

Testing Nested Phylogenetic and Phylogeographic Hypotheses in the *Plethodon vandykei* Species Group

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Abstract.—Mesic forests in the North American Pacific Northwest occur in two disjunct areas: along the coastal and Cascade ranges of Oregon, Washington, and British Columbia as well as the Northern Rocky Mountains of Idaho, Montana, and British Columbia. Over 150 species or species complexes have disjunct populations in each area, and a priori hypotheses based on phylogeography and geology potentially explain the disjunction via either dispersal or vicariance. Here, we test these hypotheses in the disjunct salamander complex *Plethodon vandykei* and *P. idahoensis* by collecting genetic data (669 bp of Cyt *b*) from 262 individuals. Maximum likelihood analysis indicated reciprocal monophyly of these species, supporting the ancient vicariance hypothesis, whereas parametric bootstrap and Bayesian hypothesis testing allow rejection of the dispersal hypothesis. The coalescent estimate of the time since population divergence (estimated using MDIV) is 3.75×10^6 years, and the 95% credibility interval of this value overlaps with the geological estimate of vicariance, but not the hypothesized dispersal. These results are congruent with the pattern seen in other mesic forest amphibian lineages and suggest disjunction in amphibians may be a concerted response to a geological/climatological event. Within *P. idahoensis*, we tested the corollary hypothesis of an inland Pleistocene refugium in the Clearwater drainage with nested clade analysis and coalescent estimates of population growth rate (*g*). Both analyses support post-Pleistocene expansion from the Clearwater refugium. We corroborated this result by calculating Tajima's *D* and mismatch distribution within each drainage, showing strong evidence for recent population expansion within most drainages. This work demonstrates the utility of statistical phylogeography and contributes two novel analytical tools: tests of stationarity with respect to topology in the Bayesian estimation, and the use of coalescent simulations to test the significance of the population growth-rate parameter. [MCMC; *Plethodon idahoensis*; *Plethodon vandykei*; post-Pleistocene expansion; stationarity; statistical phylogeography.]

Biogeography and geology have been complimentary disciplines for most of the last two centuries, and comparative phylogeography is a recent extension of this tradition to a restricted taxonomic scale. Lineages with disjunct distributions have long attracted attention (e.g., Lyell, 1832), largely because they provide data critical to our understanding of evolutionary processes associated with lineage divergence. However, the study of disjunctions within species or between sister species is impeded by the continuum of evolutionary processes acting to structure genetic variation at these levels (Fig. 1). Researchers have typically approached phylogeographic data with methodological tools developed from either end of this continuum, with either a phylogenetic background (e.g., Sullivan et al., 1997) or a population genetic background (e.g., Gysels et al., 2004), and often limit the methods used to investigate lineage divergence to those consistent with their background. The integration of methods designed to detect signal at various positions of the continuum of genetic variation is critical to understanding the full spectrum of evolutionary and ecological forces that have structured genetic diversity. Furthermore, congruent results will reinforce confidence in their accuracy, whereas incongruent results may suggest ways that the data can be improved or instances where assumptions of a particular method may be violated. Although examples of the power of this integrated approach to phylogeography exist (e.g., Wares and Cunningham, 2001), the majority of phylogeographic studies employ a limited range of analytical methods. The importance of methods that attempt to infer signal from wide portions of this continuum is underscored by recent contention with respect nested clade analysis (Knowles and Maddison, 2003;

Templeton, 2004), perhaps the only method that attempts to both account for geography explicitly and to utilize genetic information from across the continuum of genetic variation. Here, we apply a wide range of methods developed to infer signal at different levels of genetic diversity to the investigation of a lineage of salamanders with a disjunct distribution in the Pacific Northwest in order to maximize the inferences possible with the most basic phylogeographic data.

Mesic forests in the Pacific Northwest of North America (PNW) are characterized by both the late-successional dominance of western hemlock (*Psuedotsuga heterophylla*) and western redcedar (*Thuja plicata*), and by mild, wet winters, and cool, relatively dry summers. The PNW can be divided longitudinally into three provinces: the Cascades and Coastal Mountains in the west, the Northern Rocky Mountains in the east, and the intervening xeric Columbia Basin. Mesic forest ecosystems occur as two geographically separate and ecologically isolated entities in the Cascades/Coastal Mountains and in the Northern Rocky Mountains. More than 150 plant, animal, and fungi lineages are known to have disjunct distributions, with populations in both coastal and inland mesic forests of the PNW (Brunsfield et al., 2001; Neilson et al., 2001). The ecosystems of the PNW have been influenced by both ancient and recent geological events. The Northern Rocky Mountains were formed ca. 45 to 36 Mya during the Eocene and pollen records indicate the occurrence of coniferous forests on the western slopes of the Rocky Mountains since the mid-Eocene (Graham, 1993). The Cascades Range was formed during the early Pliocene (5 to 2 Mya), and this orogeny established a rain shadow that dried out the Columbia Basin, resulting in a well-developed steppe

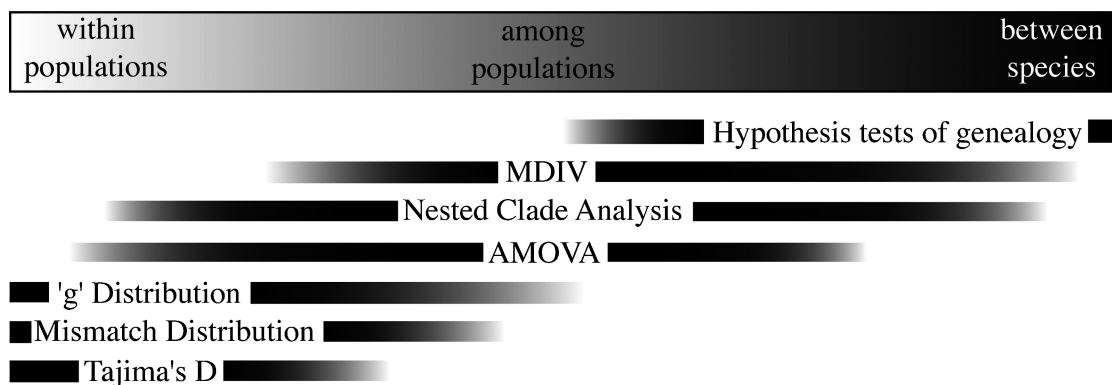


FIGURE 1. The continuum of genetic variation (top) and analytical methods used to interpret the genetic signal (bottom). The bars on either side of the method name show the approximate range to which we have applied these tests.

vegetation by the end of the Pliocene (Daubenmire, 1952; Graham, 1993). Subsequently, glaciers have repeatedly formed and retreated, especially the Cordilleran Ice Sheet in the Northern Rockies (Richmond et al., 1965; Waite and Thorson, 1983; Delcourt and Delcourt, 1993).

This complex history and the broad taxonomic range of disjunct mesic forest taxa has led researchers to propose alternative hypotheses to explain the isolation of inland mesic forest lineages from their coastal counterparts. First, inland elements of the mesic forests could be the result of an ancient vicariance caused by the Pliocene xerification of the Columbia Basin, with subsequent persistence of inland taxa in refugia throughout the Pleistocene glacial cycles (Brunsfeld et al., 2001). This 'ancient vicariance' hypothesis predicts deep divergence and reciprocal monophyly between coastal and inland populations because it allows sufficient time for completion of lineage sorting. A corollary to the ancient vicariance hypothesis is the persistence of inland mesic forests in one or more refugia throughout the Pleistocene. The Clearwater Drainage in northcentral Idaho has been proposed as one such refugium, based on both geology and phylogeography (Daubenmire, 1975). This corollary hypothesis predicts a genetic signature of an expansion from the Clearwater Refugium. An alternate hypothesis explains the populations in the Northern Rockies as the result of recent (i.e., post-Pleistocene) dispersal from the Cascades (Mack et al., 1978; Brunsfeld et al., 2001). The 'recent dispersal' hypothesis predicts shallow (post-Pleistocene) divergence between northern coastal and inland populations, absence of reciprocal monophyly, and very little genetic diversity within and among inland populations. DNA evidence supporting ancient vicariance has been identified in the Tailed frog (*Ascaphus truei*/*A. montanus*) species complex (Nielson et al., 2001; mtDNA), whereas recent dispersal is supported by mtDNA in the water vole *Microtus richardsoni* (Demboski et al., unpublished; mtDNA), and cpDNA in the whitebark pine *Pinus albicaulis* (Richardson et al., 2002).

We focus here on the *Plethodon vandykei* species group, a salamander species complex that exhibits the charac-

teristic disjunction and is most often found in affiliation with seeps or streams in areas with exposed and fractured bedrock (Wilson et al., 1995, 1997). It includes *P. vandykei* and *P. idahoensis*, species that were only recently recognized as specifically distinct (Howard et al., 1993). *P. vandykei* occurs in Washington in three isolated areas: the Olympic peninsula, the Mt. Rainier/Mt. St. Helens region, and Willapa Hills (Nussbaum et al., 1983). *P. idahoensis* is found in northern Idaho, extreme western Montana, and southern British Columbia (Wilson and Larson, 1998a; Wilson and Oshanjanian, 2002), and is isolated from *P. vandykei* by 300 km of desert sage/scrub and xeric forests (Fig. 2; Brunsfeld et al., 2001). Available data, including allozymes (Howard et al., 1993) and morphometrics (Wilson and Larsen, 1998b), indicate that each species is diagnosable. However, because each of those studies lacked a phylogenetic framework, explicit tests of alternative explanations for the disjunction have not been conducted.

Here, we combine statistical phylogenetics with coalescent simulations to assess the geographic structure of mtDNA variation in the *P. vandykei* species group. Specifically, we test the dispersal versus vicariance hypotheses, and test the Clearwater refugium hypothesis. The nested nature of the ancient vicariance/Clearwater refugium hypothesis makes the application of a range of methods that span the continuum of genetic variation particularly appropriate here. The results of this study are then interpreted in light of phylogeographic studies of other mesic forest disjuncts in an ongoing synthesis of the ecosystem genetics of mesic forests in the PNW.

MATERIALS AND METHODS

Sample Collection

We collected tail clips from 243 *P. idahoensis* from 49 localities and 18 *P. vandykei* from 3 localities (see Appendix 1 for detail, available at the Society of Systematic Biologists website, <http://systematicbiology.org>). A single *P. larselli* was collected and used as an outgroup (Mahoney, 2001). The *P. idahoensis* samples are from seven

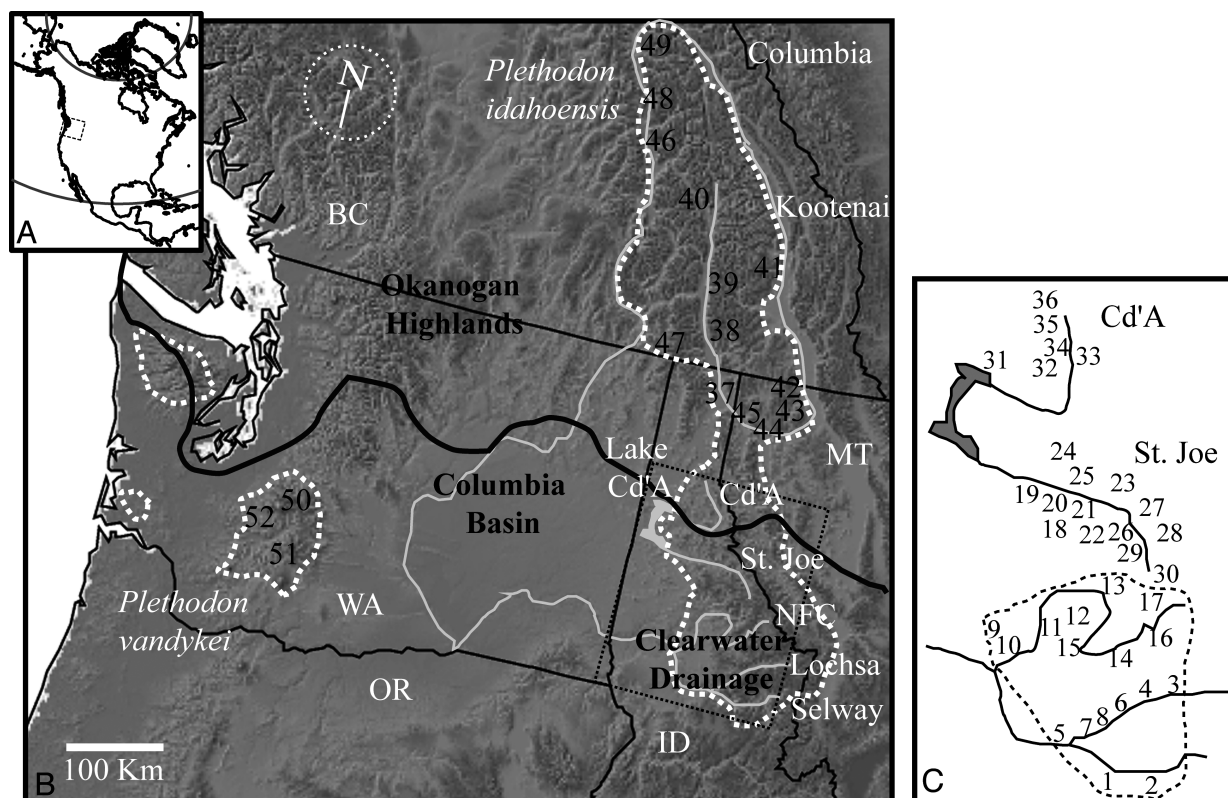


FIGURE 2. (A) Map of North America. (B) Collection localities for *P. idahoensis* and *P. vandykei*. Locality numbers correspond to those in Appendix 1. The approximate ranges of *P. vandykei* and *P. idahoensis* are shown with dotted white lines. The maxima of the Cordilleran Ice Sheet is shown with a black line. States or provinces are abbreviated as follows: Montana (MT), Idaho (ID), Oregon (OR), Washington (WA), and British Columbia (BC). (C) Details of localities in central Idaho. The approximate boundary of the Clearwater River drainage is shown with a dotted line.

river drainages and one lake basin whereas the *P. vandykei* samples are from three river drainages (Fig. 2). Sample sizes ranged from 1 to 13 per locality, with an average $n = 5$. When *P. idahoensis* samples were pooled by drainage, sample sizes ranged from 9 to 60 (Table 1), with an average $n = 30.4$. Therefore, following Saunders et al. (1984; $P = [n - 1]/[n + 1]$), there is a high probability that we have sampled the deepest coalescent event in each drainage ($P = 0.8\text{--}0.97$). However, these estimates should be conservative because ancestral haplotypes are expected to be most frequent in the population (Waterson and Guess, 1977; Crandall and Templeton, 1996).

DNA was extracted from 10 to 20 mg tail clips, which had been stored in 90% EtOH, with either the DNeasy Tissue kit (Qiagen, Inc.; Valencia, CA), following the manufacturer's instructions for rodent tails, or the method described in Nielson et al. (2001). A 669-bp fragment of the mitochondrial cytochrome *b* gene (Cyt *b*) was amplified with the forward primer MVZ 15' (5'-GAAGTAATGGCCACACWWTACGNA-3'; Moritz et al., 1992) and with H-14963 (5'-GGCAAA-TAGAARTATCATT-3'; Sullivan et al., 2000) as the initial reverse primer. For subsequent polymerase chain reaction (PCR) amplification, we designed a lineage-specific reverse primer (5'-AGGAGTGAGAGTAGAGTAAGTA-3'). Amplicons were purified using polyethylene glycol

precipitation, and sequencing reactions were performed with the BigDye Kit (Applied Biosystems, Inc., Foster City, CA) with 10 to 20 ng of PCR product in 10 μ L volumes. CentriSep columns (Princeton Separations,

TABLE 1. Diversity statistics by river drainage for *P. idahoensis*. River drainage, the number of samples collected in that drainage (n), the probability of the samples from the drainage capturing the deepest coalescent event ($Prob$), the number of haplotypes in the drainage ($No. hap$), the number of polymorphic sites (s), the nucleotide diversity (π) ± 1 standard deviation, Tajima's D (with significant values in bold), and the P -value for the mismatch distributions are shown.

Drainage	n	$Prob$	$No. hap$	s	π	D	Mismatch
Clearwater ^a	89	0.98	29	38	0.0104 \pm 0.005	<i>n/a</i>	<i>n/a</i>
Selway River	15	0.88	4	7	0.0015 \pm 0.001	-1.89	0.58
Lochsa River	23	0.92	7	10	0.0024 \pm 0.002	-1.37	0.35
NFC ^b River	51	0.96	18	23	0.0073 \pm 0.004	-0.13	0.65
St. Joe River	60	0.97	13	28	0.0031 \pm 0.002	-2.10	0.80
Coeur d'Alene River	19	0.9	2	1	0.0064 \pm 0.004	-1.81	0.69
Kootenai River	38	0.95	16	37	0.0017 \pm 0.001	-1.97	0.71
Columbia River	28	0.93	8	11	0.0002 \pm 0.000	-1.16	0.73
Lake Coeur d'Alene	9	0.8	1	0	0	0	<i>n/a</i>
<i>P. idahoensis</i> (total)	243	0.99	63	97	0.0069 \pm 0.004	<i>n/a</i>	<i>n/a</i>

^aNFC, Lochsa, and Selway are the major tributaries of the Clearwater drainage.

^bNFC = North Fork of the Clearwater.

Adelphia, NJ) were used to filter sequencing reactions, and samples were run on an ABI 377 automated sequencer using 5% Long Ranger polyacrylamide gels. Cyt *b* was sequenced in both the 5' and 3' directions, and edited and aligned with Sequencer 3.0 (GeneCodes, Ann Arbor, MI). Sequences were deposited in GenBank under accession numbers AY572046 to AY572108.

We used two methods of preliminary data exploration: descriptive diversity statistics and AMOVA. ML distances were computed between *P. larselli* and *P. vandykei*, between *P. vandykei* and *P. idahoensis*, within *P. idahoensis*, and within the Clearwater *P. idahoensis*. In addition, within *P. idahoensis*, we calculated several diversity statistics (nucleotide diversity [π], the number of pairwise differences [k], and the number of polymorphic sites [s]) using ARLEQUIN 2.0 (Schneider et al., 2000). Analysis of molecular variance (AMOVA; Cockerham, 1969; Excoffier et al., 1992) was conducted using ARLEQUIN to investigate population structure by testing the global hypothesis that haplotype variance is partitioned among drainages. We further grouped the seven drainages in which *P. idahoensis* occur into two groups, the Clearwater drainages (Selway, Lochsa, North Fork Clearwater) and the northern drainages (St. Joe, Coeur d'Alene, Kootenai, Columbia).

Phylogenetic Analyses

Phylogeny estimation.—We generated maximum likelihood (ML) and maximum parsimony (MP) estimates of phylogeny to identify the major clades and to test phylogenetic hypotheses in the *P. vandykei* species group. We pruned all redundant haplotypes, and performed searches with PAUP* 4.0 (Swofford, 2002). Under parsimony, heuristic searches included 150 random addition-sequence replicates and TBR branch swapping. Characters were equally weighted, and there was no limit to the number of trees retained. Nodal support was assessed using bootstrap analysis with 1000 replicates (Felsenstein, 1985). We used DT-MODEL (Minin et al., 2003) to select a model; this method incorporates fit, a penalty for overparameterization, and performance into model selection, and is expected to select simpler models that nevertheless provide better branch-length estimates than other alternative model-selection methods. Because model-selection methods cannot assess the adequacy of chosen models, we evaluated the absolute goodness of fit of the chosen model chosen using the parametric bootstrap approach (e.g., Goldman, 1993; Sullivan et al., 2000). We then conducted heuristic searches under ML with the chosen model, and TBR branch swapping, 10 random-addition replicates, and a random starting tree. Nodal support for the ML tree was assessed using nonparametric bootstrap values (Felsenstein, 1985), computed from 100 replicates.

Frequentist hypothesis testing.—We used parametric bootstrap analyses (Huelsenbeck et al., 1996) to test the ancient vicariance and recent dispersal hypotheses. Because ancient vicariance predicts reciprocal monophyly

of *P. idahoensis* and *P. vandykei*, we searched for the best ML topology that refutes reciprocal monophyly of *P. idahoensis* and *P. vandykei* (i.e., a converse constraint). We then used SEQ-GEN (Rambaut and Grassley, 1997) to simulate 100 data sets on the best constrained tree, with model parameters and branch lengths estimated using the ML model selected above. Each simulated data set was then searched with PAUP* for both the best tree and the best tree constrained for nonreciprocal monophyly. The distribution across all replicates of the difference in log-likelihood scores ($\ln L_{\text{unconstrained}} - \ln L_{\text{constrained}}$) served as the null distribution, which was used to assess the difference calculated from the real data. The recent dispersal hypothesis was tested in a similar fashion. This hypothesis proposes that inland populations are the result of dispersal from the Cascades through the Okanogan Highlands (Fig. 1), and predicts that haplotypes from the Columbia River Drainage will be basal to other *P. idahoensis*. Therefore the tree used to test the recent dispersal hypothesis was found by constraining the haplotypes from the Columbian drainage to be basal within the *P. idahoensis* clade.

Bayesian hypothesis testing.—We used MRBAYES (Huelsenbeck and Ronquist, 2001) to assess the posterior probability of each of the two a priori hypotheses (i.e., ancient vicariance and inland dispersal). This approach attempts to determine the posterior probability that some hypothesis is correct, given the model and the data, by using MCMC to estimate the joint posterior probability distribution. It further assumes that the Markov chain is sampling from the true posterior distribution (i.e., the chain has reached stationarity with respect to the parameter of interest; Huelsenbeck et al., 2002) and, because we are assessing conformation to topological predictions in these tests, stationarity with respect to topology is critical. Because the topology parameter may be particularly susceptible to non-stationarity (Huelsenbeck et al., 2002), we employed a stationarity test similar to one proposed by Huelsenbeck et al. (2002). We conducted four independent heated runs (each composed of four Metropolis-coupled chains), and started each run with a different random tree. We ran the chains for 10^7 generations and sampled trees every 1000 generations. If the four independent runs have each converged on the true joint posterior probability distribution, the four samples of trees should represent independent samples drawn from that distribution. In order to assess this statistically, we saved the last 5000 trees from each run and computed the symmetric-difference distance between each tree in the sample and the ML tree (found above) using PAUP*. We then conducted a standard analysis of variance (ANOVA) on the four groups of tree-to-tree distances to assess whether the four chains sampled independently from the same underlying (and hopefully true) joint posterior probability distribution. Although a nonsignificant result for this test would not guarantee that the runs have reached stationarity with respect to topology, it would provide much stronger evidence of such than would the standard examination of $\ln L$ plots. To complete the hypothesis test, we then

imported the sample of trees from the Bayesian analysis into PAUP* and filtered them with the same constraint trees used to test the ancient vicariance and recent dispersal hypotheses. The proportion of trees in the sample consistent with the topology predicted by each hypothesis is the Bayesian posterior probability of that hypothesis.

Population Genetic Analyses

MDIV analysis.—The ancient vicariance hypothesis predicts that the coalescent depth of divergence between *P. vandykei* and *P. idahoensis* will be great due to their isolation since the xerification of the Columbia basin began some 5 Mya. To investigate the depth of divergence, we analyzed the data with MDIV (Neilson and Wakeley, 2001), an MCMC-based program designed to distinguish migration from isolation. We performed several iterations of the analysis before we were satisfied that we were analyzing the data with appropriate priors on M and T . Our initial analysis of *P. vandykei* and *P. idahoensis*, with $M = 10$ and $T = 5$, indicated that the migration rate was close to zero. This was congruent with the results of the phylogenetic hypothesis testing, so we reanalyzed the data with prior values of $M = 0$ and $T = 6$. We conducted 3×10^6 generations of the Markov chain, and then repeated the analysis using a different seed value to assess stationarity. We assumed a mutation rate of 10^{-7} when converting the estimate of θ to N_e , which was then used to convert the posterior distribution of T_{MRCA} to years.

The a priori corollary hypothesis of ancient vicariance, the Clearwater refugium hypothesis (Daubenmire, 1975), predicts a post-Pleistocene population expansion from a refugium located in the Clearwater Drainage. We used two methods, nested clade analysis (NCA) and coalescent estimates of the rate of population growth (g) to investigate *P. idahoensis* for the genetic signature of recent population expansion.

Nested clade analysis.—Of all available population genetic approaches, NCA is the method that most explicitly incorporates geography. We first used NCA to assess the significance of the correlation of haplotypes and geography. Unweighted geographic distances were calculated from latitude and longitude, and TCS (Clement et al., 2001) was used to estimate a haplotype network for *P. idahoensis*. Groupings in this network were nested following Templeton (1998) and ambiguities were resolved following the guidelines in Templeton et al. (1987). GEODIS (Posada et al., 2000) was used to calculate the average geographical distance of the haplotypes to the center of each clade (D_c), the average distance of the haplotypes in each clade to the center of the clade at the next nesting level (D_n), and the difference between tip clades and interior clades for D_c and D_n . We tested for the significant association of haplotype with geography using a categorical permutation contingency analysis, and used the inference key (Templeton, 2004) to infer the likely population process that contributed to the pattern of genetic diversity observed within *P. idahoensis*.

Under the Clearwater refugium hypothesis, population expansion should result in little association of haplotype with geography because there has been insufficient time for lineage sorting to occur among populations derived from the same source, and the NCA should fail to reject the null hypothesis of no association between haplotype and geography. By contrast, upon inclusion only of samples from the putative refugium, the hypothesis predicts a significant association haplotype and geography because these samples represent populations descended from long-isolated populations. As a test of the Clearwater refugium hypothesis, we performed a second NCA using only the samples from the Clearwater drainage to test for a significant association of haplotype and geography.

Coalescent simulations.—Population expansion can also be assessed using methods that directly estimate parameters of an expanding population by including the growth rate (g) in a coalescent model. Such genealogical methods are more efficient than methods based on diversity statistics (Kuhner et al., 1998), particularly when the g is low or negative. Expanding populations produce genealogies that are dominated by the tips of the tree, because most of the coalescent events occur in the recent past after the population expansion. The coalescent model for expanding populations of Kuhner et al. (1998) rescales the total time of the coalescent process so that $g = (1 - e^{-gt})/T$, where t is the original time to the common ancestor and T is the rescaled time. A significantly large g indicates population expansion. We used the program FLUCTUATE (Kuhner et al., 1998), which implements a Metropolis-Hastings genealogy sampler to estimate g for the entire sample of *P. idahoensis* (g_T) and the Clearwater samples alone (g_C). The hypothesis predicts that g_T will be significantly greater than zero, because *P. idahoensis* has expanded into recently deglaciated areas. Genealogical samplers rely on the shape of the likelihood surface to generate confidence intervals (CIs), and under some conditions these CIs have been shown to be too narrow for parameters of structured coalescent models (Abdo et al., 2004). In order to generate a null distribution of g_T , we conducted coalescent simulations under a constant population size model (e.g., $g_i = 0$), with the effective population size (N_e) and mutation rate (μ) parameters estimated from the original data. Because these parameters are interdependent, we used the compound parameter θ , which is equal to $2N_e\mu$ for mitochondrial data, in the simulations. We used MIGRATE (Beerli, 2002) to estimate θ for the Clearwater samples (θ_C) and for all the samples (θ_T). We then simulated 100 data sets with θ_C and 100 data sets with θ_T using TREEEVOLVE (Grassly et al., 1999) and a model of constant population size. All Markov chains in FLUCTUATE were initialized on the empirically estimated values of θ , but were allowed to move into regions of higher log-likelihood, and we used the estimates of g_i from each of these simulated data sets to construct the distribution of g_T and g_C under the null hypothesis of stable populations.

Analyses within drainages.—We explored genetic structure within different populations of *P. idahoensis* by

calculating several diversity statistics for samples collected within a single river drainage. These drainages likely contain isolated populations of *P. idahoensis* because most are separated by ridge lines that exceed the recorded elevation limits of *P. idahoensis* (Nussbaum et al., 1983). Within each drainage, genetic diversity was estimated by calculating nucleotide diversity (π), the number of pairwise differences (k), and the number of polymorphic sites (s) using ARLEQUIN 2.0 (Schneider et al., 2000). These parameters were used in the calculation of Tajima's D (Tajima, 1989), which can be used to test for population expansion (Rogers, 1995). We also tested for population expansion using the demographic expansion model of Rogers and Harpending (1992) to analyze the pairwise mismatch distributions. Populations that have been stable historically are predicted to have multimodal mismatch distributions, whereas those that have undergone a recent expansion are predicted to be unimodal. We calculated the mismatch distribution with ARLEQUIN (Schneider et al., 2000) for samples collected within each of the eight river drainages in which *P. idahoensis* occurs, and used the Raggedness Index of Harpending (1994) to identify populations in which the demographic model of a unimodal mismatch distribution can not be rejected.

RESULTS

We sequenced 669 nucleotides, corresponding to positions 16294 to 16963 of the *Xenopus laevis* mitochondrial gene Cyt *b* (Roe et al., 1985). The pattern of molecular evolution was consistent with mitochondrial DNA of plethodontids and inconsistent with the presence of nuclear pseudogenes. Seven haplotypes were sampled in *P. vandykei*, and 62 haplotypes were sampled in *P. idahoensis* (Table 2). One of the haplotypes (A) was found in 45% of all *P. idahoensis* individuals sampled (109/243), and was sampled in 31 of the 49 populations.

Minimum and maximum ML distances (HKY+ Γ) were as follows: *P. larselli*–*P. vandykei* (min = 0.26411 substitutions/site; max = 0.27034), *P. vandykei*–*P. idahoensis* (min = 0.08529; max = 0.10604), within *P. idahoensis* (max = 0.02776), and within Clearwater *P. idahoensis* (max = 0.02329). With the exception of the Lake Coeur d'Alene sample (which was fixed for haplotype A), all populations of *P. idahoensis* were polymorphic for Cyt *b*, with the largest number of nucleotide polymorphism (s) occurring on the Kootenai River Drainage (Table 1). The AMOVA suggested that most of the molecular variance occurred within drainages (51.69%; $P < 0.001$) and between the southern and northern drainages (37.17%; $P < 0.001$), with only little variation among drainages (11.14%; $P = 0.084$).

Phylogenetic Analyses

Phylogeny estimation.—We selected the HKY+ Γ model of sequence evolution using DT-MODEL (Minin et al., 2003) with equilibrium base frequencies of $\pi_A = 0.305$; $\pi_C = 0.235$; $\pi_G = 0.14$; $\pi_T = 0.32$; transition-transversion ratio = 1.336; and Γ -distribution shape parameter ($\alpha = 0.716$). Furthermore, the absolute goodness-of-fit test indicated that the HKY+ Γ model could not be rejected ($P = 0.64$), indicating that the parametric bootstraps and Bayesian hypothesis tests should not be compromised by reliance on a single substitution model.

There were 91 parsimony-informative sites in the complete data set. We found 2160 MP trees (not shown), with a tree length of 278. The topology of the MP trees was similar to that of two ML trees with log-likelihood scores $-\ln L = 2511.685$. The ML topologies differed from one another in only minor ways, and one of these ML trees is shown (Fig. 3). In the ML estimate of phylogeny, *P. idahoensis* and *P. vandykei* form reciprocally monophyletic clades that are well supported (100% bs_{ML} ; 99% bs_{MP} ; 1.0 Bpp) in both ML trees and MP trees. Within *P. idahoensis*, haplotypes from the Selway River (a tributary of the Clearwater) were monophyletic in the optimal tree, but this clade was weakly supported (61% bs_{ML} ; 54% bs_{MP} ; 0.429 Bpp). Most of the haplotypes from the Lochsa River (another Clearwater tributary) formed a sister clade to the Selway haplotypes in the optimal topology, but this relationship was also poorly supported (60% bs_{ML} ; 46% bs_{MP} ; 0.261 Bpp). One haplotype (bcc66), collected two miles downstream from the confluence of the Selway and

TABLE 2. Number of haplotypes by drainage for *Plethodon idahoensis*. Haplotypes sampled in multiple individuals are labeled A to V (M and U were found in *P. vandykei*). The final column, denoted with the *, lists the number of individuals from each drainage with a haplotype not sampled in any other *P. idahoensis* individual. Haplotype letters correspond to those in Figures 2 and 3.

Drainage	Haplotype																				*
	A	B	C	D	E	F	G	H	I	J	K	L	N	O	P	Q	R	S	T	V	
Selway ^a	.	.	12	3
Lochsa ^a	.	.	.	8	10	5
NFC ^{a,b}	11	3	.	.	.	8	.	.	4	4	4	6	11
St Joe	40	3	6	3	9
Cd'A ^c	18	1
LkCd'A	9	0
Kootenai	16	1	.	2	5	2	3	.	2	10
Columbia	15	1	4	3	.	4

^aNFC, Lochsa, and Selway are the major tributaries of the Clearwater drainage.

^bNFC = North Fork of the Clearwater.

^cCoeur d'Alene River.



FIGURE 3. ML phylogeny estimate for the *P. idahoensis*/*P. vandykei* complex under a HKY+ Γ model of sequence evolution. This is one of two ML trees with a log-likelihood score of -2511.685 . ML bootstrap values from 100 replicates are shown for nodes recovered in $>50\%$ of the bootstrap data sets. Haplotypes are grouped by drainage, and shown in shaded boxes, but haplotype 'A' is found in the North Fork of the Clearwater, the St. Joe, the Coeur d'Alene, the Kootenai, and the Columbia drainages as well as the Lake Coeur d'Alene population. Haplotype 'N' is found on both the Kootenai and the Columbia Rivers, but is sister to a haplotype (haplotype R) found only on the Kootenai River. Haplotype 'O' is found on the St. Joe, but is sister to haplotypes found on the Columbia River. Haplotype 'bcc255' is found on the Kootenai River, but is sister to a clade found on the North Fork of the Clearwater River.

using data from geology, paleoecology, and paleobotany. Such hypotheses predict specific patterns of genetic variation that can be tested while simultaneously accounting for the stochastic nature of the coalescent process and the amplifications of these effects over geological time. Phylogenetic trees are extremely useful for testing the predictions of hypotheses that relate to the deepest genetic structure that may be present in a phylogeographic data set. For example, our ML estimate of phylogeny supports both the ancient vicariance and Clearwater refugium hypotheses and the statistical tests such as the parametric bootstraps or Bayesian posterior tests eliminate phylogenetic uncertainty as an explanation. The congruence between the parametric bootstrap tests and the Bayesian hypothesis tests, coupled with the greater speed of the latter, suggest that Bayesian hypothesis testing may be a more efficient use of computational resources, although power tests are needed to explore the issue. Similarly, both have been criticized as relying too heavily on the adequacy of a particular model of sequence evolution (e.g., Felsenstein et al., 2003; Buckley, 2001). This emphasizes the need to both choose a model rationally and to assess the absolute fit of the data to the model. In our case, the chosen model adequately approximates the unknown process that generated the data.

Due primarily to the stochasticity of the coalescent process, it is considerably more difficult to design phylogenetic tests of recent events; it is not possible to construct appropriate predicted topologies in order to test hypotheses regarding events in the recent past such as post-Pleistocene expansion. Such hypotheses are best evaluated with predictions derived from population genetic theory and test statistics that account for coalescent stochasticity (Knowles and Maddison, 2002). We have investigated post-Pleistocene expansion using several complementary methods. On the among-populations scale, NCA was used to test the prediction that haplotypes within the Clearwater drainage (the location of the putative refugium) would have a significant association with geography. Because NCA cannot be used to test an *a priori* null hypothesis of a rapid expansion (but can identify a signature of population expansion at specific nested clades), the significance of our empirical estimate of the growth rate parameter (g) was assessed by simulating a null distribution. On smaller geographic scales, we validated the expectation of population expansion using Tajima's D and analysis of mismatch distributions.

Perhaps the greatest difficulty of the statistical phylogeographic approach is choosing which test(s) are most appropriate to the question being asked; each of the tests used here has shortcomings that may lead to misinterpretation. One advantage to utilizing a variety of tests designed to detect signal across a wide range of genetic divergence is that the tests can be cross validated. For example, the standard interpretation of a significantly negative Tajima's D is that selection and/or population expansion are present (Tajima, 1989). Similarly, both sampling and unaccounted for genetic structure can lead

to spurious inferences of Tajima's D (Hammer et al., 2003). Nevertheless, we interpret the signal present in our data as population expansion because this result was also found when we estimated the population growth rate and calculated the mismatch distributions. In two drainages, we had non-negative values of Tajima's D but could not reject the unimodal mismatch distribution. We suspect that this discrepancy is related to the low values of π for these drainages; π is incorporated into Tajima's D but not the distribution of mismatches. The NCA results were consistent with our prediction that extant populations within the refugium would retain significant geographic structure; there was a significant association of geography and haplotypes in the Clearwater, and this association degraded when all samples were analyzed. This apparently results from insufficient time for lineage sorting to have occurred in the northern portion of the distribution. In each nested clade that had a significant association of haplotype and geography, the inference derived by use of Templeton's (2004) inference key was one of population expansion (results not shown). This may be one population process that NCA is particularly capable of inferring (Templeton, 1998). The appeal of NCA is that it represents perhaps the only analytical framework that attempts to infer genetic processes ranging from small scale population processes to those acting across a complex geography and on a longer time scale. However, we agree with Knowles and Maddison (2002) that further study is needed on the circumstances under which NCA works well and where it fails. Pending the results of that research, there is a need to cross-validate inferences made with NCA using other data (Templeton, 2004) and other methods.

Genetics of the Plethodon vandykei Species Group

We found deep divergence between *P. vandykei* and *P. idahoensis* with mitochondrial data. The depth of this divergence (corrected distances of 0.0853 to 0.106 substitutions per site), as well as the results of the parametric bootstrap Bayesian hypothesis tests, and coalescent-based estimates of T_{MRCA} all support the ancient vicariance hypothesis. The corollary of the ancient vicariance hypothesis is that *P. idahoensis* persisted throughout the Pleistocene glaciations in a mesic forest refugium, putatively situated in the Clearwater drainage, and subsequently expanded into deglaciated drainages at the close of the Pleistocene. The results of the NCA, as well as our coalescent-based estimates of the growth rate, support recent population expansion. Furthermore, our data suggest that extant populations within the refugial Clearwater drainage have undergone significant post-Pleistocene growth. One possible explanation is that refugial mesic forests within the Clearwater may have been elevationally compressed during Pleistocene glacial maxima, and the retreat of glaciation at the close of the Pleistocene led to an increase of available habitat within the drainage. The three major tributaries that comprise the Clearwater drainage (the North Fork, Lochsa, and Selway Rivers) may also have functioned as independent refugia during

Pleistocene cold periods. These results suggest that our understanding of the Clearwater refugium is preliminary, and that more refined hypotheses are needed to understand the dynamics of this complex refugium. We are currently screening multilocus SNP data to further test the hypotheses that we have refined here. Several authors have suggested that these data offer significant advantages for phylogeographic studies (Arbogast et al., 2002; Brumfield et al., 2003; Funk and Omland, 2003).

Ecosystem Phylogeography

These results increase our understanding of the assembly of the inland mesic forest ecosystem in the Pacific Northwest. Three other vertebrate lineages exhibit disjunct mesic forest distributions: tailed frogs (*Ascaphus truei*/*A. montanus*), giant salamanders (*Dicamptodon* spp.), and water voles (*Microtus richardsoni*). Studies of these taxa suggest that the vertebrate components of this ecosystem were not assembled in a concerted manner. Good (1989) studied genetic diversity in *Dicamptodon* using allozymes and found that the deepest divergence was between coastal and inland populations. He considered this divergence deep enough to recommend recognition of the inland forms as a distinct species (Good, 1989). Nielsen et al. (2001) found a similar pattern in *Ascaphus* using data from two mitochondrial genes, and also argued for the recognition of the inland *Ascaphus montanus* as a distinct species. Although the distribution of these lineages, as well as that of the *P. vandykei* species group, is best explained by an ancient vicariance and persistence through the Pleistocene glaciations, the pattern found in *Microtus richardsoni* suggests a northern dispersal (J. R. Demboski and J. Sullivan, unpublished). Demboski et al. (in preparation) have found shallow genetic divergence in the water vole, and haplotypes that are shared between populations from the northern Cascades and the northern Rocky Mountains. This northern dispersal also appears to explain the distribution of white-bark pine (*Pinus albicaulis*; Richardson et al., 2002), a subalpine disjunct.

Based on certain common life history traits, the similarity in the genetic pattern between tailed frogs and Pacific giant salamanders might be expected. Adults of each have lungs and live in the wet habitat surrounding high gradient mountain streams and both groups have larval stages that utilize the same streams. However, the life history of *Plethodon vandykei* and *P. idahoensis* is quite different; the genus *Plethodon* is characterized by direct development (no free-living larval stage), and is lungless and dependent on ambient moisture for cutaneous respiration. Although *P. vandykei* and *P. idahoensis* occupy the same broad ecosystem as *Ascaphus* and *Dicamptodon*, they primarily exploit rocky seeps and talus slopes rather than high gradient streams. They also have much lower reproductive capacities, laying 4 to 12 eggs per year, relative to *Dicamptodon*, which lays 135 to 200 eggs per year, or *Ascaphus*, which lays 44 to 75 eggs per year (Nussbaum et al., 1983). In light of

these life histories, the differences among the northern distributional limits of the three inland species is striking. *Dicamptodon aterrimus* is unknown from north of the St. Joe River Drainage (see Fig. 1), whereas *Ascaphus montanus* is known from the extreme southeastern portion of British Columbia, apparently reaching no more than 50 km into Canada (Dupruis, 2002). By contrast, *Plethodon idahoensis* has recently been found to extend 95 km north of Revelstoke, British Columbia (our locality no. 49; Wilson and Ohanianian, 2003), which is several hundred kilometers north of the international border and 300 km farther north than *Ascaphus montanus* reaches. The northern extent of our sampling indicates that the range of *P. idahoensis* has expanded over 600 km since the close of the Pleistocene. This rate of expansion, >50 m annually, is striking in comparison to the slower apparent rates of *A. montanus* and *D. aterrimus* (assuming both were present in the Clearwater Refugium; genetic data for each suggest this; Nielson et al., in preparation; Carstens et al., in preparation). The best explanation for this rapid expansion may be the ability of *P. idahoensis* to exploit ephemeral habitat created in the wake of retreating glaciers. The wet moraines deposited by retreating glaciers are in many ways ideal habitat for *P. idahoensis*, and deglaciation may have created temporary colonization routes that allowed them to expand their range to a greater degree than the other codistributed amphibians.

CONCLUSIONS

The advantages of statistical phylogeography may be most pronounced in the comparison across all the taxa that comprise a given ecosystem. Organisms as diverse as bryophytes and conifers, millipedes and salamanders, chipmunks and fungi have vastly different life histories and will necessarily be studied with data from divergent genomes. Such data can best be synthesized into ecosystem phylogeography through rigorous statistical tests of independent a priori hypotheses. It is this commonality of such testable hypotheses that will lead to a significant improvement in our understanding of regional phylogeography and the historical assembly of ecosystems.

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