Integrating Phylogenetic and Population Genetic Analyses of Multiple Loci to Test Species Divergence Hypotheses in Passerina Buntings

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ABSTRACT

Phylogenetic and population genetic analyses of DNA sequence data from 10 nuclear loci were used to test species divergence hypotheses within Passerina buntings, with special focus on a strongly supported, but controversial, sister relationship between *Passerina amoena* and *P. caerulea* inferred from a previous mitochondrial study. Here, a maximum-likelihood analysis of a concatenated 10-locus data set, as well as minimize-deep-coalescences and maximum-likelihood analyses of the locus-specific gene trees, recovered the traditional sister relationship between *P. amoena* and *P. cyanea*. In addition, a more recent divergence time estimate between *P. amoena* and *P. cyanea* than between *P. amoena* and *P. caerulea* provided evidence for the traditional sister relationship. These results provide a compelling example of how lineage sorting stochasticity can lead to incongruence between gene trees and species trees, and illustrate how phylogenetic and population genetic analyses can be integrated to investigate evolutionary relationships between recently diverged taxa.

NFERRING the species tree of recently diverged taxa L can be a challenging exercise (MADDISON 1997; Felsenstein 2004; Maddison and Knowles 2006). The difficulty arises in part because the stochastic sorting of ancestral polymorphisms after speciation can result in gene trees that are discordant with species trees (PAMILO and NEI 1988; TAKAHATA 1989; MADDISON 1997; ROSENBERG 2002, 2003; MADDISON and KNOWLES 2006). For recently diverged taxa that have not yet attained complete reproductive isolation, gene flow introduces another source of discordance between the gene trees and the species tree (TAKAHATA and SLATKIN 1990; TAKAHATA 1995; WAKELEY 1996; WAKELEY and HEY 1998). The impact of these processes on species tree reconstruction has long been recognized (GILLESPIE and LANGLEY 1979; NEI and LI 1979), but it is only recently that a modelbased phylogenetic framework has integrated one of them (ancestral sorting) into the species tree inference. Because postspeciation, interspecific gene flow has yet to be incorporated into a phylogenetic framework, nonphylogenetic methods must be used to infer species relationships indirectly. Here, phylogenetic and divergence population genetic analyses of a multilocus nuclear data set are used to examine a controversial sister species relationship in Passerina buntings.

Over most of its taxonomic history, the genus Passerina (Aves: Cardinalidae) has been composed of six species of small (13–20 g), sexually dichromatic songbirds. The

collective breeding ranges of these species encompass most of Mexico, the United States, and southern Canada (Figure 1 and supplemental Figure 1 at http://www. genetics.org/supplemental/). Within the genus, *Passerina* cyanea and P. amoena have typically been considered sister species because they are morphologically similar and form a well-known hybrid zone where their breeding distributions overlap in the western Great Plains and eastern foothills of the Rocky Mountains in North America (SIBLEY and SHORT 1959; EMLEN et al. 1975; KROODSMA 1975; BAKER and BAKER 1990; BAKER and BOYLAN 1999). While the monophyly of Passerina has never been questioned (HELLACK and SCHNELL 1977; TAMPLIN et al. 1993), there have been attempts to subsume Guiraca caerulea and some species of the genus Cyanocompsa into the genus (PHILLIPS et al. 1964; BLAKE 1969; MAYR and SHORT 1970; PAYNTER 1970).

KLICKA *et al.* (2001) addressed the evolutionary relationships of the traditional six-member genus and closely related species, including *G. caerulea* and three Cyanocompsa species, using 1143 bp of sequence data from the mitochondrial cytochrome *b* gene. An unexpected result of the study was that *P. amoena* and *P. cyanea*, the two morphologically similar species that form a hybrid zone in the Great Plains, were not sister species. Instead *G. caerulea* (hereafter, *P. caerulea*) was sister to *P. amoena*, a relationship that had never before been hypothesized. *P. cyanea* was placed as the most basal member of the now seven-member genus. As expected, the three Cyanocompsa species fell outside of Passerina.

The branching structure of the mitochondrial tree and the demography of Passerina buntings suggest that the unexpected *P. amoena/P. caerulea* sister relationship

Sequence data from this article have been deposited with the EMBL/ GenBank Data Libraries under accession nos. EU191758–EU191837.

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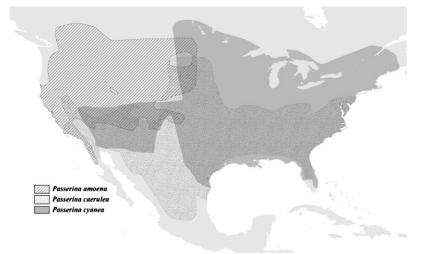


FIGURE 1.—Breeding distributions of *Passerina* amoena, *P. cyanea*, and *P. caerulea*. Digital maps (RIDGELY *et al.* 2003) for each species were downloaded from NATURESERVE (2006). Breeding distributions of the other four members of the genus Passerina are available in supplemental Figure 1 at http://www.genetics.org/supplemental/.

could reflect discordance between the mitochondrial gene tree and the species tree. The internodes in the mitochondrial phylogeny are short, possibly indicating rapid divergences, and the population sizes for some of the species, including P. cyanea, P. amoena, and P. caerulea, are large-on the order of tens of millions of individuals (PAYNE 1992; GREENE et al. 1996). This combination of short internodes and large population sizes is among the most difficult scenarios for inferring phylogenies from single locus data sets given the elevated risk of discordance between a species tree and its underlying gene trees (Figure 2). Using the calculation of HEIN et al. (2005) for the probability of species tree/gene tree discordance and realistic parameter estimates for Passerina buntings, the probability of discordance is estimated to be as high as 55% (see Comparing mtDNA and nuclear DNA signals in DISCUSSION).

Here, phylogenetic and divergence population genetics methods are used to test speciational hypotheses in Passerina buntings, with special focus on the controversial sister relationship uncovered by the mitochondrial study. Using the Passerina data set, the utility of combining phylogenetic and population genetics methods to uncover the evolutionary history of closely related species where hybridization may complicate evolutionary history is examined. Specifically, support for the following two alternative hypotheses was assessed:

- Hypothesis 1: *P. cyanea* and *P. amoena* are more closely related to each other than either is to *P. caerulea*. This is the traditional sister relationship.
- Hypothesis 2: *P. amoena* and *P. caerulea* are more closely related to each other than either is to *P. cyanea*. This is the sister relationship found in the mitochondrial tree.

MATERIALS AND METHODS

Sampling, amplification, and sequencing: From museum collections we obtained vouchered frozen tissue samples from

two individuals of each focal species, P. cyanea, P. amoena, and P. caerulea (see APPENDIX for sampling localities). We also included one individual from two additional species: P. rositae and Cyanocompsa cyanoides. C. cyanoides is a member of a genus closely related to Passerina (KLICKA et al. 2001) and was included here as an outgroup. We extracted total DNA from ~ 25 mg of pectoral muscle using a DNeasy tissue kit (QIAGEN, Valencia, CA), following the manufacturer's recommended protocol. Each individual was amplified at 10 nuclear loci (Table 1) using the following PCR conditions in a 25 µl amplification reaction: ~50 ng template DNA (2.5 µl of QIAGEN DNA extracts), 1 µl of 10 mM dNTPs (2.5 mM each dATP, dTTP, dCTP, and dGTP), 1 μ l of each primer (10 mM, Table 1), 2.5 μ l 10× buffer with MgCl₂ (15 mM), 0.1 µl Taq (5 units/µl AmpliTaq DNA polymerase, Applied Biosystems, Foster City, CA), and 16.9 μ l sterile dH₂O. The thermocycling profile was as follows: an initial 95° denaturation for 2 min followed by 35 cycles consisting of a 30-sec, 95°-denaturation step, a 30-sec, locusspecific temperature primer annealing step (Table 1), and a 2min, 72°-extension step, and a final extension of 5 min at 72°. We checked for amplification by electrophoresing 2.5 µl of each PCR amplicon on a 1% agarose gel and then used 20% polyethylene glycol (PEG) to purify the remaining 22.5 µl of the PCR product from successful amplifications.

Both strands of the PEG-purified PCR amplicons were cyclesequenced using 1.5 μl of 5× sequencing buffer (Applied Biosystems), 1 µl of 10 mM primer, 2-3 µl template (depending on the intensity of the PCR bands on the agarose gels), 0.5 µl Big Dye Terminator cycle-sequencing kit v 3.1 (Applied Biosystems), and 1–2 μ l sterile dH₂O (depending on amount of template added), for a total volume of 7 µl. The longer loci were also cycle-sequenced with internal sequencing primers using the same protocol (Table 1). We cleaned cycle-sequencing products on Sephadex (G-50 fine) columns and electrophoresed the cleaned products on a 3100 Genetic Analyzer (Applied Biosystems). In those cases where direct sequencing of purified PCR amplicons revealed more than one heterozygous site within a sequence, we phased the haplotypes by cloning the PCR products using the TA Topo cloning kit according to the manufacturer's instructions (Invitrogen, Carlsbad, CA) and sequencing five clones. All sequences were edited and assembled using Sequencher v 4.1 and 4.6 (GeneCodes, Ann Arbor, MI).

Phylogenetic analyses: Sequence alignments of all 10 nuclear loci were unambiguous and checked visually. In all phylogenetic analyses, gaps were alternately classified as a fifth state or as missing data. Phylogenetic analyses were conducted on a concatenated data set since partition homogeneity tests revealed

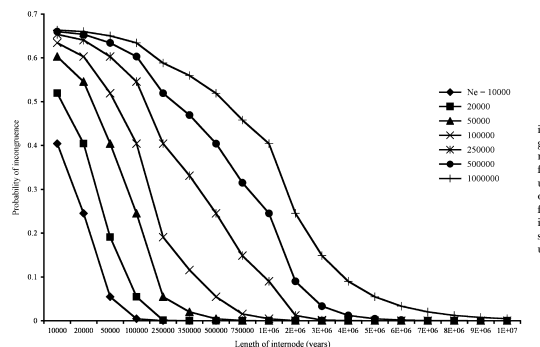


FIGURE 2.—The probability a gene tree will be incongruent with a rooted threemember species tree as a function of the effective population size and the length of the internode in years. Effective population sizes and internode lengths were chosen to bracket realistic values for Passerina buntings.

no significant conflicts between the data from each of the individual loci. Tests for recombination (see Population genetic analyses below) indicated that three loci (Bact3, MPLPR, and TGF_{β2}) had undergone recombination. Therefore, phylogenetic analyses were conducted on both the entire concatenated data set (5232 bp) and on a reduced concatenated data set containing only the largest independently segregating block of sequence for Bact3, MPLPR, and TGFB2 (4668 bp). Using PAUP* v 4.0b10 (Swofford 1998) we estimated a neighbor-joining tree from the p-distance matrix and calculated likelihood scores on a series of nested substitution models. These were then evaluated using ModelTest v 3.8 (POSADA and CRANDALL 1998). Due to its advantages over hierarchical likelihood-ratio tests (POSADA and BUCKLEY 2004), we used the Bayesian Information Criterion (BIC) to determine the best-fit model of sequence evolution for the concatenated data set as well as for each locus independently (Table 2).

To investigate species-level relationships between these taxa we conducted a variety of phylogenetic analyses on the concatenated data set using the model of sequence evolution identified by ModelTest v 3.8. First, we analyzed an 8-terminal-taxa data set composed of sequence data from either one (P. rositae and C. cyanoides) or two individual(s) per species (P. cyanea, P. amoena, and P. caerulea) where double peaks in chromatograms were inferred to be heterozygous sites and coded as polymorphisms using the standard IUPAC degeneracy codes. Second, we analyzed a 16-terminal-taxa data set that consisted of two phased haplotypes, determined by sequencing cloned PCR products from each individual for a total of four sequences per species of *P. cyanea*, *P. amoena*, and *P. caerulea* and two sequences per species for P. rositae and C. cyanoides. These data sets of concatenated sequences will be referred to hereafter as the 8-terminal-tip data set and the 16-terminal-tip data set.

Using parameters estimated on a neighbor-joining tree (SAITOU and NEI 1987), we used PAUP* v 4.0b10 (SwOFFORD 1998) to perform heuristic searches [tree bisection–reconnection (TBR) branch swapping; 10 random-addition replicates] under the maximum-likelihood (ML) optimality criterion for both the 8- and 16-terminal-tip concatenated data sets. We assessed nodal support using 100 ML heuristic bootstrap replicates with

the search parameters described above. We also performed a variety of constrained ML searches (Table 3) on the 8- and 16-terminal-tip data sets using the sequence evolution model and search parameters, except without bootstrap replicates, as described above. Using a Shimodaira–Hasegawa test (SH) (SHIMODAIRA and HASEGAWA 1999) with full optimization and 1000 bootstrap replicates, we assessed whether the likelihoods of the unconstrained and constrained trees were significantly different.

We also assessed nodal support for the 8- and 16-terminaltip topologies using posterior probabilities generated using MrBayes v 3.1.2 (HUELSENBECK and RONQUIST 2001; RONQUIST and HUELSENBECK 2003). We used the same general model (GTR + I) as in the ML searches and uninformative priors and ran two replicate analyses of four Markov chains for 1,000,000 generations. Additionally, we ran a mixed-model analysis using MrBayes in which the 8-terminal-tip concatenated data set was partitioned into the 10 individual loci, each partition with the appropriate model of sequence evolution (Table 2). In both the one-model and mixed-model searches, we sampled trees every 100 generations and calculated final posterior probabilities after excluding the first 250,000 generations as burn-in. Plots of likelihood scores during the run were examined visually to insure stationarity had been reached. In all searches, the average standard deviation of split frequencies was <0.005, further evidence stationarity had been reached.

Minimize-deep-coalescences analyses: Because well-supported species trees inferred from concatenated data sets can be discordant with the true species tree, especially if the species divergence was recent (DEGNAN and ROSENBERG 2006; EDWARDS *et al.* 2007; KUBATKO and DEGNAN 2007), we inferred a species tree using a method that considers the individual gene histories. Specifically, we assessed the fit of the gene trees onto three possible rooted species trees for *P. cyanea, P. amoena*, and *P. caerulea*, using the minimize-deep-coalescences method as implemented in the program Mesquite (MADDISON 1997; MADDISON and KNOWLES 2006). This method attempts to reconcile gene trees contained within species trees by minimizing the number of deep coalescences, those cases where the coalescence of two gene copies predates the speciation event. Using a parsimony optimality criterion,

Locus	Annealing temp	Length (bp) ^a	PCR/sequencing primers ^b	Internal sequencing primers $^{\epsilon}$	Accession nos.
AETC	°00	299	For: TGGACTTCAAATCCCCCCGATGATCCCAGC	NA	EU191758-765
			Rev: CCAGGCACCCCAGTCTACCTGGTCAAA		
Bact3	62°	423 (255)	For: GGCAATGAGAGGTTCAGGT	NA	EU191766–773
			Rev: TGGTACCACCAGACAGCAC		
GADPH	62°	303	For: ACCTTTCATGCGGGGGGGGCGCGCGCATTGC	NA	EU191774–781
			Rev: CATCAAGTCCACAACACGGTTGCTGTA		
MCIR	62.5°	774	For: CTGGCTCCGGAAGGCRTAGAT	For: SGCRTAGAAGATGGTGATGTAGC	EU191782–789
			Rev: AYGCCAGYGAGGGCAACCA	Rev: GTBGACCGCTACATCACCAT	
MPLPR	63.5°	293 (144)	For: TACATCTACTTTAACACCTGGACCACCTG	NA	EU191790–797
			Rev: TTGCAGATGGAGGAGCAGGTTGGAGCC		
Myo2	58°	684	For: GCCACCAAGCACAAGATCCC	NA	EU191798–805
			Rev: GCAAGGACCTTGATAATGACTT		
ODC	65°	609	For: GACTCCAAAGCAGTTTGTCGTCTCAGTGT	NA	EU191806-813
			Rev: TCTTCAGAGCCAGGGGAAGCCACCACCAAT		
Rho1	00°	855	For: TGCTACATCGAGGGCTTCTT	For: GTGCCTAAAAATGTCCTCT	EU191814-821
			Rev: GCAGTGACCAGAGAGCGATT	Rev: GCTGGCATGGGGGGGGGGGGGGG	
TGFB2	62°	544 (283)	For: GAAGCGTGCTCTAGATGCTG	NA	EU191822–829
			Rev: AGGCAGCAATTATCCTGCAC		
Tropo6	$^{\circ}09$	448	For: AATGGCTGCAGAGGATAA	NA	EU191830-837
			Rev: TCCTCTTCAAGCTCAGCACA		

÷ Ξ ••• --10 £ -• -Ra 4 FMRI / Cei f li i i ЪСВ et al. (2006); MPLPR, PRIMMER et al. (2002); Myo2, HESLEWOOD et al. (1998) (For) and SLADE et al. (1993) (Rev); ODC, PRIMMER et al. (2002); Rho1, PRIMMER et al. (2002); TGFβ2, PRIMMER et al. (2002); Tropo6, PRIMMER et al. (2002). ⁽¹⁾Internal sequencing primer references: MCIR, CHEVIRON et al. (2006); Rho1, this study.

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Data set	Model	Base frequencies ^a	ti/tv ratio	Rate matrix ^{<i>b</i>}	Pinv
AETC	K80	Equal	$5.8 imes10^{36}$	_	_
βact3	K80	Equal	$6.1 imes10^{36}$	_	_
GADPH	JC	Equal	_	_	_
MC1R	F81	0.1533, 0.3795, 0.2348, 0.2304		_	_
MPLPR	F81	0.2097, 0.2600, 0.3476, 0.1827		_	
Myo2	JC	Equal		_	_
ÓDC	F81	0.2790, 0.1677, 0.2127, 0.3406		_	_
Rho1	JC	Equal		_	_
TGF _{β2}	ĴС	Equal		_	_
Tropo6	F81	0.1959, 0.2227, 0.1714, 0.4057		_	_
8-terminal-taxa ^c	K81uf+I	0.2231, 0.2429, 0.2462, 0.2876		1.0, 4.424, 1.8284, 1.8284, 4,424, 1.0	0.9295
16-terminal-taxa ^c	K81uf+I	0.2364, 0.2287, 0.2616, 0.2633	—	1.0, 7.150, 2.243, 2.243, 7.150, 1.0	0.9307

Summary of model parameters used in all phylogenetic analyses as determined by the Bayesian information criterion (BIC) as implemented in ModelTest 3.8 (POSADA and CRANDALL 1998)

^a Order of base frequencies is A, C, G, T.

^bOrder of rate matrix is A to C, A to G, A to T, C to G, C to T, and G to T.

^e These data sets contain the concatenated sequences from all 10 loci.

the "best" species tree is the one that requires the fewest deepcoalescent events in contained gene genealogies. The number of deep-coalescent events, or costs, can be summed across multiple markers to accommodate multilocus data sets. Using Mesquite (MADDISON and MADDISON 2004), we compared the fit of the genealogies from each locus into the possible species trees using methods slightly modified from the recently described method of MADDISON and KNOWLES (2006).

We used MrBayes v 3.1.2 (HUELSENBECK and RONQUIST 2001; RONQUIST and HUELSENBECK 2003) to sample the tree space of the gene trees for each locus independently, using completely phased haplotypes and the same model parameters with uninformative priors as in the ML tree searches and the same MrBayes run parameters as described above. After discarding the first 250,000 generations as burn-in, we used PAUP* to calculate a majority-rule consensus gene tree from each locusspecific search. We then overlaid each unrooted locus-specific gene tree onto the three possible species trees (Table 3) and tallied the number of deep- coalescent events required to reconcile the gene trees with the species trees. The total cost of each species tree was determined by summing the locus-specific tallies (Table 4), with the species tree having the lowest cost being inferred as the correct species tree. **Calculating probabilities of gene trees given possible species trees:** Recent software developments give researchers increased ability to evaluate the variation in lineage sorting among loci and how it influences phylogenetic inferences. The program COAL (DEGNAN and SALTER 2005), which extends earlier work (TAKAHATA and NEI 1985; ROSENBERG 2002), can calculate the probability of a gene tree given a species tree for any number of taxa. This represents a significant step toward developing a method capable of estimating the likelihood of sequence data given a species tree, which can be estimated as follows:

$$\prod_{\text{loci}} \sum_{\text{gene} \atop \text{trees}} [P(\text{sequences} | \text{gene tree}) \\ \times P(\text{gene tree} | \text{species tree})]$$

(MADDISON 1997). The first probability statement is analogous to the log-likelihood scores generated by phylogenetic inference software packages such as PAUP* (SWOFFORD 1998). The program COAL can calculate the second probability statement. Unfortunately, complete evaluation of the above equation is

]	TABLE 3	
Likelihood scores for	various topological	constraints

		$Constraint^a$	
Data set	(((Pam, Pcy), Pca), Pro, Ccy) ^b	(((Pam, Pca), Pcy), Pro, Ccy)	(((Pcy, Pca), Pam), Pro, Ccy)
8-terminal-tip concatenated ^e 8-terminal-tip concatenated ^d 16-terminal-tip concatenated ^e	$-\frac{8148.9118}{-\frac{7239.1485}{-9140.8436}}$	-8151.7215 -7240.0937 -9142.4041	-8155.0551 -7243.9608 -9147.5444

Likelihood scores (ln *L*) were estimated using a heuristic search (TBR branch swapping, 10 random addition replicates). Underlining indicates topology with greatest likelihood. The constraint trees are also the species toplogies used in the minimize-deepcoalescences (MADDISON and KNOWLES 2006) and COAL analyses (DEGNAN and SALTER 2005).

^a Species name abbreviations: Pam, P. amoena; Pcy, P. cyanea; Pca, P. caerulea; Pro, P. rositae; Ccy, Cyanocompsa cyanoides.

^b This was also the topology recovered in all the unconstrained searches, which also gave identical ln L scores.

^e These data sets included all sequence data from all loci.

^d This data set included only the largest independently segregating block of sequence for those loci showing evidence of recombination.

Deep coalescent events required to reconcile unrooted locus-specific gene trees, inferred using Bayesian tree searches, with three possible species topologies

	Species topologies ^a				
Locus	(((Pam, Pcy), Pca), Pro, Ccy)	(((Pam, Pca), Pcy), Pro, Ccy)	(((Pcy, Pca), Pam), Pro, Ccy)		
AETC	5	7	7		
βact3	6	6	6		
GADPH	3	3	3		
MC1R	1	<u>0</u>	1		
MPLPR	12	11	$\underline{10}$		
Myo2	3	3	3		
ODC	7	7	<u>6</u>		
Rho1	<u>0</u>	1	1		
TGFβ2	5	7	7		
Tropo	$\overline{0}$	1	1		
Total Cost	42	46	45		

Underlining indicates species topology with fewest deep coalescent events for each locus as well as for the lowest total summed over all locus-specific gene trees.

^a Species name abbreviations: Pam, P. amoena; Pcy, P. cyanea; Pca, P. caerulea; Pro, P. rositae; Ccy, Cyanocompsa cyanoides.

hampered because the summation step requires summing over all possible gene trees, which includes both topology and branch lengths (MADDISON 1997). Currently, no method is capable of performing that summation because of the near infinite number of possible gene trees. It will likely require a method that employs some type of importance sampling of the gene tree space to effectively evaluate that step of the overall equation (J. DEGNAN, personal communication).

In light of the current methodological limitations, we focused on calculating the conditional probability of a gene tree given a species tree. Locus-specific gene trees were inferred in three ways. First, we randomly chose a single individual to represent *P. amoena, P. cyanea*, and *P. caerulea*, which along with one individual from each outgroup taxon (*P. rositae* and *C. cyanoides*), resulted in a data set with five terminal tips. For each locus, we estimated a gene tree under the maximum-likelihood search criterion as described above and then used COAL to calculate the probability of each ML gene tree given three possible species trees (Table 3) using COAL. ML inferences for Tropo6 produced >100 equally likely gene trees and so we eliminated Tropo6 from the COAL analysis.

We also inferred locus-specific gene trees from the 8-terminaltip data set. Since COAL requires fully resolved bifurcating gene trees and since locus-specific ML searches of this 8-terminal-tip data set produced numerous equally likely topologies with unresolved polytomies, we inferred locus-specific neighbor-joining gene trees, using uncorrected *p*-distances. In the Tropo6 data set, some of the sequences were identical across individuals. Accordingly, neighbor-joining analyses failed to produce a fully resolved tree for Tropo6 and we excluded it from COAL calculations.

In an attempt to incorporate a larger sample of the possible gene topologies and branch lengths, we sampled a collection of gene trees from the locus-specific Bayesian tree searches described above (see *Minimize-deep-coalescences analyses*). Our approach was to calculate the probability of each of the three possible species trees (Table 3) given each of the last 1000 gene trees from the Bayesian searches, using COAL, and average the probabilities of those gene trees to generate a more robust estimate of the likelihoods of the three species trees. Unfortunately, in each locus-specific Bayesian tree search some gene trees contained unresolved polytomies, which COAL cannot handle; therefore, we abandoned this method of inferring gene trees.

We replicated the COAL calculations, for both the five- and eight-terminal tip data sets, using three different internal: terminal branch-length ratios (1:1, 1:2, and 1:10). All internal: terminal branch-length ratios, with the exception of 1:1, correspond to topologies with internode lengths that are short, relative to the terminal branches, which appears to be the general pattern observed in Passerina buntings. To assess the difference in probabilities of the gene trees if the species tree united *P. cyanea* and *P. amoena* as sisters (hypothesis 1) with one in which *P. caerulea* and *P. amoena* were sisters (hypothesis 2), we conducted a test of proportions (SOKAL and ROHLF 1995) whereby we tallied the number of gene trees that were more probable under hypothesis 1 and compared that with a tally of the number of gene trees that were more probable under hypothesis 2.

Population genetic analyses: Both phylogenetic methods described above assume that discordance between gene trees and the species tree is due to genetic drift (*i.e.*, lineage sorting stochasticity). Because of the well-documented hybrid zone between *P. amoena* and *P. cyanea*, it would be ideal if a gene flow parameter could be incorporated into the phylogenetic analysis. This is not currently possible. We thus examined the utility of divergence population genetics for inferring species relationships indirectly. The Isolation with Migration (IM) software package (HEY and NIELSEN 2004; HEY 2005) can estimate pairwise divergence times between *P. cyanea*, *P. amoena*, and *P. caerulea*, while jointly estimating pairwise migration rates. A sister relationship is inferred to be between the species with the most recent divergence estimate.

IM combines coalescent theory with Bayesian methodologies to simultaneously estimate multiple population genetic parameters for two diverging populations. Briefly, using a Markov chain Monte Carlo (MCMC) approach, IM generates posterior probability distributions of six parameters ($\theta_{population1}$, $\theta_{population2}$, $\theta_{ancestral}$, m_1 , m_2 , t). These parameters can then be used to calculate population demographic estimates, *i.e.*, effective population sizes of the two populations (N_{e1} and N_{e2}), the ancestral population size (*i.e.*, the population size prior to initial divergence = N_{eA}), migration rates between the two populations (m_1 and m_2 , which allows for asymmetrical migration rates), and the time since divergence of the two populations (t). A more recent version of IM (Hey 2005) also allows for the inclusion of a population splitting parameter, *s*, which allows for exponential population growth or decline in each descendant population.

IM assumes selective neutrality of each marker and no intralocus recombination. We tested for neutrality by calculating Tajima's D (TAJIMA 1983) for each locus using Arlequin v 3.0 (Excoffier et al. 2005). None of the 10 loci showed a significant departure from neutrality (each P > 0.25). To investigate intralocus recombination, we used TOPALi v 2 (McGuire and WRIGHT 1998, 2000), which uses a sliding-window approach to search for putative recombination breakpoints along an alignment of DNA sequences. Using the difference in sum-of-squares (DSS) method, with a 100-bp window and a 10-bp increment, we looked for signals of recombination, manifested by a significant DSS peak along the sequence alignment. The statistical significance of DSS peaks was assessed using 100 parametric bootstraps. Significant DSS peaks were detected in 3 loci, βact3, MPLPR, and TGF β 2 (Table 1). For each of those 3 loci, we only included the longest independently segregating block of sequence from the phased haplotype data in the IM runs along with the total available phased haplotype sequences from each of the other seven markers.

Using IM, we estimated three pairwise divergence times, one between P. cyanea and P. amoena, one between P. caerulea and P. amoena, and one between P. cyanea and P. caerulea. Initial IM runs, conducted using the infinite sites (IS) model (KIMURA 1969) and employing unrealistically large maximum priors $(t = 20, q_1 = q_2 = q_a = 500, m_1 = m_2 = 50, \text{ or } m_1 = m_2 = 0 \text{ for}$ parameter estimation without gene flow) were used to establish appropriate parameter priors for all subsequent analyses (WON and HEY 2005). Once appropriate priors were identified, we ran IM three times with identical starting conditions, with the exception of the random number seed, to assess convergence. All runs began with a burn-in period of 100,000 steps and were allowed to continue until the minimum effective sample size (ESS) was >100 (Hey 2005). Because runs with the same starting conditions produced qualitatively similar results, we only present data from the longest run ($>3 \times 10^9$ steps). For each pairwise estimate of divergence time, we performed a variety of IM analyses with varying numbers of parameters. The simplest model included four parameters (θ_1 , θ_2 , θ_A , and t); other models included five $(\hat{\theta}_1, \theta_2, \theta_A, t, and m$ -equal migration between populations or θ_1 , θ_2 , θ_A , *t*, and *s*—allowing for population growth) or six parameters (θ_1 , θ_2 , θ_A , t, m_1 , and m_2 —allowing for different, directional migration between populations). All model parameter estimates in IM are scaled to the mutation rate, µ. To translate scaled divergence time estimates to real time, we included mutation rates for each locus calculated as the sequence length multiplied by 3.6×10^{-9} substitutions/site/year (AXELSSON et al. 2004).

RESULTS

Phylogenetic analyses: Maximum-likelihood analysis of the entire concatenated 8-terminal-tip data set, based on 5232 bp of sequence data from each individual, produced one strongly supported tree (Figure 3A). In it, the traditional sister relationship between *P. amoena* and *P. cyanea* (hypothesis 1) was recovered. This unconstrained tree was more likely than either of the constrained topologies, regardless of the data set, although the differences in likelihood scores were not significant (SH test, P > 0.05, Table 3). Support values from ML bootstrap analyses and

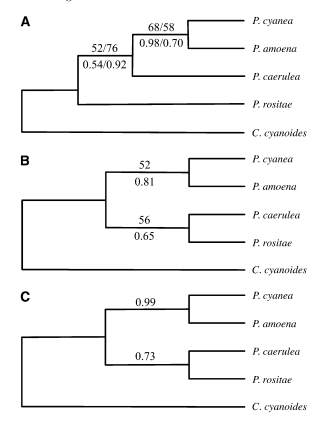


FIGURE 3.—Species trees inferred from the entire concatenated data set (5332 bp), except where noted. In all phylogenies, multiple sequences from each species were collapsed into a single "representative." Only bootstrap support values >50 (at the top of the branches) and Bayesian posterior probabilities >0.50 (at the bottom of the branches) are illustrated. (A) Phylogeny inferred from the 8-terminal-tip data set. For both bootstrap support values and Bayesian posterior probabilities the first number refers to the concatenated data set and the second to a reduced concatenated data set containing only the largest independently segregating block of sequence from those loci for which recombination was detected. (B) ML and Bayesian species tree inferred from the 16-terminal-tip data set with bootstrap support values at the top of the branches and posterior probabilities at the bottom. (C) Posterior probabilities of Bayesian phylogeny inferred from the 8-terminal-tip data set with a mixed model of sequence evolution.

Bayesian posterior probabilities, calculated under a single model of sequence evolution for the entire data set, indicated the topology recovered in the unconstrained searches was relatively well supported (Figure 3). The Bayesian analysis of the 8-terminal-tip data set using a mixed model, in which a sequence evolution model was specified for each locus, uncovered the same topology as the Bayesian analysis of the 16-terminal-tip data set (Figure 3, B and C). All of the above phylogenetic analyses were rooted using *C. cyanoides* as an outgroup. Replicate analyses with the reduced concatenated data set (4668 bp) produced the same topologies as the entire concatenated data set (Figure 3A).

Phylogenies inferred from the 16-terminal-tip data set were also similar to the ML and Bayesian phylogenies inferred from the 8-terminal-tip data set (Figure 3). Both, ML and Bayesian, 16-terminal-tip phylogenies contained a well-supported sister clade between *P. cyanea* and *P. amoena*. The placement of *P. caerulea* with respect to *P. rositae* and the *P. cyanea/P. amoena* clade varied depending on the data set, 8- or 16-terminal tip, used to infer the phylogeny. Both ML and Bayesian inferences uncovered a weakly supported clade containing *P. cyanea*, *P. amoena*, and *P. caerulea*, when tree searches were conducted on the 8-terminal-tip data set. When based on the 16-terminal-tip data set, both analyses recovered a sister relationship between *P. rositae* and *P. caerulea*, and that clade was sister to the *P. cyanea/P. amoena* clade (Figure 3C).

The topologies inferred from the tree searches of the concatenated data sets did not differ when gaps were coded as missing data or as a fifth state (topologies not shown).

Minimize-deep-coalescences analyses: Overlaying the majority-rule consensus gene trees from the Bayesian tree searches on the three possible species trees resulted in three different deep coalescent costs for each gene—one cost for each species tree (Table 4). For four locus-specific gene trees (AETC, Rho1, TGF β 2, and Tropo6) the fewest number of deep coalescent events were observed when the species topology united *P. amoena* and *P. cyanea* as sister species. The fewest deep coalescent events for one locus (MC1R) occurred when *P. amoena* and *P. caerulea* were sister species, and the *P. cyanea*/*P. caerulea* sister species tree contained the fewest deep coalescences when overlaid with the ODC and MPLPR gene trees.

Summing the number of deep coalescent events needed to fit the gene trees into the species tree across loci produced a total cost of 42 for the *P. amoena/P. cyanea* sister species tree (Table 4). Fitting the gene trees into the other two species trees required more deep coalescences: 46 for the *P. amoena/P. caerulea* tree and 45 for the *P. cyanea/P. caerulea* tree.

Calculating probabilities of gene trees given possible species trees: After randomly choosing one individual to represent P. amoena, P. cyanea, and P. caerulea, and eliminating Tropo6 from further analyses, 14 ML gene topologies were inferred from phylogenetic tree searches conducted on each locus separately (Table 5). Multiple gene trees with equal likelihoods were inferred from three loci (AETC-three gene trees, βact3-two, and MPLPRthree). Replicate COAL calculations that differed only in the ratio of internal to terminal branch lengths (1:1, 1:2, and 1:10) all gave qualitatively similar results. When terminal branch lengths were longer than internal branch lengths (1:2 and 1:10), 3 of the 14 gene topologies were equally probable whether the given species tree had P. cyanea and P. amoena as sister species or a sister species relationship between *P. amoena* and *P. cyanea* (Table 5). Those 3 gene topologies (one of two Bact3 gene trees and two of three MPLPR gene trees) were excluded from

further analyses, because with respect to these relationships, these topologies are uninformative. Of the remaining 9 ML gene topologies (one of the three equally likely AETC gene trees was chosen at random), 8 were more probable given a species tree uniting *P. cyanea* and *P. amoena* as sisters, and 1 was more probable given a *P. amoena–P. caerulea* species tree (Table 5, test of proportions, $n_{obs} = 9$, P = 0.05). When the internal:terminal branch length ratio was 1:1, 4 of the 14 gene topologies were equally probable given either species tree and were subsequently eliminated. Seven of the remaining 8 (1 AETC gene tree chosen at random) ML gene topologies were more probable with *P. cyanea/P. amoena* as sister species (Table 4, test of proportions, $n_{obs} = 8$, P = 0.1).

Nine gene topologies, one for each locus with the exception of Tropo6, were inferred from the eight-terminal tip data set using neighbor-joining analyses (Table 6). In the COAL calculations with internal:terminal branch lengths of 1:1 and 1:2, there was no difference in the number of locus-specific gene topologies that were more probable with *P. cyanea* and *P. amoena* as sisters in the species tree and the number of gene topologies that were more probable with a *P. amoena/P. caerulea* sister relationship (Table 6, test of proportions, $n_{obs} = 9$, P > 0.05). When the internal:terminal branch length ratio of the species trees was 1:10, all nine gene topologies were more probable when the species tree placed *P. cyanea* and *P. amoena* as sisters (Table 6, test of proportions, $n_{obs} = 9$, P = 0.02).

Population genetic analyses: After excluding all gaps in the sequence data and retaining only the largest independently segregating block of sequence for those loci that showed evidence of recombination, 4721 bp of sequence data were included in the IM analyses. When divergence times were estimated in the absence of migration, analyses uncovered a more recent splitting between P. cyanea and P. amoena than between either of those two species and P. caerulea (Figure 4A, Table 7). Estimates for θ_1 , θ_2 and θ_A were inconclusive; the distributions of the posterior probabilities were nearly flat in all comparisons (data not shown). Allowing for growth or decline in the descendant populations did not qualitatively alter the divergence time estimates for any of the pairwise comparisons; the most recent divergence was between P. amoena and P. cyanea, with divergence times later for both the P. caerulea/P. amoena and P. caerulea/ P. cyanea comparisons (Table 7).

When divergence time was estimated jointly with migration, the high points of the posterior probability distributions suggested a different pattern. The divergence time pattern was the same regardless of whether the direction of gene flow was constrained to be symmetrical (five-parameter model) or asymmetrical (six-parameter model); thus, only the divergence times estimated under the six-parameter model are discussed. In the *P. cyanea/ P. amoena* comparison, the divergence time (0.211), estimated jointly with migration, was nearly identical to the

		(1:1, 1:2, and 1:10)			
				Probability	
Locus ^a	ML gene tree ^b	Species topology ^b	1:1	1:2	1:10
AETC	((Pam, Pcy), ((Ccy, Pca), Pro))	((((Pcy, Pam), Pca), Pro), Ccy)	0.00082	0.00495	0.01511
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00018	0.00199	0.01157
	((Pam, Pcy), ((Pro, Ccy), Pca))	((((Pcy, Pam), Pca), Pro), Ccy)	0.00082	0.00495	0.01511
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00018	0.00199	0.01157
	((Pam, Pcy), (Ccy, (Pro, Pca)))	((((Pcy, Pam), Pca), Pro), Ccy)	0.01663	0.03091	0.02587
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00366	0.01270	0.01997
βact3	((((Pam, Ccy), Pcy), Pro), Pca)	((((Pcy, Pam), Pca), Pro), Ccy)	0.00000	0.00487	0.01103
•		(((Pam Pca) Pcu) Pro) Ccu)	0.00000	0.00191	0.00748

Probabilities of ML gene trees given two alternative species topologies with differing internal:terminal branch length ratios (1:1, 1:2, and 1:10)

		((((<i>Pam</i> , <i>Pca</i>), <i>Pcy</i>), <i>Pro</i>), <i>Ccy</i>)	0.00366	0.01270	0.01997
βact3	((((Pam, Ccy), Pcy), Pro), Pca)	((((Pcy, Pam), Pca), Pro), Ccy)	0.00000	0.00487	0.01103
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00000	0.00191	0.00748
	(((Pam, Ccy), (Pcy, Pro)), Pca)	((((Pcy, Pam), Pca), Pro), Ccy)	0.00002	0.00051	0.00647
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00002	0.00051	0.00647
GADPH	((((Pam, Pcy), Ccy), Pca), Pro)	((((Pcy, Pam), Pca), Pro), Ccy)	0.00082	0.00487	0.01103
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00018	0.00191	0.00748
MC1R	(((Pam, Pca), Pcy), (Ccy, Pro))	((((Pcy, Pam), Pca), Pro), Ccy)	0.01075	0.02075	0.02104
		((((Pam, Pca), Pcy), Pro), Ccy)	0.07336	0.07322	0.03321
MPLPR	((Pam, ((Pro, Pca), Pcy)), Ccy)	((((Pcy, Pam), Pca), Pro), Ccy)	0.00059	0.00259	0.00532
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00059	0.00259	0.00532
	(((Pam, (Pro, Pca)), Pcy), Ccy)	((((Pcy, Pam), Pca), Pro), Ccy)	0.00059	0.00259	0.00532
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00059	0.00259	0.00532
	(((Pam, Pcy), (Pro, Pca)), Ccy)	((((Pcy, Pam), Pca), Pro), Ccy)	0.06330	0.05614	0.02232
		((((Pam, Pca), Pcy), Pro), Ccy)	0.01366	0.02188	0.01605
Myo2	(((Pam, Pcy), (Pro, Pca)), Ccy)	((((Pcy, Pam), Pca), Pro), Ccy)	<u>0.06330</u>	0.05614	0.02232
		((((Pam, Pca), Pcy), Pro), Ccy)	0.01366	0.02188	0.01605
ODC	((((Pam, Pcy), Pca), Pro), Ccy)	((((Pcy, Pam), Pca), Pro), Ccy)	0.40808	0.17006	0.02217
		((((Pam, Pca), Pcy), Pro), Ccy)	0.05686	0.04358	0.01230
Rho1	((((Pam, Pcy), Pca), Pro), Ccy)	((((Pcy, Pam), Pca), Pro), Ccy)	0.40808	0.17006	0.02217
		((((Pam, Pca), Pcy), Pro), Ccy)	0.05686	0.04358	0.01230
TGFβ2	((((Pam, Pcy), Ccy), Pca), Pro)	((((Pcy, Pam), Pca), Pro), Ccy)	<u>0.00082</u>	0.00487	0.01103
		((((Pam, Pca), Pcy), Pro), Ccy)	0.00018	0.00191	0.00748

Underlining indicates probabilities that are greater with a *P. amoena/P. cyanea* species topology than with a *P. amoena/P. caerulea* species topology. Italic numbers indicate the opposite pattern. All probabilities calculated using COAL (DEGNAN and SALTER 2005).

^{*a*} Gene tree searches based on Tropo6 sequence data from one randomly chosen individual from each species resulted in >100 ML gene trees and so that locus was dropped from the COAL analysis.

^b Species names abbreviations: Pam, P. amoena; Pcy, P. cyanea; Pca, P. caerulea; Pro, P. rositae; Ccy, C. cyanoides.

divergence time estimates using a model without migration (0.2105; Table 7). For both the *P. amoena/P. caerulea* and *P. cyanea/P. caerulea* comparisons, the divergence time estimate with the highest posterior probability was 4.9975, much greater than the estimates in the absence of migration (0.7185 and 0.537, respectively; Table 7). However, each of the posterior probability distributions of divergence time estimates for the *P. amoena/P. caerulea* and *P. cyanea/P. caerulea* comparisons, which never converged to zero, also had peaks much closer to the peak of the *P. amoena/P. cyanea* distribution (Figure 4B, Table 7).

Estimates of migration rates between the species varied. The posterior probability distributions of m_1 and m_2 between *P. cyanea* and *P. amoena* were flat, but nonzero (Figure 4, C and D). In contrast, migration into *P. caerulea* from both *P. cyanea* and *P. amoena* was effectively zero (m_2 in both comparisons, Figure 4D). However, the parameter distributions for migration into *P. cyanea* and *P. amoena* from *P. caerulea* both show evidence for nonzero gene flow (Figure 4C). The posterior probability distributions for m_1 were also similar for both comparisons involving *P. caerulea*, one peak at zero and a secondary, nonzero peak indicating gene flow into either *P. cyanea* or *P. amoena* from *P. caerulea* (Figure 4C).

While the divergence time parameter estimates did change with the inclusion of the gene flow parameter (Figure 4 and Table 7), the data were not significantly more likely under the more complex model in any of the pairwise comparisons. In all three comparisons twice the difference between the likelihoods of the data under the complex model (allowing for directional migration) and the simple model (no migration) were less than the critical value ($\chi = 5.991$, d.f. = 2, P = 0.05)

DISCUSSION

Both phylogenetic and population genetic analyses of the 10-locus data set supported the traditional sister

Probabilities of neighbor-joining gene topologies, inferred from the eight terminal tip data set, given two alternative	ve
species trees with differing internal:terminal branch length ratios (1:1, 1:2, and 1:10)	

				Probability	
Locus ^a	NJ tree ^b	Species topology ^b	1:1	1:2	1:10
AETC	(Pcy1, (((Pam1, Pcy2), Pam2), ((Ccy, Pro),	((((Pcy, Pam), Pca), Pro), Ccy) ((((Pam, Pca), Pcy), Pro), Ccy)	<u>2.365E-09</u> 1.62E-10	<u>4.78789E-07</u> 4.6316E-08	<u>1.8218E-05</u> 1.82953E-06
	(Pca1, Pca2))))				
βact3	(Pcy1, (((Pam1, Pcy2),	((((Pcy, Pam), Pca), Pro), Ccy)	5.05E-10	<u>2.77914E-07</u>	<u>2.1178E-05</u>
	Ccy), ((Pro, (Pca1, Pca2)), Pam2)))	(((((Pam, Pca), Pcy), Pro), Ccy)	3.99E-10	1.07017E-07	2.62828E-06
GADPH	(Pcy1, (((Pam1, Pam2),	((((Pcy, Pam), Pca), Pro), Ccy)	3.143E-09	1.8008E-06	0.00018838
	(Pro, (Pca1, Pca2))), (Ccy, Pcy2)))	(((((<i>Pam</i> , <i>Pca</i>), <i>Pcy</i>), <i>Pro</i>), <i>Ccy</i>)	2.7306E-08	2.7271 <i>3</i> E-06	3.75984E-05
MC1R	(Pcy1, ((((Pam1, Pam2),	((((Pcy, Pam), Pca), Pro), Ccy)	2.124E-09	1.30936E-06	0.000149501
	(Pca1, Pca2)), (Ccy, Pro)), Pcy2))	(((((Pam, Pca), Pcy), Pro), Ccy)	1.02979E-07	5.10169E-06	3.50167E-05
MPLPR	(Pcy1, ((((Pam1, Pam2),	((((Pcy, Pam), Pca), Pro), Ccy)	5.136E-09	9.52899E-07	2.77842E-05
	Pcy2), Ccy), ((Pro, Pca1), Pca2)))	(((((<i>Pam</i> , <i>Pca</i>), <i>Pcy</i>), <i>Pro</i>), <i>Ccy</i>)	3.98E-10	5.10169E-06	1.74861E-05
Myo2	(Pcy1, ((((Pam1, Pam2),	((((Pcy, Pam), Pca), Pro), Ccy)	<u>1.53135E-06</u>	6.8394E-05	0.000690821
	Pcy2), Ccy), (Pro, (Pca1, Pca2))))	(((((<i>Pam</i> , <i>Pca</i>), <i>Pcy</i>), <i>Pro</i>), <i>Ccy</i>)	1.28294E-06	2.86844E-05	9.5464E-05
ODC	(Pcy1, ((((Pam1, Pcy2),	((((Pcy, Pam), Pca), Pro), Ccy)	<1.0E-12	1.393E-09	2.7787E-06
	(Ccy, Pro)), (Pca1, Pca2)), Pam2))	(((((<i>Pam</i> , <i>Pca</i>), <i>Pcy</i>), <i>Pro</i>), <i>Ccy</i>)	<1.0E-12	5.27E-10	3.8284E-07
Rho1	(Pcy1, ((((Pam1, Pam2),	((((Pcy, Pam), Pca), Pro), Ccy)	2.124E-09	1.30936E-06	0.000149501
	(Pca1, Pca2)), (Ccy, Pro)), Pcy1))	(((((Pam, Pca), Pcy), Pro), Ccy)	1.02979E-07	5.10169E-06	3.50167E-05
TGF _{β2}	(Pcy1, ((Pam1, (Ccy,	((((Pcy, Pam), Pca), Pro), Ccy)	<u>1.4E-11</u>	3.7386E-08	1.2647E-05
	(Pcy2, Pam2))), (Pro, (Pca1, Pca2))))	(((((Pam, Pca), Pcy), Pro), Ccy)	1.1E-11	6.71486E-07	1.4255E-08

Underlining indicates probabilities that are greater with a *P. amoena/P. cyanea* species topology than with a *P. amoena/P. caerulea* species topology. Italic numbers indicate the opposite pattern. All probabilities calculated using COAL (DEGNAN and SALTER 2005).

^a NJ gene tree search based on Tropo6 sequence data produced an unresolved topology and was dropped from COAL analyses. ^b Species names abbreviations: *Pam, P. amoena; Pcy, P. cyanea; Pca, P. caerulea; Pro, P. rositae; Ccy, C. cyanoides.* Numbers following species abbreviations in NJ topologies refer to the different individuals sampled. All species were constrained to be monophyletic in the species topologies.

relationship between P. amoena and P. cyanea. Interestingly, the P. caerulea/P. amoena sister relationship proposed by KLICKA et al. (2001) was supported by analyses of the MC1R data (Tables 4–6). Mutations in MC1R are known to correlate with variations in melanin-based plumage in a variety of taxa, including birds (MUNDY 2005; HOEKSTRA 2006). Within Passerina, P. amoena and P. caerulea are the only species with putatively melanin-based plumages (HILL and McGRAW 2006) and selection acting on MC1R might explain the differences we found between inferences based on MC1R and those based on other loci. There is a nonsynonymous change at amino acid residue 102 (Val \rightarrow Leu) that unites *P. caerulea* and P. amoena (Val) to the exclusion of P. cyanea (Leu), which does not have any melanistic plumage. However, P. rositae and C. cyanoides (neither of which have melanin-based plumage) share the Val allele with P. amoena and P. caerulea indicating that the allele (Leu) found in P. cyanea is likely derived and probably not related to the differences in plumage patterns between *P. caerulea*, *P. amoena*, and *P. cyanea*. Therefore, while we cannot completely rule out the possibility that selection on the melanin-based plumages of *P. amoena* and *P. caerulea* has resulted in the observed patterns, the available data suggest selection is an unlikely cause.

Below, we discuss the benefits of using the two types of analyses for resolving sister relationships in Passerina and then with respect to closely related taxa in general.

Comparing mtDNA and nuclear DNA signals: The sister relationship between *P. cyanea* and *P. amoena* supported by the multilocus analyses conflicts with a well-supported *P. caerulea/P. amoena* sister relationship inferred from mitochondrial data (KLICKA *et al.* 2001). A possible explanation for the discrepancies between the mtDNA and nuclear results is differential lineage sorting of the mtDNA with respect to the true species tree (BRAUN and BRUMFIELD 1998). Assuming ((*P. cyanea, P. amoena*), *P. caerulea*) is the correct topology, one can estimate the

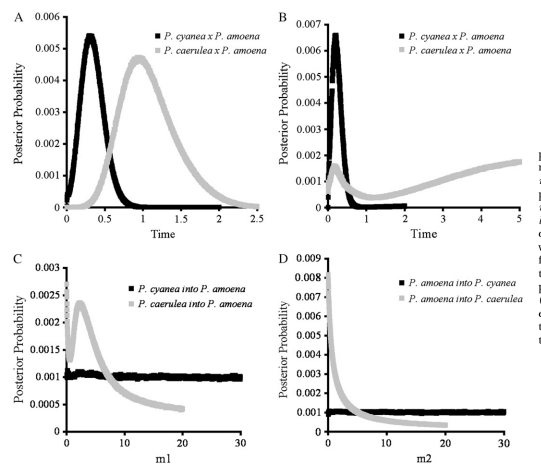


FIGURE 4.—The posterior probability distributions of model parameters $(t, m_1, and$ m_2) estimated using IM for pairwise comparisons P. cya $nea \times P.$ amoena and P. caeru $lea \times P.$ amoena. (A) Pairwise divergence time estimates with no postdivergence gene flow. (B) Pairwise divergence time estimates allowing for postdivergence gene flow. (C and D) Migration-rate estimates. All parameter estimates are scaled to the neutral mutation rate (μ) .

probability of incongruence between a gene tree and the species tree (FELSENSTEIN 2004; HEIN et al. 2005). Incorporating a range of internode values based on the divergence time estimates from IM and a range of potential effective ancestral population sizes, the chance a gene tree differs from the true species tree in the Passerina system ranges from 0% ($N_{\rm e} = 10,000$ individuals, internode length = 100,000 years) to 65% (ancestral $N_{\rm e}$ = 1,000,000, internode length = 100,000 years, Figure 2). Using an internode length of 350,000 years, which is the difference in the maximum-likelihood estimates of the divergence time for the P. cyanea \times P. amoena (~138,000 years ago) and the P. caerulea \times P. amoena (~486,000 years ago) comparisons (Table 7), and assuming no postspeciation gene flow, the probabilities of incongruence range from 0% (ancestral $N_{\rm e} = 10,000$) to 55% (ancestral $N_{\rm e} = 1,000,000$) (Figure 2). While an accurate estimate of the ancestral population size is unavailable, an effective size of 10,000 or 20,000 may be unrealistically small. Current population census estimates for P. cyanea are on the order of 20-80 million individuals (PAYNE 1992). Furthermore, while some coalescent-based ancestral population-size estimates tend to be relatively small (HEY 2005; WON and HEY 2005), an estimate in birds based on sequence data from multiple nuclear markers suggested an ancestral population size on the order of hundreds of

thousands of individuals (JENNINGS and EDWARDS 2005). Large population sizes and short internode lengths combine to increase the chance of incomplete and incorrect lineage sorting of gene copies, confusing the phylogenetic relationships of the focal taxa. The analyses of a multilocus data set presented herein suggest that the stochastic nature of lineage sorting of the mitochondrial genome resulted in a gene tree that is discordant with the species tree.

In closely related taxa, gene flow, in addition to lineage sorting problems, may limit the ability to correctly infer the species tree from gene sequences. P. cyanea and P. amoena currently hybridize where their breeding ranges overlap, although the amount of introgression is currently unknown, and hybrid specimens between other members of the genus have been reported. The samples of *P. amoena* and *P. cyanea* included in this study were collected far from the Great Plains contact zone (APPENDIX) to minimize the chance that recent introgression between the two species might be obscuring the true evolutionary relationships between P. cyanea, P. amoena, and P. caerulea. Ongoing studies of the genetic structure of the hybrid zone between P. cyanea and P. amoena suggest little mitochondrial or nuclear introgression is occurring (M. CARLING, unpublished data). The data from the hybrid zone, along with the nonsignificant difference in the likelihood of

	Constant	Constant population ^{b}	Variable	Variable population $^{\epsilon}$	With migratic	With migration (initial peak) d	With migrati	With migration (high point) ^{d}
Comparison	t	$t (yr)^e$	t	$t (yr)^{e}$	t	t (yr) ^e	t	$t (yr)^e$
P. cyanea \times P. amoena								
MLE^a	0.2105	138,000	0.207	136,000	0.211	138,000	0.211	138,000
90% HPD ^a	0.0375 - 0.4345	25,000-286,000	0.037 - 0.443	24,000-291,000	0.033 - 0.445	21,700-292,000	0.033 - 0.445	21,700-292,000
P. caerulea \times P. amoena								
MLE^a	0.7185	486,000	0.4825	327,000	0.1825	103,000	4.9975	3,380,000
90% HPD ⁴	0.3555 - 1.3125	241,000 - 888,000	0.2475 - 1.5325	167,000-1,037,000	NA	NA	NA	NAV
P. caerulea \times P. cyanea								
MLE^{a}	0.537	363,000	0.4935	334,000	0.0735	49,700	4.9975	3,380,000
90% HPD ^a	0.177 - 1.091	120,000-738,000	0.1505 - 1.3265	102,000-898,000	NA	NA	NN	NA

Maximum-likelihood estimates (MLE) and 90% highest posterior density (HPD) intervals of divergence time estimates for pairwise comparisons between P. amoend,

TABLE 7

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^{*a*} Divergence time estimates for IM runs under constant population-size model while allowing for migration. In all comparisons (except *P. cyanea* × *P. amoena*) there was an initial peak and a nonconverging, nonzero tail (high point). See Figure 4. ^{*f*} Mutation scaled estimates of divergence time (*t*) were translated into years using a mutation rate of 3.6×10^{-9} substitutions/site/year (AXELSSON *et al.* 2004). ^{*f*} These runs failed to converge so 90% HPDs are not presented.

the data to the more complex model in IM, indicate that gene flow between *P. cyanea* and *P. amoena* has probably not had a strong influence on the speciational history inferred in this study.

Implications for uncovering phylogenetic signals in closely related taxa: Integrating top-down phylogenetic methods with bottom-up population genetic methods was successful in that the same sister relationship was supported by both types of analyses. The data set of 10 loci sampled from either one (P. rositae and C. cyanoides) or two individuals per species (P. cyanea, P. amoena, and P. caerulea) generated a relatively well-supported species tree despite the low level of variation at each locus. One possible solution to increase the resolution of individual gene trees is to increase the length of the sampled sequence. Theoretically, lengthening the sampled sequence should increase the chance of detecting variable sites, which may then offer additional support for a particular topology (DOYLE 1992). In practice however, lengthening loci may not be a straightforward solution to a problem of poor resolution in phylogeny reconstruction. As the length of the sampled locus increases, so does the chance of sampling sites that have undergone recombination. We detected recombination in 3 of the 10 loci, a ratio similar to a recent multilocus study investigating the speciational history of Australian grass finches (JENNINGS and EDWARDS 2005). Both studies sampled loci between 200 and 900 bp, which demonstrates that potential effects of recombination should be considered in any population genetic or phylogenetic study employing sequence data from nuclear loci. Here, the results of the phylogenetic analyses were similar regardless of whether we analyzed the entire sequence length or just the longest independently segregating block of sequence from the 3 loci (Bact3, MPLPR, and TGFB2) that showed evidence of recombination (Table 3, Figure 3).

The sampling design used here closely mimicked the design that resulted in the greatest accuracy of species tree inference in MADDISON and KNOWLES'S 2006 investigation into the utility of using the minimize-deepcoalescences method for phylogenetic reconstruction. Using simulated data sets that differed both in the number of individuals sampled and the number of loci sampled, they assessed how often the minimize-deepcoalescences method recovered the true species tree, that is, the tree on which the sequences were simulated. A sampling design of nine loci and three individuals per species proved to be quite successful in recovering the correct species tree. The results of the modified minimizedeep-coalescences method used here provide empirical support for their best sampling design and serve as a useful reference point for researchers designing sampling protocols with the purpose of inferring phylogenies of closely related taxa.

Conclusions: Using DNA sequence data from a suite of nuclear loci, phylogenetic, and population genetic analyses supported the traditionally accepted evolution-

ary relationships between three North American passerines. In contrast to the findings of an earlier study that used data from the mitochondrial gene cytochrome b to infer the phylogeny of the genus (KLICKA et al. 2001), we found, using both phylogenetic and population geneticbased methods, P. cyanea and P. amoena to be more closely related to each other than either is to P. caerulea. These results underscore the importance of sampling multiple individuals and multiple loci when estimating multiple demographic parameters simultaneously under complicated evolutionary scenarios, e.g., when allowing for postdivergence gene flow. Unfortunately, IM and other coalescent-based methods available for inferring population genetic parameters are often limited in some ways (e.g., how many populations they can handle, the sequence evolution models employed, etc). The continued improvement of these methods will facilitate the combined analytical approach we advocate in studies with a greater number of focal taxa.

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LITERATURE CITED

- AXELSSON, E., N. G. C. SMITH, H. SUNDSTROM, S. BERLIN and H. ELLEGREN, 2004 Male-biased mutation rate and divergence in autosomal, Z-linked and W-linked introns of chicken and turkey. Mol. Biol. Evol. 21: 1538–1547.
- BAKER, M. C., and A. E. M. BAKER, 1990 Reproductive-behavior of female buntings - isolating mechanisms in a hybridizing pair of species. Evolution 44: 332–338.
- BAKER, M. C., and J. T. BOYLAN, 1999 Singing behavior, mating associations and reproductive success in a population of hybridizing Lazuli and Indigo Buntings. Condor 101: 493–504.
- BLAKE, C. H., 1969 Notes of the Indigo Bunting. Bird Banding 40: 133–139.
- BRAUN, M. J., and R. T. BRUMFIELD, 1998 Enigmatic phylogeny of skuas: an alternative hypothesis. Proc. R. Soc. Lond. Ser. B 265: 995–999.
- CHEVIRON, Z. A., S. J. HACKETT and R. T. BRUMFIELD, 2006 Sequence variation in the coding region of the melanocortin-1 receptor gene (MC1R) is not associated with plumage variation in the Blue-crowned Manakin (*Lepidothrix coronata*). Proc. R. Soc. Lond. Ser. B **273**: 1613–1618.
- DEGNAN, J. H., and N. A. ROSENBERG, 2006 Discordance of species trees with their most likely gene trees. PLoS Genet. 2: 0762–0768.
- DEGNAN, J. H., and L. A. SALTER, 2005 Gene tree distributions under the coalescent process. Evolution **59**: 24–37.
- DOYLE, J. J., 1992 Gene trees and species trees—molecular systematics as one-character taxonomy. Syst. Bot. 17: 144–163.
- EDWARDS, S. V., L. LIU and D. K. PEARI, 2007 High-resolution species trees without concatenation. Proc. Natl. Acad. Sci. USA 104: 5936– 5941.
- EMLEN, S. T., J. D. RISING and W. L. THOMPSON, 1975 A behavioral and morphological study of sympatry in the Indigo and Lazuli buntings of the Great Plains. Wilson Bull. 87: 145–179.
- EXCOFFIER, L., G. LAVAL and S. SCHNEIDER, 2005 Arlequin ver 3.0: an integrated software package for population genetics data analysis. Evol. Bioinformatics Online 1: 47–50.

FELSENSTEIN, J., 2004 Inferring Phylogenies. Sinauer Associates, Sunderland, MA.

- GILLESPIE, J. H., and C. H. LANGLEY, 1979 Are evolutionary rates really variable? J. Mol. Evol. 13: 27–34.
- GREENE, E., V. R. MUEHTER and W. DAVISON, 1996 Lazuli Bunting (*Passerina amoena*), pp. 1–24 in *The Birds of North America*, No. 232, edited by A. POOLE and F. GILL. The Academy of Natural Sciences, Washington, DC/The American Ornithologists' Union, Philadelphia.
- HEIN, J., M. K. SCHIERUP and C. WIUF, 2005 Gene Genealogies, Variation and Evolution. Oxford University Press, New York.
- HELLACK, J. J., and G. SCHNELL, 1977 Phenetic analysis of the subfamily Cardinalinae using external and skeletal characters. Wilson Bull. 89: 130–148.
- HESLEWOOD, M. M., M. S. ELPHINSTONE, S. C. TIDEMANN and P. R. BAVERSTOCK, 1998 Myoglobin intron variation in the Gouldian Finch *Erythrura gouldiae* assessed by temperature gradient gel electrophoreses. Electrophoresis **19**: 142–151.
- HEY, J., 2005 On the number of New World founders: a population genetic portrait of the peopling of the Americas. PLoS Biol. 3: 965–975.
- HEY, J., and R. NIELSEN, 2004 Multilocus methods for estimating population sizes, migration rates, and divergence time, with applications to the divergence of *Drosophila pseudoobscura* and *D. persimilis.* Genetics **167**: 747–760.
- HILL, G. E., and K. J. McGRAW, 2006 *Bird Coloration*, Vol. I. Harvard University Press, Cambridge, MA.
- HOEKSTRA, H. E., 2006 Genetics, development and the evolution of adaptive pigmentation in vertebrates. Heredity **97:** 222–234.
- HUELSENBECK, J. P., and F. RONQUIST, 2001 MrBayes: Bayesian inference of phylogenetic trees. Bioinformatics 17: 754–755.
- JENNINGS, W. B., and S. V. EDWARDS, 2005 Speciational history of Australian Grass Finches (Poephila) inferred from thirty gene trees. Evolution 59: 2033–2047.
- KIMURA, M., 1969 The number of heterozygous nucleotide sites maintained in a finite population due to steady flux of mutations. Genetics 61: 893–903.
- KLICKA, J., A. J. FRY, R. M. ZINK and C. W. THOMPSON, 2001 A cytochrome-b perspective on Passerina bunting relationships. Auk 118: 611–623.
- KROODSMA, R. L., 1975 Hybridization in buntings (Passerina) in North Dakota and eastern Montana. Auk 92: 66–80.КИВАТКО, L. S., and J. H. DEGNAN, 2007 Inconsistency of phyloge-
- KUBATKO, L. S., and J. H. DEGNAN, 2007 Inconsistency of phylogenetic estimates from concatenated data under coalescence. Syst. Biol. 56: 17–24.
- MADDISON, W. P., 1997 Gene trees in species trees. Syst. Biol. 46: 523–536.
- MADDISON, W. P., and L. L. KNOWLES, 2006 Inferring phylogeny despite incomplete lineage sorting. Syst. Biol. **55:** 21–30.
- MADDISON, W. P., and D. R. MADDISON, 2004 Mesquite: a modular system for evolutionary analysis, Version 1.01. http://mesquiteproject.org.
- MAYR, E., and L. L. J. SHORT, 1970 Species taxa of North American birds, a contribution to comparative systematics. Pub. 9, Nuttall Ornithological Club, Cambridge, MA.
- McGuire, G., and F. Wright, 1998 TOPAL: recombination detection in DNA and protein sequences. Bioinformatics 14: 219–220.
- McGUIRE, G., and F. WRIGHT, 2000 TOPAL 2.0: improved detection of mosaic sequences within multiple alignments. Bioinformatics **16:** 130–134.
- MUNDY, N. I., 2005 A window on the genetics of evolution: *MC1R* and plumage colouration in birds. Proc. R. Soc. Lond. Ser. B **272:** 1633–1640.
- NATURESERVE, 2006 NatureServe Explorer: An Online Encyclopedia of Life, Version 6.1. http://www.natureserve.org/explorer.
- NEI, M., and W. H. LI, 1979 Mathematical model for studying genetic variation in terms of restriction endonucleases. Proc. Natl. Acad. Sci. USA 76: 5269–5273.
- PAMILO, P., and M. NEI, 1988 Relationships between gene trees and species trees. Mol. Biol. Evol. 5: 568–583.

- PAYNE, R. B., 1992 Indigo Bunting, pp. 1–24 in *The Birds of North America*, No. 4, edited by A. POOLE, P. STETTENHEIM and F. GILL. The Academy of Natural Sciences, Washington, DC/The American Ornithologists' Union, Philadelphia.
- PAYNTER, R. A., 1970 Subfamily Cardinalidae, pp. 216–245 in *Checklist of Birds of the World*, edited by R. A. PAYNTER and R. W. STORER. Museum of Comparative Zoology/Heffernan Press, Worcester, MA.
- PHILLIPS, A. R., J. T. MARSHALL and G. MONSON, 1964 The Birds of Arizona. University of Arizona Press, Tucson, AZ.
- POSADA, D., and T. R. BUCKLEY, 2004 Model selection and model averaging in phylogenetics: advantages of the AIC and Bayesian approaches over likelihood ratio tests. Syst. Biol. 53: 793–808.
- POSADA, D., and K. A. CRANDALL, 1998 MODELTEST: testing the model of DNA substitution. Bioinformatics **14**: 817–818.
- PRIMMER, C. R., T. BORGE and G. P. SAETRE, 2002 Single-nucleotide polymorphism characterization in species with limited available sequence information: high nucleotide diversity revealed in the avian genome. Mol. Ecol. 11: 603–612.
- RIDGELY, R. S., T. F. ALLNUTT, T. BROOKS, D. K. MCNICOL, D. W. MEHLMAN *et al.*, 2003 Digital distribution maps of the birds of the Western hemisphere, version 1.0. NatureServe, Arlington, VA.
- RONQUIST, F., and J. P. HUELSENBECK, 2003 MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19: 1572–1574.
- ROSENBERG, N. A., 2002 The probability of topological concordance of gene trees and species trees. Theor. Popul. Biol. 61: 225–247.
- ROSENBERG, N. A., 2003 The shapes of neutral gene genealogies in two species: probabilities of monophyly, paraphyly, and polyphyly in a coalescent model. Evolution **57**: 1465–1477.
- SAITOU, N., and M. NEI, 1987 The neighbor-joining method—a new method for reconstructing phylogenetic trees. Mol. Biol. Evol. 4: 406–425.
- SHIMODAIRA, H., and M. HASEGAWA, 1999 Multiple comparisons of log-likelihoods with applications to phylogenetic inference. Mol. Biol. Evol. 16: 1114–1116.
- SIBLEY, C. G., and L. L. J. SHORT, 1959 Hybridization in the buntings (Passerina) of the Great Plains. Auk 76: 443–463.
- SLADE, R. W., C. MORITZ, A. HEIDEMAN and P. T. HALE, 1993 Rapid assessment of single-copy nuclear DNA variation in diverse species. Mol. Ecol. 2: 359–373.
- SOKAL, R. R., and F. J. ROHLF, 1995 Biometry. W. H. Freeman, New York. SWOFFORD, D. L., 1998 PAUP*. Phylogenetic Analysis Using Parsimony
- (* and Other Methods), Version 4. Sinauer Associates, Sunderland, MA. Талим, F., 1983 Evolutonary relationships of DNA sequences in fi-
- nite populations. Genetics **105**: 437–460. Таканата, N., 1989 Gene genealogy in three related populations—
- consistency probability between gene and population trees. Genetics **122**: 957–966.
- TAKAHATA, N., 1995 A genetic perspective on the origin and history of humans. Annu. Rev. Ecol. Syst. **26:** 343–372.
- TAKAHATA, N., and M. NEI, 1985 Gene genealogy and variance of interpopulational nucleotide differences. Genetics 110: 325–344.
- TAKAHATA, N., and M. SLATKIN, 1990 Genealogy of neutral genes in two partially isolated populations. Theor. Popul. Biol. 38: 331– 350.
- TAMPLIN, J. W., J. W. DEMASTES and J. V. REMSEN, 1993 Biochemical and morphometric relationships among some members of the Cardinalinae. Wilson Bull. **105:** 93–113.
- WAKELEY, J., 1996 Pairwise differences under a general model of population subdivision. J. Genet. 75: 81–89.
- WAKELEY, J., and J. HEY, 1998 Testing speciation models with DNA sequence data, pp. 157–167 in *Molecular Approaches to Ecology and Evolution*, edited by R. DESALLE and R. SCHIERWATER. Birkhauser Verlag, Basel, Switzerland.
- WON, Y. J., and J. HEY, 2005 Divergence population genetics of chimpanzees. Mol. Biol. Evol. 22: 297–307.

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APPENDIX: SAMPLING LOCALITIES FOR INDIVIDUALS INCLUDED IN THE STUDY TESTING SPECIES DIVERGENCE HYPOTHESES IN PASSERINA BUNTINGS

Species	ID no.	Locality
Passerina amoena	B-4005	Douglas County, Washington
	B-24769	San Bernadino County, California
P. cyanea	B-20575	Iberville Parish, Louisiana
2	B-20773	Cass County, Michigan
P. caerulea	B-20598	Cameron Parish, Louisiana
	B-21812	Jeff Davis County, Texas
P. rositae	69937^{a}	Monte Bonito, Chiapas, Mexico
Cyanocompsa cyanoides	B-12708	Velasco, Santa Cruz Department, Bolivia

^{*a*} Tissue obtained through loan from the University of Washington Burke Museum. All other tissue samples were obtained from the Genetic Resources Collection at the Louisiana State University Museum of Natural Science.