air sample was taken for N₂O analysis after 24, 48, and 72 h with a 60 ml syringe. The syringes were sealed by a shutoff valve and stored at 20°C until analysis (within 48 h). The N₂O concentration was analysed by a gas chromatograph (CE instruments GC8000 Top, Interscience, Breda, The Netherlands) using a Haysep Q 80–100 mesh 2 m×1/8" SS column, at a constant temperature of 60°C. N₂O was detected with an electron capture detector (ECD). Injection volume was 5.0 ml in a fixed loop.

NH₃ concentrations in the climate respiration chambers were determined twice daily (at 12.00 and 24.00 h) by means of a gas detection tube system (Kitagawa, type AP-20; Komyo rikagaku kogyo, Tokyo, Japan; type 105 NH₃ gas detector tubes with a range of 1–20 ppm).

2.4 Calculations

Production of N₂O was calculated by subtracting the N₂O concentration from the incoming air from that in the outgoing air. These differences were then used in a formula adapted from Wheeler et al (2003) [18]:

ER = Emission rate of N₂O = $[N_2O] \text{ change (ppm} \times 10^{-6}) \times VV \text{ (m}^3/\text{day}) \times 44 \text{ (g/mol)} / 0.0224 \text{ (m}^3/\text{mol)}$, where VV = ventilation volume of air in a specified time period. The average concentration difference of the three samples taken during the three-day period was used to determine the average N₂O production in a repetition.

The formula used by Wheeler (2003) was also used for the calculation of NH₃ production. A molecular mass of 17 was used and instead of a difference in concentration, the measured concentration was used, leading to a slight overestimation of the actual NH₃ production (between 0 and 0.1 mg/kg BM/day).

CO₂ equivalents were calculated by adding the multiplications of the produced amounts of CH₄ and N₂O with their global warming potential; 25 for CH₄, and 298 for N₂O [1].

Mean body mass was calculated by averaging the body mass at the start of the experiment and the body mass at the end of the experiment. Average daily gain (ADG) was calculated as follows:

$((\text{End mass} - \text{Start mass}) / \text{Start mass}) / 3 \times 100\%$, in which 3 is the number of days the experiment was running.

The ratio between CO₂ production per unit biomass per day and ADG gives an indication of the feed conversion efficiency, in which higher values indicate lower efficiencies.

To determine CO₂ production from feed and substrate, all feeds were independently tested in the same respiration chambers, without the animals. A linear time course of consumption was assumed and CO₂ production was recalculated to kg of live insect.

2.5 Statistics

The N₂O and NH₃ assay data were subjected to a two-way analysis of variance (ANOVA) with species and time of sampling (24, 48, or 72 h) as fixed factors to determine whether the time of sampling had an effect. No significant effect of the time of sampling was found for N₂O (Pillai's trace: $F = 1.467$, $P = 0.199$). Therefore, the average of the three samples taken during the 3-day trial period was used to determine the change per repetition and to calculate total production. However, NH₃ production was significantly affected by the time of sampling (day or night; Pillai's trace: $F = 4.065$, $P = 0.019$) and the day of the repetition (first, second or third; Pillai's trace: $F = 17.170$, $P < 0.001$). CO₂ and CH₄ production for all five species were analyzed by means of a one way analysis of variance (ANOVA) followed by a Tukey post hoc test. Statistical analysis of all data was done by means of SPSS 15.0.

Results

Production of CO₂ is expressed per kilogram of mean live body mass (BM) per day (24 hours) and per kilogram of mass gain (Table 2) and the average daily gain (ADG) is reported (Table 2). Production of CH₄, N₂O, CO₂ equivalents, and NH₃, are expressed per kilogram of mean live body mass (BM) per day (Table 3) and per kilogram of mass gain (Table 4).

3.1 ADG and CO₂ production

ADG varied between 4.0% (*P. marginata*) and 19.6% (*L. migratoria*) with the three other species having an ADG of 6–7%. CO₂ production among the five insect species differed significantly and ranged from 19 (*B. dubia*) to 110 (*L. migratoria*) g per kg BM/day. Also, the CO₂ production per kg of metabolic weight (i.e. the weight of metabolically active body tissue) differed greatly between

Table 2. CO₂ production (average ± standard deviation) per kilogram of bodymass per day, per kg of mass gain and average daily gain for five insect species, pigs and beef cattle.

Species	CO ₂ (g/kg BM/day)	CO ₂ (g/kg mass gain)	ADG (%)
<i>Pachnoda marginata</i> (n = 4)	50 ± 22 ^a	1,539 ± 518 ^a	4.0 ± 2.1% ^a
<i>Tenebrio molitor</i> (n = 4)	61 ± 9 ^b	1,031 ± 349 ^b	7.3 ± 2.5% ^b
<i>Blaptica dubia</i> (n = 3)	19 ± 3 ^c	337 ± 51 ^c	6.1 ± 0.7% ^c
<i>Acheta domesticus</i> (n = 4)	68 ± 10 ^d	1,468 ± 971 ^a	7.2 ± 3.4% ^b
<i>Locusta migratoria</i> (n = 6)	110 ± 21 ^e	734 ± 119 ^d	19.6 ± 2.1% ^d
Pigs	21.6–29.6	865–1,194	3.2 ± 0.53%
Beef cattle	5.3–7.0	2,835	0.3 ± 0.07%

BM = Body Mass;

ADG = Average daily gain;

Reported values for pigs and beef cattle were obtained from: [5] Aarnink et al., 1995; [49] Groot Koerkamp et al., 1998; [52] Demmers et al., 2001; [50] Nicks et al., 2003; [59] Beauchemin & McGinn, 2005; [48] Cabaraux et al., 2009 and [53] Harper et al., 2009. Mean values bearing different superscripts in a column differ significantly ($P < 0.05$).

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Table 3. CH₄, N₂O, CO₂ eq. and NH₃ production (average $\pm$ standard deviation) per kilogram of bodymass per day for five insect species, pigs and beef cattle.

Species	CH ₄ (g/kg BM/day)	N ₂ O (mg/kg BM/day)	CO ₂ eq. (g/kg BM/day)	NH ₃ (mg/kg BM/day)
<i>Pachnoda marginata</i> (n = 4)	0.16 $\pm$ 0.085 ^a	0.0 $\pm$ 0.03 ^a	4.00 $\pm$ 2.13 ^a	0.1 $\pm$ 0.16 ^a
<i>Tenebrio molitor</i> (n = 4)	0.00 $\pm$ 0.002 ^b	1.5 $\pm$ 0.13 ^b	0.45 $\pm$ 0.04 ^b	0.0 $\pm$ 0.09 ^a
<i>Blaptica dubia</i> (n = 3)	0.08 $\pm$ 0.021 ^c	0.3 $\pm$ 0.24 ^a	2.12 $\pm$ 0.57 ^c	3.0 $\pm$ 1.63 ^b
<i>Acheta domesticus</i> (n = 4)	0.00 $\pm$ 0.002 ^c	0.1 $\pm$ 0.13 ^a	0.05 $\pm$ 0.04 ^b	5.4 $\pm$ 3.40 ^c
<i>Locusta migratoria</i> (n = 6)	0.00 $\pm$ 0.017 ^c	8.0 $\pm$ 13.50 ^b	2.37 $\pm$ 4.02 ^c	5.4 $\pm$ 1.65 ^c
Pigs	0.049–0.098	2.7–85.6	2.03–27.96	4.8–75
Beef cattle	0.239–0.283	N/A	5.98–7.08	14–170

BM = Body Mass;

N/A = Not Available;

Reported values for pigs and beef cattle were obtained from: [5] Aarnink et al., 1995; [49] Groot Koerkamp et al., 1998; [52] Demmers et al., 2001; [50] Nicks et al., 2003; [59] Beauchemin & McGinn, 2005; [48] Cabaraux et al., 2009 and [53] Harper et al., 2009. Mean values bearing different superscripts in a column differ significantly ($P < 0.05$).

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species (Table 5). CO₂ production expressed per kg of mass gain was intermediary for *L. migratoria* due to the high ADG. Still, the CO₂ production of *L. migratoria* per kg of mass gain was more than double the production of CO₂ by *B. dubia*. *Pachnoda marginata* had the highest production of CO₂ per kg of mass gain (1,539 g/kg), which was more than double the amount of *L. migratoria*.

3.2 CH₄

Production of methane was detected for *P. marginata* and *B. dubia*, but not for the three other species. *Pachnoda marginata* produced more than three times as much CH₄ per kg of mass gain than *B. dubia* (4.9 vs 1.4 g). This difference was caused by a higher production of CH₄ per kg BM (0.16 g vs 0.08 g) and a lower ADG (4.0% vs 6.1%).

3.3 N₂O

N₂O was produced only in significant amounts by *T. molitor* and *L. migratoria* (1.5 and 8.0 mg/kg BM/day, respectively). Production of N₂O by *L. migratoria* per kg BM was more than 5-fold the production by *T. molitor*, this difference decreased to almost 2.5-fold when expressed per kg of mass gain, due to a much higher ADG of *L. migratoria*.

3.4 NH₃

NH₃ was produced by *A. domesticus*, *L. migratoria*, and *B. dubia* (3.0–5.4 mg/kg BM/day), and ranged from 36–142 mg/kg of mass gain (Table 3 and 4). Significant differences (Pillai's trace: $F = 4.065$, $P = 0.019$) between daytime (12.00) and night-time (24.00) NH₃ emission levels were found for *A. domesticus* (6.4 and 4.4 mg/kg BM/day), *L. migratoria* (5.6 and 3.9 mg/kg BM/day), and *B. dubia* (3.4 and 2.6 mg/kg BM/day).

Discussion

Insects, being poikilotherms, do not use their metabolism to maintain a body temperature within narrow ranges, contrary to homeothermic animals. This is expected to result in higher feed conversion efficiencies. CO₂ production related to growth, has an inverse relationship with feed conversion efficiency in a given situation. CO₂ production by insects depends on the species, stage of development [19,20], temperature [21], feeding status [22], and on activity level [23,24]. A production of 37 g CO₂/kg BM/day was reported for *Anabrus simplex* (Orthoptera, Tettigoniidae), 40 g CO₂/kg BM/day for the locust *Schistocerca americana* (Orthoptera; Acrididae) [25] and 94 g/kg BM/day for adult *Tribolium castaneum* (Coleoptera; Tenebrionidae) [26]. All five species in the current

Table 4. CH₄, N₂O, CO₂ eq. and NH₃ production (average $\pm$ standard deviation) per kilogram of mass gain for five insect species, pigs and beef cattle.

Species	CH ₄ (g/kg mass gain)	N ₂ O (mg/kg mass gain)	CO ₂ eq. (g/kg mass gain)	NH ₃ (mg/day/kg mass gain)
<i>Pachnoda marginata</i> (n = 4)	4.9 $\pm$ 1.96 ^a	1.03 $\pm$ 1.06 ^a	121.86 $\pm$ 49.09 ^a	3 $\pm$ 4.8 ^a
<i>Tenebrio molitor</i> (n = 4)	0.1 $\pm$ 0.03 ^b	25.5 $\pm$ 7.70 ^b	7.58 $\pm$ 2.29 ^b	1 $\pm$ 2.0 ^a
<i>Blaptica dubia</i> (n = 3)	1.4 $\pm$ 0.30 ^c	5.7 $\pm$ 4.05 ^a	37.54 $\pm$ 8.01 ^c	54 $\pm$ 31.1 ^a
<i>Acheta domesticus</i> (n = 4)	0.0 $\pm$ 0.09 ^b	5.3 $\pm$ 6.05 ^a	1.57 $\pm$ 1.80 ^d	142 $\pm$ 184.5 ^b
<i>Locusta migratoria</i> (n = 6)	0.0 $\pm$ 0.11 ^b	59.5 $\pm$ 104.8 ^c	17.72 $\pm$ 31.22 ^e	36 $\pm$ 10.8 ^a
Pigs	1.92–3.98	106–3457	79.59–1,130	1140–1920
Beef cattle	114	N/A	2,850	N/A

BM = Body Mass;

N/A = Not Available;

Reported values for pigs and beef cattle were obtained from: [5] Aarnink et al., 1995; [49] Groot Koerkamp et al., 1998; [52] Demmers et al., 2001; [50] Nicks et al., 2003; [59] Beauchemin & McGinn, 2005; [48] Cabaraux et al., 2009 and [53] Harper et al., 2009. Mean values bearing different superscripts in a column differ significantly ($P < 0.05$).

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Table 5. CO₂ production (g) per kilogram of metabolic weight per day for five insect species, pigs and beef cattle based on Kleiber's law ($B = aM^b$).

Species	b = 0.67	b = 0.75	b = 0.82
<i>Pachnoda marginata</i> (n = 4)	7	11	17
<i>Tenebrio molitor</i> (n = 4)	3	7	12
<i>Blaptica dubia</i> (n = 3)	2	4	6
<i>Acheta domesticus</i> (n = 4)	4	8	14
<i>Locusta migratoria</i> (n = 6)	9	17	29
Pigs	63	50	41
Beef cattle	50	31	21

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study had a fairly high production of CO₂. This might to a large extent be explained by *ad libitum* feeding during the experiment that has been reported to increase oxygen consumption fivefold [22]. Reported CO₂ production for inactive, unfed, Tenebrionid adults ranged between 5.4–13.3 g/kg BM/day [27], which is 5–10 times lower than observed for *T. molitor* in this experiment. This can partially be explained by the locomotory activities of *T. molitor* larvae in this experiment [37]. Furthermore, growing larvae are expected to have a higher CO₂ production than adults. The range of CO₂ production for *T. molitor* is comparable to the factorial metabolic scope reported for tiger beetles (*Cicindela* spp: Coleoptera; Cicindelidae) of 6.1–16.5 [28].

Size differences in animals account for a difference in metabolic rate, and thereby CO₂ production. The relation between metabolic rate (B) and body mass (M) was described by Kleiber [29] as $B = aM^b$, in which a is a constant and b = 0.75. The value of b has been much debated since [30,31,32]. For poikilotherms values between 0.67 and 1.0 have been reported and a comparison of several arthropod species suggested b approximates 0.82 [33,34]. The value chosen for b has a large impact on the metabolic weight and thereby the calculated CO₂ production (Table 5). Applying b = 0.75 for pigs and beef cattle and b = 0.82 for insects, resulted in a lower CO₂ production based on metabolic weight for the studied insect species (Table 5). For *L. migratoria* CO₂ production was only slightly lower than for beef cattle, however, for the other four species production was between 18% and 54% of that for beef cattle and between 11% and 34% of the CO₂ production of pigs.

The CO₂ production per kg BM of insect species investigated in this study was higher than for pigs or cattle (Table 3). This concurs with Prothero *et al.* (1979) [35], who reported a higher oxygen consumption per kg of BM for insects than for mammals, assuming the respiratory quotient (CO₂ production/O₂ consumption) has similar values (0.7–1.0) for both animal groups. However, the CO₂ production per kg of mass gain for the five insect species in the current study (337–1,539 g/kg) was either 39% (minimum values) or 129% (maximum values) when compared with pigs (865–1,194 g/kg) and much lower (12%–54% respectively) than cattle (2,835 g/kg). Therefore, CO₂ production per kg of mass gain suggests higher feed conversion efficiencies for insects than for mammalian livestock. These results concur with those of other authors [13,14,36,37].

A similar trend was visible for ADG; the ADG for the five insect species studied was 4.0–19.6%, the minimum value of this range being close to the 3.2% reported for pigs, whereas the maximum value was 6 times higher. Compared to cattle (0.3%), insect ADG values were much higher. In general, the rate of ADG depends,

amongst others, on life phase. Therefore, where available, literature data on growing animals were used. The fundamental biological differences in growth and development processes between pigs and cattle and the studied insects impeded further synchronization.

CH₄ production for the species studied was in agreement with Hackstein and Stumm (1994) [38]; for insects, only representatives of cockroaches, termites, and scarab beetles produce CH₄. This originates from bacterial fermentation by methanobacteriaceae in the hindgut [39].

We found large variability for the N₂O emission rates. Earlier studies in laying hens using a similar method for determining N₂O production, concluded that production was either negligible or undetectable [7,40]. However, other authors [41,42] determined a production of 28 mg N₂O/kg BM/day and 52 mg N₂O/kg BM/day, respectively, indicating the difficulty of accurately determining N₂O production [43].

In earlier studies respiration of feed was considered to have a negligible effect on utilisation of dry mass as determined gravimetrically [44] and therefore on CO₂ production. Later studies suggested that respiration by plant leaves can be an important source of error in the calculation of insect feed intake using gravimetric methods [45] and can cause major errors in energy budget studies of plant-feeding insects [46]. Our reported CO₂ production includes the respiration of the feed (Table 6). The extremely high contribution to total CO₂ production by the substrate of *P. marginata* (92.5%) was most likely due to large amounts of fungal biomass observed in the mixed feed and substrate when insects were absent in the experiments aimed to obtain correction values for CO₂-production by the substrate. No fungal growth was apparent during the experiments on feeding *P. marginata* larvae, suggesting that the contribution of the substrate to total respiration during the experiment was much lower. We conclude that the interaction between actively feeding *P. marginata* larvae and the substrate suppressed fungal growth through either consumption by the beetle larvae [47] of fungal biomass or through unknown chemical or combined chemical/mechanical mechanisms. Such interactions hinder the application of realistic corrections for the contribution of feed and substrate to the total CO₂ production and thus to quantify the CO₂ production arising from insect metabolism separately.

For all other species the relative contribution of the feed to total CO₂ production was minor, varying between 1.3% and 3.6%. Although feed respiration did have an impact on production of CO₂, still the production of CO₂ is much higher for *L. migratoria* than for the other insect species. A likely explanation for this higher production of CO₂ is the 7°C higher temperature *L. migratoria* was kept at, as a difference of 10°C is expected to double CO₂ production. Furthermore, the comparatively high ADG of *L. migratoria* is expected to result in higher production of CO₂.

In one of the repetitions for *A. domesticus*, a lower ADG and increased mortality were observed. Excluding this repetition, the emission of CO₂ per kg BM decreased slightly (68 vs 71 g/kg), but the emission of CO₂ per kg mass gain changed considerably (918 vs 1468 g/kg). This difference can for a large part be explained by a decrease in ADG (from 9.0 to 7.2%). *Acheta domesticus* did not produce CH₄, but N₂O production doubled (from 0.1 to 0.2 mg/kg BM; 1.9 vs 5.3 mg/kg mass gain). The production of CO₂ eq. also increased (0.04 vs 0.05 g CO₂ eq. /kg BM and 0.57 vs 1.57 g/kg mass gain). It is well possible that the higher N₂O production measured was caused by saprophytic bacteria utilising the dead *A. domesticus* and producing N₂O [6]. Although we included this repetition in the results, it is not clear whether this represents the practical situation best.

Table 6. Calculated CO₂ production of provided feed for five insect species recalculated per kg of animal body mass.

Species	CO ₂ production (g)/kg BM of insect	Relative contribution
<i>Pachnoda marginata</i>	46.2	92.46%
<i>Tenebrio molitor</i>	2.2	3.58%
<i>Blaptica dubia</i>	0.4	2.31%
<i>Acheta domesticus</i>	0.9	1.34%
<i>Locusta migratoria</i>	3.3	3.04%

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Large differences in NH₃ emission have been reported for conventional livestock. Pigs for example emit 4.8–75 mg/kg BM/day [48,49,50], poultry 72–436 mg/kg BM/day [41,49,51] and cattle 14–170 mg/kg BM/day [49,52,53]. Several factors influence NH₃ emission, such as temperature, relative humidity, food type, moisture content, pH, wind speed, housing type, and substrate [54,55].

In the current experiment, a clear NH₃ emission pattern was found; higher amounts of NH₃ were emitted during daytime for *A. domesticus*, *L. migratoria* and *B. dubia*, than during nighttime. Day-night rhythms for NH₃ excretion have been documented for pigs [5] and are strongly correlated with activity levels [56]. Quantitatively the differences between day and night emission levels are small; 7–10% with a maximum difference of 25% [5]. In our study this relative difference was approximately 33%. In all cases NH₃ emission levels were higher during the daytime than during the night-time. For *L. migratoria* this is the active period, for the nocturnal *B. dubia* and *A. domesticus* it is not, indicating that a different, unknown variable might influence NH₃ emission patterns in these insects.

NH₃ concentrations in the outgoing air, and consequently calculated NH₃ emission, increased from day one to day three in *B. dubia* (1.57 to 4.29 mg/kg BM/day) and *A. domesticus* (2.46 to 8.01 mg/kg BM/day). This could indicate that NH₃ emissions might be underestimated due to the relatively short time frame of our experiments. For *L. migratoria* NH₃ emission did not increase between day 1 and day 3 (5.57 and 5.05 mg/kg BM/day), suggesting that NH₃ production was stable. This might be caused by the faeces of this species that, contrary to those of *B. dubia* or *A. domesticus*, dry quickly after defecation.

We conclude that *P. marginata* and *T. molitor* probably did not emit NH₃. Poultry deep litter systems [57] have higher NH₃ emission rates than battery systems [55], which is explained by the presence of substrate.

The presence of substrates for *P. marginata* and *T. molitor* in this study corresponded with lower NH₃ emissions. A possible explanation is that gas exchange in the container is inhibited by

the substrate and therefore less emission of NH₃ was measured. However, it could also be that these species produce less NH₃.

All insect species in this study produced much lower amounts of NH₃ (3.0 to 5.4 mg/kg BM/day for *A. domesticus*, *L. migratoria* and *B. dubia*) than conventional livestock (4.8–75 mg/kg BM/day for pigs and 14–170 mg/kg BM/day for cattle). Further research is needed to determine for which insect species and to what extent NH₃ emissions increase further when a longer time frame is used.

Conclusions

To the authors' knowledge, the study presented here is the first to report on both GHG and NH₃ emissions of edible insect species. An evaluation of the GHG emissions of edible insect species is most relevant when based on CO₂ eq. per kg of mass gain. In that way a comparison of the selected species with each other and with conventional livestock is based on a cost-benefit principle, in which the GHG production (environmental cost) is directly linked to food production (benefit). GHG emission of four of the five insect species studied was much lower than documented for pigs when expressed per kg of mass gain and only around 1% of the GHG emission for ruminants.

The measured NH₃ emission levels of all insect species in this experiment were lower than reported NH₃ emission levels for conventional livestock.

The ADG of all insect species in this study was higher than for conventional livestock, while CO₂ production expressed as g/kg mass gain was comparable or lower, which indicates higher feed conversion efficiencies for insects.

This study therefore indicates that insects could serve as a more environmentally friendly alternative for the production of animal protein from the perspective of GHG and NH₃ emissions. A complete lifecycle analysis for species of edible insects is lacking at this point in time [58] and should be the focus point of further studies to allow a conclusive evaluation of the sustainability of insects as a protein-rich food source. The data presented in this study are indispensable for conducting a lifecycle analysis for edible insects.

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Author Contributions

Conceived and designed the experiments: DO HvdB JvL AvH. Performed the experiments: DO JvL. Analyzed the data: DO JvL MJWH. Contributed reagents/materials/analysis tools: MJWH. Wrote the paper: DO HvdB JvL AvH.

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An exploration on Greenhouse Gas and Ammonia Production by Insect Species Suitable for Animal or Human Consumption

Een verkenning van broeikasgassen en ammoniak productie door insecten die geschikt zijn voor dierlijk of menselijke consumptie

Dennis Oonincx

Samenvatting

Achtergrond: Broeikasgas productie, als oorzaak van klimaat verandering, wordt gezien als een van de grootste problemen waarmee de maatschappij op dit moment mee wordt geconfronteerd. De veehouderij is een van de grootste veroorzakers van de menselijke broeikasgas emissie. Daarnaast worden ook grote hoeveelheden ammoniak, wat leidt tot bodem nitrificatie en verzuring, geproduceerd door de veestapel. Hierdoor worden momenteel andere bronnen van dierlijke eiwitten, zoals eetbare insecten, onderzocht.

Voornaamste bevindingen: Er is een experiment uitgevoerd om te kwalificeren wat de productie van CO₂ is van de insecten. Daarnaast zijn de productie van methaan, stikstofdioxide (N₂O) en ammoniak (NH₃) gemeten. Dit is in absolute getallen uitgedrukt en weergegeven ten opzichte van de gemiddelde dagelijkse groei van de insecten. Voor het onderzoek zijn er naar 5 insecten onderzocht, waarbij 3 eetbaar zijn, dit zijn de Tenebrio molitor (meelworm), Acheta Domesticus (huiskrekel), Locusta migratoria (Europese treksprinkhaan), Pachnoda marginata (Kongo-rozenkever) en Blaptica dubia (Argentijnse Kakkerlak). Onderling zijn er grote verschillen gevonden tussen de soorten insecten gekeken naar de uitstoot van broeikasgassen. De insecten in deze studie hebben een hogere relatieve groei, en een vergelijkbare of lagere uitstoot van broeikasgassen dan varkens en een veel lagere uitstoot van broeikasgassen in vergelijking met koeien. Dit was ook het geval voor de CO₂ productie per kg metabool gewicht of per kg massa toename. Daarnaast was ook de productie van ammoniak bij insecten lager dan dat van conventioneel vee.

Conclusie: Deze studie laat zien dat insecten zouden kunnen dienen als meer milieu vriendelijk alternatief voor de productie van dierlijk eiwit, rekening houdend met de broeikasgas uitstoot en de ammoniak uitstoot. De resultaten van deze studie kunnen gebruikt worden als basis informatie om de vergelijking te maken tussen productie van insecten in vergelijking met de gangbare veehouderij.

Introductie

De productie van broeikasgassen wordt gezien als een belangrijke oorzaak van klimaat veranderingen. De belangrijkste broeikasgassen zijn CO₂, methaan en stikstofdioxide (N₂O). De relatieve bijdrage van CO₂ equivalenten van de veehouderij sector is groot, dit loopt op tot 18% van de totale menselijke broeikasgasemissie. Op basis van een levenscyclus analyse, dat de gehele productieproces van dierlijke producten beslaat, wordt 9% van de wereldwijde CO₂, 35-40% van de wereldwijde methaan uitstoot en 65% van de wereldwijde N₂O uitstoot veroorzaakt door de veehouderij.

Vee wordt in verband gebracht met milieuvervuiling door ammoniak emissie van mest en urine. Dit leidt tot nitrificatie en verzuring van de bodem. Al hoewel ammoniak niet wordt gezien als een broeikasgas draagt het wel degelijk bij aan de N₂O emissie, dit komt doordat bodembacteriën het ammoniak omzetten naar N₂O. De veestapel is verantwoordelijk voor 64% van alle menselijke

ammoniak emissie. De voornaamste oorzaak hiervan is bacteriële fermentatie van urinezuur in pluimveemest en bacteriële fermentatie van ureum in zoogdierenmest. Naast milieu problematiek spelen meerder factoren zoals antibiotica resistentie, zoönose en dierenwelzijn.

Al deze problemen samen illustreren de noodzaak om alternatieven te vinden voor de bestaande dierlijke eiwitten. Mini-livestock, zoals eetbare insecten, zijn voorgesteld als alternatieve bron voor dierlijk eiwit. De productie van dierlijk eiwit in de vorm van eetbare insecten hebben waarschijnlijk een lagere milieu impact dan traditioneel vee. Wanneer er naar de totale milieu impact van dierlijke eiwitten wordt geëvalueerd is er een levenscyclus analyse nodig, hierbij wordt er met alle productiefactoren rekening gehouden. Verschillen in milieubelasting in een levenscyclus analyse worden voornamelijk toegeschreven aan drie factoren: spijsvertering methaan emissie, voederconversie en voortplantingssnelheid.

Voordat er een levenscyclus analyse kan worden uitgevoerd is het belangrijk om de broeikasgasproductie van eetbare insecten te weten. Deze informatie is niet te vinden in de literatuur. In deze studie is experimenteel vastgesteld wat de directe broeikasgasproductie van methaan en N₂O is voor de vijf insecten. De CO₂ productie en de gemiddelde dagelijkse toename werden gekwantificeerd om een schatting te maken van de voederconversie. Daarbij werd ook de NH₃ emissie gekwantificeerd. De resultaten van deze studie representeren een kwantificatie van de fysiologische bijdrage van broeikasgassen van insecten en hierdoor zal er een levenscyclusanalyse kunnen plaatsvinden voor insectafgeleide producten.

Materiaal en methode

De proeven van de insecten werden drie tot zes keer herhaald voor een periode van 3 dagen. Hierbij werden de dieren gehuisvest per soort in twee kooien of containers per respiratiekamer. In tabel 1 staat vermeld wat de klimatologische omstandigheden waren voor de insecten.

Tabel 1 gemiddelde waarden en standaard afwijkingen van temperatuur, luchtvochtigheid, ventilatie, uren of zonlicht en gemiddelde start en eind gewicht van de vijf insecten soorten.

	Kongo-rozenkever	Meelworm	Argentijnse Kakerlak	Huiskrekel	Europese sprinkhaan
Temperatuur (°C)	28,0±0	25,0±0	28,0±0	28,0±0	32,0±0
Luchtvochtigheid (%)	84,3±3,3	79,8±0,2	70,0±0,0	69,9±0,1	69,7±0,2
Ventilatie (L/min)	6,46±2,06	6,82±1,31	5,16±0,05	11,18±1,80	4,98±0,39
Uren licht	0	0	12	12	12
Start gewicht (kg)	0,99	0,91	1,10	0,96	0,08
Eind gewicht (kg)	1,10	1,10	1,28	1,17	0,13

Resultaten

De productie van CO₂ is uitgedrukt per kilogram gemiddeld levend lichaamsgewicht per dag, en per kilogram gewicht toename daarnaast is de gemiddelde dagelijkse gewicht toename vermeld. De productie van methaan, N₂O en CO₂ equivalenten en NH₃ is uitgedrukt per kilogram gemiddeld levend gewicht per dag en per kilogram gewichtstoename.

3.1 gemiddelde dagelijkse toename en CO₂ productie.

De gemiddelde dagelijkse toename varieerde tussen de 4,0%(Kongo-rozenkever) en de 19,6%(Europese treksprinkhaan). De drie andere soorten hebben een gemiddelde dagelijkse toename van 6-7%. De CO₂ productie van de 5 insectensoorten verschillen significant en varieerden tussen de 19(Argentijnse kakkerlak) en de 110(Europese treksprinkhaan) gram per kg gemiddelde levend lichaamsgewicht. De CO₂ productie per kg metabool gewicht(het gewicht van metabolisch actieve weefsel) verschilde sterk tussen de verschillende soorten. De CO₂ productie uitgedrukt per kg gewichtstoename was gemiddeld voor de Europese treksprinkhaan door de hoge gemiddelde dagelijkse toename. Nog altijd was de CO₂ productie van de Europese treksprinkhaan per kg gewichtstoename meer dan het dubbele van CO₂ productie van de Argentijnse kakkerlak. De Kongo-rozenkever had de hoogste productie van CO₂ per kg gewichtstoename (1,539 g/kg. Dit was meer dan het dubbele dan die van de Europese treksprinkhaan.

Tabel 2 CO₂ productie (gemiddelde± standaard afwijking) per kilogram lichaamsgewicht per dag, per kg gewichtstoename en gemiddelde dagelijkse toename voor vijf insectensoorten, varkens en vlees vee

Soorten	CO ₂ (g/kg lichaamsgewicht/dag)	CO ₂ (g/kg gewicht toename)	Gemiddelde dagelijkse gewichtstoename
Kongo-rozenkever	50±22	1.539±518	4,0±2,1%
Meelworm	61±9	1.031±349	7,3±2,5%
Argentijnse kakkerlak	19±3	337±51	6,1±0,7%
Huiskrekel	68±10	1.468±971	7,2±3,4%
Europese sprinkhaan	110±21	734±119	19,6±2,1%
Varkens	21.6-29.6	865-1.194	3,2±0,53%
Vlees vee	5.3-7.0	2.835	0,3±0,07%

3.2 methaan

Er werd methaanproductie gesignaleerd voor de Kongo-rozenkever en de Argentijnse kakkerlak. Bij de andere drie soorten werd geen methaanproductie gesignaleerd. De Kongo-rozenkever produceerde meer dan drie maal meer methaan per kg gewichtstoename dan de Argentijnse kakkerlak(4,9 om 1,4) dit verschil werd veroorzaakt door een hogere productie per kg lichaamsgewicht (0,16 vs 0,08) en een lagere gemiddelde dagelijkse toename(4,0% vs 6,1%)

3.3 N₂O

N₂O werd alleen in significante hoeveelheden geproduceerd door de meelworm en de Europese treksprinkhaan (1,5 en 8,0 mg/kg lichaamsgewicht per dag). De productie van de Europese treksprinkhaan was per kg lichaamsgewicht meer dan vijf keer de productie van de meelworm. De verschillen namen af met een factor tweeënhalve wanneer dit wordt uitgedrukt per kg gewichtstoename. Dit komt door de hogere dagelijkse groei van de Europese treksprinkhaan.

3.4 NH₃

Ammoniak werd geproduceerd door de huiskrekel, de Europese treksprinkhaan en de Argentijnse kakkerlak(3,0-5,4 mg/kg levend gewicht per dag), en varieerde van 36 tot 142mg/kg of gewichtstoename. Significante verschillen waren gevonden tussen dag en nacht de NH₃ emissie

niveaus. Voor de huiskrekel (6,4 en 4,4) de europeese treksprinkhaan (5,6 en 3,9) en de argentijnse kakkerlak (3,4 en 2,6) mg/kg levend gewicht per dag.

Tabel 3 Methaan, N₂O, CO₂ equivalent en NH₃ productie (gemiddelde ± standaard afwijking) per kilogram lichaamsgewicht per dag voor vijf insectensoorten varkens en vlees vee

Soorten	methaan (g/kg lichaamsgewicht/dag)	N ₂ O (mg/kg lichaamsgewicht/dag)	CO ₂ equivalenten (g/kg lichaamsgewicht/dag)	NH ₃ (mg/kg lichaamsgewicht/dag)
Kongo-rozenkever	0,16±0,085	0,0±0,03	4,00±2,13	0,1±0,16
Meelworm	0,00±0,002	1,5±0,13	0,45±0,04	0,0±0,09
Argentijnse kakerlak	0,08±0,021	0,3±0,24	2,12±0,57	3,0±1,63
Huiskrekel	0,00±0,002	0,1±0,13	0,05±0,04	5,4±3,40
Europese sprinkhaan	0,00±0,017	8,0±13,50	2,37±4,02	5,4±1,65
Varkens	0,049-0,098	2,7-85,6	2,03-27,96	4,8-75
Vlees vee	0,239-0,283	Niet beschikbaar	5,98-7,08	14-170

Tabel 4 methaan, N₂O, CO₂ equivalenten en NH₃ productie (gemiddelde ± standaard afwijking) per killogram gewicht toename voor vijf insectensoorten, varkens en vlees vee.

Soorten	methaan (g/kg gewicht toename)	N ₂ O (mg/kg gewicht toename)	CO ₂ equivalenten (g/kg gewicht toename)	NH ₃ (mg/kg gewicht toename)
Kongo-rozenkever	4,9±1,96	1,03±1,06	121,86±49,09	3±4,8
Meelworm	0,1±0,03	25,5±7,70	7,58±2,29	1±2,0
Argentijnse kakerlak	1,4±0,30	5,7±4,05	37,54±8,01	54±31,1
Huiskrekel	0,0±0,09	5,3±6,05	1,57±1,80	142±184,5
Europese sprinkhaan	0,0±0,11	59,5±104,8	17,72±31,22	36±10,8
Varkens	1,92-3,98	106-3457	79,59-1130	1140-1920
Vlees vee	114	Niet beschikbaar	2850	Niet beschikbaar

Conclusie

Bij de auteurs wetens, de studie die hiervoor ligt is de eerste die bericht over de broeikasgassen en ammoniakemissie van eetbare insectensoorten. Een evaluatie van de broeikasgassen van eetbare insecten is het meest relevant als deze gebaseerd is op CO₂ equivalenten per kg of per gewichtstoename. In dat geval is een vergelijking tussen de geselecteerde soorten en van conventioneel vee mogelijk. Hierbij zou het ook kunnen gaan om een vergelijking op basis van een kosten baten principe. Waarbij de broeikasgasproductie (environmental cost) direct gerelateerd is aan de voedselproductie (benefit). Broeikasgasemissie van vier van de vijf insecten soorten, die bestudeerd zijn, zijn lager dan de cijfers die beschikbaar zijn voor varkens, wanneer de

broeikasemissie wordt uitgedrukt in gewichtstoename, dan produceert een insect ongeveer 1% van de hoeveelheid broeikasgassen voor herkauwers.

Het gemeten niveau van de ammoniak emissie van alle insectensoorten in dit experiment waren lager dan de gerapporteerde NH_3 emissie van conventioneel vee.

De gemiddelde dagelijkse gewichtstoename van de insectensoorten in deze studie was hoger dan voor conventioneel vee, terwijl de CO_2 productie uitgedrukt als g/kg gewichtstoename vergelijkbaar of lager is, welke aangeeft dat er een hoger voederconversie voor insecten mogelijk is.

Deze studie laat zien dat insecten als een meer milieu vriendelijk alternatief voor de productie van dierlijk eiwit kunnen worden beschouwt gekeken naar de broeikasgas en NH_3 emissie. De data die naar voren komt in deze studie zijn onmisbaar voor een levenscyclusanalyse.