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# Something in the air? The impact of volatiles on mollusc attack of oilseed rape seedlings

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## ABSTRACT

• **Background and Aims** Mounting concerns about balancing food security with the environmental impacts of agro-chemical use underpin the need to better understand the mechanisms by which crop plants, particularly during the vulnerable seedling stage, attract or repel herbivores.

• **Methods** The feeding preferences of the mollusc *Helix aspersa* were determined for several oilseed rape (*Brassica napus*) cultivars and a rank order of acceptability established. This was compared with glucosinolate concentrations and volatile organic compound (VOC) profiles to determine whether seedling acceptability to molluscs was linked to either form of defence.

• **Key Results** While VOC profiles for each oilseed rape cultivars could be separated by canonical discriminant analysis and associated with mollusc feeding preferences, glucosinolate profiles were unrelated to snail feeding behaviour. A mixture of monoterpenes ( $\alpha$ -pinene,  $\beta$ -myrcene and  $\delta$ -3-carene) was identified as a putative attractant, while a blend of the Green Leaf Volatiles (GLVs) 3-hexen-1-ol, 3-hexen-1-ol acetate, and the monoterpene  $\alpha$ -terpinene was identified as a putative repellent mix. Added to the VOC profile of oilseed rape seedlings, the “repellent” mix reduced mollusc selection, while the “attractant” mix had no effect.

• **Conclusions** Despite the widespread assumption that seedling selection by generalist herbivores is governed by chemical defence and taste, we show that olfactory cues may be more important. Oilseed rape may be atypical of wild plants, but our ability to identify repellent volatile organic compounds that can

influence snail olfactory selection points to new methods for crop protection using modified VOC profiles during the vulnerable seedling stage.

**Key Words:** *Brassica napus* L.; Brassicaceae; Crop Protection; Food Security; Green Leaf Volatiles; *Helix aspersa* Müller; Monoterpenes; Plant Volatiles; Seedling Herbivory; Solid-phase Microextraction

## INTRODUCTION

Human population growth, coupled with the impact of a changing climate on crop yield, has the potential to destabilise global food security (Godfray *et al.*, 2010). In addition, the likely impacts of anthropogenic climate change on agricultural pest numbers and distributions (Bebber *et al.*, 2013) may increase already significant annual global crop losses to pathogens and herbivores (Oerke, 2006) and necessitate increased agro-chemical inputs to maintain productivity. This option is undesirable for many reasons, particularly given the development of pest resistance and imposition of further carbon costs (Alyokhin *et al.*, 2014). Pesticides have adverse effects on key non-target species such as pollinators and cause wider environmental contamination (Clarke *et al.*, 2009; Whitehorn *et al.* 2012). Consequently there is a pressing need to achieve food security whilst minimising the further adverse environmental impact of agro-chemical use (Godfray *et al.*, 2010).

Fundamental to this goal is an understanding of how and why pests are attracted to crop plants and how they are able to overcome natural defences.

Initial location of host plants by pest species in agricultural and forestry systems often relies on detection of volatile organic compounds (VOCs), rather than visual cues (Leather, 1987; Bruce *et al.*, 2005). Once a plant has been located, other cues including taste (Newland, 1998) or physical characteristics such as toughness play a part in determining acceptance or rejection (Hanley *et al.*, 2007). Generalist herbivores are often attracted to plants by VOCs, but may be repelled by specific VOCs, or constitutive chemical or structural defences (Bruce, 2014). Specialist herbivores by contrast are often attracted to specific VOCs, sometimes using the very cues that repel generalists, frequently employing evolved mechanisms to metabolise or sequester defensive toxins produced by the plant. For example, larva of the small cabbage white butterfly (*Pieris rapae*) prefer food plants containing glucosinolates (Renwick & Lopez, 1999), while the monarch butterfly (*Danaus plexippus*) can locate and then sequester cardiac glycosides from milkweed species (Rothschild *et al.*, 1975).

The likelihood and consequences of herbivore attack on plants are more serious for seedlings than mature plants, typically because most seedlings are less well defended against, and thus more susceptible to, herbivore attack than their mature counterparts (Barton & Hanley, 2013). The fact that removal of even small amounts of plant tissue is proportionally more damaging, and harder to recover from, means that seedlings, whether chemically distasteful or not, can ill afford to suffer herbivore attack (Hanley *et al.*, 2004). Even if tissue loss to herbivores at the seedling stage does not result in immediate mortality, significant negative repercussions for plant growth and reproductive potential may be apparent many weeks, or even months later (Hanley and Fegan 2007;

Hanley 2012; Barton 2013). Thus it may be advantageous for a seedling to advertise its chemical defences before damage occurs.

Although many different invertebrate and vertebrates are responsible for seedling losses, the principal seedling herbivores in temperate agriculture are slugs and snails (Moens & Glen, 2002; Strauss *et al.*, 2009). While regarded as generalists (Iglesias & Castillejo, 1999), terrestrial gastropods can be highly selective when presented with a choice of food plants (Hanley, 2004)..

Although a link between seedling chemical defences and mollusc selection is often suggested (Hanley, 2004; Elger *et al.*, 2009) there is in fact little direct evidence to support this assumption (but see Hanley & Lamont, (2001)) and the reasons for seedling selection remain poorly understood (Barton & Hanley, 2013). Recent evidence for temperate grassland plants implies however that molluscs use seedling volatiles to discriminate between species (Hanley *et al.*, 2011; Hanley *et al.*, 2013) supporting the suggestion that the detection of VOCs by pest species may contribute to the location and selection of crop seedlings (see also Carroll *et al.* 2006; Carroll *et al.* 2008). Initial selection of food plants by slugs and snails is by olfaction using the sensory epithelia of the two posterior tentacles (Friedrich & Teyke, 1998) and molluscs are known to respond to the VOCs fenchone, carvone,  $\rho$ -cymene,  $\beta$ -caryophyllene, D-limonene,  $\gamma$ -coniceine,  $\beta$ -pinene and  $\beta$ -myrcene (Frank *et al.* 2002; Birkett *et al.* 2004). Glucosinolates, the constitutive defence chemicals contained in oilseed rape (OSR – *Brassica napus* L.), may also drive gustatory selection by molluscs (Falk *et al.* 2014; Moshgani *et al.*, 2014).

Despite the increased interest in seedling-herbivore interactions (Barton & Hanley, 2013), no study to date has attempted to link herbivore food plant selection with variation in seedling constitutive chemical defence and volatile chemistry. In addition to offering a valuable contribution to our wider understanding of plant-herbivore interactions, a more complete understanding of variation in seedling defence may be vital in our attempts to protect crops from pest attack over the coming decades. The aim of this study therefore is to elucidate whether selection of cultivars of the commonly cultivated crop plant, oilseed rape, by a generalist mollusc herbivore (the snail *Helix aspersa* Müller) is linked to variation in the expression of seedling VOC profiles and glucosinolate defence. We test the following hypotheses:

- 1: That *Helix aspersa* exhibit consistent feeding preferences between different seedlings (i.e. different oilseed rape cultivars)
- 2: That selection of oilseed rape seedlings by snails is linked to glucosinolate and VOC profiles
- 3: That seedlings with effective glucosinolate defences advertise defensive capability with distinct VOC profiles, to deter potentially fatal “sampling” by herbivores
4. That repellent VOC profiles identified for low-acceptability cultivars can be used to manipulate snail olfactory preferences.

To this end we quantified the feeding preferences of molluscs for seedlings of 13 oilseed rape cultivars against a lettuce standard to yield an Acceptability Index (AI) and compared this with glucosinolate and volatile profiles from six



cultivars, chosen to span the range of measured AI. To alter the attractiveness of oilseed rape, we added putative repellent or attractant volatiles to the profile of oilseed rape seedlings and tested snail preferences in a y-tube olfactometer to determine the feasibility of using VOC signals to influencing mollusc food plant selection.

## **MATERIALS AND METHODS**

### *Study species*

Oilseed rape is widely grown for food, bioenergy oils, and as cattle feedstock (Moens & Glen, 2002). The crop accounted for 715,000 ha (18% by area) of agricultural land use in the UK (DEFRA, 2014) and in 2012, 63.7 million tons was grown worldwide, the largest producers being Canada and China (DEFRA 2012; USDA 2014). Severe damage of oilseed rape seedlings by molluscs necessitates the use of metaldehyde-based molluscicides (Garthwaite *et al.*, 2013) with resulting ecotoxicological effects on non-target species (Birkett *et al.* 2004; Nicholls 2014), and contamination of drinking water (Clarke *et al.*, 2009). Oilseed rape has been the subject of breeding programmes over many decades to reduce its toxicity as an animal feed, consequently reducing concentrations of its main secondary defence metabolite, glucosinolates (Moens & Glen, 2002). This may have consequences for crop protection, especially due to cultivar-specific variation in glucosinolate profiles (Glen *et al.* 1990; but see Giamoustaris and Mithen 1995). Mature plants have a well-resolved VOC profile comprising monoterpenes, green leaf volatiles (GLVs), aromatics, and isothiocyanates (Blight *et al.* 1997; McEwan and Macfarlane 1998), but nothing is known about seedling defences.

Although slugs, particularly *Deroceras* spp. are the most common mollusc pest species of UK arable ecosystems (Moens and Glen 2002; Birkett *et al.*, 2004) we used the snail *Helix aspersa* in our experiments due to ease of collection and culture. As generalist herbivores, *Helix aspersa* and *Deroceras reticulatum* have broadly similar feeding preferences and patterns of seedling selection (Hanley, 1995).

#### *Snail feeding preference*

The oilseed rape cultivars used in these experiments were Amulet; Carnival; Fashion; Kumily; Tamarin (seeds supplied by Senova Ltd., Great Abington, UK); Avatar; Cracker; Sesame; Thorin (LS Plant Breeding, Impington, UK); Agatha; Astrid and Cubic (Grainseed Ltd., Eye, Suffolk, UK). Oilseed rape seeds were germinated in 90 mm-diameter Petri dishes containing two layers of 90mm Whatman No. 1 filter paper, 5 ml of distilled water and maintained in an incubator at 18 °C on a 12:12 light:dark cycle. Following radicle appearance, two seedlings from the same oilseed rape cultivar (cv) were planted 45 mm apart in 50 mm plastic pots containing John Innes No. 2 compost. These plants were planted with two 1-wk-old lettuce seedlings (cv Little Gem) such that the seedlings were arranged in a square with each species at opposite corners.

Lettuce cultivated in the same way as oilseed rape was used to ascertain the relative acceptability of the ‘test’ oilseed rape plants with reference to a standard ‘index’ (Fenner *et al.*, 1999), but due to differences in development time requiring an extra week to attain a similar relative size as 7-d-old oilseed rape at growth stage 1.0 (cotyledons only) to 1.1 (cotyledons plus one leaf) (Sylvester-Bradley, 1985). Seedlings were then cultivated for a further 7 d at 18

°C (12:12 light:dark cycle) to reach growth stage 1.2. Four replicate pots for each oilseed rape cv were sunk into the corners of large plastic propagator trays (375 mm × 230 mm × 60 mm deep) filled with John Innes No. 2 compost, such that the top of each pot was flush with the level of the compost (Hanley and Sykes, 2009). This arrangement was replicated ten times for each oilseed rape cultivar.

Several hundred individual snails were collected around Plymouth, UK (or for subsequent experiments, Southampton), retained in large aerated plastic containers in a constant environment ( $16 \pm 2$  °C, 12:12h day:night) containing compost (Westland Horticulture Ltd, UK), wood, and cuttlefish shells, and fed a mixed diet of lettuce and carrot. Snails were kept under this regime for 4 months before use in behavioural assays and no individual was reused in any trial. Snails were starved for 4 d prior to the start of any experiment. Two snails of uniform size (approx. 30mm diameter) were added to each tray and retained overnight (approx. 16 h) under a clear plastic propagator lid sprayed with water. To determine the Acceptability Index (AI), above-ground seedling tissue loss was quantified for each replicate tray by counting the number of seedlings removed by snails and adding to this an estimate of the proportion of tissue removed from remaining seedlings. Damage was scored as: whole seedling eaten – 1; one leaf eaten – 0.5; half a leaf eaten – 0.25; small hole – 0.02. The combined damage score was used to calculate the AI as follows (modified from (Hanley and Sykes, 2009):

$$A = D_O / (D_O + D_L)$$

Where  $A$  = acceptability index;  $D_O$  = OSR damage score;  $D_L$  = lettuce damage score.

### *Seedling glucosinolate content*

Six cultivars (Agatha, Astrid, Avatar, Cracker, Cubic and Thorin) were selected for further work as they spanned the full AI range. To test the hypothesis that glucosinolate profiles would match snail selection preferences and VOC profiles, we quantified the glucosinolate concentrations of growth stage 1.1 oilseed rape seedlings, and compared with AI by conducting a Canonical Discriminant Analysis (CDA) before testing the resultant discriminant functions (DFs) for correlation (van Dam and Poppy, 2008). Seeds from the six oilseed rape cvs were germinated as described above and after radicle emergence, 20 seedlings from each cv were transplanted into 90mm pots containing John Innes No.2 compost. These were grown on in a glasshouse under natural light, at  $25^{\circ}\text{C} \pm 10^{\circ}\text{C}$  and watered daily. All seedlings were harvested by cutting the stem just above the soil and immediately flash frozen in liquid nitrogen, before storage at  $-80^{\circ}\text{C}$ . Samples were dried in batches of 4 for 48 h using an Edwards Modulyo freeze-drier. Each sample was pulverised in a Retsch MM300 mill (Retsch GmbH, Haan, Germany) and weighed on a Mettler-Toledo AB54 balance (Mettler-Toledo Ltd., Leicester, UK) before glucosinolate extraction. 50.0 mg dry material was placed in a 2 ml Eppendorf tube and extracted as in (van Dam *et al.*, 2004). Correction factors for detection at 229 nm from Buchner (1987) and Brown *et al.* (2003) were used to calculate the concentrations of the different types of glucosinolates based on the sinigrin reference curve.

Glucosinolates were identified based on retention time, UV spectrum, LC-MS analysis of selected reference samples, and the following reference standards obtained from PhytoPlan (Heidelberg, Germany); glucoiberin (3-methylsulfenylpropylGSL), glucoerucin (4-methylthiobutylGSL), progoitrin (2-hydroxy-3-butenylGSL), sinigrin (2-propenylGSL), gluconapin (3-butenylGSL), glucobrassicinapin (4-pentenylGSL), glucobrassicin (indol-3-ylmethylGSL), sinalbin (4-hydroxybenzylGSL), glucotropaeolin (benzylGSL), gluconasturtiin (2-phenylethylGSL).

#### *VOC collection and GC-MS*

Volatiles were collected from each of the six oilseed rape cultivars to establish cv-specific variation and examine any relationship with AI. For each cultivar, 20 seedlings were grown in seven pots (90 mm diameter pots as described above). In preliminary trials we found that seedlings at growth stage 1.0 – 1.1 did not produce detectable levels of VOCs, so were allowed to grow until they reached growth stage 1.1 – 1.2. Seedlings were gently removed from their pots, soil carefully washed away to avoid damage, and up to 140 seedlings per replicate placed together in a 200 ml glass beaker (Fisher Scientific) with 100 ml of distilled water. Initial trials had established that while this eliminated volatiles from the soil and the pot, it did not alter the VOC profile of the plants (Fig. S1). All collections took place within an environment-controlled room (ECR) at 23°C. Each beaker was placed inside a 46 x 56 cm polyester (PET) oven bag (Lakeland, Cumbria, UK) with one corner cut off, through which a Teflon tube was inserted before being tied shut (Stewart-Jones & Poppy, 2006). Air was drawn from the ECR air inlet via Tygon tubing (Saint-Gobain S.A.,

Paris, France), passed through an activated charcoal filter and pumped into the bag at a rate of 1000 ml min<sup>-1</sup> using a Neuberger KNDC B pump (Neuberger, Freiburg, Germany). Three samples and one control (a bag containing a beaker with distilled water) were taken simultaneously, under two racks of compact fluorescent bulbs, giving approximately 200 μmol photons m<sup>-2</sup> s<sup>-1</sup> of photosynthetically active radiation at canopy height, equivalent to an overcast day. The open end of the bag was tied around a manual solid-phase microextraction (SPME) fibre holder (Supelco Inc., Bellefont, PA, USA) until the bag inflated, after which the SPME holder was removed. The bag was left for 1 h with the pump running to completely purge unfiltered air and allow the plants to acclimatise. The SPME holder was then replaced, the bag allowed to fully inflate and the SPME fibre exposed (Blue PDMS/DVB 65 μm fibre 57310-U, all from Lot no. 40981, Supelco Inc., Bellefont, PA, USA). The airflow to each bag was reduced to 100 ml min<sup>-1</sup> to maintain positive pressure, preventing contamination over the 4 h of VOC collection. As the volume of each tied and inflated bag was approximately 14 l, this small airflow only resulted in one change of air every 140 min over a 240 min collection time. The SPME fibres were never saturated with any particular VOC. The maximum amount collected was a TIC of around 160 000, five to ten times less than typically collected when running standards. Initial trials also established that the proportion of each volatile in the profile remained constant as collection time was increased, so there was no effect of volatiles being excluded from a saturated fibre due to a long collection time (Fig. S2).

VOCs were detected using a Hewlett Packard HP 6890 gas chromatography (GC) system fitted with an HP Innowax column (polyethylene glycol, 30.0 m x

250  $\mu\text{m}$  i.d. x 0.25  $\mu\text{m}$  film) coupled to an HP 6890 Mass Selective Detector (MS) (Hewlett Packard Inc., CA, USA) run in EI mode. Immediately following collection, the VOCs were thermally desorbed from the fibre in the injector port. The GC was operated in splitless mode, with helium carrier gas at 7.45 psi and the inlet temperature at 250°C. The oven was maintained at 50°C for the first 2 min, then increased by 5°C min<sup>-1</sup> for 4 min, followed by 10°C min<sup>-1</sup> for 17 min, ending at 240°C. The GC-MS was controlled by HP ChemStation version B.02.05 software (Hewlett Packard Inc., CA, USA) and data analysis by HP Enhanced Data Analysis version B.01.00. This protocol proved sufficiently sensitive to collect and identify a sample from 8 ng each of  $\alpha$ -pinene and  $\beta$ -limonene diluted in hexane and introduced to an oven bag on filter paper (Whatman, GE Healthcare, Little Chalfont, UK). The peak total ion count for these test samples (5000) was at the lower end of the range of the samples taken from the oilseed rape plants (approximately 4000 to 160,000). VOCs were initially identified using the NIST database, and confirmed by comparing retention time with standards (Sigma-Aldrich Ltd., UK).

#### *Modified VOC profile olfactory assay*

Olfactory preferences of snails were tested using a glass y-tube olfactometer (5 cm diameter; main tube 20 cm long; arm length 10 cm; angle between arms 90°). Twenty stage 1.1 (cotyledons and one true leaf) to 1.2 (second true leaf) oilseed rape seedlings (Sylvester-Bradley, 1985), cv Astrid or Agatha, (representing the least attractive and a moderately-attractive cultivar respectively) cultivated as above in 90 mm diameter pots, were in each of the entrainment jars, together with a filter paper with 2.5  $\mu\text{l}$  of hexane. Hexane has

no effect on feeding behaviour in a mollusc (Clark *et al.*, 1997). In the treatment jar, the hexane contained three VOCs at a concentration such that 10 ng of each were present; this concentration was around the middle of the estimated range for the detected volatiles. On the basis that none of our cvs released single VOCs and that phytophagous insects respond to blends of VOCs (Bruce *et al.*, 2005) we used a blend of three VOCs rather than single volatiles. In the putative attractive mix, these were  $\alpha$ -pinene,  $\beta$ -myrcene and  $\delta$ -3-carene; in the putative repellent mix, they were cis-1-hexen-3-ol, cis-1-hexen-3-ol acetate and  $\alpha$ -terpinene.

Air was pumped at 250 ml min<sup>-1</sup> through a charcoal filter using a Neuberger KNDC B pump (Neuberger, Freiburg, Germany) through a 100ml Dreschel bottle containing activated charcoal, deionised water in another and then to one of two glass entrainment jars (10 x 24 cm) containing the plant material. Air and VOCs then flowed from each entrainment jar to one arm of the y-tube. Airflow was maintained at 150 ml min<sup>-1</sup> using a Platon NGX regulator (CT Platon, Domont, France). All apparatus was connected using 8mm diameter Teflon tubing (Fisher Scientific, UK) and all glassware supplied by Soham Scientific (Ely, UK). Three minutes were allowed for the VOCs to reach the y-tube before snails were placed in the main tube. They were deemed to have made a choice once their shell had completely entered one of the arms and they were travelling towards the end. Each treatment was tested using 36 snails, allowed to choose without time restriction although snails were timed through the olfactometer. All equipment was thoroughly washed between trials to remove mucus and rinsed with acetone to remove any remaining VOCs. The y-



tube was flipped between runs and the position of test plants varied to eliminate potential positional bias.

### *Statistical analysis*

Snail feeding preference for different oilseed rape cvs (AI) was analysed by ANOVA using the arcsine square-root transformed AI score, and subsequently by a series of pairwise *t*-tests using a Bonferroni correction in R 3.0.2 (R Foundation for Statistical Computing, Vienna, Austria). A modelling approach was first used to establish whether any factor other than cultivar was significant.

To test whether glucosinolate content varied between oilseed rape cvs, a series of one-way ANOVAs were conducted in R 3.0.2, and correlation between individual glucosinolate and snail preferences tested using Pearson's product-moment correlations with Bonferroni corrections applied. As each sample contains multiple glucosinolates, each of which may vary independently, a Canonical Discriminant Analysis (CDA) was also carried out on the entire glucosinolate data set using IBM SPSS Statistics 21 (IBM Corp., NY, USA), to identify which glucosinolates contributed most to differences between cultivars (van Dam & Poppy, 2008). To test Hypothesis 2, that snail feeding preferences are linked to a suite of identifiable defensive chemicals, any significant Discriminant Functions (DFs) from the CDA were tested for correlation with the AI using Spearman's rank correlation in R 3.0.2.

For the VOCs, the cut off point for integration was set at a TIC area of 2500. Peaks that were artefacts of the SPME fibre were also discarded. As more than one SPME fibre was employed for volatile collection, the total abundance of VOCs could not be compared directly. Therefore, the arcsine square-root

transformed proportion of the MS-TIC peak area of each VOC out of the total VOCs for that sample were the responses used in a CDA carried out in IBM SPSS Statistics 21. As with glucosinolate profiles, any significant DFs were tested for correlation with AI values.

To test Hypothesis 4, that addition of VOCs could alter the attractiveness of oilseed rape to snails, a GLM with binomial errors was carried out on the modified VOC profile assay data to determine if cultivar or any positional factor had affected snail choice. As none of these factors were significant and the sample size was small, snail choice was analysed using separate two-tailed exact binomial tests to determine whether snails preferred the modified plant volatile profiles or unmodified controls. As the positive and negative mix tests were entirely independent, no Bonferroni correction was applied. The more stringent two-tailed test was employed, allowing no prior assumptions about the direction of preference. All analysis of the modified profile assays was carried out using R 3.0.2. In all of the above analyses except for the binomial choice assays, data were tested for a normal distribution using Shapiro-Wilks tests in R 3.0.2, and transformed if necessary.

## **RESULTS**

### *Snail feeding preferences*

Snails consistently preferred oilseed rape over lettuce (ANOVA,  $F_{1,258} = 126.6$ ,  $P < 0.001$ ) and a rank order of preference for cultivars was established (Fig. 1), with AI values ranging from the least attractive (cv Astrid), with an AI of  $0.61 \pm 0.09$  (n trials = 10) to the most preferred cv, Thorin ( $0.78 \pm 0.06$ ). Overall, of 897 seedlings attacked by snails, only 6 (0.67%) were not consumed entirely.

### *Glucosinolate content and snail feeding preferences*

Five glucosinolates were detected from the six oilseed rape cultivars investigated (Table 1): progoitrin (PRO); 4-hydroxyglucobrassicin (4OH); 4-methoxyglucobrassicin (4MeO); neo-glucobrassicin (NEO) and glucobrassicin (GBC). Of these, only PRO (ANOVA,  $F_{5,16} = 3.661$ ,  $P = 0.021$ ) and NEO (ANOVA,  $F_{5,16} = 5.867$ ,  $P = 0.003$ ) differed significantly between cultivars. GBC dominated with concentrations ranging from  $0.23 \pm 0.02$  (Astrid) to  $0.73 \pm 0.21$  (Thorin)  $\mu\text{mol g dw}^{-1}$  for PRO, and  $0.34 \pm 0.03$  to  $0.75 \pm 0.09$   $\mu\text{mol g dw}^{-1}$  for GBC. Other compounds rarely exceeded  $0.2 \mu\text{mol g dw}^{-1}$  (Table 1). A canonical discriminant analysis (CDA) was performed and, while this effectively separated the cultivars on the basis of their glucosinolate profile (with 95% of cases correctly categorised), snail feeding preferences were not correlated with any discriminant function (DF); nor were they correlated with any individual glucosinolate, indole glucosinolates nor total glucosinolates. Thus we conclude that the order of snail gustatory selection of oilseed rape cultivars was not related to their glucosinolate profile.

### *VOCs and mollusc preferences*

Twenty-four different VOCs were collected from the six oilseed rape cultivars and were mainly classed as monoterpenes and GLVs (Table 2). Most are well described in oilseed rape, with the exception of the ten unknown compounds, here labelled as Umono A to C (thought to be monoterpenes, from examination of their retention time on the GC column, and the size of fragments in the spectrum obtained from the MS) and unidentified compounds Ucomp A to G. The seedlings produced relatively low levels of volatiles overall, and these

unknown compounds were detected in such small quantities that it was not possible to make a consistent identification. Some compounds were detected in only one sample per cultivar. Monoterpenes dominated the VOC profile of most of the cultivars, particularly  $\alpha$ -pinene,  $\alpha$ -phellanderene,  $\delta$ -3-carene, D-limonene and  $\gamma$ -terpinene. There was also a substantial proportion of GLVs, notably cis-1-hexen-3-ol acetate (particularly dominant in cv Astrid) and cis-1-hexen-3-ol. Although proportions of several VOCs differed between cultivars (Table 2), there was no correlation between any one VOC and the AI preferences.

The CDA on seedling VOCs reduced the data set to five DFs, of which DF1 and DF2 were significant (Fig. 2). The analysis successfully categorised 100% of the samples into their correct cultivar, while DF1 accounted for 82.3% of the variation and DF2 explained 15.2%. The structure matrices for DF1 and DF2 (Table 3) show the correlation between individual VOCs and that discriminant factor. VOCs with a more positive value are likely to be at a higher level in samples with more positive values of that DF, and the opposite is true for negative values. Thus DF1 represents an axis with more GLVs at the negative end and monoterpenes at the positive end, although with some exceptions, while DF2 does not split by chemical category as easily, although there were more GLVs negatively than positively correlated.

There was no relationship between DF1 and AI ( $\rho = 0.2$ ,  $P = 0.7139$ , d.f. = 4), but their DF2 values were significantly correlated with AI ( $\rho = 0.943$ ,  $P = 0.0167$ , d.f. = 4). VOCs most positively associated with DF2 (Table 3) are likely to be the most attractive to snails, while negatively associated compounds

may be repellent (Fig. 3). Thus we conclude that oilseed rape seedling acceptability to snails is strongly related to VOCs (Hypothesis 2).

#### *Glucosinolates and VOCs*

To test the hypothesis that VOC profiles would reflect defensive chemicals (i.e. glucosinolates), we conducted a series of pairwise correlation tests. The monoterpene  $\rho$ -cymene was positively correlated with PRO ( $r^2 = 0.89$ ,  $P = 0.004$ ) and 4MeO ( $r^2 = 0.90$ ,  $P = 0.004$ ), while PRO was also correlated with Ucomp G ( $r^2 = 0.92$ ,  $P = 0.003$ ) and Umono A ( $r^2 = 0.95$ ,  $P = 0.001$ ). These VOCs were not identified as attractive or repellent by the CDA (Table 3) and thus there was only limited support for this hypothesis.

#### *Modified VOC profile assay*

The putative attractant volatiles identified following CDA analysis of the VOCs (“positive mix”) were  $\alpha$ -pinene,  $\beta$ -myrcene and  $\delta$ -3-carene; the “negative mix” contained cis-1-hexen-3-ol, cis-1-hexen-3-ol acetate and  $\alpha$ -terpinene. Positive or negative treatments did not affect the time taken to choose (GLM,  $z = 0.797$ ,  $P = 0.42$ , d.f. = 69). The positive mix attracted only slightly more (53%,  $n = 36$ ) of the snails than did controls, indicating that there was no difference in the attractiveness of the treatments ( $P$  (two tailed) = 0.868), but snails avoided the negative mix, with only 31% ( $n = 35$ ) choosing it over the control ( $P$  (two-tailed) = 0.041). This finding adds further weight to our conclusion and confirms that snail preferences are influenced by (repellent) VOCs. This supports the hypothesis (4) that VOCs can be used to manipulate seedling selection by snails.

## DISCUSSION

Molluscs exhibit strong preferences for seedlings of different plant species (Fenner *et al.* 1999; Hanley, 2004; Barlow *et al.* 2013) but the reasons underpinning selection remain unclear and the link with chemical defences is inferred rather than established (Barton & Hanley, 2013). Terrestrial molluscs are known to discriminate between the odours of macerated seedlings (Hanley *et al.*, 2011) and also avoid food laced with volatiles that they find unpleasant; consequently the fact that we show that snail selection of oilseed rape seedlings is highly correlated with cultivar-specific VOC profiles is perhaps unsurprising. In addition to the fact that this study is the first to show this effect for intact seedlings, it is however interesting that we were unable to locate any relationship between seedling selection by snails and cultivar-specific variation in glucosinolates. Thus seedling acceptability in oilseed rape was determined primarily by olfactory selection (VOC profile) rather than gustatory cues, likely to be a much more effective way of signalling defensive capabilities to herbivores than a process that relies on potentially lethal tissue loss at such an early ontogenetic stage (Hanley and Fegan 2007; Hanley 2012).

Nonetheless, the lack of any relationship between glucosinolates and AI was unexpected, especially given that many studies have detected a deterrent effect for (or reduced performance of) molluscs when presented with plants high in glucosinolates (Moshgani *et al.*, 2014; but see Moyes *et al.* 2000). This may be due to the low glucosinolate levels present in young seedlings; indeed, the glucosinolate concentration in our samples was around 10% of those reported for mature cultivated and wild brassica plants (Gols *et al.*, 2008; Moshgani *et*

*al.*, 2014) and selective breeding has also reduced natural glucosinolate levels (Moens & Glen, 2002). Given that biochemical constraints on young seedlings imposed by the need to develop resource-acquiring organs (true-leaves and roots) are thought to limit the ability to independently synthesize chemical defences (Boege and Marquis, 2005), it is likely that cotyledon-stage defence is the result of maternal provisioning alone. Indeed, Cole (1980) reported a positive association between glucosinolate concentrations and plant age in oilseed rape during early ontogeny. Consequently, the spectrum of possible defences available for young seedlings is limited (including a likely inability to deploy induced defences) and low glucosinolate concentrations in oilseed rape unsurprising. Given that *Helix* is also able to produce a sulfatase enzyme capable of detoxifying glucosinolates (Wittstock *et al.* 2000), snails are likely equipped to tolerate the low level of glucosinolates present in young oilseed rape seedlings.

Cultivar-specific variation in VOC profiles therefore remains as the strongest candidate to explain oilseed rape seedling selection by snails. The relationship between VOCs and seedling acceptability to snails we established, enabled us to predict which VOCs would be attractant or repellent, and demonstrate that mollusc feeding behaviour can be modified by the addition of repellent VOC blends. Many studies with gastropods have shown that single chemicals are insufficient to elicit any behavioural response (but see Chase 1982), but like many other invertebrate herbivores (Webster *et al.*, 2008) molluscs show greater response to volatiles presented as a blend (Fink *et al.*, 2006), a situation that most closely mirrors natural field conditions (Ache and Young, 2005). The VOC profiles of the least attractive oilseed rape cultivars were actually

characterised by GLVs, a group of plant VOCs indicative of herbivore damage (Shiojiri *et al.* 2006; Fürstenau *et al.* 2012) and whose increased concentrations with age have previously been linked to reduced olfactory selection of *Plantago lanceolata* by *Helix* (Hanley *et al.*, 2013). GLV repellent capabilities were further evidenced here by the fact that the “negative” VOC mix contained two GLVs (cis-1-hexen-3-ol and cis-1-hexen-3-ol acetate) and altered snail behaviour greatly. Given that we found no relationship between the expression of these two GLVs and glucosinolate profiles across our oilseed rape cultivars, it seems likely that these VOCs form the basis of early seedling defence in oilseed rape (like Schiestl (2014), we also failed to identify any isothiocyanates in seedling VOC profiles, the most plausible candidates to advertise glucosinolates to would-be herbivores). It is interesting that GLVs are derived from the same oxylipin pathway leading to jasmonic acid synthesis (Kombrink 2012; Scala *et al.*, 2013), leaving open the possibility that they might be perceived by herbivores as a signal that a plant’s defences are activated. The other main component of our ‘negative’ blend,  $\alpha$ -terpinene, is a taxonomically widespread volatile (Pherobase, 2014) and has previously been identified as a herbivore repellent (for whitefly, Bleeker *et al.*, 2009 and weevils, Wang *et al.*, 2009). Taken together these volatiles fulfil the criteria suggested by Bleeker *et al.* (2009) for accepting a blend of volatiles as actively repellent.

Crop plants are bred for various desirable characteristics, but most often prioritise increased yield and disease resistance over traits favouring herbivore resistance. We have demonstrated however that distinct VOC profiles already produced by different oilseed rape cultivars could be harnessed to lessen the attractiveness of this crop to molluscs during early establishment. Enhancement



of repellent oilseed rape volatiles might be achieved by conventional plant breeding techniques (Bleeker *et al.*, 2009) and these may be especially effective if used in tandem with other crop protection methods; for instance a “push-pull” system, where pests may be repelled by VOCs from the crop at the same time as being pulled towards nearby attractive plants as part of an integrated pest management strategy (Cook *et al.* 2007; Kergunteuil *et al.* 2015). At a time when increasing demands for food security (Godfray *et al.*, 2010) are in conflict with concern over pesticide use (Whitehorn *et al.*, 2012), we show that for one major crop species at least, plant protection could be developed without ecotoxic side effects.

#### **SUPPLEMENTARY DATA**

Supplementary data are available online at [www.aob.oxfordjournals.org](http://www.aob.oxfordjournals.org) and consist of Figure S1. Comparison of volatiles collected from oilseed rape seedlings cv Avatar with soil carefully washed away and from cv Avatar seedlings left in pots.

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## FIGURE LEGENDS

Figure 1: Mean ( $\pm$  1 SE) seedling acceptability to snails (*Helix aspersa*) of 13 oilseed rape (*Brassica napus*) cultivars determined as a proportion of the amount of damage suffered relative to a lettuce (*Lactuca sativa*) control (Acceptability Index - AI).

Figure 2: Discriminant Factor 1 (DF1) against DF2 from a canonical discriminant analysis on relative proportions of 24 volatile organic chemicals released by six cultivars of oilseed rape (*Brassica napus*) seedlings at growth stage 1.1 – 1.2. Group centroids are represented by closed circles, individual samples (n = 4) by open circles of the same colour, as standard error bars are too small to be shown. The analysis classified each sample into its correct cultivar with 100% accuracy.

Figure 3: Relationship between Discriminant Factor 2 from a canonical discriminant analysis on relative proportions of 24 volatile organic chemicals released by six cultivars of oilseed rape (*Brassica napus*) seedlings at growth stage 1.1 – 1.2, and gustatory preferences of snails (*Helix aspersa*), calculated as oilseed rape seedling consumption relative to a standard index species lettuce (*Lactuca sativa*).



Table 1: Glucosinolate (GSL) concentrations in growth stage 1.1 – 1.2 seedling oilseed rape (*Brassica napus*) cultivars. Key: progoitrin (PRO); 4-hydroxyglucobrassicin (4OH); glucobrassicin (GBC); 4-methoxyglucobrassicin (4MeO); neo-glucobrassicin (NEO).

Oilseed rape		Glucosinolate concentration $\pm$ 1 s.e. ( $\mu\text{mol gdw}^{-1}$ )											Total indoles		Total GSL	
cultivar	n	PRO	4OH	GBC	4MeO	NEO										
Agatha	3	0.327 $\pm$ 0.047	0.097 $\pm$ 0.009	0.534 $\pm$ 0.042	0.039 $\pm$ 0.002	0.158 $\pm$ 0.013							0.829 $\pm$ 0.065	1.156 $\pm$ 0.111		
Astrid	4	0.294 $\pm$ 0.053	0.070 $\pm$ 0.015	0.349 $\pm$ 0.031	0.032 $\pm$ 0.003	0.042 $\pm$ 0.003							0.493 $\pm$ 0.048	0.787 $\pm$ 0.083		
Avatar	4	0.559 $\pm$ 0.048	0.133 $\pm$ 0.030	0.752 $\pm$ 0.094	0.040 $\pm$ 0.005	0.087 $\pm$ 0.014							1.012 $\pm$ 0.126	1.571 $\pm$ 0.167		
Cracker	3	0.279 $\pm$ 0.080	0.133 $\pm$ 0.031	0.434 $\pm$ 0.023	0.030 $\pm$ 0.005	0.072 $\pm$ 0.005							0.669 $\pm$ 0.049	0.948 $\pm$ 0.128		
Cubic	4	0.236 $\pm$ 0.022	0.101 $\pm$ 0.033	0.584 $\pm$ 0.116	0.032 $\pm$ 0.005	0.073 $\pm$ 0.007							0.772 $\pm$ 0.146	1.008 $\pm$ 0.159		
Thorin	4	0.737 $\pm$ 0.213	0.201 $\pm$ 0.101	0.473 $\pm$ 0.121	0.048 $\pm$ 0.012	0.073 $\pm$ 0.024							0.795 $\pm$ 0.257	1.532 $\pm$ 0.469		

Table 2: Volatile organic compounds (VOCs) detected in 6 different cultivars of growth stage 1.1-1.2 oilseed rape (*Brassica napus*) seedlings by GC-MS. Asterisks next to names indicate whether proportions were significantly different across cultivars (\*  $P < 0.05$ ; \*\*  $P < 0.01$ ; \*\*\*  $P < 0.001$ ); figures indicate the percentage of that cultivars total made up by that VOC,  $\pm 1$  s.e.; n.d. = not detected. Ucomp A-G indicates unknown compound, Umono A-C indicate unknown monoterpenes.

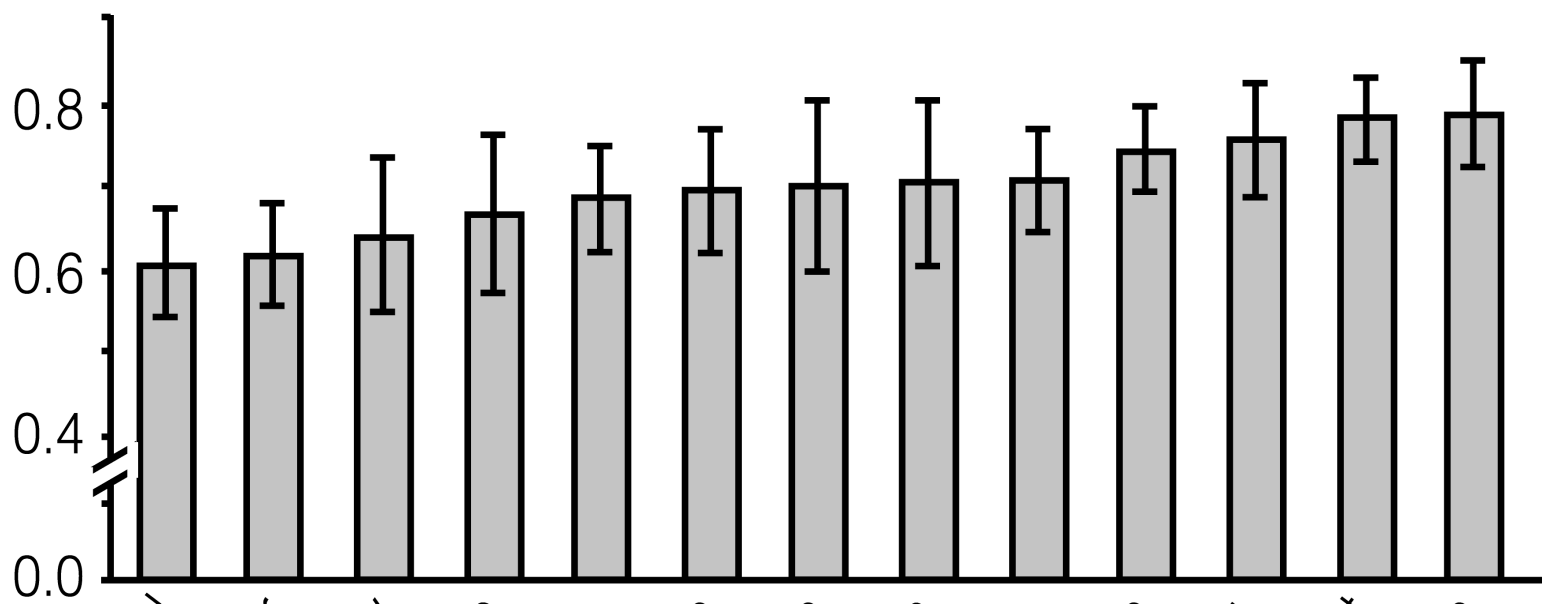
VOC	OSR cultivar			Astrid			Avatar		
	Agatha								
acetone	8.00	±	8.00		n.d.		1.90	±	1.75
Ucomp F	1.24	±	1.24		n.d.		0.19	±	0.19
α-pinene **	8.13	±	1.23	1.82	±	0.89	4.95	±	0.39
α-phellanderene	15.17	±	4.04	4.66	±	2.09	19.55	±	4.20
Ucomp G			n.d.				0.38	±	0.24
Umono C	1.43	±	0.85				3.22	±	1.09
β-myrcene	0.77	±	0.77				1.36	±	1.10
δ-3-carene **	22.71	±	3.94	4.10	±	2.21	18.41	±	2.36
α-terpinene ***			n.d.				2.18	±	0.46
D-limonene *	23.97	±	4.29	6.04	±	2.53	31.75	±	4.02
eucalyptol			n.d.				0.74	±	0.43
Ucomp C *			n.d.						n.d.
γ-terpinene	1.91	±	1.14	0.51	±	0.51	5.66	±	1.14
octadiene	1.85	±	1.85	1.70	±	1.14	0.44	±	0.31
p-cymene	0.72	±	0.72	0.51	±	0.51	1.34	±	0.48
Umono A			n.d.				0.43	±	0.25
Umono B			n.d.				0.25	±	0.25
3-hexen-1-ol acetate ***	1.09	±	1.09	75.27	±	8.65	4.98	±	4.83
anisole	0.45	±	0.45				0.36	±	0.36
3-hexen-1-ol			n.d.	0.65	±	0.40	0.12	±	0.12
Ucomp E	5.86	±	4.77	2.70	±	2.70	0.71	±	0.46
Ucomp A	2.44	±	2.44	1.04	±	1.04	0.16	±	0.16
Ucomp D	1.55	±	1.55	0.55	±	0.55	0.55	±	0.42
Ucomp B *	2.71	±	1.06	0.47	±	0.47	0.36	±	0.23

	Cracker			Cubic			Thorin		
acetone	3.81	±	3.43		n.d.		0.11	±	0.11
Ucomp F			n.d.		n.d.		1.48	±	1.31
α-pinene **	8.12	±	2.54	13.75	±	3.09	5.64	±	0.92
α-phellanderene	15.64	±	6.68	15.10	±	6.48	16.71	±	2.28
Ucomp G			n.d.		n.d.		0.40	±	0.23
Umono C	2.33	±	1.34	1.12	±	1.12	2.17	±	0.80
β-myrcene	6.63	±	5.79	2.39	±	1.85	1.41	±	0.59
δ-3-carene **	16.10	±	3.84	20.49	±	3.00	18.32	±	3.47
α-terpinene ***			n.d.		n.d.		1.38	±	0.80
D-limonene *	19.47	±	8.86	20.66	±	5.67	25.14	±	3.87
eucalyptol			n.d.		n.d.		0.25	±	0.14
Ucomp C *			n.d.		n.d.		0.20	±	0.12
γ-terpinene	2.14	±	1.29	1.41	±	0.87	4.66	±	1.88
octadiene	0.31	±	0.31	1.69	±	1.32	3.27	±	1.72
ρ-cymene			n.d.	0.37	±	0.37	1.72	±	0.65
Umono A			n.d.		n.d.		0.51	±	0.30
Umono B	1.98	±	1.22	0.25	±	0.25	0.19	±	0.19
3-hexen-1-ol acetate ***	17.51	±	8.37	10.45	±	6.56	7.35	±	3.71
anisole			n.d.		n.d.		0.25	±	0.15
3-hexen-1-ol			n.d.		n.d.			n.d.	
Ucomp E	2.08	±	0.79	3.68	±	3.22	5.60	±	3.78
Ucomp A	1.51	±	0.92	1.07	±	1.07	1.68	±	1.39
Ucomp D	0.68	±	0.68	2.54	±	0.88	0.78	±	0.28
Ucomp B *	1.70	±	0.61	5.03	±	1.53	0.79	±	0.58

Table 3: Structure matrices for Discriminant Factor 1 (DF1) and DF2 for Volatile Organic Compounds (VOCs) from 6 cultivars of growth stage 1.1 – 1.2 oilseed rape (*Brassica napus*) seedlings. A more positive value (on a scale of 1 to -1) indicates a stronger positive correlation between that VOC and the DF; a more negative value indicates a negative correlation, while a value close to zero indicates little correlation.

VOC	DF1	VOC	DF2
Ucomp D	0.233	Ucomp B	0.141
Ucomp B	0.188	Ucomp D	0.062
Ucomp A	0.042	$\alpha$ -pinene	0.061
$\alpha$ -terpinene	0.041	$\delta$ -3-carene	0.056
$\delta$ -3-carene	0.038	Ucomp A	0.044
D-limonene	0.034	Ucomp C	0.037
Umono C	0.024	$\beta$ -myrcene	0.022
$\gamma$ -terpinene	0.024	Ucomp F	0.022
$\alpha$ -phellanderene	0.022	Umono B	0.020
eucalyptol	0.021	$\alpha$ -phellanderene	0.016
$\alpha$ -pinene	0.019	octadiene	0.014
Umono A	0.017	D-limonene	0.013
Ucomp G	0.017	acetone	0.011
$\rho$ -cymene	0.011	anisole	0.006
anisole	0.011	Umono C	0.004
acetone	0.010	$\gamma$ -terpinene	-0.009
$\beta$ -myrcene	0.007	Umono A	-0.011
Ucomp F	0.007	$\rho$ -cymene	-0.012
Umono B	0.006	Ucomp G	-0.012
Ucomp C	0.001	Ucomp E	-0.015
octadiene	-0.006	eucalyptol	-0.030
Ucomp E	-0.051	3-hexen-1-ol	-0.052
3-hexen-1-ol acetate	-0.069	$\alpha$ -terpinene	-0.058
3-hexen-1-ol	-0.680	3-hexen-1-ol acetate	-0.086

Acceptability Index



OSR Cultivar

