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An 18S and 28S-based clock calibration for marine Proseriata (Platyhelminthes)

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1	A 18S and 28S-based clock calibration for marine Proseriata
2	(Platyhelminthes)
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25 **Abstract**

Geminate species are a powerful tool for calibrating the molecular clock in 26 marine organisms, and their adoption is mandatory for soft-bodied taxa, which 27 lack fossil records. The first attempt to calibrate the molecular clock in taxa 28 belonging to meiofaunal microturbellaria (Platyhelminthes: Proseriata) based 29 on geminate species is presented here. We used two species pairs from both 30 sides of the Isthmus of Panama: Minona gemella (Caribbean) and Minona cf 31 gemella (Pacific); Parotoplana sp. nov. 1 (Caribbean) and Parotoplana sp. nov. 32 2 (Pacific). The mutation rates per million years were estimated for both 33 geminate species pairs on two ribosomal regions, the complete nuclear small 34 subunit rDNA (18S) gene and the partial nuclear large subunit rDNA (28S) gene 35 fragment (spanning variable domains D1-D6). Similar values of mutation rates 36 per million years were found in both species pairs, ranging 0.12-0.16 % for 18S 37 and 0.49-0.52% for 28S. The values obtained were used as calibration points at 38 minimum age, in order to estimate the divergence times within the 39 phylogenetic tree of the whole dataset, and tested on three cases of trans-40 American (not-geminate) species from Pacific Panama and S-E Brazil, belonging 41 to the genera *Kata*, *Archimonocelis* and *Duplominona*. They consistently 42 showed higher divergence times (ranging 9.4-17.9 Myr) than geminate, trans-43 isthmian pairs. These results suggest potential usefulness of our molecular 44 clock calibration, for future research on phylogeography and evolution of 45 Proseriata. 46

47

Key words: Meiofauna; Isthmus of Panama; mutation rates; relaxed molecular
 clock; divergence time; calibration point.

52 **1. Introduction**

Interstitial meiofauna is among the most diverse and species-rich components 53 of marine biodiversity (Kennedy and Jacoby, 1999). Knowledge of many aspects 54 of the biology of these minute organisms is however limited, even in 55 comparatively well-studied areas (Curini-Galletti et al., 2012), and patterns of 56 spatial distribution are particularly poorly understood. Early studies pointed to 57 the existence of large, amphi-Atlantic or even cosmopolitan, distributions in 58 meiofaunal taxa (Sterrer, 1973; Westheide and Schmidt, 2003). Such large-59 scale ranges in species lacking any obvious means of dispersal is at the basis 60 of the so-called 'meiofauna paradox' (Giere, 2009). Ancient vicariance events, 61 followed by evolutionary stasis, were hypothesized to be responsible of the 62 observed patterns (Sterrer, 1973); this, however, would imply a surprisingly old 63 phylogenetic age for these species (see Sepkoski, 1998). Although recent 64 molecular surveys showed that, in many instances, these vast distributions are 65 the result of the lack of resolution of cryptic species complexes (i.a., Casu et al., 66 2009; Fontaneto et al., 2009; Jörger and Schrödl, 2013; Todaro et al., 1996), at 67 least some cases of large-scale distribution of meiofaunal taxa have been 68 supported by molecular data (see e.g., Dericke et al., 2008; Jörger et al., 2012; 69 Meyer-Wachsmuth et al., 2014; Tulchinsky et al., 2012), leaving open the 70 choice between great antiquity of lineages, or unsuspected capabilities for 71 dispersal. 72

In order to provide an adequate coverage of information, systematic,
biogeographic and phylogenetic studies should ideally be flanked by the
estimation of divergence time among clades (see i.a., Heads, 2005a; Ree and
Smith, 2008), which may allow inferences on the time-scale of speciation
processes. The modern molecular phylogenetic approach applied to date

evolutionary divergence is based on the molecular clock hypothesis (MCH) 78 (Zuckerkandl and Pauling, 1965), which assumes a relatively constant rate of 79 molecular evolution over time and across taxa (see Kimura, 1968, and 80 references therein). However, recent empirical studies have demonstrated the 81 existence of a significative variation in the rate of molecular evolution 82 (Bromham and Penny, 2003; Thomas et al., 2006), and the use of a more 83 sophisticated approach, such as a relaxed clock model, has been 84 recommended (see Lepage et al., 2007 and references therein). In order to 85 assign concrete dating, a molecular clock needs to be calibrated against 86 independent evidence (Benton and Donoghue, 2007). The most common 87 calibration of the molecular clock is achieved by using fossil records (e.g., 88 Blanton et al., 2013; Mulcahy et al., 2012; Ronquist et al., 2012a) or, when 89 possible, by means of both fossil records and paleogeographic events (e.g., 90 Heads, 2005a). 91

In the case of meiofaunal organisms, however, dating of divergence is 92 particularly difficult, as these taxa (and especially the soft-bodied component) 93 do not leave any fossil record that can be used as calibration points (e.g., 94 Blanton et al., 2013; Mulcahy et al., 2012). Therefore, the adoption of geminate 95 species - i.e. morphologically indistinguishable sister species that live in 96 allopatric conditions and occur at the opposite sides of a (datable) geographic 97 barrier (Jordan, 1908) - and the MCH constitute an almost inevitable strategy 98 (Coyne and Orr, 2004; Lessios, 2008). Geminate species represent a widely 99 cited model of allopatric speciation (Coyne and Orr, 2004; Lessios, 2008; 100 Vermeij, 1978), constituting a 'natural experiment' that can describe 101 evolutionary divergence and its causes (Lessios, 2008). Indeed, several studies 102 have demonstrated that geminate species may represent a suitable alternative 103

to fossil records as calibration points for a molecular clock, and can be used in 104 turn to estimate divergence times between related species (e.g., Lessios, 1998; 105 Knowlton and Weigt, 1998; Marko and Moran, 2009). For this purpose, one of 106 the most used geographical barriers is the Isthmus of Panama (see, e.g., Heads, 107 2005b). Geological literature suggests that complete isolation between the 108 Caribbean Sea and the Pacific Ocean occurred about 3.1-3.5 million years (Myr) 109 ago, due to the final emergence of the Isthmus (Allmon, 2001; Collins et al., 110 1996; Jackson and Budd, 1996; Knowlton and Weig, 1998). 111

This approach is however not without criticisms (see Heads, 2005b). A major 112 objection is that identification of a species pair as geminate species may be 113 biased by the taxonomic sampling available (Lessios, 2008). Furthermore, 114 establishing the time of separation between geminate species may be fraught 115 with problems, as the emergence of the isthmian landmass was a prolonged 116 geological process, and not all geminate pairs were simultaneously separated 117 by the emerging Isthmus (Knowlton, 1993; Knowlton and Weigt, 1998; Marko, 118 2002). Populations of intertidal meiofauna may be ideal candidates for the 119 120 calibration of the molecular clock, as they were more likely to maintain continuity of habitat and gene flow across the emerging isthmus, until the 121 separation of eastern Pacific and Caribbean was completed. However, they 122 have never been studied in this context. 123

We aimed to assess the molecular clock on species belonging to different
families of meiobenthic, intertidal free-living microturbellarians
(Platyhelminthes: Proseriata), using geminate species from the Isthmus of
Panama. Representatives of Proseriata may be numerically abundant and
characterize entire soft-bottom communities (Reise, 1988; Remane, 1933). As
most meiofaunal organisms, Proseriata lack larval stages, and, combined with

the reduced mobility of adults, their potential for dispersal is limited (Curini-130 Galletti et al., 2012). Indeed, setting the molecular clock for taxa belonging to 131 the order of Proseriata, would be of particular interest, as support for 132 phylogeographic studies (Casu et al., 2011) or setting up the evolutionary time-133 frame in cases of allopatric distributions (Delogu and Curini-Galletti, 2009; Casu 134 et al., 2014; Curini-Galletti et al., 2011). Furthermore, the existence of many 135 supra-specific taxa with anti-tropical distributions (Laumer et al., 2014) could 136 be ideally interpreted with information on the timing of their divergence. 137 Notwithstanding the current, different ecological conditions at the two sides of 138 the isthmus (Lessios, 2008), morphologically similar congeneric species were 139 found, which could be tested as potential geminate species. We sequenced 140 further morphologically similar congeners, allopatrically distributed along the 141 Pacific shores of Panama and in western Atlantic (South Brazil). These latter, 142 trans-American species were used as potential test cases, as their age of 143 divergence should pre-date that of trans-isthmian geminates. 144 We calibrated the molecular clock on two ribosomal genes, the complete 145 146 nuclear small subunit rDNA (18S) gene and the partial nuclear large subunit

rDNA (28S) gene fragment (spanning variable domains D1-D6), since their

sequences constitute the only large database available for Proseriata.

150 **2. Materials and methods**

151

152 **2.1.** The species

153 2.1.1. Trans-isthmian species pairs

- *Minona gemella* Ax and Sopott-Ehlers, 1985 (Caribbean) / *Minona* cf gemella
 (Pacific) (Proseriata: Monocelididae).

Species found at the opposite ends of the Canal (Table 1; Fig. 1), in intertidal 156 habitats. Minona gemella (type locality: Bermuda) is characterized by the 157 presence of two accessory prostatoid organs, one anterior and one posterior to 158 the copulatory organ - a unique feature for species of the genus Minona Marcus 159 1946 (Ax and Sopott-Ehlers, 1985). The Pacific counterpart appears identical in 160 morphology, as reconstructed from observations on living, semi-squeezed 161 specimens, and from histological sections, as well as for all measurable 162 characters of the sclerotized structures. The only appreciable differences have 163 been detected in their karyotypes: Caribbean specimens from Panama and 164 Puerto Rico have chromosome II metacentric, while it is more heterobrachial in 165 166 Pacific specimens (Curini-Galletti, 1991; unpubl. data). - Genus Parotoplana Meixner, 1938 (Proseriata: Otoplanidae). 167 Parotoplana sp. nov. 1 (Caribbean coast of Panama) and Parotoplana sp. nov. 2 168 (Pacific coast of Panama) are morphologically very similar, differing for minute 169 details of the sclerotized structures (unpubl. data), only detectable on strongly 170

squeezed, karyological slides, where tissues have been macerated with acetic

acid (see Curini-Galletti et al., 1989). Both species occur intertidally.

173

174 2.1.2 Trans-American species

- Genus *Kata* Marcus, 1950 (Proseriata: Otoplanidae).

The four described species of the genus *Kata* are distributed on both American 176 coasts: Kata evelinae Marcus, 1949 and Kata leroda Marcus, 1950 from South 177 Brazil (both of which were here sequenced); Kata galapagoensis Ax and Ax, 178 1974 from Galapagos Island; and Kata galea Ax and Sopott-Ehlers, 1987 from 179 Bermuda. The two new species from the Pacific coast of Panama (Kata sp. nov. 180 1 and Kata sp. nov. 2) differ from each other and the other species of the 181 genus for details of the morphology of the sclerotized pieces of the copulatory 182 organ (unpubl. data). All species occur intertidally. 183

- Genus *Duplominona* Karling, 1966 (Proseriata: Monocelididae).

The two species tested share a unique feature of the posterior end, deeply split into a 'trident' shape. *Duplominona tridens* Marcus, 1954 is a south Brazilian species (Marcus, 1954a). The Pacific counterpart (*Duplominona* sp. nov. 1) is identical in external morphology and general topography of organs, but differs for characters of the sclerotized structures of the copulatory organ (Curini-Galletti, 2014). Both species occur intertidally.

- Genus Archimonocelis Meixner, 1938 (Proseriata: Archimonocelididae).

192 The American species here sequenced, Archimonocelis marci Curini-Galletti,

193 2014 and Archimonocelis sp. nov. 1 from Brazil, and Archimonocelis sp. nov. 2,

from Pacific Coast of Panama are morphologically similar, as they share a
simple structure of the copulatory organ, with a stylet surrounded by a girdle of
short, nearly identical spines (Curini-Galletti, 2014; unpubl data). All species
occur in shallow subtidal habitats.

198

199 2.2. Sampling, DNA extraction, amplification and sequencing

200 Samples were collected manually by scooping up the superficial layer of

201 sediment. All necessary permits for samplings in protected areas were

obtained. No specific permits were required for other sites, which were not
 privately owned or protected.

Extraction of the animals from the sediment was accomplished using MgCl₂ decantation (Martens, 1984). Each specimen was studied alive by slight squeezing under the cover slip. Whenever possible, vouchers were prepared, consisting of whole mounts of posterior body regions of the specimens sequenced, and are maintained in the collections of the Zoological Museum of the University of Sassari (CZM). For information about sampling localities see Table 1 and Fig. 1.

Genomic DNA was extracted using the Macherey-Nagel NucleoSpin Tissue 211 (Macherey-Nagel GmbH and Co. KG) according to the supplier's instructions. 212 After extraction, DNA was stored as a solution at 4 °C. Complete 18S and 213 partial 28S (D1-D6) sequences were analyzed for a total of 92 individuals; 62 214 were newly obtained specifically for this study, and 30 taken from GenBank (for 215 details see Table 1). The dataset was built with 40 sequences of individuals 216 belonging to the family Otoplanidae (15 of which newly sequenced, Table 1), 217 42 to the family Monocelididae (39 of which newly sequenced, Table 1), 6 to 218 the family Archimonocelididae (all of which newly sequenced, Table 1), one to 219 the family Calviridae (from GenBank), one to the family Coeloginoporidae 220 (newly sequenced), and two to the suborder Unguiphora (one of which newly 221 sequenced). Amplifications for 18S and 28S D1-D6 regions were carried out 222 using the following primers: 18S: A (forward) GCG AAT GGC TCA TTA AAT CAG, 223 and B (reverse) CTT GTT ACG ACT TTT ACT TCC (Littlewood and Olson, 2001); 224 28S: LSU5 (forward) TAG GTC GAC CCG CTG AAY TTA AGC A, and LSUD6-3 225 (reverse) GGA ACC CTT CTC CAC TTC AGT C (Littlewood et al., 2000). 226

PCRs were carried out in a total volume of 25 μ l containing about 25 ng (5 ng/ μ l) 227 of total genomic DNA on average, 1.0 U of Tag DNA Polymerase (EuroTag by 228 Euroclone), $1 \times$ reaction buffer, 3.5 mM of MgCl₂, 0.32 μ M of each primer, and 229 200 µM of each dNTP. PCR amplifications were performed in a MJ PTC 200 230 Thermal Cycler (Biorad) programmed as follows: 1 cycle of 2 min at 94° C, 35 231 cycles of 1 min at 94° C, 1 min at 54° C (18S / 28S primers' annealing 232 temperature), and 1 min and 30 s at 72° C. At the end, a post-treatment for 5 233 min at 72° C and a final cooling at 4° C were carried out. Both positive and 234 negative controls were used to test the effectiveness of the PCR reagents, and 235 the absence of possible contaminations. Electrophoreses were carried out on 236 2% agarose gels, prepared using $1 \times$ SBA buffer (sodium boric acid, pH 8.2) and 237 stained with a 1 μ l/20 ml ethidium bromide solution. PCR products were 238 purified by ExoSAP-IT (USB Corporation) and sequenced for both forward and 239 reverse 18S and 28S strands, using an external sequencing core service 240 (Macrogen Inc., Europe). The sequencing runs were repeated twice in order to 241 verify the reliability of results. 242

243

244 2.3. Estimates of genetic distance and phylogenetic analysis

The 18S and 28S sequences were aligned separately using the algorithm Q-245 INS-I, implemented in Mafft 6.903 (Katoh and Toh, 2008), which is appropriate 246 for non-coding RNA as it considers RNA secondary structure. The best 247 probabilistic model of sequence evolution was determined after evaluation by 248 jModeltest 2.1.1 (Posada, 2008), with a maximum likelihood optimized search, 249 using the Akaike Information Criterion (AIC) and the Bayesian Information 250 Criterion (BIC). Both criterions selected the GTR+I+G (Tavaré, 1986) as the 251 best fitting model for both 18S and 28S datasets. The pairwise genetic 252

distances corrected according to the Kimura two-parameter model (*K2P*) (Kimura, 1980) were estimated between population's representatives to the trans-isthmian geminate species pairs, by means of the software Mega 6.06 (Tamura et al., 2011) with 1,000 bootstrap replications. *K2P* distances were estimated singularly for each gene in order to insert them into the formula proposed by Li and Graur (1991) (see section 2.4.1).

Phylogenetic relationships among individuals and species were investigated 259 using both Maximum Likelihood (ML) and Bayesian Inference (BI) on the 260 combined 18S and 28S sequences. We set as outgroup for the analyses the 261 species Polystyliphora novaehollandiae Curini-Galletti, 1998. ML was performed 262 using the genetic algorithm implemented in Garli 2.01 (Zwickl, 2006). In order 263 to find the best tree, the configuration file for partitioned models was set up to 264 perform 10 replicate searches (searchreps = 10). Model parameters: 265 ratematrix = $(0\ 1\ 2\ 3\ 4\ 5)$, statefrequencies = estimated, ratehetmodel = 266 gamma, numratecats = 4, corresponding to the evolution model calculated by 267 Modeltest, were used. In order to allow independent estimates of the 268 parameters for each gene, the option link was set to 0. The parameter 269 modweight was set to 0.0015, as we have two partitions. Finally, node support 270 was assessed by 1,000 bootstraps (bootstrapreps = 1000). Consensus tree was 271 computed using TreeAnnotator 1.7.4 (Drummond and Rambaut, 2007) and 272 visualised by FigTree 1.4.0 (http://tree.bio.ed.ac.uk/software/figtree/). 273 BI was carried out using the software MrBayes 3.2.2 (Ronguist et al., 2012b), 274 specifying a partitioned model and setting as model parameters: NST = 6, 275 rates = invgamma, ngammacat = 4. We allowed each partition to have its own 276 set of parameters and a potentially different overall evolutionary rate. Two 277 independent runs, each consisting of four Metropolis-coupled MCMC chains 278

(one cold and three heated chains), were run simultaneously for 5,000,000
generations, sampling trees every 1,000 generations. The first 25% of the
10,000 sampled trees was discarded as burnin.

In order to assess the convergence of chains we checked that the Average
Standard Deviation of Split Frequencies (ASDSF), approached 0 (Ronquist et al.,
2012b), and the Potential Scale Reduction Factor (PSRF) was around 1 (Gelman
and Rubin, 1992). Nodes with a percentage of posterior probability lower than
95% are considered not highly supported. Phylogenetic tree was visualized
using FigTree 1.4.0 (http://tree.bio.ed.ac.uk/software/figtree/).

288

289 2.4. Calibration of molecular clock

290 2.4.1. Estimation of the mutation rates per million years

After phylogenetic analysis showed that the trans-isthmian species (Minona 291 gemella + Minona cf gemella and Parotoplana sp. nov. 1 + Parotoplana sp. nov. 292 2) represent two pairs of sister species, and may thus be considered as 293 geminate species (see section 3.1 below and Fig. 2), the mutation rates per 294 295 million years (r) between species from both sides of the isthmus were estimated for each gene. We used the formula r = K (K2P genetic distance) / 2T 296 (time of divergence multiplied by 2 to account for the age of each lineage) (Li 297 and Graur, 1991). The obtained mutation rates per million years (two for each 298 species pair) were used for calibrating the timetree, in order to estimate the 299 divergence time throughout the whole dataset. 300

301

302 2.4.2. Estimation of divergence time

303 The software package Beast 1.7.4 (Drummond and Rambaut, 2007) was used

304 to estimate the divergence time for all of the clades evidenced by the

phylogenetic tree. Site parameters (Substitution Model = GTR; Bases 305 Frequencies = Estimated; Site Heterogeneity Model = Gamma + Invariant Sites; 306 Number of Gamma Categories = 4) have been set according to the best-fitting 307 evolution model selected by jModeltest. For the molecular clock rate variation 308 model, the lognormal uncorrelated relaxed clock was chosen because it 309 assumes independent rates on different branches. Moreover, the use of the 310 lognormal uncorrelated relaxed clock model gives an indication of how clock-311 like data is (measured by the ucld.stdev parameter). If the ucld.stdev 312 parameter estimate is close to 0, then the data is guite clock-like, while if it has 313 an estimated value much greater than 1, then data exhibits very substantial 314 rate heterogeneity among lineages. For the tree prior the Yule prior process to 315 the speciation model was applied. The priors for model parameters and 316 statistics have been set for calibrating the timetree assuming the mutation 317 rates per million years estimated separately for each of the two regions (18S 318 and 28S) on the Minona and Parotoplana species pairs. Divergence times were 319 estimated using a uniform distribution with lower and upper values set 320 321 according to the mutation rate per million years of the two species pairs (see Table 2). Operator parameters have been set following the instructions on the 322 user manual. In order to obtain the Effective Sample Size (ESS) greater than 323 200 for all of the statistic parameters, a run of 400,000,000 generations was 324 performed, sampling a tree every 40,000 generations. 325

We used Tracer 1.6 (Rambaut and Drummond, 2009) for viewing the resulting log file, in order to ensure convergence of parameter values, to verify whether ESS values exceeded 200, and to estimate node ages. TreeAnnotator and FigTree were used for drawing and visualizing the timetree, respectively. Alignments and Bayesian tree-files are deposited and available in Treebase **(TB2: S16487.**

332 **3. Results**

333 3.1. Estimates of genetic distance and phylogenetic analysis

After the alignment, sequences of 1632 *bp* and 1650 *bp* were obtained for the

18S and 28S regions, respectively (see Table 1 for the GenBank accession

numbers). For each region, the genetic pairwise distance corrected according

to the K2P model provided comparable values between the pairs Minona

338 *gemella* + *Minona* cf *gemella* and *Parotoplana* sp. nov. 1 + *Parotoplana* sp. nov.

339 2: $K2P = 0.0085 \pm 0.0022$ and $K2P = 0.0115 \pm 0.0027$, for the 18S; and K2P =

 0.0345 ± 0.0045 and $K2P = 0.0361 \pm 0.0049$, for the 28S D1-D6, respectively

341 (Table 2).

ML and BI generated consistent trees with negligible differences in topology; 342 additionally, in both trees the nodes of our interest are highly supported. We 343 therefore reported the BI tree obtained by the software MrBayes only (Fig. 2). 344 This phylogenetic tree shows that *M. gemella* (Atlantic coast) and *Minona* cf 345 gemella (Pacific coast) (GS1) are in a sister-taxon relationship, as well as 346 Parotoplana sp. nov. 1 (Atlantic coast) and Parotoplana sp. nov. 2 (Pacific coast) 347 (GS2); the corresponding nodes are highly supported both for posterior 348 probability and bootstrap values (Fig. 2). Therefore, according to Jordan's 349 definition (1908), they can be considered as geminate species, and they will be 350 used for estimating the mutation rate per million years. 351

Furthermore, the tree confirmed the sister-taxa relationship between Atlantic and Pacific clusters of species belonging to the families Otoplanidae (node A), Monocelididae (node B) and Archimonocelididae (node C) (Fig. 2). In particular: - Within Otoplanidae, species belonging to the genus *Kata* were separated into two geographic clusters, one grouping the Atlantic *K. evelinae* + *K. leroda*, and one the Pacific *Kata* sp. nov. 1 + *Kata* sp. nov. 2 (node A in Fig. 2); Within Monocelididae, node B (Fig. 2) splits the Atlantic specimens of *Duplominona tridens* from the Pacific specimens of *Duplominona* sp. nov. 1;
Within Archimonocelidae, node C (Fig. 2) splits the Atlantic *A. marci* + *Archimonocelis* sp. nov. 1 from the Pacific *Archimonocelis* sp. nov. 2. For each
of these three cases, nodes are highly supported (Fig. 2).

363

364 3.2. Mutation rates per million years and divergence time

The estimated ucld.stdev parameter amounts to 0.842 and 0.677 for the 18S 365 and 28S, respectively, indicating that our dataset is clock-like. The mutation 366 rate per million years between *M. gemella* and *Minona* cf gemella amounts to 367 0.12% for the 18S, and 0.49% for the 28S (Table 2). Slight higher values were 368 obtained between Parotoplana sp. nov. 1 and Parotoplana sp. nov. 2: 0.16% for 369 the18S, and 0.52% for the 28S (Table 2). Analysis performed by means of the 370 software Beast produced a tree whose topology is consistent to those obtained 371 by both Garli and MrBayes. On these bases, we estimated a divergence time 372 for node A (splitting Atlantic K. evelinae and K. leroda from the Pacific Kata sp. 373 nov. 1 and Kata sp. nov. 2) of about 17.9 Myr, ranging 12.9 - 23.8 Myr (Fig. 3); 374 for node B (splitting Atlantic Duplominona tridens from the Pacific Duplominona 375 sp. nov. 1) of about 13.9 Myr, ranging 8.9 - 20.0 Myr (Fig. 3); and for node C 376 (splitting the Atlantic A. marci and Archimonocelis sp. nov. 1, from the Pacific 377 Archimonocelis sp. nov. 2) of 9.4 Myr, ranging 5.5 - 14.9 Myr (Fig. 3). 378

380 **4. Discussion**

Molecular tools and the MCH have provided new clues on past evolutionary 381 processes and mechanisms driving molecular evolution (Bromham and Penny, 382 2003). However, several authors have shown perplexity about the wide 383 applicability of the MCH (see e.g., Heads, 2005b; Lessios, 2008; Palumbi, 1997), 384 385 and the use of the molecular clock to infer divergence time elicits criticisms, mostly concerning the way the clock is calibrated (Peterson et al., 2004). The 386 use of paleogeographic events, which represents the only possible alternative 387 of calibration in absence of fossil records, is a contentious issue (Coyne and Orr, 388 2004; Lessios, 2008). It should be noticed that estimates on geminate species 389 assume the final closure of a given geographic barrier as a minimum age 390 calibration; hence, time since divergence may have been underestimated if 391 taxa diverged before this date. In addition, considering merely geminate 392 species as species pairs originated after the rise of a geographic barrier could 393 be an oversimplification of their evolutionary path, since the evolutionary 394 history of many nominal geminate species potentially may be more complex 395 (see Knowlton and Weigt, 1998). 396

To overcome such limitations, the use of different genes or loci, and different 397 calibration points has been recommended (Marko and Moran, 2009). In the 398 case of Proseriata, the number of genes/loci we could use are limited, because 399 most of 'universal' primers for invertebrates, such as those for the cytochrome 400 c oxidase subunit I (COI) Folmer's region (Folmer et al., 1994) do not provide 401 satisfactory results, and specific primers are not available except for a few 402 species (see Casu et al., 2011; Sanna et al., 2009). Furthermore, since a limited 403 number of sequences of Proseriata is at present available in Genbank, the 404 number of calibration points depends on sampling's success, and the adequacy 405

of the sampling campaign can be assessed only after morphological and 406 molecular analyses in laboratory (see e.g., Casu et al., 2014). In this context, it 407 is noteworthy that an inadequate taxonomic coverage may lead to the use of 408 false geminate species for the calibration of the molecular clock, and thus to 409 the use of species pairs separated well before the last closure of the isthmus 410 which results in an overestimation of the mutation rates per million years 411 (Heads, 2005a; Knowlton and Weigt, 1998). Consequently, the use of an higher 412 rate may cause an underestimation of the divergence time among groups in 413 the timetree. 414

Albeit it might be questionable whether our taxonomic coverage is extensive 415 enough to assess sister species relationships reliably, the Atlantic M. gemella 416 and Parotoplana sp. nov. 1, and their Pacific counterparts (Minona cf gemella 417 and Parotoplana sp. nov. 2, respectively) are reciprocally monophyletic and 418 morphologically indistinguishable at the routine level of morphological 419 observation, and are thus highly suggestive of geminate lineages. Furthermore, 420 the two pairs show very similar values of mutation rate per million years in 421 both genes. It is noteworthy that these similar values have been found in 422 species pairs belonging to two different families (Monocelididae and 423 Otoplanidae), and may thus prove applicable across the Proseriata. Finally, in 424 the three trans-American species used as test cases (Kata spp., Duplominona 425 spp. and Archimonocelis spp.), the obtained divergence times are greater than 426 the final closure of the Isthmus of Panama - ranging from 9.4 Myr (time of 427 divergence between Archimonocelis spp.) and about 17.9 Myr (time of 428 divergence between *Kata* spp.) - and therefore not conflicting with the values 429 obtained with trans-isthmian species. 430

Our results are consistent to those found for other trans-American species pairs 431 (see e.g., Beu, 2001; Coates and Obando, 1996; Collins, 1996; Jackson et al., 432 1993; Roopnarine, 2001; Vermeij, 2001). For instance, the calibration on COI 433 and ITS (Internal Transcribed Spacer) sequences revealed a time of divergence 434 of 17.4-27.0 Myr, and 14.5-18.8 Myr, respectively, between trans-American 435 populations of the subgenus Acar (Bivalvia) (Marko and Moran, 2009). 436 In the cases of *Kata* spp, *Duplominona* spp., and *Archimonocelis* spp., dispersal 437 between ocean basins along the southern tip of South America would obfuscate 438 interpretation of our results. However, no member of the species pairs involved 439 was found in previous research in Chile, Uruguay, Terra del Fuego, or sub-440 Antarctic islands (Marcus, 1954b; Schockaert et al., 2009, 2011). Furthermore, 441 species of the genus Kata are only known from tropics; similarly, Duplominona 442 and Archimonocelis species, with few exceptions, occur in tropical to warm-443 temperate areas (Martens and Curini-Galletti, 1993; Tyler et al., 2006, 2012). 444 At least in recent times, therefore, the rigid conditions of extreme south of 445 South America acted as barrier to dispersal of these organisms. 446

447

448 *4.1.Conclusions*

The study of geminate species of Proseriata across the Isthmus of Panama
allowed the first calibration of the molecular clock for a meiofaunal taxon.
Results of our research open potentials for the use of intertidal meiofauna for
MCH. Among the major objections of the MCH, in fact, is that speciation among
geminate pairs may predate the final emergence of the isthmus. However,
meiofaunal, intertidal/shallow-water taxa may have shown continuity of habitat
until final emergence of the barrier, and their divergence may indeed reflect

the final stage of the isthmian formation. A similar suggestion was advanced
for species from brackish-water and mangrove habitats (see Miura et al., 2010).
Although further tests on a larger dataset and on other test-cases are deemed
necessary, data obtained (both mutation rates and divergence times) might
prove invaluable to provide further insights into the phylogenetic relationships
and evolution of Proseriata.

462

463

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702

704 FIGURE AND TABLE LEGENDS

705

706 Figure 1. Sampling localities.

Map of the four trans-American and geminate species sampling localities:
Bocas del Toro, Panama; Playa La Angosta - Colón, Panama; Naos Island Panama City, Panama; Ilhabela/São Sebastião, Brazil.

710

711 Figure 2. Phylogenetic tree.

Tree obtained by BI showing the interrelationships of the species based on 712 combined 18S+28S D1-D6. The branch length scale refers to the number of 713 substitutions per site. Nodal supports are indicated for BI as posterior 714 probability (PP). For the three test cases and the two geminate species pairs ML 715 bootstrap values are also reported at each node. 716 GS1: geminate species 1 (Minona gemella - Atlantic coast + Minona cf gemella 717 - Pacific coast). GS2: geminate species 2 (Parotoplana sp. nov. 1 - Atlantic coast 718 + Parotoplana sp. nov. 2 - Pacific coast). Node A: Atlantic Kata evelinae and 719 720 Kata leroda + Pacific Kata sp. nov. 1 and Kata sp. nov. 2. Node B: Atlantic

721 Duplominona tridens + Pacific Duplominona sp. nov. 1. Node 3: Atlantic

Archimonocelis marci and Archimonocelis sp. nov. 1 + Pacific Archimonocelis
sp. nov. 2.

724

725 Figure 3. Timetree.

Tree obtained by the software Beast showed divergence time among taxa.

Nodes indicated with A, B and C correspond to the nodes showed in Fig. 2.

728 Values within brackets represent the median values of divergence time of the

node. Only for the three test cases (nodes A, B and C) within brackets are

showed both median values and the range (underlined) of divergence time ofthe node.

732

733

Table 1. List of species sampled and sequences used for this study. Accession
 numbers refer to GenBank codes; accession numbers of new sequences are in
 italic.

737

738 **Table 2.**

- 18S and 28S mutation rates for the two geminate species pairs. *K2P*: genetic
- 740 distance corrected according to the Kimura two-parameters model (Kimura,
- 1980) and standard error; *r*: mutation rates per million years.

Table 1.

Family	Species	Locality*	185	28S D1- D6
	Archotoplana holotricha Ax, 1956	GenBank	<u>AJ24367</u> <u>6</u>	<u>AJ270165</u>
	<i>Xenotoplana acus</i> Ax, Weidemann and Ehlers, 1978	GenBank	<u>AJ27015</u> <u>5</u>	<u>AJ270181</u>
	Parotoplana ambrosolii Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>43</u>	<u>KC971066</u>
	<i>Parotoplana ambrosolii</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>44</u>	<u>KC971067</u>
	<i>Parotoplana ambrosolii</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>45</u>	<u>KC971068</u>
	<i>Parotoplana tubifera</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>46</u>	<u>KC971069</u>
	Parotoplana tubifera Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>47</u>	<u>KC971070</u>
	Parotoplana tubifera Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>49</u>	<u>KC971072</u>
	<i>Parotoplana tubifera</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>58</u>	<u>KC971081</u>
Otoplanidae	<i>Parotoplana impastatoi</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>48</u>	<u>KC971071</u>
	<i>Parotoplana impastatoi</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>50</u>	<u>KC971073</u>
	Parotoplana ambrosolii Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>56</u>	<u>KC971079</u>
	<i>Parotoplana ambrosolii</i> Curini-Galletti and Delogu, 2014	GenBank	<u>KC9710</u> <u>57</u>	<u>KC971080</u>
	Parotoplana spathifera Delogu and Curini-Galletti, 2007	GenBank	<u>KC9710</u> <u>53</u>	<u>KC971076</u>
	Parotoplana pythagorae Delogu and Curini- Galletti, 2007	GenBank	<u>KC9710</u> <u>52</u>	<u>KC971075</u>
	Parotoplana renatae Ax, 1956	GenBank	<u>AJ01251</u> <u>7</u>	<u>AJ270176</u>
	Parotoplana renatae Ax, 1956	GenBank	<u>KC9710</u> <u>62</u>	<u>KC971085</u>
	Parotoplana multispinosa Ax, 1956	GenBank	<u>KC9710</u> <u>61</u>	<u>KC971084</u>
	Parotoplana primitiva Ax, 1956	GenBank	<u>KC9710</u> <u>60</u>	<u>KC971083</u>

	-			
	Parotoplana bicupa Sopott-Ehlers, 1976	GenBank	<u>KC9710</u> <u>63</u>	KC971086
	Parotoplanella progermaria Ax, 1956	GenBank	<u>KC9710</u> <u>59</u>	<u>KC971082</u>
	Parotoplana crassispina Delogu and Curini-Galletti, 2009	GenBank	<u>KC9710</u> <u>51</u>	<u>KC971074</u>
	Parotoplana rosignana Lanfranchi and Melai, 2008	GenBank	<u>KC9710</u> <u>54</u>	<u>KC971077</u>
	Parotoplana procerostyla Ax,1956	GenBank	<u>KC9710</u> <u>55</u>	<u>KC971078</u>
	<i>Monostichoplana filum</i> (Meixner,1938)	GenBank	<u>AJ27015</u> <u>8</u>	<u>AJ270173</u>
	Archotoplana holotricha Ax, 1956	Faro (Portugal)	<u>KJ68232</u> <u>2</u>	<u>KJ682384</u>
	Kata evelinae Marcus, 1949	Ilhabela (Brazil)	<u>KJ68232</u> <u>3</u>	<u>KJ682385</u>
	Kata evelinae Marcus, 1949	Ilhabela, (Brazil)	<u>KJ68232</u> <u>4</u>	<u>KJ682386</u>
	Kata leroda Marcus, 1950	São Sebastião (Brazil)	<u>KJ68232</u> <u>5</u>	<u>KJ682387</u>
	Kata leroda Marcus, 1950	São Sebastião, (Brazil)	<u>KJ68232</u> <u>6</u>	<u>KJ682388</u>
	Kata sp. nov. 1	Naos Island (Panama)	<u>KJ68232</u> <u>7</u>	<u>KJ682389</u>
	Kata sp. nov. 1	Naos Island (Panama)	<u>KJ68232</u> <u>8</u>	<u>KJ682390</u>
	Kata sp. nov. 2	Naos Island (Panama)	<u>KJ68232</u> <u>9</u>	<u>KJ682391</u>
	Parotoplana sp.nov. 1	Bocas del Toro (Panama)	<u>KJ68233</u> <u>0</u>	<u>KJ682392</u>
	Parotoplana sp.nov. 1	Bocas del Toro (Panama)	<u>KJ68233</u> <u>1</u>	<u>KJ682393</u>
	Parotoplana sp.nov. 1	Bocas del Toro (Panama)	<u>KJ68233</u> <u>2</u>	<u>KJ682394</u>
Otoplayidae	Parotoplana sp. nov. 2	Naos Island (Panama)	<u>KJ68233</u> <u>3</u>	<u>KJ682395</u>
Otopianidae	Parotoplana sp. nov. 2	Naos Island (Panama)	<u>KJ68233</u> <u>4</u>	<u>KJ682396</u>
	Parotoplana sp. nov. 2	Naos Island (Panama)	<u>KJ68233</u> <u>5</u>	<u>KJ682397</u>
	Parotoplana primitiva Ax, 1956	Roscoff (France)	<u>KJ68233</u> <u>6</u>	<u>KJ682398</u>

	Archimonocelis marci Curini-Galletti, 2014	São Sebastião (Brazil)	<u>KJ68233</u> Z	<u>KJ682399</u>
	Archimonocelis marci Curini-Galletti, 2014	São Sebastião (Brazil)	<u>KJ68233</u> <u>8</u>	<u>KJ682400</u>
Archimonoceli	Archimonocelis sp. nov. 1	São Sebastião (Brazil)	<u>KJ68233</u> <u>9</u>	<u>KJ682401</u>
didae	Archimonocelis sp. nov. 2	Naos Island (Panama)	<u>KJ68234</u> <u>0</u>	<u>KJ682402</u>
	Archimonocelis sp. nov. 2	Naos Island (Panama)	<u>KJ68234</u> <u>1</u>	<u>KJ682403</u>
	Archimonocelis sp. nov. 2	Naos Island (Panama)	<u>KJ68234</u> <u>2</u>	<u>KJ682404</u>
	<i>Minona ileanae</i> Curini-Galletti, 1997	GenBank	<u>JN22490</u> <u>5</u>	<u>JN224910</u>
	Monocelis longiceps (Duges, 1830)	GenBank	<u>KC9710</u> <u>64</u>	<u>KC971087</u>
	<i>Monocelis longistyla</i> Martens and Curini-Galletti, 1987	GenBank	<u>KC9710</u> <u>65</u>	<u>KC971088</u>
	<i>Minona ileanae</i> Curini-Galletti, 1997	Great Bitter Lake (Egypt)	<u>KJ68234</u> <u>3</u>	<u>KJ682405</u>
	<i>Minona</i> sp. nov.	Playa La Angosta, Colón (Panama)	<u>KJ68234</u> <u>4</u>	<u>KJ682406</u>
	<i>Minona</i> sp. nov.	Playa La Angosta, Colón (Panama)	<u>KJ68234</u> <u>5</u>	<u>KJ682407</u>
	Minona cf trigonopora Ax, 1956	Palau (Sardinia, Italy)	<u>KJ68234</u> <u>6</u>	<u>KJ682408</u>
Monocelididae	Minona gemella Ax and Sopott-Ehlers, 1985	Playa La Angosta, Colón (Panama)	<u>KJ68234</u> <u>7</u>	<u>KJ682409</u>
	Minona gemella Ax and Sopott-Ehlers, 1985	Playa La Angosta, Colón (Panama)	<u>KJ68234</u> <u>8</u>	<u>KJ682410</u>
	Minona gemella Ax and Sopott-Ehlers, 1985	Playa La Angosta, Colón (Panama)	<u>KJ68234</u> <u>9</u>	<u>KJ682411</u>
	Minona cf gemella Ax and Sopott-Ehlers, 1985	Naos Island (Panama)	<u>KJ68235</u> <u>0</u>	<u>KJ682412</u>
	Minona cf gemella Ax and Sopott-Ehlers, 1985	Naos Island (Panama)	<u>KJ68235</u> <u>1</u>	<u>KJ682413</u>
	Minona cf gemella Ax and Sopott-Ehlers, 1985	Naos Island (Panama)	<u>KJ68235</u> <u>2</u>	<u>KJ682414</u>
	<i>Minona</i> sp. nov.	Boa Vista Island (Cape Verde)	<u>KJ68235</u> <u>3</u>	<u>KJ682415</u>
	<i>Minona</i> sp. nov.	Boa Vista Island (Cape Verde)	<u>KJ68235</u> <u>4</u>	<u>KJ682416</u>

	<i>Monocelis lineata</i> OF Müller, 1774	Porto Pozzo (Sardinia, Italy)	<u>KJ68235</u> <u>5</u>	KJ682417
	<i>Monocelis lineata</i> OF Müller, 1774	Charaki (Rhodes, Greece)	<u>KJ68235</u> <u>6</u>	<u>KJ682418</u>
	<i>Monocelis lineata</i> OF Müller, 1774	Pilo (Sardinia, Italy)	<u>KJ68235</u> <u>7</u>	<u>KJ682419</u>
	<i>Monocelis lineata</i> OF Müller, 1774	Colostrai (Sardinia, Italy)	<u>KJ68235</u> <u>8</u>	<u>KJ682420</u>
	Minona sp. nov. 1	Faro (Portugal)	<u>KJ68235</u> <u>9</u>	<u>KJ682421</u>
	Minona sp. nov. 1	Faro (Portugal)	<u>KJ68236</u> <u>0</u>	<u>KJ682422</u>
	<i>Minona</i> sp. nov.	Lanzarote, Canary Island (Spain)	<u>KJ68236</u> <u>1</u>	<u>KJ682423</u>
	<i>Minona</i> sp. nov.	Tenerife, Canary Island (Spain)	<u>KJ68236</u> <u>2</u>	<u>KJ682424</u>
	<i>Minona</i> sp. nov.	Tenerife, Canary Island (Spain)	<u>KJ68236</u> <u>3</u>	<u>KJ682425</u>
	Duplominona sp. nov.	Lanzarote, Canary Island (Spain)	<u>KJ68236</u> <u>4</u>	<u>KJ682426</u>
	Duplominona sp. nov.	Faro (Portugal)	<u>KJ68236</u> <u>5</u>	<u>KJ682427</u>
	Duplominona sp. nov.	Faro (Portugal)	<u>KJ68236</u> <u>6</u>	<u>KJ682428</u>
	Duplominona brasiliensis Curini-Galletti, 2014	Ilhabela (Brazil)	<u>KJ68236</u> <u>7</u>	<u>KJ682429</u>
	Duplominona sp. nov. 1	Naos Island (Panama)	<u>KJ68236</u> <u>8</u>	<u>KJ682430</u>
Managalididaa	Duplominona sp. nov. 1	Naos Island (Panama)	<u>KJ68236</u> <u>9</u>	<u>KJ682431</u>
Monocendidae	Duplominona sp. nov. 1	Naos Island (Panama)	<u>KJ68237</u> <u>0</u>	<u>KJ682432</u>
	Duplominona tridens (Marcus, 1954)	São Sebastião (Brazil)	<u>KJ68237</u> <u>1</u>	<u>KJ682433</u>
	Duplominona tridens (Marcus, 1954)	São Sebastião (Brazil)	<u>KJ68237</u> <u>2</u>	<u>KJ682434</u>
	Duplominona sp. nov. 2	Naos Island (Panama)	<u>KJ68237</u> <u>3</u>	<u>KJ682435</u>
	Duplominona sp. nov. 3	Naos Island (Panama)	<u>KJ68237</u> <u>4</u>	<u>KJ682436</u>
	Duplominona sp. nov. 3	Naos Island (Panama)	<u>KJ68237</u> <u>5</u>	<u>KJ682437</u>

	Duplominona sp. nov.	Roscoff (France)	<u>KJ68237</u> <u>6</u>	<u>KJ682438</u>
	Duploperaclistus circocirrus Martens, 1983	Roscoff (France)	<u>KJ68237</u> <u>7</u>	<u>KJ682439</u>
	Duploperaclistus circocirrus Martens, 1983	Roscoff (France)	<u>KJ68237</u> <u>8</u>	<u>KJ682440</u>
	Duplominona sp. nov.	Blanes (Spain)	<u>KJ68237</u> <u>9</u>	<u>KJ682441</u>
	Archilopsis spinosa (Jensen, 1878)	Roscoff (France)	<u>KJ68238</u> <u>0</u>	<u>KJ682442</u>
	<i>Archilopsis arenaria</i> Martens, Curini-Galletti & Pucinelli, 1989	Roscoff (France)	<u>KJ68238</u> <u>1</u>	<u>KJ682443</u>
Calviriidae	Calviria solaris Martens and Curini-Galletti, 1993	GenBank	<u>AJ27015</u> <u>3</u>	<u>AJ270168</u>
Coelogynopori dae	<i>Coelogynopora tenuis</i> Meixner, 1938	Roscoff (France)	<u>KJ68238</u> <u>2</u>	<u>KJ682444</u>
Unguinhors	Polystyliphora novaehollandiae Curini-Galletti, 1998	GenBank	<u>AJ27016</u> <u>1</u>	AJ270177
Unguiphora	Nematoplana coelogynoporoides Meixner, 1938	Roscoff (France)	<u>KJ68238</u> <u>3</u>	<u>KJ682445</u>

⁷⁴⁴ *For newly sequenced taxa only.

Table 2.

	Geminate species	К2Р	r (%)
	185		
	Minona gemella (Atlantic Coast)		
	Vs	0.0085 ± 0.0022	0.12
	Minona cf gemella (Pacific Coast)		
	Parotoplana sp. nov. 1 (Atlantic Coast)		
	Vs	0.0115 ± 0.0027	0.16
	Parotoplana sp. nov. 2 (Pacific Coast)		
	285		
	<i>Minona gemella</i> (Atlantic Coast)		
	Vs	0.0345 ± 0.0045	0.49
	<i>Minona</i> cf <i>gemella</i> (Pacific Coast)		
	Parotoplana sp. nov. 1 (Atlantic Coast)		
	Vs	0.0361 ± 0.0049	0.52
	Parotoplana sp. nov. 2 (Pacific Coast)		
747			
748			
- 10			
/49			
750			
751			
750			
/52			
753			
754			
766			
/ 55			
756			
757			
758			
/ 50			
759			

Figure 1



777 Figure 2



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788 Figure 3

