

2 Development of first linkage map for *Silphium integrifolium* (Asteraceae) enables identification
of sporophytic self-incompatibility locus

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16 SUMMARY

- 18 • *Silphium integrifolium* (*Asteraceae*) has been identified as a candidate for domestication
20 as a perennial oilseed crop and has a sporophytic self-incompatibility system—the
22 genetic basis of which is not well understood in the *Asteraceae*. To address this gap, we
24 sought to map the genomic location of the self-recognition locus (S-locus) in this species.
- 26 • We used a biparental population and genotyping-by-sequencing to create the first genetic
28 linkage map for this species. Then we developed a novel crossing scheme and set of
30 analysis methods in order to infer S-locus genotypes for a subset of these individuals,
32 allowing us to map the trait. Finally, we identified potential gene candidates using
synteny analysis with the annual sunflower (*Helianthus annuus*) genome.
- Our linkage map contains 198 SNP markers and resolved into the correct number of
linkage groups. We were able to successfully map the S-locus and identify several
potential gene candidates in the sunflower syntenic region.
- Our method is effective and efficient, allowed us to map the *S. integrifolium* S-locus
using fewer resources than previous studies, and could be readily be applied to other
species. Our best gene candidate appears to be worthy of future work in *S. integrifolium*
and other *Asteraceae* species.

34 Key words: *Asteraceae*, domestication, genetic resources, S-locus, self-incompatibility, *Silphium*
integrifolium, QTL mapping

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INTRODUCTION

38 *Silphium integrifolium* (Michx.) (wholeleaf rosinweed or silflower) is a member of the
40 *Asteraceae* family native to prairies throughout the central United States. In the early 2000s, *S.*
42 *integrifolium* was selected to be a candidate for domestication as a perennial oilseed crop by the
44 Land Institute in Salina, Kansas (Van Tassel *et al.*, 2017), attracting attention for its tolerance to
46 drought, upright growth habit, and large seeds (DeHaan *et al.*, 2016). Subsequently, *S.*
48 *integrifolium* has been found to have a seed oil composition similar to landrace sunflower
50 (Reinert *et al.*, 2019) and good winter survival and persistence in a range of climates (J.H. Price
& D.L. Van Tassel, pers. obs.). In addition, the yield potential of *S. integrifolium* populations
46 that have undergone relatively little selection is approximately 60% the yield of advanced
48 sunflower hybrids (Kandel *et al.*, 2019; Schiffner *et al.*, 2020), indicating that significant
50 improvement is likely with continued breeding efforts. These characteristics further encourage
52 the domestication of this species as a new crop. In the past, domestication occurred over long
54 periods of time, largely due to the largely unintentional nature of early selection (Rindos, 1984).
With the advantage of contemporary knowledge of genetics, genomics, and breeding techniques,
52 the amount of time necessary to domesticate a new crop could be drastically reduced. Therefore,
the development of genomic resources is a crucial step in this process (Sedbrook *et al.*, 2014). To
54 this end, we have developed the first genetic linkage map for *S. integrifolium*.

56 Among the traits for which better genetic knowledge will accelerate the domestication of *S.*
integrifolium is self-incompatibility. Although occasional *S. integrifolium* individuals have been
58 observed to produce at least some seed when self-pollinated (Reinert *et al.*, 2020), *S.*
integrifolium is self-incompatible, and as a member of the *Asteraceae* family is assumed to have
60 a sporophytic self-incompatibility (SSI) system (Hiscock, 2000). In sporophytic systems, self-
62 recognition is typically controlled by a single multi-allelic locus, known as the “S-locus”, with
rejection of self-pollen caused by stigma recognition of S-locus gene products found in or on the
64 pollen. Because these products are produced in the anther, pollen acceptance or rejection is
determined by the diploid genotype of the male parent, rather than the haploid genotype of a
66 given pollen grain (Hiscock & Tabah, 2003). SSI alleles are also expected to display complex
dominance patterns, and dominance relationships between alleles may differ from the anther to

the stigma (Hiscock & Tabah, 2003).

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The molecular mechanisms that underlie SSI are best described in the *Brassicaceae*, where the female S-phenotype is determined by a receptor kinase complex known as SRK. When pollen of the same S-phenotype lands on the stigma, this kinase binds a cysteine-rich protein (CRP) found in the pollen coat, known as SP₁₁/SCR, initiating the pollen rejection response (Fujii & Takayama, 2018). These two genes are tightly linked and rarely recombine; thus, they combine to form the S-locus, which is more properly thought of as an S-haplotype (Edh *et al.*, 2009). SRK is unable to bind SP₁₁/SCR proteins produced by different haplotypes, resulting in the acceptance of non-self pollen (Fujii & Takayama, 2018). Although the identity of the receptor protein varies, secreted CRPs are also considered a promising candidate for the male determinant of SSI in the *Convolvulaceae* (morning glory) family (Rahman *et al.*, 2007), and serve as the female determinant of gametophytic self-incompatibility (GSI) in the genus *Papaver* (poppy) (Marshall *et al.*, 2011),.

82 *Asteraceae* systems are less well understood. In *Senecio squalidis*, and subsequently in other *Asteraceae*, SRK-like sequences have been identified and cloned. However, results from *S. squalidis* and chicory (*Cichorium intybus*) indicate that they likely are not integral to S-genotype determination (Hiscock & Tabah, 2003 ; Gonthier *et al.*, 2013), and that the molecular control of *Asteraceae* SSI is quite different from the *Brassicaceae* system (Allen *et al.* 2011). Efforts to map the S-locus in chicory provided a 1.8 cM QTL region but have not yet determined a molecular basis for self-incompatibility (Gonthier *et al.*, 2013). Although efforts have been undertaken in other species, chicory represents perhaps the only example where a true *Asteraceae* S-locus has been definitively mapped, as opposed to other loci contributing to breakdowns in self-incompatibility (Gandhi *et al.*, 2005; Koseva *et al.*, 2017).

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Mapping the S-locus is important for breeding efforts and for our understanding of the genetics and evolution of this critical locus. For example, the “collaborative nonself recognition” system identified in the *Solanaceae* (Kubo *et al.* 2011) has revealed that the mechanism underlying self-incompatibility can change how new S-alleles originate (Bod'ová *et al.*, 2018, Harkness *et al.*, 2021) and migrate (Harkness & Brandvain, 2020) across populations. Identifying the basis (or

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98 bases) of SSI in the *Asteraceae* would help us better understand and predict features of its
evolution, including its maintenance in populations with very few (two to six) S-alleles (Brennan
100 *et al.* 2006).

102 To map the S-locus, researchers must determine the S-locus genotype for individuals in a
population large enough to conduct linkage mapping. In other species, S-genotype determination
104 required mating large numbers of full-sibling individuals to one or a few “tester” genotypes with
a known S-genotype (Camargo *et al.*, 1997; Tomita *et al.*, 2004). Testers may be a parent of the
106 population, or may be obtained by mating siblings in a diallel design, grouping individuals based
on their compatibility (Hiscock, 2000), and then selecting one or several of these individuals as a
108 tester for their siblings (Gonthier *et al.*, 2013). However, this process has several limitations. For
some species, pollen availability may limit the number of testcrosses that can be made, especially
110 if clonal propagation is not used to multiply tester individuals. Additionally, because the tester is
typically used as the male parent, differentiation between alleles may be difficult for SSI as
112 dominance relationship between alleles may differ from anthers to stigma. Finally, this process
often requires a large number of crosses, especially if the parents are not available. A diallel
114 requires $n(n-1)$ crosses, where n is the number of individuals to be analyzed, and multiple
crosses are likely required for each individual being tested. Even selecting a tester from amongst
116 a conservative number of siblings may require a prohibitively high number of crosses. Methods
of S-allele determination that address these issues may make S-locus mapping feasible for a
118 greater number of species.

120 In this study, we present a putative location for the *S. integrifolium* S-locus. To achieve this, we
developed a novel framework for inferring the S-genotypes of individuals within a population
122 large enough for mapping. Our method does not require tester individuals with known S-
genotypes, requires only three to four crosses per individual, and identifies alleles with sex-
124 specific dominance interactions. We used these inferred genotypes, in conjunction with the
aforementioned linkage map, to identify a QTL likely containing the S-locus. We then identify
126 regions in the sunflower (*Helianthus annuus*) genome assembly syntenic with our putative S-
locus and use this information to identify potential gene candidates for S-allele determination.

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MATERIALS AND METHODS

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Population development: A linkage map was constructed using an F₁ population of 265 *Silphium integrifolium* (Michx.) individuals, derived from the crossing of two genotypes known as “965” and “1767”, selected at the Land Institute. These parents were chosen because they expressed agronomically favorable morphologies and differed for several phenological traits. Seventy-four seeds from this cross were grown at the Land Institute, and 191 were grown at the University of Minnesota. All 265 individuals were used for map construction, but only a subset of the 191 Minnesota individuals were used for S-allele determination. Following approximately two months of growth in a greenhouse, the Minnesota seedlings were transplanted to a field at the Minnesota Agricultural Experiment Station at the University of Minnesota St. Paul campus with 1.2 meters between each plant.

Genotyping and Variant Detection: Tissue was collected from seedling leaves of the 74 Land Institute progeny, and adult leaf tissue was collected from the parents and all other progeny, with all tissue lyophilized prior to DNA extraction. Dual indexed genotyping-by-sequencing (GBS) libraries were created from genomic DNA using the restriction enzymes *SbfI* and *TaqI* and sequenced on 1.5 Illumina NovaSeq 6000 lanes (1x100 single-end reads). All extraction, library preparation, and sequencing occurred at the University of Minnesota Genomics Center.

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Demultiplexed reads were mapped to *S. integrifolium* genomic contigs using BWA-MEM (Y. Brandvain, unpublished; Li, 2013). The “ref_map” pipeline of Version 2.5 of the “Stacks” variant calling software was then used for the detection of SNP loci (Catchen *et al.*, 2013). Loci that were missing in at least 30% of the population were excluded, as were loci missing in either parent. This resulted in 935 SNP markers.

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Linkage Map Construction: Linkage analysis and map construction were carried out in *JoinMap5* (Stam, 1993). First, 715 SNP markers showing significant (Chi-square test, $P > 0.01$) segregation distortion were removed. Markers were then grouped by LOD independence score, using a threshold score of five. Any groups that contained fewer than four markers were discarded. Map order and distances were then estimated using the maximum likelihood mapping

160 algorithm, with default settings. One marker was removed from the end of linkage group three
because it created a gap of more than 50 cM. Linkage group names were assigned based on
162 estimated centimorgan length, with group one being the longest.

164 **Crossing design to determine self-incompatibility genotype:** Of the 191 members of this
mapping population planted in Minnesota, 84 were intermated for S-allele determination. To
166 cross, capitula (compound flower heads) to be used as a female, or both as a male and female,
were covered with a cotton bag prior to anthesis to prevent pollination. Capitula to be used
168 exclusively as a male parent were covered with mesh bags at least one day prior to crossing to
reduce contamination by insects depositing pollen from other plants. After stigma emergence,
170 pollen from a male parent capitulum was collected into a container and dusted onto stigma with a
pipe cleaner. Alternatively, on occasion a capitulum was removed and used to brush pollen
172 directly onto a female parent. Each female capitulum was mated with only one male parent. The
bag was then reclosed, harvested after senescence, and dried. For each capitula, the number of
174 filled seeds and total number of seeds were then counted, with filled and unfilled seeds
differentiated by visual and manual assessment. Seed set was then calculated as the ratio of filled
176 seeds to total seeds.

178 Individuals were crossed in a structured design, which we have named the “connected small
diallel design”, illustrated in Figure 1. This structure is based on groupings of four individuals
180 that are mated in a diallel. These small diallels are then linked together through reciprocally
mating single individuals. The purpose of this design was to maximize the amount of
182 information that could be derived from the mating behavior of any given individual, while
minimizing the number of matings for which it would need to be used. The design ensures that
184 all individuals may be connected to one another through pairs of matings, allowing for the entire
population to be used to predict the mating behavior each individual.

186
The design for this experiment included 20 small diallels. The design was implemented
188 incompletely, with many of the recommended crosses not completed due to time or pollen
availability constraints. To ensure that all individuals could be connected, more crosses
190 connecting diallels were conducted than indicated in the design. Finally, several additional

individuals were included in the study with only reciprocal crosses to one other individual. In
192 total 268 crosses were performed between the 84 individuals, covering 138 different
combinations of parents. Of these crosses, 126 combinations were crossed reciprocally, with
194 both individuals used as male and female. One hundred and eighty-six crosses were conducted in
2018, and an additional 82 were conducted in 2019, primarily to increase the number of
196 reciprocal crosses. In seven cases, multiple replications of a cross were performed. These
observations were combined by summing the total number of filled and total seeds from each
198 replication, and then calculating a seed set from the sums.

200 Three distinct but complementary methods were used to translate the results of the connected
small diallel experiment into an S-locus map position. In all three methods, each mated pair of
202 individuals is first determined to be compatible or incompatible, based on a threshold seed set
value. A threshold value of 20% was selected for this experiment, based on previous
204 observations of seed set in manually self-pollinated individuals (Reinert *et al.*, 2020). Matings
that resulted in a seed set value above 20% were considered compatible. These methods, along
206 with their relative strengths and weaknesses, are described below. All methods were developed
using the R language (R Core Team, 2019).

208
Direct mapping with single-marker regression: The simplest method attempted to directly
210 associate variation for compatibility with an SNP marker, without inferring the S-genotype of
any particular individual. To accomplish this, a logistic regression was conducted following the
212 formula $S_i = M_{ij} + P_{ij} + M_{ij} \cdot P_{ij}$, where the success S of the i th mating was predicted by the
maternal genotype M , the paternal genotype P , and the interaction of those genotypes, all for the
214 j th marker. The calculated P-value of the maternal by paternal genotype interaction was then
used to determine association with cross incompatibility, and thus the S-locus. Only biallelic
216 markers were used in this approach, with homozygotes recoded as -1 or 1 and heterozygotes as
0. This approach only makes limited use of available genotype data, as each SNP marker is
218 considered individually. Additionally, this approach could not be applied to species without
genetic marker data. Finally, because it does not make inferences about the S-genotype carried
220 by any particular individual or take potential dominance relationships into account, it is not able

to fully leverage the advantages of the connected crossing design used in this study. Thus, this
222 method's usefulness is likely limited to confirming the results of other methods.

224 **Inference of self-incompatibility genotype for all individuals followed by QTL mapping of
the inferred S-genotype:** As an alternative approach to mapping the S-locus, we develop two
226 methods to infer every individual's S-locus genotype from its crossing behavior, then map this
inferred phenotype with traditional QTL mapping software. Both methods assumed that the two
228 parents of the population were each heterozygous at the S-locus and did not share any S-alleles
with each other, resulting in a population with four distinct S-alleles and therefore four distinct
230 S-genotypes. Each method is described in detail below. Code to replicate all methods may be
found on GitHub (see data availability statement).

232

Approach 1: A hill-climbing algorithm: The first S-allele determination method used a simple
234 hill-climbing algorithm to fit genotypes, given a user-generated set of dominance relationships
among alleles. To start, the user hypothesizes a set of crossing relationships, dictating which of
236 the sixteen possible pairings of genotypes will and will not be able to successfully mate. A
random genotype is then assigned to each individual in the population. One arbitrarily selected
238 individual is always set to a predetermined genotype to enable comparison of different runs of
the algorithm. Each mated pair in the dataset is then scored using the user-supplied crossing
240 relationships. A mismatch between the observed and predicted outcomes results in a score
between 0.5 and 1, determined by the user. A match results in a score of one minus the
242 previously mentioned score. Scores for the entire dataset are then summed. Next, the genotype of
one individual is randomly changed, and the dataset rescored. If the total score decreases, then
244 that set of genotypes is kept; if not, the algorithm returns to the previous position.

246 The cycle is repeated, stopping if 1,000 changes are attempted without decreasing the score. For
this experiment, the algorithm was run 4,000 times for each set of dominance relationships, with
248 the solution to the lowest scoring run considered optimal genotype assignments for a given
dominance relationship. The lowest-scoring genotype assignments across all dominance
250 relationships were then used for QTL mapping. We compared the results of 48 dominance

relationships, each of which met three criteria: 1) individuals sharing both alleles must be
252 incompatible, 2) individuals sharing no alleles must be compatible, and 3) there must be at least
one combination of genotypes that was asymmetrical (compatible when one individual was used
254 as a female and incompatible when the other was used as a female), as this was observed in the
crossing data.

256
This method has both the advantage and disadvantage of being highly parameterizable. This
258 makes it flexible, and gives the user a high degree of control, but may take many attempts to find
the right combination of parameters to develop a solution. This method may be employed using a
260 personal computer, however the reduction in parallelization necessary to achieve this may make
it take too long to be useful. In addition, as a hill-climbing method there is no guarantee that this
262 approach will find a global maximum.

264 *Approach 2: A Markov Chain Monte Carlo algorithm:* The second method employed sets of
extreme gradient boosting decision trees with Markov chain Monte Carlo (MCMC) algorithms to
266 infer S-locus genotypes. For each MCMC chain, an initial set of genotypes was created by first
assigning a random heterozygous S-genotype to one random individual. The genotype of an
268 individual that had been mated to the initial individual was then determined. If the cross was
incompatible, the second individual's genotype could share one or two alleles with the initial
270 individual's genotype, with equal probability. If the cross was compatible, the second
individual's genotype could share either zero or one allele with the initial plant's genotype, with
272 equal probability. The procedure was then repeated by choosing, at random, an individual that
did not yet have a genotype assigned and that was crossed to the most recent individual assigned
274 a genotype. If all individuals that were crossed to the most recently assigned individual had an
assigned genotype, a random individual that did not yet have an assigned genotype was selected
276 and the process started again until all individual were assigned genotypes.

278 Maternal and paternal S-locus genotypes were then treated as predictive variables with cross
success used as a binary response variable. Constraints were placed on predictive variables so
280 that matching maternal and paternal genotypes were not allowed to interact with one another, as
crosses between individuals with matching genotypes should always be incompatible. At each

282 step of an MCMC chain, model performance was measured as error of a logistic regression for
classification using an extreme gradient boosting model with four-fold cross validation with 200
284 iterations and maximum tree depth of four splits, using the R package “xgboost” (Chen &
Guestin, 2016). A single genotype was altered at each MCMC chain step, using the same sets
286 of probabilities used to construct the initial set. The newly proposed set of genotypes was
accepted if the ratio of errors from the proposed genotype set to the former genotype set was
288 greater than a randomly drawn number bound by zero and one.

290 Each MCMC chain, starting from a unique set of genotypes, was allowed to explore parameter
space for 96 hours (~ 1 million steps). The set of genotypes that produced the smallest error from
292 each chain was then used as the starting condition for hill-climbing algorithms. We used the
same extreme gradient boosting model conditions described above, however at each step
294 proposed genotypes were only accepted if they produced an error that was less than the previous
set of genotypes. The hill-climbing algorithms each ran for 96 hours. For each chain, the set of
296 genotypes that produced the smallest error, calculated as the percentage of crosses classified
incorrectly, were recorded. A cross was predicted to be a success if the estimated logistic
298 regression probability was greater than 0.5. The lowest error models were considered the best
candidates for S-genotype determination. This method has the advantage of less dependence on
300 user decisions, making it more repeatable and facilitating the exploration of a wider space of
possible solutions. However, the resources required by this method mean that it generally cannot
302 be employed on a personal computer.

304 *Mapping the S-locus from inferred S-locus genotypes:* Assigned alleles from the two inference
methods were then used as phenotypes for QTL mapping, using the “qtl” R package (Broman *et*
306 *al.*, 2003), with the population treated as a four-way testcross. Marker genotypes were imputed
using the “sim.geno” function, and QTL were identified using the “scanone” function to conduct
308 interval mapping using the EM algorithm. Self-incompatibility allele was treated as a binary
trait, with one allele from each parent arbitrarily coded as “1” and the other as “0”, and the two
310 parental alleles mapped separately. For the MCMC method, all 96 of the inferred genotype sets
were mapped, with the maximum LOD score produced by each set serving as a criterion to
312 differentiate the sets. Eight genotype sets produced by the hill-climbing method were also

mapped. Significance thresholds were determined independently for each set of genotypes used
314 for mapping; only QTL with an error probability less than 5% were considered significant. This
threshold ranged from a LOD score of 3.6 to 4.2. Finally, a Bayesian credible interval (similar to
316 a confidence interval) around any identified QTL was calculated, using the “bayesint” function.

318 **Synteny with related species:** To help elucidate the relationship between the *S. integrifolium* S-
locus and other *Asteraceae*, the genomic contigs associated with the loci that comprise this
320 linkage map were aligned to the annual sunflower (*Helianthus annuus*, variety ‘HA412’, version
HOv1.1) (Badouin *et al.*, 2017) and lettuce (*Lactuca sativa*, variety ‘Salinas’, version 7) (Reyes-
322 Chin-Wo *et al.*, 2017) genome assemblies using BLASTN (Altschul *et al.*, 1990). Syntenic
regions were then identified using the R package “syntR” (Ostevik *et al.*, 2020). For synteny
324 analysis, all BLASTN alignments for a given *S. integrifolium* query sequence that had a bitscore
greater than 90% of the maximum bitscore for that query sequence were used.

326
Identification of candidate genes: For regions identified as syntenic to the *S. integrifolium* S-
328 locus, gene annotations for sunflower were obtained from the genome assembly HA412v1
(Badouin *et al.*, 2017). InterPro taxonomy was searched for each gene in the region to identify
330 plausible candidates by shared terms with S-genes in other species. The male and female
determinants from *Brassica oleracea* (*Brassicaceae*) and *Petunia hybrida* (*Solanaceae*), as well
332 as the female determinant of *Papaver rhoeas* (*Ranunculales*) chosen as representatives for
comparison (The UniProt Consortium, 2019). In addition, defensin-like proteins were considered
334 a proxy for the *Ipomoea trifida* (*Convolvulaceae*) male determinant (Rahman *et al.*,
2007). These species were selected because their self-recognition genes are relatively well
336 characterized and they represent a broad range of Eudicot diversity, including both SSI and GSI
systems. Any genes homologous to known CRPs were also considered potential candidates
338 (Marshall *et al.*, 2011). The best homologs for a subset of these candidates were then identified
in the *S. integrifolium* reference transcriptome (Raduski *et al.*, 2021), and their expression was
340 measured in a previously-published set of whole-plant seedling transcriptomes from a diversity
panel of 73 wild collected *S. integrifolium* accessions (Raduski *et al.*, 2021). Expression was
342 quantified in total transcripts per million (TPM), TPM as a percentile of total expressed genes,
and TPM as a percentile of a reference set of conserved single copy genes, known as BUSCO

344 (Simão *et al.*, 2015). We hypothesized that strong S-locus gene candidates would not be highly
expressed in seedling vegetative tissue (Williams *et al.*, 2014).

346
Additionally, the coding region sequence for the *B. oleracea* S-receptor kinase protein (Stein *et*
348 *al.*, 1991) was aligned to the lettuce and sunflower genomes using BLAST. These alignments
were then compared to the lettuce and sunflower regions syntenic to the *S. integrifolium* S-locus.

350

RESULTS

352

Linkage Map: The linkage map contains 198 markers, spanning 1,049 centimorgans and
354 divided into 7 linkage groups (Fig. 2, Table S1). This is consistent with the observation that *S.*
integrifolium has seven chromosomes (Settle, 1967). The average distance between markers is
356 5.5 cM, with 29 gaps greater than 10 cM.

358 **S-locus mapping:** Of the 268 crosses completed, 53% were incompatible, with a seed set value
less than 20%, and 47% were compatible. These frequencies were not significantly different
360 from an equal occurrence of compatible and incompatible crosses ($df = 1$, $\chi^2 = 0.956$, $P = 0.328$).
This indicates there is a hierarchy of dominance between alleles in this population—if all alleles
362 were codominant a ratio of 25% compatible to 75% incompatible would be expected. Of the 126
reciprocal pairs of genotypes crossed, 30% were compatible in both crossing directions (both
364 individuals could be used as male and female), 34% were incompatible, and 36% were
asymmetrical, or compatible when one individual was used as a female and incompatible when
366 the other was used as a female. The presence of asymmetrical crosses indicates that the
dominance relationship between some allele pairs differs from anthers to stigma. These
368 observations confirm that *S. integrifolium* has a sporophytic SI system, as dominance
relationships are not observed in gametophytic systems (Breton *et al.*, 2014). Figure 3 shows the
370 distribution of seed set values for reciprocal crosses, numerical data for all crosses may be found
in Table S2.

372

Single marker regression for male/female interaction: The interaction of male and female
374 marker genotype was found to be a significant ($P > 0.01$) predictor of crossing success for three

376 markers, located at two regions on the linkage map: marker “247873”, located at 31.3 cM on
linkage group six ($P = 0.0006$, $df = 251$), and markers “140641” and “16612”, located at 143 cM
and 130.9 cM, respectively, on linkage group 2 ($P = 0.0013$, $P = 0.003$, $df = 251$ for both).

378

Hill-climbing algorithm method:

380 Of the 48 dominance relationships tested using the hill-climbing algorithm, nine were used in
QTL mapping. Of these, seven produced at least one significant LOD peak, all located on
382 linkage group 6. The highest of these ($LOD = 5.3$) was located at 18.9 cM, and all but one of the
other six were located between 9- 31 cM.

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MCMC method: Of the 192 genotype sets produced by the MCMC method (96 models, with a
386 separate genotype set for each of the two parental alleles), 30 produced at least one significant
LOD peak when used for QTL mapping, covering five linkage groups. Eleven of these genotype
388 sets produce a significant LOD peak at either 18.9 or 27.7 cM on linkage group 6, and a further
five were located elsewhere between 0-31 cM on linkage group 6. No other linkage group
390 contained more than five of the 30 significant LOD peaks. The highest LOD peak produced by a
single MCMC model was 7.56, located at 18.9 cM on linkage group six. This model also had the
392 lowest error, and so we have selected it as the best result from the MCMC method. This set of
genotype assignments is illustrated in Figure 4.

394

Consensus map location: Of the three methods used to map the S-locus, the MCMC method
396 produced the strongest association between a genomic region and S-allele. The peak of this QTL
was located at 18.9 cM on linkage group six, with a 95% probability Bayes credible interval
398 from 0 to 27.7 cM. Broadly speaking, the results of the hill-climbing method agree with this
region, although the credible interval for several of those results extend to 31 cM. Additionally,
400 the most significant single-marker regression association between the interaction of male and
female genotype with cross success was found at 31 cM. Taken together, we conclude that the *S.*
402 *integrifolium* S-locus is located between 0 and 31 cM on linkage group 6, with the marker closest
to the locus likely located at 18.9 cM.

404

S-locus synteny with related species: Synteny was found between this the putative *S. integrifolium* S-locus region and both the sunflower and lettuce genomes. In sunflower, synteny was found with two chromosomes—the *S. integrifolium* 0-18.9 cM region was found to align with chromosome 17, from 230.7 Mb to 262.67 Mb, and the 18.9- 31cM region was found to align with chromosome 3, from 133.21 Mb to 149.9 Mb. In lettuce, the 0-9 cM region aligned with chromosome 4, from 18.13 Mb to 27.29 Mb, and the 18.9-31 cM region aligned with chromosome 9, from 144.19 Mb to 147.34 Mb. The syntenic region on sunflower chromosome 17 does not appear to overlap with the putative self-incompatibility breakdown QTL found on that chromosome by Gandhi *et al.* (2005), which we estimate was somewhere between 42 and 98 Mb, based on the alignment of SSR marker primer sequences to the sunflower genome assembly.

Identification of candidate genes: Of the 642 genes located within the sunflower genomic regions syntenic with the *S. integrifolium* S-locus, 42 shared at least one InterPro term with either the male or female determinants of self-incompatibility in *B. oleracea* or *P. hybrida*, the female determinant of *P. rhoeas*, or the male determinant of *I. trifida*. Thirty of these genes were, like *B. oleracea* SRK, protein kinases, which is not a specific enough homology to infer a gene candidate. The other 12 genes were, like the female determinant in *P. hybrida*, F-box genes. In addition, one gene was similar to “STIG1”, a CRP. The expression of this gene, as well as two of the 12 F-box genes, were then measured in the 73 diversity panel seedling transcriptomes. One of these F-box genes was expressed in 65 of the seedling transcriptomes, with TPM as a percentile of BUSCO genes ranging from 5.4 to 79.5. The other was expressed in 42 of the seedlings, with TPM as a percentile of BUSCO genes ranging from 0.09 to 41.2. The STIG1-like gene was expressed in five of the seedling transcriptomes; expression varied widely but was always relatively low (Table 1).

The best alignment for *B. oleracea* SRK in sunflower was found on chromosome 11, while the best alignment in lettuce was found on chromosome 7. Neither of these alignments are near any regions syntenic with the putative *S. integrifolium* S-locus in either species.

434

DISCUSSION

436

The described S-locus mapping methods are effective and efficient: This study represents one
438 of the few successful identifications of a genomic region containing the S-locus in an *Asteraceae*
species. This result is valuable for its own sake, as the comparison of syntenic genomic regions
440 between *S. integrifolium* and other species may help to answer questions about the evolution of
SSI systems in other *Asteraceae*. In addition, this study lays the groundwork for future efforts to
442 clone the *S. integrifolium* S-locus.

444 In this experiment, we were able to predict the S-locus genotype for 84 individuals by
conducting 268 crosses, with an average of 3.2 crosses per individual. This method did not
446 require all 84 individuals to be crossed with a single tester, and so could be applied to a pre-
existing full-sibling family without necessitating clonal propagation of any individuals. As a
448 point of comparison, we estimate that the mapping of the S-locus in chicory, which combined a
twenty-two plant diallel with testcrossing, required approximately 3,000 crosses (Gonthier *et al.*,
450 2013). That approach, which also required the clonal propagation of tester genotypes, was used
to assign an S-genotype to approximately 350 individuals (multiple replications of each cross
452 were performed). The chicory mapping effort produced a more accurate determination of S-
genotype and thus a narrower S-locus QTL region than our study, but with the trade-off of
454 requiring more crosses per individual and the clonal replication of tester genotypes. Additionally,
our method requires less time to complete, as testcrossing cannot be conducted until the diallel is
456 completed, the results interpreted, and selected testers clonally propagated. Finally, our method
does not require every cross to be replicated, as the analysis relies on inference methods that are
458 relatively robust to error in the data.

460 Our method should be readily applicable to any other species that meet three criteria: 1) They
possess an SSI system, 2) large full sibling families can be produced, and 3) individuals can
462 readily be used as both a male and female parent in three to five matings. Among numerous other
species, self-incompatible members of the genus *Helianthus* could be excellent candidates for
464 this method and could help to confirm our results. Applying this method to phylogenetically
diverse *Asteraceae* species could evaluate the conservation of the method of SSI in the family.
466 Additionally, with some modifications, our framework could be used for GSI species.

468 **Evaluation of potential S-locus candidate genes:** Of the 642 annotated or predicted genes
present in the sunflower genomic regions syntenic to the putative S-locus, 43 show potential
470 similarity to known S-genes. Of these, two groups of genes were considered particularly
interesting, and are detailed below. It is important to note that because virtually nothing is known
472 about the molecular mechanisms that underlie SSI in the *Asteraceae*, any discussion of potential
candidate genes based on annotations or similarity to S-related genes in other systems is at best
474 informed speculation. However, we see this as an important step towards identifying the genes
that control SSI in the *Asteraceae*, and therefore believe it is worth pursuing.

476
One potential candidate is the sunflower gene Ha3_00036110, which is annotated as a putative
478 member of the “Stigma Specific Protein 1” like, or STIG1-like, group of genes. In sunflower,
this gene is found on the chromosome 3 region syntenic with the 18.9-31 cM segment of the
480 putative *S. integrifolium* S-locus. In lettuce, the best BLAST alignment for Ha3_00036110 is on
chromosome 9, within the region syntenic to the putative *S. integrifolium* S-locus, implying that
482 this region may be conserved between the three species.

484 The best characterized member of this family, STIG1, encodes a small, cysteine-rich protein
(CRP). In tomato (*Solanum lycopersicum*), this protein is primarily found in stigma exudate, and
486 binds a pollen-specific kinase to promote pollen tube growth (Huang *et al.*, 2014). In tomato,
silencing either STIG1 or the pollen kinase resulted in slowed pollen tube growth and reduced
488 seed set (Huang *et al.*, 2014). More broadly speaking, as CRPs, STIG1-like genes are members
of a gene class that also includes the male determinant of SSI in the *Brassicaceae* and the
490 *Convolvulaceae*, and the female determinant of GSI in the genus *Papaver*. Based on this
similarity to other SI genes, the function of STIG1, and the location of this gene within the
492 sunflower genomic region syntenic to the *S. integrifolium* S-locus, we speculate that
Ha3_00036110 is a signaling protein that controls the female determinant of SSI in *S.*
494 *integrifolium*, with differential binding between different alleles of the STIG1-like protein and an
unidentified pollen kinase resulting in compatibility or incompatibility. If this is correct, a
496 possible candidate for this pollen kinase may be the gene Ha3_00036112, a serine/threonine
protein kinase located 109 Kbp downstream from Ha3_00036110.

498

The best *S. integrifolium* homolog for Ha3_00036110 was only expressed in five of 73 seedling transcriptomes, and Ha3_00036112 was found to be expressed in four of 73 seedlings. The two genes were never expressed in the same individual, and expression was relatively low, never exceeding the 33rd percentile of BUSCO genes (Table 1). The low level of seedling expression for these genes further supports their candidacy, as S-gene expression is expected to be primarily limited to floral tissue (Williams *et al.*, 2014). Expression in other tissues may still occur at low levels, either due to gene expression noise or as-yet uncharacterized uses for the gene product. For example, in tea (*Camellia sinensis*), an S-gene candidate was found to be expressed in leaf tissue at approximately 20% the level it was expressed in style tissue (Zhang *et al.*, 2016).

508

Another set of potential candidate genes are found on the sunflower chromosome 17 region syntenic with the 0-18.9 cM segment of the putative *S. integrifolium* S-locus. This region contains 12 F-box genes, resembling known gametophytic self-incompatibility (GSI) systems. For example, in *Petunia*, 17 tightly linked F-box genes serve as the male determinant of GSI (Williams *et al.*, 2015). However, no genes in this sunflower region share any annotation terms with known S-RNase genes (The UniProt Consortium, 2019), which serve as the female determinant in GSI systems. In addition, two of these genes that were arbitrarily selected as representatives were found to be expressed in more than half of available seedling transcriptomes, further diminishing the likelihood that they control the self-incompatibility response. Because so little is known about the molecular basis of the *Asteraceae* SSI response, these genes cannot be completely discounted, but they appear to be less likely candidates than the STIG1-like gene.

522 In addition to providing candidates for genes involved in *S. integrifolium* self-recognition, this study suggests that SRK-like genes are not involved in *S. integrifolium* SSI, as the best homologues for the *B. oleracea* SRK in the lettuce and sunflower genomes are not found in or near the regions of those genomes syntenic with the putative *S. integrifolium* S-locus. This finding adds evidence to the theory that specific genes that underlie *Asteraceae* SSI are different than those in the *Brassicaceae* (Allen & Hiscock, 2008).

528 **Implications for domestication and breeding:** We expect the results of this study to facilitate
the domestication of *S. integrifolium*. The availability of a genetic map, associated with
530 particular restriction enzymes, supports the relatively inexpensive GBS genotyping of large
numbers of progeny; both because these enzymes are now known to be useable in *S.*
532 *integrifolium*, and because it is reasonably likely that some of the same loci would be recovered
if these enzymes were applied to a different population. This may support the implementation of
534 marker-based selection, marker-based pedigree development, and genome-wide selection.
Additionally, this genetic map will assist in the anchoring and orientation of future genome
536 assemblies.

538 Identifying a map location for the S-locus may also contribute to practical breeding efforts. If
molecular marker data is routinely available for individuals within a breeding program, it may be
540 possible to predict whether any two plants will be able to successfully mate, through the direct
identification of particular S-alleles or ancestral haplotype blocks in the S-region. This
542 information could save effort by excluding crosses that would not be successful. Perhaps more
importantly, this would allow for the S-allele diversity of a given population to be monitored and
544 maximized. It is likely that at least some *S. integrifolium* cultivars will take the form of synthetic
populations, where a set of superior genotypes are intermated and their progeny form a distinct
546 variety that may be reproduced for several generations. If a synthetic population is released that
contains a low number of S-alleles, its long-term fecundity may be adversely affected by the
548 limited number of individuals that are able to intermate. Alternatively, as *S. integrifolium* is
known to express moderate to severe inbreeding depression for a number of traits (Price *et al.*,
550 2021), it is possible that S-allele characterization information could be used increase long-term
productivity by limited mating among relatives for several generations within synthetic
552 populations.

554 Overall, we anticipate that the availability of a genetic map and identification of the self-
incompatibility locus will support efforts to domesticate *S. integrifolium* as a crop that will help
556 to enable sustainable agricultural systems.

558

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572 AUTHOR CONTRIBUTIONS

The effort to develop a linkage map was initiated by KPS, DLVT, and YB, who also secured
574 initial funding. DLVT created the mapping population, contributed text and visualization, and
provided feedback in writing. JHP constructed the linkage map, initiated and designed the S-
576 locus mapping work, and developed the HC method. ARR developed the MCMC method. JHP
and ARR conducted data analysis and wrote the manuscript. KPS and YB provided supervision,
578 and feedback on analysis, writing, and visualization.

580 DATA AVAILABILITY

All code to replicate analyses may be found on GitHub, at [https://github.com/UMN-
582 BarleyOatSilphium/SilphiumSLocus](https://github.com/UMN-BarleyOatSilphium/SilphiumSLocus). All sequence data may be accessed under BioProject
PRJNA695552.

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766 SUPPORTING INFORMATION

Table S1: The linkage map in tabular form.

- 768 Table S2: All crossing data used in this experiment.

Table 1: Expression of two S-gene candidates in a set of whole-plant seedling transcriptomes from a diversity panel of 73 wild collected *Silphium integrifolium* accessions. Expression is for *S. integrifolium* homologs of the named sunflower genes.

Sunflower gene	Description	Plant ID	Transcripts per million (TPM)	TPM as percentile of all genes	TPM as a percentile of BUSCO genes
Ha3_00036110	“Stig1-like” gene	C7	0.13	0.076	0.052
		D8	2.26	0.352	0.332
		E4	0.57	0.256	0.206
		E5	0.57	0.264	0.237
		H2	0.2	0.130	0.096
Ha3_00036112	serine/threonine protein kinase	B3	0.27	0.177	0.130
		B9	1.14	0.315	0.305
		E8	1.37	0.340	0.329
		H7	0.07	0.027	0.019

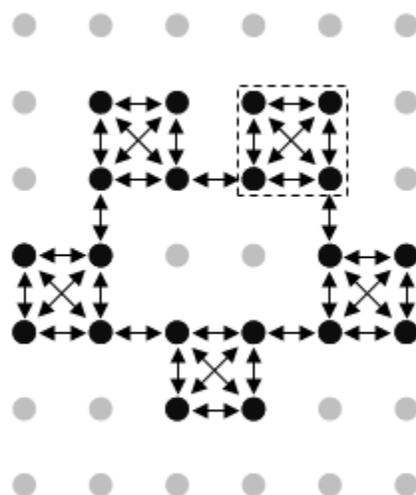


Figure 1: Visualization of the crossing design used in this experiment. Each circle represents one plant, with black circles representing plants that were used for crosses, and gray plants representing other members of the mapping population that were not selected. The dashed box represents one of the small diallels that formed the basis for the design—twenty of these were used for this experiment. Arrows represent matings, with each arrow pointing from the male parent to the female parent.

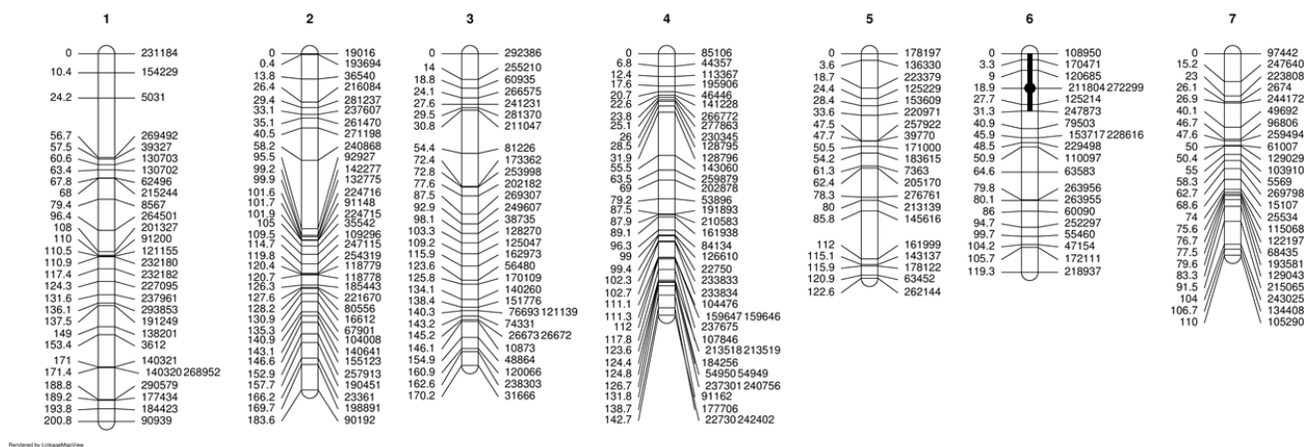


Figure 2: Linkage map for *Silphium integrifolium*. The black bar on linkage group 6 represents the putative S-locus QTL, with the circle showing the LOD peak and the extent of the bar showing a composite of the 95% Bayes credible intervals for several mapping methods. Map visualization was performed using “LinkageMapView” (Ouellette *et al.*, 2018).

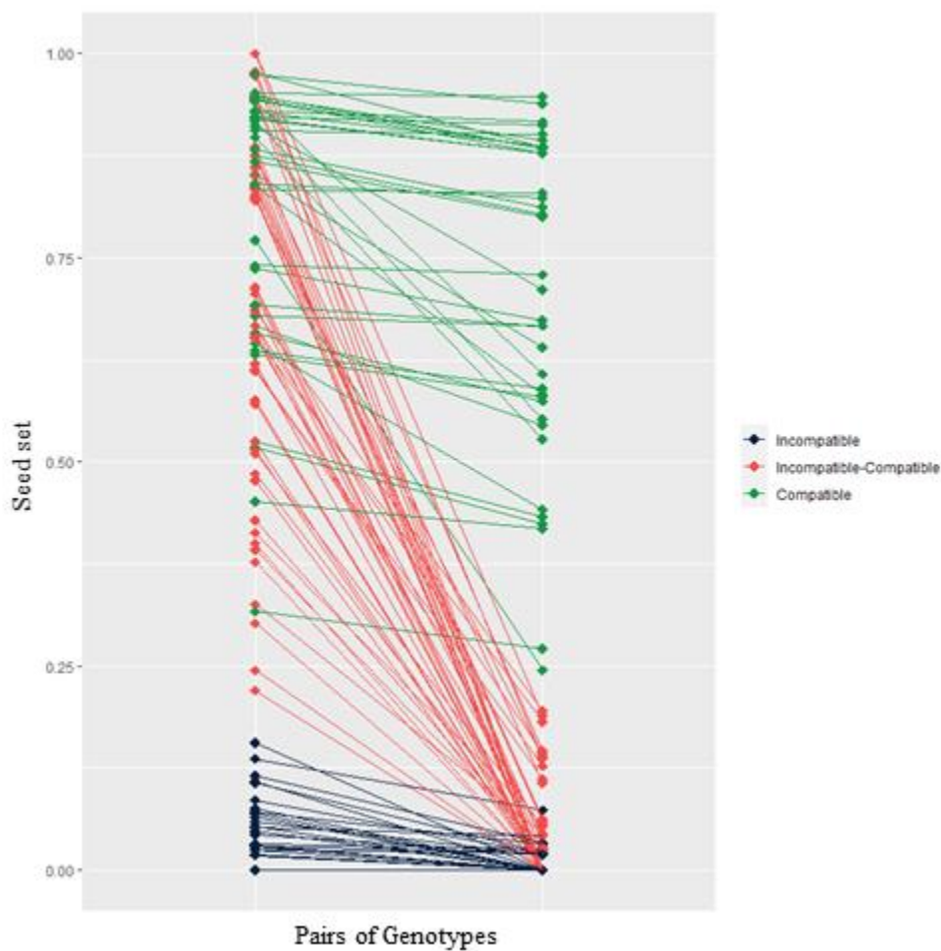


Figure 3: Distribution of seed set values for reciprocal matings (pairs of individuals mated, with each individual used as both a male and female). Each dot represents the seed set value of one plant, with lines connecting mated pairs. The individuals with the higher seed set value in the pair are placed on the left side of the chart.

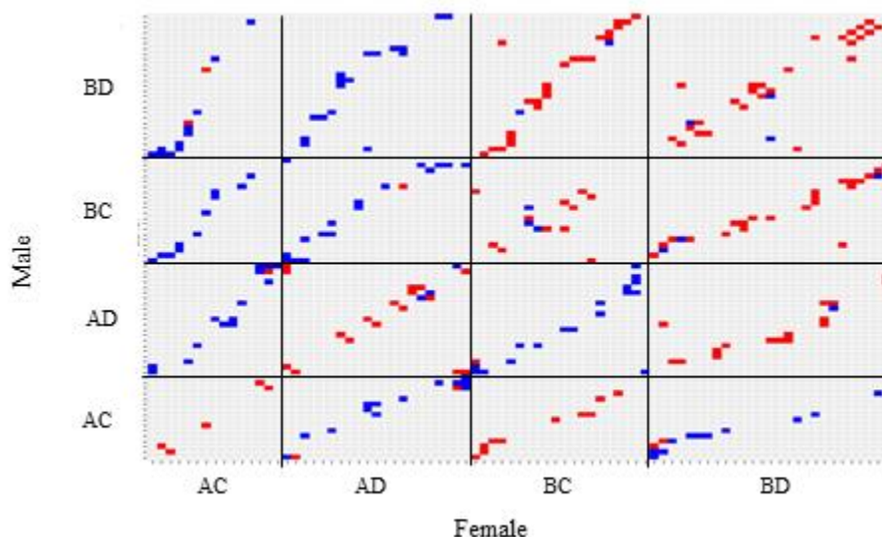


Figure 4: Representation of all possible matings that could have been completed for this study, with each square in the heatmap representing one pairing. Red or blue squares represent the crosses that were actually made, with red representing incompatible crosses, and blue representing compatible. Individuals are grouped by S-locus genotype, as assigned by the best MCMC genotype inference model. “A” and “B” refer to the S-alleles from one parent of the population, and “C” and “D” to the alleles from the other.