for Katja and Alfred
Bee pollination of strawberries on different spatial scales – from crop varieties and fields to landscapes

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Chapter 1
General introduction
INTRODUCTION. Pollination is one of the most important ecosystem services based on its contribution to more than one third of world crop production (Klein et al. 2007). Estimated 75% of world crop species (Klein et al. 2007) and 84% of European crop species (Ghazoul 2005) are dependent on or at least profit from animal pollination. Hence, pollination has been frequently confirmed as an ecosystem service of high commercial and social importance (Costanza et al. 1997; Klein et al. 2007; Gallai et al. 2009; Winfree, Gross & Kremen 2011). Growing demands for food and energy (Godfray et al. 2010) and simultaneously increasing cultivation of pollination dependent crops (Aizen et al. 2008, Lautenbach et al. 2012) highlight the need to maintain or even improve future pollination services (Aizen et al. 2008, Lautenbach et al. 2012). Summarizing the main questions of future pollination research (Mayer et al. 2011) emphasize that pollination research is still at the beginning and our knowledge about crop pollination is scarce at various scales. What do we really know about crop pollination, its main facets and drivers and where are important limits?

Current knowledge about the benefits of crop pollination is almost exclusively focused on increasing fruits set and fruit size, resulting in higher yields mainly due to bee pollination (Free 1993; Klein et al. 2007; Jauker et al. 2012; Holzschuh, Dudenhöffer & Tscharntke 2012). But the results from only few available studies (Al-Attal, Kasrawi & Nazer 2003; Cuevas, Hueso & Puertas 2003; Dag & Mizrahi 2005; Shin, Park & Kim 2007; Freihat et al. 2008; Choi et al. 2009; Gajc-Wolska et al. 2011) suggest, that the benefits of pollination, including fruit quality and shelf life, are yet far from being fully explored.

Bees, which are the most important crop pollinators (Roubik 1995; Klein et al. 2007; Dötterl & Vereecken 2010) are still endangered by anthropogenic impacts,
especially from agricultural intensification (Potts et al. 2010). It has been little explored, how bees are attracted by various crop varieties (Rodriguez-Saona et al. 2009), although the availability of varieties is rapidly increasing (CPVO 2012), simultaneously to agricultural intensification. Recent declines of honeybees (Cox-Foster et al. 2007; Anderson & East 2008) demonstrate the importance of pollination services offered by wild bees for the future, but clearly less is known about the general mechanisms of attraction for solitary wild bees (Dötterl & Vereecken 2010). In general, it is still debated whether honeybees, wild bees, or even few generalist species provide adequate pollination services on crops and how this is affected by the surrounding landscape. Previous findings are contradictory, but have so far been conducted on field scales (Albano et al. 2009) or at distinct field locations (Andersson, Rundlöf & Smith 2012) without considering effects arising from different locations on the field. This emphasizes that most studies that have been conducted on pollination so far are limited to specific scales (Potts et al. 2010). Formulating effective conservation strategies for the maintenance of pollination services requires the connection of comprehensive knowledge about pollination (Potts et al. 2010). Hence, the current work is focused on the main facets and drivers of pollination at different spatial scales, from varieties over fields to landscapes. It addresses unknown benefits of bee pollination on strawberry fruits from different varieties including the so far not considered aspect of shelf life, pollinator attraction between varieties due to chemical cues and the role of honeybees, wild bees and overall bee diversity for the pollination efficiency of strawberry flowers at different field locations.
STUDY ORGANISM. Strawberry (Fragaria x ananassa DUCH.) is the most frequently grown soft fruit in the world with an increasing production quantity worldwide (FAOSTATS 2012). More than 1000 varieties are currently available at the market (CPVO 2012) and usually a couple of different varieties are grown on each commercial field. Strawberry varieties differ in several traits as flowering and harvest time and date, resistance against diseases and taste as well as fruit size and quantity (CPVO 2012), but also in their dependence on insect pollination (Connor & Martin 1973; Connor 1975; Zebrowska 1998). In general, most strawberry varieties are self compatible and thus accessible for wind and self pollination (Free 1993). But stigmas become receptive before the antheres of the same flower releases pollen and thus allogamy is favoured (Free 1993).

During their visits, insect pollinators transfer conspecific pollen between plants as well as allocate clumped and aggregated pollen homogenous over the receptacles, increasing the number of fertilized achenes (Svensson 1991). Achenes are the true “nut-fruits” on the surface of the strawberry as an aggregated fruit (Free 1993). They are known to produce growth factors that enhance cell progeny and size (Nitsch 1950; Csukasi et al. 2011) and thus strawberries are known to have weight increases and less deformations in dependence on the variety, if insect pollination is provided (Free 1998). Strawberries are mainly pollinated by bees, with honey bees being most frequent (Free 1993). In addition, several wild bee species visit strawberry flowers with their abundance and species composition depending on the country (Free 1993). In greenhouses, the genus Osmia spp. Panz. has been confirmed as a suitable pollinator for strawberries (Chagnon, Gingras & de Oliveira 1993), which is also known to be an efficient pollinator of strawberries under field conditions (Nye & Andersson 1974; Albano et al. 2009). Strawberry is an excellent model organism for
pollination experiments. Its metabolism is well investigated (see chapters 2 and 3 as well as references therein) and as it is an aggregated fruit, pollination success can be assessed on the individual fruit level (Andersson et al. 2012).

STUDY REGION & SITES. All experiments and surveys were conducted in Germany, on an experimental strawberry field, in a greenhouse and on conventionally managed fields for commercial strawberry selling, respectively. The first study (chapter two) was conducted on an experimental field. It was located adjacent to the greenhouse of the Agroecology group (Agroecology, Department of Crop Sciences, University of Goettingen, Goettingen, Germany) in the eastern part of the city of Göttingen. It was surrounded by hedges from two sides and grassy margins at the other sides. The entire area is surrounded by hedges and dominated by open space that is partly used for experiments and surveys. Large areas remain unused all over the year with several flowering plant species. Five honey bee hives (Apis mellifera L.), several trap nests dominated by Osmia bicornis L., as well as nests of bumble bees (e.g. Bombus terrestris L.) and other ground nesting bees (e.g. Andrena spp. F., Lasioglossum spp. Curt.) were located near the strawberry field. The second study (chapter three) was partly conducted on the experimental field and on a commercial strawberry field. Latter is located in the vicinity of Göttingen with a size of more than 4 ha. The surrounding landscape is dominated by arable land and pastures with a proportion of semi-natural habitats (mostly hedges and forest strips) of about 8%. The field is divided into different parts, concerning to novel plantings each year. Alltogether, eight varieties are grown on the field with the current study focussing on the variety Yamaska.
For the third study (chapter four), the first part of the experiment was conducted in an climate controlled, isolated part at the greenhouse of the Buesgen-Institute (Department of Forest Zoology and Forest Conservation, University of Goettingen, Goettingen, Germany). The second part of this study was conducted at the same commercial strawberry field used for study two (chapter three), but on the varieties Honeoye and Sonata.

The fourth study (chapter five) was conducted on commercial strawberry fields located between the city of Northeim, Lower Saxony, the Southern Harz, Lower Saxony and the city of Kassel, Northern Hesse. This region is dominated by intensive agriculture, interspersed by variously extended areas of semi-natural habitats (see Thies & Tscharntke 1999 and Steffan-Dewenter et al. 2002 for details). The north to south extension of the study area was 50 km and 75 km from east to west. Aim of the study was to analyse landscape effects on pollen loads of bees and the effects of bee diversity and abundance on strawberries. Hence, strawberry fields were selected based on a landscape gradient of surrounding semi-natural habitats ranging from homogenous landscapes with about 2 % of semi-natural habitats to more heterogeneous landscapes with up to 14 % of semi-natural habitats on a radius of 1000 m (ArcGIS, ESRI, München, Germany). Semi-natural habitats were mainly forest, orchards and hedges and tree lines adjacent to crop fields and roads as well as successional areas (unmanaged grassland interspersed with bushes and small trees). Fallows and flowering stripes were not present in the selected landscapes. Strawberry fields were at least 2 km separated and had a minimum size of 1.5 ha.
RESEARCH QUESTIONS & CHAPTER OUTLINE. The current work is focused on the overall benefits and main drivers of crop pollination by bees. It is giving a broad perspective on the influence of bee pollination on strawberry fruits and the underlying mechanisms of bee pollination at different spatial scales such as varieties, fields and landscapes.

Bee pollination of strawberries. Relationships between bee pollination, marketability and post-harvest quality of strawberries in dependence on different varieties were analysed. In detail, the following questions were addressed (separated according to chapters 2 and 3):

- Does bee pollination improve the commercial value and post-harvest quality of strawberry fruits due to higher pollination success compared to wind and self pollination?
- How does strawberry varieties differ on the effects of the three pollination treatments?
- What is the influence of pollination mediated commercial grades on firmness and fruit decay of strawberries during storage.

Strawberry pollination by bees on different spatial scales. Effects from various disciplines influencing strawberry pollination by bees between varieties, field positions and landscapes were analysed. In detail, the following questions were addressed (separated according to chapters 4 and 5):
• Do strawberry varieties differ in the emission of flower volatile compounds?
• How do females of the solitary wild bee *O. bicornis*, a frequently strawberry pollinator, respond to the found compounds?
• Do differences in the emission of flower volatile compounds between strawberry varieties lead to diverse visitation rates of *O. bicornis* females under field conditions?

• Which of the landscape dependent bee parameters abundance, species richness, Shannon-diversity and Evenness is the best predictor of strawberry pollination?
• Does the relative importance of these parameters vary depending on the location in the field?
• How does the performance of *A. mellifera*, the most abundant pollinator species in strawberry fields, influence these parameters and their effects on strawberry yield?

**RESULTS & CONCLUSIONS.** Bee pollination improved the commercial value and post-harvest quality of strawberries depending on varieties. Bee pollinated fruits were heavier and could be sorted into higher commercial grades due a better shape than fruits resulting from wind and self pollination across all varieties. These improvements led to a generally higher commercial value of bee pollinated fruits. Most varieties showed their fruits to have an elongated shelf life, intensified red colour as well as lower sugar-acid ratios from bee pollination, thereby enhancing the
post-harvest quality of strawberries. These effects were confirmed as a result of bee pollination, by higher amounts of fertilized achenes, the true “nut”-fruits of the strawberry, compared to wind and self pollinated fruits.

Detailed analysis on the relationship between pollination and shelf life showed that bee pollinated strawberries and strawberries from higher commercial grades had a higher number of fertilized achenes. Higher commercial grades improved firmness and fruit weight and lead to less decay during storage time. Firmness, fruit weight and decay were highly correlated. Hence, bee pollination resulted in higher commercial grades and elongated the shelf life of strawberries compared to fruits from minor grades, resulting from wind pollination.

Strawberry varieties differed in the emission quantity, not quality, of various flower volatile compounds, while several, for strawberry varieties so far unknown compounds were found. Females of the solitary wild bee *O. bicornis*, a frequent strawberry pollinator, responded on all emitted compounds but in different intensity.

On a commercial strawberry field, *O. bicornis* females visited the variety Sonata more frequently than the variety Honeoye, presumably due to higher emissions of the most important flower volatile compounds.

At the field scale, strawberry pollination was dependent on either, honeybees and wild bees, with effects changing between field locations. Strawberry fruit weight was best explained by Evenness and less by Shannon-diversity, while the effects of bee species richness and abundance were inconsistent. An even bee community benefited strawberry fruit weight at the field edge, while the most dominant pollinator *A. mellifera*, presumably due to higher conspecific pollen loads, was most important at the field centre, leading to contrasting effects of Evenness. In general, Evenness was negatively correlated to the proportion of honeybees. Pollen from
oilseed rape (*Brassica napus* L.) formed the highest fraction of heterospecific pollen and was negatively correlated to the proportion of strawberry pollen.

In conclusion, bee pollination is a key factor for strawberry quantity and in particular quality and appeared to be driven by varying effects between spatial scales with context-dependent contributions of both, honeybees and wild bees. Hence, on one side, pollination is of higher importance for crop production as investigated so far and has the potential to countervail increasing demands on high quality food. On the other side, crop pollination is dependent on the entire bee community as well as influenced by various spatial factors and thus shows up to be complex process, which is highly sensitive to disturbances. This emphasizes that continuing anthropogenic threats as agricultural intensification will have extensive impacts on world food security. Hence, considering a broader perspective on the benefits and requirements of crop pollination as shown in this work, might improve so far insufficient conservation strategies to maintain this highly important ecosystem service for the future.

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Chapter 1 – General introduction


Chapter 2
Bee pollination improves crop quality, shelf life and commercial value
SUMMARY

1. Pollination is known to improve the yield of most crop species and a third of global crop production, but comprehensive benefits including crop quality are still unknown.

2. This is why the commercial value of crop pollination is underestimated, which is particularly alarming in times of agricultural intensification continuing to diminish pollination services.

3. In this study, exclusion experiments on strawberries showed bee pollination, mainly conducted by wild bees, to improve fruit quality, quantity and market value compared to wind and self pollination.

4. Bee pollinated fruits were heavier, had less malformations and reached higher commercial grades.

5. They were firmer, thus improving the commercially important shelf life and had increased redness and reduced sugar-acid-ratios.

6. These comprehensive findings demonstrate bee pollination to be a hitherto underestimated but vital determinant of fruit quality.

KEY-WORDS: colour, commercial grades, ecosystem services, firmness, market value, post-harvest quality, strawberry, wild bees, yield
**INTRODUCTION.** Awareness of global biodiversity losses and declines of ecosystem services such as pollination (Potts et al. 2010) has influenced international politics, shown by the COP strategic plan of the CBD meeting in Nagoya in 2010, but was disregarded by the common agricultural policy (CAP) of the EU. Pollination is known to increase quantity and thereby the commercial value of most crop species (Klein et al. 2007; Gallai et al. 2009). But at the same time, modern agriculture is still contributing to the decline of pollination through intensified management (Potts et al. 2010). Benefits of pollination may include several features of crop quality that have been rarely analysed. Crop features allowing longer storage and thereby, reducing postharvest losses in supermarkets and households are a major challenge worldwide (Tscharntke et al. 2012). In this study, we expand our knowledge of the underestimated benefits of bee pollination by experimentally quantifying its impacts on crop quantity, quality, shelf life and market value. We used strawberries (*Fragaria x ananassa* DUCH.), a worldwide increasing crop (FAOSTATS 2012), as a model system.

In strawberries, flowers are ordered in consecutive flowering periods, becoming smaller over time (Free 1993). Varieties are self-compatible in most cases and stigmas become receptive before the antheres of the same flower release pollen, so that allogamy is favoured. Bee pollination increases strawberry weight and shape, but effects depend on varieties (Free 1993). Recent findings about new metabolic processes in strawberries support the idea, that pollination may also impact the shelf life of strawberries (Given, Venis & Grierson 1988; Roussos, Denaxa & Damvakaris 2009; Villareal, Martinez & Civello 2009; Csukasi et al. 2011). Due to high fruit sensitivity to fungal infections and mechanical injuries, strawberry fruits have a short shelf life (Roussos et al. 2009). More than 90 % of fruits can be non-
marketable after only four days in storage (Hernandez-Munoz et al. 2006). Several studies have focused on the potential elongation of the shelf life of strawberries with modified storage procedures (Civello et al. 1999; Sanz et al. 1999; Hernandez-Munoz et al. 2006; Colla, Sobral & Menegalli 2006; Caner, Aday & Demir 2008), highlighting its economically huge importance. Shelf life and pathogenic susceptibility of strawberry fruits are mostly related to their firmness (Hernandez-Munoz et al. 2006), but also surface colour and sugar-acid-ratios are involved (Civello et al. 1999; Sanz et al. 1999; Hernandez-Munoz et al. 2006; Colla et al. 2006; Caner et al. 2008). Fruit colour further determines the first impression of consumers influencing their purchase behaviour (Caner et al. 2008). The colour of strawberry fruits results mainly from anthocyanin pigments (Given et al. 1988; Seeram 2008), which protect the fruit against UV-radiation and oxidative substances making fruits healthier for humans (Seeram 2008). The colour of fruits has never been related to animal pollination and only few studies report a relation of pollination to firmness (Al-Attal, Kasrawi & Nazer 2003; Shin, Park & Kim 2003; Gajc-Wolska et al. 2011) and sugar-contents (Shin et al. 2003; Cuevas, Huesco & Puertas 2003; Dag & Mizrahi 2005; Freihat et al. 2008; Choi et al. 2009) of fruits. Hence, comprehensive economic gains of bee pollination on strawberries and other fruits are largely unknown and in particular, the potential effect on commercially important parameters of the overall fruit quality has not yet been explored.

We set up a field experiment with nine commercially important strawberry varieties. The influence of self, wind and bee pollination on strawberry fruits was analysed using exclusion treatments. We calculated the commercial value of each fruit based on commercial grades (European Commission 2007) and fruit weight while accounting for market value differences between harvest months (AMI 2012). Further,
we assessed pollination success, quantified by the number of fertilized achenes per fruit, as well as the influence of bee pollination on fruit colour, sugar-acid ratios and firmness, affecting shelf life. We fitted linear mixed effects models with pollination treatments as fixed effect levels and with random effects allowing treatment slopes and intercept to vary among varieties. To test whether all or individual pollination treatments had a main effect across all varieties or predominating variety differences impede differences between pollination treatments, models with unpooled and successively pooled treatment levels as well as models without treatment as a fixed effect were compared (Bolker et al. 2009) using AICc and likeliness (Burnham & Anderson 2002), respectively.

MATERIAL & METHODS. Nine commercially important strawberry varieties of *Fragaria x ananassa* DUCH. (Darselect, Elsanta, Florence, Honeoye, Korona, Lambada, Salsa, Symphony, Yamaska) were planted on an experimental field in 2008. The field was subdivided in twelve plots and nine rows per plot planted with 18 plants of a single variety per row. All varieties were present in all plots. The sequence of the rows within the plots was randomized. The field was surrounded by two further rows of strawberries to weaken edge effects. Experiments were conducted in 2009 in the first yield year using exclusion treatments on two plants per variety and plot. All buds of a plant were covered with Osmolux®-bags (Pantek, Montesson, France) to allow only self pollination (self pollination treatment), gaze bags (mesh width 0.25 mm) to allow self and wind pollination (wind pollination treatment) or remained uncovered to allow additional insect pollination (bee pollination treatment), respectively. Bags were removed shortly after fruit set and at
least 50 fruits per variety and treatment were harvested at maturity. All analyses except the titratable acid content were conducted at the same day of harvesting to avoid influence on post harvest quality due to water loss and metabolic procedures. The commercial value of each fruit was calculated based on commercial grades and fruit weight (BA2001 S, Sartorius, Goettingen, Germany) while accounting for value differences between harvest months (AMI 2012).

Fruits were sorted into commercial grades, due to aberrations in shape, colour and size, following the official trade guidelines (European Commission 2007). Fruits without or with only slight aberrations were sorted into grade extra/one, whereas distinct aberrations lead to a classification in grade two. Non-marketable fruits had strong colour and shape aberrations. Following the above mentioned Commission Regulation, grades Extra and One can be treated separately, but are used combined in practice. Proportions of fruits for each commercial grade and pollination treatment were calculated across all varieties (Fig. 2A) and also separately for each variety (Table S3).

Fruits were bisected and firmness was measured at the center of each half according to Sanz et al. (1999) with the following modifications: the texture analyzer (TxT2, Stable Micro System, Surrey, England) was fitted with a 5 mm diameter probe and a 25 kg compression cell, while a maximum penetration of 4 mm was used.

Colourimetric analysis were applied according to Caner et al. (2008) at two opposite sides of the center of each fruit in the Lab-colour space using a portable colorimeter (CR-310 Chromameter, Konica Minolta, Badhoevedorp, The Netherlands).

The total soluble solids are strongly correlated to the total sugar content of a solution and were measured using a handheld refractometer (HRH30, Krüss, Hamburg, Germany). Measurements for each fruit were conducted twice and repeated when
the values differed more than 0.2 Brix. Fruit solutions were freeze-dried (Epsilon 2-40, Christ, Osterode, Germany) and all samples from the same plant were pooled and milled. To account for an average water content of 82 %, which was analysed on a sample of 250 fruits, 0.18 g of each freeze dried sample was diluted in 20 ml destilled water and titrated according to Caner et al. (2008).

At least eight fruits from each variety and treatment were used to analyse the number of fertilized achenes per fruit, which represent pollination success. Each fruit was blended in 100 ml distilled water for two minutes (Speedy Pro GVA 1, Krups, Offenbach, Germany). Fertilized achenes are heavier than water and sink to the bottom whereas aborted achenes are lighter and accumulate at the water surface. Fertilized achenes were counted (Contador, Pfeuffer, Kitzingen, Germany) after drying for 48 hours at 85 degrees Celsius.

Mean values were calculated in cases of repeated measurements per fruit. We fitted linear mixed-effects models with random effects allowing treatment slopes and intercept to vary among varieties (Bolker et al. 2009). To account for space and time errors and unbalance in the data, the random part was completed by two further terms: plot variety and plant as well as flowering period as a crossed random effect. Response variables were commercial value per fruit, fruit weight, number of fertilized achenes, firmness and surface colour values (red colour, brightness, yellow colour). In the models with sugar-acid-ratio as response variable, only plot and variety were used to complete the random part, because sugar-acid-ratios were calculated based on arithmetic means per plant.

Bee, wind and self pollination treatments were used as fixed effect levels. To test whether pollination treatments differ and whether there was a main effect of all pollination treatments across all varieties, a model with unpooled treatment levels
(full model), models with successively pooled treatment levels and a model without treatment as fixed effect were compared (Bolker et al. 2009) using second order Akaike’s Information Criterion (AICc) and likeliness (Burnham & Anderson 2002). This allowed us to test whether treatment in general, only specific treatment levels, or no treatment had an effect on the response variables. Latter case indicated that variety differences dominated treatment effects. Residuals were inspected for constant variance, and transformations were used to account for non-normality and heterogeneity, where necessary. Main effect values and parameter estimates were extracted from the model and used for plotting after back transformation.

RESULTS. Strawberry flowers were mainly pollinated by wild bees (75.8 %), while Apis mellifera L. (24.2 %) was less abundant (Table S1). The solitary wild bee Osmia bicornis L. (38.9 %) was the most abundant pollinator, while other wild bee species accounted to less than 5 % of the bee community. Bee pollination resulted in strawberry fruits with the highest commercial value (Fig. 1). On average, bee pollination increased the commercial value per fruit by 38.6 % compared to wind pollination and by 54.3 % compared to self pollination. Fruits resulting from wind pollination had a 25.5 % higher market value than self pollinated fruits. Pollination treatments were stronger than differences between varieties and thus had a main effect across all varieties (Table S2). Our results suggest that altogether bee pollination contributed more than 1.06 billion US-Dollars to a total of 2.76 billion US-Dollars made with commercial strawberry selling in the EU in 2009 (FAOSTATS 2012). Price and marketability of strawberries depend on commercial grades of fruit quality (shape, size and colour) (European Commission
2007). Especially malformations are a common problem affecting strawberry price and marketability (Ariza et al. 2010). Our experiment showed that bee pollination reduced malformations and thus enhanced marketability in all varieties except the variety Symphony (Fig. 2a; Table S3). The highest proportion of bee pollinated fruits was assigned to the best grade extra/one, whereas non-marketable fruits formed the smallest fraction. In contrast, wind and self pollination led to high proportions of non-marketable fruits. Bee pollination did not only enhance fruit shape, but also fruit weight compared to wind and self pollination (Fig. 2b). Bee pollinated fruits were on average 11.0 % heavier than wind pollinated and 30.3 % heavier than self pollinated fruits. Pollination treatments were stronger than differences between varieties and thus had a main effect across all varieties (Table S2).

According to our results, bee pollination significantly impacted the shelf life of strawberries by improving their firmness (Fig. 3a). The firmness values of each treatment and variety were related to shelf life, measured as days until 50 % of fruits
Chapter 2 – Bee pollination improves crop market value and quality

had been lost due to surface and fungal decay (Fig. S1). Higher firmness resulting from bee-pollination potentially elongated the shelf life of strawberry fruits about twelve hours compared to wind pollination and more than 24 hours compared to self pollination. Pollination treatments had a main influence on shelf life across all varieties (Table S2). Varieties producing fruits with high firmness benefitted most from bee pollination.

Bee pollinated fruits further had a more intense red colour compared to fruits resulting from wind and self pollination for most varieties (Fig. 3b). Self pollinated fruits of the varieties Lambada and Symphony showed the most intense red colour in the self pollination treatment. The bee pollination treatment differed from both other pollination treatments across all varieties, whereas strong variety differences impeded a difference between wind and self pollination treatments (Table S2). The brightness of bee and wind pollinated fruits was similar and highly correlated to yellowness (Fig. S2). Thus bee pollination resulted in bright fruits with a more intense red colour than wind pollination fruits, whereas self

**Fig. 2.** Bee pollination improved commercial grades and weight of strawberry fruits. (a) Commercial grades. Fruit proportions (values within bubbles) were calculated for each commercial grade in dependence on pollination treatments (see Table S3 for division into varieties). (b) Weight of strawberry fruits. G1/E = Commercial grade one/extra; G2 = Commercial grade two; NM = non-marketable. Further details and abbreviations are explained in the legend of Fig. 1.
pollinated fruits were darker and less red (Fig. 3b, S2). Senescence of strawberries is not only related to losses in firmness and colour changes, but also to increasing sugar-acid-ratios. Bee pollinated fruits had generally a lower sugar-acid-ratios compared to wind- and self pollinated fruits across all varieties (Fig. 3c), but fruits of the varieties Elsanta and Symphony had a higher sugar-acid-ratio with bee pollination. The difference between wind and self pollination remained variety dependent (Table S2), whereas the sugar-acid-ratio of fruits resulting from bee pollination differed to both other treatments across all varieties.

Pollination success was related to the number of fertilized achenes dependent on pollination treatments. Bee pollination was much more efficient than wind and self pollination, resulting in a higher number of fertilized achenes per fruit across all

**Fig. 3. Bee pollination improves the post-harvest quality of strawberries.** (a) Shelf life in days until 50 % fruit loss was calculated from firmness values that were related to published data on firmness decreases during storage (Hernandez-Munoz et al. 2006). (b) Red colour intensity. (c) Sugar-acid-ratios. Red lines for the main effect are dashed when pollination treatments did not significantly differ, indicating stronger variety effects than pollination treatments (see Table S2 for AICc and likeliness values). Further details and abbreviations are explained in the legend of Fig. 1.
varieties (Fig. 4; Table S2). Bee pollinated fruit on average increased the number of fertilized achenes about 26.8% compared to wind pollination and 61.7% compared to self pollination. Wind pollinated fruits had 47.7% higher number of fertilized achenes than fruits resulting from self pollination. This confirms our findings to be true effects of bee pollination.

**DISCUSSION.** We found bee pollination to play a key role for several features of the quality and quantity of marketable strawberry fruits, including more intensive colour, longer shelf life, less malformations and greater fruit weight. Our results showed strawberries to be mainly pollinated by wild bees. This contrasts to earlier findings, where honeybees were the most abundant pollinator of strawberries (Free 1993), but further highlights the importance of wild bees for crop pollination. The mechanism behind the benefits of bee pollination is based on the fertilization of the true “nut” fruits of the strawberry, the achenes (Given *et al.* 1988; Roussos *et al.* 2009; Villareal *et al.* 2009; Csukasi *et al.* 2011). During their visits, bees allocate pollen homogeneously on the receptacles, increasing the number of fertilized achenes per fruit (Svensson 1991). Unfertilized achenes resulting from insufficient pollination have no physiological functionality (Free 1993). Fertilized
achenes produce the plant hormone auxin, which mediates the accumulation of gibberellic acids (Csukasi et al. 2011). Together, these plant hormones induce fruit growth by improving cell progeny and size and enhance the weight and quality of berries (Roussos et al. 2009). Enhanced levels of auxin and presumably gibberellic acid (Villareal et al. 2009) also delay fruit softening (enhance firmness and shelf life) by limiting the expression of several fruit-softening proteins, the so-called expansins (Given et al. 1988). Whereas auxin alone reduces the accumulation of anthocyanins (Given et al. 1988), it is increased by higher levels of both plant hormones together (Roussos et al. 2009). In contrast to firmness and colour changes, sugar-acid-ratios of strawberries are not directly affected by auxin and gibberellic acid (Roussos et al. 2009). But higher firmness of fruits based on persistent cell walls might reduce respiration, which is known to limit metabolic processes affecting sugar and acid contents during storage (Caner et al. 2008), so that indirect positive effects of pollination are probable.

**CONCLUSIONS.** In conclusion, our results showed that crop pollination is of higher economic importance than hitherto thought. Quality improvements of crops can greatly affect marketability and contribute to reducing crop waste. In the industrialized countries, 40% of all crops are thrown away at retail and consumer levels (Tscharntke et al. 2012; Gustavsson et al. 2011). Under the current scenario of rapid human population increase and global food demand (Godfray 2010), achieving high quality and quantity of crops is a pressing issue. Our study suggests that comprehensive analyses of the benefits of pollination for animal-dependent crops, which comprise 70% of all major crop species, may significantly increase
estimates of the economic value of this ecosystem service. It is economically much more important than previously recognized and needs better implementation in agricultural management policies.

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REFERENCES


SUPPLEMENTARY INFORMATION

Fig. S1. Correlation between firmness and shelf life of strawberries based on published data (11). (a) The proportion of fruits lost due to fungal and surface decay correlated to firmness. At a firmness of 1.23 N, 50% of the fruits were lost due to surface and fungal decay (Spearman’s correlation = -0.87; P = 0.001). (b) Firmness loss based on storage time. Firmness decreased about 0.2 N per day (Spearman’s correlation = -0.97; P = 0.008) at a temperature of 20°C.
Fig. S2. Effects of pollination treatments on the yellow colour and brightness of strawberry varieties. (a) Brightness. (b) Yellow colour. Variety effects were stronger than differences between bee and wind pollination impeding a main effect (indicated by red lines for the main effect displayed dashed; see Table S2 for AICc and likeliness values). Further details and abbreviations are explained in the legend of Fig. 1.
Table S1. Bee pollinators visiting strawberries on the experimental field. To identify the main pollinators of strawberry flowers on the experimental field, four varieties were randomly selected and insects pollinating strawberry flowers were collected. Sweep netting was conducted for ten minutes on four transects that were randomly selected on each of four different days in 2010. Strawberries were mainly pollinated by solitary wild bees with *O. bicornis* being the most frequent species, while honey bees (*A. mellifera*) and bumble bees (*Bombus* spp.) were less abundant.

<table>
<thead>
<tr>
<th>Species</th>
<th>Abundance</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Osmia bicornis</em></td>
<td>114</td>
<td>38.9</td>
</tr>
<tr>
<td><em>Apis mellifera</em></td>
<td>71</td>
<td>24.2</td>
</tr>
<tr>
<td><em>Bombus terrestris</em></td>
<td>10</td>
<td>3.4</td>
</tr>
<tr>
<td><em>Andrena flavipes</em></td>
<td>8</td>
<td>2.7</td>
</tr>
<tr>
<td><em>Bombus lapidarius</em></td>
<td>5</td>
<td>1.7</td>
</tr>
<tr>
<td><em>Andrena sp.</em></td>
<td>4</td>
<td>1.4</td>
</tr>
<tr>
<td><em>Andrena gravida</em></td>
<td>2</td>
<td>0.7</td>
</tr>
<tr>
<td><em>Bombus pascuorum</em></td>
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<td>0.7</td>
</tr>
<tr>
<td><em>Bombus pratorum</em></td>
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<td>0.7</td>
</tr>
<tr>
<td><em>Andrena chrysosceles</em></td>
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<td>0.3</td>
</tr>
<tr>
<td><em>Bombus hypnorum</em></td>
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<td>0.3</td>
</tr>
<tr>
<td><em>Bombus sp.</em></td>
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<td>0.3</td>
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Table S2. Delta AICc values and likeliness resulting from model comparisons. AICc = 0 indicates the model with the highest explanatory power. Lower delta AICc and higher likeliness values indicate better explanatory power of a model. Likeness was calculated between models with delta AICc less than seven (Burnham & Anderson 2002). Likeness values are signed with asterisk. Best explaining models are highlighted in bold. None = no level pooled; Sans = model without fixed effect.

<table>
<thead>
<tr>
<th>Fruit parameter</th>
<th>None</th>
<th>Bee = Wind</th>
<th>Wind = Self</th>
<th>Bee = Self</th>
<th>Sans</th>
</tr>
</thead>
<tbody>
<tr>
<td>Commercial value</td>
<td>0</td>
<td>4.512</td>
<td>0.173</td>
<td>3.527</td>
<td>2.501</td>
</tr>
<tr>
<td></td>
<td>*0.403</td>
<td>*0.042</td>
<td>*0.370</td>
<td>*0.069</td>
<td>*0.115</td>
</tr>
<tr>
<td>Fruit weight</td>
<td>0</td>
<td>4.162</td>
<td>3.507</td>
<td>4.872</td>
<td>3.137</td>
</tr>
<tr>
<td></td>
<td>*0.627</td>
<td>*0.078</td>
<td>*0.109</td>
<td>*0.055</td>
<td>*0.131</td>
</tr>
<tr>
<td>Shelf life</td>
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<td>1.791</td>
<td>7.218</td>
<td>5.273</td>
</tr>
<tr>
<td></td>
<td>*0.431</td>
<td>*0.362</td>
<td>*0.174</td>
<td>–</td>
<td>*0.031</td>
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<tr>
<td>Red colour</td>
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<td>1.608</td>
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<td>2.021</td>
<td>0.323</td>
</tr>
<tr>
<td></td>
<td>*0.155</td>
<td>*0.142</td>
<td>*0.317</td>
<td>*0.115</td>
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<td></td>
<td>*0.131</td>
<td>*0.075</td>
<td>*0.378</td>
<td>*0.203</td>
<td>*0.213</td>
</tr>
<tr>
<td>Pollination success</td>
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<td>9.192</td>
<td>8.704</td>
<td>7.290</td>
</tr>
<tr>
<td></td>
<td>*0.894</td>
<td>*0.106</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Brightness</td>
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<tr>
<td></td>
<td>*0.411</td>
<td>*0.589</td>
<td>–</td>
<td>–</td>
<td>–</td>
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<tr>
<td>Yellow colour</td>
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<td>11.648</td>
<td>9.405</td>
<td>10.614</td>
</tr>
<tr>
<td></td>
<td>*0.445</td>
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Table S3. Effects of pollination on commercial grades separated for varieties.

<table>
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<th>Variety</th>
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<th>Commercial grade</th>
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<tr>
<td></td>
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<td>Extra/One (%)</td>
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<tr>
<td>Darselect</td>
<td>Bee</td>
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</tr>
<tr>
<td></td>
<td>Wind</td>
<td>40.8</td>
</tr>
<tr>
<td></td>
<td>Self</td>
<td>48.3</td>
</tr>
<tr>
<td>Elsanta</td>
<td>Bee</td>
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</tr>
<tr>
<td></td>
<td>Wind</td>
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</tr>
<tr>
<td></td>
<td>Self</td>
<td>43.9</td>
</tr>
<tr>
<td>Florence</td>
<td>Bee</td>
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</tr>
<tr>
<td></td>
<td>Wind</td>
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</tr>
<tr>
<td></td>
<td>Self</td>
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</tr>
<tr>
<td>Honeoye</td>
<td>Bee</td>
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</tr>
<tr>
<td></td>
<td>Wind</td>
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</tr>
<tr>
<td></td>
<td>Self</td>
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<tr>
<td>Korona</td>
<td>Bee</td>
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<tr>
<td></td>
<td>Wind</td>
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</tr>
<tr>
<td></td>
<td>Self</td>
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<td>Lambada</td>
<td>Bee</td>
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<td></td>
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<tr>
<td>Yamaska</td>
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<tr>
<td></td>
<td>Wind</td>
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</tr>
<tr>
<td></td>
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</table>
Chapter 3
Enhancing crop shelf life with bee pollination
SUMMARY

1. Shelf life of crops is one of the most important quality parameters. It is influenced by pollination, but effects have not been quantified so far.

2. Commercial grades of strawberries are a function of pollination. We tested how firmness, fruit weight and decay changed in consequence of commercial grades (determined by bee vs. wind pollination).

3. During storage time, firmness and fruit weight declined and the proportion of decayed fruits increased in all three commercial grades.

4. Fruits of higher commercial grades had higher firmness and weight as well as less decay, which resulted in smaller quality loss with storage time. After three days, half of the wind-pollinated fruits (lower grades), but only a fifth of the bee pollinated fruits (grade one) were decayed.

5. Synthesis and Applications. Crop pollination does not only increase yield, but also the economically most important shelf life. Hence, crop pollination has the potential to reduce food waste and pollinator conservation should be considered more seriously in international conservation strategies.

KEY-WORDS: commercial grade, crop quality, decay, degradation, firmness, food demand, quality, storage, strawberry, weight
INTRODUCTION. A rising world population during the next decades will lead to increasing demands of food (Godfray et al. 2010). While increasing importance of pollination dependent crops highlight the need to stabilize pollination services (Aizen et al. 2008; Lautenbach et al. 2012), pollinators are endangered by various anthropogenic threats (Potts et al. 2010). There is some evidence that pollination can improve the quality of agricultural products such as fruits (Al-Attal, Kasrawi & Nazer 2003; Cuevas, Hueso & Puertas 2003; Dag & Mizrahi 2005; Shin, Park & Kim 2007; Freihat et al. 2008; Choi et al. 2009; Gajc-Wolska et al. 2011). Food quality has become a major topic with increasing attention in public, policies and science, last not least because high amounts of food are wasted worldwide due to insufficient quality (Gustavsson et al. 2011; Tscharntke et al. 2012)

Fruits are the most pollination dependent agricultural products (Klein et al. 2007). An important economic factor determining the quality of fruits is their shelf life (Manning 1998). The quality of fruits is declining during storage, due to fruit softening, which leads to weight loss and decay (Toivonem & Brummel 2008). Thus the shelf life of fruits is mainly dependent on fruit softening, represented by the firmness of fruits (Toivonem & Brummel 2008).

So far only four publications reported that the firmness of fruits can be enhanced by insect pollination (Al-Attal et al. 2003; Shin et al. 2007; Gajc-Wolska et al. 2011). The importance of strawberries is increasing worldwide (FAOSTATS 2012) and strawberry yield has been shown to strongly benefit from pollination. Strawberry pollination is mainly conducted by bees and leads to fruits with less malformations and higher weight (Free 1993; Zebrowska 1998). However, a direct relationship between shelf life and pollination for strawberries or any other crop has not been shown so far.
The aim of this study was to analyze the relationship between pollination and shelf life of fruits, using strawberry as a study organism. Strawberries were classified into commercial grades based on malformations mediated by insufficient pollination. Bee pollinated fruits have a higher amount of fertilized achenes than wind pollinated fruits, showing the effectiveness of pollination conducted by bees. Firmness, fruit weight and decay as parameters determining shelf life were measured during storage. We expect those fruits pollinated by bees and of higher commercial grades to have higher amounts of fertilized achenes. Fruit quality during storage should decline due to decreasing firmness and fruit weight and increasing proportions of decayed fruits. This degradation will vary in dependence on commercial grades.

**MATERIAL & METHODS.** In 2009, exclusion treatments considering bee, wind and self pollination were conducted on an experimental strawberry field with nine strawberry varieties (Fragaria x ananassa Duch.; Chapter 2). From this experiment, bee and wind pollinated fruits of the variety Yamaska, that were sorted into commercial grades based on malformations resulting from insufficient pollination, were used for the current study. Thus commercial grades signed pollination success representing different levels of pollination. The variety Yamaska does not produce antheres, so it depends on cross pollination.

Fruits were collected at maturity, when more than 75% of the surface were coloured red and divided into commercial grades following the Regulation of the European Community (2007). Fruits were blended separately in 100 ml distilled water for two minutes (Speedy Pro GVA 1, Krups, Offenbach, Germany). Fertilized achenes are heavier than water and sink to the bottom whereas aborted achenes are lighter and
accumulate at the water surface. Fertilized achenes were counted (Contador, Pfeuffer, Kitzingen Germany) after drying for 48 hours at 85 °C. In 2012 fruits of the same variety from a conventionally managed strawberry field near the city of Goettingen, Germany, were used. As for 2009, fruits were collected at maturity and classified into three commercial grades based on malformations resulting from insufficient pollination. All selected fruits did not show any physical damage or fungal infection.

To simulate retail conditions, fruits were stored at 20 °C (Sanz et al. 1999; Colla, Sobral & Menegalli 2006; Hernandez-Munoz et al. 2006) for four days. Dependent on their availability on the field, a random set of 7 to 13 fruits from each commercial grade was used for analysis on each consecutive day. Fruits were weighted (BA2001 S, Sartorius, Goettingen, Germany) and visually inspected for physical and fungal decay. For firmness analysis, the peduncle and calyx were removed and fruits were bisected. Firmness was measured at the center of each half according to Sanz et al. (1999), using a texture analyser (TA-XT2 Textur Analyzer, Stable Micro Systems, Surrey, UK).

**Fig. 1. Differences in the mean number of fertilized achenes for pollination treatments and commercial grades.** (a) Mean number of achenes for pollination treatments resulting from exclusion experiments. (b) Mean number of achenes for commercial grades. The mean number of fertilized achenes significantly differed between pollination treatments ($\Delta$ Deviance$_{1,9}$ = 14.10; $P < 0.001$) and commercial grades ($\Delta$ Deviance$_{2,8}$ = 15.03; $P < 0.001$), respectively. G1 = grade one, G2 = grade two, NM = non-marketable. $P < 0.05$ = significant.
fitted with a 5 mm diameter probe and a 25 kg compression cell with the following adjustments: pre-test speed 6.00 mm/sec.; test speed 1.0 mm/sec; post-test speed 8.0 mm/sec; penetration distance 4 mm; trigger force 1.0 N. The maximum force in Newton reached during tissue breakage was recorded as firmness and mean values of both halves for each fruit were used for statistical analysis. Generalized linear models (package “MASS”; Venables & Ripley 2002) with negative binomial distribution were fitted in R (R Development Core Team 2012) to analyse the relationship between pollination treatments as well as commercial grades and the number of fertilized achenes. Linear models (package “stats“; R Development Core Team 2012) were used to test effects of storage time in interaction with commercial grades on fruit weight and firmness, respectively. Generalized linear models (package “base“; Venables & Ripley 2002) with quasibinomial distribution were fitted to test whether storage time and commercial grades had an effect on the proportion fruit decay. Residuals were inspected to meet model assumptions. P-values were defined as significant if smaller than 0.05.

RESULTS. Bee pollinated strawberry fruits had a significantly higher number of fertilized achenes than wind pollinated fruits ($\Delta Deviance_{1,9} = 14.10; P < 0.001$; Fig. 1). Calculating the number of achenes between commercial grades showed that fruits of grade one had a much higher number of fertilized achenes compared to fruits of grade two and non-marketable fruits ($\Delta Deviance_{2,8} = 15.03; P < 0.001$; Fig. 1). Bee pollination exclusively resulted in fruits of grade one, while all wind
pollinated fruits were of the minor grade two or even non-marketable (Fig. 1). Firmness ($F_{1,151} = 96.597; \ P < 0.001$; Fig. 2a) and fruit weight ($F_{1,154} = 41.836; \ P < 0.001$; Fig. 2b) significantly declined during storage time, while the proportion of decayed fruit was increasing ($F_{1,155} = 151.110; \ P < 0.001$; Fig. 2c). Declines in firmness and fruit weight and the increase in fruit decay did not differ in slope between commercial grades. However, commercial grades mattered. Better commercial grades showed higher values of firmness ($F_{1,151} = 8.214; \ P = 0.005$; Fig. 2a) and fruit weight ($F_{1,154} = 184.896; \ P < 0.001$; Fig. 2b) as well as lower
proportions of decayed fruits \( (F_{1,154} = 14.501; P < 0.001; \text{Fig. 2c}) \) during the entire storage time.

Fruit decay was negatively correlated to firmness (\textit{Pearson’s correlation} = -0.93; \( P < 0.001 \)) and fruit weight (\textit{Pearson’s correlation} = -0.73; \( P = 0.002 \)), while firmness and fruit weight were less, but positively correlated (\textit{Pearson’s correlation} = 0.65; \( P = 0.009 \)).

**DISCUSSION.** Our results demonstrate a relationship between pollination, commercial grades and the shelf life of strawberries. Bee pollinated fruits had a higher number of fertilized achenes and were of higher commercial grades than fruits resulting from wind pollination. Firmness and fruit weight of strawberries decreased during storage time, while the proportion of decayed fruits was increasing. The shelf life of strawberries is short, based on fast quality losses during storage, which is due to their high metabolic activity and sensitivity to fungal decay (Hernandez-Munoz et al. 2006). During fruit ripening, cell wall degrading proteins, so called expansins (Given, Venis & Grierson 1988) are produced, which lead to decreasing firmness and increasing decay during storage (Hernandez-Munoz et al. 2006), reducing firmness, due to the degradation of cell walls, higher respiration rates and the loss of water (Hernandez-Munoz et al. 2006). Thereby strawberry fruits become softer and often muddy as well as more sensitive to fungal decay (Hernandez-Munoz et al. 2006), resulting in fruit decay being correlated to firmness and fruit weight in our results.

Although commercial grades had similar rates of degradation, fruits with higher commercial grades, resulting from successful bee pollination, had a longer shelf life.
due to improved values. Bee pollination leads to a higher number of fertilized achenes (Free 1993; Zebrowska 1998), which produce the hormonal growth factor auxin (Nitsch 1950; Csukasi et al. 2011) that in turn mediates the production of another growth factor, gibberellic acid (Csukasi et al. 2011). Both enhance cell progeny and size and thus fruit weight (Roussos, Denaxa & Damvakaris 2009) and improve firmness by limiting the production of expansins (Villareal, Martinez & Civello 2009). This further prevents malformations, which are caused by areas of unfertilized and thus physiologically inactive achenes (Nitsch 1950; Ariza et al. 2010), thus improving commercial grades.

**CONCLUSIONS.** In conclusion, pollination appears to be a key factor for the shelf life of strawberries. We showed for the first time, that bee pollination essentially elongated the shelf life of crops, which is based on growth factors that are known for many important crops (Toivonem & Brummel 2008). After three days, half of the wind pollinated strawberry fruits, but only a fifth of the bee pollinated fruits were decayed. Pollination contributes to more than one third of the global crop production and has been mainly related to just fruit set, fruit weight and overall yield (Klein et al. 2007). Increased shelf life due to crop pollination also results in less food waste due to insufficient quality (Gustavsson et al. 2011; Tscharntke et al. 2012). Our results provide a new perspective on the dependence of food shelf life on pollination, emphasizing the need to protect and enhance pollination services in international conservation strategies.
ACKNOWLEDGEMENTS. The authors thank K. Mecke for the permission to collect strawberry fruits at his field as well as I. Smit and E. Pawelzik for the provision with lab space and instruments. We thank U. Kierbaum for essential assistance during fieldwork and M. von Fragstein for discussion and comments on earlier drafts of this manuscript. This work has been funded by the German Research Foundation (DFG; TS 45/29-1).

REFERENCES


Chapter 4

Flower volatiles of crop varieties and bee responses
SUMMARY

1. Pollination affects a third of global food production, improving both yield and quality of crops. Volatile compounds of crop flowers mediate plant-pollinator interactions, but differences between crop varieties are still little explored.

2. We investigated whether crop flower visitation is determined by variety-specific flower volatiles using strawberry varieties (*Fragaria x ananassa* Duch.) and the pollination services of the wild bee *Osmia bicornis* L..

3. Flower volatile compounds of three strawberry varieties were measured via headspace collection. Gas chromatographic analysis showed that the three strawberry varieties produced the same volatile compounds but with quantitative differences.

4. Electroantennographic recordings showed that inexperienced females of *O. bicornis* had higher antennal responses to all volatile compounds than to controls of air and paraffin oil, while responses differed between compounds. The variety Sonata, which emitted more of the compounds that evoked highest antennal responses than the variety Honeoye, received also more flower visits of *O. bicornis* females under field conditions.

5. Our results suggest that differences in the emission of flower volatile compounds between strawberry varieties mediate their attractiveness to females of *O. bicornis*. Since quantity and quality of marketable fruits depend on optimal pollination, the role of flower volatiles in crop production need to be better understood and more considered in crop-variety breeding.
KEY-WORDS: abundance, antennal responses, bee visitation rates, flower volatile composition, *Osmia bicornis*, pollinator attraction, scent, wild bees, volatile mediated differences
INTRODUCTION. Globally increasing food demand due to a rapidly rising world population (Godfray et al. 2010) highlights the importance of world food security (Lautenbach et al. 2012). Contributing to more than one third of world crop yield (Klein et al. 2007), pollination appears to be key factor to sustain the stability of agricultural food production (Tscharntke et al. 2012). However, our knowledge of crop pollination is still limited (Mayer et al. 2011). Recent threats on pollination services (Potts et al. 2010), due to declining populations of honeybees (Anderson et al. 2008) and wild bees (Biesmeijer et al. 2006), emphasize to expand the knowledge about bee-flower interactions to maintain pollination services (Dötterl & Vereecken 2010).

The influence of floral traits such as colour, shape and handiness on the foraging behaviour of bees has been widely analysed (Parachnowitsch & Kessler 2010). But it has recently been shown that the scent of flowers can be of higher importance than for example flower size and colour (Parachnowitsch, Raguso & Kessler 2012). Flower volatile compounds are further assumed to be the main drivers for visitation decisions of pollinators including flower constancy (Raguso 2008; Wright & Schiestl 2009; Dötterl & Vereecken 2010), but have so far mainly been reported for honeybees, bumble bees (Wright & Schiestl 2009) and few specialised wild bee species (Dötterl & Vereecken 2010). However, the foraging behaviour of the red mason bee *O. bicornis* has recently been shown to be highly influenced by floral scents (Howell & Alarcon 2007), but details about potentially compounds were missing. Thus pollination ecology is still scarcely linked to chemical traits (Kessler & Halitschke 2009; Dötterl & Vereecken 2010) and in particular our knowledge how crop varieties attract pollinators remains scarce (Rodriguez-Saona et al. 2011; Adler & Irwin 2012).
The concentration of floral volatile compounds can vary due to genetic differences as shown for subspecies (Chess, Raguso & LeBuhn 2008) and populations at different locations (Dötterl, Wolf & Jürgens 2005). Only few studies have so far reported differences between crop varieties (Beker et al. 1989; Robertson et al. 1993; Wright, Skinner & Smith 2002; Pham-Delegue et al. 1989; Cheong et al. 2011; Rodriguez-Saona et al. 2011; Soler et al. 2011). Only four of these studies investigated the influence of varieties differing in volatile emissions on pollinator attraction (Beker et al. 1989; Pham-Delegue et al. 1989; Wright et al. 2002; Rodriguez-Saona et al. 2011).

In the current study we aim to highlight the importance of volatile emissions for the attractivity of crop varieties to wild bee pollinators and thus pollination efficiency. We analyse three strawberry varieties for differences on flower volatile emission and the influences on antennal responses as well visitation rates of the solitary wild bee *O. bicornis*. Due to the high abundance and efficient pollination services, *O. bicornis* has recently been classified as important for crop pollination services (Holzschuh, Dudenhöffer & Tscharntke 2012; Jauker et al. 2012). *Osmia* bees have also been shown to be a suitable pollinator for strawberries (Chagnon, Gingras & Oliveira 1993). Strawberries benefit from pollination by enhanced fruit shape and weight (Free 1993; Zebrowska 1998). Recent findings about pollination improving the marketability and postharvest quality of several strawberry varieties including shelf life (Chapters 2 and 3) highlight the overall importance of strawberry pollination. Strawberry breeding focuses on several plant parameters differing between varieties (CPVO 2012), but the attraction to pollinators appears to be neglected. The emission of volatile composition and quantities has so far been tested for a single variety (Hamilton-Kemp, Loughrin & Anderson 1990) and female and
hermaphroditic flowers are known to differ in their emission of volatiles (Ashman et al. 2005). Neither information about the influence of volatile compounds of strawberry flowers on pollinators, nor differences between varieties are known. In detail we analysed (i) the emission of flower volatile compounds comparing three simultaneously flowering strawberry varieties. (ii) The antennal response of females of *O. bicornis* to these compounds were measured and related to (iii) differences in the visitation frequency of *O. bicornis* females on a commercial strawberry field. We expected strawberry varieties to differ in the qualitative and quantitative emission of flower volatile compounds. Antennal responses of *O. bicornis* females would differ between compounds. This would further mediate the visitation rates of *O. bicornis* females between strawberry varieties under field conditions.

**MATERIAL & METHODS.** As scents can be highly variable depending on environmental conditions (Reinhard & Srinivisan 2009), refrigerated strawberry plants were planted and grown separately in 10 liter vessels in an isolated greenhouse under standardised conditions (20 °C; 60 % RH; 12 h daylight per 24 h). Volatiles were sampled simultaneously on all plants. Varieties differed in the amount of open flowers, but produced similar flower overall mass ($F_{2,19} = 0.839; P = 0.448$).

Volatiles were sampled on a charcoal trap (CLSA-Filter, Daumazan sur Arize, France) using a modified push-pull headspace collection system (Tholl *et al.* 2006) directly from flowers. The flowers were enclosed in a plastic roasting bag (Melitta GmbH, Minden, Germany). Air was circulated through the trap by a miniature pump (Fürgut, Aichstetten, Germany) at a flow of 0.8 l min$^{-1}$. The sampling time was 2
hours. Adsorbed volatiles were eluted with 50 µl of dichloromethane/methanol (2:1). The solvents used were of analytical quality (Suprasolv quality, Merck/VWR, Darmstadt, Germany). After elution, samples were stored in an ultralow temperature freezer at -80°C.

Volatile samples were analysed with a coupled GC-MS consisting of a gas chromatograph Agilent type 6890 connected to a type 5973 quadrupole mass spectrometer (both Palo Alto, USA) with electron ionisation (EI, 70 eV). Two column types in a similar setup, a HP-5ms (Agilent, 30 m, 0.25 mm ID, and 0.25 µm film thickness, phenylmethylsiloxane), and a HP-INNOWax (Agilent, 30 m, 0.25 mm ID, and 0.25 µm film thickness, polyethylenglycol), were used to analyse the composition of the extracts. An aliquot of 1 µl was injected into the injector held at 250°C. The oven temperature program was 50°C held for 1.5 min, followed by an increase of 7.50 °C/min to 200°C, remaining at 200°C for 5 min. Helium (purity 99.999 %) was used as carrier gas (1 ml/min).

For identification of the constituents, mass spectra GC retention values and linear retention indices (Van den Dool & Kratz 1963) were compared to those of authentic standards and those of the mass spectral databases and published parameters (Table 1). Databases used, were Wiley 9 combined with NIST ´08 (McLafferty 2009) and “Terpenoids and Related Constituents of Essential Oils”, a database available from MassFinder 3.07 software (Hochmuth Scientific Consulting, Hamburg, Germany).

Experiments were conducted with antenna of *O. bicorns* females using an EAG setup as described in Weissbecker, Holighaus & Schütz (2004). The tests were carried out by manually injecting the synthetic volatile standards upstream the dissected antenna into a stream with synthetic air. To guarantee standard conditions
stimuli were supplied every 120 s. \(10^{-3}\) (w/w) dilutions of synthetic standards were prepared of benzaldehyde, 6-methyl-5-hepten-2-one, \((Z)\)-3-hexenyl acetate, d/l-limonene, nonanal, methyl salicylate, p-anisidehyde, dihydro-\(\beta\)-ionone, geranyl acetne, \(\beta\)-ionone, and \((E,E)\)-\(\alpha\)-farnesene, in paraffin oil (Uvasol\textsuperscript{®}, spectrosc. qual., high visc., Merck, Darmstadt, Germany). Approximately 100 \(\mu\)l of standard dilution or paraffin oil as a control were dropped on 2 cm\(^2\) filter paper pieces (Schleicher & Schuell, Dassel, Germany). A soaked filter paper was inserted into a 10 ml glass syringe (Poulten & Graf GmbH, Wertheim, Germany). A typical stimulus was supplied by puffing 5 ml of air over the antenna and repeated once for each compound and control. The EAG response for each compound and control was recorded for \textit{O. bicornis}.

The abundance of females of \textit{O. bicornis} on the varieties Sonata and Honeoye was assessed using standardised transect walks on a commercial strawberry field. For each strawberry variety, two adjacent rows were subdivided into nine transects and females of \textit{O. bicornis}, the most abundant wild bee, was counted while visiting strawberry flowers at morning and afternoon on 17 days (26 transect walks) at the variety Sonata and on 16 days (22 transect walks) at the variety Honeoye.

Statistical analyses were carried out using the software R, Version 2.13.2 (R Development Core Team, 2011). To test the differences of floral volatiles of the three cultivars we fitted generalized linear models (“glm”-function in package “stats and MASS”; Venables & Ripley 2002) using quasipoisson distribution with cultivar as fixed factor. Multiple comparisons among varieties were calculated using Tukey contrasts with \(P\)-values adjusted by single-step method (“multcomp”-package; Hsu 1996).
To calculate differences of EAG responses of *O. bicornis* females between synthetic compounds and paraffin oil and air control we fitted glm using quasipoisson distribution considered significant at $P < 0.05$ (glm, *F*-test).

To test abundance differences between varieties on the commercial strawberry field we fitted generalized linear models with variety as fixed effect using quasipoisson distribution considered significant at $P < 0.05$ (glm, *F*-test).

**RESULTS.** In total, strawberry flowers produced 24 volatile compounds. All three varieties emitted all volatile compounds, but differed in the quantities of several compounds (Table 1). The variety Sonata produced highest amounts of (Z)-3-hexenol, methyl salicylate, 6-methyl-5-hepten-2-one, dihydro-β-ionone, β-ionone, β-myrcene, ocimene and (E,E)-α-farnesene. The varieties Honeoye and Darselect produced similar quantities of these compounds, except (Z)-3-hexenol, 6-methyl-5-hepten-2-one and (E,E)-α-farnesene that were more produced by the variety Honeoye. Further, the variety Honeoye produced the highest amounts of lily aldehyde and α-copaene. Latter was produced least by the variety Darselect and more by the variety Sonata, while the production of lily aldehyde did not differ between these varieties. The emission of benzyl alcohol, 2-phenyl ethanol, (Z)-3-hexenyl acetate and geranyl acetone did not differ between the varieties Honeoye and Sonata, but were significantly lower or differed marginally to the variety Darselect, respectively. The variety Darselect produced intermediate but similar quantities of d/l limonene compared to the other varieties, whereas these differed significantly in the emission of this compound. Similar could be observed for the production of 1-Hexanol by the variety Honeoye that produced this compound in
similar but intermediate quantities, whereas the varieties Sonata and Darselect differed significantly. The compounds p-anisaldehyde and lily aldehyde were emitted in similar quantities by the varieties Darselect and Sonata. While p-anisaldehyde was less emitted by the variety Honeoye, lily aldehyde was emitted in higher quantities compared the other varieties. The emission of hexanal, heptanal, benzaldehyde, octanal, nonanal, decanal and phenol did not differed between varieties (Table 1).

![Diagram of antennal responses to synthetic compounds](image)

Fig. 1. Antennal responses of naïve *O. bicornis* females to synthetic compounds identified from floral volatile extracts of strawberry varieties (10⁻³ dilution). (mean ± SE, n = 10).
Antennal responses of *O. bicornis* females were significantly higher to all compounds compared to the controls synthetic air and paraffin oil (Fig. 1). The highest responses were shown on nonanal, 6-methyl-5-hepten-2-one, benzaldehyde, methyl salicylate and (Z)-3-hexenyl acetate. Responses to dihydro-β-ionone, β-ionone and \((E,E)\)-α-farnesene differed only slightly but significantly from the control treatments, whereas responses to p-anisaldehyde, d/l limonene and geranyl acetone were intermediate. The variety Sonata produced two of the compounds that induced highest bee responses (6-methyl-5-hepten-2-one, methyl salicylate) in higher quantities than the other varieties and the compound (Z)-3-hexenyl acetate) in similar quantity to the variety Honeoye. The variety Darselect produced none of the compounds in higher quantities than the other varieties. The varieties Sonata and Honeoye were further tested for their attractiveness to *O. bicornis* on a conventionally managed strawberry field. Females of *O. bicornis* were much more abundant flower visitors on the variety Sonata compared to the variety Honeoye \((F_{1,16} = 9.843; P = 0.006; \text{Fig. 2})\).
DISCUSSION. Here we show for the first time, detailed antennal responses of a generalist wild bee pollinator on a broad spectrum of crop flower volatiles and how different emissions between varieties can influence the bee’s visitation rates due to varying responses on distinct compounds.

Strawberry varieties produced the same flower volatile compounds, but differed in volatile quantities. Significant differences between varieties were found for p-anisaldehyde, lily aldehyde, 1-hexanol, (Z)-3-hexenol, benzyl alcohol, 2-phenyl ethanol, (Z)-3-hexenyl acetate, methyl salicylate, 6-methyl-5-hepten-2-one, dihydro-\(\beta\)-ionone, \(\beta\)-ionone, \(\beta\)-myrcene, d/l-limonene, ocimene, \(\alpha\)-copaene and (E,E)-\(\alpha\)-farnesene, whereas emissions of geranyl acetone and d/l limonene differed only marginally between varieties. There was no difference between all varieties for the compounds hexanal, heptanal, benzaldehyde, octanal, nonanal, decanal and phenol.

Antennal responses of *O. bicornis* females differed between compounds, while the variety Sonata produced higher amounts of two of the five compounds evoking highest responses than the other varieties and was under field conditions visited more frequently than the variety Honeoye.

Compositions and quantities of flower volatile compounds emitted by strawberries have been solely reported by Hamilton-Kemp *et al.* (1990) so far and show differences compared to our results. Almost two thirds of the compounds we found in the current study (hexanal, heptanal, octanal, nonanal, decanal, lily aldehyde, phenol, 6-methyl-5-hepten-2-one, dihydro-\(\beta\)-ionone, geranyl acetone, \(\beta\)-ionone, \(\beta\)-myrcene, \(\alpha\)-copaene, (E,E)-\(\alpha\)-farnesene) have not been found in Hamilton-Kemp *et al.* (1990) and thus are reported here for strawberries for the first time. In contrast, we did not find the compounds germacrene D and hexyl acetate in our samples.
However volatile emissions and their differences between strawberry varieties have never been reported before.

All compounds emitted by strawberry flowers are known to be generally emitted by flowers (Knudsen, Tollsten & Bergström 1993; Knudsen *et al.* 2006; Dobson 2006). Almost half of the compounds (benzaldehyde, octanal, nonanal, decanal, benzyl alcohol, 2-phenyl ethanol, (Z)-3-hexenyl acetate, 6-methyl-5-hepten-2-one, β-myrcene, limonene, ocimene) belong even to the most frequently emitted flower volatile compounds (Knudsen *et al.* 1993; Jürgens, Witt & Gottsberger 2003; Knudsen *et al.* 2006; Dobson 2006).

Honeybees are known to respond to several of the tested compounds, namely *(E,E)-α-farnesene* (Blight *et al.* 1997; Le Metayer *et al.* 1997; Dötterl & Vereecken 2010), limonene (Henning & Teuber 1992; Blight *et al.* 1997), *p*-anisaldehyde (Theis 2006), *(Z)-3-hexenyl acetate* (Henning & Teuber 1992), *methyl salicylate* (Henning & Teuber 1992), benzaldehyde (Blight *et al.* 1997), but some of the found compounds seem also to be attractive to certain wild bees. *Bombus terrestris* L. responds to ocimene (Mena Granero *et al.* 2005), *Lasioglossum* spp. Curt. to *p*-anisaldehyde (Theis 2006) and *Andrena vaga* Panz. to *(E,E)-α-farnesene* (Blight *et al.* 1997; Dötterl & Vereecken 2010) and *methyl salicylate* (Dötterl & Vereecken 2010).

Antennal responses of females of *O. bicornis* to all volatile compounds were higher as controls, while the responses differed among most compounds. Although *Osmia* spp. has been reported to respond to floral scents (Howell & Alarcón 2007), details about certain compounds as in the current study have not been published before. From the five compounds inducing the highest antennal responses (nonanal, 6-methyl-5-hepten-2-one, benzaldehyde, methyl salicylate and *(Z)-3-hexenyl acetate,*
two were produced by Sonata in highest quantities (6-methyl-5-hepten-2-one, methyl salicylate). These data suggest decreasing attractiveness of flowers to bees from Sonata over Honeoye to Darselect. This finding could be confirmed for the varieties Sonata and Honeoye growing in a commercial strawberry field, where females of the most abundant wild bee, *O. bicornis*, visited the variety Sonata much more frequently. Thus the bees’ preference appeared to be related to the emitted volatile compounds (Wright & Schiestl 2009). Although females of *O. bicornis* responded to the whole variety of compounds, different concentrations of the same compounds lead to different bee responses. This supports the idea that the relative quantity of certain compounds, creating a distinguished blend of volatiles, might be the main driver for the distinctiveness among floral scents (Sachse & Galizia 2003; Carlsson & Hansson 2006) and also among strawberry varieties in the current study.

Different concentrations of distinct volatile compounds have so far been reported to influence the visitation frequency of honeybees between varieties of sunflowers (Pham-Delegue *et al.* 1989) and oilseed rape (Wright *et al.* 2002). However, still practically nothing is known how this is affecting wild bee pollinators and thus these effects are still under discussion (Raguso *et al.* 2008). Although *(E,E)*-α-farnesene was produced by the variety Sonata in much higher quantities than all other compounds, antennal responses showed *(E,E)*-α-farnesene belonging to the compounds that were of minor importance for females of *O. bicornis* (dihydro-β-ionone, β-ionone, *(E,E)*-α-farnesene). But it is known to be highly attractive for honeybees (Blight *et al.* 1997; Le Metayer *et al.* 1997) to which the variety Sonata might be also highly attractive. This may indicate a higher level of specialisation for pollinator species in Sonata compared to the varieties Honeoye and Darselect.
Although certain strawberry varieties are preferred by bees, all varieties are visited (Skrebtsova 1957; Connor 1975). The varieties Sonata and Honeoye did not differ in the emission of Nonanal, Benzaldehyde and (Z)-3-hexenyl-acetate that we found among the five compounds evoking high responses of O. bicornis. Nonanal, benzaldehyde and (Z)-3-hexenyl acetate belong to the most frequently found flower volatile compounds (Knudsen et al. 1993; Knudsen et al. 2006; Dobson 2006), are typical for generalist flowers (Jürgens et al. 2003) and highly attractive for many pollinators (Dobson 2006).

Our findings are in line with recent reports on the importance of volatile compounds for the flower selection of O. bicornis (Howell & Alarcon 2007). Wild bees (Klein, Steffan-Dewenter & Tscharntke 2003; Greenleaf & Kremen 2006; Winfree et al. 2008; Breeze et al. 2011) and especially Osmia spp. (Holzschuh et al. 2012; Jauker et al. 2012) have been suggested to be major pollinators of crops and can affect the fitness of plants (Majetic, Raguso & Ashman 2009).

**CONCLUSIONS.** Volatile compounds of crop flowers were important in attracting wild bees for sustaining pollination services. To our knowledge, only two studies (Beker et al. 1989; Rodriguez-Saona et al. 2011) have so far been shown volatile mediated differences of pollinator attraction between crop varieties and the importance for the visitation by solitary wild bees. As varieties of strawberries and other crops differ in the emission of flower volatile compounds, differences in bee visitation rates can be expected to affect pollination success and thereby, yield and quality (Dötterl & Vereecken 2010). Different bee species improve strawberry pollination by complementary behaviour (Chagnon et al. 1993) and sufficient
strawberry pollination needs the services of honeybees and wild bees (Chapter 5). Hence, breeding strawberry varieties and also the farmers’ selection of varieties should focus more on flower volatiles triggering fruit set and market value by the attraction of several pollinator species.
Table 1. Identified floral volatile compounds of three strawberry varieties (ng g\(^{-1}\) flowers). Trace indicates average amount less than 0.1 ng g\(^{-1}\) flowers. Bold font indicates significant \(P\)-values for the calculated model (glm). Different lower-case letters indicate significant pairwise differences between respective means of different strawberry varieties at \(P < 0.05\) (Tukey test).

<table>
<thead>
<tr>
<th>compound</th>
<th>Darselect</th>
<th>Honeoye</th>
<th>Sonata</th>
<th>F-value</th>
<th>(P)-value</th>
</tr>
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<tr>
<td></td>
<td>ID</td>
<td>mean ± SE</td>
<td>mean ± SE</td>
<td>mean ± SE</td>
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<tr>
<td></td>
<td></td>
<td>(n = 7)</td>
<td>(n = 8)</td>
<td>(n = 7)</td>
<td>df = 2, 19</td>
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<td><strong>Aldehyde</strong></td>
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<td>Hexanal</td>
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<td>A,B(^{3})</td>
<td>2.5 ± 0.6</td>
<td>2.4 ± 0.1</td>
<td>3.9 ± 0.8</td>
</tr>
<tr>
<td>Heptanal</td>
<td>908 / 1198</td>
<td>A,B(^{3})</td>
<td>1.5 ± 0.4</td>
<td>1.0 ± 0.2</td>
<td>2.1 ± 0.6</td>
</tr>
<tr>
<td>Benzaldehyde</td>
<td>967 / 1546</td>
<td>A,B(^{1})</td>
<td>18.1 ± 3.4</td>
<td>14.5 ± 1.8</td>
<td>20.1 ± 3.4</td>
</tr>
<tr>
<td>Octanal</td>
<td>1006 / 1303</td>
<td>A,B(^{2})</td>
<td>1.2 ± 0.2</td>
<td>1.0 ± 0.1</td>
<td>1.4 ± 0.2</td>
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<td>Nonanal</td>
<td>1106 / 1406</td>
<td>A,B(^{2})</td>
<td>12.7 ± 3.6</td>
<td>11.3 ± 1.3</td>
<td>14.6 ± 2.4</td>
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<tr>
<td>Decanal</td>
<td>1207 / 1511</td>
<td>A,B(^{4})</td>
<td>5.5 ± 1.1</td>
<td>5.4 ± 1.0</td>
<td>6.9 ± 1.3</td>
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<td>p-Anisaldehyde</td>
<td>1273 / 2051</td>
<td>A,B(^{1})</td>
<td>3.9 ± 0.5</td>
<td>(b)</td>
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<tr>
<td>Lily aldehyde</td>
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<td>0.2 ± 0.0</td>
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<td>1.4 ± 0.0</td>
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<td>&lt; 900 / 1355</td>
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<td>0.5 ± 0.1</td>
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<td>0.8 ± 0.1</td>
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<td>(Z)-3-Hexenol</td>
<td>&lt; 900 / 1389</td>
<td>A,B(^{2})</td>
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<td>2-Phenyl ethanol</td>
<td>1122 / 1927</td>
<td>A,B(^{2})</td>
<td>0.3 ± 0.1</td>
<td>(a)</td>
<td>0.6 ± 0.1</td>
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</table>
### Chapter 4 – Flower volatiles, crop varieties and bee responses

*P*-value $< 0.0001$.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Darselect</th>
<th>Honeoye</th>
<th>Sonata</th>
<th>F-value</th>
<th>P-value</th>
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<td>Ester</td>
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<td></td>
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<td>(Z)-3-Hexenyl acetate</td>
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<td>4.2 ± 1.7 (b)</td>
<td>6.2 ± 3.0 (b)</td>
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<td>1.5 ± 0.2 (b)</td>
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<td>993 / 1349</td>
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<td>0.6 ± 0.1 (b)</td>
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<td>trace (a)</td>
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<td>Geranyl acetone</td>
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<td>0.8 ± 0.1 (b)</td>
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</tr>
<tr>
<td>(E,E)-α-Farnesene</td>
<td>1510 / 1757</td>
<td>A,B$^6$</td>
<td>0.8 ± 0.1 (a)</td>
<td>4.9 ± 1.6 (b)</td>
<td>59.0 ± 17.4 (c)</td>
</tr>
</tbody>
</table>

*: Stereochemistry not determined. Linear retention indices (LRI) were calculated from chromatograms obtained with a HP-5MS (LRI$^a$) and an HP-INNOWax (LRI$^b$) column. Identification (ID) is based upon mass spectrum matched with those of databases (Wiley 09, Nist 08, and Hochmuth, 2004). LRI is confirmed by synthetic standards. Source of synthetic standards: 1 Fluka (Germany), 2 Merck-Suchardt (Hohenbrunn, Germany), 3 Aldrich (Germany), 4 Acros (Germany), 5 Sigma-Aldrich (Steinheim, Germany), 6 TCI (Zwijndrecht, Belgium). n. d. = non detectable.
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Chapter 5
Context-dependent importance of honeybee vs. wild bee pollination from crop field edges to centres
SUMMARY

1. Crop pollination is mainly conducted by bees, but the relative importance of honeybees (*Apis mellifera* L.) or wild bees and their potential differences in responses to within-crop field heterogeneity is unclear.

2. We analysed how diversity and abundance of bees changed from strawberry field edges, to the strawberry-dominated field centre. Evenness and Shannon-diversity were hypothesized to be of higher functional importance for complementary pollination and final crop yield than abundance and richness of bees. Strawberry fruits and bees pollinating strawberry flowers were collected at transects from the edge to the centre of commercial strawberry fields.

3. Strawberries were visited by 24 bee species with *Apis mellifera* L. being most abundant. Fruit weight was best explained by Evenness, less well by Shannon-diversity and least by species richness and bee abundance.

4. The relationship between pollinator community structure and yield contrasted between field edge and centre. Strawberry pollination at the field edge was most efficient when provided by evenly distributed bee communities, whereas dominance of *A. mellifera* was most efficient at the field centre.

5. This changing importance of honeybees was supported by higher heterospecific pollen loads carried by *A. mellifera* at the field edge and higher loads of strawberry pollen in the field centre.

6. *Synthesis and applications*. Honeybees and wild bees are both needed for strawberry pollination, as their relative importance changes from the field edge to the centre. Efficient pollination of strawberries cannot be restricted to
honeybees because of this context dependency of honeybee pollination success. Protection and enhancement of wild bee species, coping with crop field heterogeneity, is needed to maintain high overall crop pollination levels as an important ecosystem service.

**KEY-WORDS:** conspecific pollen load, flower constancy, biodiversity, ecosystem service, Evenness, field heterogeneity, fruit weight, heterospecific pollen load, species richness, strawberries.
**INTRODUCTION.** Bees are the most important pollinators in most cropping systems (Roubik 1995, 2002; Klein *et al.* 2007; Dötterl & Vereecken 2010). Crop pollination often relies solely on honeybees (Klein *et al.* 2007). However, solitary and social wild bees can also provide pollination for crops (e.g. Klein, Steffan-Dewenter & Tscharntke 2003; Greenleaf & Kremen 2006). In several cases, wild bees have been shown to be even more abundant (Winfree *et al.* 2008) and more important (Breeze *et al.* 2011) than honeybees. However, recent views are contradictory whether wild bees or honeybees are most important for crop pollination services (Corbet 1991a, b; Morse 1991; Ollerton *et al.* 2012; Aebi *et al.* 2012).

Insect pollination is known to greatly increase crop yield and quality (Free 1993), and most crops are pollinated by many pollinator species (Free 1993), but the role of changing pollinator community composition is still unclear. While bee abundance and species richness can have positive effects on pollination of several crops other crops seem to be more influenced by the community composition of bees (Klein *et al.* 2007). For example, pollination and fruit set of *Coffea arabica* (Klein *et al.* 2003) and *Cucurbita moshata* (Hoehn *et al.* 2008) is best explained by the species richness, not the abundance of bees.

Complementary benefits of bees have been also shown for strawberry flowers differing in spatial positions on the plant (Chagnon, Gingras & Oliveira 1993). Kakutani *et al.* (1993) found honeybees to be more effective for strawberry pollination than solitary wild bees, because of high numbers of individuals. Following Williams & Thomson (2001), honeybees are less effective, whereas Albano *et al.* (2009) report that single visits of different bee species contribute
equally to strawberry pollination. Thus the patterns of strawberry pollination are still not fully understood.

The aim of our study was to identify the main drivers and patterns of strawberry pollination by bees in conventional strawberry fields. We used abundance, species richness, Shannon-diversity and Evenness of bees along transects from edge to centre of commercial strawberry fields to assess the influence of bee communities on fruit weight of strawberries in relation to field location. Pollen loads of the most common strawberry pollinator *A. mellifera* were collected to analyse its flower constancy in strawberry pollination. We hypothesized that the community composition of bees and its role for pollination changes in dependence on the location on the field. We expected Evenness and Shannon-diversity to be the best predictors of strawberry fruit weight due to complementary effects in strawberry pollination.

**MATERIAL & METHODS.** The study was conducted on seven conventional strawberry fields in the vicinity of Göttingen in 2010. Strawberry varieties can differ in their attraction to bee pollinators (Abrol 1992), thus only one variety – Honeoye – was used on all study sites. This variety flowers at the same time as oilseed rape (*Brassica napus* L.).

Bees pollinating strawberry flowers were collected at four transects from field edge to centre, using sweep nets. Two transects were positioned at the field edge, the first at the field corner (field position one), the second in the middle of the crop row adjacent to the field edge (field position two). The two other transects were positioned in the field, one in the field middle (field position four), the other
(position three) in equal distance between field positions two and four. Distances between field positions varied due to different field sizes. Each transect consisted of two adjacent strawberry rows at a length of 100 plants and was sampled twice for 30 minutes in the morning and 30 minutes in the afternoon. Morning and afternoon samples on the same field were applied on different days. Morning and afternoon samples were pooled and bee abundance, species richness, Shannon-diversity and Evenness were calculated for each field position and used for statistical analysis.

Pollen was sampled from honeybees. All pollen loads were sampled as follows. After removal of pollen baskets, individual bees were placed in Eppendorf tubes with distilled water and a drop of detergent. They were then vortexed to dislodge pollen from the body, and removed from the tube for pinning and identification. Samples were then centrifuged (Centrifuge 5403, Eppendorf, Engelsdorf, Germany) for five minutes at 3,000 rpm. The resulting pollen pellet was then air dried and frozen for later analysis. For counting, three subsamples were taken from each pellet, by mixing the pellet with a sterile metal dissection instrument, and scraping a small amount of pollen onto a microscope slide. A small square (c. 5 x 5 mm) of fuchsin jelly was then melted onto the pollen sample, and covered with a cover slip, thus staining and preparing it for identification and counting with a light microscope. Oilseed rape pollen was identified in addition to strawberry pollen, because oilseed rape played a major role in the surroundings of the strawberry fields. Oilseed rape pollen was not clearly distinguishable from other Brassica species, but other Brassica species were comparatively rare at the sites.

After anthesis, two plants per transect were covered with gauze-bags to protect the fruits from pest damage and hand picking by consumers. All fruits from three stems per plant were harvested when fully ripe and weighted. Mean fruit weight was
calculated per plant while considering consecutive flowering periods, which result in different fruit weight.

Statistical analyses were carried out using the software R, Version 2.13.2 (R Development Core Team, 2011). Bee abundance, species richness, Shannon-diversity and Evenness were calculated for each field and field position. The influence of each parameter on strawberry fruit weight was assessed by fitting linear mixed effects models (“lme”-function in package “nlme”; Pinheiro & Bates 2002)

Fig. 1. Evenness of bee communities in relation to strawberry fruit weight. Effects differed between field positions (from edge to centre, i.e. field positions one to four). (a) Field position one (edge). (b) Field position two. (c) Field position three. (d) Field position four (centre).
with mean fruit weight per plant versus each diversity parameter in interaction with field position. Study site was used as random effect to account for location variations. Residuals were inspected for non-normality and heterogeneity and data were transformed to meet assumptions of heterogeneity and normality where necessary. Models were simplified and bee abundance and diversity parameters were compared for their explanatory power on strawberry fruit weight using second order Akaike’s Information Criterion (“AICc”-function in package “MuMIn”; Burnham & Anderson 2002). Models which AICc values differed less than seven were further compared for their likeliness (“model.sel”-function in the package “MuMIn”; Barton 2009).

The results of the above models were refitted using restricted maximum likelihood and inspected to determine whether bee abundance or diversity parameters generally influenced strawberry fruit weight or whether their influence were dependent on field positions.

**RESULTS.** In total, 1584 fruits were harvested and 805 bees from 24 species were collected. Honeybees (*A. mellifera*) were most abundant (63.2 %), followed by *Bombus terrestris* L. (16.0 %), *B. lapidarius* L. (3.8 %) *Andrena nigroaenea* Kirby (3.7 %) and *Osmia bicornis* L. (3.6 %).

Strawberry fruit weight was best explained by Evenness, whereas Shannon-diversity (delta AICc = 2.67), species richness (delta AICc = 7.45) and abundance (delta AICc = 13.06) were less explanatory. As AICc values between models of Evenness and Shannon-diversity differed less than seven, they were compared for their likeliness. The Evenness model was almost four times as likely as the Shannon-diversity model
(0.794 versus 0.206) confirming a much higher explanatory power of Evenness on strawberry fruit weight.

Effects of Evenness on strawberry fruit weight significantly varied between field positions ($F_{3,40} = 5.021; P = 0.005$; Fig. 1). Whereas Evenness had a positive effect on fruit weight at the field edge (field position one), this effect became smaller with increasing distance to the edge in field positions two and three and became even negative in the field centre (field position four). Similar effects were found for Shannon-diversity ($F_{3,40} = 4.097; P = 0.013$; see Fig. S1 in Supplementary Information). Effects of bee species richness ($F_{3,42} = 4.754; P = 0.006$; see Fig. S2 in Supplementary Information) and bee abundance ($F_{3,42} = 2.999; P = 0.041$; see Fig. S3 in Supplementary Information) on fruit weight significantly differed between field positions, but were inconsistent. The diversity parameters did not have a general relation to fruit weight (Evenness: $F_{1,40} = 0.781; P = 0.382$; Shannon-diversity: $F_{1,40} = 0.004; P = 0.3992$; species richness: $F_{1,42} = 0.393; P = 0.534$; abundance: $F_{1,42} = 0.583; P = 0.449$) of strawberries, and could not explain the differences in fruit weight along the edge-centre field positions.

Proportions of the most abundant bee species *A. mellifera* were calculated for each field position and related to Evenness by fitting linear mixed effects models.
Increasing proportions of *A. mellifera* generally decreased Evenness ($F_{1,19} = 7.991$; $P = 0.011$; Fig. 2), without differing between field positions. There was no overall effect of the proportion of *A. mellifera* on strawberry fruit weight ($F_{1,42} = 0.044$; $P = 0.835$), but its effects were contrasting between field edge and centre ($F_{3,42} = 4.807$; $P = 0.006$; Fig. 3). Fruit weight at the field edge (field position one) was much lower

**Fig. 3. Proportion of honeybees in relation to strawberry fruit weight.** Effects differed between field positions (from edge to centre, i.e. field positions one to four). (a) Field position one (edge). (b) Field position two. (c) Field position three. (d) Field position four (centre).
in cases of higher proportions of *A. mellifera*. Field position two showed a slightly positive effect, field position three a slightly negative effect. In contrast, fruit weight at the field centre (field position four) increased with higher proportions of *A. mellifera*.

Comparisons of pollen loads of *A. mellifera* between field positions showed that the proportion of strawberry (conspecific) pollen was lowest at the field edge and successively increased with increasing distance to the field edge until it reached a maximum of 97% at the field centre (Fig. 4). The second most abundant pollen was from oilseed rape. In contrast to strawberry pollen, the proportion of oilseed rape pollen decreased with increasing distance from the field edge (Fig. 4). Proportions of strawberry pollen and oilseed rape pollen were negatively correlated between transects (*Spearman’s correlation* = -0.79; *P* < 0.001).

![Fig. 4. Pollen loads of *A. mellifera* in dependence on field positions. (a) Proportion of strawberry pollen depended on field location (from edge to centre, i.e. field positions one to four). (b) Proportion of oilseed rape pollen dependent on field location (field positions one to four). Error bars show standard errors.](image-url)
DISCUSSION. Our results showed for the first time that crop pollination is dependent on both honeybees and wild bees. Evenness was the best predictor of strawberry fruit weight. Shannon-diversity, species richness and abundance were less important. However, Evenness was positively related to fruit weight at the edge and negatively in the centre. This pattern was driven by the most abundant bee species, *A. mellifera*. Increasing proportions of *A. mellifera* resulted in decreasing Evenness due to its high abundances. The contrasting influence of Evenness on fruit weight between field positions appeared to be due to the changing proportions of heterospecific pollen collected by *A. mellifera*, preventing consistent pollination success (Morales & Traveset 2008; Mitchell *et al.* 2009; Muchhala & Thomson 2012). Context-dependency of Evenness versus dominance by honeybees on crop pollination has not been reported so far.

Relationships between Evenness and ecosystem processes have been scarcely analysed and have mainly been shown for the productivity and functional diversity of natural ecosystems (Hillebrand, Bennett & Cadotte 2008; Crowder *et al.* 2012). However, in our study effects of Evenness appeared to rely on complementarity effects of wild bees benefitting strawberry pollination at the field edge, whereas beneficial effects of honeybee dominance at the field centre gave evidence for species identity effects.

Why did *A. mellifera* have contrasting effects on yield between field edge and centre? Such effects could result from variation in pollen loads between field locations. Interestingly, the proportion of strawberry (conspecific) pollen carried by *A. mellifera* increased with distance to the field edge. In the field centre, increasing proportions of *A. mellifera* were positively related to pollination and thus fruit
weight. Higher importance of honeybees towards the field centre led to weaker or negative effects of Evenness. Foraging by \textit{A. mellifera} in crop fields is often conducted along crop rows (Pyke 1978; Cresswell 1995), which avoids the revisiting of flowers. In the current study, bees may have started foraging at the field edge and continued along strawberry rows until the field centre. Thus having visited more strawberry flowers by the time they reach flowers at the field centre than at the field edge. In general, \textit{A. mellifera} carried high percentages of strawberry pollen, which revealed high constancy to strawberry flowers. The flower constancy of \textit{A. mellifera} is known to be influenced by other foraging resources than the target one (Wells & Wells 1986). \textit{A. mellifera} is able to change flower constancy and foraging patterns due to changing environmental conditions (Well & Wells 1984, 1986), which can cause heterospecific pollen placement and reduce the productivity and fitness of plants (Morales & Traveset 2008; Mitchell \textit{et al.} 2009; Muchhala & Thomson 2012). In strawberry, even small numbers of unfertilized achenes might lead to malformations and a significantly decreased fruit weight (Free 1993). Hence, the higher proportions of strawberry pollen picked up en route to flowers in the field centre would be likely to make \textit{A. mellifera} more efficient pollinators at central locations, and less efficient pollinators at the field edge, where conspecific pollen loads were low.

Wild bees seem to improve pollination at the field edge, resulting in similar fruit weights compared to the other field positions. In general, wild bees have been found to be more efficient pollinators than honeybees (Parker, Batra & Tependino 1987, Torchio 1990, Richards 1996, Klein \textit{et al.} 2003, Kremen \textit{et al.} 2004, Greenleaf 2006, Bosch, Kemp & Trostle 2006; Tuell, Ascher & Isaacs 2009; Breeze \textit{et al.} 2011; Holzschuh, Dudenhöffer & Tscharntke 2012) for various reasons as better
performance in pollen exchange, transfer and deposition as well as interspecific interactions with honeybees (Holzschuh et al. 2012). However, these findings cannot be directly related to all crops, because the pollination success of honey bees is dependent on flower morphology (Wilson & Thomson 1991) and honeybees are known to be equally efficient strawberry pollinators as wild bees (Albano et al. 2009), which is further suggested by similar fruit weights at all field positions. In general, Evenness and diversity are important for the stability and performance of ecosystem services (Hillebrand et al. 2008). Our results showed that strawberries were pollinated by an uneven bee community, with *A. mellifera* being most abundant at all field positions. Especially in dominated bee communities, a poor performance of the dominating species lead to a higher importance of Evenness and also diversity for the efficiency of the pollination service (Hillebrand et al. 2008; Bommarco et al. 2011). Thus, higher amounts of heterospecific pollen, carried by *A. mellifera* at the field edge suggests an inferior pollination efficiency and could have resulted in a better performance of more even and diverse wild bee communities.

**CONCLUSIONS.** In conclusion, our results show that *A. mellifera* and wild bees are both important for strawberry pollination on commercial fields, but their pollination success appears to be dependent on field positions. This finding provides a new perspective on the current debate on whether honeybees or wild bees are the most important pollinators (Ollerton et al. 2012; Aebi et al. 2012). Further, contrary to assumptions of other studies (Ghazoul 2005), pollination services cannot be sustained by a few generalist species. Our results suggest that declining populations of either bee group can have serious impacts on the overall pollination and the yield
of strawberries. Bee abundance, species richness and Shannon-diversity that are usually expected to be good predictors of crop pollination (e.g. Cardinale et al. 2006), were much worse predictors of strawberry yield than Evenness. Future pollination studies should take potential effects of the environmental context into account, including the location inside fields. Rather than the traditional focus on pollinator abundance and species richness, community composition and Evenness should be considered as a major driver of pollination. Agricultural management policies will have to mitigate threats to all pollinator species to maintain future pollination services across the heterogeneity of crop fields.

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Chapter 5 – Context-dependent pollination of honey- and wild bees

SUPPLEMENTARY INFORMATION

Fig. S1. Shannon-diversity of bee communities in relation to strawberry fruit weight. Effects differed between field positions (from edge to centre, i.e. field positions one to four). (a) Field position one (edge). (b) Field position two. (c) Field position three. (d) Field position four (centre).
Fig. S2. Bee species richness in relation to strawberry fruit weight. Effects differed between field positions (from edge to centre, i.e. field positions one to four). Bee species richness seemed not or only slightly related to strawberry fruit weight at field positions one and three, but strongly contrasted between field positions two and four. (a) Field position one (edge). (b) Field position two. (c) Field position three. (d) Field position four (centre).
Fig. S3. Bee abundance in relation to strawberry fruit weight. Effects differed between field positions (from edge to centre, i.e. field positions one to four). Bee abundance was negatively related to strawberry fruit weight at the field positions one, three and four, but in contrast showed a positive relationship at field position two. (a) Field position one (edge). (b) Field position two. (c) Field position three. (d) Field position four (centre).
SUMMARY. The global majority of our crops is dependent on pollination. Hence, pollination contributes to one third of global crop production and is an ecosystem service of high commercial and social importance. Bees are the most important crop pollinators, but they are endangered by several anthropogenic impacts, in particular agricultural intensification. Facing rising global demands for food and energy in the background of a growing world population, the prevention of an impending pollinator crisis attracts increasing interest from the public, policymakers and scientists. However, despite new reports on crop pollination are frequently arising, pollination research seems still at the beginning and our knowledge on crop pollination by bees at various scales is scarce. This work aims to explore so far unknown benefits of bee pollination to highlight its overall importance. It is also focused on the main drivers of crop pollination by bees at different spatial scales, using strawberries as a study organism. The first part (chapter 2) explores the benefits of bee pollination on commercial value and post-harvest quality of strawberry fruits from different varieties. Exclusion experiments with bee, wind and self pollination treatments were conducted on nine strawberry varieties at an experimental strawberry field. Bee pollination strongly increased the commercial value of strawberry fruits across all varieties by producing well shaped fruits with higher weight. It further elongated the shelf live of strawberries from most varieties, which was calculated from firmness values. Bee pollinated fruits had a more intense red colour and lower sugar-acid-ratios in most varieties. In general, effects differed between varieties, but with mostly similar directions. These results give a positive reply on the questions wether bee polination benefits commercial value and post-harvest quality of strawberries and the differences between varieties.
The second part of this work (chapter 3) focused on the relationship between bee pollination and shelf life of strawberries in detail. From the first part (chapter 2), the number of fertilized achenes from fruits of the variety Yamaska was related to pollination treatments and commercial grades. In 2012, further fruits were harvested on a commercial strawberry field and divided into commercial grades in relation to malformations from insufficient pollination. The process of firmness, fruit weight and decay during storage was analysed in relation to commercial grades. During storage time, firmness and fruit weight were decreasing while the proportion of decayed fruits was increasing, independent from commercial grades. Commercial grades had a general effect on fruit degradation, with improved values in higher grades. Thus higher commercial grades resulting from improved pollination elongated the shelf life of strawberries.

Differences between varieties in part one (chapter 2) lead to the question whether strawberry varieties differ in their attraction to bees with consequences on their visitation rate (chapter 4). The influence of different volatile emissions between strawberry varieties, the bees’ responses on distinct compounds and resulting visitation rates on a commercial strawberry field were analysed. Three strawberry varieties were grown in a greenhouse and the flower volatile emissions from each variety as well as antennal responses of *Osmia bicornis* L. females on each compound were tested. Further the bees’ visitation rates on two varieties were counted at a commercial strawberry field. All strawberry varieties emitted the same flower volatile compounds, but differed in the quantities of most of them. Antennal reactions of *O. bicornis* females differed between most compounds and were higher than responses on controls. Under field conditions, the variety that produced higher quantities of the most attractive compounds was visited much more frequently.
Hence, the question about volatile emissions affecting the responses and visitation of solitary wild bees could be positively answered.

The last part of this work (chapter 5) was focused on the main drivers of various bee diversity parameters that explain the community composition of bees and their influence on strawberry fruit weight. As bee diversity is known to change related to landscape complexity, the study was conducted on commercial strawberry fields located in landscapes ranging in a gradient from homogenous to more heterogenous, to get ranging values of the bee diversity parameters. Bees were collected via sweep netting and strawberries were harvested at four field positions, that were located from field edge to center. In general, strawberry weight was strongly influenced by Evenness and to a lower level by Shannon-diversity of bee pollinators, whereas results from species richness and bee abundance were inconsistent. Higher Evenness and Shannon-diversity of the bee community improved strawberry weight at the field edge, whereas higher proportions of honeybees improved fruit weight at the field center, seemingly due to higher conspecific pollen loads. Evenness was negatively correlated to the proportion of honeybees. The research questions could be answered with Evenness being most influential on strawberry fruit weight, with contrasting effects between field locations, driven by the proportion of honeybees.

In conclusion, the commercial value and post-harvest quality of strawberry fruits was strongly dependent on bee pollination, which was generally conducted by both honeybees and wild bees, but further affected by diverse scale-dependent effects. The strong relationship between fruit quantity and quality emphasizes, that pollination countervails to increasing global demands on high quality food. However, against the background of serious pollinator declines due to increasing anthropogenic impacts on various scales, pollination appears to be an ecosystem
service that is strongly endangered, which may have extensive impacts on world food security. Hence, the results of the current work emphasize, that international conservation strategies will become more efficient by focusing on a broad scale of facets affecting crop pollination to maintain this highly important ecosystem service for the future.
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