1 Intraspecific genetic variation underlying postmating reproductive barriers between species

- 2 in the wild tomato clade (*Solanum* sect. *Lycopersicon*)
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22 Abstract

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24 A goal of speciation genetics is to understand how the genetic components underlying 25 interspecific reproductive barriers originate within species. Unilateral incompatibility (UI) is a 26 postmating prezygotic barrier in which pollen rejection in the female reproductive tract (style) 27 occurs in only one direction of an interspecific cross. Natural variation in the strength of UI has been observed among populations within species in the wild tomato clade. In some cases, 28 29 molecular loci underlying self-incompatibility (SI) are associated with this variation in UI, but the 30 mechanistic connection between these intra- and inter-specific pollen rejection behaviors is 31 poorly understood in most instances. We generated an F₂ population between SI and SC 32 genotypes of a single species, Solanum pennellii, to examine the genetic basis of intraspecific variation in the strength of UI against other species, and to determine whether loci underlying 33 34 SI are genetically associated with this variation. We found that F_2 individuals vary in the rate at 35 which UI rejection occurs. One large effect QTL detected for this trait co-localized with the SI-36 determining S-locus. Moreover, individuals that expressed S-RNase—the S-locus protein involved in SI pollen rejection—in their styles had much more rapid UI responses compared to 37 38 those without S-RNase protein. Our analysis shows that intraspecific variation at mate choice 39 loci—in this case at loci that prevent self-fertilization—can contribute to variation in the 40 strength of interspecific isolation, including postmating prezygotic barriers. Understanding the 41 nature of such standing variation can provide insight into the accumulation of these barriers 42 between diverging lineages. 43

44 **Key Words:** self-compatibility, QTL, reproductive isolation, postmating prezygotic, unilateral

- 45 incompatibility, S-locus
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47 Introduction

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49 Speciation involves the accumulation of genetic differences and—in sexually 50 reproducing organisms—reproductive isolation, among diverging lineages. Accordingly, loci that 51 contribute to this cumulative process between species must first arise within an individual 52 population prior to spreading to other conspecific populations within their own lineage. During 53 this process, populations of a single species are expected to show varying strengths of 54 reproductive isolation against other lineages; that is, there will be intraspecific genetic variation 55 for the magnitude of interspecific reproductive isolation from other lineages. Intraspecific phenotypic variation in the strength of hybrid incompatibility has been observed in many 56 57 systems including mammals (Good, Handel and Nachman, 2007; Vyskočilová, Pražanová and 58 Piálek, 2009), arthropods (Bordenstein, Drapeau and Werren, 2000; Kopp and Frank, 2005; 59 Reed and Markow, 2004; Shuker, et al., 2005), nematodes (Kozlowska, et al., 2012) and plants 60 (Case and Willis, 2008; Leppälä and Savolainen, 2011; Martin and Willis, 2010; Rieseberg, 2000; 61 Sweigart, Mason and Willis, 2007). Understanding the nature, origin, and accumulation of this 62 variation, including the underlying molecular genetic variants responsible, can provide insight 63 into the evolutionary dynamics of lineage divergence (Cutter, 2012), including the order in 64 which alleles contributing to interspecific reproductive isolation arise and fix within diverging 65 lineages.

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67 The genetic basis of intraspecific variation for interspecific barriers has been investigated in few cases, most of which focus on postzygotic isolating barriers. Some of these 68 69 studies have confirmed that variable reproductive isolation is due to genetic variation between 70 populations of a species, but have not identified the specific loci or gene(s) responsible [e.g. 71 (Kozlowska, et al., 2012; Machado, Haselkorn and Noor, 2007)]. In other cases, isolation 72 variation has been mapped to localized chromosomal regions (quantitative trait loci, or QTL) or 73 even individual loci, whose geographic distribution is then investigated. For example, between 74 the plant sister species Mimulus nasutus and M. guttatus, the M. guttatus hybrid male sterility 75 1 (hms1) allele interacts with the M. nasutus hms2 allele to cause male sterility in hybrids; the 76 *M. nasutus hms2* allele is common across populations, but the interacting *hms1* allele is geographically restricted within *M. auttatus* (Martin and Willis, 2010; Sweigart, et al., 2007; 77 78 Zwellig and Sweigart 2018a, b). This and other studies of hybrid sterility and inviability (e.g., 79 Leppälä and Savolainen, 2011; Reed and Markow, 2004; Shuker, et al., 2005) confirm that 80 variation in the strength of interspecific isolation between populations within species can be 81 due to standing genetic variation. 82

83 Compared to these studies of variable postzygotic isolating barriers, fewer analyses 84 address within-species polymorphism for prezygotic reproductive isolation [for an exception, 85 see (Hopkins and Rausher, 2012)]. Because prezygotic barriers act earlier in reproduction, they 86 could have a much larger role than postzygotic barriers in restricting levels of gene flow between closely related species (Lowry, et al., 2008; Rieseberg and Willis, 2007). Of these, 87 88 postmating prezygotic interactions can be particularly important for reproductive isolation 89 when species are only weakly isolated by other prezygotic mechanisms, such as pollinator 90 isolation [reviewed in (Swanson, Edland, and Pruess 2004; Moyle, Jewell and Kostyun, 2014)].

92 Unilateral incompatibility (UI) is an example of a postmating prezygotic isolating barrier 93 that shows variation in strength among populations within species. In plants, this barrier 94 manifests after pollen transfer, as the (male) pollen grains germinate and produce pollen tubes 95 that grow down the female reproductive tract (the 'pistil', composed of the stigma (the pollen receiving site), the ovary, and the style which connects them) towards individual ovules. UI 96 97 occurs between species when pollen rejection in the female style occurs in only one direction of 98 an interspecific cross (and is therefore 'unilateral'; de Nettancourt, 1977; Lewis and Crowe, 99 1958), while the reciprocal cross results in pollen tubes successfully growing down the style and 100 into the ovary. UI often follows an 'SI x SC rule' in which genetically self-incompatible (SI) 101 species reject pollen from self-compatible (SC) species but the reciprocal cross is successful (de 102 Nettancourt, 1977; Lewis and Crowe, 1958; Murfett, et al., 1996); however, there are 103 exceptions to this rule even within species (Baek, et al., 2015). In the wild tomato clade, species 104 that are largely SI and display strong UI responses can nevertheless include SC populations that 105 exhibit weakened UI. In some species, the UI response is less rapid in SC populations, but in 106 other species, SC populations fail to reject heterospecific pollen altogether (i.e. UI is lost in 107 these populations) (Baek, et al., 2015, and see Discussion).

109 These observations suggest that among-population variation in the strength of 110 interspecific UI might be mechanistically associated with molecular factors contributing to SI (Li and Chetelat, 2015; Tovar-Méndez, et al., 2014). However, the extent to which UI and SI are 111 112 consistently genetically associated remains unclear. Within the plant family Solanaceae, a 113 primary determinant of gametophytic self-incompatibility is the S-locus (McClure, Cruz-Garcia 114 and Romero, 2011; McClure, et al., 1989) which encodes at least two proteins responsible for 115 the self-rejection mechanism: an S-RNase protein (the stylar component) that recognizes one or 116 more pollen-expressed F-box protein(s) in germinated pollen tubes and arrests pollen tube 117 growth within the style (Kubo, et al., 2015; Kubo, et al., 2010; Li and Chetelat, 2015; Sijacic, et 118 al., 2004; Williams, et al., 2014). Pollen is rejected when a haploid pollen tube bears an S-119 haplotype that is identical one of the S-haplotypes of the pistil (maternal) parent (McClure, et 120 al., 1989). Loss of SI in the wild tomato clade is frequently associated with mutations in the 121 locus producing S-RNase (Bedinger, et al., 2011; Igic, Lande and Kohn, 2008; Rick and Chetelat, 122 1991). Similarly, within predominantly SI species, population-level transitions to SC are also 123 often associated with the loss of S-RNase function. Within the two wild tomato species S. 124 habrochaites and S. pennellii, for example, several SC populations have been shown to lack S-125 RNase expression in stylar tissue (Kondo et al. 2002). Despite the loss of S-RNase, however, 126 most SC populations of these wild species still exhibit UI against interspecific pollen, indicating UI can also have S-RNase-independent mechanisms. Nonetheless, the degree to which natural 127 128 intraspecific variation in genes involved in SI might simultaneously affect interspecific isolation 129 via UI remains to be directly investigated in most cases (but see Broz et al. 2017, Markova et al. 130 2016).

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In this study, we generated an F₂ mapping population between two populations within a
 single species to map QTL underlying variation in the strength of UI against a second, tester,
 species. Our two parental genotypes were drawn from an SI population (*S. pennellii* accession)

LA3778) and a conspecific SC population (S. pennellii accession LA0716) which has recently lost 135 136 SI; both these genotypes express UI, but differ in how fast they reject pollen from domesticated 137 tomato pollen and other SC species (Results). Our goal was to determine the genetic basis of 138 this standing variation for the rate of UI rejection within our target species, and its association 139 with molecular loci underlying SI. We quantified UI response, evaluated SI status, and measured 140 several floral and fertility traits in the recombinant F₂ population. We assessed 1) the number of 141 large effect QTL that contribute to variation in the UI response within S. pennellii, 2) the 142 association, if any, between these UI loci and a priori candidate loci known to contribute to 143 intraspecific SI variation, and 3) the degree of association, if any, between UI phenotypes and 144 floral or fertility traits. These data allow us to assess whether the strength of UI (and changes in 145 SI status) in different intraspecific Solanum lineages is due to changes at the same underlying 146 loci.

- 147
- 148 Materials and Methods
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150 Generating the F₂ population

151 The wild tomato clade, Solanum sect. Lycopersicon is a group within the diverse 152 nightshade family Solanaceae that consists of 13 closely related [<2.5 MY old; (Pease, et al., 153 2016; Peralta, Spooner and Knapp, 2008; Rodriguez, et al., 2009)] hermaphroditic species, 154 including the domesticated tomato and its wild relatives (Peralta, et al., 2008). In this study our 155 focal species was Solanum pennellii, a wild, herbaceous, perennial species (Supplementary 156 Figure 1). S. pennellii populations-- including the two parental accessions (populations) used 157 here (see below)--can vary in the strength of UI against other SC Solanum species (Lied), 158 McCormick and Mutschler, 1996). We generated a recombinant F_2 population in which the 159 female parent was from self-compatible S. pennellii accession LA0716, and the male parent was 160 from self-incompatible S. pennellii accession LA3778. LA0716 does not express S-RNase, likely 161 due to a deletion in the underlying gene (Li and Chetelat, 2015), however both accessions 162 exhibit UI against other SC species, including domesticated tomato.

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164 Seeds of the parental accessions were obtained from the Tomato Genomics Resource 165 Center (TGRC; tgrc.ucdavis.edu), grown to maturity and one individual from each accession was 166 chosen to make the original cross. The F_1 offspring of this cross were self-compatible and one F_1 167 was selfed by hand-pollination to generate the F_2 generation (n = 100). To cultivate all 168 experimental plants, seeds were treated with 50% bleach for 30 minutes, rinsed, placed on 169 moist blotting paper and incubated (12hr day-length, 24°C) to stimulate germination. 170 Germinated seedlings were transplanted into flats with Metro Mix 360 (Sun Gro) potting mix 171 and hand watered daily. Once well-established, the seedlings were transferred to individual 1-172 gallon pots containing 50% Metro Mix 360 and 50% Indiana University (IU) greenhouse potting 173 mix; pots were placed in a climate controlled greenhouse at IU with 14hr day-length. Plants 174 were watered twice daily, fertilized weekly, staked prior to flowering, and regularly pruned 175 thereafter. 176

177 Quantifying unilateral incompatibility

178 To assess the quantitative expression of interspecific UI, each F_2 was pollinated with the 179 same tester genotype of S. lycopersicum (accession LA3475, SC). While both SC LA0716 and SI 180 LA3778 reject S. lycopersicum pollen, the former has a slower UI response (i.e. the pollen is 181 halted after growing further down the style; *Results*). At least 3 flowers from each F_2 individual 182 were emasculated one day prior to anthesis, the styles pollinated 24hrs later, and collected 183 after an additional 24hrs, which is sufficient time for compatible (i.e., conspecific) pollen tubes 184 to reach the ovary in the parental genotypes. Styles were fixed in 3:1 ethanol:glacial acetic acid, 185 stained using aniline blue fluorochrome (Biosupplies Australia Pty Ltd) and imaged using 186 fluorescent microscopy. Because styles were too long to be captured in one image frame, 187 several images were taken along the axis of each style and then stitched (Autostitch; (Brown 188 and Lowe, 2007)). Stitched images were visualized for data collection using ImageJ (Abramoff, 189 Magalhaes and Ram, 2004). UI response (location of pollen rejection within the style) was 190 quantified by measuring the total style length, length of the five longest pollen tubes, and 191 length of the pollen tube "front" where the majority of pollen tubes stopped growing. In all 192 analyses reported here, UI response was calculated by dividing the average of the five longest 193 pollen tubes by the total length of the style. Thus, mean pollen tube growth is quantified as a 194 proportion of style length travelled and varies from 0 (representing no growth down the style) 195 to 1 (where pollen tubes reach the end of the style). UI was similarly quantified in the parents 196 and F₁ as described above.

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198 Evaluating self-incompatibility status

199 In gametophytic SI, complete pollen rejection only occurs if both S-haplotypes in the 200 pistil match both S-haplotypes in the pollen parent. Because our F₂ population was generated 201 by self-fertilization of one F₁ individual, a simple SI/SC nomenclature cannot be applied to this 202 population. Instead, we expected that F_{2s} would display 'acceptor' phenotypes, as they would 203 have at most one functional S-haplotype from the original SI parent. For example, here, if the 204 LA3778 parent is designated S_1S_2 and the LA0716 parent S_0S_0 , their F_1 could be S_1S_0 or S_2S_0 . During self-fertilization of a particular F_1 individual (e.g., S_1S_0), pollen bearing the LA3778 (SI 205 206 parent) S_1 -haplotype will be selectively arrested, leaving only pollen with LA0716 (SC parent) S_0 -207 haplotype to fertilize the F₁ ovules; the resulting F₂ individuals are therefore expected to be 208 either S_1S_0 or S_0S_0 , and none should completely reject pollen from either SI or SC parent (i.e. 209 they are 'acceptors'). To confirm this was the case, we evaluated the pollen-rejection status of individuals in several ways. To initially test self-fertility status, at least three flowers from each 210 211 F_2 individual were manually self-pollinated. Selfed F_2 s that produced fruits were designated as 212 acceptor phenotypes. Fruits from these pollinations were left to mature on the plant; at 213 maturity each was weighed and measured (length and width), and seeds extracted by hand to 214 count viable seeds per fruit. In the rare cases (i.e., three individuals) where no fruits were 215 produced from these initial hand-pollinations, individuals were provisionally designated as self-216 sterile. These individuals were further evaluated using pollen from the original SI LA3778 parent 217 used to generate the F2, first by evaluating fruit set following pollination and then by directly 218 assessing pollen rejection by visualizing pollen tube growth in styles. The latter experiments 219 were performed as for UI (see 'Quantifying unilateral incompatibility' above), except that the 220 tester pollen came from the LA3778 parent. The F_1 was evaluated for self-fertility/acceptor 221 status in the same manner.

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223 Floral morphology and fertility traits

224 To quantify additional reproductive traits that might be relevant to mating system 225 variation or to the expression of UI, six floral and six fertility traits were measured. The six floral 226 traits were: corolla diameter, style length, stigma exsertion (distance between stigmatic surface 227 to the tip of the anther cone, on an intact flower), anther length, ovary height, and ovary width 228 [(Moyle, 2007); **Supplementary Figure 1**]. Using digital calipers, three fully open flowers (day 1 229 of opening) per F_2 individual were measured for all floral traits, and replicate measures 230 averaged within each individual prior to analyses. For each parent individual and the F_1 , five 231 replicate flowers were similarly measured. For overall comparison of the two parental 232 accessions, floral traits were also quantified on five additional individuals from each accession, 233 by taking the average of measurements from three flowers per individual.

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235 For fertility traits, we quantified total pollen, proportion viable pollen, fruit weight, fruit 236 width, fruit length, and seed set. Pollen number per flower was estimated by collecting whole 237 anther cones from individual flowers one day before opening, into lactophenol aniline blue 238 histochemical stain (Kearns and Inouye, 1993; Moyle and Graham, 2005). Each anther cone was 239 homogenized and for each sample an aliquot of homogenate was examined on an inverted 240 microscope using a hemacytometer, to count total pollen grains and estimate proportion of 241 viable and inviable pollen. Pollen that fails to stain lacks functional cytoplasm and was classified 242 as inviable. At least three anther cones were collected and counted per individual; mean counts 243 for each individual were used for analysis. At least three selfed fruits per individual (where 244 possible) were hand harvested, individually weighed, and bisected to take length and width 245 measurements (see 'Self-incompatibility status' section above). All seeds were extracted and 246 the number of viable seeds was counted per fruit. At least three fruit were measured per 247 individual and trait means for each individual were used for analysis.

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249 Style Protein Expression

250 Two loci (S-RNase and HT) were directly investigated for their association with UI phenotypic variation, by assessing their protein expression in parental, F1, and F2 styles. Both 251 252 S-RNase protein and HT (which is a small asparagine-rich protein (Covey, et al., 2010; McClure, 253 et al., 1999; O'Brien M, 2002)) have been previously implicated in UI expression (Murfett et al., 254 1996, Tovar-Mendez et al., 2014)-including in QTL mapping studies in other Solanum species 255 (Bernacci and Tanksley, 1997)—making them a priori candidates for UI variation in this 256 population. Both genes are also essential for SI (McClure, et al., 1999). The HT gene was 257 duplicated in the ancestor of Solanum, giving rise to two tandemly arrayed genes 258 (Sopen12g029190, HT-A and Sopen12g029200, HT-B) on chromosome 12 (Covey et al. 2010). A 259 subset of individuals (F₁ n = 1; F₂ n = 21) was screened for protein expression of S-RNase and HT 260 using protein blotting. For each individual, flowers were emasculated 24hrs before opening; 261 styles were collected 24hrs later and weighed. At least 5mg stylar tissue was collected per 262 individual and protein was extracted using 2xLSB (Laemmli Sample Buffer; 10uL LSB / 1mg 263 tissue). Samples were boiled 5min, centrifuged (10min at 20,000 x g) and the supernatant was 264 retained for analysis. For S-RNase detection, extract equivalent to 0.2mg fresh weight per lane 265 was separated in 10% Tris-Tricine SDS PAGE, blotted to PVDF, and immunostained (1:5000) with

an antibody against the conserved C2 S-RNase motif, as described previously (Covey, *et al.*,
2010). For HT-protein detection, extracts equivalent to 1.5mg fresh weight were separated in
12.5% Tris-Tricine SDS PAGE, blotted to PVDF, and immunostained (1:5000) with an affinitypurified antibody that recognizes both HT-A and HT-B proteins. The antibody was prepared

- against the synthetic peptide LEANEIHNTELNNPTLQKKGGC-amide (21st Century Biochemicals),
- as described previously (Tovar-Méndez, *et al.*, 2017).
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273 Genotyping F₂s

274 To genotype our F_2 population, genomic DNA from 93 F_2 s and each parent was 275 extracted using Qiagen DNeasy Plant Mini Kits. Extracted genomic DNA samples were sent to 276 the Cornell University Institute of Biotechnology's Genomic Diversity Facility for genotyping-by-277 sequencing (GBS), using restriction enzyme Pstl. An unfiltered SNP marker set was generated by 278 the Cornell Institute of Biotechnology, by mapping trimmed raw sequence reads onto the S. 279 pennellii (LA0716) genome (Bolger, et al., 2014) using bwa (Li 2013), within the Cornell TASSEL 280 3.0 GBS reference pipeline (version 3.0.173). To obtain a high-quality set of markers for the 281 linkage map and QTL mapping, only markers with bi-allelic sites and that had fixed differences 282 between parental genotypes were used. For consistency, we required that a maximum of 30% 283 individuals differ in genotypes for any pair of markers that are within 500bp of each other, for 284 these markers to be retained. Segregation distortion was assessed by testing for Hardy-285 Weinberg equilibrium at each marker, and markers showing significant deviation (p < 0.05) 286 were removed from the final marker dataset. After implementing these filters and also 287 removing samples with low sequencing quality (those that had more than 15% missing 288 genotypes), the initial genetic map contained 810 markers and was significantly expanded 289 (average LG length of 332 cM) likely due to unaccounted for genotyping errors. To 290 accommodate this, we used the Genotype-Corrector tool (Miao et al., 2018) which corrects or 291 imputes genotype calls at reference-mapped markers based on a sliding-window algorithm, 292 prior to rebuilding the linkage map. The resulting dataset contained 569 high-quality markers 293 from 88 individuals; five additional individuals were removed due to high levels of missing data 294 (>15%) following correction or removal of unlikely genotype calls.

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296 Linkage Map and QTL Mapping

The linkage map was constructed using the Rqtl (Broman, *et al.*, 2003) and ASMap (Taylor & Butler, 2017) packages in R version 3.2.2 (R Core Team, 2015); ASMap implements the minimum spanning tree (MST) algorithm (Wu et al., 2008) for map construction. Markers were first clustered by chromosome (based on reference mapping) prior to inferring the marker order on each group using the MST algorithm. The final map length was 1750.48cM (average of 145.87 cM per LG), with an average of 0.276 cM between markers.

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For phenotypes that were non-normally distributed (Shapiro-Wilk test; W < 0.05), we
 transformed the trait data using the nqrank function which transforms the vector of
 quantitative values to corresponding normal quantiles and preserves the mean and standard
 deviation. Missing genotypes were imputed prior to performing genome scans with the
 multiple QTL model [MQM; (Arends, *et al.*, 2010)] for each trait. Genome-wide significance LOD
 thresholds were calculated for each trait based on permutation tests (1000 iterations) with

310 alpha = 0.05. For each trait, we included the significant QTL in a model as the main effect to 311 obtain estimates of the total phenotypic variation explained and individual contributions of 312 each QTL, as well as to test for interactions between QTL. The percent parental difference 313 explained (relative homozygous effect, RHE) was calculated for each detected QTL as the 314 additive QTL effect size divided by the parental difference. We assessed overlap of identified 315 QTL with previously identified UI QTL (Bernacci and Tanksley, 1997) by using information on 316 physical location of markers, the annotated S. lycopersicum and S. pennellii genomes, and other 317 gene position data from the Sol Genomics Network (solgenomics.net). Finally, we quantified the 318 number of loci that fell within the 1.5-LOD confidence interval (CI) of our UI QTL (see Results) by 319 identifying the two markers closest to each end of the CI and then counting all annotated genes 320 that fell between these two markers, using version 2 of the AUGUSTUS annotation of the S. 321 pennellii genome.

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323 Statistical Analyses

All analyses were run in R version 3.2.2 (R Core Team, 2015) and statistical significance was reported if p < 0.05. Shapiro-Wilks tests were performed to test for normality for each quantitative trait. T-tests were used to test for trait variation between the parental accessions, and to compare UI responses between individuals found to express S-RNase protein or not.

329 Results

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331 Rapidity of unilateral incompatibility rejection response varies among F₂s

332 The two parental accessions used to make our F_2 population both exhibited UI but 333 differed quantitatively in how rapidly they rejected *S. lycopersicum* pollen, i.e. in the position 334 within the style that their UI response manifested. The SI LA3778 parent had a rapid UI 335 response to heterospecific pollen tubes $(0.038 \pm 0.005 \text{ proportion of style length};$ Figure 1), 336 where pollen rejection was defined in terms of the proportional distance of pollen tube growth 337 down the style (so that values closer to zero indicate a rapid UI response whereas slower 338 responses have values closer to 1). The SC LA0716 parent had a less rapid response (0.32 ± 0.16 339 proportion of style length). The F_1 individual also expressed a rapid UI response (0.040 ± 0.034 340 proportion of style length). All measured (n=99) F_2 individuals expressed UI. Nonetheless, there 341 was broad quantitative variation in the rapidity of the UI rejection response, with the tester 342 pollen tube growth response ranging from 0.01 - 0.55 of the length of F₂ styles (Figure 1).

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344 All F_2 s accepted self-pollen

345 Because the F_2 population was generated by self-fertilization of one F_1 individual, we expected that all F₂s would accept self-pollen. This is because only F₁ pollen bearing the LA0716 346 347 (SC parent) S-haplotype would have fertilized F₁ ovules (due to pistil-mediated gametophytic 348 selection against the alternative, functional, pollen-side allele from the LA3778 SI parent) (see 349 Methods). We confirmed that all evaluated F_2 individuals (n = 94) accepted self-pollen 350 ('acceptors' in **Figure 1**). In the few individuals (n = 3) that did not successfully develop fruits 351 after self-fertilization, when pollen rejection was directly assessed in their styles, they showed 352 no evidence of a pollen rejection response; that is, pollen tubes were observed growing all the 353 way to the ovary. Therefore we infer that other downstream factors (e.g. low ovule fertility,

gamete isolation, or early fruit abortion) may have prevented fruit development in these threeindividuals.

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Large effect unilateral incompatibility QTL is associated with the S-locus and variation in
 functional S-RNase

359 One large effect QTL was identified for UI, explaining 32.6% of the phenotypic variance among F2s, and 23.0% of the difference observed between the parents (Table 2, Figure 2). 360 361 Located on chromosome 1, this QTL colocalizes both with the location of a UI QTL previously 362 identified in a different Solanum cross (Bernacci and Tanksley, 1997) and with the genomic 363 location of the S-locus (Figure 2), which contains genes encoding S-RNase, F-box proteins, and 364 other factors involved in self-incompatibility (Bernacci and Tanksley, 1997; Li and Chetelat, 365 2015). The 1.5 LOD CI of this QTL spans 29.47cM or ~85.08Mb, and contains 2684 gene models, 366 based on the v2 annotation of the S. pennellii genome. This large low-recombination region is 367 characteristic of the physical location of the S-locus (Figure 2), which exhibits suppressed 368 recombination in this and other species.

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370 Because S-RNase has been hypothesized, and shown, to be a significant contributor to 371 the UI response in some previous studies in Solanum (Chalivendra, et al., 2013; Covey, et al., 372 2010; Tovar-Méndez, et al., 2014), we assessed a subset of F₂ individuals (n = 22) for expression 373 of this protein in mature unpollinated styles. This subset represented individuals with the most 374 and least rapid UI responses in our F₂ population. We found that individuals that expressed S-375 RNase protein had a significantly more rapid UI response (n = 9, mean = 0.0815 of style) against 376 S. lycopersicum pollen, compared to those that did not express S-RNase (n = 13, mean = 0.227 377 of style) (t(19.885) = 3.374, p = 0.003; Figure 3), indicating that S-RNase protein 378 presence/absence is a major contributor to observed quantitative variation in UI. In comparison, 379 all individuals were found to express HT-protein in their styles; therefore the presence/absence 380 of HT-protein is not implicated in the phenotypic variation in UI segregating in our F_2 population. 381 (Note that HT expression may still be required for the expression of UI; see Discussion.) The F_1 382 expressed both HT and S-RNase and had a rapid UI response.

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Previous work in a different *Solanum* species cross also identified UI QTL on chromosomes 3 and 12 (Bernacci and Tanksley, 1997) as did a recent analysis of UI between *S. pennellii* (LA3778) and *S. lycopersicum* (LA3475) (Jewell 2016). We did not detect QTL at either of these positions. As HT-protein is thought to likely underlie the chromosome 12 UI QTL detected in other studies (Bernacchi and Tanksley, 1997; Tovar- Méndez, *et al.*, 2014; Jewell 2016) this finding is consistent with our observation (above) of no differential protein expression of *HT* in our F₂ population.

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392 Floral traits vary with mating system differences between the parental genotypes

The parent accessions differed in four of the six floral traits measured: stigma exsertion,
 style length, and ovary height and width (Table 1). There were no significant differences
 between the parent populations for corolla diameter or anther length (Table 1, Supplementary
 Figure 1). Despite large parental differences, we identified only one QTL affecting floral
 morphology (Table 2). This locus, on chromosome four, had a moderate to large effect on

stigma exsertion (percent variance explained = 17.1; percent parental difference explained =
 48.0). No significant QTL were detected for corolla diameter, anther length, style length, ovary
 height, or ovary width. Because of the limited size of the mapping population, our analyses

401 likely missed smaller effect loci that contribute to observed parental variation in these floral402 traits.

403

404 Few fertility QTL detected

We identified two QTL for fertility traits, both of which were for fruit height (**Table 2**, **Figure 2**) and were of small to moderate effect, explaining ~6-16% of the variation among F2s and 5% of the parental difference each. There was no evidence for an interaction between these QTL. Interestingly, these two QTL have opposing effects on fruit height, consistent with little difference between the parental accessions in this fruit trait. Neither of the detected fertility trait QTL colocalized with our UI QTL on chromosome 1.

- 411
- 412 Discussion
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414 Standing genetic variation across populations within species can contribute to 415 differences in the strength of interspecific isolating barriers. Understanding the nature of this 416 genetic variation can provide insight into the evolutionary dynamics that shape the 417 accumulation of these barriers among diverging species. Here we assessed the genetic basis of 418 intraspecific variation in the strength of interspecific pistil-side unilateral incompatibility (UI). 419 One goal was to assess whether variation in key components of mating system variation 420 (including genetic self-incompatibility) influences this phenotypic variation. We found one large 421 effect QTL underlying variation among populations in the rate at which UI is expressed against a 422 second species. This QTL overlaps a major player in the self-incompatibility response--the S-423 locus--and we found that the presence/absence of stylar S-RNase protein is significantly 424 associated with the rapidity of the UI response. Although we observe trait and genetic 425 differences in floral and fertility traits between these two S. pennellii populations, some of 426 which are typically associated with mating system transitions, QTL underlying these differences 427 are not associated with the major effect locus controlling variation in the strength of UI. Our 428 analysis suggests that standing variation for mate choice loci--in this case, to prevent self-429 fertilization--can directly affect variation in interspecific isolation--in this case a postmating 430 prezygotic reproductive barrier.

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UI genetic mechanisms are associated with mating system loci

433 Both our QTL mapping analysis and our protein expression assay support the inference 434 that S-RNase protein plays a major causal role in the observed quantitative variation in UI 435 responses. In Solanaceae, loss of SI often involves the loss of pistil S-RNase expression as one of 436 the first causal changes, so that individuals no longer reject conspecific pollen with which they 437 share the functional pollen-side component of the SI mechanism. Our analysis indicates that 438 this loss of pistil S-RNase protein in the SC S. pennellii accession (LA0716) has pleiotropic 439 consequences for the rate at which this genotype rejects heterospecific pollen. While all 440 individuals expressed UI within our F_2 population, the speed of UI pollen rejection was significantly decreased when S-RNase protein was absent. Thus, we infer that standing variation 441

within *S. pennellii* at a major mating system locus also directly contributes to how rapidly aninterspecific postmating prezygotic barrier is expressed.

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445 In addition, our data also imply that other molecular factors beyond S-RNase also 446 contribute to UI expression in this species. That is, while loss of functional S-RNase protein 447 reduced the speed of UI it did not abolish this response, suggesting other functional UI 448 elements are retained in the pistil. S-RNase-independent UI mechanisms have been described 449 previously in Solanum (Murfett, et al., 1996; Eberle, et al., 2013; Tovar-Méndez, et al., 2017; 450 reviewed in Bedinger et al., 2017). In particular, HT protein has been implicated as a necessary 451 molecular component of the UI response in Nicotiana and in other Solanum crosses (Bernacci 452 and Tanksley, 1997; Covey, et al., 2010; Hancock, Kent and McClure, 2005; McClure, et al., 1999; 453 O'Brien, 2002; Tovar-Méndez, et al., 2017). In addition, both SC and SI S. pennellii populations 454 have previously been shown to express HT in their styles (Chalivendra, et al., 2013, Pease et al. 455 2016), and here we found that all tested F_2 individuals also expressed HT. Together, these 456 observations suggest that the observed quantitative variation in UI expression depends on 457 variation in the functionality of S-RNase, on a background of functional HT expression. In 458 addition to HT, there might also be other S-RNase-independent factors supporting UI function 459 in these accessions. For example, other QTL studies have identified a major effect UI QTL on 460 chromosome 3 in populations generated between S. lycopersicum and SI S. habrochaites 461 (Bernacci and Tanksley, 1997) and between S. lycopersicum and the SI S. pennelii parent 462 genotype used in our cross here (Jewell 2016, Hamlin et al. 2018). Furthermore, loss-of function 463 of HT in SC S. pennellii accession (LA0716) resulted in tomato pollen rejection further down the 464 pistil, suggesting that in addition to the S-RNAse-independent UI factors there are also HT-465 independent UI factors (Tovar-Méndez, et al., 2017). The potential contribution of additional 466 factors to S. pennellii intraspecific variation in UI is testable in the future, once the specific 467 identity of these factors is known.

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469 Regardless, our observations support a mechanistic, explicitly genetic, association 470 between SI and UI, consistent with other studies in closely related species (Broz et al. 2017, 471 Tovar-Méndez, et al., 2014; Tovar-Méndez, et al., 2017). These findings in turn imply that 472 factors governing the maintenance or loss of SI can have collateral effects on the expression of 473 UI barriers among species. Ecologically, the loss of SI is often associated with strong selection 474 for reproductive assurance in low density or marginal environmental conditions, where small 475 population sizes severely restrict the availability of mating partners with different S-haplotypes. 476 Under these conditions, individuals that are physiologically capable of selfing-due to loss of 477 proteins involved in self-recognition and rejection—have a demographic and mating advantage. 478 If SI and UI share underlying genetic mechanisms, these specific ecological conditions will also 479 strongly determine the emergence and distribution of UI mating barriers between lineages, as 480 will the physiological constraints governing the genetic progression of SI to SC transitions, as we 481 discuss further below. 482

483 No evidence for an association between UI and floral trait loci across mating transitions
 484 Another possibility we examined was whether the strength or expression of UI was
 485 associated with other, non-SI, transitions that accompany mating system differences. Although

486 losing SI permits selfing, the transition from facultative to predominant self-fertilization often 487 involves additional morphological changes, especially in floral traits that affect pollinator 488 attraction and the likelihood of self-pollination. While outcrossing species typically have larger 489 flowers and greater distances between the receptive stigmatic surface of the female pistil and 490 the male pollen-bearing anthers (i.e. greater stigma exsertion) (Brunet and Eckert, 1998; 491 Motten and Stone, 2000; Rick, Holle and Thorp, 1978; Takebayashi, Wolf and Delph, 2005), 492 highly self-pollinating species tend to have smaller flowers and smaller or no stigma exsertion 493 (Lloyd and Barrett, 1996; Sicard and Lenhard, 2011). It is unclear to what extent changes in loci 494 directly involved in the breakdown of SI (i.e. S-locus and its modifiers) work in conjunction with 495 the genes controlling these morphological changes, as well as whether these morphological loci 496 are associated with the expression of UI. Here, we found some floral morphology differences 497 between the LA3778 and LA0716 parent populations that are typical of mating system 498 differences, specifically greater stigma exsertion and greater style length in the SI compared to 499 the SC parent genotype (Table 1), and we identified one QTL for stigma exsertion (Table 2). 500 However, this QTL did not coincide with the UI QTL (or with known loci involved in SI). These 501 findings provide little evidence for strong phenotypic or genetic associations between the loss 502 of gametophytic SI and/or the expression of UI, and the morphological shifts that typically 503 accompany mating system transitions.

505 The progression to UI among species, and the role of mating system transitions

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506 Finally, our findings also contribute to an emerging picture of the evolution of UI 507 between species, and the specific role of mating system transitions in the formation of this 508 post-mating prezygotic isolation barrier. First, in conjunction with mapping studies (Bernacchi 509 and Tanksley 1997, Jewell 2016) and crossing analyses (e.g. Broz et al. 2017; see further below) 510 in other closely-related Solanum species, we infer that quantitative transitions from UI 511 competence to its loss are often associated with the cumulative loss of one or more loci 512 functionally involved in self-incompatibility. For example, S. habrochaites (the sister species to S. 513 pennellii) is generally an SI species within which some populations have transitioned to SC 514 (Martin 1961), often (but not always) via the loss of functional S-RNase genes (Broz et al. 2016, 515 Markova et al. 2016). Unlike the two populations of S. pennellii examined here, several of these 516 S. habrochaites SC populations have also lost the ability to reject certain heterospecific pollen— 517 consistent with the loss of UI competence (Baek, et al., 2015; Covey, et al., 2010, Broz et al. 518 2016) via the loss of one or more additional pistil-side UI factors. This greater loss of UI 519 competence, which may be related to a longer history of self-compatibility in these S. 520 habrochaites populations, appears to involve at least one other S-RNase-independent molecular player apart from HT, as all but one of these populations continue to express HT 521 522 (Baek, et al., 2015; Covey, et al., 2010, Broz et al. 2017). Nonetheless, HT can clearly also 523 contribute to this transition from UI to non-UI styles, as Tovar-Méndez, et al., (2017) showed 524 that suppressing HT expression in S. habrochaites LA0407 completely abolished UI against (SC) S. 525 lycopersicum pollen. Together, these observations suggest that populations undergoing a 526 progressive loss of pistil-factors can proceed stepwise from rapid UI against other SC species (coincident with a fully functional SI system), through a transitional period of quantitative 527 528 reductions in the strength of UI (coincident with the loss of pistil-side factors during the

transition from SI to SC), to loss of UI against other species (coincident with the loss of both S-RNase-dependent and -independent rejection mechanisms).

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532 Notably, while the progressive loss of pistil-side factors ultimately results in genotypes 533 unable to mount a pistil-side UI rejection response, observations also indicate that transitions 534 from SI to SC can be accompanied by the gain of UI, specifically the gained ability of SI lineages 535 to reject the pollen of SC lineages (e.g. Martin 1961, Broz et al. 2016, Marcova et al. 2016). 536 However this emergence of UI against SC lineages is not due to pistil-side changes, but instead 537 to a progressive loss within these recently derived SC lineages of pollen-side function(s) that 538 would otherwise neutralize pistil rejection responses (Bedinger et al. 2017). For example, within 539 S. habrochaites, pollen from some SC populations is rejected by SI populations of the same 540 species (Broz et al., 2017), likely due to the loss of one or more pollen-side factors specifically in 541 these SC populations (Markova et al. 2016). In contrast, SI populations do not reject pollen from 542 SI populations or species, indicating that SI pollen retain mechanisms for evading UI rejection in 543 styles of other SI species. Note that the S. pennellii populations used here reciprocally accept 544 each other's pollen, indicating that the SC S. pennellii accession LA0716 has not progressed to 545 the point at which UI barriers have emerged against it.

547 This observed progression indicates a specific temporal order to the loss of pistil-side UI 548 and the gain of pollen-side UI rejection by other lineages, a trajectory that is strongly influenced 549 by the dynamics governing the loss of intraspecific SI factors during transitions from SI to SC. 550 Interestingly, in the Solanaceae (but not in other plant groups that have SI systems; Bedinger et 551 al. 2017) this transition from SI to SC usually first involves the loss of loci that contribute to 552 pistil-side function, and only subsequently the loss of pollen-side functions (Tovar-Méndez, et 553 al., 2014). This 'pistil-first' transition order likely occurs because the physiology of gametophytic 554 self-incompatibility: pollen loss-of-function mutations are incompatible on all pistils that retain 555 pistil-side function, but genotypes with pistil-side SC mutations are able to accept all pollen 556 donors (see also Markova et al. 2016). This leads to strong selection against pollen-side 557 mutations because these cannot individually permit self-compatibility, and therefore will not 558 contribute to reproductive assurance unless they are first preceded by pistil-side mutations. 559 This expectation is supported by the observation that there are no known SC populations or 560 species that lack SI pollen function but retain pistil function, in the Solanaceae (Tovar-Méndez, 561 et al., 2014). In this way, loss of S-RNase and other pistil-side factors does not contribute 562 immediately to the gain of UI, but acts as a catalyst for evolutionary changes that eventually 563 lead to the erection of an UI barrier against the evolving population, by permitting the 564 subsequent loss of pollen side factors. Although the conditions promoting this subsequent loss 565 of pollen-side factors are less clear, it is possible that these mutations reduce metabolic cost 566 (Markova et al. 2016) or increase selfing efficiency once populations have already lost pistil-side 567 SI functions.

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569 Regardless, it is clear that the dynamics of these mating system transitions play an 570 influential role in the evolution of UI as a reproductive barrier. Moreover, understanding the 571 nature of intraspecific genetic variation involved in these transitions is critical for understanding 572 the conditions that facilitate the accumulation of reproductive isolation among populations 573 within species (Good, et al., 2007; Kopp and Frank, 2005). Here, in our analysis of genetic

- 574 variation within *S. pennellii*, we have shown that one of the earliest steps in this progression
- 575 involves the large quantitative contribution of a pistil-side locus that is directly involved in
- 576 intraspecific mate choice (via self-incompatibility). This finding agrees with previous analyses
- 577 that indicate an intimate association between molecular players contributing to SI and UI. In
- 578 combination with genetic and crossing data from this and other closely related species, it also
- 579 suggests that intraspecific changes at these pistil-side loci are an essential antecedent step that
- 580 permits the subsequent accumulation of mutations that erect new intraspecific postmating
- 581 prezygotic UI barriers.582

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- 592

593 Data Availability

- 594 The primary data underlying these analyses will be deposited as follows:
- 595 morphological data: Dryad #######
- 596 raw sequence reads and SNP genotypes: NCBI BioProject: PRJNA557135.
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807 Table and Figure captions

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Table 1. Trait differences between individuals (n = 5) of the parental accessions, reported as
 means and standard deviations. T-tests (one-sided) are reported for differences between the
 parental accessions. Comparisons with p < 0.05 are denoted in bold. F₁ hybrid values are also

- 812 reported; NA = not evaluated.
- 813

Table 2. QTL associated with unilateral incompatibility, floral traits, and fertility traits. Percent
 phenotypic variance (PVE) and percent parental difference (RHE) explained are reported for

- 816 each trait, with full models including all QTL found for each phenotype.
- 817

818 **Figure 1.** Variation in the strength of unilateral incompatibility (UI) responses between *S*.

- 819 *pennellii* populations. **A.** Representative image of an F₂ with a rapid UI response. **B.** Image of an
- F_2 with a less rapid UI response. **C.** Phenotypic distribution of unilateral incompatibility
- responses across the F₂ population. Blue dashed line (0.038) is the *S. pennellii* SI parent
- 822 (LA3778); orange dashed line (0.32) is the *S. pennellii* SC parent (LA0716).
- 823

Figure 2. Map of identified QTL. Each QTL is marked with the 1.5-LOD confidence interval, with
 the peak marker indicated by a horizontal line. *A priori* hypothesized loci are marked as ellipses.

826

827 **Figure 3**. Expression of S-RNase in F₂ mature styles as determined by a protein blot. There is

significantly more rapid UI response in those styles that express S-RNase compared to those
that do not (t= 3.374, p = 0.003).

Table 1. Trait differences between individuals (n = 5) of the parental accessions, reported as

833 means and standard deviations. T-tests (one-sided) are reported for differences between the

parental accessions. Comparisons with p < 0.05 are denoted in bold. F_1 hybrid values are also

835 reported; NA = not evaluated.

Trait	<i>S. pennellii</i> (LA0716, SC)	S. pennellii (LA3778, SI)	t-test	F1	
UI	0.322 ± 0.026	0.038 ± 2.66 x 10 ⁻⁵	t = -3.34; p = 0.022	0.039 ± 3.43 x 10 ⁻⁴	
Corolla Diameter (mm)	21.22 ± 1.88	20.73 ± 0.86	t = -0.54, p = 0.612	28.31 ± 1.31	
Anther Length (mm)	7.08 ± 0.85	7.59 ± 0.33	t = 1.26, p = 0.263	9.44 ± 0.55	
Stigma Exsertion (mm)	1.50 ± 0.28	2.32 ± 0.43	t = 3.57, p = 0.0096	2.32 ± 0.52	
Style Length (mm)	8.29 ± 1.17	10.56 ± 0.87	t = 3.48, p = 0.0095	12.23 ± 0.36	
Ovary Height (mm)	1.63 ± 0.12	1.11 ± 0.05	t = -8.87, p = 0.0003	1.32 ± 0.13	
Ovary Width (mm)	1.05 ± 0.04	0.92 ± 0.05	t = -4.68, p = 0.002	1.32 ± 0.05	
Total Pollen	69.53 ± 192.8	62.40 ± 731.3	t = -0.52; p = 0.3070	NA	
Proportion Viable Pollen	0.469 ± 0.017	0.834 ± 0.003	t = 5.70; p = 0.0002	NA	

Table 2. QTL associated with unilateral incompatibility, floral traits, and fertility traits. Percent

840 phenotypic variance (PVE) and percent parental difference (RHE) explained are reported for 841 each trait, with full models including all OTL found for each phenotype

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Trait	LOD Threshold alpha = 0.05	QTL peak chromosome, position (cM), closest marker	Peak LOD	1.5-LOD CI	PVE (%)	Additive effect (S.E.)	Dominance effect (S.E.)	% parental difference explained (RHE)
Unilateral Incompatibility	2.71	1, 60 chr1_79110602	5.49	40-64.67	32.63	-0.066 (0.020)	-0.062 (0.024)	23.0
Corolla Diameter	2.19	None						
Anther Length	2.85	None						
Stigma Exsertion	2.32	4, 180, chr4_75727601	2.65	75-181.6	17	0.366 (0.082)	0.092 (0.110)	48.0
Style Length	2.45	None						
Ovary Width	2.44	None						
Ovary Height	2.19	None						
Total Pollen	2.25	None						
Proportion Viable Pollen	2.12	None						
Fruit Width	2.80	None						
Fruit Height	2.46	8, 10, chr8_2708030	2.69	0-25	16.18	-0.537 (0.149)	0.832 (0.271)	5.0
		9, 90, chr9_5441571	2.68	62.19-100	6.17	0.535 (0.173)	-0.047 (0.248)	-5.0
Fruit Weight	2.87	None						
Seed Count	2.55	None						

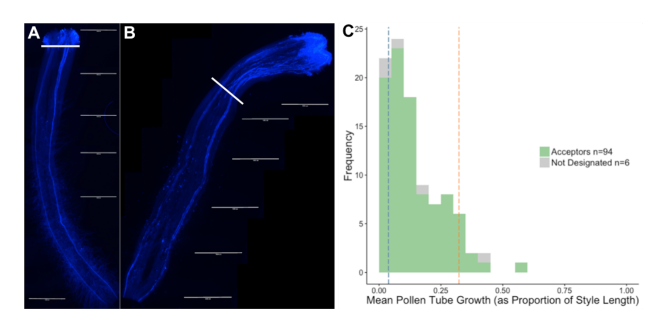


Figure 1. Variation in the strength of unilateral incompatibility (UI) responses between *S*.

pennellii populations. **A.** Representative image of an F₂ with a rapid UI response. **B.** Image of an

 $850 \qquad F_2 \text{ with a less rapid UI response. } \textbf{C.} \text{ Phenotypic distribution of unilateral incompatibility}$

responses across the F_2 population. Blue dashed line (0.038) is the *S. pennellii* SI parent

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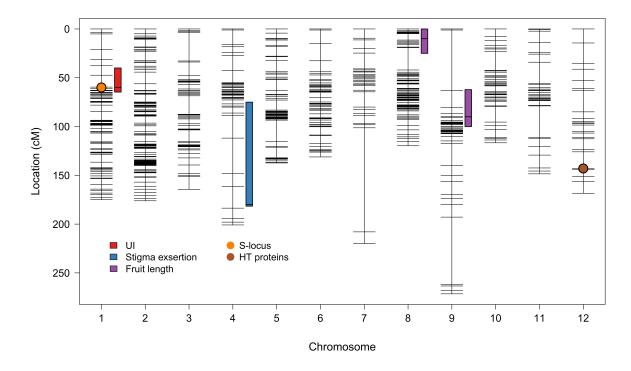
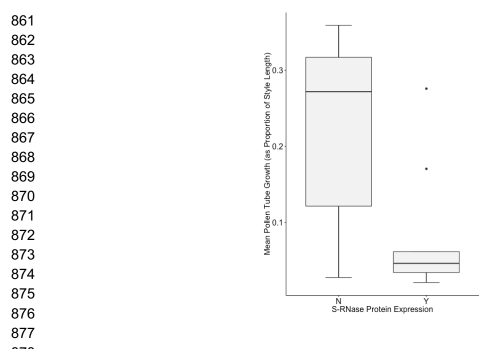


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Figure 3. Expression of S-RNase in F_2 mature styles as determined by a protein blot. There is significantly more rapid UI response in those styles that express S-RNase compared to those that do not (t= 3.374, p = 0.003).

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