Tracing the recombination and colonization history of hybrid species in space and time

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Hybrid speciation has long fascinated evolutionary biologists and laymen alike, presumably because it challenges our classical view of evolution as a ‘one-way street’ leading to strictly tree-like patterns of ancestry and descent. Homoploid hybrid speciation (HHS) has been a particularly interesting puzzle, as it appears to occur extremely rapidly, perhaps within less than 50 generations (McCarthy et al. 1995; Buerkle et al. 2000). Nevertheless, HHS may sometimes involve extended or repeated periods of recombination and gene exchange between populations subject to strong divergent natural selection (Buerkle & Rieseberg 2008). Thus, HHS provides a highly interesting setting for understanding the drivers and tempo of adaptive divergence and speciation in the face of gene flow (Arnold 2006; Rieseberg & Willis 2007; Nolte & Tautz 2009). In the present issue of Molecular Ecology, Wang et al. (2011) explore a particularly challenging issue connected to HHS: they attempt to trace the colonization and recombination history of an ancient (several MYA) hybrid species, from admixture and recombination in the ancestral hybrid zone to subsequent range shifts triggered by tectonic events (uplift of the Tibetan plateau) and climatic shifts (Pleistocene ice ages). This work is important because it addresses key issues related to the origin of the standing genetic variation available for adaptive responses (e.g. to climate change) and speciation in temperate species, which are topics of great current interest (Rieseberg et al. 2003; Barrett & Schluter 2008; de Carvalho et al. 2010).

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Wang et al. (2011) focus on three closely related conifer species. Conifers provide a very special setting for genetic studies, in which chloroplast (cp) DNA is inherited through the paternal and mitochondrial (mt) DNA through the maternal lineage. Extensive experience with the use of plant cp and mt genomes in molecular ecology (Vendramin et al. 1996; Duminil et al. 2002) implies that selected regions of these DNA molecules can easily be visualized in large numbers of individuals, which facilitates large-scale phylogeographic studies. Within the limits provided by uniparentally inherited genetic markers, this approach allowed Wang et al. to trace the colonization and recombination history of *P. densata* in space and time at a depth rarely (never?) seen in any other homoploid hybrid species system.

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Fig. 1 (a) The homoploid hybrid species *Pinus densata* in its high-altitude environment on the Tibetan plateau (up to 4200 m elevation). (b) *P. densata* is among the most dominant and ecologically successful homoploid hybrid species known to date. Photo credit: Jian-Feng Mao.
In previous studies, the authors have shown that *P. densata* is a homoploid hybrid between two other Asian pine species, *P. tabuliformis* and *P. yunnanensis* (Wang et al. 1990, 2001) (Fig. 1). The existence of a third, unidentified potential parental taxon was sometimes hypothesized in early studies (Wang et al. 2001), although a third species may not be required to explain observed allelic diversity considering that novel, private alleles frequently arise in hybrid zones (Schilthuizen et al. 1999). An important aspect of these earlier studies is that both inter- and intra-locus linkage disequilibrium (LD) in *P. densata* was found to be extremely low. This provides a good indicator for the cessation of gene flow between the homoploid hybrid taxon and its parents (Wang et al. 2001; Ma et al. 2006).

With respect to ecology, the clear elevational differences among the three taxa (*P. densata*: Tibetan plateau at 2700–4200 m; both parental species: lower elevations in north-central and south-western China) speak for strong ecological divergence with regard to temperature and other eco-variables correlated to altitude. In fact, the three species are significantly differentiated for numerous fitness-related ecological traits (Mao et al. 2009) and a recent niche modeling exercise revealed a distinct niche shift in the hybrid taxon (Mao & Wang 2011). This clear ecological divergence linked to elevation (made possible by the uplift of the Tibetan plateau initiated ca. 20 MYA ago) strongly suggests that ecological barriers were the primary drivers of HHS in *P. densata*, and this timeframe was confirmed by the coalescence times for several genetic loci. Geographic barriers may have contributed but are unlikely to be substantial in high-gene flow species such as wind-pollinated temperate trees (Lexer et al. 2005). A survey of reproductive fitness components also indicated the absence of strong fertility barriers between the hybrid and its parents (Mao et al. 2009), as one might expect for *Pinus* species. So even if minor chromosomal rearrangements are suggested by molecular cytogenetic work based on ribosomal DNA (Liu et al. 2003), their contribution to reproductive isolation among *P. densata* and its parents appears negligible.

In this study, Wang et al. greatly extend their earlier phylogeographic work (Song et al. 2003) to track the evolutionary history of this enigmatic hybrid taxon, based on an unprecedented sampling effort in the geographically challenging south-eastern Tibetan Plateau. At first sight, their results may seem inconspicuous: they find pronounced geographic patterns for maternally inherited, seed-dispersed mtDNA in the hybrid taxon (*P. densata*) and weaker structure for paternally inherited, pollen-dispersed cpDNA, as one might expect (Petit et al. 2005). Only a second, deeper look reveals the full picture of the mosaic-like colonization and hybridization history of this homoploid hybrid pine, sketched backward in time here.

Patterns of cpDNA sharing indicate that populations of *P. densata* have experienced pollen flow from both parents in different parts of their range. Notably, central populations of the hybrid taxon experienced pollen flow from its parent *P. yunnanensis* only recently, whereas the western portion of the hybrid species’ range is highly differentiated for cpDNA and has thus been isolated from both parents for much longer. Maternal mtDNA lineages provide clues on processes dating even further back in time: mtDNA variation in the hybrid taxon is structured into three major haplogroups, thus pointing to the presence of multiple differentiated lineages likely resulting from independent hybrid speciation events, as suspected previously (Wang et al. 2001). Greatly elevated mtDNA diversity and differentiation in the north-eastern range of *P. densata* points to the location of the ancestral hybrid zone from which this hybrid taxon emerged. The demographic stability of populations in this region is supported by the genetic signature inherent in mtDNA and cpDNA mismatch distributions, which are incompatible with recent population expansion.

Wang et al. also tested for mtDNA recombination using a four-gamete test (Hudson & Kaplan 1985) under simplifying assumptions. Recombination is known to occur in plant mt genomes (Houliston & Olson 2006) and has been shown in other conifers (Jaramillo-Correa & Bousquet 2005). Here, the authors used this prior knowledge in a clever way to gain further insights into the ancient hybridization history of this Asian hybrid pine. Assuming the absence of technical artefacts, they detected a likely recombination event between mtDNA haplotypes derived from each parental species, involving two ancient mitotypes found at high frequencies in the inferred ancestral hybrid zone. Thus, by combining information from two organellar genomes with contrasting modes of inheritance, Wang et al. were able to trace the evolutionary history of this hybrid pine, including identification of the putative ancestral hybrid zone and inference of recurrent westward colonization events across the Tibetan plateau. Inference of the colonization process represents in itself another highly interesting aspect of this study.

Examination of allelic diversities from east to west revealed several unexpected shifts in mtDNA haplotype frequencies, with haplotypes that were rare or even absent in the ancestral hybrid zone reaching high frequencies in more recently colonized westerly localities. These unexpected shifts were plausibly explained by the ‘surfing’ of rare alleles into newly colonized territories during range expansion (Klopfstein et al. 2006; Excoffier et al. 2009). The characteristic spatial patterns of haplotype groups, and of ‘surfing’ events of haplotypes within groups, were used to put forward the hypothesis that westward expansion of *P. densata* across the Tibetan plateau likely occurred in more than a single wave. Keeping in mind limitations in sample sizes, these findings may inspire other students interested in the origin and maintenance of biological diversity in mountain systems. Although the impact of mountain building processes on macroevolutionary patterns of diversity is increasingly well understood (Hoorn et al. 2010), we know little about its impact on exactly those population genetic processes that represent the actual ‘engine’ of divergence and speciation.

The biggest limitation of this study—the absence of information from nuclear genomic markers—also points to a potentially highly rewarding avenue for future research.
Future work on *Pinus densata* and other homoploid hybrid species should now turn to large-scale nuclear genomic surveys to trace the history of nuclear admixture and recombination in space and time. Knowledge of genomic distributions of recombination break points or junctions would enable molecular ecologists to infer the timing of the initial admixture and recombination events that triggered HHS and to unravel the migration episodes that have shaped today’s ‘genomic mosaics’ until reproductive isolation was (close to) complete. The basic analytical tools for this purpose have long been available (Fisher 1954; Chapman & Thompson 2002) and are currently being refined, driven primarily by progress in human medical genetics (reviewed by Buerkle & Lexer 2008; Winkler et al. 2010). Ultra High Throughput Sequencing (UHTS) methods to trace genome-wide recombination events in non-model species are now emerging as well (Baird et al. 2008; Hohenlohe et al. 2010). With the availability of new technologies affording ever greater genomic coverage and sequencing depth, it is likely that sufficient numbers of single nucleotide polymorphisms (SNPs) will soon be available for speciation genomic studies despite the complexity of *Pinus* genomes.

Recent verbal models and reviews of HHS (Jiggins et al. 2008; Nolte & Tautz 2009) suggest that this process is much less restricted than previously assumed (Buerkle et al. 2000). Also, there are multiple routes in addition to HHS through which hybridization and genetic exchange may impact on the ecology & evolution of wild species of animals and plants (Arnold 2006). Results on tree hybrid zones further indicate that the novel variation provided by genomic admixture not only affects the hybridizing species themselves but also the communities and ecosystems that depend on them (Whitham et al. 2006). Tracing the recombination and colonization history of homoploid hybrid lineages would thus seem like a worthwhile goal for current biology. This study on a Tibetan hybrid pine shows what can be achieved with traditional phylogeographic tools. We look forward to seeing new, exciting studies that address these issues with the complete ‘next generation’ tool box now available to students of the ecology and evolution of non-model species.

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References


