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Genes Translocated into the Plastid Inverted Repeat Show Decelerated Substitution Rates and Elevated GC Content

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Abstract

Plant chloroplast genomes (plastomes) are characterized by an inverted repeat (IR) region and two larger single copy (SC) regions. Patterns of molecular evolution in the IR and SC regions differ, most notably by a reduced rate of nucleotide substitution in the IR compared to the SC region. In addition, the organization and structure of plastomes is fluid, and rearrangements through time have repeatedly shuffled genes into and out of the IR, providing recurrent natural experiments on how chloroplast genome structure can impact rates and patterns of molecular evolution. Here we examine four loci (psbA, ycf2, rps7, and rps12 exon 2–3) that were translocated from the SC into the IR during fern evolution. We use a model-based method, within a phylogenetic context, to test for substitution rate shifts. All four loci show a significant, 2- to 3-fold deceleration in their substitution rate following translocation into the IR, a phenomenon not observed in any other, nontranslocated plastid genes. Also, we show that after translocation, the GC content of the third codon position and of the noncoding regions is significantly increased, implying that gene conversion within the IR is GC-biased. Taken together, our results suggest that the IR region not only reduces substitution rates, but also impacts nucleotide composition. This finding highlights a potential vulnerability of correlating substitution rate heterogeneity with organismal life history traits without knowledge of the underlying genome structure.

Key words: GC content, genome structure, inverted repeat, plastome, rate heterogeneity.

Introduction

Rates of molecular evolution vary dramatically among organismal lineages and across genomes (Bromham and Penny 2003) and understanding what causes this rate variation is a fundamental topic in evolutionary biology (Lanfear et al. 2010). Past studies reporting on rate variation usually focused on establishing a correlation between substitution rates and those organismal traits that might potentially be affecting the supply of mutations (e.g., generation time or metabolic rate; Wu and Li 1985; Martin and Palumbi 1993; Smith and Donoghue 2008; Korall et al. 2010; Gaut et al. 2011; Lanfear et al. 2013), or the rate at which available mutations are fixed in the population (e.g., weakened purifying selection due to small population sizes; Woolfit and Bromham 2003, 2005). Establishing such correlations is important for understanding broad evolutionary processes in relation to certain life history traits. Few studies, on the other hand, have investigated whether there are other factors that might also influence evolutionary rates.

Plant chloroplast genomes (plastomes) generally comprise a pair of inverted repeat (IR) regions and two single-copy (SC) regions. The two IR copies are identical in sequence but run in opposite directions (fig. 1a), and their sequence identity is maintained by gene conversion (Birky and Walsh 1992). Biased gene conversion, whereby new mutations are preferentially corrected back to ancestral states (Birky and Walsh 1992), is hypothesized to be responsible for a significantly lower rate of nucleotide substitution in the IR compared to the SC region (Clegg et al. 1984; Wolfe et al. 1987; Wu and Chaw 2015; Zhu et al. 2016). If the IR region does indeed suppress nucleotide substitutions, it is expected that genes moving into or out of the IR (via expansion or contraction of the region) would experience corresponding substitution rate shifts—decelerating after entering the IR and accelerating after exiting it.

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Only a few studies have examined whether the IR region can influence gene evolutionary rates. Two of these focused on situations where genes in the IR were translocated to the SC. Perry and Wolfe (2002) discovered that in legumes, where the IR region has disappeared, the former IR genes experienced accelerated substitution rates. However, Lin et al. (2012) found that when ycf2 was translocated out of the IR in *Ginkgo biloba*, its substitution rate did not increase (although it is possible that this event was too recent for a rate difference to be detectable). More recently, Zhu et al. (2016) tested whether genes translocated into the IR from the SC exhibited rate changes by examining such translocation events across a representative sample of angiosperms, gymnosperms, and ferns, and found that substitution rates did indeed decrease after gene translocation into the IR.

These three studies used pairwise synonymous substitutions to measure rate variation, which ignores the role of phylogenetic history and is unable to incorporate nucleotide substitution models to yield more accurate rate estimates. Here we estimate rate variation in a phylogenetic context, and employ a likelihood ratio test to identify significant rate shifts. In addition, we examine whether translocation into the IR alters the selective environment experienced by plastid loci, and whether plastome structure changes cause shifts in GC content. We focus on ferns (the sister group to seed plants), where plastome rearrangements have inverted the IR orientation (fig. 1b) and translocated five loci—*trnH, psbA, ycf2, rps7*, and exons 2–3 of *rps12* (henceforth simply “*rps12*”)—from the SC region to the IR (Wolf et al. 2010). This expanded and inverted IR occurs in extant Schizaeales, Salviniales, and Angiopteris.
Results and Discussions

Rate Deceleration Following Translocation into IR

We compiled *psbA, ycf2, rps7, and rps12* sequence data from published plastomes (*trnH* and other tRNA genes were not included because of their short sequence lengths; table 1), and tested the fit of two clock models: a 1-rate model and a 2-rate model. The 1-rate model applies a single substitution rate across all branches of the phylogeny, whereas the 2-rate model allows one “in-IR” rate, when these loci are in the IR (blue branches in fig. 2a and b), and another “out-IR” rate, when they are in the SC (orange branches in fig. 2a and b). For all four translocated loci, the 1-rate model was rejected in favor of the 2-rate model (*P* < 0.0000001), and within the 2-rate model the “in-IR” rates were 2- to 3-fold slower than the “out-IR” rates (fig. 2c–f). This model-testing result indicates that substitution rates decreased following gene translocation into the IR, and is consistent with the findings of Zhu et al. (2016) and also a study by Li et al. (2011) that found the *trnH–psbA* intergenic spacer has reduced levels of sequence variation in Polypodiales (where it is located in the IR), compared to the other studied fern orders (where it is in the SC). Because the translocation of *rps7* and *rps12* into the IR occurred three times (fig. 2b) within ferns, we also tested a 4-rate model that allows three separate “in-IR” rates and one “out-IR” rate. For both loci, the 4-rate model fit significantly better than the 2-rate and 1-rate models (*P* < 0.0000001), and all three “in-IR” rates were slower than the “out-IR” rate (fig. 2d), a result that is again consistent with a rate deceleration for genes translocated into the IR.

To rule out the possibility that the rate deceleration observed in *psbA, ycf2, rps7, and rps12* is part of a plastome-wide phenomenon (e.g., a change in polymerase proof-reading efficiency; Parkinson et al. 2005), or perhaps the improved fit is due simply to the extra parameter accommodating noise in the data rather than being related to the translocation (Lanfear 2011), we examined other chloroplast genes that have not translocated into or out of the IR. Based on the 1-rate model, we first confirmed that fern IR genes have significantly lower substitution rates than the non-IR genes (*P* < 10−5; two-tailed t-test), a result consistent with past studies on other plant groups (Clegg et al. 1984; Wolfe et al. 1987; Wu and Chaw 2015; Zhu et al. 2016). Next we compared the 1-rate and 2-rate models as above, but for genes that did not move into or out of the IR. The majority of these genes showed no rate shift (i.e., the 1-rate model could not be rejected in favor of the 2-rate model), and if they did, the direction of the rate change was predominantly opposite to that seen in *psbA, ycf2, rps7, and rps12* (fig. 2c–f and supplementary table S1, Supplementary Material online). The only exception is *atpB*, although it experienced no translocation, it nonetheless showed a significant deceleration (fig. 2d). However, the rate change in *atpB* is only 1.2-fold, much lower than the rate change observed in the IR-translocated loci. This comparison demonstrates that the rate deceleration in our focal loci is exceptional among chloroplast genes, and is best explained by their translocation into the IR.

Rate Deceleration Is Not Likely Due to Selection at the Protein Level

Next, we tested whether the degree of rate deceleration differs between synonymous and nonsynonymous substitutions, which would suggest a change in selection pressures upon translocation into the IR. We used RELAX (Wertheim et al. 2011) and also a study by Li et al. (2011) that found the *trnH–psbA* intergenic spacer has reduced levels of sequence variation in Polypodiales (where it is located in the IR), compared to the other studied fern orders. Because the translocation of *rps7* and *rps12* into the IR occurred three times (fig. 2b) within ferns, we also tested a 4-rate model that allows three separate “in-IR” rates and one “out-IR” rate. For both loci, the 4-rate model fit significantly better than the 2-rate and 1-rate models. The majority of these genes showed no rate shift (i.e., the 1-rate model could not be rejected in favor of the 2-rate model), and if they did, the direction of the rate change was predominantly opposite to that seen in *psbA, ycf2, rps7, and rps12* (fig. 2c–f and supplementary table S1, Supplementary Material online). The only exception is *atpB*, although it experienced no translocation, it nonetheless showed a significant deceleration (fig. 2d). However, the rate change in *atpB* is only 1.2-fold, much lower than the rate change observed in the IR-translocated loci. This comparison demonstrates that the rate deceleration in our focal loci is exceptional among chloroplast genes, and is best explained by their translocation into the IR.
Decelerated Substitution Rates and Elevated GC Content in IR

Fig. 2.—Substitution rates before and after translocation into the IR. (A, B) The resultant rate assignment schemes of the 2-rate models for psbA and ycf2 (A) and rps7 and rps12 (B). The symbols on (B) branches mark the rate assignment scheme for the 4-rate model (see D). (C, D) Rate differences of all sampled chloroplast genes observed for the 2-rate or 4-rate model. "*" indicates genes in which the 1-rate model was rejected in favor of the 2-rate or 4-rate model ($P < 0.0000001$). The rate estimates from the 4-rate model of rps7 and rps12 are shown at the bottom of (D). The symbols correspond to those in (B). The rate unit is substitutions/site/million years. (E, F) The distributions of fold rate change. The rate deceleration that is observed in ycf2, psbA, rps7, and rps12, is unique among chloroplast genes, and is due to their translocation into the IR.
2015), a codon-based, branch-site random effects method, to test for any change in selective strength. We found that between the “in-IR” and “out-IR” branches, there is no significant difference in the nonsynonymous/synonymous substitution rate ratio (ω), suggesting that protein-level selection has little or no effect on the rate decrease. The exception is ycf2, in which selection relaxed along the “in-IR” branches (P < 0.0001; table 2). However, because relaxation of selection would tend to accelerate nucleotide substitution rates, and we still observed ycf2 deceleration, this indicates its rate decrease is not due to selection.

GC Content Increases upon Translocation into the IR
Another hallmark of the IR is its high GC content relative to the SC region. Recently Wu and Chaw (2015) showed that in cycads there are more A/T to G/C substitutions in the IR (compared to the SC), and they proposed a GC-biased gene conversion mechanism for the IR. If this model is correct, we would expect to see an increase in GC content once a genomic segment translocates into the IR, particularly at the third-codon positions and in noncoding regions, which are under reduced selective constraint. Our analysis of GC content corroborates this prediction—the third codon positions of psbA, ycf2, rps7, and rps12, as well as the tmH-psbA intergenic spacer, have a higher GC content when they are in the IR than when in the SC region (fig. 3). Interestingly, GC variation is less obvious (or opposite, in psbA) in the first and second codon positions (fig. 3), implying that selection reduces this bias by limiting the substitutions that get fixed at these positions (since they usually result in amino acid changes). It should be noted that ycf2 is the only locus with a significant GC increase at the second codon position, a result consistent with the relaxed selection we detected for it (table 2). To investigate whether the rate deceleration found above was an artifact of the GC content increase, we reanalyzed our focal loci with the third codon position removed. We found similar rate decreases, with no significant difference in the fold rate change (P=0.75, paired two-tailed t-test; supplementary table S2, Supplementary Material online), suggesting that the shifts in rate and in GC content are decoupled in the IR.

Conclusion
In this study, we demonstrated that when genes are translocated into the IR, their nucleotide substitution rates dropped significantly (2- to 3-fold), and that this deceleration is not shared with other nontranslocated chloroplast genes. In addition to rate deceleration, GC content increases following translocation, indicating that the IR affects both substitution rates and GC content. Our finding also points to plastome rearrangements that can result in rate heterogeneity among lineages. This has important implications for studies trying to use genomic data to correlate rate heterogeneity together with life history traits, as well as for dating evolutionary events (e.g., Schuettpelz and Pryer 2006; Rothfels and Schuettpelz 2014). Without the knowledge of genome structure, or modeling for possible hidden rate shifts, the evolutionary inferences could be grossly misleading.

Materials and Methods
We sampled at least one plastome from each of the 11 fern orders (except for Hymenophyllales; table 1). The genome data were downloaded from Genbank, and individual gene sequences were extracted. Because some plastomes are incomplete, we focused on the genes that are present across all the sampled plastomes; a total of 48 loci were included. We inferred multiple sequence gene alignments using MUSCLE (Edgar 2004) followed by manual inspection and adjustment. We used baseml (implemented in PAML; Yang 2007) to estimate nucleotide substitution rates under the 1-rate (global clock) and 2-rate models (see fig. 2a and b). For the baseml runs, we used a GTR+G substitution model, with a fixed topology and divergence times derived from the 25-nuclear-locus study of Rothfels et al. (2015). The 2-rate models were tested against the 1-rate model using likelihood-ratio tests.

To test for differences in selective pressure among the “in-IR” and “out-IR” branches, we used a codon-based model-testing framework implemented in RELAX (Wertheim et al. 2015), available on Datamonkey (Delport et al. 2010). The key parameter in RELAX is k, which controls the degree of nonsynonymous/synonymous substitution rate ratio (ω) differences between the “in-IR” and “out-IR” branches (ωin=ωoutk). In the null model, k is fixed at 1 (i.e., no difference in selective pressures between the in-IR versus out-IR branches), whereas in the alternative model, k is a free parameter. Values of k > 1 make ω for the in-IR branches more extreme—stronger positive selection or stronger purifying selection—whereas values of k < 1 make ω closer to 1 (i.e., relaxation of selection). Both null and alternative models assume three site categories of ω. Likelihood ratio tests are then used to compare the fit of the free-k versus fixed-k models.

We calculated relative GC content for each sequence as the GC content of the focal sequence divided by the average GC content of the plastome, in order to control for genomic background variation. We used a two-tailed t-test to investigate

| Locus | Selection Intensity k | P Value |
|-------|-----------------------|---------|
| psbA  | 1.02                  | 0.8223  |
| ycf2  | 0.42                  | <0.0001 |
| rps7  | 1.96                  | 0.2013  |
| rps12 | 0.77                  | 0.1178  |

Only ycf2 showed a significant signature of selection relaxation (k = 1).
whether there is any significant GC difference between the in-IR and out-IR sequences. All the alignments and the control files for running baseml are available on figshare (dx.doi.org/10.6084/m9.figshare.3483056.v2; dx.doi.org/10.6084/m9.figshare.3483059.v1).

Supplementary Material

Supplementary tables S1 and S2 are available at Genome Biology and Evolution online (http://www.gbe.oxfordjournals.org/).

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Literature Cited

Birky CW, Walsh J. 1992. Biased gene conversion, copy number, and apparent mutation rate differences within chloroplast and bacterial genomes. Genetics 130:677–683.

Clegg MT, Rawson JR, Thomas K. 1984. Chloroplast DNA variation in pearl millet and related species. Genetics 106:449–461.

Delport W, Poon AFY, Frost SDW, Kosakovski Pond SL. 2010. Datamonkey 2010: a suite of phylogenetic analysis tools for evolutionary biology. Bioinformatics 26:2455–2457.

Gao L, et al. 2013. Plastome sequences of Lygodium japonicum and Marsilea crenata reveal the genome organization transformation from basal ferns to core leptosporangiates. Genome Biol Evol. 5:1403–1407.

Gao L, Yi X, Yang Y-X, Su Y-J, Wang T. 2009. Complete chloroplast genome sequence of a tree fern Alsophila spinulosa: insights into evolutionary changes in fern chloroplast genomes. BMC Evol Biol. 9:130.

Kim HT, Chung MG, Kim K-J. 2014. Chloroplast genome evolution in early diverged leptosporangiate ferns. Mol Cells. 37:372–382.

Karol KG, et al. 2010. Complete plastome sequences of Equisetum arvense and Isoetes flaccida: implications for phylogeny and plastid genome evolution of early land plant lineages. BMC Evol Biol. 10:321.

Lanfear R, et al. 2011. The local-clock permutation test: a simple test to compare rates of molecular evolution on phylogenetic trees. Evolution 65:606–611.

Li F-W, et al. 2011. rbcL and matK earn two thumbs up as the core DNA barcode for ferns. PLoS One 6:e26597.

Lin CP, Wu CS, Huang YY, Chaw SM. 2012. The complete chloroplast genome of Ginkgo biloba reveals the mechanism of inverted repeat contraction. Genome Biol Evol. 4:374–381.

Lu J-M, Zhang N, Du X-Y, Wen J, Li D-Z. 2015. Chloroplast phylogenomics resolves key relationships in ferns. J Syst Evol. 53:448–457.

Fig. 3.—GC content before and after genes translocate into IR. (A) Third codon position and intergenic spacer. (B) Second codon position. (C) Third codon position. ** denotes P < 0.05 in 2-tailed t-test. Relative GC content calculated as GC content of the locus divided by the whole-genome GC content.
