Copy number variation of ribosomal DNA and Pokey transposons in natural populations of Daphnia

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Abstract

Background: Despite their ubiquity and high diversity in eukaryotic genomes, DNA transposons are rarely encountered in ribosomal DNA (rDNA). In contrast, R-elements, a diverse group of non-LTR retrotransposons, specifically target rDNA. Pokey is a DNA transposon that targets a specific rDNA site, but also occurs in many other genomic locations, unlike R-elements. However, unlike most DNA transposons, Pokey has been a stable component of Daphnia genomes for over 100 million years. Here we use qPCR to estimate the number of 18S and 28S ribosomal RNA genes and Pokey elements in rDNA (rPokey), as well as other genomic locations (gPokey) in two species of Daphnia. Our goals are to estimate the correlation between (1) the number of 18S and 28S rRNA genes, (2) the number of 28S genes and rPokey, and (3) the number of rPokey and gPokey. In addition, we ask whether Pokey number and distribution in both genomic compartments are affected by differences in life history between D. pulex and D. pulicaria.

Results: We found differences in 18S and 28S gene number within isolates that are too large to be explained by experimental variation. In general, Pokey number within isolates is modest (< 20), and most are gPokey. There is no correlation between the number of rRNA genes and rPokey, or between rPokey and gPokey. However, we identified three isolates with unusually high numbers of both rPokey and gPokey, which we infer is a consequence of recent transposition. We also detected other rDNA insertions (rInserts) that could be degraded Pokey elements, R-elements or the divergent PokeyB lineage recently detected in the Daphnia genome sequence. Unlike rPokey, rInserts are positively correlated with rRNA genes, suggesting that they are amplified by the same mechanisms that amplify rDNA units even though rPokey is not. Overall, Pokey frequency and distribution are similar in D. pulex and D. pulicaria suggesting that differences in life history have no impact on Pokey.

Conclusions: The possibility that many rDNA units do not contain a copy of both 18S and 28S genes suggests that rDNA is much more complicated than once thought, and warrants further study. In addition, the lack of correlation between rPokey, gPokey and rDNA unit numbers suggests that Pokey transposition rate is generally very low, and that recombination, in combination with natural selection, eliminates rPokey much faster than gPokey. Our results suggest that further research to determine the mechanisms by which Pokey has escaped complete inactivation by its host (the usual fate of DNA transposons), would provide important insights into transposon biology.

Keywords: Daphnia, transposons, Pokey, ribosomal RNA genes, copy number variation, quantitative PCR

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Background

Transposable elements (TEs) are segments of DNA that can move or transpose around the genome [1]. Despite the fact that some have been co-opted for cellular functions by their host [2], TEs are generally considered to be detrimental because they can disrupt function when they insert into or near genes, or promote ectopic recombination, which can lead to chromosome rearrangements [3]. Moreover, their transposition may have energy costs [3] and the epigenetic mechanisms used by the host to control their expression can also alter gene function [4]. Even so, transposons are ubiquitous in eukaryotes and have become basic genomic components like exons, introns, telomeres and centromeres [5,6].

Ribosomal (r)DNA is a multigene family composed of repeated units each containing an 18S, 5.8S and 28S rRNA gene, and spacers (Figure 1). Due to the highly repetitive nature of rDNA, it usually evolves in a concerted fashion such that rDNA units are more similar within than between species [7]. The primary mechanisms responsible for this phenomenon are thought to be unequal crossing over and gene conversion [7]. Given the constant turnover of rDNA units caused by this recombination, and the strong purifying selection that operates on the rRNA genes, it is somewhat surprising that some TEs insert specifically into rDNA [7]. On the other hand, it has been argued that its repetitive nature makes rDNA an ideal niche for TEs. For example, recombination can increase the number of rDNA units and thus provide new insertion sites, or it can remove inactive TEs to reopen sites into which active elements can insert. Moreover, rRNA genes are highly transcribed providing many opportunities for expression of TE-encoded genes [8]. TEs that specifically target rDNA typically insert into either the 18S or the 28S genes, which make the genes non-functional [9,10]. However, the number of rDNA units usually exceeds the minimum required for host viability, and thus insertion of TEs into some rDNA units may have little impact on host fitness [8,11]. Indeed, some species are able to recognize and preferentially deactivate rDNA units containing insertions [12].

The most extensively studied rDNA-specific TEs are the Class I non-LTR retrotransposons, R1 and R2 [11], which are common in arthropods. R2 and other related lineages have also been found in diverse animal phyla, including Chordata, Echinodermata, Plathyhelminthes, Rotifera and Cnidaria [9,10,13]. These TEs typically undergo long periods of stable, vertical inheritance and diverge congruently with their hosts [11].

Pokey is a Class II DNA transposon in the piggyBac superfamily that ranges in size from 4 to 10 kb and contains the terminal inverted repeats (TIRs) characteristic of this element class [14,15]. Many copies of Pokey encode a transposase whose ability to excise the element has been confirmed in a yeast excision assay [15]. Pokey was first identified in the rDNA of the cladoceran crustacean, Daphnia pulex [16] and is widespread in the subgenus Daphnia [17,18]. The only other taxa in which Pokey elements have been discovered are the silk moth, Bombyx mori [19] and the tunicate, Ciona savignyi [20]. Pokey is unique among DNA transposons in that it targets a specific TTAA site in the 28S genes of Daphnia (Figure 1) where it has undergone stable, vertical inheritance for millions of years [17]. Moreover, this TTAA site is only 4 nucleotides (nt) from the insertion site of R2, and 64 nt from the insertion site of R1. However, unlike R-elements, which tend to specialize on rDNA (but see [9]), Pokey also inserts into many other locations throughout the Daphnia nuclear genome [21-23], although only a single copy has been reported from the nuclear genomes of B. mori and C. savignyi.

Daphnia are freshwater crustaceans (Cladocera, Crustacea) that inhabit standing water from shallow, ephemeral puddles to deep stratified lakes. They typically reproduce by cyclic parthenogenesis in which production of direct-developing embryos by apomictic parthenogenesis alternates with the meiotic production of ephippial (diapausing) eggs that require fertilization. Males are produced apomictically and sex-determination is environmental. Populations that inhabit temporary bodies of water that either dry up or freeze during the year are re-established from ephippial offspring annually. In contrast, adults can persist year-round in permanent ponds and lakes, and recruitment of individuals from ephippial eggs may be sporadic. In addition, some lineages have completely lost the capacity for sexual reproduction and produce their ephippial eggs apomictically (obligate parthenogenesis).
Species in the *Daphnia pulex* complex typically inhabit ponds and small unstratiﬁed lakes that lack ﬁsh. However, one species in this complex, *Daphnia pulicaria*, has invaded large stratified lakes in North America [24], and is able to tolerate predation by ﬁsh by taking refuge in the cold hypolimnion during the day [25]. The transition to lake habitats has led to substantial changes in physiology and life history [26,27]. Even so, *D. pulicaria* produces viable hybrids with other members of the *D. pulex* species complex. Hybrids typically occur in ponds or disturbed, intermediate habitats, and reproduce by obligate parthenogenesis [28]. Despite this hybridization, lake populations of *D. pulicaria* remain genetically [29] and ecologically [26,27] distinct from the other species in the complex.

The occurrence of Pokey in both rDNA (rPokey) and other genomic locations (gPokey) in *Daphnia* provides a unique opportunity to study its distribution in different genomic compartments and under different modes of reproduction (with and without meiosis). For example, using the PCR-based technique TE Display [30], Valizadeh and Crease [21] and Schaack *et al.* [23] compared the relative load of gPokey in cyclically and obligately parthenogenetic populations of *D. pulex* in North America. The results of both studies show that cyclical individuals carry, on average, more elements than obligate individuals, which is consistent with predictions about the effect of breeding system on TE dynamics [31]. However, it is not possible to estimate rPokey or rRNA gene number with TE Display, so the dynamics of rPokey have not yet been determined. On the other hand, Averbeck and Eickbush [32] measured R1 and R2 number in the rDNA of replicate laboratory lines of *Drosophila melanogaster* and found that R2 number remained relatively constant through time, but changes in R1 number were strongly correlated with changes in the overall size of the rDNA locus, which varied from 140 to 310 units after 400 generations of laboratory culture. Moreover, transposition of full-length elements accounted for most of the change in R1 number. This level of rRNA gene variation was also observed in replicate, clonally-propagated lines of *Daphnia obtusa*, in which the haploid number of 18S genes varied from 53 to 233 after only 90 generations [33].

In this study, we use quantitative PCR (qPCR) to measure the number of 18S and 28S rRNA genes, as well as rPokey and gPokey in 16 pond populations (43 isolates) of *D. pulex* and 6 lake populations (26 isolates) of *D. pulicaria* (Figure 2, Additional file 1). Our objectives are to estimate the correlation between (1) the number of 18S and 28S genes, (2) the number of 28S genes and rPokey, and (3) the number of rPokey and gPokey. In addition, we ask if life history differences between *D. pulex* and *D. pulicaria* impact Pokey frequency and distribution. We expect a positive correlation between 18S and 28S gene number if most rDNA units are complete, although this has rarely been tested. Moreover, if rates of transposition and/or elimination by the host vary in response to the number of available insertion sites, we expect to observe a positive association between rPokey and 28S gene number. We also expect a positive correlation between rPokey and gPokey number if rates of transposition and/or rates of element elimination are similar both inside and outside of rDNA. Finally, because rates of recruitment of sexually-produced offspring differ between pond populations, which are re-established each year from ephippial hatchlings, and lake populations, which often undergo extended periods of clonal reproduction, there may be more opportunities for transposition in pond populations and thus higher Pokey load.

**Results**

We calculated the haploid number of rRNA genes (18S and 28S) and Pokey relative to two single-copy reference genes, *Tif* (a transcription initiation factor) and *Gtp* (a member of the RAB subfamily of small GTPases) [33]. The number of 28S was estimated in two regions: downstream of the Pokey insertion site (total 28S or t28S) and across the Pokey insertion site (uninserted 28S or u28S, Figure 1), which allowed us to determine if there are insertions present in that position that would not amplify with our Pokey primers (for example, R2). In addition, we estimated the number of rPokey and total Pokey elements in the genome (Figure 1, Additional file 2).

We expect the number of one reference gene relative to the other to be close to 1, and this is often the case (Table 1, Additional file 2). However, values of *Tif* relative to *Gtp* (TG ratio) vary from 0.65 to 1.32 across the 69 *Daphnia* isolates, with a mean of 0.90 for *D. pulex* and 0.92 for *D. pulicaria* (Table 1). One explanation for
the extreme values is that there are three copies of one gene instead of the two expected, in which case we expect a ratio of 0.67 (3Gtp) or 1.5 (3Tif). Further analysis (Additional file 3) suggests that there are three copies of Gtp in one isolate (TG ratio = 0.65) and three copies of Tif in the four isolates (TG ratios ≥ 1.25). Estimates of gene number based on the duplicated gene were adjusted by multiplying them by 1.5 in these isolates.

Variation in rRNA gene number
The haploid number of 18S varies from 94 to 489.5 across all 69 Daphnia isolates (Table 1), and the mean value is not significantly different between the two species (t = -0.26, df = 50, P = 0.79). An even wider range of variation was observed for 28S (88 to 724.5), but again, the mean value does not differ significantly between species (t = -0.26, df = 50, P = 0.79). Similarly, there is no significant difference between the mean number of 18S and 28S within each species (D. pulicaria t = -1.52, df = 43, P = 0.14; D. pulex t = -1.64, df = 74, P = 0.10).

The estimates of 18S and 28S number within each isolate are significantly correlated (Table 2, Figure 3), but the slope of the line generated by plotting them relative to one another is 1.26, which is substantially higher than the expected value of 1. Moreover, 18S and 28S numbers are significantly different in 61% of the 69 isolates with 28S exceeding 18S in 76% of these cases (Additional files 2 and 4).

### Table 1 Pokey and rRNA gene number in Daphnia from North America

| Species   | Number of populations | Number of isolates | Tif:Gtp | 18S mean/range | 28S mean/range | gPokey | rPokey | rinserts | % Pokey in 28S | % rinserts |
|-----------|----------------------|--------------------|---------|----------------|----------------|--------|--------|----------|---------------|------------|
| D. pulex  | 16                   | 43                 | 0.90    | 221.0          | 260.0          | 2.1    | 9.6    | 21.1     | 17.2          | 6.5        |
|           |                      |                    | 0.65 to | 94 to 121      | 88 to 124      | 0.21   | 3.9    | 0.41     | 0.43          | 0.53       |
|           |                      |                    | 1.31    | 489.5          | 724.5          | 4.5    | 113.5  | 0.4       | 0.03          | 0.01       |
| D. pulicaria | 6                   | 26                 | 0.92    | 215.1          | 266.1          | 0.95   | 9.8    | 8.19     | 7.1           | 4.6        |
|           |                      |                    | 0.70 to | 97 to 109      | 0 to 45        | 0.45   | 149    | 0.0       | 0 to 792      | 0.0        |
|           |                      |                    | 1.32    | 444            | 654.5          | 113.5  | 0.0    | 0.24     | 0.0           | 0.0        |
| D. pulicaria | 6                   | 23                 | 0.93    | 2173           | 273.9          | 0.9    | 9.8    | 16.9     | 21.2          | 5.2        |
|           |                      |                    | 0.70 to | 97 to 109      | 0 to 7.5       | 0.45   | 45.4   | 0.17     | 0 to 57.7     | 0.0        |
|           |                      |                    | 1.32    | 444            | 654.5          | 113.5  | 0.0    | 0.24     | 0.0           | 0.0        |

1. Number of Tif genes relative to Gtp genes.
2. Denotes total 28S genes.
3. Denotes Pokey in 28S genes.
4. Denotes Pokey in other genome sites. Calculated as (total Pokey - rPokey).
5. Denotes other rDNA inserts, calculated as (t28S-u28S-rPokey) where u28S is uninserted 28S genes.
6. Calculated as (rPokey/total Pokey × 100).
7. Calculated as (rPokey/t28S × 100).
8. Results obtained after removing three D. pulicaria isolates with high Pokey load (see text).

### Table 2 Correlations between Pokey and rRNA gene number in Daphnia from North America

| Species   | X-axis | Y-axis | slope   | y-intercept | R²     | P-value | Figure |
|-----------|--------|--------|---------|-------------|--------|---------|--------|
| All isolates | 18S    | 28S    | 1.257   | -12.69      | 0.690  | 0.000   | 3      |
| D. pulex   | 28S    | u28S¹  | 0.850   | 15.71       | 0.976  | 0.000   | 5a     |
|           | 28S    | rinserts² | 0.150  | -17.84      | 0.556  | 0.000   | 5a     |
|           | 28S    | rPokey³ | -0.00002 | 2.14       | 0.000  | 0.995   | 5a     |
|           | gPokey⁴ | rPokey | -0.016  | 2.29        | 0.0005 | 0.884   | 4b     |
| D. pulicaria | 28S    | u28S   | 0.907   | 3.26        | 0.981  | 0.000   | 5b     |
|           | 28S    | rinserts | 0.104  | -12.80      | 0.247  | 0.0001  | 5b     |
|           | 28S    | rPokey | -0.011  | 9.54        | 0.020  | 0.493   | 5b     |
|           | gPokey | rPokey | 1.475   | -7.48       | 0.290  | 0.005   | 4b     |
|           | gPokey | rPokey | 0.123   | 1.62        | 0.031  | 0.420   | 4b     |

1. Denotes uninserted 28S rRNA genes.
2. Denotes other rDNA inserts calculated as (t28S-u28S-rPokey) where u28S is total 28S genes.
3. Denotes Pokey in 28S genes.
4. Denotes Pokey in other genome sites. Calculated as (total Pokey - rPokey).
5. Results obtained after removing three D. pulicaria isolates with high Pokey load (see text).
Variation in Pokey number

Pokey number in 28S (rPokey) varies from 0 to 44.5, with a mean of 2.1 for D. pulex and 6.6 for D. pulicaria (Table 1, Figure 4a), but these differences are not significant (t = 1.96, df = 26, P = 0.06). The higher mean for D. pulicaria is due to three isolates with 23.5, 40 and 44.5 rPokey (Figure 4a, Additional file 2). If we exclude these isolates, the mean decreases to 2.7 (Table 1). rPokey number is 7.5 or fewer in all other D. pulicaria isolates, and three have none. Similarly, we did not detect rPokey in three D. pulex isolates. Moreover, the haploid rPokey number is only 0.5 in a total of eight isolates (Additional file 2), which means they have a single copy among all of their 28S genes. We tested these results in 26 isolates using end-point PCR, and they correspond...
with six exceptions in which we were not able to amplify rPokey from isolates that have it based on qPCR. The qPCR estimate of rPokey is 0.5 in three of these isolates, 1 in one isolate and 2 in two isolates.

Pokey number outside of rDNA (gPokey) ranges from 4 to 24 with a mean of 9.6 in D. pulex and 9.5 in D. pulicaria (Table 1, Figure 4a), and this difference is not significant (t = -0.09, df = 40, P = 0.93). The same three D. pulicaria isolates with high rPokey number also have high gPokey number (8.5, 10.5, 24). If we exclude these isolates, mean gPokey decreases to 8.9, which is lower than that in D. pulex (9.6) but not significantly different. Overall, gPokey is present in all isolates, and, with the exception of the three unusual D. pulicaria isolates, it is much more numerous than rPokey (Table 1, Figure 4a).

On average, less than 28% of the variation in 28S gene insertion number is not likely to be the only explanation between t28S and u28S (as described above). However, experimental variation is due to experimental variation and rPokey does not bind to our qPCR primers. We refer to these as other rDNA inserts (rInserts). rInsert number ranges from 0 to 113.5 in D. pulex (mean = 21.1) and from 0 to 76 in D. pulicaria (mean = 14.9, Table 1), and the means are not significantly different (t = -1.06, df = 60, P = 0.29). On average, rInserts occur in less than 10% of 28S (Table 1). It is likely that some of the difference between t28S and u28S is due to experimental variation, and indeed this likely explains most cases where u28S exceeds t28S (as described above). However, experimental variation is not likely to be the only explanation because the slope of the line generated by plotting u28S relative to t28S is significantly lower than the expected value of 1 in both species, but the correlation between t28S and u28S is > 97% (Table 2, Figure 5).

There is a strong positive correlation (approximately 50%) between t28S and rInserts in both species (Table 2), although rInsert number increases at a substantially slower rate (D. pulex slope = 0.15, D. pulicaria slope = 0.10, Figure 5) than does u28S. Moreover, these results are very different from the situation with rPokey, which shows no correlation with t28S (Table 2, Figure 5) and is present in very low numbers in all but three D. pulicaria isolates (Figure 4).

### Variation in 28S gene insertion number

In general, the total number of 28S genes (t28S) exceeds the number of uninserted 28S genes (u28S), as expected. However, u28S is higher than t28S in 20 isolates. In addition, the sum of u28S plus r28S exceeds t28S in four isolates (Additional file 2). In these cases, we used the qPCR estimate of t28S and calculated u28S as (t28S-r28S).

Differences in number between t28S and u28S (Table 1, Additional files 2 and 4) suggest that some genes contain insertions other than Pokey, or their Pokey elements do not bind to our qPCR primers. We refer to these as other rDNA inserts (rInserts). rInsert number ranges from 0 to 113.5 in D. pulex (mean = 21.1) and from 0 to 76 in D. pulicaria (mean = 14.9, Table 1), and the means are not significantly different (t = -1.06, df = 60, P = 0.29). On average, rInserts occur in less than 10% of 28S (Table 1). It is likely that some of the difference between t28S and u28S is due to experimental variation, and indeed this likely explains most cases where u28S exceeds t28S (as described above). However, experimental variation is not likely to be the only explanation because the slope of the line generated by plotting u28S relative to t28S is significantly lower than the expected value of 1 in both species, but the correlation between t28S and u28S is > 97% (Table 2, Figure 5).

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### Comparison of rRNA gene and Pokey number based on qPCR and genome sequencing

Colbourne et al. [34] estimated the haploid number of rDNA units from the D. pulex genome sequence by comparing the depth of sequence coverage in the trace files across the entire rDNA repeat and the average single-copy gene. The estimate they reported is 435. In addition, Schaack et al. [22] searched the annotated genome sequence (which does not include rDNA) for Pokey using BLASTN and RepeatMasker. Their estimates

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**Figure 5 Correlation between total 28S genes and genes with and without insertions**

(a) *Daphnia pulex*. (b) *Daphnia pulicaria*. u28S genes denotes uninserted 28S genes. rPokey is inserted in 28S genes. The number of other rDNA insertions (rInserts) was calculated as (t28S-u28S-rPokey) where t is total 28S genes.

### Table 3 Pokey and rRNA gene number in two clonal lines of Daphnia pulex

| Gene Region | Isolate | 18S | t28S | u28S | rPokey | gPokey | rInserts | Tif. Gtp |
|-------------|---------|-----|------|------|--------|--------|----------|---------|
| Log50-11    | 301     | 540.5 | 445  | 2.5  | 37     | 93     | 0.84     |
| Log50-12    | 245.5   | 362.5 | 349  | 4    | 25.5   | 9.5    | 0.88     |

1. Log50 is the isolate from which the D. pulex genome sequence was generated [34], Log50-11 and Log50-12 are clonally-propagated lines derived from Log50 [22].
2. Denotes total 28S genes.
3. Denotes uninserted 28S genes.
4. Denotes Pokey in 28S genes.
5. Denotes Pokey in other genome sites. Calculated as (total Pokey - rPokey).
6. Denotes other rDNA inserts, calculated as (t28S-u28S-rPokey).
7. Number of Tif genes relative to Gtp genes.
based on these two approaches are 35 and 123, respectively.

In 2004, Schaack et al. [22] started a large number of clonally-propagated lines of the Log50 D. pulex isolate (whose genome was sequenced) from the lab culture that was established from the original female, collected in 2000. We obtained DNA extracted from two of these lines (Log50-11 and Log50-12) in 2006, approximately 40 generations after they were initiated, and estimated their Pokey and rRNA gene number using qPCR to compare them with estimates based on the genome sequence. The number of 18S is lower than 28S in both isolates (Table 3) with a ratio of 1:1.8 for Log50-11 and 1:1.5 for Log50-12. The mean number of 18S and 28S is 304 (Log50-12) and 421 (Log50-11), which is well within the range of variation observed within clonal D. obtusa lines over 90 generations [33]. Moreover, the Log50-11 estimate is very close to the value generated from the genome sequence (435).

Our estimates of gPokey number (37 and 25.5, Table 3) are also very similar to the estimate (35) obtained from the BLASTN search of the genome sequence. However, all three estimates are substantially lower than the one obtained from the genome sequence using RepeatMasker (123), suggesting that the similarity criteria used for the BLASTN search identified similar elements to the ones we detected with tPCR. Overall, our qPCR estimates are very similar to those obtained from the genome sequence.

**Discussion**

**Variation in rRNA gene number**

Variation in physiological responses to environmental heterogeneity has been linked with variation in rDNA copy number in diverse organisms (reviewed in [35]). Moreover, Paredes et al. [36] recently showed that differences in rDNA copy number have a significant effect on the level of expression of thousands of other genes in the Drosophila genome, which further reinforces the idea that rDNA copy number variation may be an important source of phenotypic and regulatory variation in natural populations. This is especially relevant for Daphnia, which is becoming an important model system for the study of genetic and physiological responses to environmental variation since publication of the genome sequence and development of other genomics tools [34].

In this study, we detected substantial variation in rRNA gene number within and between natural populations of D. pulex and D. pulicaria. The mean (approximately 220 for 18S, approximately 245 for 28S) and range is similar in both species with no values < 80 observed. However, the highest values are up to eight times larger than the smallest, which means that the diploid number of rRNA genes can vary by more than an order of magnitude among individuals of the same species. Such a large range of variation has been observed in natural populations of a diverse array of organisms, including plants, arthropods and vertebrates [35] and thus, Daphnia are not unusual in this regard.

Using computer simulations, Zhang et al. [8] showed that rDNA locus size is primarily a function of the rate of sister chromatid exchange, which creates high levels of variation among individuals and large rDNA loci. In contrast, high rates of interchromosomal exchange tend to reduce variation among individuals, and reduce the overall size of the rDNA locus. This model suggests that the highly variable number of rRNA genes in D. pulex and D. pulicaria results from higher rates of intrachromosomal than interchromosomal exchange, and this is indeed consistent with previous studies of rDNA variation in Daphnia [33,37,38]. Although rates of rDNA recombination have not been estimated in D. pulex or D. pulicaria, a rate of 0.02 to 0.06 events per generation was obtained in the congeneric species, D. obtusa [33], which is at the high end of rates estimated for rDNA in other organisms (10^{-2} to 10^{-5} [33]).

It is generally assumed that rDNA loci primarily contain canonical units consisting of one copy of each rRNA gene plus spacers (Figure 1), but this has rarely been tested. Based on this assumption, we expected our estimates of 18S and 28S number to be very similar within isolates, and they often are (Additional files 2 and 4). However, there are also a substantial number of isolates in which one gene (usually 28S) significantly outnumbers the other (Additional files 2 and 4) suggesting that experimental variation is not likely to account for all these differences. This conclusion is supported by the discovery of unusual rDNA configurations in humans. For example, Caburet et al. [39] used fluorescent in situ hybridization (FISH) and Southern blotting to show that up to one-third of the rDNA units in human fibroblast cell lines consist of head-to-head and tail-to-tail palindromic rearrangements of 18S and 28S. Moreover, these noncanonical units sometimes occur in clusters, which is consistent with their amplification by unequal sister chromatid exchange between rDNA units. In addition, Zafiropoulosa et al. [40] used qPCR to measure the number of 18S, 5.8S and 28S in adipose tissue samples from humans and found significant differences between genes within individuals. They concluded that these differences were a consequence of unequal recombination events initiating between the 18S and 5.8S and, subsequently, eliminating part of an rDNA unit. These studies suggest that our observation of large differences in 18S and 28S number in some Daphnia isolates may not be an artefact of qPCR analysis, and thus warrants further study.

**Variation in Pokey number**

On average, we found only 2 to 3 rPokey elements per haploid genome (maximum of 12 with three exceptions
discussed below, Table 1). We also found isolates in which \( r_{Pokey} \) is completely absent, which is also the case in previous studies [21,41]. In contrast, \( g_{Pokey} \) is present in all isolates that have been analyzed in this and previous studies [21,23,41]. Indeed, over 75% of \( Pokey \) elements are located outside of 28S genes in most isolates. Thus, even if all the genes containing \( Pokey \) insertions were eliminated from rDNA, active \( g_{Pokey} \) elements could eventually "recolonize" it. However, despite the higher number of \( g_{Pokey} \), they are also fairly constrained; there are usually less than 10 copies per haploid genome. These results are similar to those obtained by Valizadeh and Crease [21] using TE Display. They estimated \( Pokey \) number per diploid genome to be from 1 to 12 in 83 isolates of \( D. pulex \) from the Midwest US and Ontario. However, these are likely to be underestimates because Valizadeh and Crease [21] used an annealing temperature of 55°C. Vergilino [18] analyzed \( Daphnia \) isolates using TE Display, but used an annealing temperature of 50°C. We analyzed a few isolates from his study using an annealing temperature of 55°C, and obtained about half as many fragments as he did (data not shown). If the \( g_{Pokey} \) estimates from Valizadeh and Crease [21] were doubled, they would be even closer to the ones we obtained with qPCR (Table 1).

The low number of both \( r_{Pokey} \) and \( g_{Pokey} \) in most isolates is consistent with the hypothesis that \( Pokey \) activity is generally low. Recombination among rDNA units further increases the rate at which \( r_{Pokey} \) is eliminated, and can eventually eliminate it altogether in some isolates. However, if \( Pokey \) has been completely inactive for a substantial period of time in either species, we would not expect to find any \( r_{Pokey} \), although \( g_{Pokey} \) could persist and even go to fixation by drift in some sites if it did not have deleterious effects on gene structure or function. Nevertheless, we identified three isolates with unusually high numbers of both \( r_{Pokey} \) and \( g_{Pokey} \), and indeed, these isolates are responsible for the positive correlation between them in \( D. pulex \) (Figure 4b). It is unlikely that the high \( r_{Pokey} \) number in these isolates is a consequence of large changes in the size of the rDNA locus as the two isolates with 40 or more \( r_{Pokey} \) have less than 150 28S genes per haploid genome (Additional file 2). Moreover, such changes in rDNA would have little or no impact on \( g_{Pokey} \) number. Thus, the most likely explanation for the high \( Pokey \) number in these isolates is a recent increase in transposition activity.

There are several mechanisms that could explain the putative increase in \( Pokey \) activity in these isolates. For example, it has been shown that some TEs are activated during a response to stress ([42] and references within). Alternatively, mutations could have occurred in \( Pokey \) that increase its ability to transpose or allow it to avoid silencing by the host, or in the host that decrease its ability to silence \( Pokey \) [3,43]. A similar scenario was suggested by Eickbush and colleagues [12,44], who showed that levels of R2 transcription can vary up to 100-fold in natural populations of \( Drosophila \). They explained this by suggesting that R1 and R2 serve as foci for the formation of heterochromatin, which deactivates rDNA units. Because active rDNA units in \( Drosophila \) tend to occur in contiguous blocks [45,46], individuals are best able to silence these elements when they are clustered with one another in the rDNA array. Conversely, if they have been interspersed with uninserted, functional 28S genes by recombination, they are less likely to be silenced and thus show higher rates of transposition.

We do not know how \( Pokey \) elements are distributed in the rDNA of \( Daphnia \), but Glass et al. [47] suggested that they are likely to be clustered based on patterns of sequence variation among 28S with and without \( Pokey \) insertions. In contrast, FISH analysis of one isolate with an IGS and a \( Pokey \) probe suggests that its \( Pokey \) elements are dispersed throughout the rDNA array [34]. Thus, it is not clear if the relationship between element activity and distribution suggested for R-elements in \( Drosophila \) will also be the case for \( r_{Pokey} \) in \( Daphnia \). However, this could be tested by analyzing the same individuals (or their clonally produced offspring) using both FISH and qPCR.

**Variation in 28S gene insertion number**

Based on analysis of \( D. pulex \) rDNA cloned into a phage vector, Sullender [16] estimated that \( r_{Pokey} \) elements occupy approximately 10% of the 28S genes. Based on our qPCR analyses, this is a substantial overestimate. However, even though \( r_{Pokey} \) generally occupies about 1% of the 28S in an isolate, \( r_{Inserts} \) occupy 5 to 7% of these genes, on average. Furthermore, unlike \( r_{Pokey} \), \( r_{Insert} \) number is significantly correlated with t28S number (Table 2, Figure 5), suggesting that \( r_{Inserts} \) are amplified by the same mechanism that amplifies u28S, and/or their transposition rate increases as u28S number increases. This same pattern was observed by Averbeck and Eickbush [32] for R1 in the replicate lines of \( D. melanogaster \), although the slope of the line generated by plotting the total number of rRNA genes relative to the number of R1 insertions was 0.27, which is two to three times higher than our result of 0.10 for \( D. pulex \) (Table 2). Averbeck and Eickbush [32] argued that R2 insertions, whose number remained low (as does \( r_{Pokey} \) in \( Daphnia \)), were excluded from recombination events among rDNA units, while R1 elements were underrepresented but still included, which allowed them to increase as the size of
the rDNA locus increased (as do tIns in Daphnia). Even so, R1-inserted 28S did not increase at the same rate as u28S, suggesting that their amplification was somehow constrained. The simulation study of Zhang et al. [8] provides an explanation for this behavior. When simulations included elimination of chromosomes with low numbers of u28S by natural selection, recombination among rDNA units tended to increase u28S faster than inserted i28S.

Glass et al. [47] sequenced approximately 850 nt of 28S downstream of the Pokey insertion site in 20 isolates of D. pulex and found that variation was higher among genes with Pokey than those without. This is consistent with relaxed selection on these presumably non-functional 28S, which allows them to accumulate nt substitutions and short indels that would normally be deleterious. It is also consistent with the hypothesis that the presence of Pokey inhibits recombination between i28S and u28S. This inhibition may also explain why there is no correlation between rPokey and i28S (Table 2, Figure 5), and why selection is not as efficient at removing 28S genes with Pokey insertions, which are much more deleterious than a single nt substitution or short indel.

It is possible that some of the tIns we detected in Daphnia 28S are R1 and/or R2, but this seems unlikely. First, the insertion site for R1 is not located between the qPCR primers that span the Pokey TTAA insertion site. Second, we have not been able to amplify R2 (whose insertion site is 4 nt downstream of the TTAA) from some of the D. pulex and D. pulicaria isolates analyzed in this study using degenerate primers that have been used successfully in a wide range of arthropod species [48]. A more likely possibility is that the tIns we detected are degraded or divergent Pokey elements, and indeed, a second Pokey lineage has recently been identified in the D. pulex genome sequence [15]. Based on its similarity to a divergent lineage that was previously identified in D. obtusa [17], this lineage has been designated PokeyB. In addition, two types of miniature inverted-repeat transposable elements (MITEs) were also identified in the genome sequence. These MITEs are approximately 750 nt in length, and one of them (mPok1) has TIRs similar to the original Pokey lineage, which we now designate as PokeyA, while the other (mPok2) has TIRs similar to PokeyB [15].

Sequence alignment of the four groups of elements suggests that our qPCR primers will not amplify PokeyB or mPok2, but they should bind to mPok1. Thus, it is possible that some of the tIns are members of these two groups. This is supported by the fact that we have been able to amplify PokeyB and mPok2 from the 28S of some of our D. pulex and D. pulicaria isolates using a forward primer that is specific to them and a reverse primer in the 28S (data not shown). So far, we have not been able to amplify mPok1 from 28S, although it is usually present elsewhere in the genome.

Another possibility is that tIns are partial Pokey elements generated by recombination events that deleted part of the element and possibly part of the rDNA unit in which it resides. This would explain why numbers of PokeyA and 18S are similar for both Log50 isolates, but Log50-11 has approximately 240 more copies of 28S than 18S. These additional 28S could have inserts (Table 3), or they may be recombinants that did not amplify with the primers spanning the Pokey insertion site. The fact that Pokey contains sequences derived from the ribosomal IGS [14,15] supports the idea that aberrant rDNA units with (partial) Pokey insertions could have been created by recombination between rPokey and rDNA. If this is the case, we expect these aberrant rDNA units to be deleterious and thus eliminated by natural selection before they expand to such high number. On the other hand, they could persist within populations for a considerable period of time if (1) they are clustered, which could occur if they were amplified by unequal sister chromatid exchange between rDNA units that are offset by one or a few copies, or (2) if they are not transcribed, either because their promoters were deleted by recombination (as suggested in [39] for the human fibroblasts), or they ended up in a region of the rDNA array that is particularly “attractive” to the heterochromatinization machinery (as suggested in [44] for R1 and R2).

Whatever the nature of tIns in Daphnia 28S genes, it is clear that their dynamics are somewhat different from those of PokeyA. The latter behave like R2 in the replicate D. melanogaster lines [32], while the tIns behave more like R1. Averbeck and Eickbush [32] attributed the differences between these two elements to two factors; differences in rates of participation in recombination events and differences in transposition rate, which were estimated to be much higher for R1 than R2. If Daphnia tIns are indeed PokeyB or mPok2, we would expect them to have a higher transposition rate than PokeyA. We are in the process of testing this prediction using yeast excision assays, which have shown that the PokeyA transposase is active [15]. If PokeyB does have a higher excision rate, we would also expect it to be more numerous than PokeyA in other genomic locations as well as rDNA. We also predict that mPok2 will be more numerous in 28S than are full length PokeyA or PokeyB for two reasons. First, MITEs are often found to occur in higher number than the full-length elements whose transposase they use to move [49]. Second, the much shorter length of mPok2 may reduce or eliminate the bias against recombination between i28S and u28S. This could contribute to the
significant correlation between t28S and rInserts (Table 2, Figure 5). In order to test these predictions, we are now in the process of developing qPCR primers to quantify each of the Pokey and MITE groups separately, both in and outside of 28S genes.

Impact of life history variation on rDNA and Pokey dynamics

Previous studies [21,23,37,41,47] have shown that loss of sexual reproduction in obligately parthenogenetic lineages of *D. pulex* does impact both rDNA and gPokey dynamics. In this study, we compared two *Daphnia* species whose life histories differ in several ways, including timing and frequency of sexual reproduction, brood size, juvenile growth rate and life span [26]. Overall, patterns of rDNA and Pokey number variation are virtually indistinguishable in the two species (Table 1, Figures 4 and 5). Thus, it seems that their life history differences have little, if any, impact on Pokey dynamics, most likely because “a little bit of sex is nearly as good as a lot” [50]. The only exception is the three *D. pulicaria* isolates with unusually high rPokey and gPokey loads, which we suggest is a consequence of recent transposition activity. Further analysis of isolates from these populations would likely provide additional insights into Pokey dynamics.

Conclusions

It is clear that rDNA is very dynamic and much more complicated than once thought. In this study, we observed substantial variation in rDNA copy number among *Daphnia* isolates, which is consistent with previous studies suggesting that sister chromatid exchange is more frequent than interchromosomal exchange in *Daphnia* rDNA. Moreover, we sometimes found substantial differences in 18S and 28S number within isolates, which is not likely to be explained entirely by experimental variation and thus warrants further study. In general, there are less than 20 Pokey elements per haploid genome and most are gPokey. This suggests that transposition rates are generally very low and that recombination, in combination with natural selection, eliminates rPokey faster than gPokey. Even so, three isolates of *D. pulicaria* have unusually high numbers of both rPokey and gPokey, which we suggest is due to a recent increase in transposition activity. We also detected other rDNA insertions (rInserts) that could be degraded Pokey elements, R- elements or members of the divergent PokeyB lineage that was recently detected in the *D. pulex* genome sequence. Although rPokey number is not correlated with t28S number, rInsert number is, suggesting that they are amplified by the same mechanisms as rDNA units while rPokey is not. Overall, we observed no impact of life history differences between *D. pulex* and *D. pulicaria* on the dynamics of either rPokey or gPokey.

Methods

*Daphnia* samples and DNA extractions

A total of 69 *Daphnia* isolates collected from 16 ponds (*D. pulex*) and 6 lakes (*D. pulicaria*) were included in this study (Figure 2, Additional file 1). Ponds were sampled by skimming a dip net just under the surface of the water. Lakes were sampled by towing a net behind a small boat. Clonally-propagated lines were established from single females and grown in 200 mL of dechlorinated tap water at room temperature. Cultures were fed either live *Scenedesmus* or frozen *Nannochloropsis* algae (Landlocked Mariculture LLC, Toronto, Ontario, Canada). Genomic DNA was extracted from multiple individuals from each line using the phenol:chloroform method [51] or the AquaGenomics extraction kit with the manufacturer’s tissue protocol (MultiTarget Pharmaceuticals LLC, Salt Lake City, Utah, USA). DNA concentrations were estimated using a NanoDrop® ND-8000 spectrophotometer (Wilmington, Delaware, USA) and ranged from 1 to 1,500 ng/μL.

Lab isolates were identified as *D. pulex* or *D. pulicaria* by PCR-amplifying and sequencing an approximately 700 nt fragment of the mitochondrial NADH dehydrogenase 5 (ND5) gene [52]. The breeding system (cyclic or obligate parthenogenesis) of most lab-reared isolates was determined by examination of diapausing egg cases (ephippia) produced in the absence of males. While cyclical parthenogens often release empty ephippial cases unless the eggs have been fertilized, obligate parthenogens deposit eggs into ephippia even in the absence of males [53].

qPCR

We used the ΔC_T quantitative (q)PCR method to estimate haploid copy number by comparing the rate of amplification of a multicopy gene to that of two single-copy genes as in McTaggart *et al.* [33]. The copy number of 18S, t28S, u28S, rPokey, total Pokey, Tif and Gtp were estimated using seven pairs of primers (Figure 1, Additional file 5). We generated standard curves (Additional file 6) to validate each primer pair and determine its percent amplification efficiency (PAE).

qPCR reactions were 20 μL in volume, with 1X Power SYBR® Green PCR Master Mix (Applied Biosystems, Foster City, California, USA), 0.25 pmol of each primer, and approximately 10 ng of template. Reactions were run on the StepOnePlus™ Real-Time PCR System (Applied Biosystems). The PCR program was as follows: 95°C for 10 minutes, 40 cycles of 95°C for 15 sec and 60°C for 1 minute. After the 40 cycles were complete, a dissociation curve was created during one additional
cycle by increasing the temperature from 60°C to 95°C in increments of 0.3°C. All reactions were run in triplicate.

The baseline was set automatically by the StepOne v2.0 software (Applied Biosystems). The threshold was set based on amplicon size as larger amplicons bind more SYBR Green and thus produce more fluorescence. This has little effect when gene number is low, but the effect is substantial at high gene numbers. Thus, we set the threshold to 0.2 for the smallest amplicon of 50 bp, and calculated the threshold for larger amplicons as 0.2 × 2^(1-(50/length in bp)). If the standard deviation of the triplicate mean CT value was larger than 0.2, we excluded the most extreme replicate. However, if there was no clear outlier, we used all three CT values in the analysis.

We calculated gene number according to [33] with the PAE correction of [54] as 2^ΔCT where ΔCT is ((CTT × PAEmulticopygene) - (CT × PAEsingle-copygene)). We used all combinations of CTT values from multi-copy and single-copy gene triplicates to generate a total of 18 estimates ((3 × 3) + (3 × 3)) when all values were included (Additional file 7). These 18 estimates were averaged to give the mean and standard deviation of haploid copy number for each multi-copy gene in each isolate. The haploid numbers were rounded up or down to the nearest 0.5. We used Microsoft Excel (Richmond, Washington, USA) to do correlation and regression analyses, and t-tests. We used the sequential Bonferroni technique of Rice [55] to adjust the significance level (0.05) for t-tests comparing 18S and 28S number within isolates.

Additional material

Additional file 1: Population location and sample size. This PDF file provides sample size, latitude and longitude for the 22 *Daphnia* populations sampled.

Additional file 2: Haploid rRNA gene and Pokey number. This PDF file provides estimates (and standard deviation) of rRNA gene and Pokey number for all *Daphnia* isolates in this study. *Daphnia pulicaria* were collected from ponds (P) and *D. pulicaria* lakes (L).

Additional file 3: *Tif* and *Gip* cloning experiment. This PDF file describes the cloning and sequencing of *Tif* and *Gip* genes from four *Daphnia* isolates with a range of *TifGip* ratios. The purpose of this work was to determine if isolates with very low or very high *TifGip* ratios possess three alleles at one of the loci.

Additional file 4: Histograms of haploid rRNA gene and insertion number. This is a PDF file. (a) Haploid 18S and 28S gene number in each *Daphnia* isolate. Vertical lines are standard errors. Differences that are not significant after sequential Bonferroni correction are indicated by "ns". (b) Haploid number of rPokey and 28S with and without inserts in each *Daphnia* isolate. u28S are uninserted 28S genes, rPokey is inserted in 28S, rinserts are inserts other than rPokey in 28S. The number of rinserts was calculated as (total 28S-uninserted 28S-rPokey).

Additional file 5: qPCR primers. This PDF file provides sequences for qPCR primers, as well as the threshold value and the percent amplification efficiency (PAE) for each primer pair.

Additional file 6: Standard curve analysis for qPCR primers. This PDF file provides details of the standard curve experiments used to validate the qPCR primers, including a description of the plasmid clones used as templates.

Additional file 7: CTT values for all qPCR reactions. This Excel spreadsheet provides CTT values for all *Daphnia* isolates and qPCR amplicons. Replicates that were omitted from the analyses are highlighted in grey. A template for calculating gene number using the ΔCT method is also provided, with sample calculations for isolate P6.7.

**Abbreviations**

18S: 18S rRNA gene; 28S: 28S rRNA gene; bp: base pair; CTT: threshold cycle; ETS: external transcribed spacer; FISH: fluorescent in-situ hybridization; gPokey: Pokey elements found outside of rDNA; 28S-inserted 28S rRNA gene; IGS: intergenic spacer; ITS: internal transcribed spacer; kb: kilobase pair; LTR: long-terminal repeat; MITE: miniature inverted-repeat transposable element; nt: nucleotide; PAE: percent amplification efficiency; qPCR: quantitative PCR; rDNA: ribosomal DNA; rinserts: rDNA inserts other than rPokey; rPokey: Pokey elements found in rDNA; rRNA: ribosomal RNA; t28S: total 28S rRNA genes; TE: transposable element; TG ratio: Tif to Gip ratio; TIR: terminal inverted repeat; u28S: uninserted 28S rRNA gene.

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**Authors’ contributions**

SHCE collected and analyzed the data. TJC conceived the study and supervised the work. Both authors wrote the manuscript and approved the final version.

**Competing interests**

The authors declare that they have no competing interests.

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