New markers for new species: microsatellite loci and the East African cichlids

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The radiation of the East African cichlid fishes has engaged biologists for over a century. Because so much taxonomic diversity has evolved recently, they are an ideal natural system in which to study the process of speciation. Hypervariable microsatellite loci have been used to verify multiple paternity and maternity in cichlid broods, to quantify the fitness of cooperative breeders and reproductive parasites, to estimate effective population sizes in captive populations, and to illuminate the spatial and temporal scale of gene flow among natural populations. The patterns that have emerged from these studies often reflect important biological differences among taxa. The cichlid species of East Africa represent a large amount of taxonomic and adaptive diversity all neatly packaged into a single lineage and confined to a modest geographical area. Data from microsatellite loci are now providing us with the means to understand one of the world's most intriguing and instructive comparative evolutionary systems.

While cichlids share most aspects of their basic bauplan, the lineage contains a diversity of ecological adaptations and reproductive strategies. Furthermore, each of several behavioral, ecological and biogeographical descriptors encompasses numerous species, many of which transcend phylogenetic or biogeographical boundaries. Thus, the diversity of the East African cichlids provides an ideal and replicated comparative system in which to test the predictions of speciation theories, and might serve as a paradigm for evolutionary biologists and ecologists studying other systems.

Before the mid-1990s, the available genetic markers provided insufficient resolution to adequately describe cichlid mating strategies, population genetics and phylogeny. Although the "rich, speculative" cichlid literature contains a wealth of research on the phenotypic raw material present, it has until recently lacked evidence for the phylogenetic and population genetic underpinnings needed to understand the divergence of lineages. Here, we focus on recently obtained microsatellite data, which have overcome many of the persistent difficulties inherent in the application of other markers.

The trouble with cichlids

The evolution of taxonomic diversity has been prolific and recent in the East African cichlids, with thousands of endemic species having evolved in the three Great Lakes since the late Pleistocene.

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rampant splitting of lineages provides numerous opportunities for studying the divergence process in situ. Ironically, certain aspects of cichlid evolution are difficult to resolve, precisely because speciation is so recent. In species from the two youngest Great Lakes (Malawi and Victoria), allozymes and mtDNA contain low levels of allelic or sequence polymorphism. The brief periods between speciation events cause what little variation exists at these loci to be shared among taxa. These types of marker are not definitive in studies of assortative mating, only mildly informative for population studies, and are phylogenetically informative in only the oldest lineages. By contrast, simple sequence repeat loci (SSRs, also known as microsatellites), with their high mutation rates and high levels of polymorphism, have clarified aspects of the reproductive behaviour and population genetics in this most speciose vertebrate family.

Mating systems and microsatellites

**Paternity and maternity**

A high level of parental care is characteristic of the cichlids. In Lakes Malawi and Victoria, and in many Tanganyikan lineages, care is provided by the female who incubates eggs and fry in her mouth until the juveniles are well developed. Generally males form leks on the open sand or in the water column or defend small, spatially clustered territories among the rocks, from which they court passing females. Behavioral observations suggested that gravid females are courted by multiple males in sand-dwelling species, but not the reproductive success of these males, nor whether rock-dwelling females mate with more than one male, was known. Kellogg et al. and Parker and Kornfield used SSR loci to estimate the number of males contributing to a brood in several sand- and rock-dwelling species from Lake Malawi. Most broods surveyed from both habitat guilds are sired by two or more males. More surprisingly, brood mixing was documented in the Malawi cichlid Protomelas cf. spilopterus in four of six broods surveyed, an example of multiple maternity.

This was found to be true even for broods that were apparently too young to have been released on brief foraging trips, thus suggesting brood mixing might be under maternal control. The adaptive significance of multiple maternity and resistance in Lake Malawi has yet to be explored experimentally.

**Mate recognition**

Mate recognition is widely thought to be of fundamental importance to animal speciation. In the East African Cichlidiae, putative sibling morphospecies are often distinguished by striking differences in male breeding coloration, whereas shape differences are often subtle. In laboratory settings, hybrids between taxa from the same lake are often fertile. Behavioral experiments suggest that females mate assortatively on the basis of male color patterns. Using a combination of behavioral observations and SSR genotypes, van Oppen et al. tested mating preferences between sympatric color morphs (putative species) in the field. High and statistically significant interspecies fixation indices allowed the authors to reject the null hypothesis of genetic panmixia between sympatric color forms and to conclude that premating isolation is well developed among these morphospecies. Many alleles are shared between taxa, but it is not obvious whether this is a result of shared ancestry or a result of low levels of ongoing gene flow (sensu Wang et al.).

**Brood-care helpers and reproductive parasites**

Microsatellite and other genetic fingerprinting methods have been used to document both reproductive parasitism and cooperative breeding in two substrate-spawning species from Lake Tanganyika. Lamprologus callipterus females incubate their eggs inside snail shells, which have been gathered into nests by the larger of two male morphs. Smaller ‘dwarf’ males act as nest parasites by hiding inside shells and fertilizing eggs. Analysis of SSR alleles indicates that these ‘satellite’ males are successful, siring up to 90% of the progeny when they are present in a nest (P. Meidl, PhD thesis, University of Vienna, 1999).

Fitness benefits for ‘helpers at the nest’ have been documented in the social Tanganyikan cichlid Neolamprologus pulcher. Breeding units in this species consist of a pair of large reproductive adults and several smaller individuals who assist with brood care and defense. DNA fingerprinting techniques in laboratory experiments demonstrate that when a mature male helper is present, he sires about 10% of the offspring. Microsatellite analysis of wild N. pulcher populations further demonstrated that these family groups are largely matrilinesal and that levels of inbreeding within a group are minimized as a result of high levels of male exchange between groups (M. Taborsky, pers. commun.).

**Population structure**

**Levels of migration**

It has long been suspected that gene flow between populations in many cichlid species is low and that this has an important influence on the speciation process, either as a result of distinct selective environments or through sexual selection. Among the endemic and colorful rock-dwelling ‘mbuna’ species of Lake Malawi, lack of larval dispersal, heavy predation pressure, narrow depth tolerance and the patchy distribution of preferred habitat are all believed to contribute to high levels of population subdivision. Early biochemical and molecular studies were consistent with limited gene flow, but the markers employed could not adequately define the scale of genetic structure or the nature of any barriers to migration. Recently, SSR loci have allowed a more detailed understanding of the scope...
Box 1. Mbuna and the problem of excess polymorphism

Based on their highly significant but small \( F_{ST} \) estimates, van Oppen et al.\(^a\) speculated that mbuna species can be divided into ‘thousands’ of semi-isolated genetic units and that this isolation might be an important aspect of the speciation process in cichlids, a statement echoed in subsequent papers\(^b\)–\(^d\). Citing difficulties with the interpretation of migration indices when data from highly polymorphic loci are used, Hedrick\(^e\) has recently questioned this prevailing view. \( F_{ST} \) values are constrained by within-population homozygosity, which is often rather low at microsatellite markers. These markers also provide researchers with large amounts of statistical power, permitting the detection of allele frequency differences that are statistically real, but which might have little biological importance.

Are levels of genetic difference detected in these recent mbuna studies biologically important? The migration estimates in these studies are high enough to prevent fixation of alternative alleles between adjacent populations through drift alone. However, an alternative estimator of migration, Barton and Slatkin’s \( Nm \) (Ref. 1), suggests that some, more widely separated mbuna populations are indeed genetically decoupled\(^f\). Moreover, Danley et al.\(^g\),\(^h\) have discussed the relevance of divergence with gene flow models to the evolution of the East African cichlids.

Reduced migration between habitat patches is important biologically if selective forces are strong enough to overcome the homogenizing effects of interdemic migration (sensu Endler\(^i\)). Selective forces might include different modes of sexual selection or adaptation to variable community structures or abiotic factors that can vary widely between habitat patches\(^j\).

The sampling formulae of Jin and Chakraborty\(^k\) can be used to place migration estimates between recently separated populations in context for a given level of heterozygosity (following van Oppen et al.\(^\lambda\)). \textit{Pseudotropheus trophoe} ‘mauve’ populations have an estimated \( F_{ST} \) of 0.016. Figure I shows the null expectation of \( F_{ST} \) in the complete absence of gene flow over time for four different \( N_m \) values. The \( F_{ST} \) value estimated by van Oppen et al. for this species is consistent with an \( N_m \) of 1750 or fewer individuals in populations that have been completely isolated for 67 generations. A recent re-evaluation of geological data\(^j\) suggests that the populations could have been isolated for 2000 generations instead of the 67 generations that were assumed by van Oppen et al., consistent with an \( N_m \) of 50 000 individuals, or far fewer individuals than this assuming extensive gene flow. In such a large population, deterministic forces might become more important than stochastic forces.

It is difficult to make inferences about the evolution of cichlids from survey data alone. The problem is compounded because divergence times, census and effective population sizes, SSR mutation rates and generation times – each crucial components of the balance between selection and migration – are so far a matter of informed conjecture. Although our present knowledge does not permit us to reliably estimate absolute levels of gene flow, the absence of panmixia between proximate populations indicates that philopatry is an aspect of cichlid evolution worthy of more formal exploration.

References

\(^d\) Markert, J. A. et al. (1999) Biogeography and population genetics of the Lake Malawi cichlid \textit{Melanochromis auratus}; habitat transience, philopatry and speciation. Mol. Ecol. 8, 1013–1026

and nature of gene flow in both Lake Malawi and the Lake Victoria Region.

At Lake Malawi, van Oppen et al.\(^28\) sampled four mbuna species from habitat patches along a 3 km stretch of the western shore. They calculated small but statistically significant \( F_{ST} \) values. The estimated number of migrants (\( N_m \)) exchanged between populations ranges from 6.3 to 8.3 individuals per generation, on a scale of less than 1 km. On the basis of these estimates, the authors speculated that mbuna populations can be divided into ‘thousands’ of semi-independent genetic subunits (Box 1). If these preliminary estimates are roughly correct, modest selective pressures could lead to phenotypic divergence in long-lived populations, assuming that effective population sizes (\( N_e \)) are in the order of tens of thousands.

The biogeographical aspects of population structure in mbuna species (Fig. 1) were explored in surveys conducted in southern Lake Malawi\(^29\)–\(^31\).
Samples were collected from each major rocky habitat patch along a 42 km stretch of shoreline adjacent to the Nankumba Peninsula. All four species showed evidence of restricted gene flow between populations. Melanochromis auratus has the highest level of population differentiation, with an \( F_{ST} \) estimate of 0.15 overall. In fact, the \( N_m \) between the two most distant populations sampled is estimated to be below the 0.5 individuals per generation necessary to allow the fixation of alternate alleles through drift alone. The other three species were estimated to have higher levels of gene flow, with statistically significant \( F_{ST} \) estimates ranging from 0.063 in Labeotropheus fuelleborni populations to 0.016 in the three Metriaclima sandaracinos populations (the latter species has a limited distribution in the study area). A comparison of estimates of gene flow between adjacent habitat patches suggests ecologically important differences between these taxa (Fig. 2). For example, in M. auratus and L. fuelleborni the lowest migration estimates were obtained between populations separated by deep water. This finding is consistent with physiological studies that have demonstrated that these benthic fishes are not capable of rapidly adjusting to changes in water depth\(^7\). These same deep water channels did not restrict migration in Metriaclima zebra to the same degree, leading Danley et al.\(^{20}\) to speculate that M. zebra might cross them while feeding in the water column.

Although a picture of limited migration in mbuna is emerging, species-specific factors might influence the relative levels of migration. Barrier and habitat types were not well replicated within the surveyed transect; thus these observations merit further study. Nevertheless, a picture of high levels of mbuna philopatry regulated by species-specific dispersal is intriguing, given that each of the Great Lakes has a large endemic assemblage of rock-fish and a patchy distribution of habitats (but see Box 2).

A Lake Victoria Region (LVR) species, surveyed with different SSR loci, illustrates a different pattern of population structure. Wu sampled several populations of the LVR cichlid Astatoreochromis alluaudi, a habitat generalist found in several LVR lakes and streams (L. Wu, PhD thesis, Ohio State University, 1999). No clear evidence of population structure was found within lakes. However, moderate levels of genetic structure were observed in all of the between-lake comparisons. Overall, the scale of genetic subdivision in A. alluaudi is much larger than in the Malawi mbuna. Wu notes that A. alluaudi has an extremely broad habitat distribution and that it belongs to one of the least speciose haplochromine genera. Indeed, this genus does not seem to have speciated at all within the LVR, suggesting that philopatry might be a precondition for high levels of divergence among the East African cichlids.

Evidence for philopatry also exists within habitat patches. Knight et al.\(^{32}\) sampled Metriaclima from a 20 x 20 m grid. Using SSR-based estimates, they found a weak correlation between proximity and relatedness for females but not for males, providing evidence of male-biased dispersal and female philopatry. The authors reason that in a sexually selected species this could lead to prolific speciation. Migrating males would suffer greatly reduced mating success in foreign demes if females preferred male morphotypes from their own deme, and interdemic gene exchange would be relatively low. By contrast, male philopatry and female dispersal would result in females 'settling' for heterotypic males, who would readily mate with them, leading to the collapse of population-specific mating patterns. They hypothesized that inbreeding avoidance is a consequence of male dispersal behavior in this species.

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**Box 2. Sympatry, speciation and cichlids**

The importance of geographic isolation to the speciation process has been a long running and controversial theme in cichlid studies. Mayr⁴, for example, eventually supported an older ‘microallopatric’ model in which speciation occurs at isolated habitat patches within a lake, abandoning a stricter view in which divergence originated in separate bodies of water.

The apparent monophyly of many lineages within the Great Lakes⁵ (but see Nagl et al.⁶), the patchy distribution of rocky, sandy and muddy habitats within the lakes⁷, and recent studies of population structure in Lake Malawi, which suggest that inter-population migration is restricted⁸⁻¹⁰, are all compatible with the microallopatric view of speciation. However, none of these documented patterns exclude alternative modes of speciation.

In 1994, Schliewen et al.¹¹ presented data supporting strictly sympatric speciation in cichlid lineages within two small crater lakes in West Africa. Unlike the East African Great Lakes, the substrate in the crater lakes is homogeneous, presumably providing few opportunities for microgeographical isolation. mtDNA sequence data suggest that the several species within each crater lake arose recently from a single common ancestor, leading the authors to infer that these taxa evolved in the absence of geographic barriers.

Dominey¹² and other authors have stressed the importance of sexual selection as a driving force in the cichlid speciation process. Seehausen and van Alphen¹³ have suggested that sexual selection might permit speciation, even in the absence of microgeographical isolation. In Lake Victoria, fish species with nearly identical morphologies but with distinct male breeding coloration often coexist in the same habitat patches. Based on experiments that document the importance of these color differences in mate recognition, and on the overlap of geographic distributions of congeners, Seehausen and van Alphen¹³ argue that much speciation occurs in strict sympathy as a result of sexual selection. Although the distribution overlap observed by Seehausen and van Alphen¹³ is consistent with sympatric speciation, Bouton¹⁴ has argued that a similar pattern would result from speciation in allopatry followed by parallel colonization of linearly arrayed habitats.

At present, we are left with several unresolved questions and few excluded hypotheses. Does speciation occur primarily in allopatry on fine geographic scales as a result of divergent selection, either in the form of adaptation or through the evolution of site specific mating systems? Is there a role for reinforcement in these taxa that appear to have few post-mating reproductive barriers? Does rampant speciation in sympathy through sexual selection explain the rapid evolution of thousands of cichlid species? Or are a variety of mechanisms operating in parallel to generate this extraordinary taxonomic diversity?

**References**


**Genetic variation, habitat age and secondary contact**

As a result of fluctuations in lake water level, shoreline habitats in the East African Great Lakes are chronically unstable. The rocky habitats in southern Lake Malawi would have been dry land during the most recent major drop in lake water level, which might have occurred in the past few hundred years⁷ (but see Ref. 33). Habitats that today have rocks extending to a great depth would have become available for colonization earlier, as the lake basin refilled. A correlation between heterozygosity and the depth of the rock–sand interface (a proxy for habitat age and stability) was observed in both M. zebra and M. auratus⁳⁰,⁴⁵. The reduction in heterozygosity at shallower sites is not large. However, it does suggest that each new habitat patch was founded by a moderate number of individuals in a pattern of serial colonization from deeper to shallower sites. Evidence of the occurrence of genetic bottlenecks during this process is lacking, an interpretation consistent with previous studies⁴⁶. Deep habitats are located in the northern half of the transect surveyed and shallower habitats are found to the south. Because deeper habitats might be capable of supporting larger populations, alternative explanations for this pattern are possible.

Secondary contact might be responsible for acline in male breeding coloration in the early colonizer L. fuelleborni, which was observed along the same transect. Based on a qualitative assessment of allele frequencies at a single loc...
Box 3. Microsatellite mutation models and estimates of divergence

There has been an explosion of theoretical and empirical treatments of microsatellite mutation processes over the past decade. Currently, estimates of levels of genetic divergence or patterns of relatedness between populations and species rely on statistics that have been developed from two classic models.

Stepwise Mutation Model (SMM)

SMM has emerged as the ‘null model’ of microsatellite evolution\(^2\). The strict SMM assumes that mutations involve the loss or gain of a single repeat unit (thereby retaining some ‘memory’ of previous allelic states) and that allele sizes are unconstrained, which might hold true over short time spans\(^3\). SMM-based statistics require that a stepwise mutation process has more influence on allele frequency distributions than genetic sampling does. Based on evidence of non-stepwise mutational events at some microsatellite loci, several investigators have called for the identification and selection of mutationally ‘well behaved’ microsatellites\(^4\) for which to apply estimators of genetic divergence based on the SMM.

Infinite Alleles Model (IAM)

IAM was developed by Kimura and Crow\(^5\), and assumes that every mutational event creates a unique allele. As with the SMM, the IAM also assumes that there are no constraints on allele size. Statistics based on this mutational model violate a priori the well supported notion that there is some memory in the microsatellite mutation process. However, they might still perform well when the time since divergence is extremely short, when stochastic sampling of alleles far outweighs mutation in importance, and when the frequency of mutational events involving relatively large changes is high\(^6\).

More complex microsatellite mutation models

Evidence is accumulating that there are several complications involved in the mutation of microsatellites, including: larger mutational events, a dependence of mutation rates on the structure of the tandem repeat motif, a predisposition for microsatellite expansion over contraction, allele size constraints (e.g., reflecting or absorbing boundaries), and the possibility of heterozygote instability (reviewed by Amos\(^4\) and Estoup and Cornuet\(^7\)). Di Rienzo et al.\(^1\) introduced the two-phase mutation model (TPM) in which a mutation has a probability \(p\) of being a one step change and a probability \(1−p\) of being a larger change, drawn from a symmetric geometric distribution with specified variance. Numerous additional models have been formulated more recently. Difficulties associated with the estimation of required parameters impede the development of estimators of genetic divergence and population demographic statistics from these more complex mutation models.

References


Phylogeny quest

In Lakes Malawi and Victoria, short divergence times and the problem of retained ancestral polymorphisms have made the determination of phylogenetic relationships difficult\(^8\). The situation in these lakes contrasts with Lake Tanganyika, which appears to be a reservoir of ancient lineages\(^9\). Amplified fragment length polymorphism (AFLP)-based techniques have been used to compensate for these problems in Lake Malawi by generating an abundance of characters, but they appear to have little power to resolve intergeneric relationships in this system, perhaps because many genera evolved so rapidly\(^9\). Kornfield and Parker\(^9\) and Sültmann and Mayer\(^3\) have suggested that microsatellite loci might have the potential to overcome this difficulty owing to their high mutation rates and the potential information content of SSR alleles evolving under a stepwise mutation model (Box 3). In a limited pilot study using two species in each of three morphologically distinct genera, Kornfield and Parker\(^9\) found that they could cluster species into generic dyads for two out of the three genera. Further, they proposed that microsatellite ‘saltines’ (widely disjunct clusters of alleles) might be interpretable in a cladistic manner as synapomorphies. Sültmann and Mayer\(^3\) generated a phylogeny that roughly reflects two trophic guilds within Lake Victoria. Additionally, Wu used 14-locus SSR genotypes to estimate phylogenetic relationships among 23 taxa from the LVR (L. Wu, PhD thesis, Ohio State University, 1999). Although encouraging, these results are sensitive to mutation model assumptions. Unfortunately, independent estimates of phylogenetic relationships against which to compare these results are not available.

Microsatellite loci might ultimately be useful in estimating phylogenies, although the analytical and sampling challenges are considerable. The complexities of microsatellite mutations violate the strict assumptions of the genetic distance measures.
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References


Glossary

Amplified fragment ‘length’ polymorphism (AFLP): a technique for finding genetic variation at many loci simultaneously. Widely used in genome mapping, it has recently been used in phylogenetic estimation and might be useful in population studies. In AFLP, genomic DNA is cut with restriction enzymes. Restriction site specific oligonucleotide adapters are then ligated onto the entire population of restriction fragments. Next, PCR is performed using labeled primers that recognize the adapters and several additional base pairs. This permits rapid visualization of a subset of restriction sites using high resolution polyacrylamide gel electrophoresis.

FST: an index of population differentiation, commonly used to infer levels of migration between populations. FST ranges between zero and one. Higher FST values are interpreted to indicate a lower level of gene exchange between populations. Originally defined by Sewall Wright as $F_{ST} = (H_r - H_o) / H_o$, where $H_r$ is the average expected heterozygosity within each population and $H_o$ is the expected total heterozygosity calculated from the average of population allele frequencies. Wright’s original example was for a single locus with two alleles. Numerous multi-locus, multi-allelic analogues have been developed.

Homoplasy: evolutionary convergence. In molecular and biochemical data sets homoplasy refers to identity by character state that is not a result of evolutionary descent. Homoplasy may plague microsatellite studies where individual alleles are typically defined solely by the size of a PCR product. "Homoplasy", which is expected to increase with the mutation rate and the time since divergence, is another major problem. In addition, retained ancestral polymorphisms would result in phylogenetic noise at microsatellite loci, as with other classes of molecular markers, when closely related taxa are being compared. The high mutation rate and putative mutational ‘memory’ encoded in allele sizes might provide a partial solution to these problems. If these methods are to succeed in revealing relationships between recently diverged taxa, homology between frequently employed complex repeat arrays must be better established by sequence analysis, and the appropriateness of distance estimators needs to be evaluated on a taxon- and locus-specific basis (Box 3).

Conclusions

Why are there so many cichlids? The question remains largely unanswered, but after a century of study the theoretical and genetic tools necessary to address it might at last be available. The data derived from microsatellite markers are contributing to the framework needed for a more comprehensive understanding of cichlid evolution, revealing tantalizing differences between these closely related taxa. New insights into reproductive behavior and patterns of assortative mating have been obtained using these markers, and reproductive success and relatedness have been estimated with far higher levels of confidence than could have been achieved with behavioral observations alone.

The implications of the mating system characteristics quantified using SSR loci are not yet clear. Multiple paternity might imply that polyandry decreases the variance in male reproductive output (and thus increases the effective population size, $N_e$) by providing more mating opportunities for males. However, the element of female choice implied by these findings and supported by behavioral studies might mean that mating success is skewed in favor of a small number of highly successful males, in spite of the leveling effects of multiple paternity within this elite group. This second interpretation is compatible with data presented by Fiutera et al. who used SSR loci to estimate $N_e$ in captive breeding populations of two endangered Lake Victoria species. In these populations, $N_e$ was found to be only 1–15% of census size, although it is not clear whether these results are applicable to wild populations. A better understanding of effective population sizes will be crucial for gauging the relative influence of selection and drift in this system, and it is also needed to determine whether ephemeral populations have reached a selection–migration–drift equilibrium.

The relative importance of allopatric and sympatric speciation mechanisms remains an active theme in cichlid studies. Because the East African lakes contain closely spaced patches of habitat (and some of the smaller lakes contain only a single kind of habitat) it could be more practical to consider allopatry and sympatry as points along a gene flow continuum, which might or might not have an easily identifiable geographic component. Allopatric populations that...
References


