Letters to the Editor

Ancient Species Flocks and Recent Speciation Events: What Can Rockfish Teach Us About Cichlids (and Vice Versa)?

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The large assemblage of northeastern Pacific species of rockfish (genus Sebastes) has been proposed to be the modern result of an ancient explosive speciation event (Johns and Avise 1998). This radiation has been compared to the more recent radiation of cichlids in the Great African lakes (Greenwood 1991), and by analogy, the Sebastes assemblage was termed the “ancient species flock” (Johns and Avise 1998). An increasing number of molecular evolutionary studies focusing on cichlids from the Great African Lakes have uncovered unique modes of speciation. Sexual selection has been shown to play a crucial role in all three Lakes. In the oldest lake, Lake Tanganyka, sexual selection was evoked to explain the comparatively large genetic divergence observed between congeneric Tropehus species (Sturmbauer and Meyer 1992). Indeed, these species did not show any morphological differences besides coloration, although there was significant genetic divergence at the mitochondrial control region level [14.5% sequence divergence (Sturmbauer and Meyer 1992)]. On the other hand, haplochromine species from the most recent lake, Lake Victoria, and the species flock from the evolutionary intermediate lake, Lake Malawi, were found to exhibit a vast array of color patterns but little genetic divergence (Crapon de Caprona and Fritsch 1984; Meyer et al. 1990; Seehausen et al. 1997; McElroy and Kornfield 1990; McKaye et al. 1993; Deutsch 1997). Furthermore, lineage sorting was shown to be incomplete in the Malawi species flock, locally called mbuna, using mitochondrial control region sequences (Bowers et al. 1994; Kornfield and Parker 1997; Parker and Kornfield 1997). Recent microsatellite studies, however, show that mbuna species could be distinguished using fast-evolving molecular markers (Kornfield and Parker 1997; van Oppen et al. 1997; Markert 1998; Arnegard et al. 1999).

Thus, the relationship between speciation and divergence at the morphological and genetic level in cichlids (as in most organisms) is not simple (Meyer 1993). While few morphological differences besides coloration patterns were found in Victoria haplochromines, Malawi mbunas (within genera) and Tanganyka Tropehus species, levels of genetic divergence at the mitochondrial control region level were either large—14.5% sequence divergence for the genus Tropehus—or very limited—2.3 and 8.2% sequence divergence for the whole Victoria and Malawi species flocks, respectively. A similar situation appear to exits in the Sebastes species flock, where examples of species groups that exhibit little or no morphological difference also exist [i.e., Sebastes rubrivinctus and S. babcocki; S. rosenblatti, S. eos, and S. chlorosticus; S. carnatus and S. chrysomelas; S. helvomaculatus and S. simulator (Rocha-Olivares et al. 1999a,b)]. What levels of genetic divergence should we expect in these species? If these species groups have been maintained for an extensive period of time, as the idea of an ancient species flock would suggest, we

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should expect significant genetic divergence levels. Conversely, if the evolutionary origin of these groups is recent, we should expect these species to exhibit little genetic divergence.

In order to investigate speciating events in rockfishes, we have selected a pair of closely related species, Sebastes carnatus, the gopher rockfish, and Sebastes chrysomelas, the black-and-yellow rockfish. Gopher rockfish and black-and-yellow rockfish are morphologically and meristically identical (Phillips 1957; Miller and Lea 1972). Their color patterns, however, are different. Over an identical dark background, gopher rockfish have pink blotches, while black-and-yellow rockfish have yellow blotches. More importantly, ecological characteristics differ in these two species (Larson 1980a, b). Black-and-yellow rockfish are restricted mostly to water shallower than 10–16 m, while gopher rockfish are generally found in waters deeper than 16 m (Hallacher 1984; Larson 1980a, b). This difference in depth preference, however, does not result in different growth rates or significant differences in diet (Larson 1980a–c). After comparing depth segregation, competition, and habitat usage among these two species of rockfish, Larson (1980a–c) concluded that there was no evidence of social dominance between these fish and suggested that light intensity has a direct or indirect effect on the depth distributions of these species. Thus, gopher rockfish and black-and-yellow rockfish show similarities to cichlid species in several ways. Their morphological and meristic similarity is reminiscent of several Malawi mbuna species (which are, ironically, also called rockfish) (Ribbink et al. 1983; Bowers et al. 1994; Kornfield and Parker 1997). Furthermore, their segregation by depth is also similar to the ecological segregation that was observed in haplochromine Victoria cichlids (Goldsmith et al. 1990).

Considering the comparison between gopher rockfish and black-and-yellow rockfish sequence divergences with similar divergences in cichlids, we sequenced a 251-bp portion of the mitochondrial control region for 47 individuals (35 gophers and 12 black-and-yellows) which were sampled over their distribution range (Fig. 1). Control regions in these species were found to be fairly variable, as the average sequence divergence between gopher rockfish and black-and-yellow rockfish was 3.8%, and the haplotype diversity was 0.95 (40 haplotypes of 47 individuals). However, phylogenetic reconstructions using the neighbor-joining and the maximum-parsimony methods did not result in the separation of gopher rockfish and black-and-yellow rockfish into distinct clades (Fig. 2). Furthermore, the alternative constrained topology that would place gopher rockfish and black-and-yellow rockfish individuals into two separate monophyletic assemblages, which would require an additional 17 steps, was rejected by a Kishino and Hasegawa test ($p < 0.02$) and a T-PTP test ($p < 0.01$) (Kishino and Hasegawa 1989; Faith 1991). Therefore, our data indicate that Sebastes carnatus and S. chrysomelas lineages cannot be sorted using mitochondrial control region sequences.

Previous attempts to separate these two species using slower molecular markers such as allozymes (Seeb 1986) and nuclear ITS sequences (Hunter 1994) failed to find fixed differences between black-and-yellow rockfish and gopher rockfish. Taken together, all the evidence points toward two possible hypotheses. One possibility is that these two species are simply color morphs of a single species. Since their description in 1880 (Jordan and Gilbert 1880a, b), scientists have questioned the taxonomic status of these two species (Hubbs and Schultz 1933; Phillips 1957; Chen 1986). However, some elements negate this possibility. The color morph separation by depth is difficult to explain (Alesandrini 1997). Although phenotypic plasticity may be evoked, no intermediate color morphs have been described. An alternative explanation is that gopher rockfish and black-and-yellow rockfish are genuine species.

Since mitochondrial control regions are among the fastest molecular markers available, why can’t we separate black-and-yellow rockfish and gopher rockfish? One possibility is that there is currently active gene flow between species. While gene flow between black-and-yellow and gopher rockfish is possible, our data suggest that this is unlikely. Indeed, no haplotypes are shared between species, while several haplotypes are shared among individuals within a species. This is the case for black-and-yellow rockfish individuals CB2 and CB3 and for gopher rockfish individuals CB1 and BS1, individuals CB12 and BS10, individuals BS9 and CB8, and in-
individuals CB9, SB1, CB10, and PB1. An alternative hypothesis is that speciation occurred recently, from a large and genetically diverse ancestral population, in the absence of recent bottleneck at each speciation event (Avise 1994). Our data are compatible with this hypothesis, as the genetic diversity of each species (0.98 for G and 0.98 for BY) is similar to the genetic diversity of the combined species (0.95). Furthermore, if this hypothesis is correct, it is possible to estimate the minimum time of species divergence by determining the sequence divergence between the most closely related individuals of the two species. Sequence divergence between our two species ranges from 0.4 to 7.8%. Using currently accepted ranges for D-loop substitution rates [2 to 10%/Myr (Grant and Bowen 1998)] and assuming an absence of gene flow between species, the minimum divergence time between black-and-yellow rockfish and gopher rockfish is approximately 40,000 years.

The concept of the “ancient species flock” coined by Johns and Avise (1998) may suggest that a main burst of Sebastes speciation happened in the distant past and that we are now witnessing the results of this original "big
bang.” Our data show that speciation mechanisms may still be active in *Sebastes*. As speciation in marine organisms is still poorly understood (Palumbi 1992, 1994), it is difficult to pinpoint the mechanisms of speciation in rockfish. However, as in several cichlid assemblages, depth segregation and coloration differences suggest that ecological and sexual selections may play important roles in this system. Extensive analysis of closely related *Sebastes* species using fast-evolving markers, such as microsatellites, should determine if recent speciation events are a common feature of the *Sebastes* species flock.

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