Major Transcriptome Reprogramming Underlies Floral Mimicry Induced by the Rust Fungus *Puccinia monoica* in *Boechera stricta*

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Abstract

*Puccinia monoica* is a spectacular plant parasitic rust fungus that triggers the formation of flower-like structures (pseudoflowers) in its Brassicaceae host plant *Boechera stricta*. Pseudoflowers mimic in shape, color, nectar and scent co-occurring and unrelated flowers such as buttercups. They act to attract insects thereby aiding spore dispersal and sexual reproduction of the rust fungus. Although much ecological research has been performed on *P. monoica*-induced pseudoflowers, this system has yet to be investigated at the molecular or genomic level. To date, the molecular alterations underlying the development of pseudoflowers and the genes involved have not been described. To address this, we performed gene expression profiling to reveal 256 plant biological processes that are significantly altered in pseudoflowers. Among these biological processes, plant genes involved in cell fate specification, regulation of transcription, reproduction, floral organ development, anthocyanin (major floral pigments) and terpenoid biosynthesis (major floral volatile compounds) were down-regulated in pseudoflowers. In contrast, plant genes involved in shoot, cotyledon and leaf development, carbohydrate transport, wax biosynthesis, cutin transport and L-phenylalanine metabolism (pathway that results in phenylethanol and phenylacetaldehyde volatile production) were up-regulated. These findings point to an extensive reprogramming of host genes by the rust pathogen to induce floral mimicry. We also highlight 31 differentially regulated plant genes that are enriched in the biological processes mentioned above, and are potentially involved in the formation of pseudoflowers. This work illustrates the complex perturbations induced by rust pathogens in their host plants, and provides a starting point for understanding the molecular mechanisms of pathogen-induced floral mimicry.

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Introduction

Many phytopathogens have evolved the ability to manipulate host plants to acquire nutrients and evade host defenses. These include microbes that possess the ability to cause dramatic morphological and physiological changes in their hosts. These changes can even lead to behavioral manipulation of a third organism, which is often an insect vector. Organisms with this "long-reach" phenotype include several species of obligate plant pathogenic bacteria and fungi. One example is the Aster Yellows phytoplasma strain Witches’ Broom (AY-WB), which infects a broad range of plant hosts [1,2] and induces a variety of morphological changes. These include the conversion of floral organs into leaves (phyllody), clustering of stems and branches (witches’ broom), green pigmentation of non-green flower tissues (virescence) and growth of elongated stalks (bolting) [1]. These morphological changes are thought to entice egg-laying insects to visit infected plants where they feed on contaminated tissue and transmit bacteria to new host plants [3,4]. Another example is the rust fungus *Puccinia monoica* that manipulates its plant host *Boechera stricta* (syn. *Arabis drummondii*) to create
elaborate pseudoflowers. These structures are completely novel to the plant’s native architecture [5] and act to lure pollinators from co-blooming plant species by offering olfactory incentives and a sugary reward [5]. Pollinator visits are essential for the completion of the sexual reproductive cycle of the fungus as they transfer spores of opposite mating types between pseudoflowers [5].

Despite recent advances in our understanding of the molecular mechanisms underlying pathogen-derived host manipulation, little is known regarding the transcriptional changes that occur in plants upon infection by pathogens that cause developmental reprogramming. To address this, we examined the effect of *P. monoica* infection on its host plant *B. stricta*, a close relative of *Arabidopsis*. *B. stricta* belongs to the Brassicaceae and grows mainly in the alpine regions of western North America [6]. In late summer, wind-borne basidiospores of *P. monoica*, produced on an unknown primary host grass, systematically infect the apical meristem of *B. stricta*, its secondary host [5,7]. *P. monoica* infection inhibits flowering and radically transforms *B. stricta* morphology, manipulating it to produce yellow flower-like structures that mimic true flowers of the unrelated co-blooming buttercups, *Ranunculus inamoenus* [8,9]. Although these pseudoflowers are visually similar in size, shape, color, and nectar production to true buttercup flowers, they produce a distinct sweet fragrance that attracts insect visitors [8,10,11]. Due to the heterothallic nature of *P. monoica*, these spore-laden insect visitors are critical to completion of the pathogens life cycle [5,12].

*B. stricta* is a close relative of the model plant *Arabidopsis thaliana*, therefore we can utilize the extensive genomic resources available for *A. thaliana* to study *P. monoica*-B. stricta interactions [13,14]. To this aim, we employed *A. thaliana* whole-genome microarrays to analyze transcriptional changes in *B. stricta* gene expression upon *P. monoica* infection. We used a NimbleGen microarray to determine the expression levels of transcripts isolated from *P. monoica*-induced pseudoflowers (‘PF’), uninfected *B. stricta* flowers (‘F’), and uninfected *B. stricta* stems and leaves (‘SL’). To compare relative gene expression levels we used Rank Products (RP) protocols [15]. This analysis identified 1036 and 910 genes that showed significant changes in expression in comparisons between ‘PF’ vs. ‘SL’ and ‘F’ vs. ‘SL’ respectively. Next, we performed an enrichment analysis of Gene Ontology terms describing Biological Processes (GOBP) using BiNGO in Cytoscape on these gene sets [16]. We found a total of 256 and 199 GOBP terms significantly enriched in ‘PF’ vs. ‘SL’ and ‘F’ vs. ‘SL’ comparisons, respectively. Among 256 key biological processes, we identified 31 gene candidates (20 up-regulated, and 11 down-regulated) showing significant alterations in expression between pseudoflowers ‘PF’ and uninfected *B. stricta* stems and leaves ‘SL’. These included genes involved in (i) leaf, stem and flower development, (ii) organ symmetry, (iii) metabolism of sugars, (iv) transport of sugars and lipids, and (v) wax and volatiles synthesis.

Our findings point to major reprogramming of the *B. stricta* transcriptome during infection, with several key biological processes acting as targets that could account for *P. monoica-*induced pseudoflower formation. This study is a crucial step towards understanding how this rust fungus manipulates its host plant at the molecular level and how such “long-reach” pathogens act to indirectly manipulate insect vectors to achieve sexual reproduction.

**Results and Discussion**

**Gene expression profiling of pseudoflowers**

To identify changes in *Boechera stricta* gene expression in *Puccinia monoica*-induced pseudoflowers we hybridized cDNA to a customized NimbleGen expression array covering all predicted coding genes in the *Arabidopsis thaliana* genome, a close relative of *B. stricta*. Samples were collected as follows: (i) uninfected plant stems and leaves (‘SL’), (ii) uninfected plant flowers (‘F’) (Figure 1A) and (iii) pseudoflowers from *P. monoica*-infected plants (‘PF’) (Figure 1B). Using a Rank Products (RP) analysis, we identified 1036 genes (Table S1) and 910 genes (Table S2) showing significant differential expression in the ‘PF’ vs. ‘SL’ and ‘F’ vs. ‘SL’ comparisons, respectively (Figure 2A-B). Among these, a total of 790 genes showed differential expression in the ‘PF’ vs. ‘SL’ comparison alone, and 664 showed differential expression only in the ‘F’ vs. ‘SL’ comparison (Figure 2C).

RP analysis generated two RP FDR values for each gene indicating the probability of being up or down-regulated [15]. Of the 1036 genes present in the ‘PF’ vs. ‘SL’ comparison, we determined 513 to be up-regulated and 523 to be down-regulated (with RP FDR values < 0.05) (Figure 2A and Table S1). In the ‘F’ vs. ‘SL’ comparison, out of the pool of 910 differentially expressed genes, 458 genes were up-regulated and 452 down-regulated (Figure 2B and Table S2).

**Validation of a subset of genes differentially regulated in pseudoflowers by qRT-PCR**

To test for the robustness of the gene regulation patterns we observed by microarrays analysis, we validated by qRT-PCR the expression of a subset of seven genes that are differentially regulated in *Puccinia monoica*-induced pseudoflowers (‘PF’) compared to uninfected *Boechera stricta* stems and leaves (‘SL’) (Figure 3). We selected *TEOSINTE BRANCHED1, CYCOIDEA, and PFC TRANSCRIPTION FACTOR3 TCP3* (At1g53230), *ALTERED MERistem PROGRAMMING1 AMP1* (At3g54720), *KNOTTED-LIKE1 KNAT1* (At4g08150) and *FLOWERING LOCUS T FT* (At1g65480) genes as representatives of host cell developmental processes altered in pseudoflowers (‘PF’), such leaf morphogenesis, pattern and cell specification, respectively (Table 1, see discussion below). Also, we selected *SUGAR TRANSPORTER1 SWEET1, At1g21460* and *SUGAR TRANSPORTER15 SWEET15, At5g13170* genes involved in carbohydrate transport and *TYROSINE TRANSAminase* enzyme-encoding gene (TT, At4g23550) that participates in organic volatile compounds synthetic pathway (Table 1). These seven genes were also selected as examples of the different modes of regulation in gene expression observed in ‘PF’. TCP3, SWEET1, SWEET15 and TT genes are up-regulated in pseudoflowers whereas AMP1, KNAT1 and FT are down-regulated and these
observations support the expression patterns found by microarray analysis (Figure 3). Even though small differences were found in the strength of the changes of gene expression in pseudoflowers compared to uninfected B. stricta stems and leaves, the overall expression patterns from qRT-PCR were very similar and matches the data obtained from the microarray analysis (Figure 3).

**Biological processes altered in pseudoflowers**

To identify and annotate biological processes altered during the formation of pseudoflowers, we performed Gene Ontology enrichment analysis for terms describing Biological Processes (GOBP). We identified 256 (Table S3) and 199 (Table S4) GOBP terms significantly enriched in the ‘Pf’ vs. ‘SL’ and ‘F’ vs. ‘SL’ comparisons, respectively. These GOBPs for both
comparisons are shown in Figure 4A in a network map as circular nodes that are color-coded according to the average expression (red for up-regulated and green for down-regulated).

*P. monoica*-infected plants develop an elongated stem with modified leaves instead of flowers [5]. In accordance, we identified genes involved in maintenance and development of floral organs amongst those down-regulated in *P. monoica*-induced pseudoflowers. Functional groups overrepresented among the genes down-regulated specifically in the ‘Pf’ vs. ‘SL’ comparison, but not in the ‘F’ vs. ‘SL’ comparison, included: (1) reproduction (GO:0000003), (2) floral organ development (GO:0048437), (3) carpel development (GO:0048440), (4) stamen development (GO:0048443), (5) cell fate specification (GO:0001708), (6) maintenance of floral meristem identity (GO:0010076), (7) anthocyanin biosynthesis (GO:00009718), (8) water transport (GO:0006833), (9) pattern specification (GO:0007389), (10) xylem and phloem pattern formation (GO:0010051), (11) regulation of transcription (GO:0045449), and (12) monoterpenoid biosynthesis (GO:0016099) (Figure 4B). Among these processes we highlight the down-regulation of monoterpenoid biosynthetic genes (Figure 4B), which is consistent with the observation that *P. monoica* induces the synthesis of chemical attractants unrelated to the native floral scent production of the host [5,10]. The distinct fragrance of pseudoflowers contains both phenylacetaldehyde and phenylethanol; compounds that are chemically different to the terpenoids produced in uninfected flowers, but possess the same ability to efficiently attract pollinators [5,10].

Functional groups over-represented and specifically up-regulated in the ‘Pf’ vs. ‘SL’ comparison included (1) shoot development (GO:0048367) (2), cotyledon development (GO:0048825) (3), leaf development (GO:0048366) and (4) leaf morphogenesis (GO:0009965) (Figure 4B). We also identified a few processes that are not overrepresented but include key candidate genes: (5) L-phenylalanine metabolism (GO:0006558), (6) carbohydrate transport (GO:0034219), (7) lipid transport (GO:0006869), (8) transmembrane transport (GO:0055085), (9) wax biosynthesis (GO:0006633) and (10) fatty acid biosynthesis (GO:0010025) (Figure 4B). Altogether our results suggest that pseudoflower development involves extensive reprogramming of shoot and leaf development, synthesis of volatiles, and changes to the host cell surface. These modifications are consistent with the phenotypes noted in...
in pseudoflowers, notably stem elongation, clustering of morphologically altered leaves that are covered by nectar-like substances and that emit a distinct scent [5].

We propose that the differentially regulated biological processes mentioned above constitute key processes involved in the remarkable developmental changes that take place in *P. monoica*-induced pseudoflowers. Our data confirms previous observations that suggest *P. monoica* manipulates the host to generate novel pseudofloral structures rather than exploiting existing floral machinery [5]. Among the key biological processes related to host cell development and metabolism that are altered in *P. monoica*-induced pseudoflowers (‘PF’) compared to uninfected stems and leaves (‘SL’), we selected 31 candidate genes (20 up-regulated, and 11 down-regulated) and classified them into nine groups for detailed discussion in the following sections (Table 1).

**De-differentiation of infected mesophyll cells**

Our findings indicate that in order to alter leaf development and produce pseudoflowers, *P. monoica* appears to induce the de-differentiation of host cells. We observed up-regulation of the *TEOSINTE BRANCHED1*, *CYCLOIDEA*, and *PCF TRANSCRIPTION FACTORS* (*TCP*) genes in pseudoflowers (‘PF’) compared to *B. stricta* stems and leaves (‘SL’), while *ALTERED MERISTEM PROGRAMMING1* (*AMP1*, At3g54720), *KNOTTED-LIKE1* (*KNAT1*, At4g08150) and *FLOWERING LOCUS T* (*FT*, At1g65480) genes was down-regulated (see Table 1). In addition, *SWEET15* and *FT* genes were confirmed to be down-regulated in uninfected *B. stricta* flowers (‘F’) compared to ‘SL’ as shown by microarray analysis (see Table S2). To indicate the mode of regulation we used two symbols: ‘*‘ for significant up-regulation and ‘#’ for significant down-regulation. The number of symbols indicates level of significance: one for *P < 0.05*, two for *P < 0.01* and three for *P < 0.001*. The error bars represents standard error of the mean.

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Figure 3. qRT-PCR validation of differentially expressed genes in pseudoflowers. Quantitative Real Time PCR (qRT-PCR) on a panel of seven genes was used to verify the transcriptional changes observed by microarray analysis. Consistent with the microarray results, expression of *TEOSINTE BRANCHED1*, *CYCLOIDEA*, and *PCF TRANSCRIPTION FACTORS* (*TCP3*, At1g53230), *SUGAR TRANSPORTER1* (*SWEET1*, At1g21460), *SUGAR TRANSPORTER15* (*SWEET15*, At5g13170) and *TYROSINE TRANSAMINASE* enzyme encoding gene (TT, At4g23590) genes was up-regulated in *Puccinia monoica*-induced pseudoflowers (‘PF’) compared to *B. stricta* stems and leaves (‘SL’), while *ALTERED MERISTEM PROGRAMMING1* (*AMP1*, At3g54720), *KNOTTED-LIKE1* (*KNAT1*, At4g08150) and *FLOWERING LOCUS T* (*FT*, At1g65480) genes was down-regulated (see Table 1). In addition, *SWEET15* and *FT* genes were confirmed to be down-regulated in uninfected *B. stricta* flowers (‘F’) compared to ‘SL’ as shown by microarray analysis (see Table S2). To indicate the mode of regulation we used two symbols: ‘*‘ for significant up-regulation and ‘#’ for significant down-regulation. The number of symbols indicates level of significance: one for *P < 0.05*, two for *P < 0.01* and three for *P < 0.001*. The error bars represents standard error of the mean.

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Table 1. *Arabidopsis thaliana* homologs of *Boechera stricta* genes with altered expression in pseudoflowers.

| Gene ID  | Gene name                                               | Common name                                                | Classification used in this study                                                                 | Expression in pseudoflowersa | Log$_2$ value$^b$ | GOBP$^c$ GOBP description$^c$                  |
|----------|---------------------------------------------------------|------------------------------------------------------------|--------------------------------------------------------------------------------------------------|------------------------------|-----------------|------------------------------------------------|
| At4g18390| TEOSINTE BRANCHED1, CYCLOIDEA, and PCF TRANSCRIPTION FACTOR2 | TCP2                                                       | De-differentiation of infected mesophyll cells                                                   | Up-regulated                 | 1.25            | 9.44E-03 Leaf morphogenesis                     |
| At1g53230| TEOSINTE BRANCHED1, CYCLOIDEA, and PCF TRANSCRIPTION FACTOR3 | TCP3                                                       | De-differentiation of infected mesophyll cells                                                   | Up-regulated                 | 1.17            | 1.38E-02 Leaf morphogenesis                     |
| At3g54720| ALTERED MERistem PROGRAMMING1                           | AMP1                                                       | De-differentiation of infected mesophyll cells                                                   | Down-regulated               | -1.07           | 4.21E-02 Pattern specification process          |
| A2g29125 | ROTUNDIFOLIA-LIKE2                                      | RTFL2                                                      | Alteration of the rate of cell proliferation                                                     | Up-regulated                 | 1.44            | 3.39E-03 Shoot development                      |
| At1g13710| ALTERED MERistem PROGRAMMING1                           | NGA3                                                       | Alteration of vascular patterning and phyllotaxy                                                | Up-regulated                 | 1.17            | 1.11E-02 Shoot development                      |
| At1g30490| PHAVOLUTA                                               | PHV                                                        | Alteration of vascular patterning and phyllotaxy                                                | Down-regulated               | -1.07           | 4.22E-02 Xylem and phloem pattern formation     |
| A2g15190 | FILAMENTOUS FLOWER                                      | FIL                                                        | Alteration of vascular patterning and phyllotaxy                                                | Up-regulated                 | 2.23            | 9.33E-05 Leaf development                       |
| A1g01030 | NGATHA3                                                | NGA3                                                       | Alteration of vascular patterning and phyllotaxy                                                | Up-regulated                 | 1.17            | 1.11E-02 Shoot development                      |
| A1g52150 | INCURVATA4                                              | ICU4                                                       | Alteration of vascular patterning and phyllotaxy                                                | Down-regulated               | -1.12           | 3.27E-02 Xylem and phloem pattern formation     |
| A2g09790 | QUARTER2                                               | QRT2                                                       | Inhibition of flower differentiation and maturation                                             | Up-regulated                 | 0.93            | 4.14E-02 Cellular developmental process         |
| A1g08150 | KNOTTED-LIKE1                                           | KNAT1                                                      | Inhibition of flower differentiation and maturation                                             | Down-regulated               | -1.06           | 4.54E-02 Cell fate specification               |
| A2g27990 | POUND-FOOLISH                                          | PNF                                                        | Inhibition of flower differentiation and maturation                                             | Down-regulated               | -1.18           | 2.93E-02 Maintenance of floral meristem identity |
| A1t65480 | FLOWERING LOCUS T                                       | FT                                                         | Inhibition of flower differentiation and maturation                                             | Down-regulated               | -1.28           | 2.65E-02 Reproduction                           |
| A2g03710 | SEPATALLA4                                              | SEP4                                                       | Inhibition of flower differentiation and maturation                                             | Down-regulated               | -1.40           | 9.58E-03 Floral organ development              |
| A4g37390 | INDOLE-3-ACETIC-AMIDO SYNTHASE2                         | GH3.2                                                      | Alteration of auxin homeostasis                                                                 | Up-regulated                 | 4.40            | 0.00E+00 Response to hormone stimulus           |
| A1t59500 | INDOLE-3-ACETIC-AMIDO SYNTHASE4                         | GH3.4                                                      | Alteration of auxin homeostasis                                                                 | Up-regulated                 | 2.64            | 2.86E-05 Response to hormone stimulus           |
| A1t70560 | TRYPTOPHAN AMINOTRANSFERASE OF ARABIDOPSIS1             | TAA1                                                       | Alteration of auxin homeostasis                                                                 | Up-regulated                 | 1.47            | 4.72E-03 Cotyledon development                  |
| A2g14370 | SERINE/THREONINE KINASE                                 | WAG2                                                       | Alteration of auxin homeostasis                                                                 | Up-regulated                 | 1.09            | 2.12E-02 Cotyledon development                  |
| A4g25960 | P-GLYCOPROTEIN2                                         | PGP2                                                       | Alteration of auxin homeostasis                                                                 | Up-regulated                 | 1.04            | 2.59E-02 Transmembrane transport               |
| A1t51480 | ATP-BINDING-CASSETTE (ABC) TRANSPORTER SUPERFAMILY G13  | ABCG13                                                     | Activation of wax biosynthesis and cutin transport                                               | Up-regulated                 | 2.80            | 0.00E+00 Lipid transport                        |
| A2g15090 | 3-KETOACYL-COA SYNTHASE8                                | KCS8                                                       | Activation of wax biosynthesis and cutin transport                                               | Up-regulated                 | 1.31            | 7.20E-03 Fatty acid biosynthesis               |
Table 1 (continued).

| Gene ID | Gene name                           | Common name | Classification used in this study                                                                 | Expression in pseudoflowers | Log2 value | RP FDR value | GOBP | GOBP description               |
|---------|------------------------------------|-------------|-------------------------------------------------------------------------------------------------|-----------------------------|------------|---------------|------|---------------------------------|
| At5g12420 | WAX ESTER SYNTHASE/ACYLCOA: DIACYLGLYCEROL ACETYLTRANSFERASE7 | WSD7        | Activation of wax biosynthesis and cutin transport                                             | Up-regulated                | 0.97       | 4.51E-02     | 10025 | Wax biosynthesis                |
| At5g23940 | CUTICULAR RIDGES                    | DCR         | Activation of wax biosynthesis and cutin transport                                             | Up-regulated                | 0.94       | 4.48E-02     | 6633  | Fatty acid biosynthesis         |
| At3g13790 | CELL WALL INVERTASE1               | cwiINV1      | Subversion of sugar metabolism                                                                | Up-regulated                | 2.44       | 4.29E-05     | 6950  | Response to stress              |
| At1g21460 | SUGAR TRANSPORTER1                 | SWEET1       | Subversion of sugar metabolism                                                                | Up-regulated                | 1.50       | 1.99E-03     | 34219 | Carbohydrate transmembrane      |
| At5g13170 | SUGAR TRANSPORTER15                | SWEET15      | Subversion of sugar metabolism                                                                | Up-regulated                | 1.38       | 5.09E-03     | 34219 | Carbohydrate transmembrane      |
| At1g68130 | INDETERMINANT DOMAIN14             | IDD14        | Subversion of sugar metabolism                                                                | Down-regulated              | -1.18      | 2.67E-02     | 45449 | Regulation of transcription     |
| At3g43190 | SUCROSE SYNTHASE4                  | SUS4         | Subversion of sugar metabolism                                                                | Down-regulated              | -2.32      | 4.39E-04     | 16051 | Carbohydrate biosynthesis       |
| At4g23590 | TYROSINE TRANSMAMASE               | TT           | Alteration of volatile organic compounds synthesis                                             | Up-regulated                | 2.50       | 1.82E-05     | 6558  | L-phenylalanine metabolism      |
| At2g24210 | TERPENE SYNTHASE10                 | TPS10        | Alteration of volatile organic compounds synthesis                                             | Down-regulated              | -2.22      | 7.44E-04     | 16099 | Monoterpenoid biosynthesis      |
| At5g23960 | TERPENE SYNTHASE21                 | TPS21        | Alteration of volatile organic compounds synthesis                                             | Down-regulated              | -2.65      | 1.90E-04     | 16099 | Monoterpenoid biosynthesis      |

a. Expression in Puccinia monoica-induced pseudoflowers (‘Pf’) relative to uninfected Boechera stricta stems and leaves (‘SL’).
b. Rank Product (RP) False Discovery Rate (FDR) values used to estimate differentially expressed genes in Puccinia monoica-induced pseudoflowers (‘Pf’) compared to uninfected Boechera stricta stems and leaves (‘SL’). Genes with RP FDR value < 0.05 are considered significant.

gene ontology terms describing biological processes (GOBP) from Arabidopsis thaliana TAIR database version 10.
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maintaining undifferentiated cells in the shoot apical meristem (SAM) and in coordinating differentiation in leaf organs [17,18]. In addition, we noted the down-regulation of ALTERED MERistem PROGRAMMING1 (AMP1, At3g54720) that acts to promote cell differentiation [19,20] (Table 1 and Figure 4B). Mutations in AMP1 in A. thaliana leads to increased leaf initiation, reduced leaf and stem size, and apical dominance [19]. Therefore, up-regulation of TCPs and down-regulation of AMP1, could promote de-differentiation of infected mesophyll cells. De-regulation of these genes may also prevent branching of leaves growing in the upper part of the stem (cauline leaves) by suppression of lateral shoot development, thereby maintaining apical dominance in stems bearing pseudoflowers (Figure 1B).

Alteration of the rate of cell proliferation
Pseudoflowers consist of modified leaves differing in size and shape relative to uninfected B. stricta leaves (Figure 1). Cell proliferation and cell expansion processes of leaf morphogenesis are required to produce the final leaf shape [21], and these processes might be altered by P. monoica to achieve pseudoflower morphogenesis. In accordance, we found up-regulation of the ROTUNDIFOLIA-LIKE2 (RTFL2, At1g29125) gene in pseudoflowers (Table 1 and Figure 4B). Overexpression of ROTUNDIFOLIA4 (ROT4), a homolog of RTFL2 decreases cell numbers specifically in the leaf-length direction resulting in shortened and rounded leaves in A. thaliana [22,23] suggesting that this gene may regulate cell proliferation during pseudoflowers morphogenesis.

Alteration of coordinated organ growth and symmetry
Many plants have an outcrossing (allogamy) reproduction strategy and their reproductive success depends on pollinators [24]. The size and architecture of floral organs determines flower shape and attractiveness towards insect visitors [24]. Analogous to allogamous plants, the rust fungus P. monoica also relies on pollinators [5,8]. We observed symmetrically arranged flower-like leaves at the top of the pseudoflowers (Figure 1B), suggesting that P. monoica alters the growth of individual leaves to produce the characteristic pseudofloral structures. Consistent with this hypothesis, the cytochrome P450 monooxygenase KLUH/CYP78A5 (At1g13710) gene was up-regulated in pseudoflowers (Table 1 and Figure 4B). KLUH/ CYP78A5 promotes leaf and floral organ growth [25],
Figure 4. Overview of biological processes altered in pseudoflowers. (A) Simplified gene ontology biological processes (GOBP) network showing processes enriched among genes with expression altered in Puccinia monoica-induced pseudoflowers ('Pf') compared to uninfected Boechera stricta stems and leaves ('SL') and B. stricta flowers ('F') compared to 'SL'. Node size with average GOBP fold induction (average of log_2 ratios of all genes within a GOBP in 'Pf'/SL' and 'F'/SL', respectively) from green for average induction folds < 0 that indicates down-regulation to red for average induction folds > 0 that indicates up-regulation (node color). Some nodes and edges have been omitted for clarity. (B) Detailed GOBP network showing processes enriched among genes with expression altered in 'Pf' vs. 'SL'. Genes highlighted in the text are indicated with diamonds connected to dashed lines to the processes they are involved in. This network shows same topology as in (A).

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particularly by coordinating the growth of individual flower organs, and contributing to uniformity of flower size and symmetry [26]. Our results suggest CYP78A5 may coordinate the symmetry of pseudoflower leaf clusters (Figure 1B), an important visual cue known to attract pollinators [27,28].

**Alteration of vascular patterning and phyllotaxy**

The dramatic morphological alterations in pseudoflowers (Figure 1B) imply the establishment of specific vascular bundle and leaf patterns. Several transcription factors controlling these processes were differentially expressed in pseudoflowers: the INCURVATA4 (ICU4, At1g52150) and PHAVOLUTA (PHV, At1g30490) genes were down-regulated, whereas the FILAMENTOUS FLOWER (FIL, At2g45190) and NGATHA3 (NGA3, At1g01030) genes were up-regulated (Table 1 and Figure 4B). ICU4 encodes ATHB-15, an HD-ZIP III transcriptional factor required for shoot apical meristem patterning and stem vascular differentiation [29]. A. thaliana icu4 mutants display an abnormal arrangement of leaves due to impaired shoot apical meristem development, producing paired leaves along the stem and axillary shoots [29]. PHV encodes another HD-ZIP III transcriptional factor involved in specification of the upper (adaxial) surface of leaves [30]. FIL is a member of the YABBY family of transcriptional regulators required for vascular differentiation, specifically in the abaxialization of leaves [31]. Transgenic A. thaliana plants expressing FIL form filamentous leaves with mainly abaxial looking tissues [31]. Over-expression of NGA3 in transgenic plants results in apical dominance and alters flower phyllotaxy. It also causes abnormal arrangement of leaves in the stem axis, longer, darker, and narrower rosette leaves, as well as a flattened stem [32]. Overall, the down-regulation of master transcriptional regulators of leaf development, ICU4, PHV and up-regulation of FIL and NGA3 could contribute to the altered vascular patterning and morphology of leaves in pseudoflowers [5] (Figure 1B).

**Inhibition of flower differentiation and maturation**

The formation of pseudoflowers is likely to involve the inhibition of floral signals and floral organ development in the host. Accordingly, five genes involved in the flowering transition were differentially expressed in pseudoflowers: FLOWERING LOCUS T (FT, At1g65480), KNOTTED-LIKE1 (KNOT1, At4g08150), POUND-FOOLISH (PNF, At2g27990) and SEPATALLA4 (SEP4, At2g03710) genes were down-regulated, whereas the QUARTER2 (QRT2, At3g07970) gene was up-regulated (Table 1 and Figure 4B). FT produces a mobile floral activator signal protein that moves through the phloem from induced leaves to the shoot apex where it interacts with the FLOWERING LOCUS D (FD) bZIP transcription factor to initiate transcription of floral specification genes [33,34]. Down-regulation of FT in pseudoflowers suggests interference with the activation and transmission of floral cues, probably leading to inhibition of floral organ development in infected plants (Figure 1B). Loss of KNOT1, a member of the class1 Kotted1-like homeobox (KNOX) family of transcriptional regulators, results in reduced growth of floral pedicles, internodes and the style during reproductive growth [35,36]. PNF and its paralog PENNYWISE (PNY, At5g02030) encode for BEL1-like homeobox (BLH) proteins that regulate inflorescence internode patterning [37,38] and are also required for floral formation mediated by FT [39]. Arabidopsis pny pnf double mutants initiate compact shoots that fail to respond to flowering signals and subsequently never form flowers [37]. Together with other MADS-box transcription factors, SEP4 plays a central role in floral meristem and floral organ identity [40]. A. thaliana sep4 single mutants do not exhibit visible phenotypes, but when all four members of the SEP gene family (sep1 sep2 sep3 sep4) are mutated, plants show a conversion of floral organs to leaf-like organs [40]. As opposed to these four transcription factors, QRT2, which encodes a polygalacturonase involved in cell division, was up-regulated in pseudoflowers (Table 1 and Figure 4B). Plants over-expressing QRT2 have flowers that do not open, atypical petals, and anthers that fail to dehisce normally [41]. Together these results suggest that the regulation of several genes involved in floral organ differentiation and maturation could potentially act in conjunction to inhibit flower formation in infected plants (Figure 1B). We hypothesize that inhibition of flowering could prolong the lifespan of P. monoica infected plants, benefiting the parasite at the expense of host plant fitness and reproductive success [5].

**Alteration of auxin homeostasis**

Pseudoflowers consist of clusters of elongated stems that bolt from infected rosettes and almost never reach flowering. Changes in the regulation of plant host hormones involved in organogenesis may contribute to the formation of these dense flower-like clusters. Accordingly, we found that genes involved in various mechanisms that control auxin homeostasis were up-regulated in pseudoflowers. Among these are the TRYPTOPHAN AMINOTRANSFERASE OF ARABIDOPSIS1 (TAA1, At1g70560) gene that is essential for Trp-dependent indole-3-acetic acid (IAA) biosynthesis [42] and the IAA-AMIDO SYNTHASE2 (GH3.4, At4g37390) and 4 (GH3.4, At1g59500), which are involved in the production of IAA conjugates to regulate the level of active auxin inside the plant [43] (Table 1 and Figure 4B). Also, we identified two genes that were up-regulated as involved in auxin-transport and auxin-mediated organogenesis processes: P-GLYCOPROTEIN2 (PGR2, At4g25980) that encodes a protein with homology to PGP1 known to mediate hypocotyl growth [44,45] and protein serine/threonine AGC KINASE WAG2 (At3g14370) that positively regulates cotyledon formation [46] (Table 1 and Figure 4B). Up-regulation of genes involved in auxin-mediated organogenesis could contribute to stem elongation and growth of leaves in pseudoflowers [5] (Figure 1B).

**Activation of wax biosynthesis and cutin transport**

Plants under water deficit show decreased stem elongation [47] and restricted formation and number of leaves due to increased leaf senescence [48]. To help prevent water loss under stressful conditions some plants secrete and accumulate waxes in the surface of the leaves [49]. We found in pseudoflowers the up-regulation of key genes involved in wax and cutin biosynthesis and transport in A. thaliana leaves and...
substances: WDS7 (At5g12420), a homolog of the Wax Ester Synthase/AcylCoA:diacylglycerol acyltransferase1 (WSD1), 3-ketoacyl-CoA synthase8 (KCS8, At2g15090), and DEFECTIVE IN CUTICULAR RIDGES/PERMEABLE LEAVES3 (DCR/PEL3, At5g23940) (Table 1 and Figure 4B). In A. thaliana, WSD7 homolog, WSD1, is a wax synthase required for stem wax ester biosynthesis [50], KCS8 is a component of the fatty acid elongase complex required for the synthesis of epicuticular waxes in leaves [51], and DCR encodes a putative acyltransferase required for the incorporation of the monomer 9(10),16-dihydroxy-hexadecanoic acid into the cutin polymeric ketoacyl-CoA synthase8 Arabidopsis thaliana (SUCROSE SYNTHASE4 (SUS4, At5g20830) are down-regulated in pseudoflowers (Table 1 and Figure 4B). IDD14 shares homology with IDD8, an A. thaliana protein with a zing finger ID-domain (IDD) that indirectly promotes flowering via transcriptional activation of SUS4 [61]. Activation of SUS4 regulates photoperiodic flowering through the modulation of sugar metabolism and transport [62]. IDD14 and SUS4 down-regulation therefore suggesting transcriptional modulation of host sugar metabolism leading to the repression of flowering, that may prolong the infected plant vegetative phase and favour P. monoica.

Alteration of volatile organic compounds synthesis

The attraction of insect pollinators by pseudoflowers involves volatile compounds that significantly differ from the host flower fragrances [10, 11]. The fragrance of Arabidopsis spp. flowers is predominantly composed of monoterpens and sesquiterpenes volatile organic compounds (VOCs) [63]. We identified two genes involved in terpene biosynthesis in Arabidopsis, TERPENE SYNTHASE10 (TPS10, At2g24210) and 21 (TPS21, At5g23960), that were down-regulated in pseudoflowers (Table 1, Figure 4B and Figure 5A-B). TPS10 encodes a β-myrcene/(-)-β-caryophyllene synthase expressed almost exclusively in flowers [63,65]. Down-regulation of these two TPS genes suggests the absence of terpene blends in pseudoflowers as shown by Raguso and Roy [10].

In contrast one gene involved in phenylalanine degradation was up-regulated: Tyrosine transaminase (TT, At4g23590) (Table 1, Figure 4B and Figure 5C). This pathway ultimately produces the VOCs phenylacetaldehyde and phenylethylethanol. This finding is consistent with a previous study that identified phenylacetaldehyde and phenylethylethanol as the most abundant volatiles in Puccinia-induced pseudoflowers [10]. Phenylacetaldehyde is a well-known attractant for foraging insects [66] and has been proposed to attract pollinators in certain Arabidopsis ecotypes [67]. Our results are therefore consistent with the hypothesis by Raguso and Roy [10] that pseudoflowers produce fragrances by modifying host-plant metabolites.

Conclusions

Using whole-genome expression profiling, we identified a large number of plant genes that with altered expression in P. monoica-induced pseudoflowers (PF) compared to uninfected B. stricta stems and leaves (‘SL’). We report nine major host processes related to cell development (de-differentiation of mesophyll cells, rate of cell proliferation, coordinated organ growth and symmetry, of vascular pattering and inhibition of floral organs) and to host metabolism (auxin homeostasis, host cell wax surface compound synthesis, sugar metabolism and volatile organic compounds synthesis) that appear to be re-programmed at the transcriptional level in pseudoflowers. Alterations in the expression of regulators of cell fate and auxin metabolism (auxin homeostasis, host cell wax surface compound synthesis, sugar metabolism and volatile organic compounds synthesis) that appear to be re-programmed at the transcriptional level in pseudoflowers. Alterations in the expression of regulators of cell fate and auxin homeostasis genes may account for the distinctive morphology of pseudoflowers, which are essentially modified leaves [5]. Up-regulation of genes controlling stem and leaf development and individual organ symmetry may contribute to the characteristic phenotype in pseudoflowers (infected plants having elongated stems bearing modified short leaves with symmetrically arranged flower-like leaves at the top) (Figure

Subversion of sugar metabolism

We noted that the expression of several B. stricta genes involved in sugar metabolism is altered in pseudoflowers. Two SWEET genes encoding sugar transporters, SWEET1 (At1g21460) and SWEET15 (At5g13170) [55], are up-regulated (Table 1 and Figure 4B), suggesting that sugar transporters might be co-opted during infection by P. monoica for nutritional gain. CELL wall invertase1 (cwINV1, At3g13790) (Table 1 and Figure 4B), which in A. thaliana is induced upon fungal infection [56], is also up-regulated. Cell wall invertases control plant metabolism by hydrolyzing sucrose and providing apoplastic glucose and fructose to the cells [57]. Among the A. thaliana cell wall invertase family, cwINV1 and its paralog cwINV4, which are required for sugar accumulation during nectar production [58], are highly expressed in flowers [59]. We hypothesize that up-regulation of cwINV1 could contribute to the production of the nectar-like substance over the surface of pseudoflowers (Figure 1B). Nectar is the principal floral reward for pollinators [60], and its production in pseudoflowers should benefit the rust pathogen [5]. Unlike the flower-expressed cwINV4, cwINV1 is highly expressed in leaves [59], perhaps facilitating manipulation by the rust fungus.

Conversely, INDETERMINATE14 (IDD14, At1g68130) and SUCROSE SYNTHASE4 (SUS4, At5g20830) are down-regulated in pseudoflowers (Table 1 and Figure 4B). IDD14 shares homology with IDD8, an A. thaliana protein with a zing finger ID-domain (IDD) that indirectly promotes flowering via transcriptional activation of SUS4 [61]. Activation of SUS4 would promote flowering independently of the photoperiodic pathway and gene expression in pseudoflowers is consistent with this hypothesis.
1B). Down-regulation of genes involved in floral organ development and floral transition may prevent the switch from the vegetative phase to flowering. Genes associated with cuticular wax production that are up-regulated in pseudoflowers could help to protect and maintain leaf longevity in stressful conditions and this could benefit *P. monoica* nutrient supply. Up-regulation of genes involved in sugar metabolism and transport could provide carbon sources for *P. monoica*, and contribute to the synthesis of nectary substances attractive to pollinators. Finally, down-regulation of terpene VOCs synthesis genes and up-regulation of phenylacetaldehyde synthesis may contribute to the distinct pseudoflower fragrance resulting in odorant cues to attract insects.

Our expression profiling experiments of *P. monoica*-infected pseudoflowers has provided a number of new insights into how obligate fungal pathogens manipulate both their plant and insect hosts. Our findings contribute significant insight into the dramatic morphological and physiological changes that occur in *B. stricta* plants infected by this rust fungus.

### Materials and Methods

#### Plant material and RNA extraction

In this study we collected naturally *Puccinia monoica*-infected *Boechera stricta* plants for gene expression analysis. Artificially infection of *B. stricta* plants with *P. monoica* in the laboratory is not feasible as its alternate host grass, on which *P. monoica* must grow to complete its life cycle, is unknown. We extracted total RNA from *P. monoica*-induced pseudoflowers (‘PF’) (3 samples), uninfected *B. stricta* plant stems and leaves (‘SL’) (3 samples), and uninfected *B. stricta* flowers (‘F’) (2 samples). Tissue was collected in Gothic (2900 m) (Colorado, USA), and stored in RNAlater® tissue collection solution (Invitrogen, Cat No. AM7020). Total RNA was extracted using TRizol® Reagent (Invitrogen, Cat No. 15596-026) according to the manufacturer’s instructions. RNA quality and integrity were assessed prior to cDNA synthesis using the Bioanalyzer (Agilent 2100). NimbleGen microarray services were used for cDNA preparations, chip hybridizations to an *Arabidopsis thaliana* NimbleGen array design 4 x 72K (4-plex format with 72000 probes, two probes per target gene and a total of 30362 target genes, Cat No. A4511001-00-01) and
subsequent normalization of the probe sets using Robust Multichip Average (RMA) [68].

**Ethics statement**

Samples were collected from Gothic in a location that is about 5 miles away from the Rocky Mountain Biological Laboratory (RMBL) near Gunnison (Colorado, USA). The location used in this study is not private owned or protected and has been previously reported for sampling of *Boechera stricta* [69]. *B. stricta* is not a protected plant species in Colorado, USA. No specific permissions (for both the location and plant material) were required for the collection of the samples.

**Gene expression analysis**

For the analysis of microarray data, we estimated a False Discovery Rate (FDR) for differential gene expression using the Rank Products (RP) program [15]. This program performs permutations with no data distribution assumptions and is recommended for samples obtained outside of controlled laboratory conditions [70]. We applied the RP analysis with 5000 permutations on two comparisons of samples: (i) *Puccinia monoica*-induced pseudoflowers 'PF' vs. uninfected *Boechera stricta* plant stems and leaves ('SL') and (ii) uninfected *B. stricta* flowers 'F' vs. 'SL'. A threshold of TP FDR value < 0.05 was used to identify differentially regulated genes (up-regulated and down-regulated) in each comparison.

**Accession numbers**

The NimbleGen microarray data used in this publication have been deposited on the Gene Expression Omnibus (GEO) (GEO, http://www.ncbi.nlm.nih.gov/geo/) and are accessible through GEO series accession number GSE41165.

**Quantitative real time PCR**

Seven candidate *Arabidopsis thaliana* genes that exhibited changes in expression in *Puccinia monoica*-induced pseudoflowers ('PF') compared to uninfected *Boechera stricta* stems and leaves ('SL') in the microarray experiment were selected for validation via quantitative real time PCR (qRT-PCR). Primers were designed wherever possible to anneal at 60°C and to either sit on or amplify across an exon boundary to avoid contamination from genomic DNA using amplify v3.1 software (© Bill Engels, University of Wisconsin) (see Table S5). All amplicons were initially confirmed by agarose electrophoresis to determine if the amplicon was the predicted size and a single product. The qRT-PCR analyses were carried out in a CFX96™ real-time PCR system (Bio-Rad) instrument using the SYBR Green JumpStart Taq ready mix (Sigma, Cat No. S4438) on cDNA samples produced with the SuperScript III first-strand synthesis system kit (Invitrogen, Cat No. 18080-051) following the manufacturer's instructions. The qRT-PCR experiments were performed with cDNAs of three tissues: pseudoflowers ('PF'), *B. stricta* flowers ('F') and *B. stricta* stems and leaves ('SL'). To quantify the relative expression level of a particular gene, three independent qRT-PCR reactions (technical replicates) were performed on biological duplicate samples for each tissue analyzed. All data points were normalized to the internal control gene *ELONGATION FACTOR1 ALPHA (EF-1alpha, At1g18070)* and relative to samples collected *B. stricta* stems and leaves ('SL') using the comparative CT (ΔΔCT) method. ΔCT is defined as the difference in the cycle threshold (CT) between the gene of interest and the *EF-1alpha* control. Relative expression levels were calculated as the log₂ ratio = log₂ [(ΔΔCT of 'PF' at gene X)/(ΔCT of 'SL' at gene X)], with 'X' corresponding to the gene of interest. The statistical T-test was performed using R software to determine differences between 'PF' vs. 'SL' and 'F' vs. 'SL' group. A P-value < 0.05 was defined as statistically significant. To indicate the mode of regulation we used two symbols: '*' for significant up-regulation and '*' for significant down-regulation. The number of symbols indicates level of significance: one for P < 0.05, two for P < 0.01 and three for P < 0.001. Data is presented as average ± SEM.

**Gene Ontology (GO) enrichment and pathway analysis**

A list of GO annotations for *Arabidopsis thaliana* was extracted from The Arabidopsis Information Resource (TAIR) database [71]. Using the BINGO plug-in available for Cytoscape [16], over-represented groups of GO terms and functional domains were identified using a hypergeometric test with Benjamin & Hochberg False Discovery Rate (FDR) correction and a P-value threshold of 0.05. This test identified significantly enriched GO categories by comparing *Arabidopsis thaliana* 27822 GO annotated genes with the 1036 and 910 genes that showed significant changes in gene expression in (i) *Puccinia monoica*-induced pseudoflowers ‘PF’ vs. uninfected *Boechera stricta* stems and leaves ‘SL’ and (ii) uninfected *B. stricta* flowers ‘F’ vs. ‘SL’ comparisons, respectively. Using Cytoscape visualization tools we constructed a network map to illustrate significantly enriched Gene Ontology terms describing Biological Processes (GOBP) in (i) ‘PF’ vs. ‘SL’ and (ii) ‘F’ vs. ‘SL’ comparisons, respectively. The size of the node in the network map corresponds to -log₁₀ of the corrected P-value of enrichment within a GOBP term. In addition, snapshots of *A. thaliana* AraCyc and PlantCyc browsers (http://plantcyc.org/) were used to visualize specific metabolic pathways that were significantly regulated in ‘PF’ vs. ‘SL’.

**Supporting Information**

Table S1. List of 1036 significantly differentially expressed genes in pseudoflowers. ‘PF’: *Puccinia monoica*-induced pseudoflowers. ‘SL’: *Boechera stricta* uninfected stems and leaves. *False Discovery Rate (FDR) estimated using Rank Products (RP) to detect genes that are differentially expressed. Genes with RP FDR value < 0.05 are considered significant. (XLSX)

Table S2. List of 910 significantly differentially expressed genes in uninfected *Boechera stricta* flowers. ‘F’: *Boechera stricta* uninfected flowers. ‘SL’: *Boechera stricta* uninfected stems and leaves. *False Discovery Rate (FDR) estimated using Rank Products (RP) to detect genes that are differentially expressed. Genes with RP FDR value < 0.05 are considered significant. (XLSX)
Table S3. List of 256 gene ontology biological processes (GOBP) enriched in pseudoflowers. 'PF': *Puccinia monoica*-induced pseudoflowers. 'SL': *Boechera stricta* uninfected stems and leaves. Number of genes within found within the biological process. P-value showing the significance for enrichment of genes within the biological process.

Table S4. List of 199 gene ontology biological processes (GOBP) enriched in uninfected *Boechera stricta* flowers. 'F': *Boechera stricta* uninfected flowers. 'SL': *Boechera stricta* uninfected stems and leaves. Number of genes within found within the biological process. P-value showing the significance for enrichment of genes within the biological process.

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