Reproductive biology of free-living and commensal polynoid polychaetes at the Lucky Strike hydrothermal vent field (Mid-Atlantic Ridge)

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ABSTRACT. We examined the reproductive biology of a polynoid polychaete commensal in mussel mantle cavities (Branchipolynoe cf. seepensis) and of a polynoid polychaete that is free-living among those same mussels (Opisthotrochopodus n. sp.). Specimens of each species were collected from 2 different sites (Eiffel Tower and Sintra; -400 m apart) within the Lucky Strike hydrothermal field. Both species exhibit sexual dimorphism, with varying numbers of pairs of nephridial papillae. In B. cf. seepensis, sexual dimorphism also includes a difference in size, with females larger than males. Contrary to our expectations, the reproductive biology of both species is very similar. Sperm heads are elongate, suggesting some mode of sperm transfer or storage. Females of both species contain mature sperm and there is evidence of internal fertilization. Maximum oocyte diameters are large (>390 μm), from which we infer non-planktotrophic development; oogenesis is intraovarian and suggests rapid oocyte development. Oocyte size-frequency distributions and population size structure suggest a pattern of asynchronous gametogenesis. We found no substantive evidence for site-specific variations in reproductive biology in either species.

KEY WORDS: Hydrothermal vent · Polychaete reproduction · Polynoid

INTRODUCTION

Soon after the discovery of hydrothermal vent communities in the deep sea, biologists recognized that the ephemeral and insular characteristics of this environment do not dictate a single reproductive strategy among invertebrate types (e.g. Lutz et al. 1984, Van Dover et al. 1985), despite early expectations of r-type ecological strategies including small and abundant progeny with significant dispersal capabilities (e.g. Desbruyères & Laubier 1983). For the limited number of vent taxa investigated to date, most species are inferred to have lecithotrophic development, based primarily on egg size (reviewed in Jollivet 1996). Apparent exceptions include mytilid mussels (Bathymodiolus spp.; Berg 1985, Le Pennec & Benninger 1997) and some bythograeid crabs and shrimp (e.g. Bythograea thermydron; Van Dover et al. 1985) which produce large numbers of small eggs that presumably develop into planktotrophic larvae. While lecithotrophy is often associated with limited dispersal in shallow-water species, the uniformly cold environment of the deep sea may retard larval development, making it difficult to be certain that lecithotrophic development restricts dispersal (Lutz et al. 1980, 1984). A few vent species undergo direct development, forgoing a dispersive larval stage altogether (e.g. amphipod species; France et al. 1992).

As in other ecosystems, phylogenetic constraints within some taxonomic groups at vents (e.g. galatheid squat lobsters [Van Dover & Williams 1991] and archaeogastropod mollusks [Lutz et al. 1984]) seem to prescribe certain inflexible reproductive characteristics among species, especially with regard to planktotrophic development. Other reproductive characteristics, such as temporal patterns of gametogenesis, or even direct versus lecithotrophic development, may be more flexible.

Even within higher taxa (families and subfamilies) that are exclusive to hydrothermal vent ecosystems,
there is no consistent reproductive pattern that might
indict a particular environmental aspect of hydrother-
mal vents as an overwhelming selective pressure. The
best example of this may be within the polychaete fam-
ily Alvinellidae, where reproductive ecologies may
have radiated within sympatric and allopatric species
of the genus Paralvinella to include pseudocopulation
and broadcast fertilization, synchronous and asynchro-
nous gametogenesis, and direct and lecithotrophic
development (McHugh 1989, Zal et al. 1995). We use
the terms 'synchronous' and 'asynchronous' gameto-
genesis to describe the condition of developing
gametes within individuals and among individuals
within a population without making inferences regard-
ing spawning patterns (i.e. sensu Eckelbarger & Watling 1995). Polychaetes in general seem more flexi-
ble in adapting variations on reproductive themes than some other invertebrate groups (Wilson 1991, Eckelbarger & Watling 1995). Apart from siphonos-
tome copepods in the family Dinuvulidae, the poly-
chaete family Polynoidae demonstrates the greatest
radiation at the species level within the vent ecosys-

tem, with 33 species described to date (Tunnicliffe et al. 1998). This radiation seems likely to accommodate a
variety of reproductive ecologies; indeed, reproductive
flexibility may in part account for the radiation.

To initiate a long-term study of the comparative
reproductive biology of the Polynoidae, we compare
the reproductive characteristics of 2 polynoid poly-
chaete species expected to demonstrate adaptations
associated with their particular life habits rather than
with the more general condition of habitat ephemeral-
ity or insularity. One of these species, Opisthotro-
chopodus n. sp. (Pettibone pers. comm.), is a small
(max. length ~10 mm), free-living polychaete that for-
ages among mytilid mussels (Bathymodiolus sp.) that
dominate the Lucky Strike hydrothermal field on the
Mid-Atlantic Ridge. The other species, Branchipoly-
noe cf. seepensis, reaches much larger sizes (max. length
~30 mm) and lives as a commensal within the mantle
cavity of the Lucky Strike mussels. Opisthotrachopod-
us n. sp. and B. cf. seepensis belong to 2 different
subfamilies of the Polynoidae (BranchinotONGLuminae
and Branchiopolyloinae, respectively). These subfami-
lies are so far restricted to deep-sea reducing environ-
ments, with so little known about their reproductive
biology that it is impossible to make any inferences
about phylogenetic constraints at the subfamilial level.
Other subfamilies in the Polynoidae include represen-
tatives that have free-spawning, planktotrophic larvae
(Bhaud & Cazaux 1987, Resh 1980). At least 1 shallow-
water polynoid species (Harmothoe imbricata) is re-
ported to brood its young on the outside of its body and
then release them to disperse as planktotrophic larvae
(Blake 1975). Our sampling strategy of multiple repli-
cates from 2 discrete sites within the Lucky Strike
hydrothermal field allows us to investigate both site-
specific population structure and spatial variation in

gametogenesis within each polynoid species.

Based on preliminary measures of size differences in
2 morphs of the commensal polynoid species (reported
here), we suspected that this species was likely to be a
protandric hermaphroditic. Small males, we reasoned,
would be adept at moving between mussels while
females would benefit from being egg machines, grow-
ing large and fecund within mantle cavities of
mussels and expending little energy on foraging given
the ready supply of food and experiencing low risk of
predation. This would be a classic example of the 'size-
advantage' hypothesis which suggests that the ability
to change sex is favored when the relation between
age/size and reproductive success differs between the
2 sexes (e.g. Berglund 1990). We also expected a priori
that gametogenesis in the commensal species would
be asynchronous and that females would invest energy
in large, yolky eggs, since they live in a buffered envi-
ronment with a continual supply of food. In contrast,
we expected separate sexes in the free-living polynoid
species and, because of the need by the female to
expand energy in foraging and predator avoidance,
small eggs with relatively little yolk. Synchronous ver-
sus asynchronous gametogenesis was harder to predict
in the free-living species, but, given the steady supply
of primary production in the vent environment, we
anticipated that asynchrony would be favored.

MATERIALS AND METHODS

Study site. The Lucky Strike hydrothermal field is
located at 37°18.5'N, 32°16.5'W on the Mid-Atlantic
Ridge (Fig. 1). The field is comprised of several areas
of active venting and sulfide mounds following a roughly
N-S line across the summit of the Lucky Strike
Seamount (Langmuir et al. 1997). Two discrete sulfide
mounds separated by ~400 m were sampled (Fig. 1):
Eiffel Tower, a 20 m high tapered spire in the southern
area of the field (1587 m) and Sintra, a 5 m spire in the
northern area (1618 m). Both vents are dominated by a
new species of mussel belonging to the genus Bathymo-
diolus (Craddock et al. 1995). Mussel lengths were
not significantly different in our samples from the 2
sites. The overall mean mussel length (n = 1500) was
48 ± 18 mm (SD). The 2 species of polynoid polychaetes
examined in this study are associated with the mussel
clumps.

Sampling. All specimens were collected in July 1996
during several ROV (remotely operated vehicle)
'Jason' dives during the LUSTRE '96 cruise to Lucky
Strike. Mussel clumps were harvested from sulfide
substrates with the manipulator and placed in collection buckets until the mussel volume reached the 2 l mark, at which point the bucket was closed to retain sample integrity. Five replicate samples were collected from Eiffel Tower and 5 from Sintra. Samples were retrieved via free-ascent elevator to minimize the length of time between collection and processing.

Mussels in each sample were rinsed with filtered seawater over a 63 µm sieve to collect all organisms living within interstitial spaces. Maximum lengths of all mussels >5 mm were measured. Careful sorting of mussel washings resulted in relatively large numbers (total n = 265 individuals) of the free-living polynoid polychaete species (*Opisthotrochopodus* n. sp.). Commensal polynoids (*Branchipolynoe* cf. *seepensis*) were collected by opening all mussels and removing the worms. Additional specimens of the commensal species were found in mussel washings and are included in analyses. These are interpreted as mostly being worms that slipped out of their host mussels during collection and processing. A total of 707 *B. cf. seepensis* individuals were collected. All polynoid specimens were preserved in 10% seawater formalin and transferred to 70% ethanol for storage.

Sex determination, sex ratios and population structure. Sexual dimorphism, expressed by the number of pairs of nephridial papillae projecting ventrolaterally from specific segments, was confirmed in a preliminary manner in each species by histological examination of 5 individuals of each morphological type. Numerous (>30) additional putative females per species were examined histologically for oocyte measurements described below and provided further confirmation of this sexual dimorphism. Once sexual dimorphism was confirmed, sex ratios were determined based on examination of nearly all individuals collected within a species (*Branchipolynoe* cf. *seepensis*, total n = 703; *Opisthotrochopodus* n. sp., total n = 264). Site-specific ratios were also examined. Significant differences from 1:1 sex ratios were evaluated using chi-square tests (α = 0.05).

Analysis of population structure in each species was based on total lengths (mm) of all specimens collected. Measurements were made using digital calipers and are accurate to within ±2 mm. Sample sizes for subgroups of worms are indicated in figure captions. For *Branchipolynoe* cf. *seepensis*, size-frequency histograms could be developed with replicates at each site for worms with 1 (male) and 2 (female) pairs of nephridial papillae. Due to the small numbers of

![Fig. 1. Lucky Strike hydrothermal field on the Mid-Atlantic Ridge. Inset: regional context](image)

*Opisthotrochopodus* n. sp. in each replicate, worms from all replicates at both sites were pooled to generate size-frequency histograms for worms with 0 (female) and 1 (male) pair of nephridial papillae. An a priori test for cohort phenomena was conducted by analyzing the size-frequency distributions for normality using the Kolmogorov-Smirnov test (with Lilliefors' correction; α = 0.05). Our assumption is that normal distributions in size frequencies will preclude the existence of significant, discrete cohorts.

Histology. Histological examination of gonadal material was carried out on whole specimens. Prior to histological processing, large specimens were slit ventrally to enhance infiltration. Worms were processed through an alcohol dehydration series and embedded in paraffin. Serial cross-sections 8 µm thick were stained using Papanicolaou OG-6, hematoxylin and eosin.

Oocyte diameters and fecundity. To ensure analysis of oocytes over the range of worm sizes representing the sample populations, subsamples of 15 females per site for each species were selected according to the rel-
Table 1. *Branchipolynoe cf. seepensis*. Number of oocytes measured per individual. Because of the non-random distribution of oocyte diameter along the length of the worm, serial sections were systematically sampled as described in the text. The number of oocytes measured per individual thus depends on the length of the worm. Size-frequencies based on these oocyte measures are plotted in Fig. 9A. Random subsamples of 100 measurements per individual were used for the ANOVA statistical comparison of oocyte diameters between individuals as reported in the text. At \( n < 100 \) (i.e. \( n = 6, l \)), individuals were considered to be immature; these also corresponded to the smallest individuals (see Fig. 10A,B).

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<th>Specimen no.</th>
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atative proportion of worm size classes. Feret diameter (\( \mu m \)) — the diameter of a hypothetical circular object with the same area as the measured object — was used as an estimate of oocyte diameter due to the irregular shape of most oocytes. Feret diameters were measured only of oocytes in which the nucleolus was observed to avoid duplicate measures of the same oocyte and to standardize as much as possible the cross-section of the oocyte. Measurements were made using image processing software (Sigma Scan Pro), with an error of \( \pm 1 \) \( \mu m \). Most worms contained a number of oocytes <25 \( \mu m \) in feret diameter that could not reliably be measured or counted and are therefore not included in this analysis. Within-worm variation in oocyte size was compared to between-worm variation using analysis of variance (ANOVA) components.

Because *Branchipolynoe cf. seepensis* are large and relatively fecund worms (thousands of immature oocytes [<300 \( \mu m \) in diameter] per individual), we chose to subsample the oocyte population. We found that oocyte size distributions are not random along the length of the worms. To reduce bias in our estimate of mean oocyte diameter for each individual, 1 or 2 cross-sections were selected randomly from each slide of 25 to 50 serial cross-sections. All slides with sections containing oocytes were subsampled in this manner (i.e. 6 to 19 slides, depending on the length and reproductive condition of the worm). At least 100 oocytes were measured from each worm, unless the worm was immature (Table 1). For ANOVA comparisons, subsamples of 100 oocyte measures were taken randomly from the measured sets for each individual.

In *Opisthotrochopodus n. sp.*, all oocytes >25 \( \mu m \) were measured (see Table 2 for numbers of oocytes per individual) and a direct tally of oocytes >25 \( \mu m \) was used as an estimate of fecundity. Correlations between oocyte diameters and abundance versus body length were tested using the Spearman Rank Order method (\( \alpha = 0.05 \)). Comparable measures of fecundity in *Branchipolynoe cf. seepensis* were not obtained.

**RESULTS**

**Sexual dimorphism**

*Branchipolynoe cf. seepensis*. Three morphotypes of *B. cf. seepensis* are readily distinguished under a dissecting microscope: worms with 0, 1 or 2 pairs of nephridial papillae (Fig. 2A,B). Histological examination of 5 individuals with 1 pair of papillae on segment 12 and 51 individuals with 2 pairs of papillae (on segments 11 and 12) indicate that individuals with 1 pair are males, and those with 2 pairs are females. (Note that the first segment of polynoids is lateral to the prostomium, fused with it basally, bears dorsal and ventral tentacular cirri, and may be with or without setae. This segment is followed by the second or buccal segment which bears paired elytrophores, elytra, and ventral buccal cirri.) Individuals with 2 pairs of papillae (females) were on average 12.7 mm larger than worms with 1 pair of papillae (males) (Table 3).

Table 2. *Opisthotrochopodus n. sp*. Total number of oocytes per individual. All oocytes were measured in this species

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Sixty-six worms with no papillae were small and sexually undifferentiated (Fig. 3A). In between-site comparisons, individuals with 1 or 2 pairs
of nephridial papillae at Eiffel Tower were significantly larger than like individuals at Sintra (t-test, \( p < 0.001 \)) by a mean difference of 1.7 to 2.5 mm (Table 4).

*Opisthotrochopodus* n. sp. Two morphotypes of *Opisthotrochopodus* n. sp. are readily separated under a dissecting microscope: worms with 1 pair of nephridial papillae on segment 12 and those with no papillae (Fig. 2C,D). Histological examination of 7 individuals with 1 pair of papillae and 45 individuals with no papillae reveals that worms with nephridial papillae are male, those without nephridial papillae are female (Fig. 3B). In addition to the elongate pair of nephridial papillae in male worms, there are 5 posterior pairs of variously elongate ventral processes of unknown function. Female *Opisthotrochopodus* n. sp. from combined Lucky Strike samples were significantly (\( p < 0.001 \); Table 4) smaller (average body length 5.8 mm) than males (average body length 6.6 mm).

### Sex ratios

*Branchipolynoe cf. seepensis*. The estimated sex ratio for the Lucky Strike population based on all samples collected was 0.6:1 (males:females). Chi-squared analysis shows this to be a significant difference (\( p < 0.001 \)) from a 1:1 sex ratio. Analysis of Eiffel Tower and Sintra samples separately resulted in 0.5:1 and 0.7:1 ratios, respectively. These ratios were significantly different (\( p < 0.01 \)) from a 1:1 ratio and from each other.

*Opisthotrochopodus* n. sp. The estimated sex ratio for the Lucky Strike population based on all samples collected was 0.7:1 (males:females). Chi-squared analysis shows
Fig. 3. Size-distribution and sex of polynoids based on number of pairs of nephridial papillae. (A) Branchipolyne cf. seepensis; (B) Opisthotrochopodus n. sp. Centerline indicates the median, box ends indicate 25th/75th percentiles, whisker lines indicate 10th and 90th percentiles, and dots indicate 5th and 95th percentiles. Based on 707 individuals of B. cf. seepensis and 264 individuals of Opisthotrochopodus n. sp.

no significant difference from a 1:1 sex ratio. Analysis of Eiffel Tower and Sintra samples separately resulted in 1.1:1 and 0.6:1 ratios, respectively, but these ratios were not significantly different from 1:1 or from each other.

Gametogenesis

Spermatogenesis

Both commensal and free-living Lucky Strike polynoids exhibit filiform sperm heads (lengths: 50 µm in Branchipolyne cf. seepensis, 30 to 40 µm in Opisthotrochopodus n. sp.; Fig. 4). Testes in B. cf. seepensis are

Fig. 4. Male reproductive structures. (A to C) Branchipolyne cf. seepensis. (D) Opisthotrochopodus n. sp. sg = spermatogonia, sp = sperm packets, ss = sperm sac with filiform sperm, fs = filiform sperm. Scale bars: A = 100 µm, B–D = 50 µm
fused, with 3 distinct regions of sperm production: (1) developing spermatogonia (4 to 6 μm) and spermatocytes (Fig. 4A–C), (2) spermatids and packets of filiform sperm (comprising approximately 20 to 50 sperm in a given longitudinal section) and (3) an area of enclosed sperm storage containing dense aggregates of presumably mature sperm. In Opisthotrochopodus n. sp., sperm occur in a large region extending the length of 6 segments. Developing sperm in this production region (i.e. spermatogonia [3 to 5 μm], spermatocytes, and spermatids) are all contained between the gut and the ventral and lateral body walls, while filiform sperm occur in tubules running through the region (Fig. 4D).

Oogenesis

The ovaries of Branchipolynoe cf. seepensis occur ventral to the gut, and seem to be attached laterally. In Opisthotrochopodus n. sp., the ovaries attach ventrolaterally. Early developing oocytes can generally be seen quite far toward the sides of the body (Fig. 5D).

Both species in this study exhibit intraovarian oogenesis with oocytes surrounded by follicle cells (Fig. 5C), which is consistent with observations on other polynoid species (e.g. Harmothoe imbricata; Garwood 1978, 1981). The functional role of follicle cells in the development of oocytes of these species cannot be determined from the samples available. Developing oocytes of Branchipolynoe cf. seepensis maintain a close association with branching blood vessels (Fig. 5A), which has also been observed in H. imbricata (Garwood 1981). Blood vessel-oocyte associations are not seen within Opisthotrochopodus n. sp., but it is possible that these delicate associations may have been destroyed during histological processing.

Both species exhibit 3 distinct stages of oocyte development: previtellogenic oocytes, mature oocytes with a germinal vesicle, and mature oocytes without a germinal vesicle (Fig. 5A–D). The smaller, previtellogenic oocytes (Fig. 5A,C,D) are numerous, while larger,

![Fig. 5. Female reproductive structures. (A) Branchipolynoe cf. seepensis; previtellogenic oocytes in close association with blood vessels (bv); vo = vitellogenic oocyte. (B) B. cf. seepensis; egg sac with mature oocyte (o) lacking a germinal vesicle and filiform sperm (fs). (C) Opisthotrochopodus n. sp.; paired ventrolateral ovaries with previtellogenic and vitellogenic oocytes; arrows point to follicular cells that surround developing oocytes in both species. (D) Opisthotrochopodus n. sp.; filiform sperm (fs) in tubules with developing oocytes; vnc = ventral nerve cord, pv = previtellogenic oocytes. Scale bars = 100 μm.](image-url)
mature, vitellogenic oocytes with well-developed yolk granules and conspicuous germinal vesicles (Fig. 5A,C) are present in fewer numbers. Maximum oocyte diameter was comparable for the 2 Lucky Strike species. *Branchipolynoe cf. seepensis*, 395 μm; *Opisthotrochopodus* n. sp., 420 μm. For *B. cf. seepensis*, where oocytes were subsampled in individuals, mature oocytes ≥300 μm in diameter represented 0 to 3% of the subsample populations. *Opisthotrochopodus* n. sp. ranged from 0 to 5% mature oocytes per individual out of the total number of oocytes examined. Lowest percentages of mature oocytes (0 to 1%) were found exclusively in immature worms of small size. No intermediate oocyte stages were apparent, which suggests that both species undergo rapid vitellogenesis. Large, yolky oocytes in which the germinal vesicle is absent, probably broken down, always occur in specialized sacs or tubules running along the ventral wall of the coelom (Fig. 5B). In *B. cf. seepensis*, this sac is lined with filiform sperm (Fig. 5B). This sperm-lined egg sac splits anteriorly into 2 smaller ducts or tubules, both filled with dense filiform sperm. Filiform sperm found in females are comparable in head length (50 μm) to sperm found in males of the species. In *Opisthotrochopodus* n. sp., a smaller ventro-central tube is present, which is sometimes filled with sperm (Fig. 5D). This duct occasionally branches off into tubules that run to the body wall on either side. The nature of these sperm-lined sacs, tubes, and ducts is unknown, although they may be sites of sperm storage and egg fertilization.

Fig. 6. Relationship between log(wet weight [mg]) and log(length [mm]). (A) *Branchipolynoe cf. seepensis*. (B) *Opisthotrochopodus* n. sp. A constant of 0.1 was added to all wet weights to adjust for samples with weights less than the detectable limits; n = sample size; \( r^2 \) = correlation coefficient.

Fig. 7. *Branchipolynoe cf. seepensis*. Size-frequency distributions of total length (mm) by sex and site. (A, B) Eiffel Tower; (C, D) Sintra. Vertical bars represent mean frequencies with 1 standard deviation for a given size class, determined from 5 replicate samples from each site. Sample sizes (number of individuals): A, 192; B, 201; C, 110; D, 204.
Population structure

For size-frequency analysis of the population structure of the 2 species of polynoids, we use body length as a measure of total body size. The relationship between length and wet weight for each species is highly correlated ($r^2 > 0.8, p < 0.001$, Fig. 6), indicating that length is a reliable estimator of total body size.

*Branchipolynoe* cf. *seepensis*. Size-frequency histograms of worm length are normally distributed within sexes at both Sintra and Eiffel Tower, suggesting that recruitment is a continuous or frequently intermittent event rather than an event at discrete and resolvable intervals (Fig. 7).

*Opisthotrochopodus* n. sp. As in *Branchipolynoe* cf. *seepensis*, size-frequency histograms of worm length are normally distributed in both sexes within sites and when both sites are combined (Fig. 8). This is consistent with continuous or frequent recruitment to the population. Mean length of Eiffel Tower females was significantly larger than that of Sintra females ($p < 0.01$; mean difference $= 0.6 \text{ mm}$), but mean male size did not differ between the 2 sites.

Oocyte size-frequency distributions

*Branchipolynoe* cf. *seepensis*. Smallest oocyte size-classes were the most abundant in all individuals regardless of site (Fig. 9A). The smallest oocyte size-class measured (25 to 50 $\mu$m) also had the greatest variance about the mean. Plots of oocyte diameter versus worm length (Fig. 10A) reveal that female individuals < 15 mm in total length were reproductively immature, lacking mature, yolk-rich oocytes (>250 $\mu$m). Mean oocyte diameter was significantly correlated with worm length for combined Eiffel Tower and Sintra individuals ($r = 0.38$; Fig. 11A). No significant difference was found in mean oocyte diameter among worms collected from Eiffel Tower and Sintra ($p = 0.3$). ANOVA components indicate that, although variation of oocyte diameters within worms is greater than variation between worms, there are significant differences among worms within each site ($p < 0.001$).

*Opisthotrochopodus* n. sp. As in *Branchipolynoe* cf. *seepensis*, smallest oocyte size-classes were the most
abundant in all individuals regardless of site (Fig. 9B). The smallest oocyte size-class measured (25 to 50 μm) also had the greatest variance about the mean. Plots of oocyte diameter versus worm length (Fig. 10B) reveal that one of the smallest individuals examined (<4 mm total length) contained what we infer to be yolk-rich oocytes (cytochemical analysis was outside the scope of this project). Mean oocyte diameter was significantly correlated with worm length for combined Eiffel Tower and Sintra individuals ($r = 0.82; \text{Fig. 11B}$). We also found significant ($p < 0.001$) correlations between number of oocytes per individual and number of mature oocytes per individual versus worm length in *Opisthotrochopodus* n. sp. (Fig. 12). ANOVA indicates that there is no significant difference in mean oocyte diameter between the 2 sites and that, although variation of oocyte diameters within worms is greater than between worms, there are significant differences among worms within each site ($p < 0.001$).

**DISCUSSION**

Sexual dimorphism of the type observed in Lucky Strike polynoid polychaetes is described in detail for *Harmothoe imbricata* by Daly (1972). Gametes in *H.*
imbricata are released through the nephridial papillae and, in polytelic species, these papillae elongate during sexual maturation and regress following spawning to near-juvenile conditions. Lack of regressed papillae in specimens from Lucky Strike is consistent with asynchronous gametogenesis in these species. In addition to sexually dimorphic variation in numbers of pairs of papillae, females of Branchipolynoe cf. seepensis are typically larger than males (by ~5 to 10 mm body length). Sexual maturation in females of this species is not reached until they approach the mean size of males (~15 mm). The size differential between sexes in this species seems likely to be an accommodation to the commensal lifestyle of the worm, where small males might move more adeptly between mussels while females grow large and fecund within the mussel mantle cavity. There is also a significant size difference between sexes in Opisthotrochopodus n. sp., but in this species females are slightly smaller than males.

We examined our specimens carefully for evidence of hermaphroditism, especially since Hourdez (in Jolivet 1996) suggests protandric hermaphroditism for Branchipolynoe cf. seepensis. While sexual dimorphism in size and the female bias in the population sex ratio are consistent with protandric hermaphroditism in this species and abundant, mature sperm were observed in every mature female, we observed no instance of male germinal tissue giving rise to female germinal tissue, nor did we see any developing sperm stages (spermatogonia, spermatocytes, or spermatids).
in the females. Further, small, immature females contain no recognizable remnant of male reproductive tissue based on our histological sections, as might be expected if the species was a protandric hermaphrodite. Sperm of both Lucky Strike polynoid species have elongate heads (50 μm), a morphology that is suggestive of specialized transfer. In comparison, mature spermatozooa of several polynoid species (*Lepidonotus* sp., *Harmothoe imbricata*, *H. impar*) and other polychaete species with external fertilization have head lengths on the order of 2.7 to 3.6 μm (Rouse 1988, Bentley & Sermes 1992). Sperm with elongate heads are in general associated with species that exhibit a modified method of sperm transfer or storage, in contrast to ectaquasperm (*sensu* Rouse & Jamieson 1987) which are most often shed freely into the water (Rice 1992). Ultrastructural studies are needed to elaborate details of sperm morphology in the vent polynoid species. It is not possible to determine if sperm transfer in the Lucky Strike polynoid species occurs by release of spermatoorphes into the water or by copulation. Our observations at present are most consistent with sperm storage and internal fertilization in both *B. cf. seeepensis* and *Opisthotrochopodus* n. sp., especially given the presence of mature oocytes without germinal vesicles in sperm-lined sacs of *B. cf. seeepensis*.

Maximum oocyte sizes of both Lucky Strike polynoid species exceed 300 μm. Given that paraffin techniques can result in up to 30% shrinkage in tissues, our oocyte measurements are likely to be underestimates of actual egg size. Polynoid polychaetes are generally free-spawning and have small eggs (X = 117 μm) that undergo planktotrophic development (Wilson 1991, Giangrande 1997). Although Olive (1985) suggests that larvae deriving from eggs with diameters >180 μm should rarely be planktotrophic, egg size is not always a good indicator of developmental mode (Giangrande 1997). Nevertheless, large eggs (>300 μm) almost always give rise to either lecithotrophic larvae or direct development (Giangrande 1997). We found polynoid juveniles among our samples, but were unable to identify them to species or to determine if these represent direct development or post-larval stages. We observed no evidence of brooding beneath the elytral scales, as is the case in at least 1 polynoid (*Harmothoe imbricata*; Blake 1975).

Intraovarian oogenesis, as observed in both Lucky Strike polynoid species, and close association of developing oocytes with blood vessels (observed in *Branchipolyneae* cf. *seeepensis*) are characters sometimes indicative of rapid vitellogenesis (Eckelbarger 1983). In fast egg-producing species, oogenesis can be compressed to only hours or days (versus months in slow egg-producers), as long as food supply is maintained (Eckelbarger & Watling 1995). As already emphasized, the lack of intermediate stages of oocyte development in Lucky Strike polynoids provides further evidence that oogenesis is rapid in these species. While we cannot assess the actual rate of oogenesis in Lucky Strike polynoids, morphological indications of rapid oogenesis suggest that in situ experimental methods might be applied during the course of a single dive series (usually 2 to 3 wk) to provide a measure of this rate. According to Eckelbarger & Watling (1995), fast egg-producing species are effectively decoupled from the environmental cues that regulate egg production in synchronous spawners—as long as there is food, there are eggs.

In any 1 individual, only a small percentage of the total number of oocytes are mature in either species of Lucky Strike polynoid. We were able to identify immature specimens, but found no evidence for 'spent' individuals or for overwhelmingly 'ripe' individuals. In both species, oocyte diameters were highly variable and broadly distributed within individuals, and there were significant differences in mean oocyte diameters among worms within sites. These observations lead us to conclude that gametogenesis is asynchronous in both species, both at the individual and at the population level. This condition, combined with the style of vitellogenesis, putative sperm storage capabilities, and habitat conditions of presumably unlimited food resources lead us to infer frequent spawning in these species, with fertilized eggs released to the surrounding environment ad libitum. The combination of 110 juvenile polynoids [unidentified to species] in our samples and normal distributions of size-frequency histograms within sexes for *Branchipolyneae* cf. *seeepensis* and within both sexes combined for *Opisthotrochopodus* n. sp. is consistent with frequent rather than episodic recruitment in these species.

We found no substantive evidence for site-specific variations in the reproductive biology of either polynoid species from Lucky Strike. Thus, despite reported differences in the fluid chemistries of these 2 sites (Von Damm et al. 1998) and in the carbon and nitrogen isotopic composition of the mussels which dominate the invertebrate biomass (Trask & Van Dover 1999), these sites appear to be homogeneous at the level of resolution of polynoid reproduction.

Not unexpectedly, mean oocyte diameters, total number of oocytes and number of mature oocytes may all be correlated with body size. This is especially well-documented for *Opisthotrochopodus* n. sp., where we were able to count and measure all oocytes >25 μm in each individual. There is thus clearly a requirement for systematic sampling by size for studies of reproductive biology in polynoid polychaetes. We chose to sample with probability proportional to size frequency to
obtain measures of the reproductive biology within populations, but other strategies may also be useful (e.g. restricting specimens to multiple individuals in a single size class).

To our surprise, we find little evidence of different reproductive strategies in the commensal and free-living polynoid species, apart from the much greater size and fecundity of female *Branchipolynoe cf. seepensis* and their skewed sex ratio favoring females. Large and fecund female polynoid polychaetes at vents are not restricted to the commensal mien, however, nor are free-living species in the genus *Opisthotrochopodus* restricted to small size and low fecundity (based on unpublished observations by C.L.V.D. on polynoid polychaetes from the East Pacific Rise).

Within the context of the reproductive biology of hydrothermal vent polychaetes, Lucky Strike polynoids are remarkable in the large size of their oocytes. Their sexual dimorphism, elongate sperm, asynchronous gametogenesis within individuals and among individuals within a population, apparent sperm storage and specialized mode of fertilization are attributes shared with various alvinellid and ampharetid species (Table 5).

Acknowledgements. Dr. Marion Pettibone assisted us with polynoid identifications. We are especially grateful to Kevin Eckelbarger for his gracious and enthusiastic tutorial on polychaete reproduction. We are indebted to the personnel of the Deep Submergence Laboratory (Woods Hole Oceanographic Institution) for collections of specimens and to Dan Fornari and Susan Humphris who were Chief Scientists of the LUS-TRE Expedition. We are also grateful to anonymous reviewers who helped us clarify our prose and interpretations. This work was supported by NSF-GEO-9505579.

| Table 5. Reproductive characteristics of hydrothermal vent polychaete species (all gonochoric). EPR = East Pacific Rise; JdF = Juan de Fuca Ridge; Ex = Explorer Ridge; MAR = Mid-Atlantic Ridge |
|---------------------------------|---------|------------------|------------------|------------------|---------------------------------|
| Species                        | Egg diam. (μm) | Inferred mode of development | Gametogenesis (M:F) | Sex ratio | Comments and inferences |
| Alvinellidae                   |          |                  |                  |          |                          |
| *Alvinella pompejana*          | 200      | Lecithotrophic   | Not available    | -        | -                      |
| EPR                            |          |                  |                  |          | Chevaldonné & Jollivet (1993), Chevaldonné et al. (1997) |
| *Paralvinella grasslei* EPR    | 275      | Direct           | Synchronous      | -        | 4000 ± 1400 oocytes in coelomic cavity; sexual dimorphism; compulsory passage of oocytes through inseminated spermathecae; suggestions of pairing and brooding |
| *Paralvinella pandorae pandorae* JdF, Ex | 215 | Brooder | Asynchronous | 1:1 | 4500 oocytes per worm; sperm morphology suggests specialized mode of fertilization |
| *Paralvinella paimulormis* JdF, Ex | 260 | Lecithotrophic | Synchronous | - | 18000 oocytes per worm; demersal larvae |
| Ampharetidae                   |          |                  |                  |          |                          |
| *Amphipisamytha galapagensis* JdF | 240 | Lecithotrophic | Asynchronous | 1:1 | 5600–9600 oocytes per worm on average (max. = 12 500); early maturation; external fertilization; demersal larvae |
| Polyxidiae                     |          |                  |                  |          |                          |
| *Branchipolynoe cf. seepensis* MAR | 395 | Lecithotrophic or direct | Asynchronous | 1:2 | 0–300 mature oocytes (free in coelom) per worm; sexual dimorphism (no. of nephridial papillae and size); rapid oogenesis; sperm storage and internal fertilization |
| Opisthotrochopodus n. sp. MAR | 420 | Lecithotrophic or direct | Asynchronous | 1:1 | 100–600 oocytes (>25 μm) per worm; 0–16 mature oocytes per worm; sexual dimorphism (no. of nephridial papillae); rapid oogenesis; sperm storage and internal fertilization |
| Opisthotrochopodus n. sp. AR | 420 | Lecithotrophic or direct | Asynchronous | 1:1 | 100–600 oocytes (>25 μm) per worm; 0–16 mature oocytes per worm; sexual dimorphism (no. of nephridial papillae); rapid oogenesis; sperm storage and internal fertilization |

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Submitted: June 15, 1998; Accepted: November 21, 1998

Proofs received from author(s): May 4, 1999

Editorial responsibility: Lisa Levin (Contributing Editor), La Jolla, California, USA