EVOLUTIONARY HISTORY OF NORTHERN HEMISPHERE NUCELLA
(GASTROPODA, MURICIDAE): MOLECULAR, MORPHOLOGICAL,
ECOLOGICAL, AND PALEONTOLOGICAL EVIDENCE

TIMOTHY M. COLLINS,1,2 KENNETH FRAZER,1,3 A. RICHARD PALMER,4
GEERAT J. VERMEIJ,5 AND WESLEY M. BROWN1,6
1Department of Biology, University of Michigan, Ann Arbor, Michigan 48109-1048
E-mail: WBrown@Biology.LSA.UMich.edu
2Department of Zoology, University of Alberta, Edmonton, Alberta T6G 2E9, Canada
and Banff Field Marine Station, British Columbia V0R 1B0, Canada
E-mail: Rich.Palmer@UAlberta.CA
3Department of Geology, University of California, Davis, California 95616
E-mail: GJVermeij@UCDavis.edu

Abstract.—By combining data from a variety of sources we explore patterns of evolution and speciation in Nucella, a widely studied genus of shallow-water marine neogastropods. We present a hypothesis of phylogenetic relationships for all of the currently recognized species of northern hemisphere Nucella, based on an analysis of 718 base pairs of nucleotide sequence from the mitochondrial cytochrome b gene. The order of appearance of species in the fossil record is congruent with this hypothesis. The topology of the inferred phylogeny of Nucella, coupled with ecological, morphological, and fossil evidence, was used to address three main questions: (1) At what time and by which route was the North Atlantic invaded from the North Pacific compared to prior studies of the trans-Arctic interchange? (2) Do patterns of molecular variation within species corroborate the importance of climatic cycles in driving speciation in temperate marine animals? (3) Was radiation in the direction of increased or decreased ecological specialization, body size, or vulnerability to predation?

Molecular evidence confirmed that the sole North Atlantic species, *N. lapillus*, arose from a North Pacific ancestor. Biogeographic and paleontological evidence supported the dispersal of *Nucella*, and perhaps other intergrade species, via the Eurasian Arctic. Rather intriguingly, the linkage of *N. lapillus* to a western as opposed to eastern Pacific clade, and the biogeographic origins of the eastern Pacific species, parallel closely similar patterns observed in another genus of rocky-shore gastropods, *Littorina*. This congruence, in conjunction with information on the climatic and geographic histories of the region, as well as the geographic arrangement of mtDNA haplotypes within *Nucella* species, supports a model of speciation in *Nucella* driven by cycles of climatic amelioration and deterioration that began during the Miocene. Calibrations from the fossil record of *Nucella* suggest that third position transitions and transversions accrue at a rate of 3–4%/my and 0.5%/my respectively. This supports an early participation by *Nucella* in the trans-Arctic interchange, as suggested by paleobiogeographic studies.

Consistent with the unstable taxonomic history of species of *Nucella*, we found few nonmolecular traits to be phylogenetically informative. Among North Pacific species, more recently derived species (*N. camaliculata* and the *N. emarginata* clade) were more ecologically specialized (narrower diet and habitat range). Consistent with extensive intraspecific variation, shell traits were quite labile evolutionarily, neither overall size nor development of antipredatory traits exhibited consistent evolutionary trends over the history of the genus. Nacreous eggs (unfertilized eggs consumed by developing embryos) were an ancestral trait that was lost evolutionarily in the two clades that also exhibited increased body size, suggesting that these two life-history traits may be coupled. The reduced number of chromosomes in *N. lapillus* is clearly a derived state and is consistent with White's (1978) observations on chromosome evolution in other clades.

Key words.—Bering Strait, biogeography, character evolution, climate, cytochrome b, invasion, mtDNA, phylogeny, rate of molecular evolution, speciation, vicariance.

Received February 15, 1995. Accepted June 20, 1996.

Species in the neogastropod genus *Nucella*, the dog whelks, offer excellent opportunities to test hypotheses about geographical, ecological, and temporal aspects of speciation. This genus comprises six living North Pacific and one North Atlantic species of abundant predatory snails whose ecology, genetics, shell variation, reproductive biology, and behavior have been extensively studied on rocky shores (Crothers 1985; Hughes 1986; Strathmann 1987; Palmer 1990 and references therein). Its fossil record, spanning an interval of about 25 million yr from the Early Miocene to the Recent, includes all the living species as well as several extinct forms. We add to this a hypothesis of relationships for *Nucella*, based on a comparison of 718 base pairs (bp) of nucleotide sequence from the mitochondrial cytochrome b gene.

The strength of a multidisciplinary approach to speciation lies both in the more complete picture of the evolutionary milieu, and in the increased confidence in inferences supported by independent data sets. A phylogenetic hypothesis, in this case generated from sequence data, is necessary for the analysis of patterns of species distribution, speciation, the evolution of morphological and ecological characters, and adaptation (Maddison and Maddison 1992). In addition, for the first time, we characterize patterns of molecular variation
in the mitochondrial cytochrome \( b \) gene of numerous marine gastropod species.

Several authors (e.g., Golikov and Tzvetkova 1972; Reid 1990a,b) have suggested that fluctuations in climate leading to shifts in geographica levels during the Late Cenozoic have provided conditions favorable for genetic isolation and subsequent speciation in temperate marine organisms. *Nucella*, along with hundreds of other marine animals and plants, probably invaded the Atlantic Ocean from the North Pacific through the Bering Strait after the latter seaway opened during the Early Pliocene (see Vermeij 1991 and references therein). This inference was based on fossil occurrences and on qualitative similarities between the shells of the North Atlantic *N. lapillus* and several Pacific species. We map the distributions of living species onto a molecular phylogeny to illuminate the biogeographic history of the genus in the North Pacific, to test the hypothesis that *Nucella* participated in the so-called trans-Arctic interchange, and to corroborate the direction and timing of the interchange inferred from the fossil record. Molecular data are also used to discover patterns of geographic variation within *Nucella* species, which are then related to processes thought to be driving differentiation in *Nucella* and similarly distributed taxa.

Calibration of molecular rates with the fossil record, or other biogeographic events, gives a rough time scale to both within and between species processes in *Nucella*, and allows checks of their correspondence, on an absolute time scale, to independently dated climatic or biogeographic events. The fossil record, in combination with the phylogeny, yields minimum divergence times for cladogenetic events, gives some insights into the temporal and geographic origin of species, and presents us with information about the numbers of extinct species and their characters.

These independent data sets result in inferences that may be supported or contradicted by inferences from the other data sets. Inferences supported by several independent data sets may be considered highly corroborated, and are likely to reflect the true evolutionary history of a clade. For example, the study of character state transformations can reveal the likely history of a character of interest. The phylogenetic hypothesis presented here makes predictions about the probable direction of character state transformations and combinations of characters in basal versus derived species. We test these predictions by examination of the characters and temporal distributions of fossil species. The phylogeny of living *Nucella* species also makes predictions about the order of stratigraphic appearance of taxa, which we compare to the known fossil record. Time scales derived from the combination of fossil and molecular data can be compared to independently derived calibrations from other events, in this case the Pliocene emergence of the Isthmus of Panama, as an independent check on both.

**Materials and Methods**

**Taxonomy of Nucella**

The genus *Nucella* as used here in the restricted sense contains Miocene to Recent species from the North Pacific and North Atlantic Oceans. On the basis of anatomical studies and a phylogenetic analysis of muricid groups, Koel (1993a,b) has shown that *Nucella* is a member of the muricid subfamily Oceanenbrinae and is therefore subfamilially distinct from the rapainian genus *Thais*, in which *Nucella* was included by many previous authors. South African species referred to *Nucella* (Kilburn and Rippey 1982) differ from northern species in characters of the egg capsule and shell, and have been excluded from *Nucella* (Vermeij 1993a). The relationship between the South African taxa formerly referred to as *Nucella* and Northern Hemisphere *Nucella* will be the subject of a later paper. Several authors have considered the taxa *Acanthina* and *Acanthuncella* from the Pacific coasts of North and South America as subgenera of *Nucella*. *Nucella* may share a common ancestor with *Acanthina* and *Acanthuncella*, but there is no evidence at present that the latter two genera are derived from any member of *Nucella* (Amano et al. 1993; Vermeij 1993a).

Most species of *Nucella* are well defined, and can be recognized based on apertural shape and the number and pattern of spiral cords on the penultimate and body whorls. However, we and others (Dall 1915) had difficulty confirming the identity of western Pacific species. Russian authors (Golikov and Kussakin 1978; Egorov 1992) generally recognize four species: *N. lima* (Gmelin 1791), *N. freycinetii* (Deshayes 1841), *N. heyseaana* (Dunker 1882), and *N. elongata* (Golikov and Kussakin 1962). In southern Hokkaido, where *N. lima* does not occur, Habe (1958) recognized only two species, *N. freycinetii* and what he called *N. lamellolosa* (Gmelin 1791). This latter taxon was subsequently named *N. elongata* by Golikov and Kussakin (1962). Identified shells sent by A. N. Golikov to GIV and ARP overlap so much in form and sculpture that we are unable to associate diagnostic shell traits with specific names. In view of the extensive intraspecific variation known to occur in species of *Nucella* and the overzealous naming of shell variants by early malacologists (reviewed in Palmer et al. 1990), we have opted to pool information for these three nominal species under the name that has taxonomic priority, *N. freycinetii*.

The status of *Nucella emarginata* (Deshayes 1839) also remains unsettled. Palmer et al. (1990) showed that this name applies to two species, a southern form from California and Baja California and a northern form ranging from southeastern Alaska to California. Deshayes's syntypes are clearly of the "southern" species (ARP pers. obs.), but the identity of the "northern" species remains in doubt. We thus adhere to the recommendations of Palmer et al. (1990) and refer to these as the "northern" and "southern" species of *Nucella emarginata*. Recent collections by one of us (GIV) suggests the possibility that a southern variant of *N. canaliculata* exists, paralleling the pattern found in *N. emarginata*. This form, referred to by Dall (1915) as *Thais (Nucella) canaliculata var. compressa*, exhibits some unique morphological characteristics, and could represent a distinct species most closely related to typical *N. canaliculata*, but more study will be required to test this hypothesis.

To ensure consistent identification of samples used for molecular analysis, voucher specimens for all but *N. lapillus* were inspected by ARP. Finally, we examined specimens (usually including type material) of nearly all fossil species of *Nucella*. Difficulties in the interpretation of fossil taxa are dealt with in the Fossil Record section of the Results.
DNA Sequences

All species were sampled over a significant portion of their geographic ranges (from approximately 250 km for *N. freycinetii* to over 3000 km of coastline for *N. lima*), except for *N. emarginata-S*, which was sampled from a single locality (Table 1; Fig. 1). DNA samples were prepared from individual animals by extraction of total DNA from fresh, frozen, or 70% ethanol-preserved tissues following a slight modification of the protocol of Doyle (1990). Heart tissue samples (approximately 0.01–0.10 g) were homogenized in disposable microfuge tissue grinders (Kontes Scientific Instruments cat 749520) with 700 μL 2 x CTAB buffer, after which 5 μL aliquots of 20 mg/ml Proteinase K were added and the samples incubated at 65°C for 1 h. Samples were then extracted with phenol:chloroform:isoamyl alcohol (25:24:1), followed by a chloroform:isoamyl alcohol (24:1) extraction and precipitation with 1 volume of isopropanol at −20°C overnight. DNAs were pelleted in a microfuge, resuspended in 100 μL TE (10 mM Tris pH 8.3, 1 mM EDTA), brought to a final concentration of 2.5 M ammonium acetate and precipitated by adding 2.5 volumes of ice-cold ethanol. After a second pelleting in a microfuge, samples were air dried and resuspended in 1/10 TE. Amplifications were performed in 100 μL volume of a solution containing 10 mM Tris (pH 8.3), 50 mM KCl, 1.5 mM MgCl2, 0.01% gelatin (Difco), each primer at 0.5 μM, each dNTP at 200 μM, 10–1000 ng template DNA, and 2 μL of Taq Polymerase (Perkin-Elmer/ Cetus). The primers used were designed by alignment of the complete cytchrome b sequences of the neogastropod, *Plicoperpus columellaris* (Collins and Brown unpubl. data), a chiton, *Katharina waukata* (Boore and Brown 1994), a fly, *Drosophila yakuba* (Clary and Wolstenholme 1985), and two urchins, *Strongylocentrotus purpuratus* (Jacobs et al. 1988) and *Paracentrotus lividus* (Cantatore et al. 1989). Primers were chosen from regions of high sequence conservation among these taxa, with any differences being resolved in favor of the gastropod sequence. A HindIII site and three additional A nucleotides were added to the 5' end of each primer to facilitate cloning. The primer sequences and the position of the 3' end of the primer in the human mitochondrial sequence (Anderson et al. 1981) are, for primer 1: 5' - AAAAAAATCCTTCCCTTCAATCTCCTGATTGATGAAA-3' (14.841), primer 2: 5' - AAAAAAGCTTAAATGCGCAGGATAGCATAAGCAA-3' (15.573). These primers amplify a 731 base pair portion of the cytchrome b gene in *P. columellaris*.

PCR protocols are a slight modification of Kreitman et al. (1989). The PCR cycling parameters for double stranded amplification were 30 sec to 1 min at 94°C, 45 sec to 1.5 min at 55°C, and 1.25 to 1.5 min at 72°C, for 24 to 28 cycles. Amplified products were chloroform extracted, precipitated, resuspended in TE, electrophoresed on 2–3% Nu-Sieve TAE agarose gels (FMC BioProducts), and stained with ethidium bromide. DNA bands were visualized under long wavelength (305 nm) UV, excised with glass cover slips, and purified with the GeneClean kit (Bio 101 company). One-quarter to one-third of the double-stranded product was then used in a second, single-stranded amplification identical to the first amplification except for the addition of a single primer and the following cycling parameters: 30 sec to 1 min at 94°C, 45 sec to 1.5 min at 65°C and 1 to 1.5 min at 72°C, for 19 to 20 cycles. This single-stranded product was chloroform extracted, concentrated and purified with Ultrafree-MC UFC3 TTK 30,000 NMWL microconcentrators (Millipore). Sequencing was performed with Sequenase 2.0 (United States Biochemical) according to the manufacturer's suggestions and using the PCR primer that was not used in the single-stranded amplification. Internal sequencing primers were designed based on the sequence information from these first sequencing reactions.

Phylogenetic Analyses

The outgroup taxa used for all analyses were *P. columellaris* and *Plicoperpus patula*, two Neotropical representatives of the Rapaninae subfamily (sensu Kool 1993a,b) of the Muricidae that form a gerrate pair in the western Atlantic and eastern Pacific (Vermeij 1978; Collins 1989). Within the Muricidae, the subfamily Rapaninae is the sister clade to the subfamily Ocenebrinae, which includes *Nuclaea*. The use of these taxa as outgroups has the additional advantage of allowing an independent temporal calibration based on the emergence of the Isthmus of Panama. Estimates of rates of sequence divergence were derived from comparisons of pairwise and patristic differences between taxa calibrated by fossil occurrences and biogeographic events. Calibration points are listed with fossil occurrences in the Appendix.

 Parsimony analyses were performed with PAUP (Ver 3.0q, Swofford 1990). Characters were designated as unordered when equally weighted, and multiple states within taxa for a given character were treated as polymorphisms. Phylogenetic analyses using parsimony were limited to potentially informative characters and were carried out on two differently compiled data sets. In the first data set, referred to as the INDIV data set, single individuals, representing all of the different haplotypes found in each of the currently recognized species, were analyzed in different combinations and in toto. In the second set, hereinafter referred to as the POLY data set, composite sequences for each of the currently recognized species were created, representing all of the polymorphisms found within the individuals sampled for that taxon. The only exception to this was *N. lima*, for which the magnitude and geographic patterning of molecular variation suggested the possibility that two distinct lineages were combined under one species name. These lineages were treated separately as the eastern (Kočiak and Sitka) and western (Shemya and Amchitka) forms of *N. lima*. These two types of data set were analyzed to evaluate the effect of intraspecific polymorphism on phylogenetic reconstruction among closely related species.

Transversions were weighted relative to transitions at three levels: 8:1 weighting (the value used unless otherwise specified), equal weighting, and transversions only. The rationale for these weighting schemes will be discussed below (see Molecular Evolution and the Phylogeny of Nuclaea). Weighting was achieved by appending a stepmatrix to the data file. The PAUP program does not provide basic statistics other than tree length for stepmatrix characters. Trees are based on exhaustive or branch and bound searches. A decay index
<table>
<thead>
<tr>
<th>Taxon</th>
<th>Geographic range</th>
<th>Samples for DNA analysis§</th>
<th>Collected by (year)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. lima</td>
<td>N. Japan to N. Vancouver Is.1,2</td>
<td>Shemya Is. (1 C), Amchitka (1 C), Kodiak Is. (1 B), and Sitka (1 A), AK</td>
<td>GJV &amp; ARP (1987)</td>
</tr>
<tr>
<td>N. freycineti</td>
<td>N. Houttu and Hokkaido to Bering Sea, absent from Aleutians1,2</td>
<td>Tsukuskikoi, E. Hokkaido (49, 1 E)</td>
<td>GJV (1988)</td>
</tr>
<tr>
<td>N. lamellosa</td>
<td>Aleutian Is. to Santa Cruz, CA1,4, absent from Krale and Commander Is.</td>
<td>Cattle Pt., San Juan Is., WB (1 J, 1 K, 1 L)</td>
<td>FBM (1991)</td>
</tr>
<tr>
<td>N. canaliculata</td>
<td>Aleutian Is. to Monterey, CA1,4</td>
<td>Kodiak Is. (1 J), Cordova (21), AK</td>
<td>GJV &amp; ARP (1987)</td>
</tr>
<tr>
<td>N. emarginata-N</td>
<td>Yakutat, AK to Pt. Conception, CA1,5</td>
<td>Queen Charlotte Is., BC, Canada (1 Q)</td>
<td>ARP (1985)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Dutch Harbor, AK (1 N)</td>
<td>GJV &amp; ARP (1987)</td>
</tr>
<tr>
<td>N. emarginata-S</td>
<td>Half Moon Bay, CA to Baja California1,5</td>
<td>Toch Bay, AK (1 Q)</td>
<td>ARP (1982)</td>
</tr>
<tr>
<td>N. lapillus</td>
<td>Sole Atlantic species, Novaya Zemlya to Portugal in the east: Greenland to Long Land in the west</td>
<td>Norris Pt., Bonne Bay, Nfld., Canada (1 G)</td>
<td>GJV (1990)</td>
</tr>
</tbody>
</table>

§ See Vermeij et al. (1990) for details of Aleutian Is. collecting sites; items in parentheses are the number of samples for which 718 bp of contiguous sequence were obtained and the letter of the haplotype(s) found (see Fig. 1 for map of localities; GenBank for haplotype designations).


Latitude and longitude: 1. 43°21'N, 160°57'E; 2. 48°21'N, 122°58'W; 3. 58°19'41"N, 136°47'56"W.

(Bremer 1988; Donoghue et al. 1992) was calculated to indicate the relative support for clades found in the most parsimonious tree.

Maximum-likelihood distance trees were generated with PHYLIp version 3.4 (Felsenstein 1991). Distance matrices were constructed by the DNADIST81 program on the following settings: maximum-likelihood distance, equilibrium frequencies of the nucleotides assumed to be the observed frequencies in the input sequences, transition/transversion ratios of 1:1, 8:1, 100:1, and three categories of substitution rates for first, second, and third positions of codons in the ratios of 1:1.1 and 6:1:44. Ratios for categories were derived from empirical results. A tree was then constructed from the distance matrix with the FITCH81 program with the power

---

**Fig. 1.** North Pacific localities referred to in Table 1 and in text.
TABLE 2. Ecological and morphological characteristics of *Nucella* species.

<table>
<thead>
<tr>
<th>Trait</th>
<th><em>lima3</em></th>
<th><em>freycineti</em></th>
<th><em>lapillus</em></th>
<th><em>langsdorfi</em></th>
<th><em>canaliculata</em></th>
<th><em>emarginata-N</em></th>
<th><em>emarginata-S</em></th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ecological traits*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wave exposure regime</td>
<td>0–4</td>
<td>0–7</td>
<td>0–5</td>
<td>0–3</td>
<td>3–6</td>
<td>2–5</td>
<td>2–5±</td>
<td>1</td>
</tr>
<tr>
<td>Max. height on shore</td>
<td>5</td>
<td>4–52</td>
<td>6*</td>
<td>4</td>
<td>5</td>
<td>6</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>Subtidal populations</td>
<td>common</td>
<td>common</td>
<td>common</td>
<td>common</td>
<td>rare</td>
<td>none</td>
<td>none</td>
<td>2</td>
</tr>
<tr>
<td>Gastropods &gt; 5% of diet</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>yes</td>
<td>3</td>
</tr>
<tr>
<td>Nonmytilid bivalves in diet</td>
<td>cockles</td>
<td>oysters</td>
<td>oysters</td>
<td>cockles</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>3</td>
</tr>
<tr>
<td>Nurse eggs per embryo</td>
<td>~20</td>
<td>polymorphic*</td>
<td>&gt;15§</td>
<td>6</td>
<td>≤1</td>
<td>~30</td>
<td>&gt;20†</td>
<td>4</td>
</tr>
<tr>
<td>Shell traits*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ave./max. adult shell length (mm)</td>
<td>30/60</td>
<td>35/(20)†</td>
<td>27.4/60</td>
<td>45/(30)†</td>
<td>30/50</td>
<td>20/35</td>
<td>20/35</td>
<td>5</td>
</tr>
<tr>
<td>Well-developed varices</td>
<td>no</td>
<td>common</td>
<td>rare</td>
<td>common</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>6</td>
</tr>
<tr>
<td>#Maj. spiral ribs/body whorl</td>
<td>8–10</td>
<td>7–10</td>
<td>7–8§</td>
<td>7–8</td>
<td>10–12</td>
<td>5–6</td>
<td>5–6</td>
<td>5, 6</td>
</tr>
<tr>
<td>#Maj. spiral ribs/penultimate</td>
<td>3–4</td>
<td>2</td>
<td>2§</td>
<td>2</td>
<td>4–5</td>
<td>2</td>
<td>2</td>
<td>5, 6</td>
</tr>
<tr>
<td>Well-developed apertural teeth</td>
<td>no</td>
<td>not</td>
<td>yes</td>
<td>yes</td>
<td>no</td>
<td>no</td>
<td>no</td>
<td>6</td>
</tr>
</tbody>
</table>

*The molecular data suggest *N. lima* may be composed of two cryptic species, hence characteristics reported here may be for either species.

All information on *N. freycineti* includes information gathered for *N. freycineti*, *N. baynesi*, and *N. elongata* (see Taxonomic Considerations section in methods) from Gelinsky and Kussakin (1978) except †.

1 All information on *N. lapillus* from Corineth (1985) except § and *.

4 Details of ecological traits: wave exposure: qualitative ranking; 0 = protected bay; 1 = exposed; 2 = exposed secondarily; 3 = exposed, thirdarily; 4 = extreme exposed.

5 Details of shell traits: shell length was measured from the apex to the tip of the apical canal. Only those greater than 2 mm wide were considered well developed. Major spiral ribs were scored by counting only the most prominent ribs exhibited on the body whorl (between the tip of the apical canal and the first scroll around the second whorl); the penultimate whorl between the first and second scroll adjacent to the apertural margin. Only apertural teeth > 0.1 mm high were considered well developed.

Sources: T Palmer (1980); ARP (unpubl. obs.); * West (1986); Palmer (1988); Spiget (1976); Rivest (1981); Strathmann (1985); * Dall (1915); A Abbott (1974); ARP and GSV (unpubl. obs.); * Ahmed and Sparks (1970), Bannock and Cockayne (1973); § source for the trait; † ARP (unpubl. obs.); * Menge (1976); ** none (ARP unpubl. obs.) Ymany (Kawar, 1995).

set to 2 (Fitch-Margoliash method), no negative branch lengths allowed, global rearrangements, and randomized entry of sequences.

Ecological, Reproductive, and Morphological
Attributes of Living *Nucella*

Nearly all information on ecological and reproductive traits in *Nucella* was obtained from the literature. We report these values as precisely as possible, given the availability of published data. Information on most shell traits was compiled by ARP and GSV based upon shells in the Vermeij collection at the University of California, Davis. Details of traits, measurement protocols, and sources are in the footnotes to Table 2.

These data were used to analyze evolutionary trends and patterns of character evolution in *Nucella*, but were not used in the phylogenetic analyses. We excluded the morphological and ecological traits for two reasons. First, several traits are continuous and highly variable within, as well as between, species and we could not assign discrete character states with confidence. Second, some species (e.g., *N. freycineti*) are not nearly as well known as others (e.g., *N. lapillus*). Thus, while we feel that the coarse categorizations presented here are sufficient for outlining broad evolutionary patterns within *Nucella*, they are not yet refined or complete enough for all species to be included in the phylogenetic analysis or used in sensitivity tests (Maddison and Maddison 1992). At such time as these characters become more fully known, their inclusion in the phylogenetic analysis will be appropriate, although the potential problem of circularity or bias in the study of character evolution (Brooks and McLennan 1991; Maddison and Maddison 1992) needs to be considered.

RESULTS

Phylogenetic Analyses

The following analyses are based on variation within 718 bp of contiguous cytochrome b sequence, deposited in GenBank under the accession numbers U69710–U69728. Average nucleotide composition on the coding strand of *Nucella* species was 40% T, 25% A, 18% C, and 17% G. Individual species differed by less than 2% from these averages, indicating that marked shifts in overall nucleotide bias have not occurred among these taxa. Nucleotide bias was most pronounced at third positions of codons, 83% of which were either A or T (Table 3). Nucleotide composition varied most markedly among species at the third positions of codons, and in some cases deviated more than 5% from the average values of 49% T, 34% A, 11% C, and 6% G. Amino acid differences in this region of cytochrome b, as inferred using the "invertebrate" mitochondrial genetic code (MacVector Vers. 3.5, International Biotechnologies), were too few to be of use for phylogenetic studies within *Nucella*.

A total of 165 nucleotide sites (23% of total sites) varied among the *Nucella* individuals sequenced. These differences were unequivocally distributed among the three codon positions, a pattern typical of mitochondrial protein-coding genes in other animals (Brown 1985). Variation at third positions, which are predominantly silent, accounted for 86.5% of the differences, with first and second positions accounting for 11.5% and 2% of the remaining variable sites, respectively. The type of nucleotide difference was also biased: 79.5% of the variable sites differed by transitions, 8.5% by transversions, and 12% by both. Pairwise sequence differences ranged
Table 3. Nucleotide composition by codon position for Nucella and the outgroup species. Numbers listed are percentages of each nucleotide for first, second, and third positions of codons, as well as the composition for the total region sequenced.

<table>
<thead>
<tr>
<th>Nucleotide-</th>
<th>T</th>
<th>C</th>
<th>A</th>
<th>G</th>
</tr>
</thead>
<tbody>
<tr>
<td>Taxon</td>
<td>1 2 3 1 2 3 1 2 3 1 2 3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P. columellaris</td>
<td>32.6 42.7 44.6 40.9 16.7 22.2 13.3 17.4</td>
<td>21.8 20.5 32.5 24.9</td>
<td>38.9 14.6 0.96 17.7</td>
<td></td>
</tr>
<tr>
<td>P. pelta</td>
<td>32.2 43.3 50.8 41.8 18.0 22.6 12.3 16.4</td>
<td>21.3 20.3 33.8 25.2</td>
<td>28.5 14.6 0.67 16.6</td>
<td></td>
</tr>
<tr>
<td>N. lima</td>
<td>28.5 41.8 43.8 38.0 20.9 22.2 15.0 19.4</td>
<td>20.1 20.5 32.1 24.2</td>
<td>30.5 15.5 0.92 18.4</td>
<td></td>
</tr>
<tr>
<td>N. freycineti</td>
<td>28.9 41.8 47.9 39.6 20.9 22.2 12.1 18.4</td>
<td>21.3 20.5 37.1 26.3</td>
<td>28.9 15.5 0.92 15.7</td>
<td></td>
</tr>
<tr>
<td>N. lapillus</td>
<td>30.1 41.8 52.1 41.4 19.7 21.8 10.1 16.2</td>
<td>20.9 20.5 35.8 25.8</td>
<td>29.3 15.9 0.90 16.7</td>
<td></td>
</tr>
<tr>
<td>N. lamellosa</td>
<td>28.9 41.8 51.7 40.8 20.9 22.2 18.7 17.3</td>
<td>21.3 20.1 35.0 25.5</td>
<td>28.9 15.9 0.46 16.4</td>
<td></td>
</tr>
<tr>
<td>N. canaliculata</td>
<td>29.3 41.8 48.8 40.0 20.5 22.2 18.0 17.8</td>
<td>21.3 20.5 34.2 25.8</td>
<td>28.9 15.5 0.62 16.9</td>
<td></td>
</tr>
<tr>
<td>N. emarginata-S</td>
<td>30.5 41.4 50.0 40.7 19.2 22.6 11.7 17.8</td>
<td>20.9 20.5 30.8 24.1</td>
<td>29.3 15.5 0.75 17.4</td>
<td></td>
</tr>
<tr>
<td>Average Nucella</td>
<td>29.4 41.7 49.1 40.1 20.4 22.2 10.9 17.8</td>
<td>21.0 20.4 34.2 25.2</td>
<td>29.3 15.6 0.59 16.9</td>
<td></td>
</tr>
</tbody>
</table>

from 0.0% to 1.8% within currently recognized species and from 0.83% to 12.3% between species of Nucella (Table 4). Nucella species differed by 21.3% on average from the outgroup species.

The single most parsimonious tree obtained from the analysis of the POLY data set, using an 8:1 weighting of transitions relative to transversions, is shown in Figure 2. The same data set weighted so that transversions alone were informative also yielded a single most parsimonious tree with an identical topology. Equal weighting of transitions and transversions resulted in two equally parsimonious trees, one of which was topologically identical to Figure 2 and one that differed in the shifting of N. lapillus to the N. lamellosa branch.

Decay analyses were performed by constructing strict consensus trees of all trees less than or equal to one step longer than the single most parsimonious tree, then two steps longer, etc., until an iteration was reached in which all cladest except the eastern and western N. lima and the northern and southern N. emarginata had collapsed, at 1154 steps (Fig. 2). This analysis was carried out with 8:1 weighting, so that eight steps can represent eight transitions or one transversion.

The INDIV data set, which included all 17 haplotypes, yielded a most parsimonious tree topology identical to Figure 2 under all three weighting schemes. The increased resolution of the INDIV data set relative to the POLY data set with equal weighting is an example of the advantage of breaking polymorphic terminal taxa into monomorphic component parts (Nixon and Davis 1991; Maddison and Maddison 1992). Each of the currently recognized species was represented by three haplotypes, except N. canaliculata and N. emarginata-S, which had two haplotypes, and N. emarginata-N, which had one haplotype. Analyses of all 324 (3^x 2^x 1) possible single-haplotype-per-species combinations for the seven currently recognized species yielded a tree topology identical to Figure 2. The trees varied in length from 1093 to 1117 steps.

The Fitch-Margoliash trees generated from the maximum-likelihood distance matrices were identical in topology to the maximum parsimony tree shown in Figure 2 for all six possible combinations of transition-transversion ratios and categories of substitution rates described above.

Rates of mtDNA Evolution

Because nucleotides at first and second codon positions varied little among the Nucella sequences, we focused on transitions and transversions at third positions of codons for rate calibrations. Transversions have been found to accumulate most uniformly in the cytochrome b gene of mammals (Irwin et al. 1991). The evolutionary dynamics of first and second codon positions will be dealt with in the broader context of cytochrome b evolution within the caenogastropods (Collins et al. 1992 unpubl.). Absolute time calibration points are based on the fossil occurrences in the Appendix.

Third position transitional differences accrete quite rapidly and begin to approach saturation in the range of five to 10 million yr (Fig. 3a). These differences accumulated at roughly 3–4% per million yr over the zero- to five-million-yr timescale. The average number of transitional differences peaked sometime after 20 million yr and then declined in the most

Table 4. Pairwise number of sequence differences for selected individuals, all sites, all changes (below diagonal), and mean distances (above diagonal, adjusted for missing data).

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td>0.0</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>0.1</td>
<td></td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>0.1</td>
<td>0.1</td>
<td></td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td></td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>0.1</td>
<td>0.1</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td></td>
</tr>
</tbody>
</table>
ancient comparisons. Mammals exhibit a similar pattern, which appears to result from the accumulation of transversions, which obscures the record of transitions (Brown et al. 1982; Aquadro et al. 1984; Irwin et al. 1991). Third position transversions accumulated more slowly and more uniformly than transitions (Fig. 3b), with an average rate of about one transversion per million yr over the time scale of these comparisons for the 239 third positions of cytochrome b sequenced (0.42% per million yr). A similar value, 0.5% per million yr, was reported for cytochrome b over a comparable time scale for placental mammals (Irwin et al. 1991).

The overall pattern of sequence divergence (Fig. 3c) is typical of animal mitochondrial DNAs, with a steep early rate of divergence that gradually approaches an asymptote with increasing age (Brown et al. 1979). A similar pattern is obtained whether pairwise distances or patristic distances are used (Fig. 3c). The difference between the patristic distance and the pairwise distance becomes progressively greater as the minimum time of divergence for each node on the most parsimonious tree is estimated as the first occurrence of a fossil representing either of the lineages that descend from that node. The minimum divergence time of the Thaidine lineage leading to *Plicopurpura* and the Ocenebrine lineage leading to *Nucella* (55 M.Y.B.P., basal Bocene) is from Kool (1989). The divergences between the eastern and western forms of *N. lima* and northern and southern forms of *N. emarginata* are assumed to have taken place sometime during severe Pleistocene glaciations. The midpoint of the Pleistocene (8 × 10^3 yr bp) is used as the calibration point for these divergences.
the age of the divergence (and the number of intervening nodes on the cladogram) increases. This is to be expected, as patristic distances for unordered multistate characters cannot be less than, but may be more than, pairwise distances.

In addition, the number of opportunities for discovering homoplasy (resulting in greater branch lengths) typically increases with an increase in the number of intervening nodes (Sanderson 1990).

**Fossil Record**

The known fossil record of *Nucella* is generally consistent with the branching order of the phylogeny depicted. Basal species appear earlier in the fossil record than more recently derived species (compare Fig. 2 with Fig. 4; see the Appendix for details of the fossil record). In addition, using the methods outlined by Norell and Novacek (1992), age rank (order of appearance of species in the fossil record) was highly correlated with clade rank (branching sequence from the base of the cladogram, for the pectinate components of the tree), indicating that the fossil record of *Nucella* is consistent with the molecular phylogeny derived here (Fig. 5).

**Ecological, Reproductive, and Morphological Attributes of Living *Nucella***

Ecological characteristics also varied among species of *Nucella* in a pattern largely consistent with the molecular phylogeny (Table 2). The four more basal species (*N. lima, N. freycineti, N. lapillus, and N. lamellissa*) all occur commonly in subtidal habitats as well as in quiet water environments, whereas the three more derived species (*N. canaliculata, N. emarginata-N, and N. emarginata-S*) occur there only rarely. The two most derived species (*N. emarginata-N, N. emarginata-S*) are also species whose vertical range extends high on the shore, although the sole North Atlantic species (*N. lapillus*) and the geographically widespread *N. lima* extend nearly as high. Only the three most basal species (*N.
lima, N. freycineti, and N. lapillus) and one of the most derived species (N. emarginata-S) include gastropods in their diet more than 5% of the time. Finally, all Nucella species produce egg capsules from which juvenile snails emerge (Strathmann 1987), and all except N. lamellosa and members of the N. freycineti clade provision these egg capsules with nurse eggs (unfertilized eggs consumed by developing embryos).

Shell traits varied substantially among species of Nucella (Table 2). The two most derived species (N. emarginata-N, N. emarginata-S) exhibited the smallest average and maximum adult sizes, whereas N. lamellosa exhibited the largest average and maximum adult sizes. Well-developed varices (bladelike axial sculpture) occur in only three species (N. freycineti, N. lapillus [rarely], and N. lamellosa), and in all three their expression varies dramatically among habitats. The four most basal species exhibit seven to 10 major spiral ribs on the body whorl, but in the more derived species these have either increased (N. canaliculata) or decreased (N. emarginata-N, N. emarginata-S) in number. Complete apertural teeth, which are expressed most consistently in adults (Crothers 1985), occur only in N. lapillus and N. lamellosa. Unfortunately, information on chromosome numbers was only available for four species (Table 2). Three have N = 35 chromosomes. Nucella lapillus not only has a smaller number of chromosome, but is also polymorphic in parts of its range (N = 13 to 18).

**DISCUSSION**

**Molecular Evolution and Phylogeny of Nucella**

**Weighting.** In mitochondrial DNA sequence comparisons an excess of transitions relative to transversions is commonly observed, both among genes within the same taxa (Brown et al. 1982; Hixson and Brown 1986), and among taxonomic groups (Brown 1985; Moritz et al. 1987; Irwin et al. 1991; Simon 1991). A similar transition excess was observed in the cytochrome b sequence of Nucella. Overall, transitions outnumbered transversions by approximately 4:1 to 9:1, depending on whether sites having both transitions and transversions were included. As expected, this bias was even more pronounced among closely related terminal taxa, with values of 12:1 for the E and W forms of N. lima to 7:0 for the N and S forms of N. emarginata and 8:0 between the haplotypes of N. freycineti.

For the phylogenetic analyses reported here, we chose an intermediate value of 8:1 as a conservative estimate, although the higher values of comparisons from the most closely related taxa are probably closer to the actual values (Brown et al. 1982). For Nucella, the same tree (Fig. 2) was obtained over the range of possible transition-transversion ratios, although the choice of weights often affects tree topology (e.g., Dixon and Hillis 1993).

**Decay Analyses.** Decay analysis estimates the level of support for a given clade by determining the maximum number of steps beyond the single most parsimonious tree at or below which that clade is found in all trees (Fig. 2). For our data, the strongest support is for the basal position of the N. lima clade within the ingroup, and for the N. canaliculata-N. emarginata northern and southern clade. The weakest support is for the position of N. lamellosa and the sister status of N. freycineti and N. lapillus. We are currently collecting more sequence data from these species as well as from several additional outgroup species to test further the phylogeny presented here. The single most parsimonious tree in Figure 2 is the current best estimate of relationships within Nucella.

**Intraspecific Molecular Variation in Nucella.**—Although intraspecific polymorphism (both ancestral and in extant populations) among recently diverged taxa may confound phylogenetic studies based on single specimens per taxon (Neigel and Avise 1986; Smouse et al. 1991), the intraspecific variation uncovered in Nucella had no effect on our tree topology. The analysis of all combinations of single haplotypes per species was potentially biased in favor of finding an effect, since we treated the eastern and western forms of N. lima as one species, but in each case we obtained the same tree topology, regardless of the haplotypes chosen to represent a species. At the sequence level we can compare the kinds and amounts of differences within and between species. Where intraspecific polymorphism is restricted to one kind of change, such as transitions, it should not affect tree topologies supported by, and weighted with respect to, other types of changes, such as transversions.

The cryptic species status of the northern and southern forms of N. emarginata is supported by reproductive, morphological, and genetic (allozyme) evidence (Palmer et al. 1990). The sequence divergence between the N. emarginata-N and N. emarginata-S individuals sampled in this study (0.83%) was the smallest of the between-species comparisons and was unique in being composed entirely of transitional differences. This level and type of sequence divergence was found within single localities of some of the other species of Nucella (e.g., N. lamellosa; Table 4). Between-species divergence values approximately equal to within-locality levels of intraspecific polymorphism suggest that this speciation event occurred very recently relative to rates of mitochondrial DNA evolution.

The sequence divergence between the eastern and western forms of N. lima (1.8%) was more than twice that observed between the northern and southern forms of N. emarginata. This hints at the possibility of cryptic species within N. lima. An alternative is that these forms represent a deep phylogeographic break within a single species (Avise et al. 1987). More extensive sampling is in progress to verify the geographic pattern of sequence variation and to look for possible correlated morphological variation.

Finally, the distribution of the three haplotypes of N. freycineti suggests some geographic patterning. All four individuals from Usujiri are a single haplotype that is approximately 1% divergent from four of the five individuals sampled from Tsukusuki. The last individual from Tsukusuki has a unique haplotype, intermediate in distance from the other two haplotypes; it is 0.55% divergent from the Usujiri haplotype and 0.7% divergent from the common Tsukusuki haplotype. More sampling is needed, but the presence of an intermediate haplotype indicates that we are probably sampling continuous variation along the range of a species with limited gene flow, the type III pattern of Avise et al. (1987).

**Calibrating Rates of Molecular Evolution in Marine Gastropods.**—The sister species used as outgroups in this study,
*P. patula* and *P. columellaris*, occur in the Caribbean and eastern Pacific respectively, and are considered to be geminate, a species pair isolated by the Pliocene emergence of the central American Isthmus about 3.5 million yr before present (M.Y.B.P.) (Coates et al. 1992). This independent calibration yields rates of 7.2% per million yr for third position transitions and 2.4% per million yr for third position transversions between the *Nucella purpura* geminates, values considerably higher than those found among *Nucella* species when calibrated against their fossil record.

Several explanations for this apparent discrepancy are possible. Perhaps the simplest is that the tropical geminate snails are evolving more rapidly than *Nucella*. If we assume, for the moment, that rates are similar between *Nucella* and the tropical geminates, the implication is that either the fossil record of *Nucella* has been misinterpreted, or the geminates were isolated prior to 3.5 M.Y.B.P. The concordance of rate estimates for third position transversions between *Nucella* and the placental mammals (Irwin et al. 1991) over a similar time scale, suggests that the former explanation is unlikely. Moreover, recent studies of Caribbean Neogene molluscan and foraminiferal assemblages indicate that environmental changes and divergence of populations may have significantly preceded the final emergence of the Central American Isthmus (Jackson et al. 1993; Collins 1993) and molecular work on alpheid shrimp geminates also hints that some geminates may have been isolated before 3.5 M.Y.B.P. (Knowlton et al. 1993).

If we apply the rate calibration based on the fossil record of *Nucella* to the average slow geminate divergence, a Middle Miocene isolation is indicated. Clearly, more data are needed to distinguish among these possibilities.

**Biogeographic History and Speciation of Nucella**

The combination of molecular and paleontological data, as well as the summary of morphological, life history, and ecological data, allow us to infer the evolutionary history of *Nucella*. The genus apparently originated during the Early Miocene in the warm-temperate Pacific (see also Amano et al. 1993). By the early Late Miocene, three species (*N. lima*, *N. canaliculata*, and *N. lamellosa*) living today in cold waters had arisen, so the evolution of cold-water species can be traced back at least to this interval. Expansion of the genus into the Atlantic basin began during the Pliocene. In the following discussion we treat three aspects of the evolution of *Nucella* in detail: the directionality and timing of the trans-Atlantic interchange, parallels between the inferred biogeographic history of *Nucella* and other North Pacific genera, and evidence that climatic cycling has been important in the diversification of *Nucella*.

**Direction and Timing of the Trans-Atlantic Interchange.**—The phylogeny of *Nucella*, in which the North Atlantic species *N. lapillus* is nested within the clade of North Pacific *Nucella* species, confirms the inference from the fossil record that the direction of the interoceanic movement in this genus was from the Pacific to the Atlantic. A possible path for this migration is also indicated. Trans-Atlantic migrations via the Canadian Archipelago are usually assumed to have been the probable route of dispersal from the Pacific to the Atlantic during the Pliocene (e.g., Hopkins 1967; Reid 1990b). Nevertheless, the western Pacific sister taxa to the North Atlantic *Nucella*, the occurrence of *Nucella* on the Asian Arctic island of Novaya Zemlya (Cook 1915), and the older fossil record of *Nucella* in the eastern Atlantic relative to the western Atlantic all suggest that a trans-Eurasian route is at least as likely an alternative for *Nucella*, and perhaps for other species, as a trans-Canadian dispersal.

Paleontological evidence places the initiation of the trans-Arctic interchange in the Early Pliocene (5.2–3.4 M.Y.B.P., Hopkins 1967; Vermeij 1989b, 1991). This is based primarily on the sudden appearance of large numbers of mollusks and other marine benthos of North Pacific affinities in Pliocene fossil beds of the North Atlantic region. If our calibration presented in Figure 35 is accepted as a rough approximation, the invasion of the Atlantic by *Nucella* at 7–8 M.Y.B.P. occurred a little earlier than most estimates of the initiation of the trans-Atlantic interchange. Similar dates were obtained when the *N. freycinet-N. lapillus* divergence time was removed from the calibration. These inferred divergence times should be regarded as provisional, as they are subject to a number of possible sources of error including fossil misidentification, erroneous estimates of divergence time from fossils, uneven temporal sampling (gap between 20 and 55 million yr), and the possibility of underestimation of molecular distances because of multiple substitutions at a site. A conservative conclusion is that the molecular evidence is consistent with an early participation of *Nucella* in the trans-Atlantic interchange during Late Miocene or Early Pliocene time. The occurrence of *Nucella* in Late Pliocene deposits of Europe is also consistent with an early participation in the trans-Atlantic interchange.

Several other molecular studies also suggest the trans-Arctic interchange may have begun earlier than previously believed. First, allozyme comparisons of two separate invasions of the North Atlantic by the rocky intertidal snail *Littorina* (Zaslavskaya et al. 1992) indicated an entry between 3.45–4.05 M.Y.B.P. Second, three different studies of ray-finned fishes thought to have participated in the trans-Arctic interchange gave estimates of 3.1, 5.1, and 7.9 M.Y.B.P. (Grant et al. 1984; Grant 1986; Grant and Stahl 1988). Unfortunately, calibrations of *D* with time vary by almost a factor of four between these studies (a D-value of one accumulates in 5 × 10⁶ yr for snails, but requires 1.9 × 10⁶ years for ray-finned fishes) tempering somewhat our confidence in the similarity of these values. Other studies indicating significant divergence between Atlantic and Pacific populations include a DNA-DNA hybridization comparison between samples of the seaweed *Cladophora alpina*, in which an estimated sequence divergence equal to that commonly seen between genera of angiosperms was found (Bot et al. 1989), and an allozyme comparison of Atlantic and Pacific *Mystus trossulus* in which allele frequency differences at most polymorphic loci were reported (McDonald and Koehn 1988).

In contrast, other comparisons between Atlantic and Pacific populations of taxa indicate low levels of genetic differentiation, implying more recent interchange. Studies of variation in mtDNA restriction endonuclease sites and fragments for *Strongyluraeus droebachiensis* (Yater and Brown 1986; Palumbi and Wilson 1990) and in mtDNA sequence
comparisons for *Strongylocentrotus pallidus* (Palumbi and Kessing 1991) revealed remarkable homogeneity of Atlantic and Pacific populations of these species. Allozyme frequencies of Atlantic and Pacific populations is the Greenland Halibut differed only slightly (*D* = 0.013, Fairbairn 1981; Grant 1987). Many other examples of low Atlantic-Pacific molecular divergence estimates (Graves et al. 1984; Wilson and Waples 1984; Smith 1986; Bonhomme et al. 1987; Bowen et al. 1989; Graves and Dizon 1989) are possibly the result of continuing gene flow within these species through southern oceans (Palumbi and Kessing 1991) or recent introduction resulting from human commerce (McDonald and Koehn 1988; Meehan et al. 1989).

When viewed collectively, these studies of Atlantic-Pacific differences suggest a bimodal distribution of divergence times. Species have either been isolated since early in the trans-Arctic interchange, or appear to have undergone genetic exchange very recently. The reasons for this apparent bimodality are unclear, but may relate to the sensitivity of these species to extinction and reinvansion during periods of climatic fluctuation (Palumbi and Kessing 1991) or to their relative dispersal ability through cold Northern seas.

**Biogeographic Parallels between *Nucella* and Other Gastropods.—**The biogeographic history of *Nucella* exhibits at least two striking parallels with that of another genus of rocky-shore gastropods, *Littorina*. First, the linking of the North Atlantic *N. lapillus* to the western Pacific *N. freycinetii* parallels the case in *Littorina*, where at least two independent invasions have occurred from primarily western Pacific lineages to the Atlantic (Reid 1990a,b). This congruence is all the more striking because the mode of larval dispersal in *Nucella* and the littorinid subgenus *Neritrena* (by benthic larvae or adults) is different from that of the paralyphetic subgenus *Littorina* (by planktonic larvae). Second, species with more restricted vertical ranges on the shore appear to be of more recent origin in both *Nucella* (*canaliculata*, *emarginata*-N and *emarginata*-S) and in Pacific and Atlantic representatives of the littorinid subgenus *Neritrena* (Reid 1990a; Vermeij 1992b). Hence in both taxa we see a trend from ecological generalist to ecological specialist.

From another perspective, however, the radiation of *Nucella* appears to contrast with the pattern found in *Littorina* and in the buccinid genus *Neptunea*. In both of these latter genera, the oldest and least derived species are thought to have occurred in warm-temperate conditions, whereas more derived and geologically younger species are distributed in cooler shores in the cold-temperate and polar zones (Golikov and Tsvetkova 1972; Reid 1990b). This pattern does not appear to hold for *Nucella*. Although the oldest species of *Nucella* (*N. tokudai*) was apparently a warm-temperate species, other geologically old species (*N. lima, N. freycinetii satoe*) are generally cold-temperate species. Furthermore, the warm-temperate *N. emarginata*-S appears to be the most derived and geologically youngest Pacific species of *Nucella*.

**Climatic Vicariance.——**The diversification of *Nucella* in the North Pacific coincides with the beginning of pronounced climatic fluctuations in the Miocene. Cenozoic climatic changes have often been invoked to explain patterns of distribution and speciation in molluscs of the North Pacific (Golikov and Tsvetkova 1972; Valentine and Jablonski 1983; Valentine 1984; Vermeij 1989b; Reid 1990a,b). The general model presented is one in which fluctuating climatic conditions induce alternating contraction and expansion of species ranges on east-west or north-south trending coastlines, occasionally producing isolated subpopulations that found new species. During cooler intervals, for example, northern populations may become sequestered in pockets of warmer water as surrounding populations of a southern species become locally extinct. Aspects of intraspecific variation, phylogeny, species distribution, and the fossil record of *Nucella* are compatible with a model of speciation by climatic vicariance.

The boundaries of mollusc biogeographic provinces along the Pacific coast of North America have shifted in response to changing climate (Valentine 1961; Addicot 1966). Ranges of *Nucella* species were affected by these climatic shifts (e.g., Addicot 1966, 1969), indicating that opportunities existed for isolation of populations of *Nucella* species during climatic fluctuations. The northern and southern forms of *N. emarginata* and eastern and western forms of *N. lima* may be the most recent products of a process that has been occurring for much of the Neogene in North Pacific *Nucella*. The low levels of mtDNA sequence divergence between the northern and southern forms of *N. emarginata* imply that this event probably occurred during severe late Pleistocene climatic cycles.

**Patterns of mtDNA sequence variation in *N. lima* discussed above suggest a possible case of east-west isolation by extinction of more northerly intermediate populations. *Nucella lima* currently ranges continuously from northern Japan through the Kurile, Commander, and Aleutian Islands to the North America continent. During cooler Pleistocene intervals, however, glaciers covered the Alaskan peninsula and large portions of the Aleutian Islands (Hopkins 1967). These Pleistocene glaciers would have effectively bifurcated the range of *N. lima* into more southerly populations in the eastern and western Pacific. These populations would have come into secondary contact as they reclaimed the lost intervening habitat with the retreat of the glaciers during interglacial periods. Whether the current populations of eastern and western *N. lima* are coalescing or have produced new species has not been determined. In general, the results of these cyclical isolation events will depend on a variety of factors, such as the length of time of isolation and population genetic histories during the glacial phase.

**Character Evolution within *Nucella*.

Phylogenetic analyses among taxa above the level of species lose important information about rates of speciation, rates or directions in which particular traits may evolve, correlations between phenotypes and environments that may suggest adaptation, and the phylogenetic origins of new characters (Brooks and McLennan 1991). Thus phylogenetic analyses of intrageneric and intraspecific variation can yield significant insights into rates, correlates, and potential causes of speciation (Avise 1994). The genus *Nucella* offers an unusual opportunity for such an analysis, because so much is already known about nearly all the extant species. Unfortunately, this analysis is also constrained by the limited number of taxa. Although extensive, prior studies of *Nucella* have
generally proceeded within an ahistorical framework because phylogenetic analyses have been thwarted by highly variable morphology (Kincaid 1964; Palmer et al. 1990). The molecular phylogeny offered here allows us to examine evolutionary trends within this well-studied, cosmopolitan genus.

Ecological and Life-History Traits.—An outstanding question in studies of the speciation process is whether founders of new clades are more or less ecologically specialized than the descendants to which they give rise (Mayr 1963). In other words, are ecological generalists more likely to give rise to specialized species that exploit a subset of habitats more effectively, or are ecological specialists more likely to evolve novel attributes that promote subsequent adaptive radiation, including the origin of more generalized species? Similarly, in light of the frequent evolutionary trend towards increased body size observed in fossil invertebrates (Newell 1949), is such a trend apparent within Nucella?

Phylogenetic variation in three ecological traits suggest an evolutionary trend towards increased ecological specialization in Nucella (Fig. 6). First, gastropods have effectively disappeared from the diet in the more derived, northeastern Pacific species (N. lamellosa, N. canaliculata, and N. emarginata-N; trait 1, Fig. 6). Their reappearance in the diet of N. emarginata-S is clearly a reversion toward the primitive state. Second, nonmytilid bivalves have also disappeared from the diet in this clade (trait 2, Fig. 6). Why gastropods and nonmytilid bivalves are effectively absent from the diet of these three species remains a mystery (Palmer 1988). Third, the most derived, northeastern Pacific species (N. canaliculata, N. emarginata-N, and N. emarginata-S) are found in a narrower range of habitats. Unlike their inferred ancestors, they are rare or absent in subtidal or very quiet water environments (trait 3, Fig. 6 and Table 2, respectively). Thus diet and habitat breadth became progressively more circumscribed in the northeastern Pacific species. The expansion of the diet in one of the most recently derived eastern Pacific species (N. emarginata-S), though, reveals that such changes are not unidirectional.

The two life-history traits examined exhibited a correlated pattern. First, average adult size and range of adult size (as measured by shell length) increased independently in two lineages (N. freycineti and N. lamellosa) and decreased in a third (the N. emarginata clade) (Table 2, trait 5, Fig. 6); thus, it appears to be rather labile evolutionarily and no long-term trends are evident. Second, the loss of nurse eggs in N. lamellosa and within the freycineti clade are clearly derived states (trait 4, Fig. 6). Nurse eggs are widespread among muricacean gastropods, where they are more commonly associated with direct development than with planktonic larvae (Spight 1976). They also have an ecologically significant impact upon size at hatching: the more nurse eggs, the larger the hatching size (Rivest 1981). Significantly, the independent loss of nurse eggs occurred in the two clades that also exhibited independent increases in body size (N. freycineti and N. lamellosa), suggesting that increases in adult size are correlated with a novel reproductive pattern that decreases hatching size (P = 0.02, binomial test).

Morphological and Chromosomal Traits.—Vermeij (1987) argues that much morphological evolution in marine invertebrates can be viewed as the result of an "arms race" between predators and prey. Since we know a modest amount about the functional significance of shell traits in gastropods in general (Vermeij 1993b), and about the benefits and costs of shell traits in Nucella in particular (Palmer 1981, 1985, 1992; Crothers 1985), the Nucella phylogeny allows us to ask how evolutionarily labile morphological traits are. For example, are traits that reduce vulnerability to shell-breaking predators ever lost evolutionarily? In addition, for shell traits we can also confirm inferences of ancestral state and patterns of evolution against the fossil record.

Both varices and apertural teeth reduce the vulnerability of shells to peeling predators (Vermeij 1978). However, based on their distribution in extant species of Nucella, both appear quite labile evolutionarily. The common development of conspicuous varices by N. freycineti and N. lamellosa represent independently derived states (trait 6, Fig. 6), even though varices are widespread among Northern Hemisphere muricacean gastropods (Abbott 1974). Similarly, well-developed apertural teeth arose independently in both N. lapillus and N. lamellosa (trait 7, Fig. 6). Neither of these traits, however, exhibits a consistent trend to be more well developed in more recently derived species. This result is not surprising. Shell-damaging predators as well as the incidence of repair of shell injuries are greatly reduced in cool-temperate as compared to warm-temperate areas, as evidence for both Nucella and Littorina strongly indicates (Vermeij 1992a). The loss of shell defenses in cool-temperate descendant species of Nucella relative to warm-temperate ancestors is consistent with this difference. Moreover, the phenotypic induction of shell defenses in Nucella in the presence of predatory crabs (Appleton and Palmer 1988; Palmer 1990) strongly underscores the evolutionary lability of predation-related shell traits in this genus.

For the most part, the fossil record confirms the inferred ancestral form and lability of Nucella shell traits. The oldest known species, the Early Miocene N. tokudai (as N. packi in Addicott 1970), was of intermediate size (25 mm shell length), and lacked well-developed varices and well-developed apertural teeth. Although prominent apertural teeth persist in only two modern species, N. lapillus and N. lamellosa (Table 2, Fig. 6), they were well developed in the latest Miocene N. collomi, and the Late Pliocene N. trancosana (Addicott 1969), which may be a close relative of N. emarginata. Thus, in spite of their defensive roles, varices and apertural teeth have been lost and regained one or more times over the evolutionary history of Nucella, hence an evolutionary trend toward species with more well-defended shells since the Miocene does not seem evident.

The role that chromosome variation plays in speciation is a source of ongoing controversy (Sites 1995). The reduced number of chromosomes in N. lapillus is derived from an ancestral state of N ≥ 30, when the Nucella data are combined with those of other muricids as well as most neogastropods (numbers vary from N = 30–35 [Nishikawa 1962; Ahmed and Sparks 1970], character tracing not shown). Reductions in chromosome number seem to occur more commonly than increases in other taxa as well (White 1978). In addition to having fewer chromosomes, N. lapillus exhibits a latitudinal cline in chromosome number, with N = 13 in northern populations and N = 18 in southern populations. In regions of
Fig. 6. Phylogenetic variation in ecological and shell traits in Nucella mapped onto the molecular phylogeny of Figure 2. All figures from Dall (1915) except *N. lapillus* (from Berry and Crothers 1974) and *N. freycineti* (from Habe and Hoo 1971). Character tracing was conducted with MacClade (vers. 3.05, Sinauer) using the criterion of parsimony to infer ancestral states (i.e., inferred ancestral states were those that yielded the fewest state changes for the tree topology of Fig. 2). All possible optimizations were examined using equivocal cycling. Where inferred ancestral states were equivocal, the unsolvable alternative states are indicated in parentheses. All state changes were considered to be equally probable, except those noted by *, where state changes were treated as ordered in the order illustrated in the legend.
overlap, along the south coast of England and the north coast of France (Staiger 1954; Bantock and Cockayne 1975), clear evidence exists for a Robertsonian polymorphism, a rather common form of polymorphism in hybridizing species (White 1978). Curiously, this latitudinalcline is the reverse of that expected if southern populations were derived from northern ones, which in turn were derived from the ancestral state. Hence either the southern populations experienced a reversion towards the neogastropod norm, or the northern populations are the more derived, perhaps because they arose from a southern stock having $N = 18$ chromosomes during the range expansion following Pleistocene glaciations.

CONCLUSIONS

In this study we have attempted to integrate molecular, morphological, ecological, and paleontological data to understand the evolutionary history of speciation within Nucella. These data have given us a more complete and accurate view of the history of this genus than would have been possible if only one of these approaches had been followed. The congruence of inferences from these approaches is a powerful affirmation of the results of each.

The molecular phylogeny derived here is congruent with the order of appearance of taxa in the fossil record. The phylogeny and geographic distribution of living species confirms the inference from the fossil record of a Pacific to Atlantic directionality and early participation of Nucella in the trans-Atlantic interchange. Phylogenetic, biogeographic, and fossil data indicate that a trans-European dispersal of species into the Atlantic is a viable alternative to the trans-Canadian hypothesis. Patterns of intraspecific molecular variation and fossil data on shifting biogeographic provinces confirm that climatic cycles initiated during the last Cenozoic can form effective barriers along north-south and east-west coastlines, creating conditions for geographic speciation. The patterns found in Nucella are paralleled in other North Pacific taxa.

Character state reconstruction of morphological and ecological characters of living species suggest an evolutionary trend toward increasing ecological specialization in Nucella. Finally, earlier fossil taxa tend to show the characters inferred by parsimony to be plesiomorphic, although both the phylogeny, as well as fossil and living species, indicate that shell characters are quite labile.

ACKNOWLEDGMENTS

We thank S. Goshima, K. Kawai, P. B. Marko, L. West, and N. Wheelwright for sending samples of Nucella for DNA analyses, and N. Gollvik, S. Goshima, and K. Kawai for sending shells of western Pacific Nucella species. C. Cunningham, R. DeSalle, R. Grosberg, R. Harrison, S. P Kool, P. Kraus, P. B. Marko, and D. Reid commented on earlier drafts of the manuscript. This research was funded by National Science Foundation grant BSR-9107306 to WMB and TMC, and NSERC Operating Grant A7245 to ARP.

LITERATURE CITED


EVALUATION OF NUCELLA


Corresponding Editor: R. DeSalle

APENDIX

Geological occurrences, distributions, and characteristics of fossil species of Nucella and dates used for calibration of rates of molecular evolution of Nucella species.

Nucella tokudati

The earliest recognized species of Nucella is N. tokudai (Yokoyama 1932) (see Amano et al. 1993). This species has pronounced uniform spiral cords and lacks axial ribs, which are characteristic states expected to be primitive (see Evolutionary Trends within Nucella). Addicott (1970) synonymized N. carriigenus (Loel and Corey 1932) (= N. tokudai; see Amano et al. 1993) with Thais paccs Clark 1918, which was described from the earliest Miocene of latest Oligocene San Ramon Sandstone of California. Amano et al. (1993), however, have shown that T. paccs possesses the trace of a labral spine on the edge of the shell's outer lip and belongs in the related genus Acmaenellina.

Nucella tokudai (Yokoyama 1932) (= N. paccs Clark 1918 of Addicott 1970) not N. paccs Clark 1918; and = N. carriigenus (Loel and Corey 1932). Middle Miocene Vaqueros Formation and Inkoust sandstone in California; early Middle Miocene Chikubetsu and Middle Okada beds in Hokkaido; Kalet Formation of Kamchatka, Middle Miocene Tenbador Formation, Sobane Sandstone and Flores Lake Sandstone USA; Ishizawa Formation of Hokkaido; Late Miocene Togezhita Formation (upper part) in Hokkaido and Ceroso Sandstone in California (K. Amano et al., unpubl.); earliest occurrence 23-21 M.Y.B.P.

Nucella lima

The oldest specimens attributed to the living, N. lima were described by Etherington (1931). Our examination of Etherington's material (Museum of Paleontology, University of California, Berkeley, number 32090) revealed that these shells resemble N. lima in overall shape, in the absence of axial sculpture, and in the pattern of spiral sculpture of alternating large and small cords. However, the number of cords on the penultimate whorl and last whorl is less on the fossil shells (2 and 16, respectively) than on shells of living N. lima (3 and 18 to 20, respectively). We therefore regard these Miocene specimens as belonging to the N. lima lineage, but a definitive assignment must await the collection of additional material. In any case, the N. lima lineage seems to have been present at an early stage in the history of Nucella.

Nucella lima (Genmal 1971) s.l.: Late early Miocene to early middle Miocene Astoria Formation of southwestern Washington (Etherington 1931, Addicott 1976a,b, Ammen et al. 1984) to Recent; earliest occurrence 23-17 M.Y.B.P. Calibration date 20 M.Y.B.P.

Nucella freycineti

The N. freycineti lineage is recorded from early Middle Miocene onward. The Miocene forms, which have been assigned to N. freycinetti sutori Hatai and Kotaika, 1959 (see Amano et al. 1993), have shells that lack axial ribs and therefore resemble Deshayes's types of the living N. freycineti rather than the axially sculptured forms that have been named N. hysteron and N. elongata. Typical N. freycineti is recorded from Placooe to Recent deposits in Japan (Amano et al. 1993). Dall (1920) named specimens from Placooe deposits near Nome, Alaska, as N. nolimm, but these specimens probably represent N. freycineti (MacNeil et al. 1943) since they exhibit seven widely spaced major spiral cords on the body whorl and lack apertural teeth.

Nucella freycineti sutori Hatai and Kotaika 1959; early middle Miocene Ginzan and Teshikarashumi formations (K-AR dated, 13.7 to 13.8 M.Y.B.P) to late Miocene Togezhita formation (upper part) of Japan and Kamchatka (Amano et al. 1995); Calibration date 14 M.Y.B.P.

Nucella freycineti (Deshayes 1841); Placooe to Recent of Japan and Kamchatka (K. Amano et al. unpubl.); Placooe of Nome, Alaska as N. nolimm (Dall 1920); earliest occurrence 5.2-3.4 M.Y.B.P. Calibration date 4 M.Y.B.P.

Nucella lamellosa

Fossils identified as N. lamellosa by Gladenkova et al. (1984) from the early late Miocene Etioln Suite of Kamchatka may represent either an unnamed ancestral taxon or a lamellosa form of N. f. sutori similar to the living, axially ribbed forms of N. freycineti. In the living form, N. lamellosa is confined to the eastern Pacific (Vermejju et al. 1994).


Nucella canaliculata

Nucella canaliculata (Dall 1832); early late Miocene Grassy and Wisskhan stages of Oregon and Washington (as N. precursor Dall 1909) (Weaver 1942, Addicott 1976b, 1978, 1981, Marinovich 1983) to Recent; earliest occurrence 12-7 M.Y.B.P. Calibration date 16 M.Y.B.P.

Nucella collici

Nucella collici Carson 1926 is a distinctive species from the late Miocene of California characterized by numerous scalary spiral cords,
which may be uniform or alternating in strength, and a long siphonal canal.

*Nucella californi* (Carson 1791): latest Miocene San Joaquin Formation of California (Durham and Addicott 1965; Addicott 1978); 8–5 M.Y.B.P.

**Nucella funkeana**


**Nucella shiwa**

*Nucella shiwa* (Chinzei 1961) (~ 7N. freycineti): Pliocene Tagawa Formation of northern Honshu, Japan (Chinzei, 1961); 5–3.5 M.Y.B.P.

**Nucella transversa**

*Nucella transversa* is a low-spired thick-shelled species with strong apertural teeth that extend into the aperture as ribs, a short siphonal canal, and numerous spiral cords (up to 24) on the body whorl.

*Nucella transversa* (Arnold 1908): Pliocene Meced Formation of California and Cantil Costero Formation of Baja California (Addicott 1969); 5–2.5 M.Y.B.P.

**Nucella emarginata**

The *N. emarginata* clade is the last of the living *Nucella* to appear in the fossil record of the eastern Pacific and is known from Pliocene time onward.


**Nucella lapillus**

*Nucella* is first recorded in the North Atlantic in deposits of Late Pliocene age. The earliest Atlantic form is *N. lapillus incrassata* (Sowerby 1825). Its high-spired shell resembles that of the northeastern Pacific *N. lima*, but it lacks the axial sculpture of that species. Typical *N. lapillus*, whose shell is generally low-spired and more weakly sculptured, appears in the latest Pliocene and in the overlying Early Pleistocene. The western Atlantic population of *N. lapillus*, which is morphologically indistinguishable from that in Europe, is recorded from the Late Pliocene onward (Richards 1962). *Nucella tetragona* (Sowerby 1825) was regarded as the earliest Atlantic species of the genus, but this species belongs in another genus *Spinucella* that neither descended from nor gave rise to *Nucella* (Vermeij 1995a).

*Nucella lapillus incrassata* (Sowerby 1825): Late Pliocene Red Crag of England and Merxemian stage of Belgium (Gilbert 1959, 1965; Gir- bault et al. 1991); 3.4–1.8 M.Y.B.P.

*Nucella lapillus* (Lamarck 1798): Early Pleistocene Norwich Crag of England to Recent in Europe, Mediterranean, and Morocco; Late Pleistocene (40 M.Y.B.P.) to Recent in eastern North America (Gil- bault 1959, 1965), (Richards 1962; Cambridge and Kitching 1982).