

8 Abstract

9 Phylogenetic analysis has led to significant insights into the evolution of early life-history stages of marine invertebrates. 10 Although echinoderms have been a major focus, developmental and phylogenetic information are relatively poor for ophiuroids, the most species-rich echinoderm class. We used DNA sequences from two mitochondrial genes to develop a phylogenetic hypothesis for 11 14 brittlestar species in the genus Macrophiothrix (Family Ophiotrichidae). Species are similar in adult form and ecology, but have 12 diverse egg sizes and modes of larval development. In particular, two species have rare larval forms with characteristics that are inter-13 mediate between more common modes of feeding and non-feeding development. We use the phylogeny to address whether interme-14 diate larval forms are rare because the evolution of a simplified morphology is rapid once food is no longer required for 15 16 development. In support of this hypothesis, branch lengths for intermediate forms were short relative to those for species with highly 17 derived non-feeding forms. The absolute rarity of such forms makes robust tests of the hypothesis difficult.

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19. Keywords: Ophiuroid; Brittlestar; Facultative feeding; Pluteus; Mode of development; Life-history transition; Mitochondrial DNA

20 1. Introduction

The evolutionary analysis of early life cycle diversity in 21 marine invertebrates has advanced significantly over the 22 last decade from the use of phylogenetically based com-23 parative methods (Cunningham, 1999; Hadfield et al., 24 1995; Hart et al., 1997; Jeffery and Emlet, 2003; Levitan, 25 2000; Rouse and Fitzhugh, 1994; Villinski et al., 2002). 26 These analyses have confirmed or challenged a number of 27 assumptions about the order, timing, and frequency of 28

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1055-7903/\$ - see front matter 2004 Published by Elsevier Inc. doi:10.1016/j.ympev.2004.09.011

change in developmental and life-history characters 37 (reviewed in Hart, 2000). 38

Certain echinoderm taxa feature prominently in this 39 work (Arndt et al., 1996; Emlet, 1990; Hart et al., 1997; 40 Jeffery and Emlet, 2003; Smith et al., 1995; Wada et al., 41 1996; Wray, 1996). Gametes and embryos of sea stars 42 (Asteroidea) and sea urchins (Echinoidea) are easy to 43 obtain and culture (Strathmann, 1987), and some larval 44 traits can be inferred indirectly from egg size (Emlet et al., 45 1987) or adult traits (Emlet, 1989). In contrast, the study 46 of development and larval biology in brittlestars (Ophiu- 47 roidea), the most-species rich echinoderm class, has been 48 hindered by the cryptic lifestyles of adults, a lack of reli- 49 able methods for inducing spawning and oocyte matura- 50 tion (Selvakumaraswamy and Byrne, 2000; Strathmann, 51 1987), and the absence of adult characters that can 52 diagnose larval traits (Emlet, 1989; Hendler, 1978). As in 53 better documented classes, egg size in ophiuroids is corre- 54

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55 Table 1

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56 Modes of larval development and maximum likelihood branch lengths for *Macrophiothrix* species with obligate non-feeding (group 1) or intermediate (group 2) larval forms and feeding modes

Species	Egg volume (nl)	Larval morphology	Can feed?	Must feed?	Branch length							
					No clo	ock		With clock				
					16S	COI	COI Both		COI	Both		
M. belli	35.1	Non-pluteus	No	No	.129	.239	.208	.123	.286	.214		
M. lampra	12.6	(non-pluteus)	(No)	(No)	.110	na	na	.111	na	na		
M. nereidina	9.7	Non-pluteus	No	No	.154	na	na	.123	na	na		
M. caenosa	7.4	Pluteus	No	No	.082	.097	.086	.071	.148	.115		
M. rhabdota ^a	6.3	Pluteus	Yes	No	.023	na	na	.018	na	na		
(to <i>lorioli</i>)					.062	.089	.084	.042	.081	.062		
M. lorioli	2.4	Pluteus	Yes	Yes		na			na			
M. longipeda	1.9	Pluteus	Yes	Yes		na			na			
M. koehleri	1.6	Pluteus	Yes	Yes		na	7		na			
M paucispina	<1.0	(pluteus)	(Yes)	(Yes)		na			na			

71 127 Also shown are obligate feeding species (group 3) that are important to the analysis of branch lengths. Larval character states are known from larval 72 128 culture (R. Podolsky, unpublished data) except for those of M. lampra and M. paucispina, which are inferred from egg size relative to other species. 73 Of nine additional Macrophiothrix species for which we have egg size data, all are likely to be obligately feeding plutei. Because COI data were miss- 129 74 ing for some species that are relevant to the branch length analysis (*M. lampra*, *M. nereidina*) and COI genetic distances were saturated for the two 130 outgroups, branch lengths in units of expected change per nucleotide site are given for 16S (TVM + G model, 18 taxa), COI (GTR + G + I, 12 ophio-131 75 trichid taxa only), and both genes analysed together (GTR + G, 12 ophiotrichids plus Ophionereis) under maximum likelihood substitution models 132 76 estimated independently for each analysis. na, not available or applicable. 77 133

¹⁷⁷ ^a The relevant branch length for estimating persistence time of the intermediate larval form of *M. rhabdota* depends on whether the closest relative
 ¹⁷⁵ with feeding larval development is *M. paucispina* (inferred from egg size, first line) or *M. lorioli* (known from larval culture, second line); see Fig. 1.
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lated with mode of development (Hendler, 1991) and
developmental patterns are diverse (sensu McEdward and
Janies, 1997). However, no species-level phylogenies have
been published for ophiuroids that would aid the analysis
of early life cycle evolution (McEdward and Miner, 2001).

88 The brittlestar genus Macrophiothrix (Clark; F. 89 Ophiotrichidae) is common to certain Indo-west Pacific 90 coral reef habitats and especially diverse (21 species) in 91 tropical Australia (Hoggett, 1990). Despite the occur-92 rence of exceptionally high local diversity, adults of 93 different Macrophiothrix species are similar in size, form, 94 and ecological habits (Hoggett, 1991). The genus, how-95 ever, exhibits remarkable variation and specificity in 96 early life cycle characters, involving species-specific sperm chemotaxis (Miller, 1997) and extensive variation 97 98 in egg size and development mode. At Lizard Island, 99 Australia, 12 Macrophiothrix congeners vary more than 100 60-fold in average egg volume (Podolsky, unpublished data) and produce larval forms that include a typical 101 and morphologically complex feeding pluteus, a mor-102 103 phologically simple non-feeding larval form, and two 104 forms that are functionally intermediate (Table 1). Larvae of *M. rhabdota* are facultative planktotrophs that do 105 not require food to complete development, whereas 106 107 larvae of *M. caenosa* do not feed but retain the pluteus 108 morphology (R. Podolsky, unpublished data). The rarity of these intermediate larval forms-relative to the large 109 110 number of evolutionary transitions from feeding to

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non-feeding larval forms (Duda and Palumbi, 1999; 139 Hart et al., 1997; Jeffery and Emlet, 2003; Lieberman et 140 al., 1993; Wray, 1996)–led Wray (1996) and others to 141 propose that intermediate forms are evolutionarily 142 unstable and short-lived. Phylogenetic information can 143 be used to address this hypothesis (Hart, 1996). 144 We use information from mitochondrial DNA 145

we use information from mitochondrial DNA 145 sequences to resolve relationships among 14 species clas- 146 sified in the genus *Macrophiothrix*, two species in the 147 genus *Ophiothrix* not recently referred to *Macrophiothrix* 148 (Hoggett, 1990), and two outgroups from related families. 149 We use the phylogeny to infer relationships between spe- 150 cies with functionally intermediate, putatively transitional 151 larval forms and species with non-feeding larval morpho- 152 logies, and to test the hypothesis that the former species 153 are more recently derived (Hart, 1996). 154

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We obtained new mtDNA sequences from the large 159 subunit ribosomal RNA (16S) and cytochrome *c* oxidase 160 subunit I (COI) genes for 16 ophiotrichid brittlestar spe-161 cies in the genera *Macrophiothrix* and *Ophiothrix* and 162 for two outgroup taxa (the ophiocomid *Ophiarthrum* 163 *pictum* and the ophionereid *Ophionereis porrecta*; see 164 Appendix A for details of laboratory methods). We ana-165 lyzed phylogenetic information from both genes 166

2. Materials and methods

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167 together by maximum likelihood (ML) methods, and
168 used bootstrapping and posterior probabilities as esti169 mates of clade support (see Appendix B for details of
170 analytical methods).

171 To illustrate the utility of the phylogeny for testing 172 hypotheses about life history evolution, we inferred and 173 compared times of persistence for species with highly 174 derived, non-feeding larval forms and for species with 175 more intermediate modes of larval development. The 176 intermediate forms develop from relatively large eggs and 177 do not require food, but retain phenotypic traits of sus-178 pension feeding larvae that are absent from the more 179 derived forms. If branch lengths leading to intermediate 180 forms are short relative to those leading to highly derived 181 non-feeding forms, this pattern would support the 182 hypothesis that intermediate phenotypes are short-lived 183 steps in a transition from obligate feeding to simplified 184 non-feeding development, rather than stable develop-185 mental modes (Hart, 1996). (This hypothesis assumes 186 that rates of molecular evolution are similar across lin-187 eages with different derived modes of development.)

188 We asked whether the ML branch lengths leading to 189 two ophiotrichids with intermediate larval phenotypes 190 (M. caenosa and M. rhabdota) are shorter than those 191 leading to three species that are known (M. nereidina, M. 192 belli) or inferred (M. lampra) to have obligate non-feed-193 ing larvae. Other species (M. longipeda, M. koehleri, M. 194 paucispina, and M. lorioli) that are known or inferred to 195 produce obligately feeding plutei (R. Podolsky, unpub-196 lished data) were used to establish the earliest possible 197 times of divergence from species with obligate feeding 198 larval forms (Table 1). 199

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201 3. Results

203 General sequence characteristics (including missing
204 COI sequence data for some species) are summarized in
205 Appendix A. Within- and among-species genetic dis206 tances are summarized in Appendix B.

208 3.1. Tree topology

210 We found a single ML tree with a $-\ln(L)$ score of 211 7788.9 (Fig. 1). Seven clades were moderately or strongly 212 supported by high bootstrapping percentages and posterior probabilities: M. leucosticha + M. megapoma; this 213 214 species pair in a larger clade with M. caenosa, M. koeh-215 *leri*, and *M. longipeda*; *M belli* + M. *nereidina*; *M*. 216 demessa + O. trilineata; and the clade (M. lampra) (M. lorioli (M. rhabdota + M. paucispina). The three most 217 218 basal nodes, and some other interior branches in the tree, 219 were short and poorly supported, which could reflect low 220 resolving power of these genes for these taxa or rapid 221 divergence of clades. We were unable to identify the sis-222 ter groups of three species: M. robillardi, M. propiniqua,

O. caespitosa. Only one of these (*M. robillardi*) lacks 223 COI data (see Appendix A), which suggests that includ- 224 ing these taxa with missing data did not greatly impede 225 our ability to find some well-supported relationships 226 based on 16S only (see also Appendix B).

Ophiothrix and Macrophiothrix species did not appear 228 to form reciprocally monophyletic groups: O. trilineata 229 was strongly inferred to be sister species to M. demessa 230 (Fig. 1). The unresolved relationship between O. caespi- 231 tosa and other ophiotrichids is consistent with some 232 Ophiothrix as sister group to a paraphyletic Macrophio- 233 thrix. However, the low resolution of basal relationships 234 prevents us from drawing a firm conclusion about 235 monophyly for these two genera: a tree in which O. 236 caespitosa + O. trilineata was the sister group to Macro- 237 phiothrix species (with other relationships among Macro- 238 phiothrix as shown in Fig. 1) had a $-\ln(L)$ score of 7799.5 239 and was not significantly less likely than the ML tree 240 (7788.9) by the Shimodaira-Hasegawa test in PAUP* 241 (Swofford, 2002, P = 0.177). We therefore cannot reject the 242 hypothesis that Macrophiothrix (sensu Hoggett, 1991) is a 243 monophyletic clade with respect to Ophiothrix. 244

3.2. Branch lengths of derived larval forms

247 The terminal branches leading to species with inter- 248 mediate larval forms (M. rhabdota, M. caenosa) were rel- 249 atively short (e.g., 0.023, 0.082, for 16S without a clock) 250 compared to terminal branches for species with simpli- 251 fied, non-feeding larval forms. We obtained qualitatively 252 the same result when we allowed sister lineages to differ 253 in rates of molecular evolution and when we enforced a 254 molecular clock (Table 1). Patterns were similar for 16S 255 sequences alone, for COI, and for both genes analyzed 256 together. Both species with intermediate forms were 257 close relatives of species known to develop as feeding 258 larvae (M. lorioli; M. longipeda, and M. koehleri), so the 259 intermediate form is assumed to have arisen along the 260 short terminal branches (Table 1). If M. paucispina 261 (whose mode of larval development is uncertain) were 262 also a transitional non-feeding pluteus larva, then the 263 persistence time for that transitional form could be as 264 long as the sum of the terminal and internal branches 265 separating M. rhabdota and M. paucispina from their 266 common ancestor with M. lorioli. Measurements of egg 267 size from a single museum specimen of M. paucispina 268 (Australian Museum J22076)-which was dried before 269 ethanol preservation, and could have experienced a 270 reduction in egg dimensions-indicated an egg volume 271 (<1 nl) characteristic of species with planktonic, feeding 272 pluteus larvae (Table 1). This observation suggests that 273 the transitional larval form of M. rhabdota evolved after 274 its recent divergence from M. paucispina, and supports 275 the inference of the shorter branch lengths in Table 1. 276

In contrast, the terminal branches leading to sister spe-277 cies (*M. belli*, *M. nereidina*) with obligate non-feeding lar-278

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Fig. 1. Phylogram for 16 Macrophiothrix and Ophiothrix brittlestar species and an outgroup (Ophionereis) based on maximum likelihood analysis of 311 367 combined 16S rDNA and COI sequences (six-rate GTR + G model). Numbers beside branches are bootstrap percentages (bold) and Bayesian pos-312 terior probabilities (italics); only values >50% are shown. Heavy lines show branch lengths relevant to the analysis of modes of larval development 368 (shown as abbreviations to the right of some taxon names): feeding planktonic larvae (P), non-feeding lecithotrophic larvae (L), transitional pluteus 313 369 larvae that cannot feed (T_1) , transitional pluteus larvae that can feed (T_p) (see Table 1). The mode of larval development for *M. paucispina* is inferred 370 314 from egg size in a museum specimen and is considered uncertain but probably a feeding planktonic larva (P?). 315 371

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317 val development were longer (e.g., 0.129, 0.154 for 16S 318 without a clock). The terminal branch lengths for these 319 sister species give a minimum estimate of persistence time 320 for species with obligate non-feeding development if this 321 larval form and life history evolved once in their common 322 ancestor. A third, independent terminal branch for a spe-323 cies assumed to have non-feeding larvae (M. lampra) gives 324 a maximum estimate of this persistence time that is also 325 longer (0.110) than branches leading to species with inter-326 327 mediate larval forms. Branch lengths for M. belli based on COI (which were missing for *M. lampra* and *M. nereidina*) 328 and the combined sequences were also longer than the 329 comparable branch lengths for species with intermediate 330 331 forms. Of course, any of these derived larval forms could 332 have evolved at any time since the divergence of the species from an ancestor with obligate feeding larvae, and the 333 number of such lineages is small. In addition, some long 334

373 terminal branches might be shortened by the discovery of 374 sister group relationships with other Macrophiothrix spe- 375 cies that we were not able to sample. 376

4. Discussion

380 4.1. Taxonomic and phylogenetic resolution in 381 *Macrophiothrix*

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Analysis of mtDNA sequences from Macrophiothrix 384 and *Ophiothrix* species identified some sister-group rela- 385 tionships that are statistically well-supported, but left 386 other relationships unresolved. Improved taxonomic 387 sampling of more Macrophiothrix species using more 388 slowly evolving molecular characters is needed to fully 389 resolve these basal relationships. Nevertheless, our 390

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391 current results provide a useful context for comparative392 analyses of developmental evolution among ophiotri-393 chids.

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394 Our mtDNA sequence comparisons add to earlier 395 efforts of Hoggett (1991) to resolve generic boundaries 396 involving species in Macrophiothrix and related sub-397 genera of Ophiothrix. Among the 16 species for which 398 we generated mtDNA sequences, we found some well-399 supported clades of species that Hoggett had assigned 400 to Macrophiothrix. However, two other Macrophio-401 *thrix* were not reliably grouped with other members of 402 the genus. First, morphological characters reliably 403 placed M. propingua and M. robillardi with other 404 Macrophiothrix (Hoggett, 1990), but 16S and COI 405 characters could not reliably identify the sister group 406 of either. Second, analyses of mtDNA tend to group 407 M. demessa with O. trilineata, though Hoggett (1990) 408 assigned each species with confidence to its respective 409 genus on the basis of adult morphological and allo-410 zyme evidence.

411 We also found examples of strong conflict between 412 our mtDNA phylogeny and cladograms based on mor-413 phology or allozymes (Hoggett, 1990). Some species that 414 share numerous adult morphological apomorphies, such 415 as M. belli, M. caenosa, and M. paucispina (Hoggett, 416 1990; Fig. 2.17), are distant relatives in the mtDNA trees. 417 Other species with highly divergent allozyme alleles, 418 such as M. leucosticha and M. megapoma (Hoggett, 419 1990; Fig. 4.3), are strongly supported as sister species in 420 the mtDNA trees. In general, the mtDNA data more 421 finely and robustly resolved species relationships for the 422 clades identified in common by the three analyses.

423 Our analyses of mtDNA and life-history variation 424 support other taxonomic revisions. For example, Hogg-425 ett (1990) reversed a long history of classification (Clark, 426 1938) by distinguishing *M. caenosa* (sp. nov.) from *M*. 427 longipeda. The main diagnostic character (shape of 428 papillae on the adult dental plate) is difficult to resolve 429 with live specimens. In contrast, the species are easily 430 distinguished by a 3.8-fold difference in egg volume and 431 qualitative differences in mode of development (R. Pod-432 olsky. unpublished data). Mitochondrial DNA 433 sequences extend this conclusion by suggesting that 434 these taxa might not even be sister species (Fig. 1). Simi-435 lar surveys of reproductive traits could be useful for 436 identifying cryptic species diversity in putative morphospecies such as *M. demessa* that have broad geographic 437 438 ranges (Philippines through Australia and several oce-439 anic islands), long mtDNA terminal branches (Fig. 1), 440 and high within-species mtDNA sequence variation 441 (Appendix B).

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443 4.2. Evolutionary analysis of early life cycle characters

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445 Efforts to understand the origins of developmental 446 diversity have focused on processes that underlie transitions between modes of development (Wray, 447 1995a). In marine invertebrates, one widespread transi- 448 tion involves the loss of a complex feeding larval mor- 449 phology in association with the evolution of large, 450 nutrient-rich eggs (Emlet et al., 1987; Strathmann, 451 1985). Intermediate phenotypes, such as facultative 452 planktotrophy, could represent short-lived steps on the 453 path toward non-feeding development and morphol- 454 ogy (Wray and Raff, 1991) or advantageous and evolu- 455 tionarily stable modes of larval development (Emlet, 456 1986; McEdward, 1997; Levitan, 2000). Estimates of 457 relative persistence times for these larval forms are a 458 powerful and novel way to evaluate these two hypothe- 459 ses (Hart, 1996; Wray, 1996). 460

Intermediate larval forms are taxonomically rare and 461 phylogenetically scattered (Alatalo et al., 1984; Crump, 462 1989; Emlet, 1986; Hart, 1996; Kempf and Hadfield, 463 1985; Kempf and Todd, 1989; Perron, 1981). As the first 464 genus documented to include more than one species 465 with an intermediate larval form-in addition to several 466 species with more derived, non-feeding forms-Macro- 467 phiothrix offered a unique opportunity to apply phylo- 468 genetic information to analysis of this transition. 469 Branch lengths from our phylogeny provided estimates 470 of maximum persistence times for intermediate forms. 471 All of these estimates (from two species, and two genes 472 analyzed separately or in combination, with and with- 473 out rate variation among lineages) were shorter than the 474 relevant branch lengths for species or clades with 475 obligate non-feeding development and simple larval 476 morphology. 477

These data are consistent with predictions of the 478 rapid transitions hypothesis (Hart, 1996; Wray, 1995b). 479 Notably, comparison between the two transitional lar- 480 val forms is also consistent with this hypothesis: the spe- 481 cies (*M. rhabdota*) with larvae that retain the capacity to 482 feed occurs on a shorter branch than the species (M. 483)caenosa) with larvae that retain a feeding morphology 484 but have lost the capacity to feed (e.g., 0.081 versus 485 0.148 for COI with a molecular clock). This observation 486 provides indirect support-because these two species are 487 part of separate lineages-for the concept of an ordered 488 transformation series (Wray, 1996) in which feeding 489 capacity and feeding morphology are lost in sequential 490 steps (R. Podolsky, unpublished data). Whether persis- 491 tence times for these intermediate forms are transient 492 enough to explain their absence in most well-sampled 493 taxa that include both feeding and simplified non-feed- 494 ing forms is uncertain, but we have failed to reject this 495 hypothesis for the most extensive phylogenetic example 496 available. As the first species-level phylogeny to address 497 early life cycle evolution in ophiuroids, the analysis 498 presented here is a starting point for improved under- 499 standing of a large, diverse, and ecologically important 500 class for which developmental and life-history data are 501 relatively scarce. 502

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503 Acknowledgment

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505 We thank J. Allen and J. McAlister for discussions 506 and contributions to larval culture and measurement, J. 507 Mader and K. Kemp for collection of samples from 508 museum collections, J. Addison for assistance with 509 sequencing, P. Berents (Australian Museum) and J. Fro-510 mont (Western Australian Museum) for access to and 511 assistance with collections, D. Posada for software assis-512 tance, K. Beckenbach and M. J. Smith for sequence data, 513 and B. Schierwater and four anonymous reviewers for 514 constructive criticism. A. Hoggett and L. Vail provided 515 research space and logistical support at Lizard Island Research Station. Supported by National Science Foun-516 517 dation grant OCE-9811121, the Canada Foundation for Innovation, and the Natural Sciences and Engineering 518 519 Research Council.

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522 Appendix A. Laboratory methods and sequence 523 characteristics

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525 The table below shows reference information for the 526 mtDNA data. Australian Museum accession numbers for 527 tissue samples are in parentheses after the species name; 528 other samples were collected from fresh specimens at Liz-529 ard Island, Australia, and fixed in 70% ethanol. Identifi-530 cations of live material were based on Hoggett (1990) and 531 verified by A. Hoggett (personnal communication). Two 532 GenBank numbers are given for species in which 16S 533 sequences were obtained from two individuals. We used 534 standard digestion and extraction methods (e.g., Gros-535 berg et al., 1996) to obtain genomic DNA from gonads, 536 gametes, or arms. From serial dilutions of these DNA 537 samples, we amplified part of each of the 16S and COI 538 genes using 'universal' PCR primers (Folmer et al., 1994; 539 Palumbi, 1996). For six species (M. caenosa, M. demessa, 540 M. koehleri, M. longipeda, M. lorioli, and M. nereidina) 541 we obtained 16S sequences from two individuals. Only 542 four species (M. belli, M. demessa, M. longipeda, and 543 Ophionereis) could be reliably amplified and sequenced 544 using the universal COI primers; all other COI amplifica-545 tions required some variable combination of ten other 546 oligonucleotide sequences (available from the authors on 547 request) that we designed from preliminary sequence data and comparison with complete COI sequences for two 548 549 other brittlestars (Scouras et al., 2004). We were unable to 550 amplify and sequence COI from M. lampra, M. nereidina, 551 M. paucispina, and M. robillardi using the universal or 552 modified primers. We sequenced these PCR products 553 directly using the PCR primers, Thermosequenase (USB), 554 and infrared dye-labeled dideoxy terminators (Li-Cor), 555 and we visualized the fragments on a Li-Cor 4200L-2 556 DNA sequencer. We used the manufacturers' standard 557 cycle sequencing and electrophoresis conditions. Comple-558 mentary sequences were compared and edited in AlignIR

(Li-Cor). Most PCR products were sequenced from both 559 strands through the end of the opposite primer, but in 560 some cases we obtained only partial sequences of the 561 product or sequences from just one strand, so that the 562 lengths of sequences for each gene varied among taxa. 563

The 24 aligned 16S sequences were 596 bp long; 184 of 564 these sites were parsimony-informative (with gap sites 565 coded as missing). Many alignment gaps corresponded to 566 insertions in the *Ophiarthrum* sequence relative to *Ophi*- 567 *onereis* and the ophiotrichids. Some gaps were large (e.g., a 568 37-base insertion near the 3' end of the 16S sequence of 569 *M. propinqua*). The 14 aligned COI sequences were 654 570 bp; 225 of these sites were parsimony-informative. We 571 found one amino acid deletion: a Gly/Gln codon near the 572 5' end of the *Ophiarthrum* sequence. 573

Taxon names, sequence lengths, and GenBank accession 575 576

numbers			57
Species	16S	COI	57
Ophiotrichidae			579
Macrophiothrix belli	492 (AY365143)	571 (AY365144)	580
M. caenosa	492 (AY365145)	654 (AY365146)	581
	491 (AY365147)		582
M. demessa	483 (AY365148)	654 (AY365149)	583
	508 (AY365150)		584
M. koehleri	490 (AY365151)	654 (AY365152)	583
	489 (AY365153)		580
M lampra	494 (AY365154)		30 J
(J14001)	· · · · · ·		580
M leucosticha	511 (AY365155)	565 (AY365156)	590
M. louginada	491 (AY365158)	654 (AY365159)	591
M. longipeda	480 (AV365160)	001 (111 000 107)	592
Maniali	511 (AV365161)	654 (AV365162)	593
M. lorloll	402 (AV265162)	054 (111 505 102)	594
14	493 (A 1 303103) 494 (A V 365165)	654 (AV365166)	595
M. megapoma	404 (AY265167)	054 (711 505 100)	596
M. nereidina	494 (A ¥ 303107)		597
	492 (AY 365169)		598
M. paucispina	498 (AY 365170)		599
M. propinqua	542 (AY365172)	654 (AY365173)	600
M. rhabdota	493 (AY365174)	654 (AY365175)	601
M. robillardi	491 (AY365176)		602
(J19445)			604
Ophiothrix	442 (AY365179)	654 (AY365180)	604
caespitosa (J16397)			606
O. trilineata	491 (AY365182)	654 (AY365183)	607
(J19233)			608
o 1			609
Ophionereidae			610
Ophionereis porrecta	507 (AY365184)	571 (AY365185)	611
Onhiocomidae			612
Ophiarthrum nictum	523 (AY365186)	651 (AY365187)	613
Spinarin an pictum	525 (11 505 100)	051 (11 505107)	614

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615 Appendix B. Phylogenetic methods and genetic distance

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results

618 The table below shows pairwise genetic distances 619 based on maximum likelihood (ML) models. 16S nucleo-620 tide sequences were aligned in ClustalX using the default 621 multiple alignment parameters (Jeanmougin et al., 1998). 622 COI sequences were translated using the echinoderm 623 mtDNA translation table in GeneJockey (http:// 624 www.biosoft.com), checked to confirm that they con-625 tained no frame shifts, nonsense codons, or stop codons, 626 and aligned as amino acid sequences in ClustalX.

627 We analyzed phylogenetic information from both 628 genes together (total evidence; Kluge, 1989). We favored 629 this approach because we preferred to include some data 630 for taxa where we lacked COI sequences (see Appendix 631 A) rather than to leave these taxa out of some analyses 632 entirely. Other recent studies show that adding taxa with 633 missing sequence characters tends to improve the quality 634 of total evidence analyses (Hughes and Vogler, 2004; 635 Wiens, 1998). As a quantitative check on this approach, 636 we used the incongruence length difference (ILD) test in 637 PAUP* (Swofford, 2002). We inferred a preliminary maximum parsimony (MP) tree in PAUP* with equal 638 639 weighting for transitions and transversions, then esti-640 mated likelihood scores and the Ti/Tv ratio for sequence 641 data mapped onto that MP tree under a two-parameter 642 ML substitution model. We estimated the appropriate 643 transversion weighting separately for three classes of 644 characters: 16S, COI 3rd codon positions, and all other 645 COI nucleotides. We applied these estimated transver-646 sion weightings in the ILD test by use of step matrices 647 with accelerated character transformations, heuristic 648 searching, TBR branch swapping, and 1000 iterations of 649 the test. Although we are aware of the uncertainty sur-650 rounding use of the ILD test for identifying significant 651 conflict between data partitions (e.g., Darlu and Lecoin-652 tre, 2002; Dowton and Austin, 2002; Hipp et al., 2004; 653 Yoder et al., 2001), we found no indication that the 16S 654 and COI sequences gave conflicting information 655 (P=0.722), and in subsequent analyses we used both 656 genes together with COI characters coded as missing for 657 M. lampra, M. nereidina, M. paucispina, and M. robil-658 lardi.

Our analyses of phylogenetic relationships among
ophiotrichids emphasized ML methods rather than parsimony methods because our specific goal was to infer
branch lengths leading to species with different larval
phenotypes under a biologically realistic model of DNA
sequence evolution. We used hierarchical likelihood
ratio testing in Modeltest (Posada and Crandall, 1998)

to choose the appropriate substitution model (six-rate 666 GTR + G, $\alpha = 0.2136$), then estimated the tree topology 667 using this model in PAUP* with stepwise addition of 668 taxa, TBR branch swapping, and no molecular clock. 669 We estimated nodal support for the ML tree as boot- 670 strap percentages in PAUP* (100 replicates) and as the 671 posterior probabilities of clades under the same likeli- 672 hood model in coalescent simulations using MrBAYES 673 (Huelsenbeck and Ronquist, 2001). We ran this ML sim- 674 ulation for 110,000 generations, and discarded the first 675 10,000 generations as a burn-in (in preliminary analyses 676 this number was sufficient to avoid sampling trees and 677 parameter values in the non-stationary part of the simu- 678 lation). We set the model parameter values (six substitu- 679 tion rates, alpha value for among-site rate variation, and 680 nucleotide frequencies) as priors using the "prset" com- 681 mand. We used default values for other Bayesian param- 682 eters, including number of chains (4), temperature (0.2), 683 and sampling frequency (100). 684

Because some branch length comparisons involved 685 species for which we lacked COI sequence data, we esti-686 mated these branch lengths (with and without a molecular 687 clock enforced) under the appropriate substitution model 688 identified for 16S alone (five-rate TVM+G, α =0.2823), 689 COI alone (six-rate GTR+G+I, α =0.5544, I=0.4301), 690 and for the combined data (parameters above). 691

Genetic distances between 16S sequences of conspe- 692 cific pairs (bold values below) were smallest in M. lorioli 693 (0.017) and largest in M. demessa (0.042). These dis- 694 tances are less than or comparable to conspecific genetic 695 distances based on 16S sequences of other ophiotrichids 696 (Baric and Sturmbauer, 1999) or other brittlestar fami- 697 lies (Sponer et al., 2001). Most among-species distances 698 were larger, but two sister species pairs were separated 699 by genetic distances within this range: M. leucosticha 700 and M. megapoma (0.030); M. paucispina and M. rhab- 701 dota (0.032). Genetic distances between COI sequences 702 were substantially larger, and reflected more rapid evo- 703 lution of this gene relative to 16S. COI distances 704 between the outgroup taxa and the ophiotrichids were 705 very large or undefined and are not reported in the 706 table. 707

In some preliminary phylogenetic analyses, we found 708 very long terminal branch lengths leading to *Ophiar*- 709 *thrum* in comparison to all other internal and terminal 710 branches. This result reflects the unexpectedly large 711 genetic distances between this species and all other 712 sequences. To avoid artifacts associated with this excep- 713 tionally long branch, we removed *Ophiarthrum* from the 714 analyses reported in the main text, and we used *Ophione*- 715 *reis* as the sole outgroup. 716

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Genetic distances among brittlestar mtDNA sequences for 16S rDNA (below diagonal) and COI (above diagonal). Undefined distances are indicated with an asterisk																								
	bel	cae1	cae2	dem1	dem2	koe1	koe2	lam	leu	lon1	lon2	lor1	lor2	meg	ner1	ner2	раи	pro	rha	rob	csp	tri	por	pic
belli		0.658	*	0.308	*	0.654	*	*	0.544	0.529	*	0.572	*	0.564	*	*	*	0.467	0.585	*	0.697	0.863	*	*
caenosa1	0.236		*	0.695	*	0.388	*	*	0.259	0.262	*	0.591	*	0.253	*	*	*	0.638	0.523	*	0.980	0.859	*	*
caenosa2 ^a	0.240	0.028		*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
demessa1	0.263	0.321	0.307		*	0.842	*	*	0.612	0.592	*	0.571	*	0.659	*	*	*	0.593	0.569	*	0.684	0.655	*	*
demessa2 ^a	0.265	0.316	0.293	0.042		*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
koehleri1	0.269	0.172	0.163	0.371	0.322		*	*	0.429	0.369	*	0.707	*	0.443	*	*	*	0.809	0.620	*	1.018	1.072	*	*
koehleri2 ^a	0.257	0.146	0.137	0.334	0.331	0.041		*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
<i>lampra</i> ^a	0.297	0.277	0.276	0.291	0.311	0.341	0.286		*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
leucosticha	0.291	0.141	0.149	0.415	0.361	0.185	0.162	0.362		0.268	*	0.495	*	0.017	*	*	*	0.647	0.576	*	0.741	0.759	*	*
longipeda1	0.260	0.113	0.119	0.321	0.317	0.128	0.110	0.325	0.148		*	0.485	*	0.258	*	*	*	0.612	0.457	*	0.671	0.872	*	*
longipeda2ª	0.273	0.124	0.114	0.321	0.297	0.145	0.121	0.316	0.157	0.020		*	*	*	*	*	*	*	*	*	*	*	*	*
lorioli1	0.245	0.214	0.226	0.319	0.333	0.259	0.234	0.149	0.267	0.258	0.247		*	0.465	*	*	*	0.644	0.157	*	0.622	0.727	*	*
lorioli2ª	0.249	0.239	0.234	0.334	0.351	0.244	0.217	0.148	0.269	0.255	0.247	0.017		*	*	*	*	*	*	*	*	*	*	*
megapoma	0.251	0.106	0.126	0.356	0.350	0.158	0.126	0.309	0.030	0.113	0.120	0.224	0.233		*	*	*	0.643	0.564	*	0.688	0.832	*	*
nereidina1 ^a	0.214	0.259	0.259	0.326	0.340	0.280	0.237	0.344	0.282	0.283	0.244	0.311	0.302	0.230		*	*	*	*	*	*	*	*	*
nereidina2ª	0.223	0.272	0.270	0.362	0.377	0.269	0.236	0.314	0.288	0.249	0.208	0.268	0.267	0.235	0.030		*	*	*	*	*	*	*	*
paucispina ^a	0.324	0.287	0.277	0.400	0.400	0.261	0.241	0.185	0.340	0.319	0.295	0.068	0.057	0.296	0.370	0.344		*	*	*	*	*	*	*
propinqua	0.410	0.416	0.345	0.408	0.408	0.424	0.422	0.356	0.527	0.458	0.450	0.418	0.386	0.500	0.519	0.487	0.385		0.609	*	0.541	0.720	*	*
rhabdota	0.308	0.316	0.285	0.393	0.356	0.260	0.245	0.180	0.344	0.325	0.318	0.073	0.063	0.307	0.379	0.352	0.032	0.314		*	0.709	0.773	*	*
robillardi ^a	0.220	0.223	0.191	0.287	0.249	0.190	0.185	0.269	0.247	0.220	0.204	0.227	0.207	0.225	0.284	0.289	0.272	0.351	0.271		*	*	*	*
caespitosa	0.342	0.410	0.375	0.382	0.360	0.299	0.270	0.390	0.407	0.328	0.301	0.315	0.303	0.372	0.359	0.362	0.311	0.572	0.320	0.365		0.835	*	*
trilineata	0.409	0.417	0.371	0.381	0.344	0.404	0.418	0.435	0.508	0.467	0.467	0.393	0.435	0.450	0.461	0.518	0.425	0.524	0.463	0.381	0.360		*	*
ophionereis	0.797	0.764	0.774	0.817	0.774	0.788	0.757	0.725	0.981	0.802	0.701	0.637	0.648	0.869	0.863	0.809	0.705	0.954	0.715	0.818	0.867	0.751		*
ophiarthrum	1.817	1.807	1.808	1.994	2.048	1.773	1.814	2.019	1.930	1.822	1.687	1.922	1.997	1.732	1.768	1.987	1.973	2.284	1.957	1.782	2.262	2.326	1.863	

Distances between conspecifics are shown in bold face. ^a 16S sequence only

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