

ORIGINAL ARTICLE

Interspecific crossing and genetic mapping reveal intrinsic genomic incompatibility between two *Senecio* species that form a hybrid zone on Mount Etna, Sicily

AC Brennan^{1,2,3}, SJ Hiscock⁴ and RJ Abbott¹

Studies of hybridizing species can reveal much about the genetic basis and maintenance of species divergence in the face of gene flow. Here we report a genetic segregation and linkage analysis conducted on F₂ progeny of a reciprocal cross between *Senecio aethnensis* and *S. chrysanthemifolius* that form a hybrid zone on Mount Etna, Sicily, aimed at determining the genetic basis of intrinsic hybrid barriers between them. Significant transmission ratio distortion (TRD) was detected at 34 (~27%) of 127 marker loci located in nine distinct clusters across seven of the ten linkage groups detected, indicating genomic incompatibility between the species. TRD at these loci could not be attributed entirely to post-zygotic selective loss of F₂ individuals that failed to germinate or flower (16.7%). At four loci tests indicated that pre-zygotic events, such as meiotic drive in F₁ parents or gametophytic selection, contributed to TRD. Additional tests revealed that cytonuclear incompatibility contributed to TRD at five loci, Bateson–Dobzhansky–Muller (BDM) incompatibilities involving epistatic interactions between loci contributed to TRD at four loci, and underdominance (heterozygote disadvantage) was a possible cause of TRD at one locus. Major chromosomal rearrangements were probably not a cause of interspecific incompatibility at the scale that could be examined with current map marker density. Intrinsic genomic incompatibility between *S. aethnensis* and *S. chrysanthemifolius* revealed by TRD across multiple genomic regions in early-generation hybrids is likely to impact the genetic structure of the natural hybrid zone on Mount Etna by limiting introgression and promoting divergence across the genome. *Heredity* (2014) 113, 195–204; doi:10.1038/hdy.2014.14; published online 5 March 2014

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INTRODUCTION

Although strong, divergent natural selection can maintain population divergence in the face of gene flow (Nosil, 2012), intrinsic genetic barriers can also evolve under such conditions (Rundle and Nosil, 2005; Agrawal *et al.*, 2011) or may already be in place to varying degrees between hybridizing populations that diverged in allopatry (Coyne and Orr, 2004; Bierne *et al.*, 2011; Feder *et al.*, 2012; Abbott *et al.*, 2013). Intrinsic incompatibility between species is generally detected by crossing studies, which can also reveal the nature of such incompatibility at both genetic and genomic levels. For example, transmission ratio distortion (TRD), which is often observed at segregating loci among progeny of interspecific crosses or crosses between divergent lineages within species, can result from selection against particular hybrid genotype combinations at these loci (TRDLs; Fishman *et al.*, 2001; Moyle and Graham, 2006). TRD may result from Bateson–Dobzhansky–Muller (BDM) incompatibilities at haploid and/or diploid stages of the life cycle caused by negative epistatic interactions between nuclear loci showing polymorphisms within and between populations of the same or different species (Fishman *et al.*, 2008; Cutter, 2012; Bomblies, 2013; Corbett-Detig *et al.*, 2013; Ouyang and Zhang, 2013). It may also arise from

cytonuclear incompatibility (Levin, 2003; Fishman and Willis, 2006; Turelli and Moyle, 2007) or from 'selfish' meiotic drive of alleles in a new genomic background (Fishman and Willis, 2005). Incompatibility caused by cytonuclear or haploid–diploid incompatibilities may depend on cross direction leading to asymmetric incompatibility, which can influence patterns of introgression following hybridization (Fishman *et al.*, 2001; Turelli and Moyle, 2007; Tang *et al.*, 2010). Genetic incompatibilities of all kinds accumulate with increasing phylogenetic distance until complete reproductive isolation is evident (Matute *et al.*, 2010; Moyle and Nakazato, 2010; Levin, 2012; Corbett-Detig *et al.*, 2013).

Interspecific crossing, when used in linkage analysis and mapping, may also reveal chromosomal rearrangements between species that can reduce the fitness of hybrids and suppress recombination that, in turn, will affect rates of interspecific gene flow (Rieseberg, 2001; Ortiz-Barrientos *et al.*, 2002; Kirkpatrick and Barton, 2006). Studies of hybridizing annual sunflowers have revealed that differences in genetic architecture due to chromosomal inversions and translocations are an important cause of hybrid sterility (Lai *et al.*, 2005; Yatabe *et al.*, 2007), whereas in other groups of hybridizing species such as irises (Taylor *et al.*, 2012) and sculpin fish (Stemshorn *et al.*, 2011)

¹School of Biology, Harold Mitchell Building, University of St Andrews, St Andrews, UK; ²Estación Biológica de Doñana (EBD-CSIC), Avenida Américo Vespucio s/n, Sevilla, Spain;

³School of Biological and Biomedical Sciences, University of Durham, Durham, UK and ⁴School of Biological Sciences, University of Bristol, Bristol, UK

Correspondence: Dr AC Brennan, School of Biological and Biomedical Sciences, University of Durham, South Road, Durham DH1 3LE, UK.

E-mail: a.c.brennan@durham.ac.uk

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such rearrangements appear to be absent or minor. In contrast, the presence of TRDLs in genetic maps of interspecific crosses appears to be common, if not the rule (Fishman *et al.*, 2001; Lu *et al.*, 2002; Tang *et al.*, 2010). Either way, the cumulative action of multiple intrinsic incompatibilities with diverse modes of action distributed across the genome will act as a potent barrier to gene flow between species and seems to be a common, if not universal, intermediate stage in the process of speciation (Bierne *et al.*, 2011; Feder *et al.*, 2012; Abbott *et al.*, 2013; Bomblies, 2013).

In the present study, we utilize interspecific crossing and linkage analysis to investigate the genetic nature of intrinsic incompatibility between two diploid, ragwort species, *Senecio aethnensis* ($2n=20$) and *S. chrysanthemifolius* ($2n=20$) (Asteraceae), that form a hybrid zone on Mount Etna, Sicily (James and Abbott, 2005; Brennan *et al.*, 2009). These two species are self-incompatible, short-lived, herbaceous perennials, which grow at high and low altitudes, respectively, on Mount Etna. They are connected by a series of hybrid populations, which potentially provides a corridor for high levels of interspecific gene flow to occur. Interestingly, material collected from this hybrid zone and introduced to Britain in the late 17th century subsequently gave rise to a highly invasive homoploid hybrid species, *S. squalidus*, which spread through much of Britain in the 19th and 20th centuries (James and Abbott, 2005; Abbott *et al.*, 2009).

Previous analyses of the hybrid zone on Mount Etna showed that while extrinsic environmental selection is important in determining the ecological differences and relative distributions of the two *Senecio* species (Brennan *et al.*, 2009; Ross *et al.*, 2012; Chapman *et al.*, 2013; Muir *et al.*, 2013; Osborne *et al.*, 2013), intrinsic selection against hybrids was predominant in determining changes in allele frequencies and quantitative trait expression in the hybrid zone (Brennan *et al.*, 2009). Indeed, based on indirect measures of assessment, Brennan *et al.* (2009) showed that the hybrid zone was characterized by strong selection against hybrids, high dispersal rates and few loci differentiating quantitative traits. The evidence for strong selection against hybrids was somewhat surprising, as both Hegarty *et al.* (2009) and Brennan *et al.* (2013) have reported that fertile hybrids are easily produced from crosses between the two species, whereas Chapman *et al.* (2005) showed that *S. aethnensis* exhibits no conspecific pollen advantage when pollinated with mixtures of pollen from both species and *S. chrysanthemifolius* showed only a small conspecific pollen advantage when treated similarly. However, Hegarty *et al.* (2009) noted that a marked decline in germination rate and survival occurred in the F_2 generation of crosses they examined, indicating that post-mating incompatibility between the two species becomes apparent after the F_1 generation, i.e., as a consequence of hybrid breakdown.

Clearly, the two *Senecio* species and the hybrid zone they form on Mount Etna comprise a very useful system for investigating adaptive divergence and mechanisms of reproductive isolation between hybridizing, diploid species. In the present study, we investigate further the nature of intrinsic reproductive isolation between *S. aethnensis* and *S. chrysanthemifolius* by examining genetic segregation in an F_2 population derived from a reciprocal cross between the two species. This enabled us to (i) construct a genetic linkage map based on segregation in this F_2 population, (ii) identify if large-scale linkage group (chromosomal) rearrangements exist between the two species, (iii) determine the occurrence and extent of TRD across the linkage groups identified and (iv) examine some of the possible causes of TRD at particular loci. Overall, the aim of our study was to obtain a better understanding of the occurrence and genetics of intrinsic incompatibility between these two species.

MATERIALS AND METHODS

F_2 population

Parent plants were raised from seed collected from wild populations located at ~2600 m (*S. aethnensis* population VB) and 600 m (*S. chrysanthemifolius* population C1) altitude, respectively, on Mount Etna, Sicily (see James and Abbott (2005) for further details of populations). Reciprocal crosses (Brennan *et al.*, 2013) were made between one representative of each of *S. aethnensis* (A) and *S. chrysanthemifolius* (C) to produce 16 F_1 individuals. The F_1 s were grown to flowering and inter-crossed in a partial diallel design. The most compatible reciprocal cross between a pair of F_1 s, i.e., producing highest seed-set in both directions, was chosen to found an F_2 reciprocal cross family (hereafter referred to as F_2 AC). The maternal and paternal origin of each F_2 individual and its F_1 progenitors were recorded to permit testing of cytoplasmic effects. A total of 120 F_2 AC seed was sown, half coming from each direction of a cross between the selected pair of F_1 parents. Following germination, F_2 AC individuals were grown, one per pot (15 cm diameter, containing a 3:1 compost to grit mix), in a glasshouse. Pots were randomized on a bench and re-randomized weekly until commencement of flowering. Twenty F_2 AC individuals (16.7%) failed to flower because they either failed to germinate, experienced early mortality or exhibited stunted growth and remained vegetative during the course of the study. As plants comprising this F_2 family were also to be used in another study aimed at examining the quantitative genetics of morphological and life history differences between *S. aethnensis* and *S. chrysanthemifolius*, non-flowering individuals were excluded from further analysis. This left 100 F_2 plants for genotyping and use in the construction of linkage maps.

DNA isolation and genotyping

DNA was extracted from young fresh leaves of all plants, i.e., the two parents, 16 F_1 progeny, 100 F_2 AC progeny (Brennan *et al.*, 2009). For a subsample of 9.4% of randomly chosen plants (41 out of 436 plants from a larger genetic study), two independent DNA extracts were made to test for genotyping reliability.

AFLP genotyping

A protocol modified from that described by Wolf (<http://archive.is/LCe6>) was used to generate amplified fragment length polymorphisms (AFLPs; Supplementary Methods). Fluorescently labelled forward primers were multiplexed in the final selective PCR step by adding 0.05 μ l of each of two labelled primers per sample and adjusting the total 20 μ l PCR volume accordingly.

The AFLP protocol was initially tested on a panel of individuals comprising the two parents and their F_1 progeny. AFLPs were detected by running samples on a Beckman Coulter CEQ 8000 capillary sequencer (Brennan *et al.*, 2009) and analysing output with CEQ v9.0 (Beckman Coulter Inc., Fullerton, CA, USA, 2004). In total, 28 paired combinations of five selective forward and eight selective reverse AFLP primers with different combinations of three final 5' bases were tested on these individuals. Samples were genotyped first using the automatic AFLP binning options of CEQ v9.0 with a minimum relative fluorescence unit (r.f.u.) cut-off of 100 r.f.u. and bin widths of 1 bp. Manual checks of the AFLP genotypes of parents, repeat extracts, and F_1 plants were then conducted to assess what subset of AFLP bands were suitable for genotyping F_2 AC individuals based on reliable amplification and inheritance pattern. Eight of the 28 primer combinations (E1M3, E1M5, E1M7, E4M7, E5M3, E5M6, E8M5, E8M7) were chosen for genotyping F_2 AC plants because they produced many reliably scored amplified polymorphic bands showing high levels of repeatability across replicate extracts (>95% across all scored loci) and the expected inheritance pattern from parents to F_1 s, and also because they could be used in multiplexed combinations (Supplementary Table S2a). AFLP bands were scored in F_2 AC individuals from the automatic AFLP CEQ v9.0 binning output with reference to parent controls and then checked manually across all samples to correct for genotyping errors.

SSR, EST and INDEL genotypes

A screen of 340 existing and newly developed single-locus molecular genetic markers was initially carried out on the two parent individuals. These loci comprised 37 simple sequence repeat loci (SSRs) developed previously by

repeat motif enrichment of genomic DNA of *S. aethnensis*, *S. chrysanthemifolius*, *S. squalidus* and *S. vulgaris* (Liu *et al.*, 2004), 25 newly developed SSRs (A Brennan and G-Q Liu, unpublished), 10 SSRs developed for *S. madagascariensis* (Le Roux and Wiczorek, 2006), 216 newly developed expressed sequence tag (EST)-SSRs and 45 EST-indels (SSR markers labelled as ES and NES, indels labelled as EC; Hegarty *et al.*, 2008; www.seneciodb.org/), and 7 additional indel markers comprising five derived from non-duplicated Asteraceae genes (Chapman *et al.*, 2007) and two from published polymorphic *Senecio* gene sequences (*SSP*, which encodes a stigma-specific peroxidase involved in pollination (McInnis *et al.*, 2005), and *Ray2a*, which encodes a cycloidea-like transcription factor involved in the control of ray floret development in *Senecio* (Kim *et al.*, 2008)). A single PCR protocol was used to amplify all markers (Brennan *et al.*, 2009). This PCR protocol consisted of a three-primer system with universal fluorescently labelled M13 primers. Multiplexed PCR products of all F₂ plants, wild sampled plants, control repeats, and both parents were run on a Beckman Coulter CEQ 8000 capillary sequencer and genotypes were assessed and scored using CEQ v9.0. Fifty-four of these marker loci were appropriate for genetic mapping in the current study based on reliable PCR amplification, fragment length scoring and the presence of fragment length polymorphisms in the F₂AC family (Table 1 and Supplementary Table S1).

Genetic mapping

A genetic linkage map was constructed from the segregation of alleles at marker loci in the F₂AC family using the demonstration version of Joinmap v4.0 (Van Ooijen, 2001). In the analysis, the F₂AC was treated as an outcrossed mapping family (CP type) because many AFLP and codominant marker genotypes were heterozygous in the F₀ parents (*S. aethnensis* and *S. chrysanthemifolius*) and therefore did not exhibit the type of F₂ inheritance that assumes parents are homozygous for alternative alleles at polymorphic loci. The outcross mapping family option (CP type) in Joinmap v4.0 allows mapping of loci with a variety of parental genotypes showing different segregation patterns. The software automatically assigns the most likely linkage phase of heterozygous parental alleles to each of the two parental chromosomes for each group of linked loci, thus enabling more loci to be mapped. A genetic map was assembled using Joinmap's default regression mapping algorithm parameters. Linkage groups were identified at greater than four logarithm of odds (LOD) score with less than 20 Kosambi centiMorgan (cM)

map distance units between loci. Map quality was assessed by examining goodness of fit G^2 statistics and markers responsible for incompatible linkage interactions were removed to generate linkage groups with high map support. In some cases, removal of suspect loci led to the splitting of large linkage groups. Diagrams of linkage groups were constructed using MapChart v2.2 (Voorrips, 2002).

Summary statistics describing map characteristics were calculated as follows. Genome length was estimated by adding twice the mean marker distance to the length of each linkage group to account for ends beyond the terminal markers (Fishman *et al.*, 2001) and also by multiplying the length of each linkage group by the correction factor (marker number + 1)/(marker number - 1) (Chakravarti *et al.*, 1991). Map coverage in terms of the percentage of the genome that is within 5 or 10 cM of a mapped marker was assessed according to the formula: $1 - \text{exponent}((-2 \times \text{distance} \times \text{marker number})/\text{map length})$ (Fishman *et al.*, 2001). The extent of marker clustering was tested using a chi-square dispersion test against a null Poisson distribution of evenly distributed markers separated by mean marker distance.

Transmission ratio distortion

Genotype frequencies at each mapped marker locus in the entire F₂AC mapping family were tested for Mendelian segregation of genotypes using chi-square tests with Microsoft Excel 2003 (Microsoft Corp). Associations between chi-square statistics for genotype segregation and marker characteristics, including marker type (classified as dominant or codominant) and AFLP fragment length, were tested with linear models after natural log transformation using R v2.13 (R Development Core Team, 2011).

The distribution of loci showing TRD, suggestive of the number of independent TRDLs, was examined by plotting per locus chi-square values onto their genetic map positions. The number of distinct TRDLs across the genetic map was assessed as the number of clusters of distorted markers at a single-locus 95% confidence level. Clustered distorted markers within each TRDL were located at a map distance of less than 10 cM from each other. The most likely map position of a TRDL was interpreted as the map position of the locus exhibiting the greatest TRD within each cluster of distorted markers. Some markers showing TRD were isolated by more than 10 cM from the nearest marker also exhibiting TRD. These markers were considered as possibly representing different TRDLs, although with weaker supporting evidence. Clustering of distorted markers across the entire genetic map was tested with a

Table 1 Summary of genetic markers screened and mapped in the *Senecio aethnensis* × *S. chrysanthemifolius* F₂ genetic mapping family

Marker type	No. of screened	No. of developed	No. of genotyped	No. of mapped	Unlinked	Problematic
<i>Codominant markers</i>						
EST indel	45	15	10	8		EC77, EC1687
EST SSR	216	48	33	31	ES91	ES19
Indel	7	4	3	3		
SSR	72	9	8	8		
Total	340	76	54	50	1	3
<i>Dominant markers</i>						
E1M3, CAAC/ACAG	—	—	19	15	179, 219	
E1M5, CAAC/ACTA	—	—	15	10	168, 213	88, 160, 204
E1M7, CAAC/ACTG	—	—	13	12	275	
E4M7, CACT/ACTG	—	—	9	8	113	
E5M3, CACC/ACAG	—	—	8	7	302	
E5M6, CACC/ACTC	—	—	9	9		
E8M5, CAGG/ACTA	—	—	11	9	114	196
E8M7, CAGG/ACTG	—	—	7	7		
Total	—	—	91	77	8	6

Abbreviations: EST, expressed sequence tag; SSR, simple sequence repeat.

Marker types are divided into codominant or dominant markers. Codominant markers are further divided into markers derived from ESTs, SSRs or insertion-deletions (indel). Dominant markers are amplified fragment length polymorphisms (AFLPs). Each AFLP primer combination is shown as the primer names used in this study followed by the three selective bases for the *EcoRI* and *MseI* primers, respectively (Supplementary Methods). No. of screened is number of codominant marker primer pairs tested. No. of developed is number of codominant markers that could be scored in *Senecio*. No. of genotyped is number of markers showing polymorphism in the F₂AC mapping family. No. of mapped is number of markers that were included in the final genetic map. Unlinked names markers that were unlinked at a >4 logarithm of odds score or >20 cM linkage threshold limits for mapping. Problematic refers to markers that caused problems with linkage group marker order or were present in multiple linkage groups. Dominant marker names are approximate fragment base-pair lengths used to label AFLP loci.

binomial test of the hypothesis that the observed number of neighbouring pairs of distorted loci per chromosome was greater than the expected number of pairs of such loci given their observed frequency. A Poisson test of TRDL clustering was not conducted, as the underlying marker distribution was significantly clustered according to a dispersion test. A possible bias of TRDLs being located at the ends of linkage groups, due possibly to weak linkage to all other mapped markers, was investigated using a binomial test of whether loci at linkage group ends more often exhibited segregation distortion relative to the overall likelihood of loci showing TRD.

Causes of TRD

Post-zygotic mortality or failure to flower. Chi-square tests were performed on the F₂AC family of segregating offspring to test the extent to which selective post-zygotic mortality or failure to flower (exhibited by 16.7% of F₂ individuals examined) could be responsible for generating the observed TRD. For these tests, expected genotype and allele segregation patterns were based on the minimum frequencies expected assuming that all non-genotyped F₂ individuals possessed the under-represented allele or genotype.

Pre-zygotic events. TRD caused by pre-zygotic events, such as biased gamete production (meiotic drive), gametophyte selection or other maternal effects (Turelli and Moyle, 2007; Fishman *et al.*, 2008), was tested at 36 codominant loci where the parental origin of alleles could be identified unambiguously.

These early reproductive events occur at the haploid stage of the life cycle and require tests on allelic rather than genotype segregation. Because such modes of incompatibility may show unilateral effects and therefore be visible in only one cross direction, allelic segregation tests were also performed on subsets of F₂ individuals containing different parental cytoplasm.

Cytoneuclear incompatibility. Because the F₂AC family was generated from a reciprocal cross, it was possible to examine whether TRD of genotypes at particular loci resulted from cytonuclear incompatibility by testing if TRD was dependent on the direction of the cross (Fishman and Willis, 2006; Turelli and Moyle, 2007). Thus, we tested TRD of genotypes in separate subsets of 49 and 51 individuals in the F₂AC mapping family that had inherited either the *S. aethnensis* or *S. chrysanthemifolius* cytoplasm, respectively (Supplementary Figures S2 and S4).

BDM incompatibilities. Hybrid incompatibilities are often thought to be due to epistatic BDM incompatibilities between parental alleles at two or more interacting genetic loci. To test for this, interactions between all pairs of identified TRDLs were examined using Fisher's exact tests of contingency tables of paired genotype counts using R v2.13 (R Development Core Team 2011).

Deficiency or excess of heterozygotes. Tests of whether TRD was caused by either a deficiency or excess of heterozygous genotypes were conducted at 36

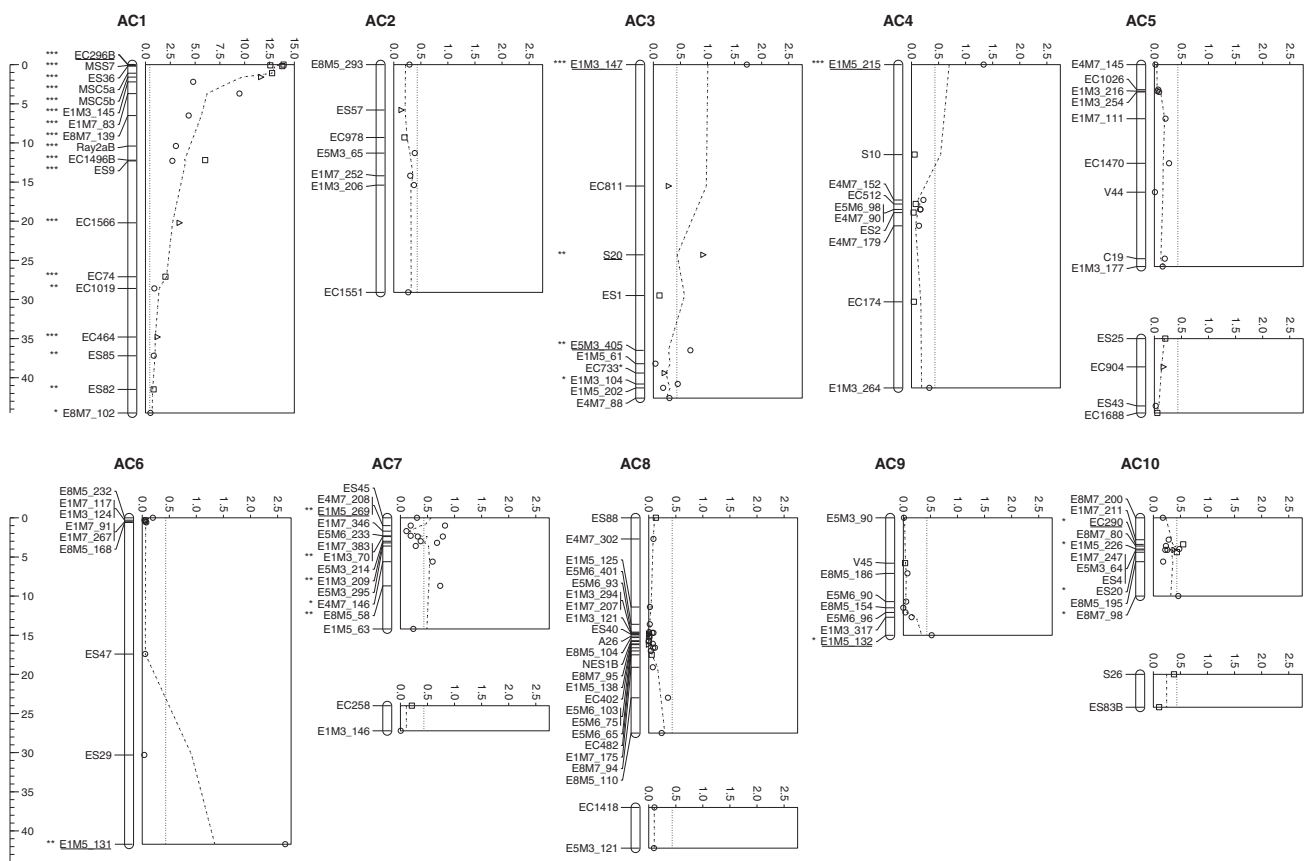


Figure 1 Genetic map of a reciprocal F₂ *S. aethnensis* and *S. chrysanthemifolius* mapping family showing mapped marker positions and associated TRD. Map distances in Kosambi centiMorgans are shown in the scale to the left of linkage groups. Linkage groups are represented by vertical bars with mapped locus positions indicated with horizontal lines. Weakly linked linkage groups (<4 LOD or >20 cM) that are thought to belong to the same chromosome are aligned vertically. Locus names are listed to the left of linkage groups. Asterisks to the left of locus names indicate significant transmission ratio distortion of genotype frequencies at a single locus χ^2 test 5% confidence limit. Underlined locus names indicate a most strongly distorted locus within a particular cluster of distorted loci. Plots to the right of each linkage group show $-\log_{1000} P$ values for single locus χ^2 tests for transmission ratio distortion of genotype frequencies. Plot symbols indicate the number of degrees of freedom for the χ^2 tests; circles = 1 df, squares = 2 df, and triangles = 3 df. The dotted lines indicate the moving average of three neighbouring loci while dashed lines indicate the 5% significance level, to the right of which, loci show significant transmission ratio distortion. The plot scale for linkage group AC1 is larger than the other linkage groups.

codominant loci where the heterozygous state of progeny in terms of parental alleles could be identified. A deficiency of heterozygous genotypes might reflect negative allelic interactions at a locus (underdominance). In contrast, an excess of heterozygous genotypes could reflect inbreeding depression (due to the expression of deleterious recessives) as a consequence of the full-sib cross used to generate the F₂ family (Remington and O'Malley, 2000; Schwarz-Sommer *et al.*, 2003), heterosis exhibited by heterozygous genotypes generated from parents homozygous for different alleles (Latta *et al.*, 2007), or the occurrence of recessive BDM incompatibilities across two or more loci (Fishman *et al.*, 2008).

Locally reduced recombination. Genomic regions exhibiting limited local recombination might show enhanced associations with TRD because of the effects of chromosomal rearrangements on chromosomal segregation during meiosis, or because they are more likely by chance to show linkage to nearby loci under selection in hybrids (Fishman *et al.*, 2013). We investigated this by testing for significant associations between local recombination rate and TRD using logistic regression of markers scored for various categories of TRD against marker distance. Different TRD categories were chosen to reflect different stringencies in defining TRDLs and TRDL map locations. These included categories where markers exhibited TRD detected by chi-square tests at (i) $P=0.05$ or (ii) $P=0.001$, (iii) where markers exhibited greatest TRD within each of the nine identified TRDLs and (iv) where markers showed greatest TRD within each of the four multi-locus TRDLs.

RESULTS

Molecular genetic markers

Details of the AFLP, and codominant markers selected to genotype *S. aethnensis* and *S. chrysanthemifolius* parents, and their F₁ and F₂ offspring for construction of linkage maps are presented in Table 1 and Supplementary Table S1.

Genetic linkage map

A genetic linkage map constructed from the segregation of markers in the F₂AC mapping family comprised 127 marker loci distributed across 14 linkage groups of average 22.4 cM length (s.d., 15.6 cM)

Table 2 Summary genetic linkage map statistics for the *S. aethnensis* × *S. chrysanthemifolius* F₂ genetic map

Linkage group	Length	Total marker no.	Dominant marker no.	Codominant marker no.	Add2s length	Method4 length
1	44.5	18	4	14	50.05	49.74
2	29.1	7	4	3	34.65	38.80
3	42.6	10	6	4	48.15	52.07
4	41.3	10	6	4	46.85	50.48
5A	25.8	9	5	4	31.35	32.25
5B	9.5	4	0	4	15.05	15.83
6	41.7	9	7	2	47.25	52.13
7A	14.2	15	14	1	19.75	16.57
7B	3.2	2	1	1	8.75	9.60
8A	27.5	22	16	6	33.05	30.12
8B	5.2	2	1	1	10.75	15.60
9	15	8	7	1	20.55	19.29
10A	10	11	8	3	15.55	12.00
10B	4.2	2	0	2	9.75	12.60
Total	313.8	127	77	50	391.56	407.06
Mean	22.41	9.07	5.50	3.57	27.97	29.08
s.d.	15.56	5.88	4.55	3.37	15.56	16.65

Map distance measures are in Kosambi centiMorgan units. Add2s length is an estimate of chromosome length calculated as linkage group length plus twice mean linkage group distance. Method4 length is another estimate of chromosome length calculated as linkage group length times (marker number + 1)/(marker number - 1).

giving a total genetic map length of 313.8 cM (Figure 1 and Table 2). Markers showing TRD were included in the construction of this map because such distortion has been shown not to bias recombination statistics nor the resulting linkage maps (Xu, 2008). Nine markers were not mapped because they were unlinked at the <4 LOD and >20 cM thresholds for inclusion in linkage groups (Table 1). Another nine markers were excluded from the map because they caused problematic linkage interactions within linkage groups (Table 1) or, in the case of two of these markers, showed linkage to multiple different linkage groups (EC77 linked AC1 and AC5A, and EC1687 linked AC5A, AC5B, and AC6). Weak linkage (<4 LOD or >20 cM distances) between markers located on four pairs of linkage groups was attributed to them belonging to distant ends of the same chromosome. Thus, ten independent linkage groups were inferred, which matches the haploid chromosome number ($n=10$) of both species (Alexander, 1979). Distance between mapped markers averaged 2.8 cM, but was highly variable (s.d. = 3.6 cM). The dispersion index for markers was large (4.66) and highly significant (dispersion test, $P=5.85e-46$), which indicated a clumped distribution of markers across the map. The total predicted map length was estimated to be 391.6 cM (Fishman *et al.*, 2001) or 407.1 cM (using the method of Chakravarti *et al.* (1991)) and ~96% and >99% map coverage of the genome was predicted to be within 5 and 10 cM of a mapped marker, respectively.

Transmission ratio distortion

TRD was common for markers included in the F₂AC genetic map with 34.0% of codominant markers and 22.1% of dominant markers showing significant deviations from Mendelian expectations of genotype segregation according to a per-locus 95% confidence level (26.8% overall percentage frequency; Figure 1 and Supplementary Table S2). Mapped codominant markers exhibited TRD more frequently than mapped dominant markers ($F_{1,125}=10.41$, $P=0.0016$), probably reflecting the fact that AFLPs, but not codominant markers, were included on the basis of inheritance patterns in the F₁ progeny panel. No association was found between AFLP fragment length and TRD ($F_{1,75}=0.67$, $P=0.4173$).

The map locations of markers were investigated to better understand the genetic architecture of incompatibilities between the two parent species, *S. aethnensis* and *S. chrysanthemifolius*. Markers showing genotypic segregation distortion were clustered in the genetic map (one-way binomial tests $P=0.0007$ and $3.5e-6$, markers showing TRD at individual chi-square test 95% and 99.9% confidence levels, respectively) to form four distinct groups located in linkage groups AC1, AC3 (distal position), AC7A, and AC10A, and also occurred individually in linkage groups AC3 (proximal and central widely separated positions), AC4, AC6 and AC9 (Figure 1 and Table 3). The widespread distribution of clusters and individual markers showing TRD across several linkage groups indicates that multiple loci contributed to the frequent TRD observed in the F₂ of this cross (Figure 1, Supplementary Figure S1, Table 3 and Supplementary Table S2). Within each cluster, all markers showed the same TRD direction favouring either alleles from the same parent or heterozygous or homozygous parental genotypes (Supplementary Figures S1–S4). Markers located at the ends of six linkage groups exhibited significant TRD, which raised the possibility that such markers that are weakly linked to other mapped markers might be erroneously interpreted as distinct TRDLs. However, a binomial test for overrepresentation of linkage group ends exhibiting TRD (6 out of 28) relative to the overall frequency of markers showing segregation distortion (26.4%) was not significant ($P=0.788$). We conclude that at least four TRDLs

Table 3 Summary of transmission ratio distortion loci and tests of different transmission ratio distortion mechanisms for the *S. aethnensis* × *S. chrysanthemifolius* F₂ genetic mapping family

Linkage group	Reference marker	Map position (cM)	Late-acting sufficient	Asymmetric	Pre-zygotic	Heterozygote	Epistasis (minority genotype)
AC1 cluster	EC296	0.0	No	No	aeth	Yes	No
AC3 singleton (proximal)	E1M3_147	0.0	Yes	No	—	—	AC3 central (CA), AC6 (CD)
AC3 singleton (central)	S20	24.3	No	chrys	chrys	No	AC3 proximal (AC), AC3 distal (AC), AC6 (AC)
AC3 cluster (distal)	E5M3_405	36.5	Yes	chrys	—	—	AC3 central (CA), AC6 (DC)
AC4 singleton	E1M5_215	0.0	No	No	—	—	No
AC6 singleton	E1M5_131	41.7	No	No	—	—	AC3 proximal (DC), AC3 central (CA), AC3 distal (CD)
AC7 cluster (central)	E1M5_269	12.5	No	No	—	—	No
AC9 singleton	E1M5_132	15	No	chrys	—	—	No
AC10A cluster	EC290	3.4	No	aeth	aeth	No	No
AC5B ^a singleton	ES25	0.0	—	aeth	—	—	—
AC10B ^a singleton	S26	0.0	—	No	aeth	No	—

Abbreviations: TRD, transmission ratio distortion; TRDL, transmission ratio distortion loci.

Linkage group names correspond to Figure 1. Cluster or singleton refers to whether a cluster or a single marker within 10 cM was observed to exhibit genotypic TRD.

The 'reference marker' is the marker with strongest TRD where a cluster of distorted markers was observed. 'Late-acting sufficient' tests if selective removal of the minority genotype according to observed post-mortality and reproductive failure would be sufficient to explain observed TRDLs. 'Asymmetric' tests if subsets of the mapping family divided according to parental cytotype showed asymmetric TRD in one cross direction only with 'aeth' and 'chrys' indicating if asymmetric TRD was observed in an *S. aethnensis* or *S. chrysanthemifolius* cytoplasmic background, respectively. 'Pre-zygotic' tests for allelic TRD with 'aeth' and 'chrys' indicating the parental alleles showing significant deficiencies and '—' indicating that no suitable codominant markers were available for testing at this TRDL. 'Heterozygote' tests for significant excesses or deficiencies in heterozygosity of genotypes scored according to parental allelic state with '—' indicating that no suitable codominant markers were available for testing at this TRDL. 'Epistasis' tests for non-independence of paired TRDL reference marker genotypes, with significantly interacting TRDLs listed. Letter codes in parentheses indicate minority genotype for the reference TRDL followed by the interacting TRDL. Genotype codes: A, homozygous *S. aethnensis* alleles; C, not homozygous for *S. aethnensis* alleles; D, not homozygous for *S. chrysanthemifolius* alleles.

^aIndicates that the last two TRDLs were only observed when specific mechanisms were investigated, so are not presented in Figure 1 or counted as primary TRDLs in the text.

influencing multiple markers were present in the mapping family or as many as nine TRDLs if additional unlinked markers showing TRD were also included.

Possible causes of TRD

Post-zygotic mortality or failure to flower. The observed TRD could have been due to selective mortality or selective failure to flower of some individuals in the mapping family (16.7% of F₂ progeny). Tests for TRD greater than could be accounted for by these causes showed that seven of the nine TRDLs (excluding the proximal and distal TRDLs on AC3) required additional mechanisms to explain the observed TRD in these regions (Table 3, Supplementary Table S2 and Supplementary Figure S1).

Pre-zygotic events. Tests of TRD caused by events at early stages of reproduction, such as meiotic drive and gametophytic selection, were conducted on codominant loci where the parental origin of alleles could be identified. These tests showed that for TRDLs in linkage groups AC1 and AC10A, and also for a new TRDL in AC10B, F₂ progeny lacked *S. aethnensis* alleles, whereas for the central TRDL in linkage group AC3, F₂ individuals lacked *S. chrysanthemifolius* alleles (Table 3, Supplementary Table S2 and Supplementary Figure S3). When cross direction was also tested for, TRDLs in AC5B and AC10A were shown to express TRD in the *S. aethnensis* maternal background only (Supplementary Figure S4 and Supplementary Table S2).

Cytoneuclear incompatibility. Tests of TRD of genotypes in subsets of F₂ individuals that reflected differences in cross direction showed an overall similar genomic distribution of TRD irrespective of parental cytoplasm (Supplementary Figures S2 and S4). However, the TRD of several TRDLs was cytoplasm dependent. This was the case for TRDLs located in linkage groups AC3 (central and distal positions) and AC9 (all dependent on *S. chrysanthemifolius* cytoplasm), and also the TRDL located in AC10A (dependent on *S. aethnensis* cytoplasm)

(Table 3, Supplementary Table S2 and Supplementary Figure S2). A further instance of TRD in linkage group AC5B was detected only in progeny possessing *S. aethnensis* cytoplasm (Table 3 and Supplementary Figure S2). Taken overall, these tests found asymmetric TRD occurring at five TRDLs dependent on individuals possessing either *S. chrysanthemifolius* or *S. aethnensis* cytoplasm.

Epistatic BDM incompatibilities. Multi-locus BDM incompatibilities were investigated by looking for negative interactions between particular genotype combinations of pairs of markers close to the nine TRDLs identified. Of the 36 paired TRDL combinations tested, five significantly non-independent pairs were found that affected four of the nine TRDLs (all three TRDLs on AC3 and one on AC6; Table 3). However, two of the five interacting TRDLs were present in the same linkage group indicating that physical proximity might contribute to their non-independence. The under-represented genotype combinations typically included one of the loci homozygous for *S. aethnensis* alleles and the other locus homozygous for *S. chrysanthemifolius* alleles.

Deficiency or excess of heterozygotes. Tests of a deficiency or excess of heterozygotes in terms of parental alleles at codominant loci where the parental state of alleles could be identified, detected a significant deficiency of heterozygous genotypes at the TRDL of large effect located in linkage group AC1 (Table 3, Supplementary Table S2 and Supplementary Figure S1). However, this deficiency is more likely caused by selection against *S. aethnensis* alleles, than underdominance (heterozygote disadvantage), because there was an even stronger bias against genotypes homozygous for *S. aethnensis* alleles across this particular region of AC1 (Supplementary Figure S3). No TRDLs showing an excess of heterozygotes relative to homozygotes were identified, thus discounting inbreeding depression, heterosis, or recessive BDM incompatibilities, as important causes of TRD in this F₂ family.

Locally reduced recombination. In no instance was an association detected between locally reduced recombination and TRD for any of the four categories of markers tested. Thus, for markers that showed TRD at 95% and 99.9% confidence levels, the probability of a stronger association with recombination rate than the null hypothesis of no association was $P=0.300$, and $P=0.291$, respectively, whereas the probability was $P=0.261$ for the most strongly distorted marker within each of the nine identified TRDLs, and $P=0.178$ for the most strongly distorted marker within each of the four multi-locus TRDLs treated separately. Thus, these results do not support the hypothesis that chromosomal regions with reduced recombination are associated with hybrid incompatibility in this system.

DISCUSSION

Studies of closely related species that form hybrid zones can reveal much about the genetic basis and maintenance of species divergence in the face of frequent interspecific hybridization and gene flow. Our previous analysis of the hybrid zone between *S. aethnensis* (occurring at high altitude) and *S. chrysanthemifolius* (occurring at low altitude) on Mount Etna, Sicily, compared clines for molecular variation with those for phenotypic trait variation, and indicated that both extrinsic and intrinsic selection against hybrids act to maintain the hybrid zone despite high levels of gene flow (Brennan *et al.*, 2009). The study reported here has expanded on these previous results and provided insights into the genetics of intrinsic reproductive isolation between *S. aethnensis* and *S. chrysanthemifolius* on Mount Etna, Sicily. We found intrinsic genomic incompatibility between these two species caused by a variety of genetic mechanisms at multiple genetic loci. However, large-scale genomic rearrangements or translocations between the species did not appear to contribute greatly to this incompatibility.

F₂ linkage map structure

To investigate the genomic architecture of these hybridizing *Senecio* species, we constructed a genetic linkage map from the segregation of dominant and codominant molecular markers in the F₂ of a reciprocal cross between the two species. The resulting F₂AC linkage map comprised ten distinct linkage groups (taking account of four weakly linked linkage group pairs), which corresponds to the haploid chromosome number of the two species (Alexander, 1979). If the parental species were distinguished by chromosomal translocations, the affected regions would link different linkage groups in genetic maps of hybrids and reduce the number of independent linkage groups to below the haploid chromosome number of the species investigated (Fishman *et al.*, 2013). Therefore, the ability to distinguish ten distinct linkage groups in the F₂AC map and the removal from maps of only two markers with suspect linkage to multiple different linkage groups indicates that large-scale genomic translocations between chromosomes probably do not distinguish the two species in contrast to what has been found in hybridizing annual sunflowers (Lai *et al.*, 2005; Yatabe *et al.*, 2007).

Overall, the map showed good coverage with an average locus distance of just 2.8 cM and >99% of the genome predicted to be within 10 cM of a mapped marker. However, markers exhibited a highly clumped distribution within the map, for which there are several possible explanations. First, there may be technical reasons for marker clustering, such as marker position uncertainty due to a limited number of recombination events observed in the relatively small F₂ family examined, and/or to genotyping error, which though estimated to be reasonably low (2.2% based on duplicated samples tested) translates to a 2.2 cM uncertainty in the position of loci within

maps. Second, some marker clusters could be explained by sequence heterogeneity across the genome. For example, clusters of AFLP loci could signal repetitive genomic regions containing many closely spaced repeated restriction enzyme cut sites, whereas clusters of EST and gene loci could signal highly expressed, gene-rich genomic regions. Third, the occurrence of clusters could reflect variable recombination rates across the genome caused by intrinsic features such as low recombination near centromeres or in regions where local chromosome rearrangements, such as inversions, exist between species.

Transmission ratio distortion

A notable feature to emerge from the current study was the large number of molecular marker loci that showed TRD in the F₂AC mapping population, i.e., 34 (26.8%) of 127 markers tested. These loci were non-randomly distributed across the genetic map and generally formed clusters in which all loci showed the same bias against alleles from one parent or against heterozygous combinations of parental alleles. We found strong multi-locus evidence for the occurrence of four TRDLs and weaker single-locus evidence for five additional TRDLs located in seven of the ten linkage groups identified. Individual markers showing TRD were sometimes located at the ends of linkage groups raising the possibility that they were technical artifacts; however, such markers were not over-represented relative to the overall observed numbers of markers exhibiting TRD.

One TRDL, located at the proximal end of linkage group AC1, showed particularly strong segregation distortion, both in terms of the extent to which allele and genotypes frequencies were distorted away from Mendelian expectations for *S. aethnensis* alleles and genotypes, and the length of the genome affected (all of AC1, that is, 44.5 cM). The fact that this TRDL, and also other multi-locus TRDLs, affected large genomic regions was likely due to both their strong effects on segregation distortion and the limited post-hybridization recombination that had occurred in the F₂AC mapping family. Strong TRD in early-generation hybrids is likely to bias patterns of introgression across large genomic regions in later generation hybrids because hybrid genotypes at linked loci are eliminated before they have an opportunity to recombine away from the TRDL (Bierne *et al.*, 2011).

Causes of TRD

Through further analyses of the patterns of segregation in the F₂AC family, we obtained some insights into the causes of TRD at different TRDLs. Twenty of the original 120 progeny that comprised the F₂AC mapping family failed to flower either because of a failure to germinate, the occurrence of early mortality, or an inability to develop to the flowering stage. These individuals were not genotyped and their absence may have contributed to the TRD observed at marker loci. The low intrinsic fitness of these individuals might stem from several causes (see below), including the disruption of coadapted gene complexes (hybrid breakdown) following recombination. We did not examine survival and flowering in the F₁ generation in our study, however a previous study (Hegarty *et al.*, 2009) recorded a marked drop in intrinsic fitness (measured in terms of seed germination and seedling survival under glasshouse conditions) between the F₁ and F₃ generations of a reciprocal cross between the same *S. aethnensis* and *S. chrysanthemifolius*, parental individuals as this study indicating the occurrence of hybrid breakdown.

Although post-zygotic selective mortality or inability to flower of some F₂ individuals would contribute to TRD at certain marker loci, we found that the level of TRD exhibited by seven of the nine TRDLs identified could not be entirely explained in this way. However,

it remains possible that earlier-acting post-zygotic incompatibility in the form of a failure of seed development (e.g. due to negative interactions between zygote and endosperm) could have contributed to the observed TRD. Alternatively, there may be pre-zygotic causes of TRD either in the production of gametes containing particular alleles (meiotic drive) or selection against gametes containing particular alleles (gametophytic selection) (Fishman and Willis, 2005; Fishman *et al.*, 2008). We tested for these possibilities at loci where the parental origin of alleles could be identified unambiguously allowing TRD of alleles rather than genotypes to be examined and found three TRDLs where there was a bias against *S. aethnensis* alleles and one TRDL where there was a bias against *S. chrysanthemifolius* alleles. Thus, pre-zygotic factors of the type mentioned above could have contributed to the TRD recorded at these loci.

Incompatibility between diverging genomes is often asymmetric when the underlying causes of incompatibility include either pre-zygotic haploid stages of the life cycle, such as meiotic drive or pollen fitness, or cytonuclear incompatibilities between nuclear and organelle genomes (Levin, 2003; Fishman and Willis, 2006; Turelli and Moyle, 2007). Strong crossing asymmetry or unilateral incompatibility is most likely observed when few asymmetric incompatibilities of large effect are involved (Turelli and Moyle, 2007). The extent to which TRD was asymmetric and dependent on cross direction was tested by comparing TRD across loci in F_2 progeny possessing cytoplasm inherited from either the *S. aethnensis* or *S. chrysanthemifolius* parent. These tests revealed five TRDLs exhibiting asymmetric differences in TRD dependent on parental cytoplasmic background. We may conclude that the asymmetric TRD at these loci reflects cytonuclear incompatibility and/or the effects of meiotic drive and/or gametophytic selection.

There is growing evidence for the widespread occurrence of BDM incompatibilities between allopatric or parapatric populations and their role in limiting subsequent hybridization and possibly promoting further divergence during speciation (Coyne and Orr, 2004; Corbett-Detig *et al.*, 2013). These BDM incompatibilities are frequently caused by deleterious interactions between different parental alleles occurring at two or more loci. Significantly non-independent, paired-locus genotype frequencies were observed for five pairs of TRDLs affecting four of the nine TRDLs (Table 3). However, three of the five interacting TRDL pairs were present in the same linkage group indicating that physical proximity might contribute to their non-independence. The under-represented genotype combination typically included one of the loci homozygous for *S. aethnensis* alleles and the other locus homozygous for *S. chrysanthemifolius* alleles.

We also tested if TRD could be due to (i) underdominance (reflected by a deficiency of heterozygotes at certain loci) or (ii) inbreeding depression, heterosis or recessive BDM incompatibilities (reflected by an excess of heterozygotes). Inbreeding depression could be caused by the expression of deleterious recessive alleles as a result of using a full-sib F_2 mapping family from two outcrossed self-incompatible parents. Tests of heterozygosity revealed that underdominance could have contributed to a significant deficiency of heterozygous genotypes at only one TRDL, in the AC1 linkage group. However, the heterozygote deficiency at this locus was more likely caused by selection against *S. aethnensis* alleles, rather than underdominance, because there was an even stronger bias against genotypes homozygous for *S. aethnensis* alleles across the AC1 TRDL (Supplementary Figure S1). An excess of heterozygotes was not evident at any TRDL and thus it is concluded that inbreeding depression, heterosis or recessive BDM incompatibilities have not contributed to the TRD recorded in the F_2 AC mapping family.

As mentioned earlier in the discussion, reduced recombination near centromeres or due to local inversions may be causes of marker clustering observed in the genetic map. These causes of reduced recombination may also contribute to TRD at marker loci. For example, selfish drive elements are typically found near centromeres where they directly influence chromosome segregation patterns in their favour during meiosis (Henikoff *et al.*, 2001), whereas chromosomal rearrangements may also influence inheritance patterns because recombination within the rearranged region is typically selected against (Ortiz-Barrientos *et al.*, 2002; Kirkpatrick and Barton, 2006; Lowry and Willis, 2010). To examine whether genomic regions showing limited recombination could be a cause of TRD in the F_2 AC family, we tested for, but did not find associations between, marker clustering and TRDLs. Therefore, genomic regions showing limited recombination do not seem to have a strong role in reinforcing genomic divergence in *Senecio*. However, our current F_2 genetic map lacks sufficient marker density to adequately test this association, as the clumped marker map distribution could be due to a variety of reasons other than variation in recombination rates and the genomic scale of TRDL-low recombination associations could be highly localized (Yatabe *et al.*, 2007; Jones *et al.*, 2012; Renaut *et al.*, 2013). Future studies aimed at determining whether differences in chromosomal rearrangement may be a cause of genomic incompatibility between *S. aethnensis* and *S. chrysanthemifolius* should involve cytogenetic comparisons of karyotypes and/or detailed comparisons of high marker-density genetic maps of the two species.

Overall, it is clear from our analyses that a variety of genetic mechanisms at multiple genetic loci across the genome are likely to contribute to the intrinsic incompatibility existing between *S. aethnensis* and *S. chrysanthemifolius*. Further fine-scale genetic mapping studies of *S. aethnensis* and *S. chrysanthemifolius* involving quantitative trait locus mapping of traits of adaptive relevance, larger families to better estimate recombination, more markers for better genome coverage, and families from intraspecific crosses, will be necessary to further investigate the various mechanisms contributing to hybrid incompatibility and ecological divergence in this species pair.

CONCLUSIONS

While a number of studies have indicated that *S. aethnensis* and *S. chrysanthemifolius* are highly interfertile (Chapman *et al.*, 2005; Brennan *et al.*, 2013), the present study has revealed evidence of intrinsic incompatibility between these two species in the form of multiple genomic regions showing TRD. Our results, therefore, support the findings of our previous clinal analysis of the natural hybrid zone on Mount Etna, which indicated that intrinsic selection against hybrids was an important factor maintaining species differences in the face of gene flow (Brennan *et al.*, 2009). The present study shows that hybrid incompatibility between these two diverging plant lineages is more cryptic than previously considered, and is manifested in the F_2 generation rather than in the F_1 (Hegarty *et al.*, 2009). Some studies of other diverging plant lineages have yielded similar findings (Fishman *et al.*, 2001; Moyle and Graham, 2006; Fishman and Willis, 2006).

It follows that the effects of TRDLs with multiple modes of action at multiple, relatively large, genomic regions in early-generation hybrids between *S. aethnensis* and *S. chrysanthemifolius* are likely to impact the genetic structure of the natural hybrid zone on Mount Etna by limiting introgression and promoting divergence across the genome (Feder *et al.*, 2012; Abbott *et al.*, 2013). However, large-scale genomic translocations or other rearrangements between the species

do not seem to contribute in any major way to this process. It has been estimated that *S. aethnensis* and *S. chrysanthemifolius* are of relative recent origin (<1 million years ago, Comes and Abbott, 2001; ~108 000 to 150 000 years ago, Chapman *et al.*, 2013, Osborne *et al.*, 2013), which indicates that the various forms of intrinsic hybrid incompatibility that clearly exist between these two species must have evolved relatively rapidly.

DATA ARCHIVING

Mapping family genotype data available from the Dryad Digital Repository: doi:10.5061/dryad.7b56k. Other results can be found in Supplementary Information.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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Supplementary Information accompanies this paper on Heredity website (<http://www.nature.com/hdy>)