Rapid Chromosome Evolution in Recently Formed Polyploids in *Tragopogon* (Asteraceae)

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Abstract

**Background:** Polyploidy, frequently termed “whole genome duplication”, is a major force in the evolution of many eukaryotes. Indeed, most angiosperm species have undergone at least one round of polyploidy in their evolutionary history. Despite enormous progress in our understanding of many aspects of polyploidy, we essentially have no information about the role of chromosome divergence in the establishment of young polyploid populations. Here we investigate synthetic lines and natural populations of two recently and recurrently formed allotetraploids *Tragopogon mirus* and *T. miscellus* (formed within the past 80 years) to assess the role of aberrant meiosis in generating chromosomal/genomic diversity. That diversity is likely important in the formation, establishment and survival of polyploid populations and species.

**Methodology/Principal Findings:** Applications of fluorescence in situ hybridisation (FISH) to natural populations of *T. mirus* and *T. miscellus* suggest that chromosomal rearrangements and other chromosomal changes are common in both allotetraploids. We detected extensive chromosomal polymorphism between individuals and populations, including (i) plants monosome and trisome for particular chromosomes (perhaps indicating compensatory trisomy), (ii) intergenomic translocations and (iii) variable sizes and expression patterns of individual ribosomal DNA (rDNA) loci. We even observed karyotypic variation among sibling plants. Significantly, translocations, chromosome loss, and meiotic irregularities, including quadrivalent formation, were observed in synthetic (S₀ and S₁ generations) polyploid lines. Our results not only provide a mechanism for chromosomal variation in natural populations, but also indicate that chromosomal changes occur rapidly following polyploidisation.

**Conclusions/Significance:** These data shed new light on previous analyses of genome and transcriptome structures in *de novo* and establishing polyploid species. Crucially our results highlight the necessity of studying karyotypes in young (<150 years old) polyploid species and synthetic polyploids that resemble natural species. The data also provide insight into the mechanisms that perturb inheritance patterns of genetic markers in synthetic polyploids and populations of young natural polyploid species.

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Introduction

Polyploidy has played a major role in generating angiosperm biodiversity. Chromosome counts suggest that between 30 and 80% of angiosperm species are polyploid, while genomic studies of selected model and crop species reveal evidence of extensive ancient genome-wide multiplications. Indeed recent genomic investigations indicate that most, if not all, angiosperm species have undergone at least one genome duplication event in their evolutionary history, and several have evidence of multiple polyploidy-diploidisation-polyploidy cycles [1,2,3].

Angiosperm genomes are astonishingly plastic in their ability to tolerate considerable karyotypic (e.g. chromosome number variation, translocations), genetic (mutations, retroelement transposition, deletions) and epigenetic (DNA methylation, histone methylation/acetylation) variability. This tolerance enables polyploids to form and establish and has contributed significantly to their widespread occurrence [4]. Large-scale genetic changes induced by polyploidy are thought to influence the transcriptome, metabolome and proteome, which can concomitantly alter the phenotype and ecology of the individuals. Most of the new genetic changes are probably maladaptive, but in a few rare instances individuals arise that are able to outcompete the parental diploids or colonize new niches.

There are several examples of recent speciation via polyploidy that occurred within the last 150 years: *Spartina anglica* [5], *Senecio*
cambrensis and S. eboracensis [6], Cardamine Schulzi [7], Tragopogon mirus and T. miscellus [6]. Studies on the genetic consequences of allopolyploidy in these de novo polyploid species reveal some significant differences. In S. anglica allopolyploidy induced few changes in genome structure, but there is epigenetic reprogramming [5,9], while in Senecio [10,11] and Tragopogon [12] allopolyploids there are substantial genetic changes including loss of sequence (genomic DNA profiles) and perturbations to the transcriptome (cDNA profiles). However, little attention has been paid to the chromosomal content of any of these allopolyploids, and it is unknown what mechanisms drive the observed genetic changes. For example, genetic change could be driven by local mutation, small-scale deletion or insertion of a particular sequence, or via major chromosomal changes, including whole-arm transposition and chromosome losses or duplications. Because there is little or no understanding of chromosomal variation in these recently formed, natural polyploids, we have embarked on a characterization of karyotypic variation between individuals and populations of the two Tragopogon allopolyploids from North America.

Tragopogon mirus and T. miscellus have proved to be excellent evolutionary model systems for understanding early allopolyploid formation. These allopolyploids are derived from three diploid progenitors (each 2n = 2x = 12), T. pratensis, T. porrifolius and T. dubius, the latter being shared by both allopolyploids (T. mirus derived from T. dubius × T. porrifolius, 2n = 4x = 24; T. miscellus derived from T. dubius × T. pratensis, 2n = 4x = 24, Figure 1) [13]. The diploid parents of both polyploids are in well separated clades [14] and are not closely related based on ITS/ETS sequences, allozymes and other genetic markers [15,16]. There are many reports of natural F1 hybrids involving these diploids and we have also produced them in the glasshouse, but these hybrids are highly sterile suggesting minimal homeologue pairing at meiosis [8]. In contrast the allopolyploids T. mirus and T. miscellus are fertile and expanded their ranges rapidly after their initial formations, in large part via multiple origins [17]. The two allopolyploids now occupy a large geographic area of eastern Washington and adjacent Idaho, USA, and comprise many thousands of individuals in several populations. Molecular analyses have revealed that T. mirus has recurrently formed at least 13 times and T. miscellus possibly as many as 21 times [13,16,18,19,20, Symonds et al. unpubl., 21], reviewed in Soltis et al. [8]. Furthermore, there are genetic differences between populations of each tetraploid, most of which likely reflect variation found in the diploids and inherited in the polyploid populations through recurrent formation, and others of which may have arisen through subsequent divergence of the polyploids. Examples of the latter include the divergence of rRNA gene copy number, sequence homogeneity and expression patterns [22,23]. Here we show that population differences are also reflected in substantial variability in karyotypes among individuals, differences that appear correlated with irregular meiosis.

Results

Genome structure of Tragopogon allotetraploids

The chromosome sizes, numbers and centromere indices are similar between the parental diploid species, and three tandem repeats characterized by us do not generate differences in chromosomal distribution. For these reasons we are unable to identify the parental origin of the chromosomes in the derived allotetraploid species via morphology alone [24]. We therefore

Figure 1. Tragopogon triangle with the flowers of the diploid Tragopogon species at the apices of the triangle. The flowers of the polyploids T. miscellus and T. mirus are shown between their respective diploid parents. The synthetic polyploid described here has the same parents as T. mirus. doi:10.1371/journal.pone.0003353.g001
used FISH with total genomic DNA probes (called Genomic In Situ Hybridization or GISH) to determine the genomic composition of karyotypes of T. mirus and T. miscellus individuals (Figures 2 and 3). We analysed 12 plants, nine of T. mirus from three populations (Table 1) and three of T. miscellus, two from one population and a third individual from a second population (Table 2). Five of these plants, including representatives of each tetraploid, were chosen because we had previously observed [22,23] that they had particular 45S nuclear ribosomal DNA (45S rDNA) compositions and expression characteristics (Table 3). GISH labelling enabled the genomic origin of the chromosomes to be determined by fluorescence colour: digoxigenin-labelled genomic DNA of T. dubius labelled chromosomes of T. dubius origin green or yellow (D-genome) and biotin-labelled genomic DNA of either T. pratensis (to T. miscellus) or T. porrifolius (to T. mirus) labelled the P-genome (either T. pratensis or T. porrifolius) orange or red. However, the distinction between genomes was only possible after electronic merger of the images because there was considerable cross-hybridisation of probes. The homologous group assigned to each chromosome (A–F) was determined using size, arm ratio, position of 5S and 45S rDNA, as described in Pires et al. [24], and by using DAPI bright bands that were revealed after denaturation in some metaphases.

Previous cytogenetic analyses revealed that both T. mirus and T. miscellus had 24 chromosomes; aneuploidy has not been previously reported in these taxa [13,24,25]. This chromosome number was also observed here in all but one individual of T. mirus, which had 23 chromosomes. It was assumed that GISH would partition the chromosome sets into 12 chromosomes of either T. dubius origin or T. pratensis origin depending on the tetraploid being analysed. However, only four had an entirely balanced additive karyotype of that expected from the chromosomes of the diploid parents (Tables 1 and 2 and Figures 2 and 3). We were surprised to observe that five of the eight T. mirus karyotypes and two of the three T. miscellus karyotypes with 2n = 24 chromosomes had unbalanced genomic contributions. This was manifest by some chromosomes occurring in one or three copies (monosomic and trisomic, respectively; see Tables 1 and 2). In addition, four plants, including individuals of both T. mirus and T. miscellus, carried intergenomic translocations (Figures 2A, 3A, arrows).

The T. mirus individuals 2603-33A and 33B are siblings from different flower heads of the same plant. Significantly, their intergenomic translocations (Figures 2A, 3A, arrows). Previous cytogenetic analyses revealed that both T. mirus and T. miscellus had 24 chromosomes; aneuploidy has not been previously reported in these taxa [13,24,25]. This chromosome number was also observed here in all but one individual of T. mirus, which had 23 chromosomes. It was assumed that GISH would partition the chromosome sets into 12 chromosomes of either T. dubius origin or T. pratensis origin depending on the tetraploid being analysed. However, only four had an entirely balanced additive karyotype of that expected from the chromosomes of the diploid parents (Tables 1 and 2 and Figures 2 and 3). We were surprised to observe that five of the eight T. mirus karyotypes and two of the three T. miscellus karyotypes with 2n = 24 chromosomes had unbalanced genomic contributions. This was manifest by some chromosomes occurring in one or three copies (monosomic and trisomic, respectively; see Tables 1 and 2). In addition, four plants, including individuals of both T. mirus and T. miscellus, carried intergenomic translocations (Figures 2A, 3A, arrows).

The T. mirus individuals 2603-33A and 33B are siblings from different flower heads of the same plant. Significantly, their karyotypes exhibit substantial differences; 2603-33B has an expected karyotype assuming complete additivity of parental chromosome sets, while 2603-33A is monocentric for chromosome F0 and trisomic for chromosome E0. Meiotic and mitotic aberrations in synthetic reconstruction of T. mirus

Meiosis in newly formed, synthetic allotetraploids (S0 and S1) (see Figure 1) was examined to determine if meiotic aberrations occurred in early allotetraploid generations and could be a source of genomic imbalance observed in root tip metaphases. Meiotic cells of developing anthers at diplotene were assessed (Figure 4). In many instances regular bivalent formation occurred (Figure 4A), although frequently bivalents overlapped (Figure 4C, D), making it difficult to be certain if they were in fact multivalents. However, in a number of cases resolution was sufficient to determine the presence of quadrivalents (Figure 4B). Using GISH, the genomic origin of the chromosomes could be resolved, although the distinction was less clear than at metaphase. Figures 4E–F show GISH labelling of a quadraivalent with two chromosomes of T. dubius origin and two chromosomes of T. porrifolius origin physically linked via chiasma in a large, twisted ring.

An analysis of root tip metaphases in 12 allotetraploids of the subsequent S1 generation revealed one plant that was 2n = 25, the rest were 2n = 24, as expected. There were cytogenetic abnormalities in two additional plants. One of these plants (73-14) had unusually large 45S rDNA loci occurring on both copies of chromosome A00 (Figure 5 A, B). This plant also had an rDNA locus on chromosome A00 of T. porrifolius origin, but the expected site on chromosome D00 was absent. The second plant (134-16-3) had a translocation of T. porrifolius origin to chromosome C00. Since there is no missing chromosome segment of T. porrifolius origin, it must be assumed that this was a non-reciprocal translocation induced during the preceding meiosis (Figure 5 C, D, E).

45S rDNA decondensation

Previous cytogenetic analyses of diploid Tragopogon species revealed that T. dubius and T. pratensis carry a 45S rDNA locus on chromosome A00 and A0, respectively, while T. porrifolius has two loci on chromosomes A00 and D00. To determine if these sites were inherited in the respective allotetraploid species we used FISH with digoxigenin-labelled probe pTa71 (yellow fluorescence, Figures 2 and 3). In T. miscellus, 45S rDNA loci occurred on chromosomes A00 and A0 (Figure 3). At metaphase both chromosomes carried secondary constrictions (Table 3). In T. mirus, 45S rDNA loci occurred on chromosomes A00, A0, and D0 (Figure 2). These loci were also frequently identifiable after GISH labeling, without the use of the rDNA probe pTa71, appearing with a brighter green (to T. dubius origin loci) or yellow fluorescence (Figure 2). In all cells, both mitotic and meiotic, the number of 45S loci and their parental distribution were as expected, i.e. pairing and segregation of rDNA-carrying chromosomes is probably regular. This was not the case, however, for 55 rDNA-carrying chromosomes (5S probe, red fluorescence). We would expect T. mirus to carry 3S rDNA loci on chromosomes A00, A0, and D00. But here we observed one T. mirus individual (2602-03-10) trisomic for chromosome F0 and each of these chromosomes carried a single 5S rDNA locus (Figure 2). It was a surprise that the chromosomes carrying the 45S rDNA were balanced because at diplotene they were frequently in close proximity (Figure 4 E, F), probably reflecting transcriptional activity and nucleolar function. The absence of aberrant numbers of 45S rDNA-carrying chromosomes at metaphase suggests that they paired regularly at meiosis.

An analysis of 45S rDNA-carrying chromosomes in 14 diplotene nuclei of T. mirus revealed that chromosome A00 was always associated with the nucleolus (Figure 4E-H). Often this chromosome was associated with a chromosome from the T. porrifolius genome (when identifiable it was chromosome D00, Figure 4G, H). Similar results were observed at metaphase (Table 3, Figure 2), where chromosome A00 and sometimes chromosome D0 carried secondary constrictions (the exception is plant 2603-33A–below). In contrast, chromosome A00 was always condensed and unassociated with the nucleolus. Interestingly, in two individuals of population 2602 only one of the two homologues of chromosome A00 carried secondary constrictions. In plant 2603-33A, there was no secondary constriction on chromosome A00 and the 45S rDNA locus was unusually small. Instead, chromosome D0 had a secondary constriction. Similarly in two synthetic T. mirus, plants 134-16-3 and 73-14, there were secondary constrictions on A00 and A0, with the locus on chromosome D0 missing in the latter individual (Figure 5 A, C).

In natural T. mirus and T. miscellus, the presence of secondary constrictions correlated well with patterns of condensation/decondensation at interphase (Figure 2D, E); for example, chromosomes A00 and D00 have no secondary constrictions in
Figure 2. (A-C) Karyotype analyses of *T. mirus* from individuals in three populations (A) 2601, (B) 2603 and (C) 2602. Homeologous chromosome group nomenclature (A-F) follows Ownbey and McCollum [25] and Pires et al. [24]. Fluorochrome colours: yellow/green (FITC, digoxigenin-labelled probes), orange/red (Cy3, biotin-labelled probes), blue (DAPI staining). Each karyotype is shown with DAPI staining, sometimes also simultaneously labelled for 45S rDNA (yellow fluorescence) or 5S rDNA (red fluorescence), and after GISH with *T. dubius* (green fluorescence) and *T. porrifolius* (red fluorescence) total genomic DNA probes. Monosomic or trisomic chromosomes are underlined. Note: (1) intergenomic translocations to chromosome Cpo/du (arrows) in individuals 2601-7 and 2601-8 and Ddu/po in individual 2602-0-3-4 (A). (2) The 45S rDNA locus on the two homologues of chromosome Adu in individuals 2602-4 and 2602-0-3-4 show different levels of decondensation, one being condensed and the other with a secondary constriction (C). (3) A large reduction in size of the 45S rDNA locus on chromosome Adu of individual 2603-33A (D) compared with other individuals (D). (4) No secondary constrictions of 45S rDNA loci on *T. porrifolius* origin chromosomes (Apo, Dpo) in individual 2603-33B. (D-E) Root-tip interphase nuclei of *T. mirus* after FISH with digoxigenin-labelled pTa71 for 45S rDNA. (D) Individual 2601-4, showing four large condensed 45S rDNA loci (arrows) and (E) Individual 2602-0-3-10 with two large condensed 45S rDNA loci (arrows). Scale bar (top right) is 10 μm.
T. mirus 2601-4 (not shown) and exhibit four sites of condensed rDNA chromatin (Figure 2D) while in T. mirus (2602-0-3-10) chromosome Apo is without a secondary constriction (Figure 2C) and at interphase there are two highly condensed rDNA loci (Figure 2E).

Southern blot analysis of 5S rDNA loci

Given the imbalance observed in the inheritance of individual chromosomes, we would expect the copy number of 5S rDNA units to vary among plants. For example, individual 2602-3-10, showing trisomy of the Fpo chromosome, had seven 5S rDNA signals instead of the expected six (Figure 2C). On the other hand, if there was Fpo monosomy, we might expect a reduction in gene copy numbers. Using Southern blot hybridisation, we determined the parental T. porrifolius/T. dubius 5S rRNA gene copy number ratio in the progeny of a single T. mirus plant, 2602-0-3. Several progeny were studied, including those analysed by GISH (plants 2602-0-3-10 and 2602-0-3-4). To ascertain the parental origin of the 5S rDNA units, we digested genomic DNA with TaqI restriction enzyme. This enzyme takes advantage of a restriction site present in T. porrifolius units that is not found in T. dubius units. Consequently, after digestion the T. porrifolius 5S rDNA was digested mostly to monomers while there was little digestion of T. dubius units that remained at high molecular weight (Figure 6). In T. mirus, the 5S rDNA probe used for Southern hybridisation revealed signals from both parental gene families. However, the distribution of signal intensities between genes of T. porrifolius and genes of T. dubius origin differed between progeny. For example, the trisomic individual 2602-0-3-10 had a T. porrifolius/T. dubius 5S rDNA unit ratio close to 1.0, while the “normal” disomic plant 2602-0-3-4 had a ratio that was 0.82. Thus, trisomy of chromosome Fpo resulted in an approximately 10% increase in the number of 5S genes in the genome. This value is in a good theoretical agreement with the expected increase in number of 5S arrays from six to seven (per diploid cell). Three other plants (2602-0-3-5, -7 and -8) had a 5S rDNA gene ratio corresponding to a trisomic genotype, suggesting meiotic aberrations were frequent in this lineage. On the other hand, two plants (2602-0-3-2 and -3) had a decreased number of T. porrifolius genes compared with the expectation (plant 2602-0-3-4, with balanced chromosomal sets). Perhaps these two individuals are monosomic for Fpo.

Discussion

Effectiveness of GISH

Phylogenetic analyses of internal (ITS) and external transcribed spacer (ETS) sequences of nuclear 45S rDNA suggest that the diploid progenitors of both T. mirus and T. miscellus are distantly related, with T. dubius in one major clade of Tragopogon and both T. pratensis and the populations of T. porrifolius that served as parents in another major clade [26,27]. Earlier allozyme studies also suggested that the three diploid progenitors were well differentiated genetically [16,28]. Likewise, comparisons of cDNA-AFLP genetic markers between the diploid species reveal that T. pratensis and T. dubius share only between 30-40% of markers [12], again suggesting considerable genomic divergence.

Recently, Markova et al. [29] used GISH with genomic DNA probes from diploid species of Silene onto metaphases of related diploids to show that labeling strength was inversely correlated to genetic distance, i.e. there was strongest labeling to the most closely related species. Given these data and an apparently large
genetic distance between *Tragopogon* diploids, we might expect GISH to work effectively on the derived allopolyploids. Instead, we observed that it worked rather weakly, with much cross-hybridisation of probes, and the genomic distinction was only resolvable after electronic merging of images. Indeed previously we had reported that GISH had not worked at all in *Tragopogon* [24]. Our success here is due to improved quality of the electronics on the microscope’s camera and improved facilities to electronically merge images using Openlab software, but it remains an enigma why GISH does not work more effectively in *T. mirus* and *T. miscellus*.

**rDNA inheritance and expression patterns**

We expect *T. miscellus* to inherit two 45S rDNA loci (carried on chromosome A<sub>du</sub> from *T. dubius* and A<sub>pr</sub> from *T. pratensis*), while *T. mirus* inherits three loci (carried on chromosomes A<sub>du</sub> from *T. dubius* and A<sub>pr</sub> and D<sub>po</sub> from *T. porrifolius*). The observations of only two 45S rDNA in synthetic *T. mirus* 73-14 (Figure 5A, B) can be explained in one of two ways. First, the enlargement of the rDNA locus on chromosome A<sub>du</sub> and the locus loss on D<sub>po</sub> occurred through an rDNA translocation/fusion involving the loci on D<sub>po</sub> and A<sub>du</sub>. However, given that the karyotype is balanced and the material is the S<sub>2</sub> generation, this scenario seems unlikely because it would suggest the union of two gametes carrying the same abnormality, or arising via restitution of segregating chromosomes in meiosis II. The second and alternative explanation is that the rDNA locus number may reflect the situation in the *T. porrifolius* parent used for the cross, although our past analyses of *T. porrifolius* never revealed such a polymorphism [24]. Unfortunately the precise *T. porrifolius* parent used in the cross is now deceased (the plants are annuals or biennials), preventing us from determining which of the competing hypotheses is correct.

Previously we showed that natural *Tragopogon* allopolyploids also do not have fixed patterns of 45S rDNA inheritance, with some individuals showing a balanced distribution of rDNA sequences originating from each of the parental diploids and some showing biased inheritance, typically with the number of rDNA units from *T. dubius* being underrepresented [22]. However, from molecular studies it was not clear whether non-Mendelian rDNA inheritance was caused by a decrease in copy number (elimination) or unit replacement (e.g. via homogenisation mechanisms). In one individual of *T. mirus*, 2603-33 (referred to here as 2603-33A),

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**Table 1. Karyotype organisation of individuals of *T. mirus* from three populations (localities of each population are indicated in column 1).**

| *T. mirus* population | Plant number | Chromosome No. (2n) | Nature of karyotype imbalance | Translocations |
|-----------------------|--------------|---------------------|-------------------------------|---------------|
|                       |              |                     | *T. dubius* origin | *T. porrifolius* origin           |
| 2601 Pullman, Washington | 4           | 24                  | E                            | E             | –                          |
|                       | 7           | 24                  | E                            | E             | 2 × C<sub>6</sub><sup>du</sup> |
|                       | 8           | 24                  | 3 × B<sup>du</sup>          | 1 × C<sup>po</sup> | 1 × C<sub>6</sub><sup>du</sup> |
| 2602 Palouse, Washington | 0-3-4       | 23                  | E                            | 1 × C<sup>po</sup> | –                          |
|                       | 0-3-10      | 24                  | 1 × C<sub>6</sub>            | 3 × E<sup>po</sup> | –                          |
|                       | 4           | 24                  | 1 × D<sub>du</sub>          | 3 × E<sup>po</sup> | –                          |
| 2603 Rosalia, Washington | 33A         | 24                  | 1 × E<sub>du</sub>          | 3 × E<sup>po</sup> | –                          |
|                       | 33B         | 24                  | E                            | E             | –                          |
|                       | 2           | 24                  | E                            | E             | –                          |

The parental origins of the chromosomes were determined by GISH. Chromosome nomenclature of homeologous groups (A–F) followed Ownbey and McCollum [25] and Pires et al. [24]. Superscript letters indicate the genome origins of the chromosomes, du = *T. dubius*, pr = *T. pratensis*. Chromosomes carrying translocations are indicated by naming the chromosome according to the genomic origins of the centromeres, followed by the genome origins of the translocated segments. E indicates chromosome set as expected from the diploid parents.

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**Table 2. Karyotype organisation of individuals of *T. miscellus* from two populations (localities of each population are indicated in column 1).**

| *T. miscellus* population | Plant number | Chromosome No. (2n) | Nature of karyotype imbalance | Translocations |
|---------------------------|--------------|---------------------|-------------------------------|---------------|
|                           |              |                     | *T. dubius* origin | *T. pratensis* origin              |
| 2604 Moscow, Idaho        | 7            | 24                  | 3 × E<sup>du</sup>          | 1 × E<sup>pr</sup> | A<sup>du/pr</sup>           |
|                           | 2605 Pullman, Washington | 3       | 24                  | E                            | E             | –                          |
|                           | 14           | 24                  | 1 × D<sub>du</sub>          | 3 × E<sup>po</sup> | –                          |

The parental origins of the chromosomes were determined by GISH. Chromosome nomenclature of homeologous groups (A–F) followed Ownbey and McCollum [25] and Pires et al. [24]. Superscript letters indicate the genome origins of the chromosomes, du = *T. dubius*, pr = *T. pratensis*. Chromosomes carrying translocations are indicated as in Table 1. E indicates parental chromosome set as expected from the diploid parent.

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there was a considerable reduction in the number of *T. dubius* units (to ∼100 copies/diploid) from an expectation of ∼700 units (typical number for any given tetraploid population). An analysis of this individual’s karyotype (Figure 2) revealed that the loss in copy number can be accounted for by a reduction in the size of the rDNA locus on chromosome Adu. The sibling to this plant, 2603-33B, does not show this rDNA copy number deletion. Therefore, the locus size reduction in 2603-33A may have arisen from a deletion event occurring as a consequence of meiotic instability in the parent.

Despite the relatively small size (perhaps only 100 copies/diploid of the 45S rDNA unit) of the *T. dubius* 45S rDNA locus, it usually dominates rRNA expression in leaf material of *T. mirus* [23]. Even in the 2603-33A individual, with extremely reduced numbers of 45S rRNA genes of *T. dubius* origin, the locus accounts for 97% of the rRNA transcripts (see also Table 3). Nevertheless, in root-tip metaphases a secondary constriction is observed on chromosome Dpo, suggesting some transcription of the *T. porrifolius* locus occurs in that tissue, perhaps to compensate for the reduced 45S rDNA gene copy number when there is a high demand for ribosomes in metabolically active meristematic cells. In plant 2603-33B, with substantially higher numbers of *T. dubius* derived 45S rRNA genes, the units of *T. dubius* origin are transcribed, and those rDNA units of *T. porrifolius* origin are silent [23]. This is reflected in the lack of a secondary constriction at the rDNA locus on chromosomes Adu and Dpo in this individual (Figure 2, Table 3). The presence of secondary constrictions correlates strongly with levels of decondensation at interphase and almost certainly reflects transcriptional activity at the preceding interphase. Two individuals of *T. mirus* (2602-4 and 2602-0-3-4) show different decondensation of the two Adu homologues, probably reflecting genetic or epigenetic differentiation between the two homologues.

In population 2605 of *T. miscellus*, all individuals investigated show only partial dominance in the expression of rDNA units of *T. dubius* origin over those of *T. pratensis* origin [23]. This is also seen

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**Table 3.** Genomic origin of decondensed rDNA compared with previously published data from Kovarik et al. [22] and Matyasek et al. [23] using genomic–cleaved amplified polymorphic sequence (g-CAP) and reverse transcription-cleaved amplified polymorphic sequence (RT-CAP) to determine the genomic origin of rDNA sequences in the genome and the cDNA, respectively.

| Population          | Figure Number | Plant number | decondensed rDNA | condensed rDNA | Parental origin of genomic rDNA units determined by g-CAPS (% of *T. dubius* origin units) | Parental origin of expressed rDNA units determined by RT-CAPS, (% of *T. dubius* origin rRNA transcripts) |
|---------------------|---------------|--------------|------------------|---------------|------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------|
| *T. mirus*, 2601 Pullman, Washington | Figure 2D (interphase only shown) | 0-4 | 2×A<sub>D</sub> | 2×A<sub>P</sub> | 35 | 99 |
| | | | | | 2×D<sub>P</sub> |
| | | | | | 2×D<sub>P</sub> |
| | Figure 2A | 0-7 | 2×A<sub>D</sub> | 2×A<sub>P</sub> | 28 | 94 |
| | | | | | 2×D<sub>P</sub> |
| | Figure 2A | 0-8 | 2×A<sub>D</sub> | 2×A<sub>P</sub> | 29 | 96 |
| | | | | | 2×D<sub>P</sub> |
| *T. mirus*, 2602 Palouse, Washington | 0-3-4 | 1×A<sub>D</sub> | 2×A<sub>P</sub> | 40 | 34 |
| | | | | | 1×A<sub>P</sub> |
| | Figure 2C | 0-3-10 | 2×A<sub>D</sub> | 2×A<sub>P</sub> | 41 | 4 |
| | | | | | 2×D<sub>P</sub> |
| | Figure 2C | 0-4 | 1×A<sub>D</sub> | 2×A<sub>P</sub> | 47 | 50 |
| | | | | | 2×D<sub>P</sub> |
| *T. mirus*, 2603 Rosalia, Washington | Figure 2B | 33A | 2×D<sub>P</sub> | 2×A<sub>P</sub> | 6 | 97 |
| | | | | | 2×A<sub>P</sub> (loci of reduced size) |
| | Figure 2C | 33B | 2×A<sub>D</sub> | 2×A<sub>P</sub> | 27 | 97 |
| | | | | | 2×D<sub>P</sub> |
| | | | | | 2×D<sub>P</sub> |
| | 2 | 2×A<sub>D</sub> | 2×A<sub>P</sub> | 27 | 99 |
| | | | | | 2×D<sub>P</sub> |
| *T. miscellus*, 2604 Moscow, Idaho | 7 | 2×A<sub>D</sub> | – | 23 | 100 |
| | | | | | 2×A<sub>P</sub> |
| *T. miscellus*, 2605 Pullman, Washington | 3 | 2×A<sub>D</sub> | – | 25 | 54 |
| | | | | | 2×A<sub>P</sub> |
| | 14 | 2×A<sub>D</sub> | – | nd | nd |
| | | | | | 2×A<sub>P</sub> |

Ns–FISH image not shown

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Figure 4. Pollen meiosis in synthetic *T. mirus*. (A,B) Feulgen staining showing in (A) 12 regular bivalents and (B) one quadrivalent (arrow) and bivalents in close association with each other and a nucleolus. (C–H). Fluorochrome colours and GISH probes are as in Figure 2. (C–D) Note chromosome A\textsuperscript{du} in association with the nucleolus. Two bivalents, one from the *T. dubius* genome (yellow) and one from the *T. porrifolius* genome (orange), overlapped (arrow), perhaps occurring as a multivalent. (E–F). Note the rDNA and the bivalents carrying these genes associated with the nucleolus. A quadrivalent with two chromosomes of *T. dubius* origin and two of *T. porrifolius* origin occurring in a ring (arrow). (G–H) Note two bivalents (A\textsuperscript{du} and D\textsuperscript{po}) with rDNA associated with the nucleolus and the A\textsuperscript{por} bivalent with condensed rDNA unassociated with the nucleolus. N = nucleolus. Scale bar (C–H) is 10 μm.
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in the occurrence of secondary constrictions on both rDNA-carrying chromosomes Adu and Apr (Figure 3, Table 3).

An analysis of 45S rDNA evolution in *Nicotiana* polyploids indicates that parental loci are initially maintained in young polyploids, although the sequences within a locus may be subject to concerted evolution, and over time frames of >1 million years individual loci are lost [30]. In *Tragopogon* polyploids we do not know which of the different karyotype variants will survive selection and become fixed. Perhaps over longer evolutionary timescales interlocus homogenisation and new rDNA variants will occur and spread across all rDNA loci.

Our study shows that more than one pathway can lead to non-Mendelian inheritance of rDNA units in allotetraploids: (i) elimination or amplification of repeats within an array can occur without changes in locus number (as in the case of 45S rDNA locus in 2601-33A); (ii) a change in the number of rDNA-bearing chromosomes (and loci) without any material change in the number of genes at each locus (as with the 5S rDNA locus—Figure 6); or (iii) a combination of both, although this situation was not detected in this study.

**Meiotic irregularities**

The chromosome multiplication step of polyploidy is thought to establish species isolation barriers between the newly formed polyploid and its diploid parents, whilst providing a homologous partner for each of the chromosomes. However, analyses of newly synthesised allopolyploids reveal that early-generation individuals are often infertile, or have highly reduced fertility, due to problems with meiosis including irregular pairing of homologous chromosomes [31,32]. Selection for fecundity in synthetic polyploids is associated with generation-by-generation increased fertility [32]. Nevertheless even after many thousands of years of evolution, meiotic irregularities can still occur, as observed in *Triticum aestivum* (wheat), an allohexaploid where meiotic misdivision has been exploited in the formation of wheat aneuploid lines [33,34].

An analysis of meiosis in the newly synthesised *Tragopogon* allotetraploids revealed the frequent occurrence of multivalents (Figure 4). Such aberrant pairing patterns may result in imbalanced chromosome contribution in subsequent generations as well as intergenomic translocations. Both abnormalities were observed in several of 12 synthetic (*S*1 generation) *T. minus* plants analysed at metaphase. One plant was 2n = 23 and lacked a *T. dubius* chromosome and one plant carried a non-reciprocal translocation from a *T. porrifolius* chromosome to chromosome Cdu (Figure 5E). Multivalents were also observed at a low frequency in natural populations of *T. minus* and *T. miscellus* [13] and in the few *F*2 plants resulting from diploid *F*1 hybrids of *Tragopogon* [35]. Meiotic abnormalities can cause unequal segregation of homoeologous chromosomes and are a likely driver of the chromosomal imbalance between genomes observed in both *Tragopogon* allotetraploids. The two sibling plants of *T. minus* 2603-33 had different karyotypes: one had a balanced karyotype of 12 chromosomes from each parental genome (2603-33B), while the other (2603-33A) was monosomic for chromosome Fdu and trisomic for chromosome Epo. Given these data, it is likely that the parent plant had a balanced karyotype of 12 chromosomes from each parental genome (2603-33B), while the other (2603-33A) was monosomic for chromosome Fdu and trisomic for chromosome Epo. Given these data, it is likely that the parent plant had a balanced karyotype of 12 chromosomes from each parental genome (2603-33B). Multivalents were also observed at a low frequency in natural populations of *T. minus* and *T. miscellus* [13] and in the few *F*2 plants resulting from diploid *F*1 hybrids of *Tragopogon* [35]. Meiotic abnormalities can cause unequal segregation of homoeologous chromosomes and are a likely driver of the chromosomal imbalance between genomes observed in both *Tragopogon* allotetraploids. The two sibling plants of *T. minus* 2603-33 had different karyotypes: one had a balanced karyotype of 12 chromosomes from each parental genome (2603-33B), while the other (2603-33A) was monosomic for chromosome Fdu and trisomic for chromosome Epo. Given these data, it is likely that the parent plant had a balanced karyotype of 12 chromosomes from each parental genome (2603-33B), while the other (2603-33A) was monosomic for chromosome Fdu and trisomic for chromosome Epo. Given these data, it is likely that the parent plant had a balanced karyotype of 12 chromosomes from each parental genome (2603-33B).

Multivalent pairing can arise either through (1) synapsis and recombination between homoeologous chromosomes in meiosis I,
Figure 6. Genomic analysis of 5S rDNA repeats. (A) Southern blot hybridisation of 5S rDNA probe to TaqI-digested genomic DNA of the progeny of 2602-0-3 individual and diploid parental accessions. (B) Quantitative representation of the parental 5S gene families determined by a phosphorimager scanning of radioactivity signals on blots (three independent experiments).

or (2) synapsis between chromosomes carrying intergeneric translocations. Four out of 12 mitotic karyotypes in plants from natural populations had intergeneric translocations visible at mitotic metaphase following GISH (Tables 2 and 3), and additional smaller translocations, not resolvable by GISH, may also be present. Nevertheless, the quadrivalent indicated in Figure 4F (see arrow) does not show intergeneric translocations at the resolution obtained using GISH. Our analyses of plants from different populations, although sample sizes were small, showed that aberrant chromosome numbers were more prevalent for certain chromosome types. For example, chromosome A was not involved in any of the aneuploidy events detected, whereas chromosome F accounted for almost 40% of all subgeneric chromosome imbalances. There might be several explanations for these results. First the homology between F-type chromosomes may be larger than between other chromosomes. It is likely that the greater sequence and morphological similarities between homeologous chromosomes, the more likely there will be homeologous and multivalent pairing. In support of this, Nicolas et al. [36] observed in *Brassica napus* haploid hybrids that chromosomes with the highest synteny had the highest frequency of homeologous pairing. Nevertheless, in *T. mirus*, the presence of a 5S rDNA locus on Fpo but not on Fdu indicates some divergence between the homeologues. Secondly, all chromosomes may be equally likely to form multivalents, but aberration in copy number of some chromosomes, e.g. homeologous group A, may not be favored by selection.

The surprisingly high incidence of trisomy and monosomy in highly fertile plants with 2n = 24 suggests that “compensating trisomy” may be operating in natural populations of both *Tragopogon* allotetraploids. Compensating trisomy was first described by Blakeslee [37] to refer to a situation in which the loss of a normal chromosome is compensated by the presence of the two arms in new translocated associations (secondary chromosomes). That concept was extended to include the replacement of the primary chromosome by two tertiary chromosomes, or by a secondary and a tertiary chromosome [38]. Compensatory trisomy has been reported in *Datura stramonium* from progeny of a plant exposed to radium [37,38], and from crops, including from Poales [39,40] and tomato [41]; all were produced experimentally [42]. The putative compensatory trisomy observed here may be an example from natural plants.

The allotetraploids *T. mirus* and *T. miscellus* formed from their diploid progenitors within the last 80 years (perhaps even within the last 60 years) and given their biennial habit, the number of generations to present is likely to be less than 40 [8]. The long-term outcome of meiotic irregularities in these species is not easily predicted. On the one hand, the genomic imbalance will reduce fitness and perpetuate cycles of meiotic irregularities. This may lead to a cascade of reduced fitness, generation upon generation. Such a phenomenon was observed in synthetic *Brassica* allotetraploids maintained by single seed descent [43]. In the wild, cryptic karyotypic instability manifest by aberrant ratios of homeologous chromosomes might ultimately lead to a slow reduction in fitness and ultimately extinction. Perhaps this accounts for the loss of some local populations of both *T. mirus* and *T. miscellus* [8] and of some recently formed *Senecio* polyploid populations [6]. Nonetheless, it is noteworthy that despite meiotic irregularities in the initial synthetic *S0* plants and subsequent *S1* generations, pollen fertility and seed set were generally high in the synthetic *Tragopogon* polyploid lines (Tate et al., in prep.). In addition, selection will favour the most fertile individuals, likely to be those with the most regular chromosome pairing. Thus, if the population can expand through early bottlenecks of reduced fertility, the derived populations are likely to be more fertile with regular bivalent pairing. Certainly in well-established allotetraploids (10^4–10^5 myrs old), e.g. of *Nicotiana* and *Triticum*, no major imbalances in chromosome numbers or the distribution of chromosomes to subgenomes are normally observed [44,45].

**Genomic instability**

The angiosperm genome is characterised by its plasticity to genetic change, including large-scale chromosome number changes, aneuploidy and polyploidy [4]. However, it may be significant that in *Tragopogon* polyploids, all the imbalances in parental chromosome dosages between individuals occurred within a near-regular karyotype of 2n = 24 (one exception at 2n = 23). Given the frequency of plants showing genomic imbalance we might expect to find more plants with unexpected
chromosome numbers. Perhaps there is selection against plants that deviate from 2n = 24 and against those that are nullisomic for a particular chromosome, or perhaps our sample size was too small to find a representative range of abnormalities.

GISH data revealed that in two out of three individuals of T. miscellus and four of nine individuals of T. mirus, there is an imbalance in the parental contribution of the chromosomes. However, the imbalance observed resulted from monosomy or trisomy, and no individual was nullisomic for a particular chromosome. An analysis of 10 genes in T. miscellus using genomic and cDNA CAPS revealed that 65% of individuals displayed losses of one of the two homeologues and a further 5% of individuals showed silencing of one of the two homeologues in leaves [12]. The missing alleles were interpreted as sequences that had been stochastically eliminated from the T. miscellus genome. The results here may suggest that chromosome loss or non-reciprocal translocations may contribute to the loss of alleles, although we did not find any example of a homozygous deletion for a chromosome segment. The differential expression of rDNA on the A′h chromosomes of two T. mirus individuals (Figure 2) points to epigenetic or genetic heterozygosity between the homologous chromosomes.

Large-scale genetic changes caused by parental genome imbalance will influence the inheritance of genetic markers, which will in turn influence the transcriptome, proteome and metabolome. Clearly genetic analyses of young or synthetic polyploids require in-depth cytogenetic studies to assess the contribution that chromosomal changes play in the inheritance of genetic markers. Unfortunately in recent years that work has seldom been standard practice. Such cytogenetic data are clearly needed even if chromosome counts appear regular [see also 43], since a deeper analysis of the genome and chromosome substructure can reveal substantial chromosome dosage deviation from expectation.

Materials and Methods

Plant material

Seeds of Tragopogon were collected from natural populations in Idaho (ID) and Washington (WA) (USA) (Table 1) and planted either in a greenhouse at the Department of Botany, University of Florida, or in field plots at the Institute of Biophysics, Academy of Sciences of the Czech Republic, Brno. Root tips from young, healthy, vigorously growing plants were harvested and placed in ice cold and saturated aqueous 2M 8-hydroxyquinoline (Sigma-Aldrich Company Ltd, Poole, Dorset, UK). After 60 min incubation on ice, the roots were fixed in ethanol-acetic acid (3:1) at room temperature overnight (several washes) and stored in the same solution at −20°C until use. Developing anthers of 1 mm length or less, that contain diplotene nuclei, were excised from young buds of length 1 cm or less. Root tip and meiotic material were fixed in freshly prepared 3:1 ethanol: glacial acetic acid for two days. Root tip material was then transferred to 90% ethanol at −20°C for long-term storage.

Making synthetic Tragopogon allotetraploids

Descriptions of the methods used to generate synthetic polyploidy lines are given in detail in a separate paper reporting the formation and availability of these lines for research (Tate et al. in prep). Briefly, numerous repeated T. dubius × T. portuloides crosses were made, and seeds from successful crosses germinated on moist filter paper. The chromosome number was doubled to resynthesise polyploids that closely resemble T. mirus by placing seedlings with fully emerged cotyledons in 0.1% or 0.25% colchicine solution overnight. After washing with water for two-three days, the seedlings were transferred to 2.5” pots with soil (and grown in the glasshouse at the University of Florida under standard conditions). Plants coded N197-3.132-1 and N197-4.90-1 were used for meiotic analyses (S0 generation) and root tips of selfed progeny (S1 generation) analysed in root tip mitosis, metaphases of plants coded 134-16-3 and 73-14 are shown.

Feulgen staining of meiotic cells

For meiotic squashes, small developing inflorescences (1 cm or less in length) were collected from the greenhouse and fixed in 3:1 (as above, for root tips). Individual anthers were then removed and macerated in 60% glacial acetic acid, stained in aceto-orcein [46], spread under a coverslip by warming over a naked flame, and observed using a Zeiss Photomicroscope III. At least 15 meiotic cells per plant were scored.

Preparing cell spreads for in situ hybridisation

Chromosome preparations for FISH, using either cloned probes or total genomic DNA probes (called GISH) were made using modifications of established methods [47,48]. Briefly, root tips or developing anthers were digested in 0.5% (w/v) cellulase Onozuka R-10 (Apollo Scientific Ltd, Stockport, Cheshire, UK), 0.3% (w/v) pectolyase Y23 (MP Biomedicals, Solon, Ohio, USA) and 0.3% (w/v) drieselase (Sigma-Aldrich Company Ltd., Poole, Dorset, UK) for 28 min and transferred to 1% citrate buffer for 2 h. The meristematic cells behind the root cap were isolated in a drop of 60% acetic acid and squashed onto a glass slide. For meiotic preparation, fixed anthers were dissected and meiotic cells gently dispersed into a drop of 60% acetic acid, a coverslip applied and gently warmed. Coverslips were removed following freezing with dry ice.

In situ hybridisation

Fluorescence in situ hybridisation followed standard protocols, Figures 2–4, Leitch et al. [48], Figure 5, Telgmann-Rauber et al. [47]. Genomic DNA from T. dubius, T. portuloides, and T. portuloides for labelling by GISH was extracted using DNeasy Plant mini kit (Qiagen Ltd, Crawley, West Sussex, UK) following manufacturer’s instructions. Genomic DNA was labelled with biotin-16-dUTP (Sigma-Aldrich Company Ltd., Poole, Dorset, UK) or digoxigenin-11-dUTP (GE Healthcare, Chalfont St Giles, Buckinghamshire, UK) using nick translation following standard protocols. The probe against 5S ribosomal DNA (rDNA) was prepared by amplifying the gene using primers described in Fulneck et al. [49] and biotin-16-dUTP-labelling protocol as described in Leitch et al. [48]. The probe against 45S rDNA was the clone pTa71, which includes the 18- 26S rDNA subunit isolated from Triticum aestivum, which was labelled with digoxigenin-11-dUTP as described in Leitch et al. [48]. Briefly, slides were denatured in 70% (v/v) formamide in 2× SSC (0.3 M sodium chloride, 0.03 M sodium citrate) at 70°C for 2 min and the hybridisation mix added (4 μg/ml labeled probes and 50% (v/v) formamide, 10% (w/v) dextran sulphate, 0.1% (w/v) sodium dodecyl sulphate in 2× SSC). In situ hybridisation was carried out overnight at 37°C, after which the slides were given a stringent wash (20% (v/v) formamide in 0.1× SSC at 42°C). Sites of probe hybridisation were detected using 20 μg/ml−1 fluorescein-conjugated anti-digoxigenin IgG (GE Healthcare, Chalfont St Giles, Buckinghamshire, UK) and 5 μg/ml−1 Cy3-conjugated avidin (Roche Pharmaceuticals, Lewes, East Sussex, UK) in 4× SSC containing 0.2% (w/v) Tween 20 and 5% (w/v) bovine serum albumin. Chromosomes were counterstained with 2 μg/ml DAPI (4,6-diamidino-2-phenylindole, Sigma Aldrich Company Ltd, in 4× SSC) and stabilised in Vectashield medium (Vector Laboratories Ltd, Peterborough, UK) prior to data acquisition using either: 1) Leica DMR2 epifluorescent microscope fitted with an Orca ER camera and
Open Lab software® (Improvement, Coventry, UK) (Figures 2–4) or 2 Olympus BX51 epifluorescent microscope using Olympus Microsuite 5 software® (Olympus America Inc, Center Valley, PA, USA) (Figure 5). The images were analysed with Adobe Photoshop® version 7 and treated for contrast variation and uniform brightness only. At least 5 mitotic or meiotic cells per plant were scored with each probe used.

Southern hybridisation

Tel restriction enzyme digestion was carried out on genomic DNA extracted from leaves using standard protocols [50] with modifications as in [51]. After fractionation in 1% agarose by gel electrophoresis, the DNA was transferred to a Hybond N membrane (GE Healthcare, Chalfont St Giles, Buckinghamshire, UK). The 32P-labelled 5S rDNA probe was from a 120-bp XhoI/EcoRI fragment of 5S rDNA cloned from N. tabacum [49]. Probe hybridisation was conducted under high-stringency conditions in the Church-Gilbert hybridisation buffer at 65 °C overnight. The radioactivity signals were quantified by phosphorimager scanning (Storm, GE Healthcare, UK).

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Author Contributions

Conceived and designed the experiments: DS PS AK AL. Performed the experiments: KJY DS JT RM HS ZL AL. Analyzed the data: KJY PS RM HS AK CP ZL AL. Contributed reagents/materials/analysis tools: DS PS AK CP AL. Wrote the paper: DS PS JT AK CP AL.

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