

Dear Emmanuelle Porcher,

We revised our preprint submitted to PCI Evol Biol entitled “Potential adaptive divergence between subspecies and populations of snapdragon plants inferred from $Q_{ST} - F_{ST}$ comparisons”. My apologies for the time it took. Reviewers gave many constructive comments. We compared snapdragon subspecies and found differences. We therefore rerun all the analyses for each subspecies separately. We also rewrote several parts of the text to accommodate all the reviewers’ comments, which improved greatly the quality of the manuscript. We hope that you – and the reviewers – will agree to follow up on this revised manuscript submission and that you will find this revised version of the manuscript more suitable for recommendation.

Please find below a point by point response to your comments and the comments by Sophie Karrenberg and Santiago Gonzalez (text of our response is in blue).

Dear Sara Marin,

I would first like to apologize for the time needed to return this decision, which is partly attributable to the difficulty of finding available reviewers in the middle of summer. Your preprint has now been read carefully by two experts (including Sophie Karrenberg (SK), who signed her review) and by me. We all agree that your manuscript addresses an important evolutionary question (patterns of adaptation to altitude), and that the experimental design was appropriate to address this question, such that I would ultimately be willing to recommend this preprint provided you can address the issues raised by the reviewers. Here are the most critical issues I identified, some of which were raised independently by both reviewers:

(1) Definition of local adaptation: both reviewers and I were lost with, or at the minimum confused by, your distinction between local adaptation (the meaning of which seems to vary throughout the paper, see comments by SK) and adaptation to latitudinal gradients, which involves local adaptation in the broad sense, such that individuals with the highest fitness in one location are on average those originating from this location. It is really important that this point is clarified. This may include for example specifying whether you consider local adaptation as a pattern or as a process or both. Another option for clarification would be to follow the first suggestion of SK, which is to focus more on biology and the question of adaptation to altitude, and how to disentangle it from adaptation to other environmental variables, which may vary in space at a finer grain than altitude (or not, see comment 7 by reviewer #2)

We now start the manuscript by defining local adaptation. We defined it as a process: “the evolutionary response to selection that makes populations fitter in their own local habitat than in other populations’ local habitats”. We modified the terms that we use throughout the text following SK and SG recommendations to avoid confusion between the local adaptation to the environmental conditions of their local sites of origin and to the altitude. Following SK comments, we gave more weight to the divergence between the two subspecies and to the biology rather than the methods in terms of adaptation.

(2) SK is worried that pooling the two subspecies might bias your analyses. There again, it is central to provide a solid justification for this pooling

We separated the two subspecies. This led to statistical power limitations for the study of some traits. We dropped these traits and kept those that were reliable.

(3) Methodological / statistical issues: both reviewers had several additional questions and suggestions regarding the statistical analyses, which I will not fully list here (although they all deserve attention), but which include for example: (i) How could you estimate the within population additive genetic variance, when the number of individuals and the number of families seem to be similar (i.e. about one individual per family)? (ii) How was the significance of the relationship between F_{st} (or Q_{st}) and altitudinal differences tested in Figure 3? (iii) On the same figure, can the statistical significance of the differences in slopes be tested? More generally, the two reviewers often requested more details on the methodology (e.g. Q_{st} - F_{st} comparisons, quantitative genetics analyses...), and I agree with them that some key information is sometimes missing.

(i) This was a misunderstanding caused by awkward writing. Sorry about that. We modified the text to clarify this point. The text now states precisely: “Nine to 42 seed families from each of the 13 study populations were grown outdoor in spring 2014 in a common garden at ENSFEA (Toulouse, France). Two plants per family were grown. Some plants died before measurements were done, which resulted in some families being represented by only one plant”. We added the detail of the number of families and the total number of plants grown for each population in Table S1.

(ii) We had conducted a regression analysis for each relationship with altitude (not appropriate because the data was not independent) and a mantel test. The presentation of the results and the figure was unclear and confusing in the previous version. We have removed the regression results, clarified what was tested by using a mantel test, and conducted partial mantel tests following the reviewer’s comments.

(iii) Unfortunately, it is not possible to estimate the statistical significance of slope differences for pairwise population differentiation data. The partial mantel tests, which we recognise are not perfect, are the best approach we can use for this type of data. It provides results close to those you were asking for. We agree that the information on methods and results was often confusing and that some information was missing. We have modified the text throughout the manuscript and hope that it will be clear.

(4) While the “Methods” section could be expanded, the two reviewers and I agreed that the text could also be shortened at places, particularly in the discussion.

We have shortened the text of the discussion.

Thank you very much for submitting your preprint for recommendation in *PCI Evolutionary Biology*. I hope you will find this feedback, and particularly the very thorough reviews of the experts, useful to revise your manuscript.

We appreciated the thorough reviews. They improved the quality of the manuscript. It was very pertinent to look into the subspecies divergence. It is generally acknowledged by the snapdragon plants research community that the two subspecies do not differ but for the flower colour and we are glad that the reviewers pushed us to investigate subspecies differences. The results of our indirect approach suggest that adaptation might have proceeded differently with

the phenotypes of the two subspecies, which opens the way for testing several hypotheses by using direct tests in the future.

Best regards Emmanuelle

Kind regards,

for the authors, Benoit Pujol

Sophie Karrenberg

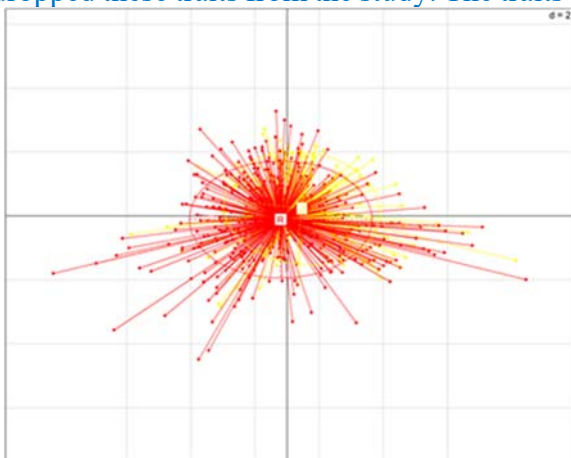
Your manuscript on altitudinal divergence in snapdragon plants treats an interesting and timely question mostly with appropriate methods. However, I do have several difficulties to evaluate the quality of this research because important information appears to be missing from the manuscript. This concerns: I), most importantly, whether there were genetic or phenotypic differences between two subspecies that were apparently pooled here, according to Material and Methods and II) the experimental design such as spatial arrangement in the garden (how plants were arranged in how many containers) that should be taken account of as a random effect in the analyses, as well as crossing design and number of individuals per family used (see comments below).

Dear Sophie Karrenberg, thank you for your thorough review of our manuscript. We have taken into account all your comments, which improved the quality of our manuscript. The research community working on snapdragon plants generally considers that snapdragon subspecies do not differ but for the flower colour without really testing for it. Following your comment, we have compared the subspecies and found differences. We therefore analysed subspecies data separately. We had to drop a few traits in the process because the statistical power was not sufficient for them. Our results suggest that adaptation shaped differentially the phenotype of the two subspecies. This was a game changer for this paper. We modified the text and reran the analyses to accommodate this perspective. We also put the biology and the effect of altitude forward in the revised version of the manuscript. We took into account the blocks – containers – where plants were located randomly in the analyses, and clarified the crossing design. As you noticed, there was some confusion in the presentation of the numbers. More than one progeny was obviously present per family (we now give the number of plants and families for each population in Table S1). We are sorry that the presentation of many aspects was awkward in the methods and results. We have attended to it in this new version. We reduced the size of the discussion and gave clearer explanations in the methods and results. We have modified the text in many places. We hope that these modifications will have sorted out the problems that you mentioned. Please find below a more detailed, point by point response to your comments.

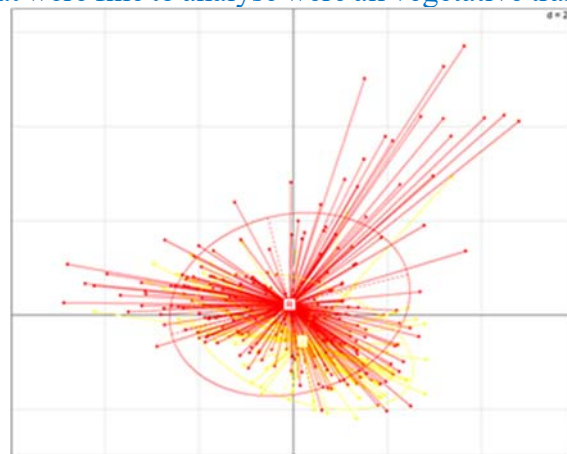
Regarding the subspecies, an explicit analysis would be most helpful (phenotypic and genetic differentiation, for example PCoA or structure analysis, potentially presented in the supplementary material). Pooling of the subspecies would probably only be appropriate if there are no considerable phenotypic differences or genetic differentiation between them. F_{st} in these analyses is used to reflect putatively neutral genetic differentiation -- this might be difficult

assumption if the dataset includes subspecies that are partially reproductively isolated from each other. The description of the subspecies in the method section as well as the citations do fuel this concern. Differences in habitat and flower color between subspecies are described — even pollinator-mediated reproductive isolation could be present in this system. Very careful and convincing analyses and explanations would be needed to motivate the pooling. And —if there are no differences between subspecies, would that not suggest that there might just be a flower color polymorphism? It is important to note that, without this potentially troublesome pooling of subspecies, the number of populations would probably be too low for the present analysis and this is why this issue is so critical. Provided that pooling can be justified, and the design clarified, this manuscript could become a good one, however, further improvements in the analysis and presentation are also needed.

We conducted a PCA analysis, see graphs below. No contrast could be observed between subspecies based on the graphs. We nevertheless conducted statistical analyses to test for phenotypic differences between subspecies. Hierarchical generalized linear models were conducted with population nested in subspecies. We found differences. This was surprising but very interesting to find out. The subspecies mostly share the same ecological niche. They are interfertile but are reproductively isolated by pollinators. Based on this information, we decided to consider them separately in all the following local adaptation analyses. By doing so, we lost statistical power and some traits were not reliable any longer for QST-FST comparisons. We dropped these traits from the study. The traits that were fine to analyse were all vegetative traits.



PCA on vegetative traits. Yellow = *A. m. striatum* Red = *A. m. pseudomajus*



PCA on reproductive traits. Yellow = *A. m. striatum* Red = *A. m. pseudomajus*

The main changes that I believe are necessary are: 1) Focus on biology. The paper would be more interesting if focus was shifted from methods and general interpretations to the biology of altitudinal adaptation. This would be facilitated by presenting the trait-altitude correlations first. These results are interesting for all traits, currently they are only presented for 3 out of 13 traits. From that you could move on to more complex analyses to identify the underlying causes of such trait clines. In the current version, there is overall rather little on HOW the traits change along altitude, for example, there are no details on leaf traits, even though there is a lot of literature of variation in SLA (see citations in literature mentioned below). It would also be very good if explicit expectations for all traits or for trait groups, based on the literature, could be added in the introduction. I would also appreciate an introduction ending in specific questions.

We agree that QST-FST approaches are well-known and made changes accordingly. We now focus more on the biology. We reduced the text on the QST-FST approaches. These approaches are now presented more briefly. We present their limits but removed the unnecessary details on

the method itself and any text that could read as if it praised their use because they are far from perfect, and there is no novelty there. We now present the biological expectations in terms of trait adaptive change with altitude in the introduction and discuss the results that we found in *A. m. striatum* in the light of these expectations.

2) In the methods part, you write that you used the between family variance component to estimate the within population additive genetic variance for a trait, as is commonly done. However, Table S1 states, for reproductive traits, the total number of families is 372 and the number of individuals is 380, such that you cannot have had more than one individual per family in most cases, unless you used a subset of families with more individuals (I could not find information on the number of individuals per family). With 1 individual per family it should be very difficult or impossible to estimate the between family variance component. In Table S3 you do not give the within family variance component so I wonder whether what you state as the within population variance component really is the between family variance component. This issue is in need of clarification. It would also be good if you could explain why you have fewer measurements for reproductive traits than for vegetative traits.

We agree that the presentation of the setting was awkward and misleading. As mentioned in our response to Emmanuelle Porcher: << We modified the text to clarify this point. The text now states precisely: “Nine to 42 seed families from each of the 13 study populations were grown outdoor in spring 2014 in a common garden at ENSFEA (Toulouse, France). Two plants per family were grown. Some plants died before measurements were done, which resulted in some families being represented by only one plant”. We added the detail of the number of families and the total number of plants grown for each population in Table S1.>>. We had fewer reproductive measures because all plants grew but not all plants produced flowers, which is why reproductive traits harboured less statistical power. Consequently, when we separated the data set in two parts, one for each subspecies, reproductive traits could not reliably be analysed and we had to remove them from the study.

3) Methods, QST - FST. I found the description of the QST - FST comparisons, using expected distributions, unclear and in part contradictory (see comments in the text). In particular, it is not clear to me how you arrived at the confidence intervals for QST on Figure 2. It would fit better if you presented the distribution of expected values for QST together with the observed values, this is the method you stated in the methods section. Alternatively, you could change the methods section.

Our presentation was indeed incomplete. In particular, the presentation of the two methods that we used independently was confusing. We clarified the text to explain each method separately: “Comparisons between F_{ST} and overall Q_{ST} values were performed for each trait based on two methods: i) a comparison of confidence intervals, the Q_{ST} is considered non significantly different from neutral differentiation when the confidence interval of the overall Q_{ST} for a trait overlaps the mean F_{ST} value, ii) a bootstrapping method developed by Whitlock and Guillaume (2009). This last approach aims at comparing the observed difference between the overall Q_{ST} and the F_{ST} values with the expected simulated distribution of this difference under a scenario of neutral evolution. We generated 100 000 bootstrap replicates of the expected $Q_{ST} - F_{ST}$ difference under the neutrality hypothesis for each trait, and built the corresponding distribution. In this approach, P values were estimated by assessing whether the observed value of the $Q_{ST} - F_{ST}$ difference overlapped its expected distribution under neutrality. We used the modification by Lind *et al.* (2011) of the approach of Whitlock and Guillaume (2009) to estimate the variance components of the simulated values of the $Q_{ST} - F_{ST}$ difference”.

“Confidence intervals of Q_{ST} values were calculated following a parametric bootstrap method adapted from O’Hara and Merilä (2005).”

4) Analysis of pairwise data: the Methods section describes the commonly used and suitable analyses for correlation tests among matrices (pairwise data), Mantel tests and partial Mantel tests, controlling for the effect of neutral genetic divergence. In the results however, a regression analysis is presented (Fig. 3) that is not suitable for this type of non-independent data points. The text states that the Mantel test was used to assess significance of the regression — this is not possible. The actual data points are also not displayed, this is a bit worrisome due to the subspecies issue. My suggestion is to re-work this analysis according to the methods described and to display the data points on Fig. 3.

Sorry about that. Awkward wording lead to some confusion about which of these methods we used in the paper. We modified the text and approach accordingly to your comment. The regression was indeed not adapted and we removed it. We have clarified the text and now present results for the mantel and partial mantel tests only.

5) (Local) adaptation, interpretation. The interpretation of what constitutes local adaptation is highly variable throughout the manuscript. At some places, the common cautious interpretation of QST - FST analyses, that they can provide hypotheses on traits that may have adaptively diverged due to selection (see for example review by Leinonen et al. 2013, Nature Reviews Genetics) is used. At quite some other places, however (abstract, introduction, discussion, marked in detailed comments), it appears that local adaptation is inferred whenever trait divergence cannot be attributed to either the altitudinal gradient or neutral divergence. I cannot be sure that the text is really meant that way, of course, but I testify here that it reads that way. This interpretation would be incorrect (genetic drift and many other processes could be responsible for trait divergence instead). Any text that can potentially be misunderstood is probably better changed.

We have modified the text accordingly to restore the coherence of our interpretation of what constitutes local adaptation throughout the manuscript. Right away from the start of the manuscript, we define local adaptation as a process: “the evolutionary response to selection that makes populations fitter in their own local habitat than in other populations’ local habitats”. We modified the text accordingly to tone down the conclusions that we obtained from QST-FST comparisons. We now say throughout the text that significant results do not confirm the presence of local adaptation but rather suggest local adaptation, reflect a potential for local adaptation, or identify candidate traits for local adaptation. We hope that this will read right. Please do not hesitate to point out specific sentences that would still be a problem.

6) Interpretation of the results on germination. The corresponding paragraph in the discussion unfortunately appears contradictory and the two types of results, overall QST - FST and pairwise QST - FST have not been compared (see comments in the text). Moreover, the text also appears as if it was not clear to the authors that a lack of quantitative genetic differentiation in traits does not exclude phenotypic plasticity (i.e., this is not really an alternative interpretation). It seems that this part needs adjustment.

We clarified the text to avoid any confusion. Plasticity is not expected to drive the overestimation of QST because the experimental environment is similar between populations. If populations express a convergent phenotypic response through plasticity to a similar

environment, then a lower QST than expected might be found. This latter mechanism might be an alternative hypothesis to standardizing selection when explaining $QST < FST$.

7) Interpretation of genetic divergence. You invoke reproductive isolation, when it appears that isolation by distance would be a sufficient interpretation, unless you refer to the subspecies differences here (see comments above and in the text). However, this cannot be evaluated from the data currently presented, as details on genetic differentiation are missing (see above). I suggest that this interpretation is re-worked once a more detailed analysis is presented.

We have clarified these aspects in the text. We now present the corresponding results to accommodate this comment in the paper. In brief, we did not find any isolation by distance. Mountains were a plausible ecological barrier to gene flow between populations. Subspecies are differentiated and pollination is an acknowledged pre-zygotic reproductive barrier between subspecies. For more details, please have a look at our previous paper on the ecological factors influencing the spatial genetic structure of snapdragon populations (Pujol et al, 2017 Botany Letters, which we cite in the paper).

8) Literature. The manuscript would much benefit by adding more literature on the analysis of altitudinal gradients and transplant experiments, for example: Luo, Y., A. Widmer, and S. Karrenberg. 2015. The roles of genetic drift and natural selection in quantitative trait divergence along an altitudinal gradient in *Arabidopsis thaliana*. *Heredity* 114:220. Halbritter, A. H., S. Fior, I. Keller, R. Billeter, P. Edwards, R. Holderegger, S. Karrenberg, A. R. Pluess, A. Widmer, and J. M. Alexander. 2018. Trait differentiation and adaptation of plants along elevation gradients. *J. Evol. Biol.* [Disclaimer: these are papers I am an author on-- but they really are related here... as are many of the citations in them].

We now cite Halbritter et al. 2018.

9) Methods are not really new. Throughout the intro and the discussion, the paper reads as if this mainly is a method advancement. This appears somewhat exaggerated, as similar methods have been used before and since a longer time (>5 yrs). I feel that the work of others should be acknowledged much more here, only few citations on such studies that are able to identify selective agents are given (for example, Hangartner et al).

We modified the text accordingly. We toned down the originality and the interest of the approach (altitudinal gradient effect on QST-FST comparisons). We also made sure that the work of others is better acknowledged by adding references (for example, Halbritter et al. 2018).

10) Shortening. I find that the text could likely benefit from shortening, in particular in the introduction and discussion, probably to at least 2/3 of its present length. I have indicated several parts that appear repetitive in the detailed comments.

We have provided more details for the sake of clarity in the Material and methods and the results sections. We reduced the size of the text in the introduction and the discussion by removing redundancy, shortening the text about the methodological approach to draw a better focus on the biology.

11) More comments. I have made comments similar to the above comments as well as many further detailed comment directly on the text using pdf commenting tools in Adobe Acrobat. I

am not repeating these detailed comments here. Such a list of details would be very difficult to write and read, because the text did not include any line numbers to refer to.

Thank you for taking the time to comment directly on the text. We have followed all your suggestions.

I hope that you will find my comments clear and helpful for further revisions of your work.

Your recommendations were particularly useful and much appreciated to improve the quality of the manuscript.

With best wishes, Sophie Karrenberg

Kind regards,

for the authors, Benoit Pujol

Santiago Gonzalez

Evolutionary inference from Qst-Fst comparisons: disentangling local adaptation from altitudinal gradient selection in snapdragon plants

This manuscript provides new and relevant insights on local adaptation of snapdragons based on quantitative genetic data from a common garden and previously published nuSSR data. However, I found confusing the way the authors constructed the paper, contrasting what they called ‘local adaptation’ (i.e. populations standing alone) versus ‘adaptation to altitudinal gradients’. From my perspective, adaptation to an environmental gradient involves also local adaptation (of each of the population of the gradient). In addition, other traits with $Q_{st} > F_{st}$ but not associated with altitudinal gradients (and thus part of the ‘local adaptation’ group) may still be associated to other environmental gradients (that have not been measured in this study, e.g. for soils).

We clarified the text accordingly. As mentioned above in response to EP’s comments: <<We now start the manuscript by defining local adaptation. We defined it as a process: “the evolutionary response to selection that makes populations fitter in their own local habitat than in other populations’ local habitats”. We modified the terms that we use throughout the text following SK and SG recommendations to avoid confusion between the local adaptation to the environmental conditions of their local sites of origin and to the altitude.>>

In addition, the strongest evidence of local adaptation along the altitudinal gradient does not come from the overall Qst-Fst comparison but from the pairwise analyses and the simple correlations between phenotypic means and altitude (Figures 3 and 4).

We now discuss the biological significance of the correlation between phenotypic mean and altitude in accordance with hypotheses formulated in the introduction. This reinforces our discussion of the biology rather than discussing the pros and cons of QST FST methods that already widely acknowledged in the literature. The number of traits that we discussed had to be modified after following up on the suggestion to look into subspecies divergence. Since we

found divergence, we conducted QST-FST comparisons in each subspecies separately. Reproductive traits (measured in less plants than vegetative traits because not all plants flowered) had to be dropped from the paper for lack of statistical power when the analysis is conducted in each species separately. We found evidence for potential adaptation to altitude in only one subspecies (*Antirrhinum majus striatum*).

Finally, please, have a careful read of the manuscript to correct errata, uncomplete sentences, unfinished citations, etc.

We agree that the text was unclear in many places and could lead to some confusion, in particular with the methods. We have clarified the text throughout the manuscript, detailed the methods and results for the sake of clarity and synthesized the intro and the discussion to put biology forward and tone down the interest of this type of approach because it is already well described in the literature.

Some more specific comments follow:

Introduction

1. The Qst-Fst method, as applied here, has also important flaws compared to reciprocal transplantation approaches, which I think should be presented and briefly discussed in the Introduction. Perhaps the most important one is the bias produced by the common garden testing environment. Parameters like heritability or Qst may differ greatly when computed in different environments, and in particular field estimates are very different from those obtained in common gardens. This should be acknowledged.

We agree and now acknowledge this clearly in the text : “In plants, reciprocal transplants directly comparing fitness between the native habitat and the foreign habitats are often preferred to QST-FST approaches conducted in common gardens because they allow to evaluate the effect of environmental conditions (Etterson, 2004; Angert and Schemske, 2005; Kim and Donohue, 2013). When the conditions for the reciprocal transplant cannot be easily met, QST-FST comparisons represent an opportunity for exploring local adaptation hypotheses.”

2. Despite being common to use adaptation to altitude as a first approach to understand adaptation to climate change, there are some fundamental differences that make the two processes very different, in particular for traits related to photoperiod where, for example, day length is also relevant. These differences should be acknowledged and discussed in the Introduction and in the Discussion.

We agree that using the environmental conditions associated with altitude to assess the potential effect of climate change is far from perfect. In the revised version of the manuscript, we do not mention climate change. We only mention that climate variables are associated with altitudinal differences amongst other environmental variables: “we investigated whether quantitative genetic differentiation increased with altitudinal difference, with the hypothesis that environmental changes associated with altitude, which include a suite of climatic variables, drove adaptive responses. Other environmental variables (e.g., atmospheric pressure) can also change with altitude. Our study also ultimately participate in evaluating whether Q_{ST} - F_{ST} comparisons can be used as a tool to identify candidate traits involved with the potential adaptation of populations to altitudinal gradients, and thereby climate differences.”

3. I like frogs, but still, why to provide examples of frog adaptation to altitude in the Introduction? There is a rich literature in plants that could be used instead, to illustrate patterns

of genetic adaptation and plasticity along altitudinal (and latitudinal, which are perhaps more relevant to discuss climate change) gradients.

We agree and made sure to cite more references that are specific to plants adaptive changes with altitude (e.g., Halbritter et al. 2018, in JEB, which reviews this field of research)

Material and Methods

4. Key information is missing about the quantitative genetics models used by the authors, in particular about sample sizes. Also, it would be good to write down the model itself, as the experimental design is not very clear. For example, did the experiment follow any block design? How many families? How many full-sibs per family? I have seen Table S1, but number of samples seem similar to number of families, which I don't understand...

The text was indeed confusing. We modified the text to clarify this aspect and now give the exact number of plants and sib-families for each population in a table: "Nine to 42 seed families from each of the 13 study populations were grown outdoor in spring 2014 in a common garden at ENSFEA (Toulouse, France). Two plants per family were grown. Some plants died before measurements were done, which resulted in some families being represented by only one plant (Table S1, Supporting Information)." We also precise that plants were located randomly into containers and that the potential effect of these containers was taken into account into the statistical analyses.

5. In my opinion, the most original test presented in the manuscript is the pairwise analysis of Q_{st} versus altitudinal differences (Figure 3). I am aware that the authors designed the study to include different altitudinal gradients so that dependence with distance is broken, but still some distance effects may remain and I think pairwise distance between populations should be included in these analyses (by means of a partial Mantel test or by using it as covariate).

We have modified this approach in the revised manuscript. Regressions used in the previous version were not suitable for this data (pairwise population differentiation because data points are not independent). We now use partial mantel tests. We do not have enough statistical power to fit pairwise distance in these analyses. We are nevertheless confident with our approach because (i) no isolation by distance was found on the basis of putatively neutral markers (we now present this finding in the paper) and (ii) as you mention, our sampling scheme (populations from different valleys and mountains) aimed at avoiding such spatial distance effect.

6. "Potential effects on population genetic differentiation... for by using AMOVA". Not clear how the authors did this... and also, should not Q_{st} estimates be also corrected in the same way? I find a bit weird, for example, that different subspecies are pooled to estimate heritability and Q_{st} .

Following this comment and comments by SK, we have investigated the phenotypic divergence between subspecies and found some. We therefore ran analyses in each subspecies separately and used different types of analyses: comparison between Q_{ST} CI and mean F_{ST} , Q_{ST} - F_{ST} difference simulation tests, mantel tests, mantel partial tests. Text, table and figure modifications related to these changes appear throughout the paper. We modified the text accordingly to incorporate the question on subspecies. As mentioned above in our response to EP: "It was very pertinent to look into the subspecies divergence. It is generally acknowledged by the snapdragon plants research community that the two subspecies do not differ but for the

flower colour and we are glad that the reviewers pushed us to investigate subspecies differences. The results of our indirect approach suggest that adaptation might have proceeded differently with the phenotypes of the two subspecies, which opens the way for testing several hypotheses by using direct tests in the future.”

Results

7. The manuscript focuses on the power of the Qst-Fst method to detect adaptation to gradients, but still most of the traits with $Q_{st} > F_{st}$ did not correlate with altitudinal variation but with other, unknown factors. The story the paper tells is, of course, the author’s own business, but still it makes me wonder whether these other traits with $Q_{st} > F_{st}$ do not deserve more attention (for example in the Discussion) and a substantial change of the main argumentation line.

We agree. QST FST methods and their interest and limits are thoroughly documented in the literature. We therefore removed text throughout the paper on methodological aspects to keep what was only necessary and focus more on the biological dimension of the study.

8. Figure 3. Differences in slope are pretty clear indeed, but I still think that they should be tested formally, using a test for differences of regression slopes.

As mentioned earlier, the regression approach was not adapted and was removed. We now use partial mantel tests.

Discussion

9. Overall I found the discussion a bit repetitive and unbalanced towards supporting the Qst-Fst approach as a means to detect adaptation along gradients. In my opinion, the advantages of the approach are not so well supported by the results. I appreciate the pairwise approach (Figure 3), which is relatively original, but still, as commented before, it does not take advantage of sampling along the gradients to produce more robust estimates.

We agree and modified the text accordingly. We removed redundancy and synthesized the content of the discussion to go straighter to the point. As mentioned earlier, we now focus on the biology rather than the approach so that our revised text should not read anymore as unbalanced towards supporting this type of approach.

10. “...trait homogenisation caused during the experiment by phenotypic plasticity might be another plausible explanation.” I really don’t see this point... I would say that is the other way around, that the differences you see in the field are the ones that are caused by phenotypic plasticity (please, notice that substantial additive variance and differences among families are still present in the common garden, as shown by high heritability for time to flowering and germination rate).

The same point was raised by SK. We have modified the text to clarify this aspect. “Plasticity is not expected to drive the overestimation of QST because the experimental environment is similar between populations. If populations express a convergent phenotypic response through plasticity to this similar environment, then a lower QST than expected might be expected. This latter mechanism might be an alternative hypothesis to standardizing selection to explain that $Q_{ST} < F_{ST}$.”

11. I also found weird that germination rate does not have $Q_{st} > F_{st}$ overall but it is significant in the pairwise test, i.e. $Q_{st} > F_{st}$ along altitude for pairs of populations.

The regression results were removed as they were not suitable to pairwise population data. Our results – that are now separated between subspecies – show a Germination date $QST < FST$ in *A. m. striatum* but not in *A. m. pseudomajus*. Germination date QST did not show a significant relationship with altitudinal differences when using the partial mantel test approach.

Tables

12. TABLE S3. Please add the standard error to the heritability estimates.

We agree that this is important information. We added the 95% confidence intervals though, rather than SE, because we find them a lot more informative and conservative when it comes to assessing the statistical reliability of heritability estimates.

Several of your comments overlapped with comments by SK. Please read our response to her comments, as they might be complementary. We thank you for having thoroughly reviewed our paper and for your constructive comments that proved useful for improving the quality of this manuscript.

Kind regards,
for the authors,
Benoit Pujol

1 **Potential adaptive divergence between subspecies and populations of snapdragon plants**
2 **inferred from Q_{ST} – F_{ST} comparisons**

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17

18 **ABSTRACT**

19 Phenotypic divergence among natural populations can be explained by natural selection or by
20 neutral processes such as drift. Many examples in the literature compare putatively neutral (F_{ST})
21 and quantitative genetic (Q_{ST}) differentiation in multiple populations to assess their potential
22 evolutionary signature and identify potential traits involved with local adaptation. Investigating
23 these signatures in closely related or recently diversified species has the potential to shed light
24 on the potential divergence processes acting at the interspecific level. Here, we conducted this
25 comparison in two subspecies of snapdragon plants (eight populations of *Antirrhinum majus*

26 *pseudomajus* and five populations of *A. m. striatum*) in a common garden experiment. We also
27 tested whether altitude was potentially involved with population phenotypic divergence. Our
28 results identified candidate phenological and morphological traits involved with local
29 adaptation. Most of these traits were identified in one subspecies but not the other. Phenotypic
30 divergence increased with altitude for a few biomass-related traits, but only in *A. m. striatum*.
31 These traits therefore potentially reflect *A. m. striatum* adaptation to altitude. Our findings imply
32 that adaptive processes potentially differ at the scale of *A. majus* subspecies.

33

34 **KEYWORDS**

35 Local adaptation, altitudinal gradient, quantitative genetics, subspecies, *Antirrhinum majus*

36

37

38 **INTRODUCTION**

39 Local adaptation - the evolutionary response to selection that makes populations fitter in their
40 own local habitat than in other populations' local habitats - is widespread in both plant and
41 animal species (Kawecki and Ebert, 2004; Leinonen et al., 2013; Halbritter et al., 2018). There
42 is evidence for its role in the adaptive divergence of plant species (Leimu and Fischer, 2008;
43 Hereford, 2009; Halbritter et al., 2018). For example, empirical studies have demonstrated
44 differential adaptation in plant sister species or hybridizing species, for instance between pairs
45 of *Silene* species (Favre et al., 2017), *Senecio* species (Abbott and Brennan, 2014), *Mimulus*
46 species (Angert and Schemske, 2005). These studies compared local adaptation for sister
47 species confronted to different ecological requirements. Different species may also respond
48 similarly to a same type of environmental gradient. Recently, Halbritter et al. (2018) combined
49 studies of multiple plant species along elevation gradients. They found significant evidence for
50 adaptation to different elevations in terms of survival and biomass. Their results also showed
51 variation across species in plant responses to elevation. The study of local adaptation in
52 populations of closely related taxa exposed to environmental gradients, e.g. altitude, is an
53 opportunity to investigate the conditions promoting or impeding the consistency of adaptive
54 responses.

55
56 An indirect approach to investigate whether local adaptation might potentially be involved in
57 the phenotypic divergence of populations is the Q_{ST} - F_{ST} comparison (Spitze, 1993; Merilä and
58 Crnokrak, 2001; McKay and Latta, 2002). The comparison of population genetic differentiation
59 estimated for putatively neutral molecular markers with the population quantitative genetic
60 differentiation estimated for phenotypic traits can be used to identify potential candidate traits
61 playing a role in local adaptation (Whitlock, 2008). This is done by estimating whether trait
62 quantitative genetic differentiation among populations is more likely the potential result of

63 divergent selection ($Q_{ST} > F_{ST}$), stabilizing selection ($Q_{ST} < F_{ST}$), or neutral evolutionary
64 divergence ($Q_{ST} = F_{ST}$, e.g., as a result of drift). Some debate around the accuracy of $Q_{ST} - F_{ST}$
65 comparisons resulted in a variety of methodological adjustments (Whitlock, 2008; Edelaar et
66 al., 2011; Ovaskainen et al., 2011; Whitlock and Gilbert, 2012). In plants, reciprocal transplants
67 directly comparing fitness between the native habitat and the foreign habitats are often preferred
68 to $Q_{ST} - F_{ST}$ approaches conducted in common gardens because they allow to evaluate the effect
69 of environmental conditions (Etterson, 2004; Angert and Schemske, 2005; Kim and Donohue,
70 2013). When the conditions for the reciprocal transplant cannot be easily met, $Q_{ST} - F_{ST}$
71 comparisons represent an opportunity for exploring local adaptation hypotheses.

72

73 In our study, we investigated patterns of potential local adaptation in two closely related plant
74 subspecies by using $Q_{ST} - F_{ST}$ comparisons estimated in a common garden experiment, and
75 evaluated whether altitudinal gradients might play a role in the potential adaptive divergence of
76 populations. This evaluation was conducted in snapdragon plants (*Antirrhinum majus* L.,
77 Plantaginaceae). We studied eight populations of magenta-flowered *A. m. pseudomajus* and five
78 populations of yellow-flowered *A. m. striatum* sampled along altitudinal gradients. These two
79 species are interfertile (Andalo et al., 2010). They are distributed parapatrically, with the
80 geographic range of *A. m. striatum* surrounded by the range of *A. m. pseudomajus*, and come
81 frequently into contact at the margins of their ranges where there is evidence for gene exchanges
82 (Khimoun et al., 2011; Ringbauer et al., 2018). Their geographic separation is not explained by
83 actual climatic differences, as illustrated by the substantial overlap of environmental conditions
84 between the two species (Khimoun et al., 2013). This system is therefore promising to explore
85 potential differential adaptive responses between closely related subspecies, in particular
86 regarding the potential role played by altitude in their adaptive divergence.

87

88 There is poor support in the literature for adaptive changes in reproductive traits along
89 altitudinal gradients (Halbritter et al., 2018). In contrast, adaptive differentiation along
90 altitudinal gradients is expected for biomass-related traits and height, with a trend toward
91 smaller plants for populations from high altitude compared to plants from lowland sites
92 (Halbritter et al., 2018). We tested this hypothesis for five morphological traits (the basal stem
93 diameter, the number of branches on the plant, the number of vegetative nodes on the main
94 stem, and the total height of the plant). We also studied three additional traits: a phenological
95 trait (the germination date), a developmental trait (the average internode length) and a
96 functional trait (specific leaf area, SLA). We expected populations from higher altitudes to
97 germinate later, over a shorter period (Gimenez-Benavides et al., 2006; Donohue et al., 2010).
98 This is because germinating later over shorter periods allows plants to track the late arrival and
99 the shorter-term availability of suitable climatic conditions for growth at higher altitudes
100 (Körner, 1999). Because the internode length is a trait related to both plant height and growth
101 rates, we had no clear expectations. Finally, SLA refers to leaf construction cost and captures
102 information about leaf economic strategies (Wright et al., 2004); low SLA suggests high leaf
103 construction cost and high stress tolerance. Selective pressures associated with lower
104 temperatures at higher elevations are expected to promote leaf trait syndromes associated with
105 superior stress tolerance but inferior competitiveness (Read et al., 2014). These relationships
106 are generally stronger among species than among populations of the same species (Read et al.,
107 2014). Therefore, we expected no correlation or a negative correlation between SLA and
108 elevation among populations.

109

110 In this study, we estimated neutral genetic differentiation (F_{ST}), and quantitative genetic
111 differentiation (Q_{ST}) based on trait heritability (h^2) in *A. majus*. Previous studies of genetic
112 differentiation between populations and subspecies at putatively neutral microsatellite markers

113 brought evidence that gene flow was limited between populations (Debout et al., 2012; Pujol et
114 al., 2017), which sets the stage for local adaptation. We then tested for the hypothesis that traits
115 were potentially involved with local adaptation by comparing Q_{ST} and F_{ST} . Finally, we
116 investigated whether quantitative genetic differentiation increased with altitudinal difference,
117 with the hypothesis that environmental changes associated with altitude, which include a suite
118 of climatic variables, drove adaptive responses. Other environmental variables (e.g.,
119 atmospheric pressure) can also change with altitude. Our study also ultimately participates in
120 evaluating whether Q_{ST} - F_{ST} comparisons can be used as a tool to identify candidate traits
121 involved with the potential adaptation of populations to altitudinal gradients, and thereby
122 climate differences.

123

124 **MATERIAL AND METHODS**

125 **Study system**

126 *Antirrhinum majus* L. (Plantaginaceae) is a hermaphroditic, self-incompatible, short-lived
127 perennial species, characterized by a patchy distribution in southern Europe centred over the
128 Pyrenees Mountains (Khimoun et al., 2011). This species occurs from sea level to an altitude
129 of 1900 m (Andalo et al., 2010), on limestone or siliceous substrates and in habitats with
130 contrasted moisture regimes (rainfall 500-1000 mm per year), where it forms restricted patches
131 mostly in rocky outcrops and screes. *A. majus* thrives in disturbed habitats, and is especially
132 common along roadside and railway embankments (Khimoun et al., 2011).

133

134 **The subspecies level**

135 *A. majus* produces annual inflorescences with zygomorphic flowers. The colour of flowers is
136 either magenta or yellow and distinguishes two interfertile subspecies *A. m. ssp. pseudomajus*
137 and *A. m. ssp. striatum* respectively (Andalo et al., 2010). At the genetic level, ~1% genetic

138 differentiation was found between *A. m. ssp. pseudomajus* and *A. m. ssp. striatum* on the basis
139 of putatively neutral microsatellite loci, which was one order of magnitude lower than the ~10%
140 differentiation found among these populations (Pujol et al., 2017). There is evidence for gene
141 exchange between subspecies in multiple populations across contact zones (Khimoun et al.,
142 2011). Genome scans across a particular contact zone in the Pyrenees also revealed little to
143 negligible differentiation between the two subspecies, with the exception of loci underlying
144 flower colour differences between the two subspecies that were characterized by high
145 differentiation (Whibley, 2006; Tavares et al., 2018). At the environmental level, the separation
146 between the geographic distribution of *A. m. ssp. pseudomajus* and *A. m. ssp. striatum* is not
147 explained by habitat differences, as illustrated by the substantial overlap of environmental
148 conditions between the two species (Khimoun et al., 2013).

149

150 **Collection sites and plant material**

151 Thirteen wild populations of *A. majus* were sampled in 2011 across the geographic range
152 (between north-eastern Spain and south-western France) to represent the overall diversity of
153 the species, with eight populations of *A. m. ssp. pseudomajus* and five populations of *A. m. ssp.*
154 *striatum* (Figure 1, Table S1, Supporting Information). For each subspecies, we sampled
155 populations from low and high altitude habitats in different parts of the species geographic
156 range. Populations sampled along elevation gradients are likely to be confronted to contrasted
157 environmental conditions. Fifty-year averages (1950-2000) of mean annual temperature and
158 annual average rainfall were extracted from the WorldClim database (resolution 1km²,
159 www.worldclim.org, Hijmans et al., 2005). They ranged from 14.8°C and 52mm (BAN, 61m
160 above sea level) to 6.1°C and 94mm (MON, 1564m above sea level) (Figure S1, Supporting
161 Information). The sampling of populations in different valleys or on different summits limits

162 spatial autocorrelation in the data and shared phylogeographic history between populations
163 from similar altitudes.

164

165 In each wild population, seeds were randomly collected from plants across their entire spatial
166 distribution in October 2011. These seeds sampled in the wild were used to grow plants in 2012,
167 in a greenhouse at the CNRS Experimental Ecology Station in Moulis, France. Seeds were sown
168 in spring in individual pots ($9 \times 9 \times 10$ cm) filled with universal compost. Plants germinated
169 and grew with no nutrient addition under an average temperature from 15 to 28°C and weekly
170 watering. Mature plants were hand-pollinated during the summer 2012. These plants were not
171 measured. Crosses were conducted within populations where mates from different families
172 were assigned randomly. The seed collection of full sib families produced by these plants was
173 stored at room temperature, in the dark, under dry conditions until they were used to produce
174 the plants measured in our experiment. This intermediate generation of plants grown in
175 controlled conditions from seeds collected in the wild allowed us to reduce potential maternal
176 effects caused by natural habitat differences on trait measurements.

177

178 **Common garden experiment**

179 Nine to 42 seed families from each of the 13 study populations were grown outdoor in spring
180 2014 in a common garden at ENSFEA (Toulouse, France). Two plants per family were grown.
181 Some plants died before measurements were done, which resulted in some families being
182 represented by only one plant (Table S1, Supporting Information). Plants were grown in
183 individual pots ($9 \times 9 \times 10$ cm) filled with universal compost, with no nutrient addition, under
184 outdoor climatic conditions (average month temperatures ranging from 20.6 to 21.5°C and
185 cumulative monthly rainfall ranging from 28.3 to 73.4mm). Plants were arranged in a
186 randomized block design (40 plastic containers, $600 \times 400 \times 120$ mm) with each containing 24

187 randomly chosen plants. The bottom of each container was covered with an irrigation sheet
188 (400 g.m⁻²) that allowed to regulate the moisture of the compost. Plants were supplied with
189 water in case of prolonged drought. Damage caused by herbivorous insects were contained by
190 using a wintering veil. This veil also limited pollination.

191

192 **Phenotypic data**

193 We measured several vegetative traits on each individual: a phenological trait (the germination
194 date), a functional trait (the specific leaf area, SLA), a developmental trait (the average
195 internode length) and multiple morphological traits. Morphological traits included the basal
196 stem diameter, the number of branches on the plant, the number of vegetative nodes on the main
197 stem, and the total height of the plant. The SLA refers to leaf construction cost and captures
198 information about species leaf economic strategies (Wright et al., 2004). It was calculated as
199 the ratio between the cumulated area of five mature but non-senescent fresh leaves and their
200 oven-dried mass (Pujol, Salager, et al., 2008; Pérez-Harguindeguy et al., 2016). Leaf area was
201 measured by using the R package Momocs v. 1.2.9 (Bonhomme et al., 2014).

202

203 **Molecular analyses**

204 To infer genetic diversity estimates in each population and to compute F_{ST} , we genotyped the
205 637 plants. DNA was extracted from silica gel dried leaf samples using the Biosprint 15 DNA
206 Plant kit (Qiagen, Hilden, Germany) following the manufacturer's instructions. Individuals
207 were genotyped for 23 putatively neutral microsatellite markers that were developed for
208 population genetic studies (Debout et al., 2012; Pujol et al., 2017). To compute F_{ST} , we used
209 population pairwise F_{ST} estimates and the overall F_{ST} estimate amongst populations from the
210 study by Pujol et al. (2017). We used the GenoDive 3.0 software (Meirmans and Van Tienderen,

211 2004) to compute the complementary parameters required for this study, e.g., the genetic
212 diversity at each locus.

213

214 **Statistical analysis**

215 All statistical analyses were performed using the R.3.5.0 software (R Core Team, 2018).

216

217 *Phenotypic traits*

218 First, to test for phenotypic differences between subspecies, hierarchical generalized linear
219 models were conducted with population nested in subspecies. Second, for each subspecies
220 linear mixed models were conducted to test for phenotypic differences among populations, with
221 population as a fixed effect and the plastic container as a random effect. Estimates of marginal
222 means for each trait in each population were extracted using the emmeans package (Lenth et
223 al., 2019). These linear mixed-effects models were implemented in R via the lme4 package
224 (Bates et al., 2015). Traits change with altitude was analysed by using a linear regression of the
225 marginal means by altitude. Finally, mean phenotypic traits were also generated, and provided
226 in the Supplementary materials (Fig S1).

227

228 *Calculation of h^2 and phenotypic differentiation indices (Q_{ST}).*

229 For each subspecies, narrow-sense heritabilities (h^2) were estimated for each phenotypic trait
230 across all populations using a model with population, family and plastic containers as random
231 factors as $h^2 = 2 V_w / (V_w + V_{res})$, with V_w is family variance component and V_{res} is the residual
232 variance component corresponding to the within-population variance component. We
233 multiplied V_w by two in the calculation of h^2 because we used a full-sib crossing design (Roff,
234 1997). Caution must be taken when considering h^2 values because its estimation can be biased
235 by the estimation of $2V_w$. Indeed, h^2 was calculated based on all the families, without

236 considering the differences of h^2 between different populations. Confidence intervals of h^2 were
237 calculated following a parametric bootstrap method adapted from O'Hara and Merilä (2005).

238

239 For each trait and each subspecies, quantitative trait divergence indices (Q_{ST}) were generated
240 among populations (overall Q_{ST}) and for each population pair (population pairwise Q_{ST}), based
241 on mixed model analyses. In these models, population, family and plastic containers were
242 random factors. Variance components were extracted from these analyses for each trait and
243 used for estimating Q_{ST} using the following formula (Spitze, 1993): $Q_{ST} = V_b / (V_b + 2 h^2 V_w)$
244 with V_b being the trait genetic variance among populations. h^2 was calculated based on all the
245 families and populations by subspecies. Here, no environmental sources of phenotypic variance
246 due to the ecological conditions of the location of origin of populations could in theory bias Q_{ST}
247 estimates because data was obtained from a common garden experiment (Pujol, Wilson, et al.,
248 2008). When a variance component was non-significant, it was considered as null in further
249 calculations. When necessary (as for population pairwise Q_{ST} calculation), data was linearized
250 by using a square root transformation. All variance components were estimated by using the
251 linear mixed model approach implemented in the R package lme4 v. 1.1.17 (Bates et al., 2015).
252 Confidence intervals of Q_{ST} values were calculated following a parametric bootstrap method
253 adapted from O'Hara and Merilä (2005).

254

255 *Overall Q_{ST} - F_{ST} comparisons.*

256 We compared overall Q_{ST} and F_{ST} for each trait to investigate if divergence was compatible
257 with a scenario of genetic drift (overall $Q_{ST} = F_{ST}$), or whether it was more likely explained by
258 directional selection (overall $Q_{ST} > F_{ST}$) or by stabilizing selection (overall $Q_{ST} < F_{ST}$).
259 Comparisons between overall Q_{ST} and F_{ST} values were performed for each trait based on two
260 methods: i) a comparison of confidence intervals, the Q_{ST} is considered non significantly

261 different from neutral differentiation when the confidence interval of the overall Q_{ST} for a trait
262 overlaps the mean F_{ST} value, ii) a bootstrapping method developed by Whitlock and Guillaume
263 (2009). This last approach aims at comparing the observed difference between the overall Q_{ST}
264 and the F_{ST} with the expected simulated distribution of this difference under a scenario of
265 neutral evolution. We generated 100 000 bootstrap replicates of the expected $Q_{ST} - F_{ST}$
266 difference under the neutrality hypothesis for each trait, and built the corresponding
267 distribution. In this approach, P values were estimated by assessing whether the observed value
268 of the $Q_{ST} - F_{ST}$ difference overlapped its expected distribution under neutrality. We used the
269 modification by Lind *et al.* (2011) of the approach of Whitlock and Guillaume (2009) to
270 estimate the variance components of the simulated values of the $Q_{ST} - F_{ST}$ difference.

271

272 *Mantel tests*

273 Mantel tests (Mantel, 1967) were used to analyse correlations between geographic distances,
274 environmental distances (altitudinal), neutral genetic differentiation (population pairwise F_{ST}),
275 and quantitative genetic differentiation (population pairwise Q_{ST}). First, a correlation test
276 between population pairwise F_{ST} and population pairwise geographic distance matrices was
277 performed to test for an isolation by distance relationship. Second, a correlation test between
278 population pairwise F_{ST} and population pairwise Q_{ST} was performed for each trait to test if
279 neutral genetic differentiation explained divergence in quantitative traits. Third, a correlation
280 test between population pairwise Q_{ST} and population pairwise altitudinal differences was
281 performed for each trait to test whether divergence in quantitative traits was related to altitudinal
282 differences. Finally, we conducted partial mantel tests to test for the association between
283 population pairwise Q_{ST} and population pairwise altitude differences while controlling for
284 neutral genetic differentiation (F_{ST}). All mantel and partial mantel tests were performed in R,
285 with a significance $\alpha=0.05$, using the vegan package (Oksanen et al., 2009).

286 RESULTS

287 Phenotypic differentiation between subspecies and populations

288 The two subspecies - *A. m. pseudomajus* and *A. m. striatum* - were significantly differentiated
289 by several phenotypic traits (Table 1 a, Figure S2). When grown in a common garden, plants
290 of *A. m. pseudomajus* were on average taller, with more branches and nodes than plants from
291 *A. m. striatum*. Their average germination date, internode length and SLA were however similar.
292 Differentiation between subspecies (c. 2.8 %) was lower than the differentiation estimated
293 among populations (c. 9.3 %, see mean R^2 in Table 1 a). Most of the traits showed a phenotypic
294 divergence among populations within each subspecies (see LRT in Table 1 b). Germination
295 date was the only trait that showed no significant difference among populations of *A. m.*
296 *pseudomajus* (see LRT in Table 1 b).

297

298 Neutral genetic differentiation

299 Population neutral genetic differentiation was low but significant. Overall F_{ST} among
300 populations of *A. m. pseudomajus* was 0.109 ($P < 0.001$), and ranged from 0.06 to 0.159 across
301 population pairs (see Table S2, and see Pujol et al., 2017 for more details on population pairwise
302 neutral genetic differentiation). F_{ST} among populations of *A. m. striatum* was 0.097 ($P < 0.001$),
303 and ranged from 0.055 to 0.131 (Table S2). There was no significant relationship between
304 population pairwise F_{ST} and population pairwise geographic distance for both subspecies
305 (Figure 2 a and b, *A. m. pseudomajus* Mantel $r = 0.04$, $P = 0.405$, *A. m. striatum* Mantel $r = -$
306 0.18 , $P = 0.595$). Similarly, there was no significant relationship between population pairwise
307 F_{ST} and population pairwise altitude difference for both subspecies (Figure 2 c), although the
308 Mantel tests showed a relationship close to significance levels in *A. m. pseudomajus* (*A. m.*
309 *pseudomajus* Mantel $r = 0.23$, $P = 0.052$, *A. m. striatum* Mantel $r = -0.3$, $P = 0.943$).

310

311 **Changes in phenotypic traits with altitude**

312 We found significant correlations between trait values (i.e. population estimates of marginal
313 means) and altitude for two traits across *A. m. striatum* populations. Plants from populations at
314 low altitude had more nodes and branches than plants from populations at high altitude of *A. m.*
315 *striatum* (Figure 3, see population arithmetic means in Figure S2 and population estimates of
316 marginal means for other traits in Figure S3). No phenotypic changes associated with altitude
317 were found significant in *A. m. pseudomajus*.

318

319 **Inheritance of quantitative traits**

320 Heritability estimates were comprised between 0.07 to 0.58 for *A. m. pseudomajus*, and 0.01 to
321 0.56 for *A. m. striatum* (Table S3, Supporting Information). The highest heritability estimates
322 were found for the internode length in *A. m. pseudomajus* (0.58) and the number of nodes in *A.*
323 *m. striatum* (0.56). Several traits had close to similar heritabilities between subspecies (stem
324 diameter, number of nodes), as illustrated by their overlapping confidence intervals. However,
325 other traits appeared to be different, with no CI overlap (germination date, number of branches,
326 plant height, internode length, SLA, Table S3).

327

328 **Q_{ST} - F_{ST} comparisons.**

329 Overall Q_{ST} was higher than mean F_{ST} for one trait in *A. m. pseudomajus* (number of branches,
330 Figure 4 a), and for three traits in *A. m. striatum* (number of branches, plant height and internode
331 length, Figure 4 b), as illustrated by their non-overlapping confidence intervals. Overall Q_{ST}
332 was lower than mean F_{ST} for the germination date in *A. m. pseudomajus* (Figure 4 a). We also
333 rejected the hypothesis of neutrality for these two traits in *A. m. pseudomajus* (number of
334 branches, germination date), and for three traits in *A. m. striatum* (number of branches, plant
335 height and internode length) on the basis of the bootstrapping method developed by Whitlock

336 and Guillaume (2009). For these traits, observed values of overall $Q_{ST} - F_{ST}$ differences were
337 either in the tail of the expected probability distribution under the hypothesis of neutrality, or
338 did not overlap with this distribution (Figure S4 et S5).

339 Mantel tests showed that population pairwise quantitative genetic differentiation (Q_{ST}) was not
340 correlated with population pairwise neutral genetic differentiation (F_{ST}) for all traits excepted
341 the germination date in *A. m. striatum* (Table 2). Population pairwise Q_{ST} for the germination
342 date was significantly correlated with population pairwise F_{ST} . They showed no support for the
343 overall $Q_{ST} < F_{ST}$ for the germination date in *A. m. pseudomajus*, almost certainly because of
344 three comparisons diverging abnormally from the overall pattern (Figure 5 b).

345

346 **Increased quantitative genetic differentiation with altitude difference**

347 Mantel tests showed a significant correlation between population pairwise Q_{ST} and population
348 pairwise altitudinal difference for two traits in *A. m. striatum*: the number of nodes and the
349 number of branches (Table 2, Figure 5). For both traits, the increase in pairwise population
350 differentiation associated with an increase in altitudinal difference was higher for the Q_{ST} than
351 for the F_{ST} (Figures 5 c and e). Partial mantel tests showed that population pairwise Q_{ST} was
352 significantly correlated with differences in altitude for the number of nodes (and marginally
353 significant for the number of branches, see Q_{ST} vs Alt. diff. / F_{ST} in Table 2) while controlling
354 for neutral genetic differentiation (F_{ST} matrix). This result is expected under the hypothesis that
355 the divergence among populations of *A. m. striatum* in the number of nodes is a result of
356 altitude-mediated divergent selection. In contrast, none of the seven traits showed a significant
357 correlation between population pairwise Q_{ST} and population pairwise altitude difference in *A.*
358 *m. pseudomajus*.

359

360 **DISCUSSION**

361 Our results support the hypothesis of differential adaptation between *A. m. pseudomajus* and *A.*
362 *m. striatum* subspecies. We detected phenotypic differentiation in a common garden among the
363 populations of *A. m. pseudomajus*, among the populations of *A. m. striatum*, and among
364 subspecies. For both subspecies, local adaptation and neutral evolution explained the extent to
365 which populations diverged over their geographic range, with slight differences between
366 subspecies. Potential divergence along altitude was also detected, but only for one subspecies:
367 *A. m. striatum*.

368

369 Our findings comforted the idea that Q_{ST} - F_{ST} comparisons are a good first step for exploring the
370 potential roles of divergent natural selection and neutral evolutionary processes in phenotypic
371 divergence (Whitlock, 2008; Edelaar et al., 2011; Ovaskainen et al., 2011; Whitlock and
372 Gilbert, 2012). They highlighted how traits can be used to identify the potential ecological
373 pressures underlying natural selection, with some traits potentially involved with *A. majus*
374 adaptation to the conditions of populations' local sites of origin, and a subsample of these traits
375 potentially playing a role in *A. m. striatum* adaptation to altitude.

376

377 **Adaptive evolution of *A. m. striatum* populations along the altitudinal gradient**

378 Our results brought indirect evidence supporting potential adaptive divergence between *A. m.*
379 *pseudomajus* and *A. m. striatum*. They imply that the quantitative genetic basis of two of the
380 seven traits under study (number of nodes, and marginally significant for the number of
381 branches) was shaped by divergent selection between populations from different altitudes in *A.*
382 *m. striatum* but not in *A. m. pseudomajus*. Most studies on plant adaptation to altitude report
383 the selection of smaller plants at higher altitudes (Körner, 1999; Halbritter et al., 2018). In
384 agreement with this expectation, we found that *A. m. striatum* plants at higher altitudes had less

385 branches and less nodes. There is also evidence for changes in leaf traits with elevation (Read
386 et al., 2014; Halbritter et al., 2018), with a decrease of SLA with elevation gradients. Our results
387 did not support a potential scenario of selection based on SLA at play in *A. m. striatum*.

388

389 **Support for different subspecies scenarios of adaptation to local sites of origin**

390 Our results showed that quantitative genetic differentiation was higher than what could be
391 explained by neutral evolutionary divergence among *A. m. pseudomajus* populations for one of
392 the seven studied traits (number of branches), and among *A. m. striatum* populations for three
393 of the seven traits (number of branches, plant height and internode length). They imply that
394 adaptation to local sites of origin potentially shaped the phenotypic diversity of populations for
395 both subspecies across their geographic range, with potentially different intensities reflected by
396 different signatures between subspecies. We used classical overall $Q_{ST} - F_{ST}$ comparisons to
397 detect potential adaptation to local sites conditions (Leinonen et al., 2008) and also more recent
398 methods to insure that our findings were robust against a range of neutral evolution scenarios
399 for these traits (Whitlock, 2008). Furthermore, our approach minimized the possibility that
400 phenotypic differences between populations were generated by environmental effects by using
401 a common garden experiment, and including trait heritability estimates in Q_{ST} calculations
402 (Spitze, 1993; Pujol, Wilson, et al., 2008). In contrast, four of the seven studied traits
403 (germination date, diameter, number of nodes and SLA) did not show departure from plausible
404 baseline scenarios of neutral evolutionary divergence, using overall $Q_{ST} - F_{ST}$ comparisons. One
405 particular trait (germination date) was in fact more similar among populations than expected
406 under neutrality in *A. m. pseudomajus*. A scenario of stabilizing selection is classically
407 extrapolated in the case of similar results (Lamy et al., 2012) but another plausible explanation
408 is that population similarity might have been caused by convergent phenotypic responses to the
409 common garden environmental similarity. Caution must be taken when interpreting different

410 Q_{ST} - F_{ST} patterns between subspecies as the signature of different adaptive processes. Here we
411 found different patterns between subspecies, which supports the hypothesis of their potential
412 adaptive divergence. Our results cannot be interpreted as direct proof for their adaptive
413 divergence, but only as evidence that this hypothesis has some potential.

414

415 **The ecological significance of adaptation to local sites of origin in *A. majus***

416 In the absence of environmental measures included in the overall Q_{ST} - F_{ST} analysis, it is
417 impossible to identify the potential environmental agents of local selection that shape the
418 quantitative genetic variation of traits. The functions behind the traits that have diverged can
419 nevertheless be used to discuss plausible evolutionary scenarios of natural selection. Our results
420 imply that adaptation to local sites of origin has potentially shaped the vegetative architecture
421 of plants that is specific to each *A. majus* population. The quantitative genetic variation of
422 several phenotypic traits characterising the vegetative growth and development of plants (plant
423 height, internode length, number of branches) has likely diverged among populations as a result
424 of adaptation to local sites of origin. Divergence in the genetic variation underlying the shape
425 and size of plants was already found at the level of *Antirrhinum* species but its adaptive
426 significance was not tested for (Langlade et al., 2005). In southern France and northern Spain,
427 under the Mediterranean climate, dryer locations are expected to select for plants with a bushier
428 vegetative architecture, i.e. plants with smaller leaves and more branches that have a better
429 water use efficiency and resilience to drought stress (Langlade et al., 2005). It is difficult to
430 identify exactly which environmental pressures underlay selection at local sites because several
431 combinations of environmental parameters (vegetation cover, wind, disturbance, temperature,
432 water availability, etc.) can interact to affect phenotypic traits.

433

434 **Gene flow, ecological and reproductive isolation**

435 Our findings imply that the most likely evolutionary scenario applying to *A. majus* requires
436 invoking a history of adaptation to local sites in a complex background of gene flow, ecological
437 heterogeneity and reproductive isolation. Pyrenees mountains are widely acknowledged to
438 constitute a heterogeneous landscape promoting complex patterns of population connectivity
439 and prone to generate local adaptation (Alberto et al., 2010). Q_{ST} - F_{ST} comparisons reflected a
440 potential scenario of population divergent adaptation to contrasting environmental conditions
441 between their local sites of origins. Our findings also suggested that evolutionary signatures of
442 local adaptation differed between *A. m. pseudomajus* and *A. m. striatum*, which includes the
443 potential adaptation to altitude of *A.m. striatum* populations. One might speculate that this
444 divergence might be related to the distribution of *A. m. striatum* populations across a narrower
445 range of climatic conditions, even if both subspecies share to a large extent the same ecological
446 niche (Khimoun et al., 2013). However, caution must be taken with this explanation because
447 the state of the environment in the past, when divergence might have occurred, is unknown and
448 might have differed. Contrasting hypotheses might be interesting to consider, e.g., different
449 evolutionary potentials in the presence of similar environmental pressures. These scenarios are
450 not exclusive and can reinforce each other through a feedback loop between reproductive
451 isolation, neutral divergence and selection.

452

453 Restricted gene flow or strong selection pressures are required for evolutionary divergence.
454 Genetic drift, or foundation events by different gene pools, might have shaped differentially the
455 genetic background of *A. majus* populations and to some extent subspecies at the scale of their
456 global geographic range. There is evidence for the genetic signature of restricted gene
457 exchanges in *A. majus* (Pujol et al., 2017). No genetic isolation by distance was found but
458 ecological barriers characterizing the mountain landscape of the Pyrenees likely participate to
459 isolate populations (Pujol et al., 2017). At first sight, *A. majus* subspecies divergence might not

460 be expected because both subspecies are interfertile (Andalo et al., 2010), and no genome wide
461 barrier to gene flow was found between them at the scale of a hybrid zone across c. 2km in the
462 Pyrenees (Ringbauer et al., 2018). There is also evidence for gene exchanges between the two
463 subspecies in several contact zone locations across at the periphery of their geographic ranges
464 (Khimoun et al., 2011). Yet, subspecies flower color differences attest that flower color genes
465 are under frequency dependent selection and generate reproductive isolation between
466 subspecies (Tastard et al., 2012; Ringbauer et al., 2018). This reproductive isolation might
467 participate to the subspecies phenotypic divergence of other traits that we detected here.

468

469 **CONCLUSION**

470 Our findings corroborate the utility of $Q_{ST} - F_{ST}$ approaches conducted in common garden
471 experiments to explore potential adaptive evolutionary divergence among populations and
472 between subspecies in plants. They also illustrate the limit of this approach that identifies traits
473 that might be involved with local adaptation but does not bring direct evidence for their
474 response to selection. Here, our common garden results for *A. m. pseudomajus* and *A. m.*
475 *striatum* populations identified vegetative traits that might play a role in the local adaptation
476 and the differential adaptation of *A. m. pseudomajus* and *A. m. striatum* along altitudinal
477 gradients. They suggest that the adaptation to climate variables of otherwise interfertile
478 subspecies might differ as a result of reproductive isolation.

479

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488 LABX-25-01).

489

490 TABLE 1
 491 Effects of subspecies and populations on phenotypic traits. a) R² and P-value from hierarchical
 492 generalized linear models (GLM) with subspecies alone and populations nested in subspecies,
 493 implemented in JMP® (Version X ,SAS Institute Inc., Cary, NC, 1989-2019). b) Likelihood
 494 Ratio Tests (LRT) comparing the maximum-likelihood fit between a model where populations
 495 were pooled and a model estimating the effect of the population of origin. A significant P- value
 496 means the model including populations effect fitted the data better than the null model.
 497 Significant results (P-value < 0.05) are in bold.

a)	Subspecies		Populations in subspecies	
	R ²	P-value	R ²	P-value
Germination date	0.0005	0.587	0.02	0.260
Diameter	0.007	0.00028	0.05	0.00068
Nodes	0.06	<0.0001	0.13	<0.0001
Branches	0.03	<0.0001	0.06	0.00001
Plant height	0.09	<0.0001	0.21	<0.0001
Internode length	0.0002	0.708	0.13	0
SLA	0.005	0.066	0.05	0.0003
Mean	0.028		0.093	

b)	<i>A. m. pseudmomajus</i>		<i>A. m. striatum</i>	
	LRT	P-value	LRT	P-value
Germination date	6	0.570	12	0.021
Diameter	18	0.001	23	<0.0001
Nodes	30	<0.0001	20	<0.0001
Branches	70	<0.0001	26	<0.0001
Plant height	32	<0.0001	81	<0.0001
Internode length	64	<0.0001	37	<0.0001
SLA	21	0.004	15	0.004

498

499

500 TABLE 2

501 Mantel tests and partial Mantel tests on pairwise Q_{ST} vs F_{ST} and Q_{ST} vs difference in altitude of
 502 origin (Alt. diff.), as well as partial Mantel tests on Q_{ST} vs Alt. diff. controlled for F_{ST} , for
 503 phenology traits in a) eight populations of *A. m. pseudomajus* and b) five populations of *A. m.*
 504 *striatum*, that were grown in a common garden. Significant values are indicated in bold.

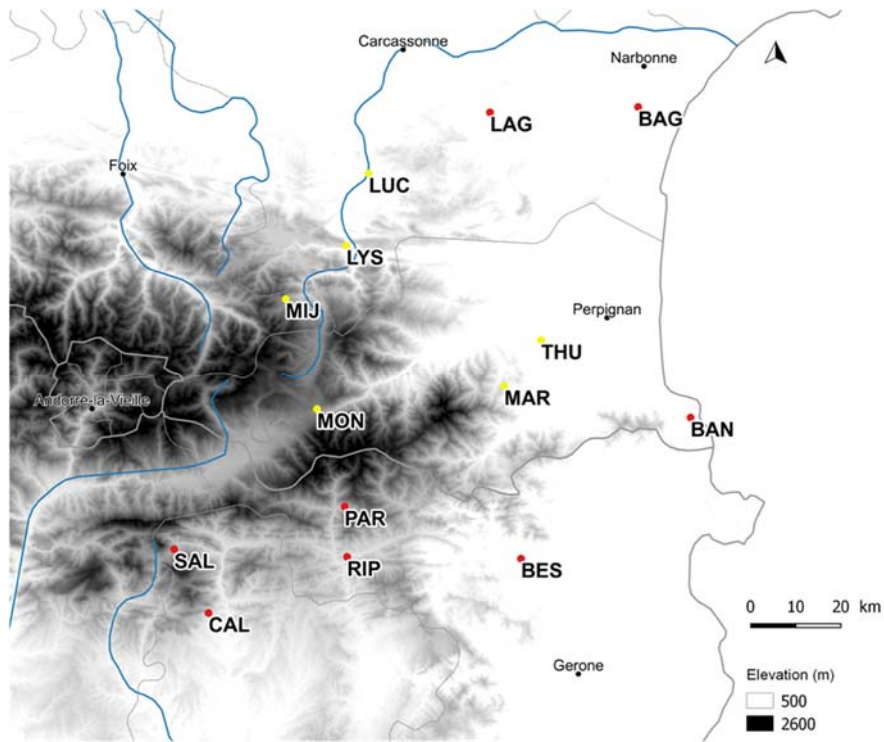
Traits	a) <i>A. majus pseudomajus</i>					
	Q_{ST} vs F_{ST}		Q_{ST} vs Alt. diff.		Q_{ST} vs Alt. diff. / F_{ST}	
	Mantel r	P-value	Mantel r	P-value	Mantel r	P-value
Germination date	-0.37	0.931	-0.13	0.737	-0.06	0.593
Diameter	-0.09	0.636	-0.15	0.812	-0.13	0.780
Nodes	0.07	0.426	-0.16	0.820	-0.18	0.888
Branches	0.07	0.329	-0.14	0.750	-0.17	0.820
Height	0.23	0.191	-0.13	0.751	-0.19	0.911
Internode length	0.24	0.184	0.03	0.335	-0.02	0.442
SLA	0.23	0.229	0.02	0.379	-0.04	0.529

Traits	b) <i>A. majus striatum</i>					
	Q_{ST} vs F_{ST}		Q_{ST} vs Alt. diff.		Q_{ST} vs Alt. diff. / F_{ST}	
	Mantel r	P-value	Mantel r	P-value	Mantel r	P-value
Germination date	0.53	0.042	0.05	0.333	0.26	0.267
Diameter	-0.01	0.508	-0.02	0.458	-0.03	0.517
Nodes	-0.3	0.842	0.95	0.008	0.94	0.008
Branches	-0.18	0.750	0.89	0.033	0.90	0.058
Height	-0.58	0.883	0.08	0.283	-0.12	0.6
Internode length	0.34	0.267	-0.02	0.492	0.09	0.258
SLA	-0.69	0.883	0.07	0.367	-0.2	0.858

505

506

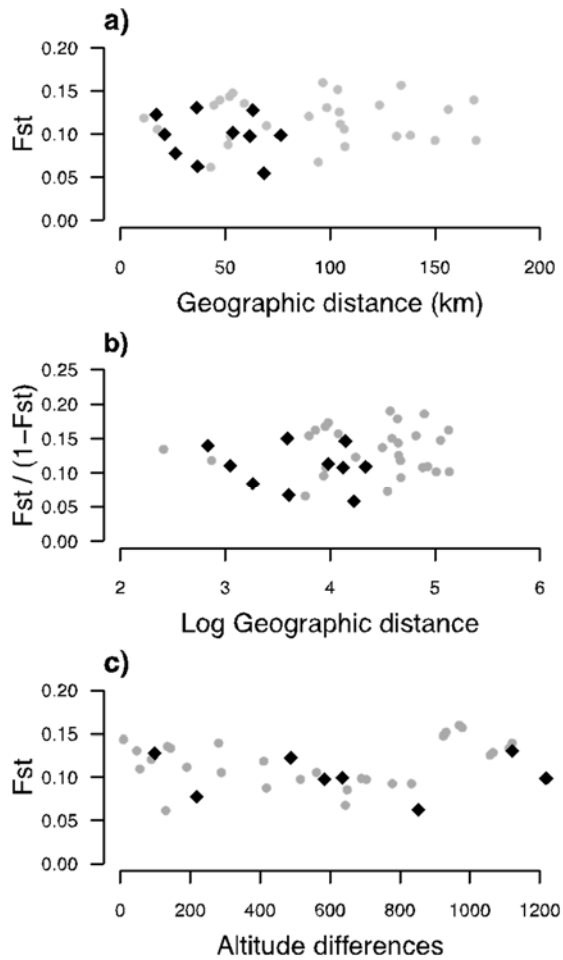
507 **FIGURE**



508

509 **FIGURE 1**

510 Map of *Antirrhinum majus* populations that were sampled across the geographic range of the
511 species in Southern France. Red dots represent *A. m. pseudomajus* populations, yellow dots
512 represent *A. m. striatum* populations. Population names and description can be found in Table
513 S1.

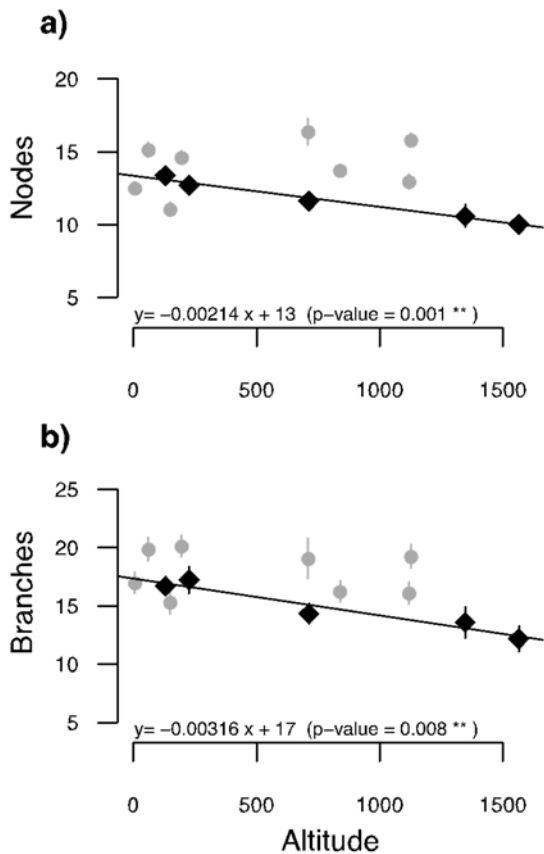


514

515 FIGURE 2

516 Pairwise neutral genetic differentiation F_{ST} plotted against pairwise geographic distances or
 517 altitudinal differences among eight *Antirrhinum majus pseudomajus* populations pairs (grey
 518 dots), and five *A. m. striatum* populations pairs (black diamonds). There were non-significant
 519 linear regression between a) F_{ST} and geographic distance in *A. m. pseudomajus* ($y= 1e-05 x +$
 520 0.116 , $P\text{-value} = 0.92$ ns) and in *A. m striatum* ($y= -0.00019 x + 0.106$, $P = 0.67$ ns), b) $F_{ST} / (1-$
 521 $F_{ST})$ and the log of geographic distance in *A. m. pseudomajus* ($y= 0.0022 x + 0.124$, $P = 0.83$
 522 ns) and in *A. m striatum* ($y= -0.0105 x + 0.148$, $P = 0.63$ ns), c) F_{ST} and altitude differences in
 523 *A. m. pseudomajus* ($y= 2.e-05 x + 0.109$, $P = 0.26$ ns) and in *A. m striatum* ($y= -2e-05 x + 0.111$,
 524 $P = 0.39$ ns).

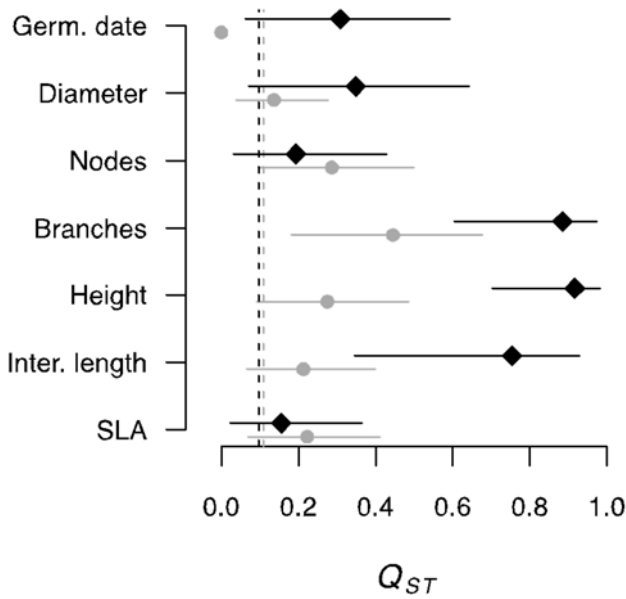
525



526

527 FIGURE 3

528 Population estimates of marginal means with standard errors of two phenotypic traits (a. number
 529 of nodes, b. number of branches) in populations of two subspecies of *Antirrhinum majus* grown
 530 in a common garden. Means are plotted against altitude of origin. Lines refer to the linear
 531 regression between trait mean estimates and altitude. Grey dots represent *A. m. pseudomajus*
 532 populations, black diamonds represent *A. m. striatum* populations. Equation of non-significant
 533 linear regressions were a) $y = 0.00125x + 12$ ($P = 0.43$ ns) and b), $y = -0.00048x + 18$ ($P = 0.78$
 534 ns) for *A. m. pseudomajus*.



535

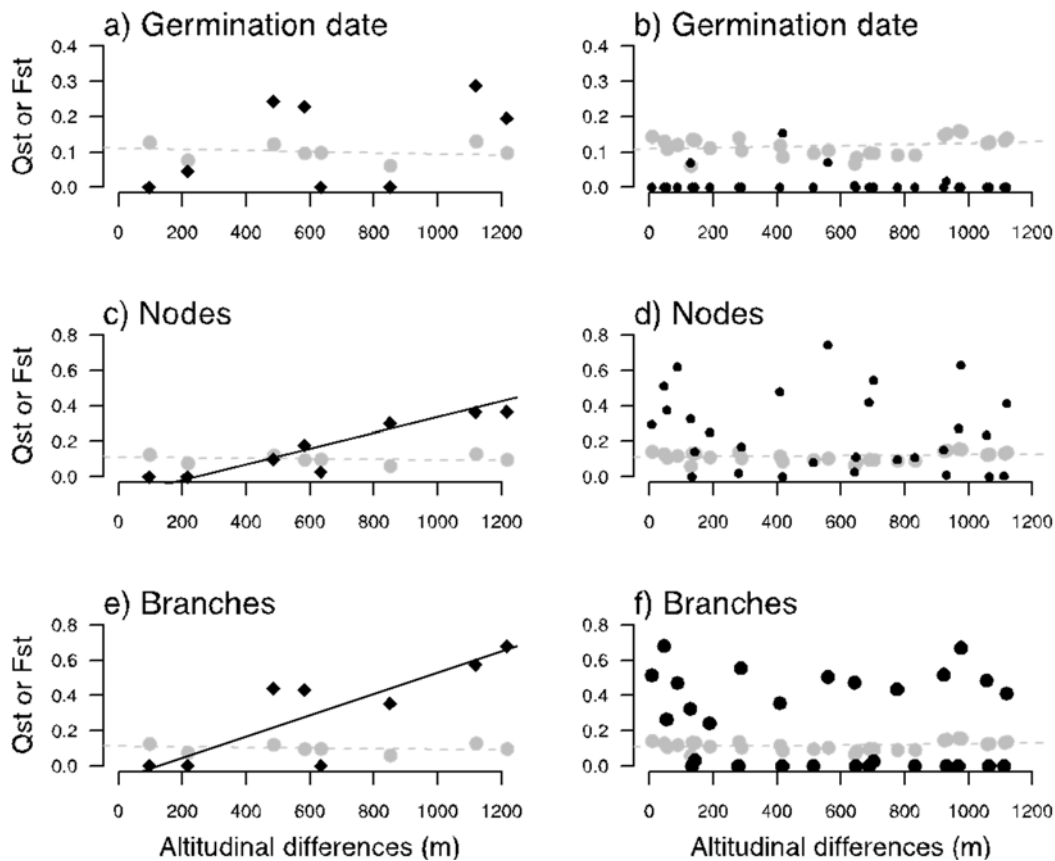
536 FIGURE 4

537 Overall Q_{ST} estimates with their 95% CI characterizing seven phenotypic traits in eight
 538 *Antirrhinum majus pseudomajus* populations (grey dots) and five *A. m. striatum* populations
 539 (black diamonds) that were grown in a common garden. Average population F_{ST} is represented
 540 by the dashed grey line for *A. m. pseudomajus*, and the dashed black line for *A. m. striatum*.

541 Germ.date = germination date, Diameter= stem diameter, Nodes = number of nodes, Branches
 542 = number of branches, Height= plant height, Inter. Length= internodes length, SLA= specific
 543 leaf area.

544

545



546

547 FIGURE 5

548 Population pairwise quantitative trait differentiation (Q_{ST}) for the germination date, the number

549 of branches and the number of nodes in *Antirrhinum majus striatum* (a, c and e, black diamonds)

550 and *A. m. pseudomajus* (b, d and f, black dots). Dashed line indicates the trend and P the level

551 of significance of the linear regression between the population pairwise Q_{ST} and population

552 pairwise altitudinal differences (m). Grey dots and dashed line refer to population neutral

553 genetic differentiation (F_{ST}). Equation of linear regressions were a) $y = 1e-05 x + 0.09$ ($P = 0.9$

554 ns), b) $1e-05 x + 0.018$ ($P = 0.5$ ns), c) $y = 0.00044 x - 0.107$ ($P = 0$ ***), d) $y = -9e-05 x + 0.303$

555 ($P = 0.42$ ns), e) $y = 6e-04 x - 0.076$ ($P = 5e-04$ ***), f) $y = -1e-04 x + 0.302$ ($P = 0.46$ ns).

556

557

558

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678

679

680 **Supplementary material**681 **TABLES**682 TABLE S1 Description of *Anthrimum majus* populations grown in the common garden683 experiment. *Nfam*= number of families, *N*= number of plants

Acronym	Latitude	Longitude	Location	Elevation (m)	Subspecies	Description	<i>Nfam</i>	<i>N</i>
BAG	43.10	2.98	Bages	6	<i>pseudomajus</i>	Dunes on seaside (rocky / herbaceous)	40	67
BAN	42.49	3.12	Banyuls-sur-Mer	61	<i>pseudomajus</i>	Rockside bank (rocky)	32	54
THU	42.64	2.72	Thuir	130	<i>striatum</i>	Roadside bank (herbaceous)	34	60
LAG	43.09	2.58	Lagrasse	149	<i>pseudomajus</i>	Roadside bank (rocky / herbaceous)	32	55
BES	42.21	2.67	Besalú	195	<i>pseudomajus</i>	Stone walls in village	39	69
LUC	42.97	2.26	Luc-sur-Aude	227	<i>striatum</i>	Roadside bank and river-side bank (rocky)	19	29
RIP	42.21	2.20	Ripoll	709	<i>pseudomajus</i>	Roadside bank (herbaceous)	9	16
LYS	42.83	2.20	'Pierre-Lys' gorge	713	<i>striatum</i>	Roadside bank (rocky / herbaceous)	32	53
CAL	42.10	1.83	Berga	838	<i>pseudomajus</i>	Roadside bank (herbaceous)	42	69
PAR	42.31	2.20	Pardines	1118	<i>pseudomajus</i>	Roadside bank (herbaceous)	32	58
SAL	42.23	1.74	Saldes	1126	<i>pseudomajus</i>	Banks in pasture (herbaceous)	30	55
MIJ	42.73	2.04	Mijanès	1347	<i>striatum</i>	Roadside bank (herbaceous)	10	18
MON	42.51	2.12	Mont-Louis citadelle	1564	<i>striatum</i>	Stone walls on fortifications	21	34
<i>All populations</i>							372	637

684 TABLE S2

685 Population pairwise F_{ST} for a) *Anthirinum majus pseudomajus* and b) *A. m. striatum*.

a) *Anthirinum majus pseudomajus*

Obs.	BAG	BAN	BES	CAL	LAG	PAR	RIP	SAL
BAG	0.000000	0.109661	0.111560	0.093282	0.134302	0.134404	0.097873	0.140489
BAN	0.109661	0.000000	0.136375	0.093343	0.120889	0.125510	0.086353	0.128994
BES	0.111560	0.136375	0.000000	0.068186	0.131472	0.147507	0.098488	0.152424
CAL	0.093282	0.093343	0.068186	0.000000	0.099317	0.139897	0.062174	0.106419
LAG	0.134302	0.120889	0.131472	0.099317	0.000000	0.159517	0.106148	0.156761
PAR	0.134404	0.125510	0.147507	0.139897	0.159517	0.000000	0.119122	0.144271
RIP	0.097873	0.086353	0.098488	0.062174	0.106148	0.119122	0.000000	0.088219
SAL	0.140489	0.128994	0.152424	0.106419	0.156761	0.144271	0.088219	0.000000

b) *Anthirinum majus striatum*

Obs	LUC	LYS	MIJ	MON	THU
LUC	0.000	0.123	0.131	0.102	0.128
LYS	0.123	0.000	0.100	0.063	0.098
MIJ	0.131	0.100	0.000	0.078	0.099
MON	0.102	0.063	0.078	0.000	0.055
THU	0.128	0.098	0.099	0.055	0.000

686

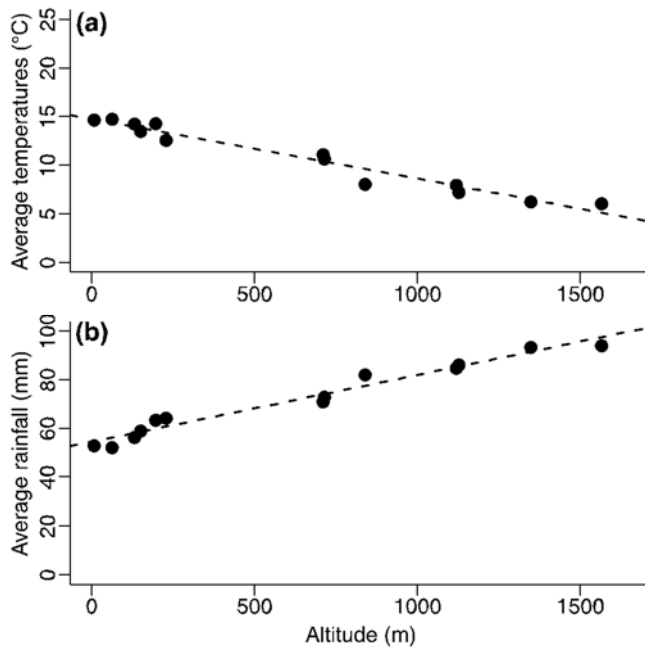
687

688 TABLE S3

689 Quantitative genetics parameters for phenotypic traits among eight populations of *Anthrimum*
 690 *majus pseudomajus* and five populations of *Anthrimum majus striatum* grown in a common
 691 garden. Values for trait heritability (h^2), family variance (V_w), among-population variance (V_b),
 692 residual variance corresponding to the within-population variance (V_{res}). The degrees of
 693 freedom used in the bootstrapping procedures are seven for the among-population component
 694 (V_b) for *A.m. pseudomajus* and four for *A. m. striatum*. Degrees of freedom are given in this
 695 table for the within-population component (dfVw).

Traits	h^2	$h^2.CI$	V_w	V_b	V_{res}	dfVw
a) <i>A. m. pseudomajus</i>						
Germination date	0.35	0.3; 0.42 0.14;	14.05	0	65.13	246
Diameter	0.17	0.20 0.39;	0.06	0.02	0.69	248
Nodes	0.45	0.52 0.06;	3.03	2.62	10.37	248
Branches	0.07	0.09 0.26;	1.54	2.83	42.11	248
Plant height	0.32	0.38 0.51;	19.71	16.1	105.45	184
Internode length	0.58	0.67 0.10;	0.08	0.05	0.19	248
SLA	0.12	0.15	128.26	78.01	1973.77	248
b) <i>A. m. striatum</i>						
Germination date	0.11	0.09; 0.15	4.37	4.49	71.98	110
Diameter	0.20	0.26 0.46;	0.06	0.08	0.57	111
Nodes	0.56	0.69 0.01;	2.94	1.55	7.6	111
Branches	0.01	0.01 0.04;	0.14	3.52	26.14	111
Plant height	0.06	0.08 0.04;	2.48	92.94	84	82
Internode length	0.05	0.06 0.27;	0.01	0.05	0.23	111
SLA	0.34	0.43	397.54	155.9	1942.69	111

696 **Figures**

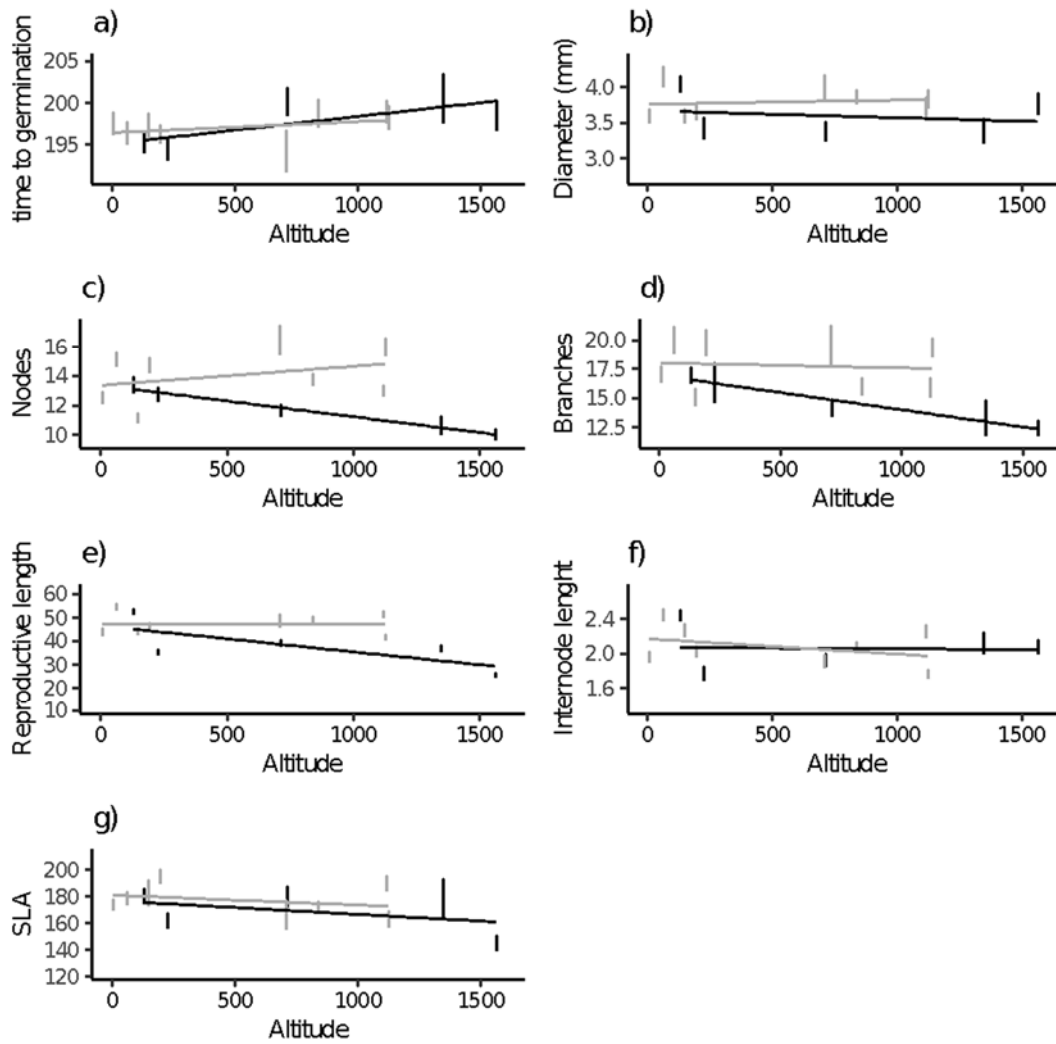


697

698 **FIGURE S1.**

699 Annual average temperatures and rainfall of 13 *Antirrhinum majus* populations from the
700 Southern France. Population average temperature (a) and average rainfall (b) as a function of
701 altitude. Bioclimatic data was extracted from the *WorldClim* database (www.worldclim.org).

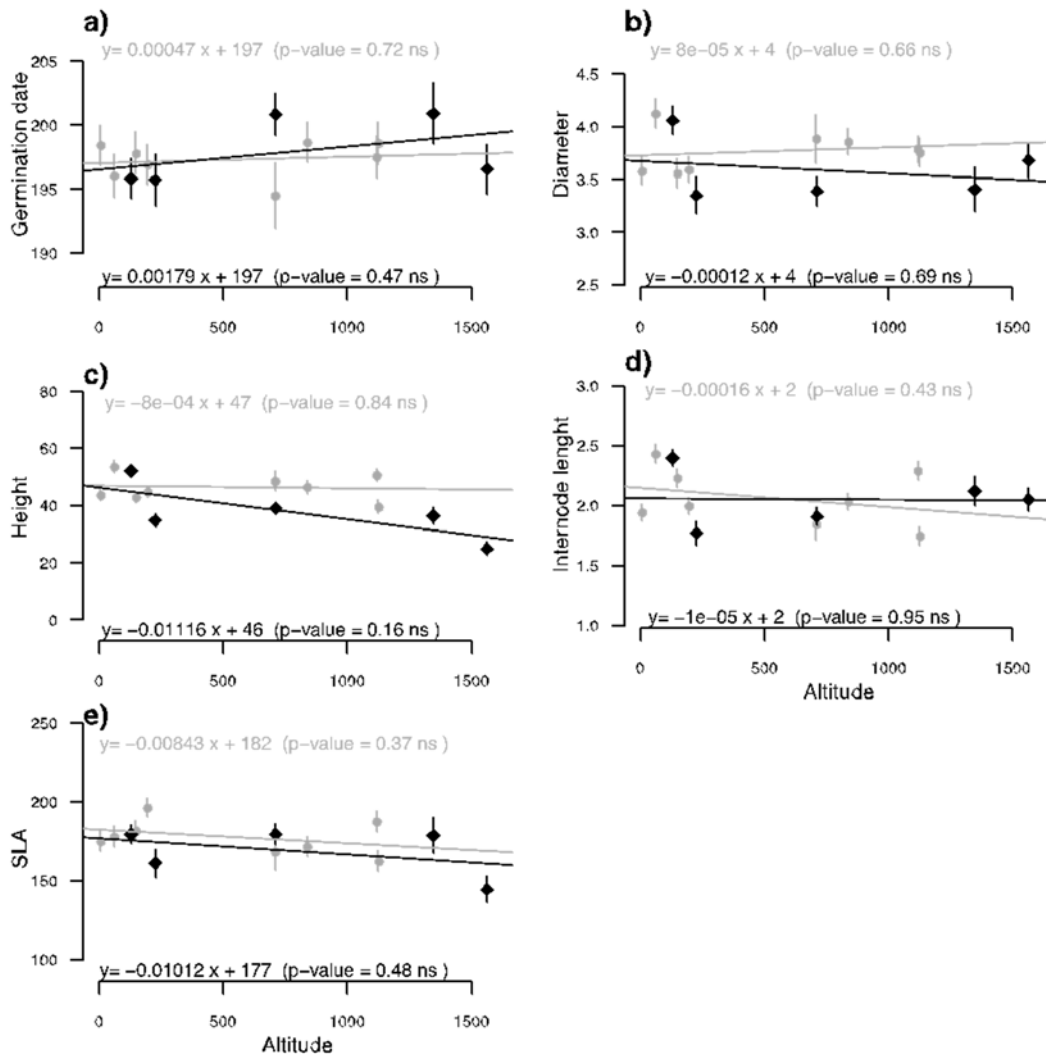
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704 FIGURE S2.

705 Population arithmetic means with standard errors of seven phenotypic traits in populations of
 706 two subspecies of *Antirrhinum majus* grown in a common garden. Means are plotted against
 707 altitude of origin. Grey dots represent *A. m. ssp. pseudomajus* populations, black diamonds
 708 represent *A. m. ssp. striatum* populations.

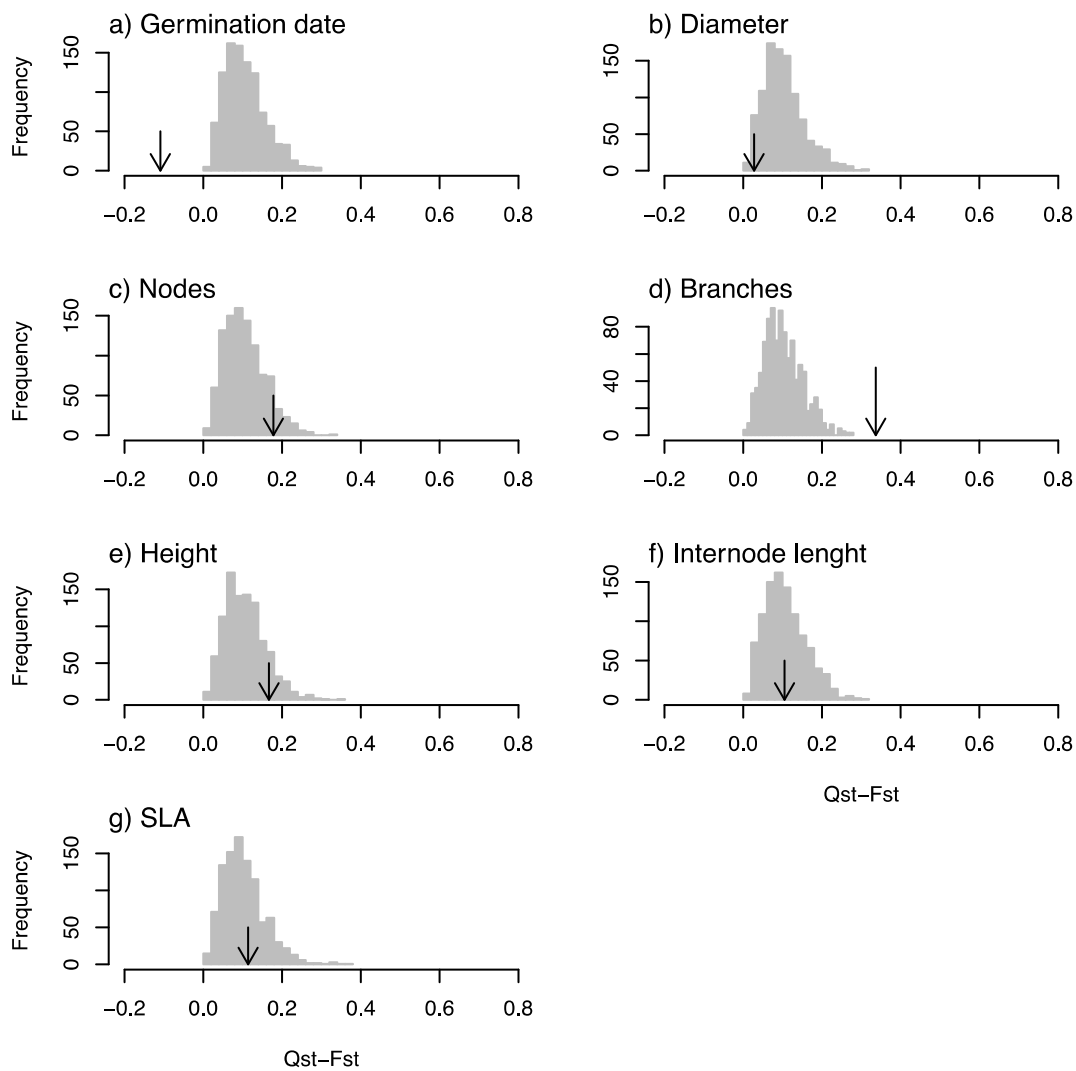


709

710 FIGURE S3. Population estimates of marginal means with standard errors of five phenotypic
 711 traits in populations of two subspecies of *Antirrhinum majus* grown in a common garden. Means
 712 are plotted against altitude of origin. Lines refer to the linear regression between traits means
 713 estimates and altitude. Grey dots and lines represent *A. m. pseudomajus* populations, black
 714 diamonds and lines represent *A. m. ssp. striatum* populations.

715

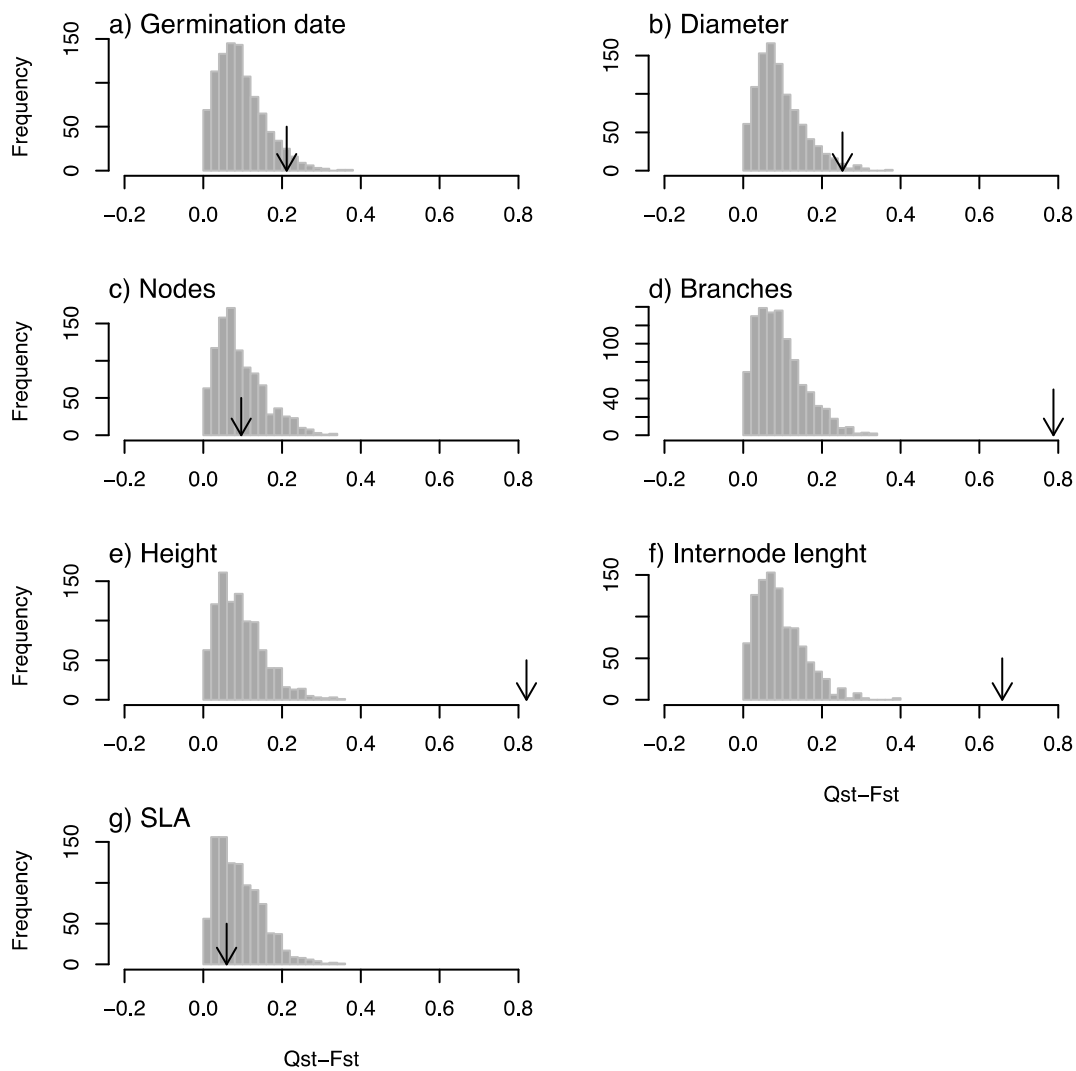
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717

718 FIGURE S4. The simulated distribution of $Q_{ST} - F_{ST}$ for a neutral trait, and the observed point
 719 estimates of $Q_{ST} - F_{ST}$ differences in seven phenotypic traits measured for the eight *Antirrhinum*
 720 *majus pseudomajus* populations from the Southern France. The distribution of $Q_{ST} - F_{ST}$
 721 differences for a neutrally evolving trait was simulated following Whitlock and Guillaume
 722 (2009) based upon the observed population differentiation in neutral markers (F_{ST}) and the
 723 within-population variance in each trait. The arrow indicates the observed $Q_{ST} - F_{ST}$.

724



725

726 FIGURE S5. The simulated distribution of $Q_{ST} - F_{ST}$ for a neutral trait, and the observed point
 727 estimates of $Q_{ST} - F_{ST}$ differences in seven phenotypic traits measured for the five *Antirrhinum*
 728 *majus striatum* populations from the Southern France. The distribution of $Q_{ST} - F_{ST}$ differences
 729 for a neutrally evolving trait was simulated following Whitlock and Guillaume (2009) based
 730 upon the observed population differentiation in neutral markers (F_{ST}) and the within-population
 731 variance in each trait. The arrow indicates the observed $Q_{ST} - F_{ST}$.

732