

RESEARCH ARTICLE

Possible mechanisms of pollination failure in hybrid carrot seed and implications for industry in a changing climate

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Abstract

Approximately one-third of our food globally comes from insect-pollinated crops. The dependence on pollinators has been linked to yield instability, which could potentially become worse in a changing climate. Insect-pollinated crops produced via hybrid breeding (20% of fruit and vegetable production globally) are especially at risk as they are even more reliant on pollinators than open-pollinated plants. We already observe a wide range of fruit and seed yields between different cultivars of the same crop species, and it is unknown how existing variation will be affected in a changing climate. In this study, we examined how three hybrid carrot varieties with differential performance in the field responded to three temperature regimes (cooler than the historical average, average, and warmer than the historical average). We tested how temperature affected the plants’ ability to set seed (seed set, pollen viability) as well as attract pollinators (nectar composition, floral volatiles). We found that there were significant intrinsic differences in nectar phenolics, pollen viability, and seed set between the carrot varieties, and that higher temperatures did not exaggerate those differences. However, elevated temperature did negatively affect several characteristics relating to the attraction and reward of pollinators (lower volatile production and higher nectar sugar concentration) across all varieties, which may decrease the attractiveness of this already pollinator-limited crop. Given existing predictions of lower pollinator populations in a warmer climate, reduced attractiveness would add yet another challenge to future food production.

Introduction

Insect-pollinated crops comprise approximately one-third of the global food supply [1]. Many of these plants owe their present uniformity [2], disease resistance [2], and high yields [3–6] to hybrid production systems, including carrot, tomato, onion, melons, squash, brassicas, and eggplant [7]—together totaling nearly 20% of global crop production [8]. Because these production systems rely on crossing two parent lines, one of which is rendered male-sterile by

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hand-emasculature or genetic techniques, they are even more reliant on insect pollinators than their open-pollinated counterparts, do not require insects to cross from one parent line to the other [9–12]. Global reports of declines in many pollinator communities [13; 14], changing climate shifting pollinating insects' active time away from peak bloom [15], and that pollinator reliance has been linked with reduced yield stability [16], indicate that hybrid systems may be at greater risk from additional disturbances than open-pollinated systems.

Yields of any given hybrid crop can vary significantly between varieties and also from year to year. In itself, this generates economic uncertainty, but it also makes determining which cultivars are best suited to the changing environment a difficult task. When hybrid crops that are grown for seed fail to produce adequate yields, pollinators are often blamed [10], particularly as male-sterile plants, which have no pollen reward, are notoriously unattractive to honey bees [9; 12; 17]. However, it is unclear if the observed poor yields are due to the lack of pollen, other characteristics that have inadvertently been selected for during the process of crop breeding and selection, or a combination of plant traits and environmental variables.

Traits that affect the yield of insect-pollinated plants can broadly be placed into two categories: plant fertility and plant attractiveness. Characters surrounding plant fertility are intrinsic to the plant, such that increased pollinator activity will not improve yield. For example, if flowering of the male-fertile and male-sterile lines is poorly synchronized, yields can suffer as the cross-pollination window may not overlap sufficiently. In such cases, cultural measures to synchronize the lines even by a few days can substantially increase yield [18]. Pollen viability has been a recurrent problem in hybrid crops as well, with inbreeding depression often causing pollen viability to drop to 50% or less before it ever leaves the flower [19–21]. This initial difficulty can be further exacerbated by heat and water stress during critical periods of plant development, which can further degrade pollen fertility [20; 22]. Female receptivity is also important, as flowers that are too young, too old, or damaged by suboptimal temperatures will set little seed even if pollen viability is high [23].

Characters surrounding plant attractiveness, in contrast, are those which influence pollinator visitation. The quality of the nectar reward is foremost for ensuring return visits [24–26]. Honey bees prefer nectar rewards between 30 and 50% sugar w/w [25], but lower concentration sources at higher volumes per flower may be chosen over many low-volume, high-concentration sources [25]. Nectar production is altered in high-temperature conditions, typically resulting in lower volumes and higher concentrations [25], but hot conditions can also alter the production of secondary compounds, such as phenolics [27], which can result in a very different flavor palette to potential insect visitors. Volatile compounds emitted by the plant are also important, as they play a key role in attracting pollinators to the flower initially [25]. Although the volatiles comprising the floral bouquet of numerous plant species have been cataloged, little is known about how these compounds are perceived by pollinators, and less still is known about how they respond to changes in temperature, individually or in aggregate.

In order to achieve successful pollination, a plant must be both fertile (able to receive pollen and set seed), and attractive to pollinators. To understand how climate change may affect pollination, we must therefore look at factors relating to both overarching categories. To address these broad questions, we chose to focus on hybrid carrot production because, despite being a generalist flower pollinated by hundreds of insect species [28–31], hybrid carrot is known for its low seed set [32] and lack of pollinator attractiveness [10; 33]. In addition, seed production for carrot occurs in areas not optimal for carrot growth in order to avoid genetic contamination from wild carrot [34; 35], which can readily cross into cultivated varieties and reduce agronomic quality of progeny [34; 35]. As a number of major carrot seed producing regions are located in temperate, semi-arid areas [36–38], additional temperature variability may negatively affect seed production. The combination of environmental stress and poor pollination in

present-day hybrid carrot make it a promising model for future conditions experienced by hybrid crops, and examining the mechanisms of current pollination failure may highlight future vulnerabilities both carrot and hybrid crops in general.

The objectives of this study were to 1) test the effect of temperature on temporal patterns of plant traits that might predict performance in the field, including bloom phenology, seed set, pollen viability, nectar quality (both sugars and phenolic compounds), and floral volatiles and 2) examine if these effects differ across varieties with a range of historical yields to test whether varietal differences are reduced or exaggerated by warming, and 3) use this information to determine which factors are important in present-day pollination failure, and which may be important given a changing climate.

Methods

This study was conducted in New Zealand as it is one of the world's largest producers of carrot seed [39]. We exposed carrot (*Daucus carota* L) plants to experimental temperature treatments, and measured characteristics relating to their innate ability to produce seed ('Plant Fertility Metrics', below) as well as several metrics that may affect their attractiveness to pollinators in the field ('Plant Attractiveness Metrics', below). To assess the extent to which these characters contribute to differences in yield and how they may respond to climate warming, we examined the correlations between each one and plant variety, temperature, and time-of-day in generalized linear mixed-effects models (GLMMs), generalized additive mixed-effects models (GAMMs) or ordination-based tests.

Plant material

In order to determine how floral receptivity and pollen viability vary with time-of-day (a known source of variation in floral traits [25; 40]) and temperature, we grew the male-fertile and cytoplasmically male-sterile (brown anther type) parents of three lines of Nantes-type hybrid carrot for hand-pollination trials and measures of pollen viability. These three lines had previously been observed to perform poorly (172 ± 43 kg/ha), average (377 ± 17 kg/ha), and well (607 ± 87 kg/ha) in the field (hereafter referred to as 'poor', 'average' and 'excellent' lines); we chose this gradient to attempt to tease out the cause(s) of the differential performance in the field, and to have a range of yields to assess the effects of temperature. Seeds for each line were sown in trays in February 2015, during the southern hemisphere summer. When plants had germinated, we transplanted them individually into 3L pots filled with potting mix and slow-release fertilizer (Canterbury Landscape Supplies). Each line had 100 male-sterile plants and 75 male-fertile plants potted out, and these were kept outdoors in ambient conditions until flowering began.

In order to minimize the effect of temperature on plant physiological processes other than flowering, we moved plants to temperature treatments after the umbels had formed, and just prior to flower opening. The three temperature treatments simulated cool, average, and warm seasons via shade houses, unheated glasshouses and heated glasshouses, respectively. Two separate buildings were used for each temperature treatment, and the plants were equally divided between them. To accurately record conditions experienced by the carrot flowers, we placed temperature and relative humidity probes (onset HOBO Prov2 temp/RH meters) at chest height in each of the six locations. Temperatures were extracted from the data loggers and recorded as temperature at the sample time (S2 Fig), average temperature during the 24-hour period prior to the sample time, and average temperature during each plant's time in the glasshouse prior to sampling. Models were run with each measure of temperature sequentially, but in all cases, the 24-hour period had the highest predictive value, so only this measure was used.

Every day, we checked plants for floral stage, and, when the petals had turned white and the outer whorl of flowers had just begun to open, they were randomly assigned to a temperature treatment. Male-sterile plants were also assigned to one of seven time treatments (4am, 8am, 11am, 2pm, 5pm, 8pm, and 11pm) for hand-pollination. Selection of treatments was done without replacement, so that male-sterile plants were always equally distributed between the three temperature treatments and each time-temperature combination received the first replicate before proceeding to the second, third, and fourth (252 male-sterile plants total). Male-fertile plants were preferentially assigned to locations where male-sterile plants required pollination.

Once selected, we bagged the primary umbel of each male-sterile plant with 1mm mesh to prevent insect pollination and, as an extra precaution, placed plants into a 1.5m³ fine mesh cage in each of the six locations. Male-fertile plants were left unbagged inside the cage. As a further precaution, we placed yellow sticky cards in each cage to trap any flying insects that entered.

Plant Fertility Metrics

Phenology. In order to properly hybridize in the field, both the male-sterile and male-fertile lines must bloom simultaneously, and the male-fertile lines should, ideally, produce pollen for the duration of the male-sterile line's bloom time. To quantify this bloom synchrony, we checked the primary umbel of each potted carrot plant daily. Just prior to the opening of the outer whorl of umbellets, we recorded the date and assigned the plant to its treatment. We then calculated the number of days between seed sowing and flowering. Any plants that had not flowered after 365 days were recorded as having failed to bloom.

Seed set. To quantify changes in stigma receptivity across lines, temperature treatments, and at different times of day, we conducted a hand pollination experiment. Once pollen from male-fertile plants started to dehisce, hand-pollinations began. Every morning at 8am, we surveyed caged plants and any plant where the stigmas appeared receptive throughout the umbel was pollinated that day at its pre-selected time slot. To assist with visually identifying receptive stigmas, a photo guide was prepared the season prior by staining stigmas with alpha-naphthyl acetate [41]. The primary umbel of each plant to be pollinated was unbagged and three umbellets were selected and tagged, one from each of the three whorls, as previous studies indicate that there may be differences in female fertility between the inner and outer umbellets [42]. As the 'medium' male-fertile line was the only one blooming throughout the sample period, we used pollen from this line for hand-pollinations of all male-sterile lines. For each time slot where a flower needed to be pollinated, we bulked together pollen from all the 'medium' male-fertile plants in the building. We took a subsample of this pollen, put it in a cryotube, immediately placed it in liquid nitrogen for later pollen viability assessment, and applied the remainder with a paintbrush to each stigma of each floret of the tagged umbellets of all flowers in the building needing pollination at that time. Plants were then rebagged and left in the glasshouse for a further 72 hours, to allow pollen tubes to reach the ovaries (typically 24–48 hours [10]), before being brought back outside to complete seed set.

Once the seed heads dried, we brought them back into the lab, and the seeds of the three tagged umbellets of each flower were counted. In addition, three untagged umbellets, one from each whorl, were examined as an unpollinated control for each plant.

Pollen viability. Subsamples of bulked pollen were stored in cryotubes in liquid nitrogen until the final pollination for each day (typically 11pm or 4am), when we transferred it to a -80°C freezer. When all pollen samples were collected, we transferred them to a second facility on dry ice and then immediately placed them in a second -80°C freezer until processing.

Pollen viability was assessed with fluorescein diacetate (FDA), which has previously been shown to have a strong correlation with *in-vivo* germination in carrot [43]. We thawed the cryotubes for 2–5 minutes and then washed the tube with 50 μ l FDA-sucrose solution (0.25% w/v FDA, 20% w/v sucrose) via a pipettor, with as much liquid as possible collected and slide mounted. We examined samples with a UV light microscope, counting 200 pollen grains across longitudinal transects of each slide. We categorized pollen as viable if it fluoresced bright green [44].

Plant Attractiveness Metrics

Nectar quality. After volatile collection (if applicable, see below), but prior to pollination, we sampled each male-sterile plant for nectar. We followed the protocol described by Gaffney [31], dipping half of each umbel (~30 umbellets, with the umbel diameter being controlled for in analyses) into 40mL of distilled water twenty times, ensuring that the umbel was shaken off afterward to recover as much water as possible. We then immediately placed the dilute nectar in a freezer until further processing. To prepare the samples for high-performance liquid chromatography (HPLC), they were thawed, filtered to remove any large contaminants, and freeze-dried in 50mL falcon tubes. We then resuspended the samples in 1mL of methanol:water at a ratio of 1:1 and divided it in two parts: a 600 μ L aliquot for nectar sugar analysis and a 400 μ L aliquot for nectar phenolic analysis.

Sugar identifications were made via HPLC using a modified combination of the methods of Ruperez [45], Knudsen [46], and Knudsen and Li [47]. The 600 μ L aliquot was centrifuged at 14,000 rpm for 10 minutes. A 250 μ l aliquot of the supernatant was placed directly into an HPLC vial. We then carried out the HPLC analysis using a refractive index (RI) detector (Waters™ Alliance 2690 HPLC with Waters™ 2414 RI detector). HPLC-RI analysis was carried out by injecting 10 μ l of sample into an isocratic mobile phase of 70% acetonitrile in water with chromatographic separation (Econosphere™, Amino, 5 μ m, 4.6 x 250mm, Grace™) at 30°C and RI detection at 40°C. Unknown sugars were identified using retention times and response factors of known sugars (Sigma).

Phenolic composition of the nectar was also analyzed via combined liquid chromatography-mass spectrometry (LC-MS). The previously prepared 400 μ l aliquots were filtered with a Single Step[®] vial 0.22 μ m PVDF (Thompson™ Part No. 65531–200) filter. The LC-MS system consisted of a Thermo Electron Corporation (San Jose, CA, USA) Accela UHPLC pump, Thermo Accela Open Auto sampler (PAL HTC-xt with DLW), Finnigan Surveyor PDA plus detector and a ThermoSphere TS-130 column heater (Phenomenex, Torrance, CA, USA). Each of the 48 extracts was analyzed by two ion formation modes creating 96 data files, as follows. A 2 μ L aliquot of each prepared extract was separated with a mobile phase consisting of 0.1% formic acid in water (A) and 0.1% formic acid in acetonitrile (B) by reverse phase chromatography (Kinetex guard cartridge and Kinetex C18, 2.6 μ m, 100 Å, 100 x 2.1 mm, Phenomenex, Torrance, CA, USA) maintained at 30°C with a flow rate of 200 μ l/min. A gradient was applied: as 0–10 min/95%A, 13 min/60%A, 15–20min/5%A, 23–28min/95%A. The eluent was scanned by API-MS (LTQ, 2D linear ion-trap, Thermo-Finnigan, San Jose, CA, USA) with electrospray ionisation (ESI) in the negative mode. Data were acquired for precursor masses from m/z 120–1000 with up to MS3 product spectral tree formation. All data were processed with the aid of Xcalibur[®] 2.20 (Thermo Electron Corporation) and an in-house Plant and Food Research database of chemical signatures.

Floral volatiles. We collected volatile organic compounds (VOCs) from the headspace of plants after the outer whorl of florets opened, but prior to pollination. Due to resource limitations, we could not collect volatiles from every plant, so a subsample was taken across the

treatments. To achieve a good cross-section of the experimental treatments, we collected three separate datasets; one across varieties, one across temperatures, and one across times of day. For the variety dataset, we took 24-hour headspace collections for twelve plants (6 male-sterile, 6 male-fertile) from each of the three varieties in the average temperature treatment, totaling 36 samples. For the temperature dataset, we took 24-hour headspace collections from 6 'medium' male-sterile plants each in the cool and hot treatments, which were analyzed together with the 'medium' male-sterile samples in the previous dataset, totaling 18 samples (12 unique to this dataset). For the final dataset, in order to capture variation throughout the course of the day, we sampled six further 'medium' plants beginning at each of the seven time periods (3–5 hour headspace collection), for 42 total time-of-day samples.

Each headspace sample was collected *in situ* using the active sampling apparatus in Fig 1. The primary umbel of each flower was fitted with a nylon oven bag and, insofar as it was possible, leaf material was excluded from the bag. In order to ensure floral volatiles rather than green leaf or ambient compounds in the air were being collected, each set of collections included a control where the bag was secured around a leaf. Each bag was fitted with a charcoal filter at the base to remove ambient VOCs. We used a pump with an airflow rate of 500mL/min, split four ways so that 125mL/min of air was pulled through each headspace collection apparatus and into a Tenax[®] filter, which adsorbed the floral VOCs. The Tenax[®] filter was constructed from a 15mm long, 10mm diameter glass tube containing 60mg of Tenax[®] 35/60 (Grace Davidson Discovery Sciences, VIC, Australia) held in place with silane-treated fiberglass (Grace Davidson Discovery Sciences, VIC, Australia). Tenax tubes were conditioned prior to use by heating for 3 hours at 250°C under a stream of nitrogen gas, and the charcoal filters were baked overnight at 150°C in a filtered-air oven. After VOC collection, each tenax was desorbed by solvent extraction with 1mL of n-hexane (Sigma-Aldrich, 99% purity).

To obtain quantitative values of each VOC identified in the headspace samples, we added an internal standard of nonadecane (10 µg) to the 1 mL of n-hexane (Sigma-Aldrich, 99% purity) used to elute the tenax tubes. A sample containing 10 µg/mL of each of the identified compounds was run as an external standard.

We used one microliter of each headspace extract for gas chromatography coupled with mass-spectrometry (GC/MS). The subsample was injected into a Varian 3800 gas chromatograph (Varian Walnut Creek, CA, USA) with the injector port set at 250°C, and then run through a DB5-MS non-polar column (J&W Scientific Folsom, CA, USA) with dimensions 30m x 0.25mm id x 0.25µm film thickness. The column was raised from 40°C up to 280°C at a rate of 4°C/min, and then held at 280°C for 5 minutes. We used a constant flow of helium as a carrier gas (1 mL/min). Injections were splitless for 36 seconds. A Saturn 2200 mass spectrometer (MS, Varian Walnut Creek, CA, USA) ionized the molecules from a mass range of 29 m/z to 399 m/z, after the GC separated each compound present in the extract according to their volatility. The abundance of each compound was then recorded as the area under each peak. After each sample was run, we identified compounds by comparing MS results to a database (NIST MS Search 2.2). Synthetic compounds were injected with the same GC-MS protocol to confirm identification.

To compare the compounds' abundance in each extract, we used the internal standard method to calculate quantities, multiplying the known amount of the internal standard nonadecane (10 ng) by the area of the compound of interest and the response factor (measured from an external standard run containing a known quantity, 10 ng, of both nonadecane and the compound of interest), then dividing by the area of the internal standard. Volatile collections were done over a different time period for each treatment (variety: 24 hours, time: 3-5h), so for comparison of each compound across all the experiments, we calculated the emission

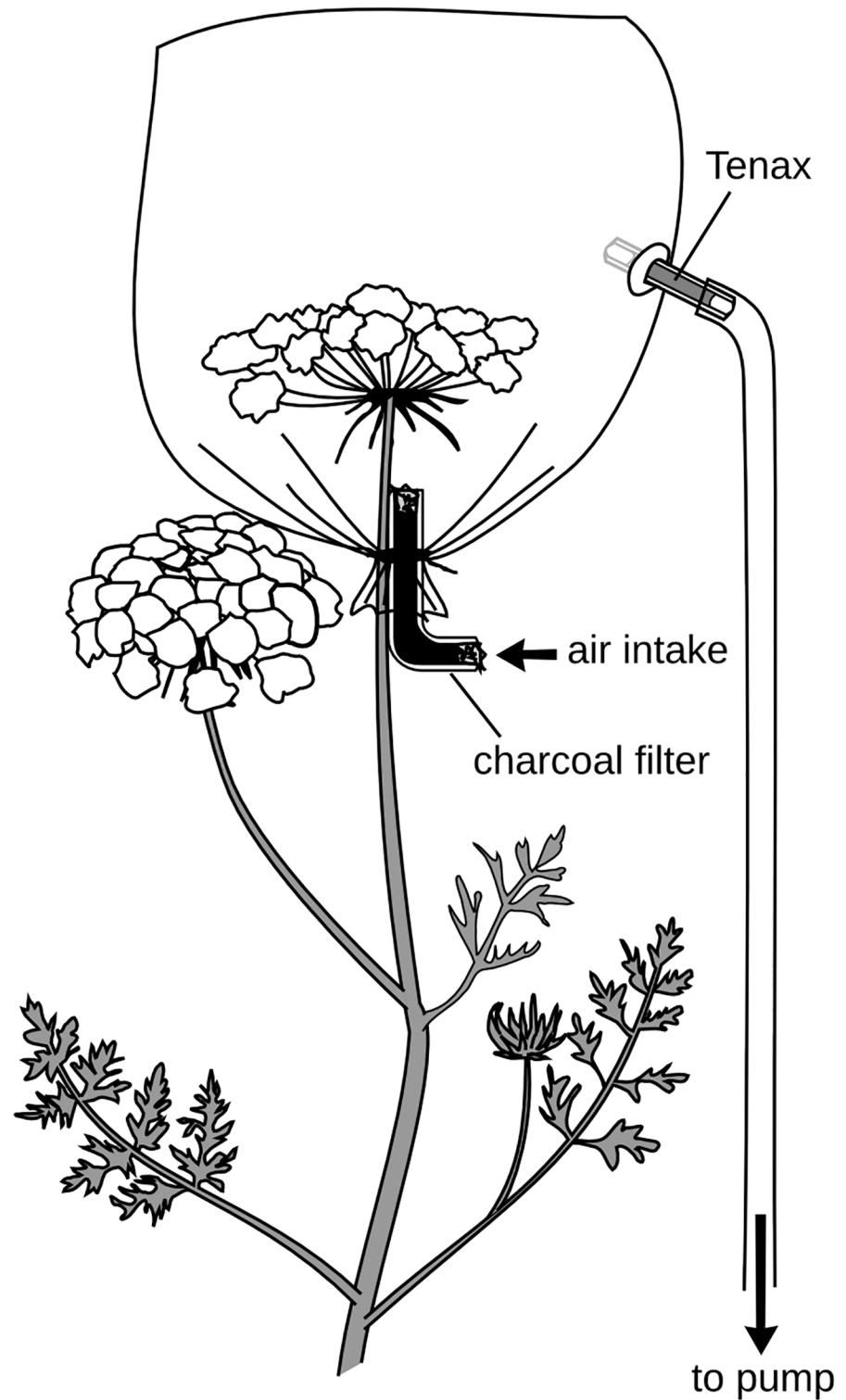


Fig 1. Apparatus for collecting carrot headspace volatiles.

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rate by dividing the quantity of each peak by the amount of time of collection in order to obtain value in $\mu\text{g}/\text{h}$.

Statistical analysis

Phenology data were recorded as a count variable—the number of days between seed sowing and flowering. Because all plants were kept under the same conditions prior to blooming, we could compare them in a generalized linear model (GLM). We used a gamma error distribution to account for the observed variance-mean relationship. The predictor variables were plant variety and plant line.

Nectar, pollen viability, and seed set data were collected over multiple days, creating the risk of temporal autocorrelation. As the temperature treatments were conducted in two locations each, there was also the potential for spatial autocorrelation within each glasshouse. In order to account for this variance and non-independence, we used generalized linear mixed-effect models (GLMMs) with date sampled and location as crossed random effects. We used the *lme4* library [48] in the R statistical programming language [49] to perform most of our analyses. We used the following model selection process to determine our best-fitting model. All permutations of the predictor variables (plant variety, temperature, and time-of-day, and their interactions) were used to create candidate models, and final models were selected if their AIC scores were within two points of the best-fit model; if multiple models met this criteria, we took a model average [50] using the *MuMIn* package 1.15–6 [51]. In order to obtain p-values for the final models, we used the Satterthwaite method of denominator synthesis, implemented within the *lmerTest* package [52]. It should be noted that this method calculates non-integer degrees of freedom. Where relevant, models were checked for over-dispersion (where error distributions were not Gaussian) or normality of residuals and homoscedasticity (for Gaussian models). In addition to differences in the mean response across treatments, initial examination of the data suggested that there may be differences in the variance of the response variables. To test for these differences, we used Levene's test.

Seed set data were recorded as a count variable, as it was not possible to count the number of initial florets once the seed heads dried to establish a proportion. Plants that failed to set seeds introduced numerous zeros to the data set. While not over-dispersed, the dataset did not conform well to the Poisson distribution and so was analyzed with a zero-inflated negative binomial GLMM in R, with the *glmmADMB* package [53]. Predictor variables in the initial model were the pollen viability (as a covariate to control for the quality of pollen used in each hand-pollination), temperature at the time of pollination, plant variety, and time of day.

Pollen viability data were recorded as a binomial variable, where individual pollen grains were either viable or not. The proportion of viable and inviable pollen grains were tested in a GLMM with binomial errors and a logit link function. Because pollen stored at -80°C slowly loses viability over time [54; 55], we included the number of days between collection and processing as a fixed covariate in final model to account for the between-sample variation in storage time. Other predictor variables in the initial model were the plant variety and temperature and time of day at the time of pollen harvest. As this model was significantly over-dispersed, we also included individual sample as a random effect [56].

Nectar sugars were measured as the concentration of fructose and glucose; no sucrose was found. As the concentrations of the two sugars were tightly correlated ($R^2 = 0.983$), only glucose was examined in a GLMM, with a gamma error distribution to account for the observed variance-mean relationship. Flowers varied in umbel diameter, which could influence the amount of nectar collected with the dipping methodology. Therefore, we controlled for flower size by adding the diameter of the umbel as a fixed covariate. Other predictor variables in the

initial model were temperature at the time of nectar collection, plant variety, and time of day nectar samples were taken.

Nectar phenolics were expressed as concentrations. Three phenolic compounds were found in carrot nectar: caffeic acid, coumaric acid, and ferulic acid. As initial plots revealed that each compound reacted differently to different conditions, it was necessary to examine all three. To avoid multiple single-response models and to capture shifts in the combined phenolic bouquet, we conducted a nonmetric multidimensional scaling (NMDS) ordination within the R package *vegan* [57] with Bray-Curtis dissimilarity to account for the large differences in mean concentration. To test whether the three compounds varied across variety, temperature, and time of day, we used a permutation multivariate ANOVA (PERMANOVA) procedure (function 'adonis') from the same package.

Floral volatiles were expressed as the concentrations of methyl salicylate, nonanal, and phenylacetaldehyde—the three compounds found in carrot floral volatiles which bees are able to sense (measured in previous work as a consistent electroantennograph response; S1 Fig). To account for the three compounds simultaneously, we ran a NMDS ordination for each of the volatile datasets with variety, temperature, and time of day as predictor variables. As with the phenolic data, we used a PERMANOVA procedure to test for significance. To aid with interpretation of these multivariate results, we then conducted univariate analyses on each volatile. Data exploration revealed that there was a non-linear relationship between time-of-day and volatile concentration, so we conducted a generalized additive mixed-effects model (GAMM), which allows for non-linear relationships between predictor and response variables [58]. In the GAMM, time sampled was a smooth term, while plant ID was a random effect. The amount of smoothing was determined in the model using maximum-likelihood within the *gamm4* package [59].

Results

We found numerous effects of plant variety, temperature, and time-of-day on measures of both plant fertility and attractiveness to pollinators (see Fig 2 for a summary).

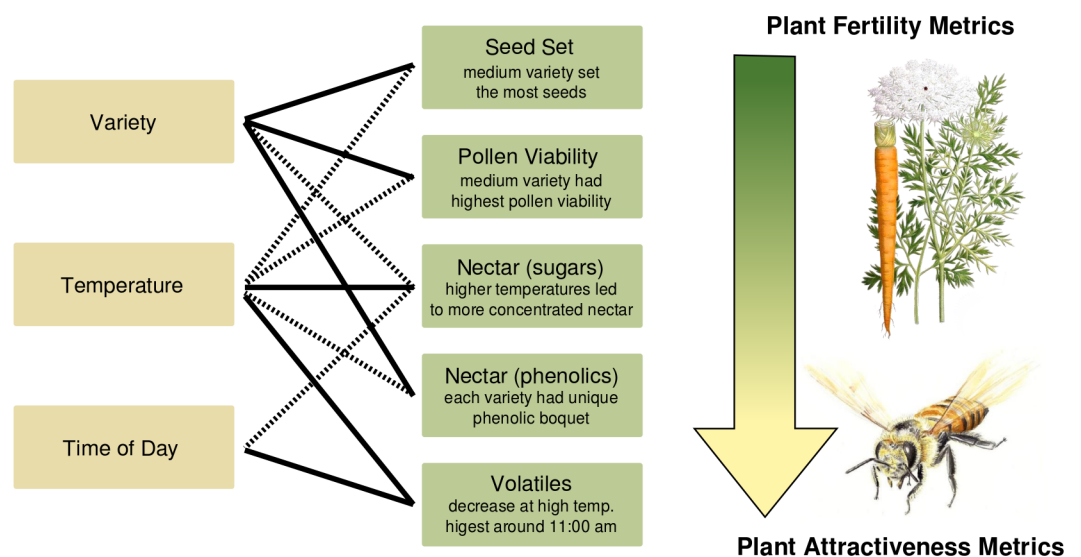


Fig 2. Relationships between different factors examined in this study. Solid lines indicate a statistically significant relationship. Dotted lines indicate factors that were conserved in the final selected models, but were not statistically significant. We did not find any significant interaction effects.

<https://doi.org/10.1371/journal.pone.0180215.g002>

Plant Fertility Metrics

Phenology. A number of plants failed to flower at all: 5% of male-sterile and 22% of male-fertile carrots did not send up a flowering stalk after one calendar year. The rate for male-fertile plants was heavily influenced by the poor line, which accounted for 76% of failures (51% of these plants did not flower). Of the plants that did bloom, there was considerable spread in flowering time, with the first plant blooming on day 285 and the last on day 341. Male-fertile lines bloomed significantly later than male-sterile lines, and there was an interactive effect between variety and line, with the poor variety having the widest gap between the bloom time of the male fertile and male-sterile lines (Fig 3, Table 1). The male-sterile excellent variety had the most tightly grouped flowering time ($P < 0.001$; $F = 8.560$; Levene’s test for differences in variance)

Seed set. The rate of seed set was low, with fewer than half of the hand-pollinated umbels setting seed. This may have been due to poor weather, poor pollen viability, and, potentially, the presence of the brown shield bug (*Dictyotus caenosus* (Westwood), Hemiptera:Heteroptera), which was able to enter the seed heads through the exclusion mesh. Nearly every mesh pollinator exclusion bag contained at least one *D. caenosus*, and though, to our knowledge, there is no record of *D. caenosus* feeding on carrot seed, we cannot exclude the possibility that this generalist plant-feeder used the seed heads as a food source. In total, less than half of the hand-pollinated umbels set seed. Despite the overall low seed set, we still found differences between the carrot varieties, such that the medium-performing variety set significantly more seed than the other two varieties (Fig 4), with an average of 1 additional seed per three umbellets ($P = 0.005$; $z = 2.788$; GLMM). Temperature ($P = 0.995$; $z = 0.010$; GLMM) and proportion of viable pollen ($P = 0.383$; $z = 0.872$; GLMM) were both retained in the best-fitting model,

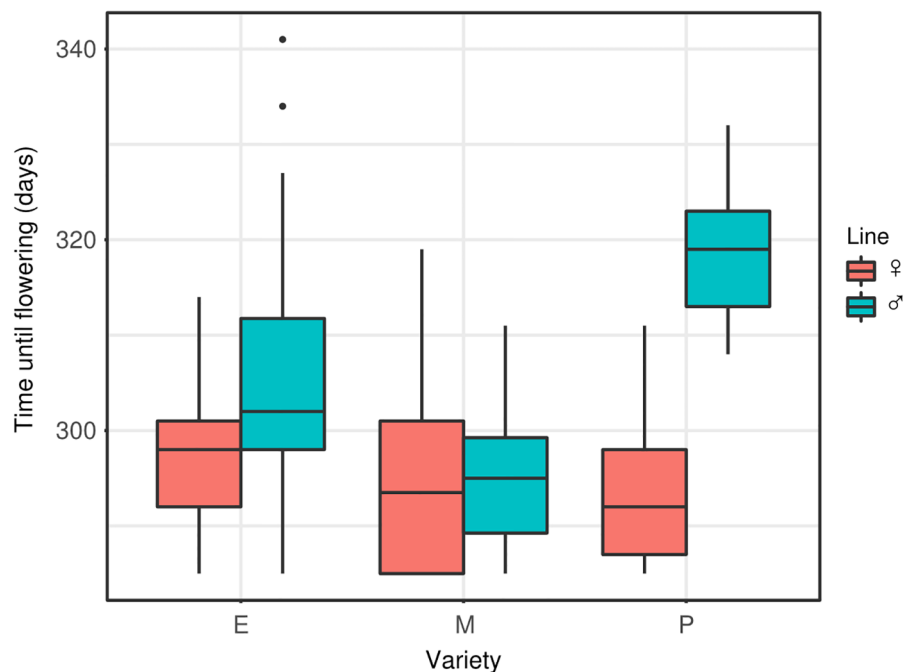


Fig 3. Number of days between seed sowing and flowering between the three carrot varieties. Variety: E = excellent, M = medium, P = poor. Line: male sterile (♀) and male fertile (♂) lines broken out for each variety. Data ceased being collected at 365 days. Boxes represent the middle 50% of the data, lines within boxes are the median.

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Table 1. Coefficients table of GLM for carrot phenology. Variety: E = excellent, M = medium, P = poor. Line: male sterile (♀) and male fertile (♂). The intercept condition is the male-sterile, excellent line.

	Estimate	SE	t statistic	P value
intercept	3.373×10^{-3}	8.855×10^{-5}	380.908	< 0.001 ***
Variety (M)	2.675×10^{-5}	1.274×10^{-5}	2.099	0.036 *
Variety (P)	4.577×10^{-5}	1.267×10^{-5}	3.611	< 0.001 ***
Line (♂)	-9.521×10^{-5}	1.345×10^{-5}	-7.076	< 0.001 ***
Variety (M): Line (♂)	8.971×10^{-5}	1.945×10^{-5}	4.612	< 0.001 ***
Variety (P): Line (♂)	-1.783×10^{-4}	2.102×10^{-5}	-8.483	< 0.001 ***

Significance codes:

* < 0.05,

** < 0.01

*** < 0.001

<https://doi.org/10.1371/journal.pone.0180215.t001>

though neither was a significant predictor of seed set. The medium variety was also more variable in seed set than the other two varieties ($P = 0.001$; Levene’s test), as would be expected for count data, where the variance often increases with increasing means.

Pollen viability. Overall pollen viability was low, with a median of 14.5%. This is comparable to previous estimations of carrot pollen viability in New Zealand (S1 Text), but is low compared to the viability of cultivated carrot pollen elsewhere in the world [21; 43; 60], and much lower than wild carrot pollen (~80%) [35]. Temperature was retained in the final model, but was not a significant predictor of viability ($P = 0.825$; $z = 0.22$; GLMM). There was a significant difference in pollen viability between varieties (Fig 5; $P = 0.004$; $z = 2.866$; GLMM), with the medium variety being the highest, about 45% higher than either of the other two (observed mean of 21.2% versus 14.3% for excellent and 15.9% for poor). The medium variety was also more variable than either the poor- or excellent-performing varieties ($P = 0.023$; Levene’s test).

Plant Attractiveness Metrics

Nectar quality. Glucose and fructose were found in a close to 1:1 ratio (1.019:1; $R^2 = 0.983$; LM). No sucrose was detected. There was a significant effect of temperature ($P = 0.002$; $t = 3.091$; GLMM) on glucose concentrations, with increasing temperature being correlated with higher concentrations of sugars (Fig 6). Time-of-day, variety, and an interactive effect between time-of-day and variety were all retained in the final model, though none of them were significant predictors of sugar concentration. For phenolic compounds in the nectar, each of the three varieties had a different composition ($P = 0.012$; $F = 3.393$; PERMANOVA), with the excellent variety having high concentrations of caffeic acid, moderate concentrations of coumaric acid, and low concentrations of ferulic acid; the medium variety had low concentrations of caffeic acid, moderate concentrations of coumaric acid and moderate concentrations of ferulic acid; the poor variety had low concentrations of caffeic acid, high concentrations of coumaric acid and moderate concentrations of ferulic acid (Fig 7). The excellent variety was more variable in its concentration of caffeic acid than the other two varieties ($P = 0.017$; Levene’s test). Temperature was not a significant predictor of nectar phenolic composition ($P = 0.080$; $F = 2.566$), but was retained in the final model.

Floral volatiles. Of the three compounds contained in the male-sterile carrot flowers’ floral bouquets that honey bees are capable of sensing, only nonanal was present in all samples. There was no effect of variety on floral bouquet ($P = 0.671$; $F = 0.467$; PERMANOVA), but there was a significant effect of temperature ($P = 0.028$; $F = 4.743$; PERMANOVA), with higher

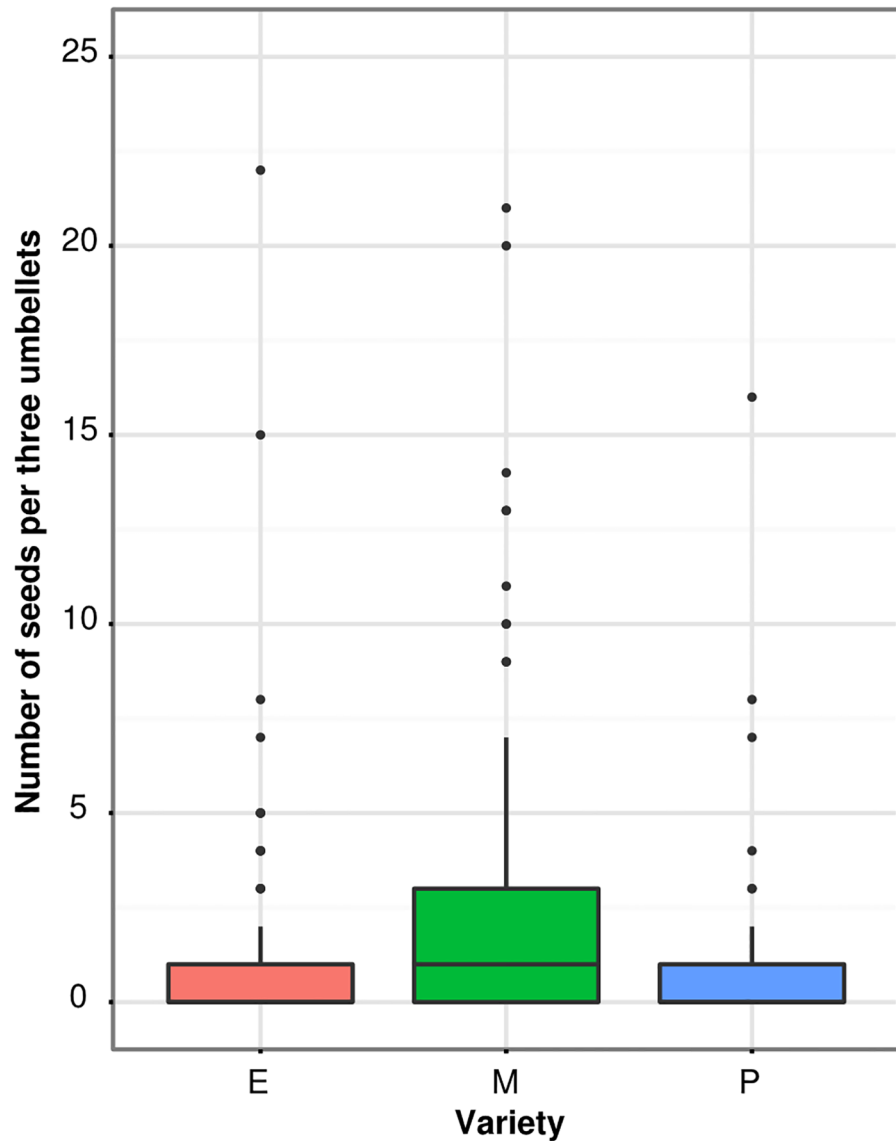


Fig 4. Seed set amongst the three carrot varieties. E = excellent, M = medium, P = poor. Boxes represent middle 50% of data, lines within boxes are the median.

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temperatures corresponding to lower volatile emissions. As plants were tracked through the course of a 24-hour day, there was a significant spike in nonanal concentration at around 11:00am (Fig 8; $P < 0.001$; $t = 17.460$; GAMM), just prior to the afternoon heat. Although there was no significant effect of variety, the medium variety was more variable than the other two ($P = 0.009$; Levene’s test), meaning that the significant time-of-day result despite this background variability, which was tested using the medium variety, is likely robust.

Discussion

While environmental conditions at flowering time affected some plant fertility and attractiveness measures, we did not find any interaction effects between plant variety and temperature.

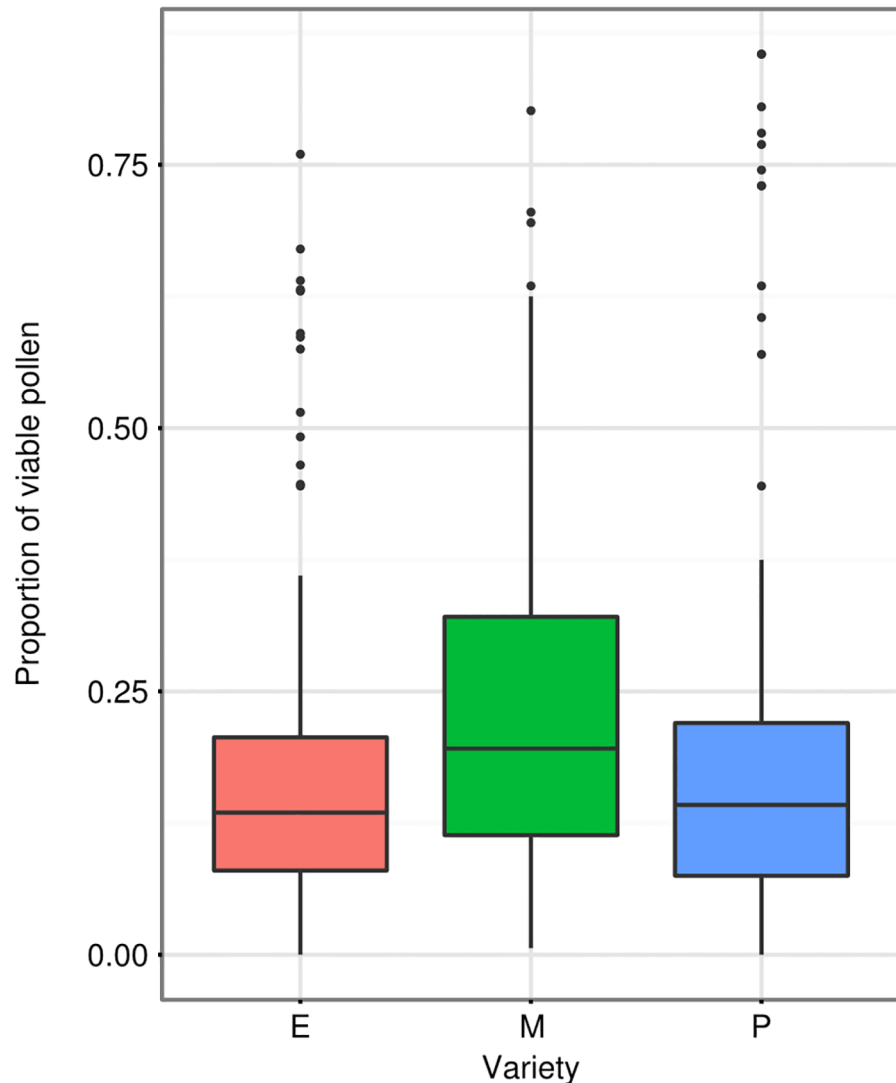


Fig 5. Pollen viability amongst the different varieties. Each sample point represents all male-fertile flowers with pollen at the time of sampling (evenly spread throughout the 7 time and 3 temperature treatment combinations). Viability was calculated as a proportion viable out of 200 grains. E = excellent, M = medium, P = poor. Boxes represent middle 50% of data, lines within boxes are the median.

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This implies that the differences observed between the poor-, medium-, and excellent-performing carrots were due largely to innate plant characteristics (Fig 2), which did not respond to temperature. Our data suggest numerous mechanisms of poor performance in the field, as the 'poor' variety consistently underperformed the other two: it bloomed late, 50% of the male-fertile plants failed to even initiate flowering, had the worst synchronization between male-sterile and male-fertile lines (Fig 3), relatively low seed set (Fig 4), and pollen viability typically below 20% (Fig 5). Additionally, the nectar phenolic profiles showed a wide gap between the varieties. The poor line's nectar was high in coumaric acid, which has been found to upregulate bee detoxification pathways [61], and ferulic acid (Fig 7), which, while commonly found in honey bee propolis [62], is thought to be an insect feeding deterrent [63].

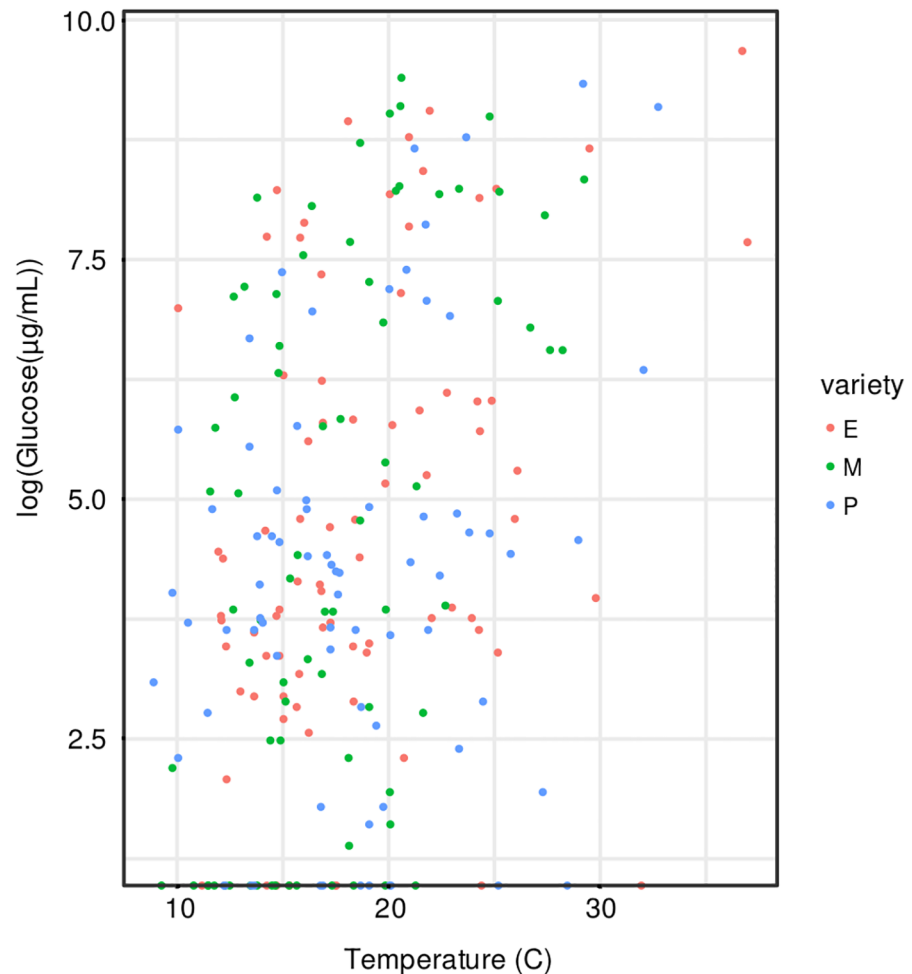


Fig 6. Log glucose concentration in nectar at different temperatures for the female lines of all three carrot varieties. E = excellent, M = medium, P = poor.

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Similarly, the poor line is low in caffeic acid, which is highly attractive to bees at modest concentrations [64].

Underperformance in multiple categories, from purely physiological characters to factors which influence plant attractiveness to pollinators, is a satisfying explanation for why a particular variety is observed to have low yield, but we have found that there is not a similarly easy explanation for the high yield of the best-performing variety. In fact, the excellent variety was observed to perform worse than the medium variety under controlled conditions: the excellent variety had somewhat worse synchronization between the male-sterile and male-fertile lines than the medium variety, and seed set and pollen viability on par with the poor variety. The excellent variety, however, distinguishes itself in its nectar phenolics, where it has high concentrations of the attractive caffeic acid, and is more consistent overall, with lower variances than the medium line in nearly every aspect, including bloom duration of the male-sterile line. Although there may be other differences that we did not capture, these two may result in increased pollinator attractiveness and better conditions for those pollinators to cross-pollinate the hybrid lines under field conditions.

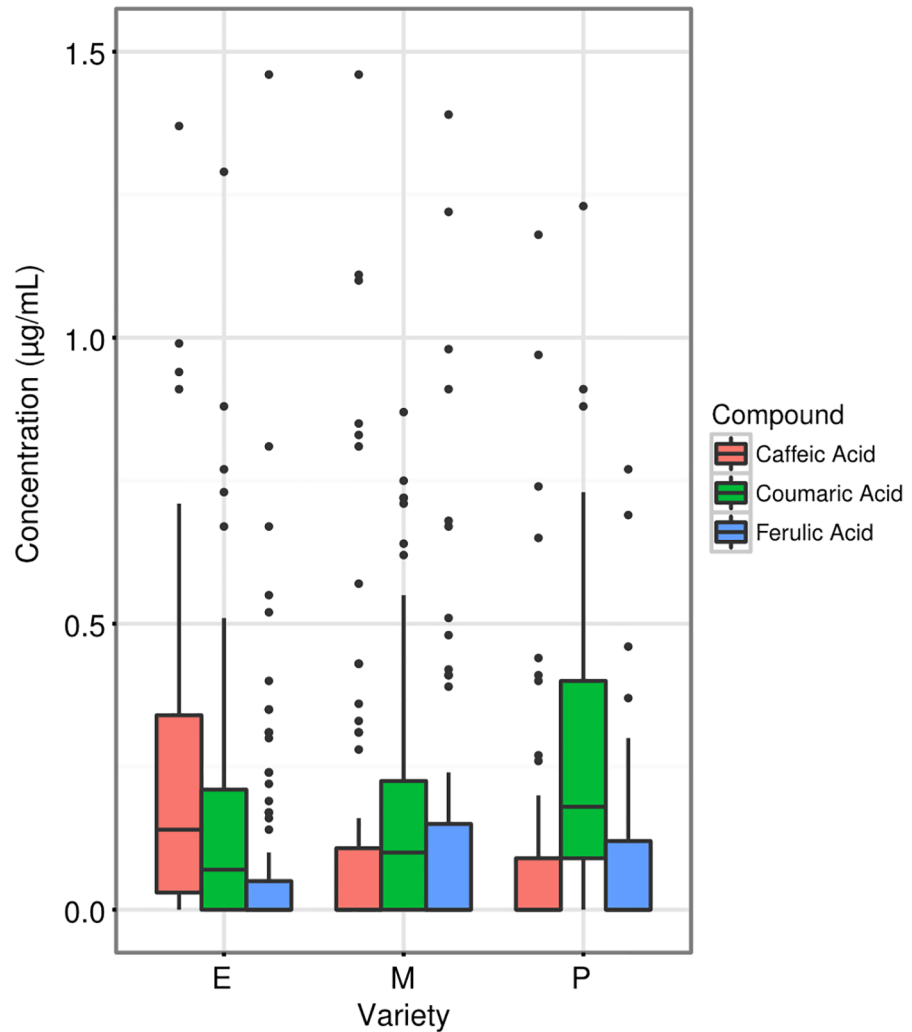


Fig 7. Concentrations of nectar phenolic compounds by carrot variety. E = excellent, M = medium, P = poor. Boxes represent middle 50% of data, lines within boxes are the median.

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Implications for industry

Our results are particularly important as previous studies have identified poor pollination as a major cause of low seed yields in hybrid carrot [18; 43; 65]. Varieties that are better able to attract pollinators may be better able to fare annual fluctuations in pollinator populations as they could draw in pollinators from the surrounding environment, a trend already observed in mass-flowering crops [66]. The importance of pollinator attractiveness for determining which varieties succeed or fail is magnified here by the very low pollen viability of these commercial hybrid carrot lines—typically less than 30%. If viability rates were closer to wild carrot (~80% [35]), the required pollen deposition would be reduced by ½, and thus pollination could be achieved with fewer insect visits and this would reduce the effect of differential attractiveness between the varieties.

We might join the numerous other authors that encourage the breeding of crops for increased insect attractiveness [10; 18], or higher capacity for seed production [67], or suggest that pollen viability be selected for [43]. However, it is important to keep in mind that

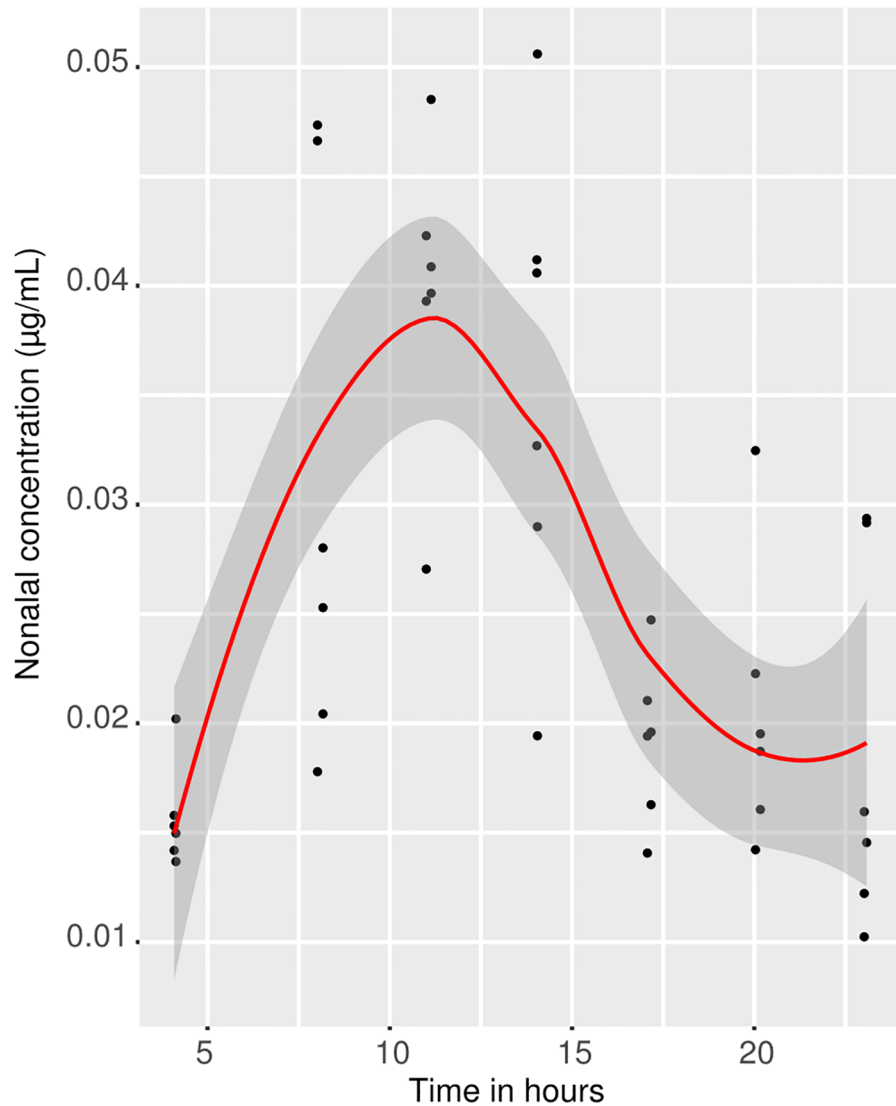


Fig 8. Concentration of nonanal versus time-of-day. Values are from six 'medium' plants sampled repeatedly over the course of 24 hours. The red line is the trend line created by the GAMM, the dark grey area is the 95% confidence interval.

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vegetable seed crops are not bred for seed production, they are bred for the desirable characteristics of the plants raised from that seed. Indeed, these two goals are often at odds with each other. For example: an onion that produces two flower spikes will produce much more seed, but it will also produce low-grade onions with doubled hearts [68]. The result of this compromise between seed set and plant characters has been hybrid production systems, which produce vigorous, uniform progeny, but may set less seed than open pollinated systems, if for no other reason than some portion of the field must be occupied by the pollinizer line, from which seed is not collected. As a result, hybrid carrots often yield less than 50% of the seed produced by open-pollinated carrots [10; 32].

Although the yield of hybrid carrot seed is likely to remain lower than open pollinated seed, there is obviously latitude for higher yield. In our experiment, synchrony of bloom was poor, as is common with hybrid crops [43; 69; 70]. Male-sterile plants of all three varieties began

blooming before the male-fertile line, meaning that some primary umbels, which typically have high seed yield [42], would have failed to set seed due to absence of available pollen. One cultural solution already underway in industry is to cut carrot plants early in the season to delay the flowering of one of the lines in order to synchronize bloom, which then increases yield [18; 43]. Cutting the plants at an early stage of flowering has been shown to delay bloom by 10–14 days [18]—however, this means that growers would need multiple cuttings to successfully synchronize poor male-fertile and male-sterile lines. Planting the male-fertile line even one month earlier than the male-sterile line isn't enough to fully align the two (South Pacific Seeds, Methven, New Zealand, pers. comm).

Another cultural solution is to increase the number of pollinators in the field. Carrot has a generalist flower type [10; 30] and is visited by hundreds of insect species [28–31; 71; 72], many of which can contribute significantly to successful pollination [31; 43; 71; 73]. Honey bees have traditionally been used to pollinate the crop with a stocking density of 5–8 hives/ha in New Zealand [37; 74] and Australia [43]. This is considerably lower than the hive density used in the United States, which has stocking densities 2–4 times that (15–20 hives/ha [75]). Increasing the stocking density of honey bees may improve yield, but the discrepancy may represent the different pollinator communities in the two localities as honey bees do not appear to favor carrot as a forage source [10] and preferentially visit other attractive floral resources where possible [33]. Developing practices in which increase the numbers of other species shown to efficiently pollinate carrot, such as *Megachile rotundata* [76] (Hymenoptera), *Calliphora vicina* [77], and *Eristalis tenax* [78] (Diptera) may prove better options for New Zealand, though additional field trials would be necessary to verify a benefit.

In addition, there may be some room for improvement of the carrot varieties. Surprisingly, none of the cultivars we examined contained any detectable quantities of sucrose, which is much more attractive to bees than glucose or fructose alone [79; 80], and has been detected in hybrid carrot varieties in other parts of the world [17; 31]. Additionally, all three cultivars examined in this study have very low pollen viability (<30%) compared to elsewhere in the world (~50% [21; 43; 60]). Inbreeding depression has been observed for numerous other agronomically important traits in carrot [67], and it may be the case that New Zealand hybrid carrots have poor pollen viability because of this. However, there is considerable genetic diversity within cultivated carrot globally [81]—including within groups sharing the same agronomic characters [75]. This being the case, introducing breeding stock from elsewhere in the world may alleviate some of the stress in the New Zealand hybrid carrot production system while still selecting for marketable qualities, and the added genetic variability could result in varieties more robust to changes in climate and weather patterns. Although hybrid seed crops are not typically bred for the fitness of the parent lines, it may become necessary to do so if poor plant vigor, poor attractiveness, lower pollinator populations, and increased stress from a warming climate lead to seed sets much lower than they are today.

Implications for pollination under climate change

In the carrot seed producing region of New Zealand, climate change is forecast to decrease rainfall and increase surface temperature, leading to seasonal shifts and an increase in droughty conditions [38]. Climate change has been linked to negative crop plant outcomes, including: increased susceptibility to insects and disease [82; 83], decreased competitiveness versus weeds [82; 84], decreased effectiveness of herbicides on weed control [84; 85], reduced overlap between bloom and pollinators [85], reduced pollen viability [22; 23], changes in volatile emissions [86], and quantity and quality of nectar which may affect plant attractiveness [27; 87]. Previous work in New Zealand has identified that higher temperatures may result in

increased foraging by honey bees, while reducing species richness [88]—primarily native and introduced flies. As a recent meta-analysis has found that crop yields tend to increase with pollinator richness, independent of honey bee abundance [89], this may result in an increased number of visits, but a decrease in average visit quality.

We found that there was an effect of temperature on nectar concentration and volatile emission, but no other plant characteristics in our study, although temperature was retained in the models for seed set and pollen viability (Fig 2), meaning it added explanatory power. It is important to remember that we attempted to expose plants only at the pollen formation stage during flowering, rather than throughout development. Temperature has been shown to affect plant development at numerous critical periods [22], and exposure to high temperatures at an earlier point may have reduced plant vigor beyond the effects we observed here. However, the temperatures we exposed plants to in the hot treatment are slightly higher than the future temperature projections for the region [38], so there are unlikely to be further effects on pollen viability in a warmer climate, which is fortunate given the already low viability of the hybrid varieties. Hybrid crops such as carrot are highly susceptible to pollination disruption, due to their requirement for pollen transmission across pollinizer lines. Therefore, if a warming climate leads to fewer non-managed pollinators, this could potentially reduce yield. Given carrot's already modest attractiveness to pollinators compared with weedy species [10; 33], the potential increase in nectar concentration to above the attractive range of 30–50% [25] and change floral scent could further limit its competitiveness. The situation may be exacerbated by the warming-induced increase in weed vigor predicted by other studies [82–84], as it would increase competition for a more limited pool of pollinators, with a net negative effect on seed set.

Conclusions

The combination of lowered attractiveness with higher competition for pollinators and higher losses to weeds could prove to be a difficulty for future hybrid carrot seed production. If other carrot seed growing regions of the world experience a decline in unmanaged pollinators, as New Zealand is expected to, it could lead to a fragile production system through over-reliance on honey bees [89; 90]. As insurance against adverse pollination conditions, future hybrid production systems may have to balance agronomic traits with the plant's ability attract to pollinators and set seed or, potentially, domesticate currently unmanaged pollinator species.

Supporting information

S1 Fig. Traces from the flame ionization detector (FID) of the gas chromatograph (top line) coupled with the electro-antennogram detector (EAD) responses (bottom line) from a honey bee antenna. Three electrophysiological responses were detected from the EAD trace, circled in red: phenylacetaldehyde, nonanal, methyl salicylate, from left to right. The antenna was exposed to a carrot flower headspace sample that had been collected over a period of 24 hours.

(TIF)

S2 Fig. Temperatures experienced by each shadehouse and glasshouse treatment. Histogram of temperatures collected at chest height from data loggers (onset HOB0 ProV2 temp/RH meters) from the two enclosures for each of the three temperature treatments during the course of the experiment.

(TIF)

S3 Fig. Stages of development in carrot florets. Petals and anthers were removed from florets for photography.

(TIF)

S1 Table. Coefficients table of GLM for flower phenology. Relationship between time between sowing and blooming and the plant variety, and the plant line (male sterile vs. male fertile), with interactions and a gamma distribution. The intercept condition is the excellent variety, male sterile.

(DOCX)

S2 Table. Coefficients table of zero-inflated binomial GLMM for seed set. The final model retained temperature at the time of pollination, plant variety, and pollen viability as predictors of observed seed set per three umbellets. The intercept condition is the excellent variety.

(DOCX)

S3 Table. Coefficients table of binomial GLMM for pollen viability. The final model retained temperature at the time of pollination, plant variety, and the number of days pollen was stored prior to processing as predictors of observed pollen viability in male fertile lines. The intercept condition is the excellent variety.

(DOCX)

S4 Table. Coefficients table of LM for nectar glucose:fructose ratio. The intercept condition is nectar glucose (μg) per $\frac{1}{2}$ umbel.

(DOCX)

S5 Table. Coefficients table of GLMM for nectar sugar composition. The final model retained time-of-day, temperature at the time of pollination, plant variety, and the interaction between time-of-day and variety. The intercept condition is the excellent variety at the peak nectar emission time of 11:00am.

(DOCX)

S6 Table. Coefficients table of ADONIS for nectar phenolic bouquet. The final model retained plant variety, temperature at the time of pollination, and time-of-day.

(DOCX)

S7 Table. Coefficients table of ADONIS for floral volatiles; variety trial.

(DOCX)

S8 Table. Coefficients table of ADONIS for floral volatiles; temperature trial.

(DOCX)

S9 Table. Coefficients table of ADONIS for floral volatiles; time-of-day trial.

(DOCX)

S1 Text. Preliminary carrot pollen viability sampling method and results.

(DOCX)

S1 Data. Raw data from which analyses were generated.

(CSV)

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References

1. Klein A-M, Vaissiere BE, Cane JH, Steffan-Dewenter I, Cunningham SA, Kremen C, et al. Importance of pollinators in changing landscapes for world crops. *Proceedings of the Royal Society B: Biological Sciences* 2007; 274:303–313. <https://doi.org/10.1098/rspb.2006.3721> PMID: 17164193
2. Dowker B and Fennell J. The relative performance of inbreds and open-pollinated populations of spring-sown onions. *The Journal of Agricultural Science* 1981; 97:25–30.
3. Gonzalez JL, Schneiter AA, Riveland NR and Johnson BL. Response of hybrid and open-pollinated safflower to plant population. *Agronomy Journal* 1994; 86:1070–1073.
4. Niemelä T, Seppänen M, Jauhiainen L and Tulisalo U. Yield potential of spring turnip rape synthetics and composite hybrids compared with open-pollinated commercial cultivars. *Canadian journal of plant science* 2006; 86:693–700.
5. Mahli S, Brandt S, Ulrich D, Lafond G, Johnston A and Zentner R. Comparative nitrogen response and economic evaluation for optimum yield of hybrid and open-pollinated canola. *Canadian journal of plant science* 2007; 87:449–460.
6. Rogers MA and Wszelaki AL. Influence of high tunnel production and planting date on yield, growth, and early blight development on organically grown heirloom and hybrid tomato. *HortTechnology* 2012; 22:452–462.
7. Tay D. 2002. Vegetable hybrid seed production, in: *Seeds: Trade, Production and Technology*, pp. 128–139.
8. FAOSTAT F. Statistical databases. Food and Agriculture Organization of the United Nations 2009;.
9. Parker FD. Efficiency of bees in pollinating onion flowers. *Journal of the Kansas Entomological Society* 1982;: 171–176.
10. Delaplane KS, Mayer DF. *Crop pollination by bees*. Cabi, Wallingford, UK. 2000.
11. Steffan-Dewenter I. Seed set of male-sterile and male-fertile oilseed rape (*Brassica napus*) in relation to pollinator density. *Apidologie* 2003; 34:227–235.

12. Greenleaf SS and Kremen C. Wild bees enhance honey bees' pollination of hybrid sunflower. *Proceedings of the National Academy of Sciences* 2006; 103:13890–13895.
13. Vanbergen AJ. Threats to an ecosystem service: pressures on pollinators. *Frontiers in Ecology and the Environment* 2013; 11:251–259.
14. Potts SG, Imperatriz-Fonseca V, Ngo HT, Aizen MA, Biesmeijer JC, Breeze TD, et al. Safeguarding pollinators and their values to human well-being. *Nature* 2016; 540:220–229. <https://doi.org/10.1038/nature20588> PMID: 27894123
15. Memmott J, Craze PG, Waser NM and Price MV. Global warming and the disruption of plant—pollinator interactions. *Ecology letters* 2007; 10:710–717. <https://doi.org/10.1111/j.1461-0248.2007.01061.x> PMID: 17594426
16. Garibaldi LA, Aizen MA, Klein AM, Cunningham SA and Harder LD. Global growth and stability of agricultural yield decrease with pollinator dependence. *Proceedings of the National Academy of Sciences* 2011; 108:5909–5914.
17. Erickson E, Peterson C and Werner P. Honeybee foraging and resultant seed set among male-fertile and cytoplasmically male-sterile carrot inbreds and hybrid seed parents. *Journal* 1979;.
18. Gracie A. Improving the Reliability of Vegetable Seed Yields. 2011;.
19. Ockendon D and Gates P. Reduced pollen viability in the onion (*Allium cepa*). *New Phytologist* 1976; 76:511–517.
20. Abdul-Baki AA and Stommel JR. Pollen Viability and Fruit Set of Tomato Genotypes under Optimum and High-temperature Regimes. *HortScience* 1995; 30:115–117.
21. Geard A, Spurr CJ, Brown PH. 2006. Improving Hybrid Carrot Seed Yields, in: 2006 Abstracts 27th International Horticultural Congress & Exhibition, pp. 42–42.
22. Young LW, Wilen RW and Bonham-Smith PC. High temperature stress of Brassica napus during flowering reduces micro- and megagametophyte fertility, induces fruit abortion, and disrupts seed production. *J. Exp. Bot.* 2004; 55:485–495. <https://doi.org/10.1093/jxb/erh038> PMID: 14739270
23. Hedhly A. Sensitivity of flowering plant gametophytes to temperature fluctuations. *Environmental and Experimental Botany* 2011; 74:9–16.
24. Wolf S, Lensky Y and Paldi N. Genetic variability in flower attractiveness to honeybees (*Apis mellifera* L.) within the genus *Citrullus*. *HortScience* 1999; 34:860–863.
25. Nicolson SW, Thornburg RW. 2007. Nectaries and Nectar, in: Nicolson SW, Nepi M, Pacini E (Eds.), Springer, pp. 215–264.
26. Cakmak I, Song DS, Mixson TA, Serrano E, Clement ML, Savitski A, et al. Foraging response of Turkish honey bee subspecies to flower color choices and reward consistency. *Journal of insect behavior* 2010; 23:100–116.
27. Petanidou T and Smets E. Does temperature stress induce nectar secretion in Mediterranean plants?. *New Phytologist* 1996; 133:513–518.
28. Willis JC and Burkill IH. Flowers and insects in Great Britain. Part I. *Annals of Botany* 1895;: 227–273.
29. Willis JC and Burkill IH. Flowers and insects in Great Britain. Part II. *Annals of Botany* 1903;: 313–349.
30. Proctor M, Yeo P, Lack A, others. The natural history of pollination. HarperCollins Publishers,. 1996.
31. Gaffney A. 2011. Insect floral visitors and their behaviour in hybrid carrot seed crops in Tasmania. (Masters dissertation). University of Tasmania.
32. Hart JM, Butler MD. 2004 Hybrid seed carrot: central Oregon..
33. Galuszka Tworek H and Goral M. Studies of pollination by bees (*Apis mellifera* L.) in flowers of male-sterile and male-fertile lines of carrot (*Daucus carota* L.). *Biuletyn-Warzywinczy* 1989; 33:35–58.
34. Small E. 1984. Plant Biosystematics, in: Grant W (Eds.), Academic Press, Toronto, pp. 195–210.
35. Hauser TP and Bjørn GK. Hybrids between wild and cultivated carrots in Danish carrotfields. *Genetic Resources and Crop Evolution* 2001; 48:499–506.
36. 2010. Principles and practices of organic carrot seed production in the Pacific Northwest.
37. Howlett B, Lankin-Vega G and Pattermore D. Native and introduced bee abundances on carrot seed crops in New Zealand. *New Zealand Plant Protection* 2015; 68:373–379.
38. 2016 Climate Change Projections for New Zealand: Atmospheric projections based on simulations undertaken for the IPCC 5th Assessment,..
39. Hampton J, Rolston M, Pyke N, Green W and others. Ensuring the long term viability of the New Zealand seed industry. *Agronomy New Zealand* 2012; 42:129–140.

40. Dudareva N, Martin D, Kish CM, Kolosova N, Gorenstein N, Fäldt J, et al. (E)- β -ocimene and myrcene synthase genes of floral scent biosynthesis in snapdragon: function and expression of three terpene synthase genes of a new terpene synthase subfamily. *The Plant Cell* 2003; 15:1227–1241.
41. Kearns CA, Inouye DW. *Techniques for pollination biologists*. University Press of Colorado, 1993.
42. Koul P, Koul A and Hamal I. Reproductive biology of wild and cultivated carrot (*Daucus carota* L.). *New Phytologist* 1989; 112:437–443.
43. Spurr C. 2003. Identification and management of factors limiting hybrid carrot seed production in Australia. (Masters dissertation). University of Tasmania.
44. Heslop-Harrison J and Heslop-Harrison Y. Evaluation of pollen viability by enzymatically induced fluorescence; intracellular hydrolysis of fluorescein diacetate. *Stain Technol.* 1970; 45:115–120. PMID: [4192549](#)
45. Rupérez P. Oligosaccharides in raw and processed legumes. *Zeitschrift für Lebensmitteluntersuchung und-Forschung A* 1998; 206:130–133.
46. Knudsen IM. High-performance liquid chromatographic determination of oligosaccharides in leguminous seeds. *J. Sci. Food Agric.* 1986; 37:560–566.
47. Knudsen KEB and Li BW. Determination of oligosaccharides in protein-rich feedstuffs by gas-liquid chromatography and high-performance liquid chromatography. *J. Agric. Food Chem.* 1991; 39:689–694.
48. Bates D, Maechler M, Bolker B, Walker S, others. 2014. lme4: Linear, generalized linear, and nonlinear mixed models: version 1,1–12.
49. R Core Team. 2014. R: A Language and Environment for Statistical Computing. <http://www.R-project.org/>
50. Burnham K, Anderson D. *Model selection and inference: a practical information-theoretic approach*. Springer-Verlag, New York. 1998.
51. Barton K. 2014 MuMIn: Multi-model inference. R package version 1.15–6..
52. Kuznetsova A, Brockhoff PB, Christensen RHB. 2015. lmerTest. R package version 2.0–3.
53. Bolker B, Skaug H, Magnusson A, Nielsen A. 2014. Getting started with the glmmADMB package.
54. Towill LE. Low temperature and freeze-/vacuum-drying preservation of pollen. *Cryopreservation of plant cells and organs* 1985;: 171–198.
55. Hanna WW and Towill LE. Long-Term Pollen Storage. *Plant Breeding Reviews* 1995; 13:179–207.
56. Browne WJ, Subramanian SV, Jones K and Goldstein H. Variance partitioning in multilevel logistic models that exhibit overdispersion. *Journal of the Royal Statistical Society: Series A (Statistics in Society)* 2005; 168:599–613.
57. Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin PR, O'Hara R, et al. 2013. 'vegan': Community Ecology Package: Ordination, Diversity and Dissimilarities, R package version 2.4.0.
58. Zuur AF, Ieno EN, Walker NJ, Saveliev AA, Smith GM. 2009. Mixed effects models and extensions in ecology with R, in: (Eds.), Springer, pp. 323–341.
59. Wood S, Scheipl F. 2009. gamm4: Generalized additive mixed models using mgcv and lme4. R package version 0.2–3.
60. Song H-x, Liu Y-l, Wu Z and Zhang G-x. The Study on Pollen Germination in Vitro of Carrot. *Journal of Shanxi Agricultural Sciences* 2010; 4:009.
61. Mao W, Schuler MA and Berenbaum MR. Honey constituents up-regulate detoxification and immunity genes in the western honey bee *Apis mellifera*. *Proceedings of the National Academy of Sciences* 2013; 110:8842–8846.
62. Cao Y, Wang Y and Yuan Q. Analysis of flavonoids and phenolic acid in propolis by capillary electrophoresis. *Chromatographia* 2004; 59:135–140.
63. Arnason J, Gale J, De Beyssac BC, Sen A, Miller S, Philogene B, et al. Role of phenolics in resistance of maize grain to the stored grain insects, *Prostephanus truncatus* (Horn) and *Sitophilus zeamais* (Motsch.). *Journal of Stored Products Research* 1992; 28:119–126.
64. Hagler JR and Buchmann SL. Honey bee (Hymenoptera: Apidae) foraging responses to phenolic-rich nectars. *Journal of the Kansas Entomological Society* 1993;: 223–230.
65. Hawthorn L, Bohart GE, Toole E. 1956. Carrot seed yield and germination as affected by different levels of insect pollination, in: *Proceedings of the American Society for Horticultural Science*, pp. 384.
66. Holzschuh A, Dormann CF, Tschamntke T and Steffan-Dewenter I. Expansion of mass-flowering crops leads to transient pollinator dilution and reduced wild plant pollination. *Proceedings of the Royal Society B: Biological Sciences* 2011; 278:3444–3451. <https://doi.org/10.1098/rspb.2011.0268> PMID: [21471115](#)

67. Stein M and Nothnagel T. Some remarks on carrot breeding (*Daucus carota* ssp. *sativus* Hoffm.). *Review of Plant Breeding* 1995; 114:1–11.
68. Krontal Y, Kamenetsky R and Rabinowitch H. Flowering physiology and some vegetative traits of short-day shallot: a comparison with bulb onion. *The Journal of Horticultural Science and Biotechnology* 2000; 75:35–41.
69. Verdial MF, Cardoso All, Lima MáSd and Chaves FCéM. Coincidence of flowering time and the productivity and quality of cauliflower hybrid seeds. *Scientia Agricola* 2001; 58:533–539.
70. Sugimoto A, Terajima Y, Terauchi T, Ponragdee W, Ohara S, Tagane S et al. Developing new types of sugarcane by hybridization between commercial sugarcane cultivars and wild relatives. *Plant Genetic Resources for Food and Agriculture in Asia and the Pacific: Impacts and Future Directions* 2012; 11.
71. Bohart GE and Nye WP. Insect pollinators of carrots in Utah. 1960;.
72. Ahmad M and Aslam M. Pollinators visiting carrot (*Daucus carota* L.) seed crop. *Journal of Research (Science)* 2002; 13:31–35.
73. Howlett B, Walker M, Rader R, Butler R, Newstrom-Lloyd L and Teuton D. Can insect body pollen counts be used to estimate pollen deposition on pak choi stigmas?. *New Zealand Plant Protection* 2011; 64:25–31.
74. Goodwin M. Pollination of Crops in Australia and New Zealand. RIRDC, 2012.
75. Rubatzky V, Quiros C, Simon P. Carrots and related vegetable Umbelliferae. CABI, New York. 1999.
76. Davidson M, Butler R and Howlett B. *Apis mellifera* and Megachile rotundata: a comparison of behaviour and seed yield in a hybrid carrot seed crop. *New Zealand Journal of Crop and Horticultural Science* 2010; 38:113–117.
77. Howlett B. Hybrid carrot seed crop pollination by the fly *Calliphora vicina* (Diptera: Calliphoridae). *Journal of Applied Entomology* 2012; 136:421–430.
78. Pérez-Bañón C, Petanidou T and Marcos-García MÁ. Pollination in small islands by occasional visitors: the case of *Daucus carota* subsp. *commutatus* (Apiaceae) in the Columbretes archipelago, Spain. *Plant Ecology* 2007; 192:133–151.
79. Waller GD. Evaluating responses of honey bees to sugar solutions using an artificial-flower feeder. *Ann. Entomol. Soc. Am.* 1972; 65:857–862.
80. Hagler JR, Cohen AC and Loper GM. Production and composition of onion nectar and honey bee (Hymenoptera: Apidae) foraging activity in Arizona. *Environmental entomology* 1990; 19:327–331.
81. Bradeen JM, Bach IC, Briard M, le Clerc V, Grzebelus D, Senalik D, et al. Molecular diversity analysis of cultivated carrot (*Daucus carota* L.) and wild *Daucus* populations reveals a genetically nonstructured composition. *Journal of the American Society for Horticultural Science* 2002; 127:383–391.
82. Patterson DT. Weeds in a changing climate. *Weed Sci.* 1995; 685–701.
83. Juroszek P and Von Tiedemann A. Potential strategies and future requirements for plant disease management under a changing climate. *Plant Pathology* 2011; 60:100–112.
84. Ziska LH, Faulkner S and Lydon J. Changes in biomass and root: shoot ratio of field-grown Canada thistle (*Cirsium arvense*), a noxious, invasive weed, with elevated CO₂: implications for control with glyphosate. *Weed Sci.* 2004; 52:584–588.
85. Ziska LH and Teasdale JR. Sustained growth and increased tolerance to glyphosate observed in a C3 perennial weed, quackgrass (*Elytrigia repens*), grown at elevated carbon dioxide. *Functional Plant Biology* 2000; 27:159–166.
86. Sagae M, Oyama-Okubo N, Ando T, Marchesi E and Nakayama M. Effect of temperature on the floral scent emission and endogenous volatile profile of *Petunia axillaris*. *Biosci. Biotechnol. Biochem.* 2008; 72:110–115. <https://doi.org/10.1271/bbb.70490> PMID: 18175901
87. Pacini E, Nepi M and Vesprini J. Nectar biodiversity: a short review. *Plant Systematics and Evolution* 2003; 238:7–21.
88. Howlett B, Butler R, Nelson W, Donovan BJ. Impact of climate change on crop pollinator in New Zealand., 2013.
89. Garibaldi LA, Steffan-Dewenter I, Winfree R, Aizen MA, Bommarco R, Cunningham SA et al. Wild pollinators enhance fruit set of crops regardless of honey bee abundance. *Science* 2013; 339:1608–1611. <https://doi.org/10.1126/science.1230200> PMID: 23449997
90. Winfree R, Williams NM, Dushoff J and Kremen C. Native bees provide insurance against ongoing honey bee losses. *Ecology Letters* 2007; 10:1105–1113. <https://doi.org/10.1111/j.1461-0248.2007.01110.x> PMID: 17877737