Molecular diet analysis of Anguilliformes leptocephalus larvae collected in the western North Pacific

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Abstract

Natural diets of leptocephalus larvae have been enigmatic. In this study, we collected DNA samples from the gut contents and body surface of leptocephali belonging to the five Anguilliform families (Anguillidae, Chlopsidae, Congridae, Muraenidae, and Serrivomeridae) from the northwest Pacific and performed next-generation 18S rDNA sequencing. Wide variety of eukaryotes was detected in both samples, from which eight eukaryotic groups (jellyfish, conoid parasite, tunicate, copepod, krill, segmented worm, fungi, and dinoflagellate) were selected on the basis of abundance. All groups except conoid parasites were common in both the samples. Cnidarian 18S rDNA reads were the most abundant in both the samples; however, the number of samples having cnidarian reads and the read counts were significantly higher in the body surface scraping samples than in the gut content samples, regardless of careful rinsing of the body surface. These results indicate that the cnidarian DNAs are most likely found because of cross contamination from the body surface and/or environment. 18S rDNA read counts of copepod and tunicate in the gut contents were greater than or comparable with those in the body surface scraping samples, which may correspond to the previous observations of fecal pellets and larvacean houses in the leptocephali gut. Thus, the present study supports previous implications that leptocephali utilize detritus materials, so called marine snow.

Introduction

Marine and freshwater eels belonging to the order Anguilliformes have peculiar transparent leaf-like larvae called leptocephalus. Natural diets of the leptocephali have been enigmatic. Earlier studies have reported a poorly differentiated gut in the leptocephali containing amorphous
and no identifiable food material [1–3]. Dissolved organic compounds were suspected to be directly absorbed through the epidermis, and it was hypothesized that the leptocephali utilized its sharp teeth to puncture other organisms and swallow the body fluid [2]. However, later histological investigation indicated that the alimentary tract of the Japanese eel (Anguilla japonica) leptocephali is functional for the uptake and digestion of intact macromolecules [4]. In captive condition, the wild-caught leptocephali of pike conger (Muraenesox cinereus) and white-spotted conger (Conger myriaster) showed no active preference toward zooplanktons, such as copepod larvae and ctenophore; however, these leptocephali visually located and actively engulfed squid paste [5]. Fecal pellets of zooplankton, aloricate protozoa, phytoplankton-like small spherical cells, and detrital aggregates have been observed in the gut of leptocephali [6–11], and the trophic positions of C. myriaster leptocephali have been observed to be nearly equal to the particulate organic matter [6].

Molecular analyses have been recently used to determine the gut contents of the leptocephali. Although 18S rDNA sequences from a wide variety of plankton organisms have been detected from the gut of the European eel (A. anguilla) leptocephali, gelatinous zooplankton (hydrozoan jellyfish) was suspected to be the major diet [12]. However, no animal ribosomal DNA (internal transcribed spacer 1) was found in the gut samples of the Japanese eel leptocephali, and already-degraded material was suspected in the diet [13]. Recently, a more advanced metagenomic analysis using next-generation sequencing (NGS) was applied, revealing that 76% of 18S rDNA reads recovered from the gut of European eel leptocephali belongs to the phylum Cnidaria [14]. However, the consumption of cnidarian jellyfish contradicts with the results of stable isotope analysis, in which the trophic positions of the leptocephali have been reported to be low [6,15–19]. An inherent problem in the previous molecular studies [12–14] is that no specimen from the body surface of the leptocephali was analyzed, which could be a major source of cross contamination.

We performed 18S rDNA-based metagenomic analysis using NGS not only for the gut content samples but also for the body surface scraping samples of the leptocephali, in which cnidarian 18S rDNAs in the body surface scraping samples predominated over those in the gut content samples.

**Materials and methods**

**Ethics statement**

Larval samples captured with plankton nets deployed from research vessels were dead on retrieval and sampled at this time, and all plankton net operations were carried out in high seas outside the Exclusive Economic Zone. Therefore, the approval of coastal states was not required under the United Nations Convention on the Law of the Sea (UNCLOS).

**Leptocephali sampling and identification**

Isaacs-Kidd Midwater Trawl (IKMT) net (8.7 m² opening, 13 m long, 0.5 mm mesh, and canvas made cod-end) was used to collect the leptocephali. Oblique tows from the depth of 200 m to the surface were performed at night in the northwest tropical and subtropical Pacific from September to October of 2017. The leptocephali were sorted after collection and placed on ice-cold petri dish. Leptocephali having gut content (GC) were visually identified, and one side of the body surface was alternately rinsed three to four times using sterilized and refrigerated artificial sea water. Sterile swab was used to scrape the body surface (body surface scraping sample: BSS) and the tip of a swab was placed in separate sterile 1.5 mL Eppendorf tube. Subsequently, GC were squeezed out using sterilized inoculating loop and pipetted into separate sterile 1.5 mL Eppendorf tube. A small piece of the dorsal muscle was dissected and placed in a separate
sterile 1.5 mL Eppendorf tube. All these tubes were kept at –60˚C and transferred to the laboratory. DNA samples of the muscle, GC and BSS were extracted using a DNA extraction kit (Genomic Prep Cell and Tissue DNA Isolation Kit, Amersham Bioscience). DNA extracted from the muscle was used for partial amplification of mitochondrial 16S rDNA and nuclear 18S rDNA segments, in which a universal primer pair (16Sar-L and 16Sbr-H) [20] was used for the former and a primer pair (18S30F: 5’–GTCTCAAGATTTAGCCCATGC–3’ and 18S580R: 5’–CACCAGACTTGCCTCAAAT–3’) for the latter. PCR amplification conditions are described previously [21] using annealing temperature of 55˚C for the former and 58˚C for the latter. Direct nucleotide sequencing for the amplified fragments was performed using the PCR primers.

Peptide nucleotide acid (PNA) directed PCR clamping

We adopted PNA directed PCR clamping to selectively inhibit amplification of the host 18S rDNA [13,22]. A PNA probe was designed to anneal to the sequence near 5’ region in the 18S rRNA gene, and the nucleotide sequence was NH₂-ACGGCCGGTACAGTGCONH₂ having 80.7˚C Tm. Versatility of a primer pair for 18S rDNA mentioned above was tested by using a wide range of eukaryotes: Japanese eel (A. japonica), Japanese pufferfish (Takifugu rubripes), Japanese pilchard (Sardinops melanostictus), broadbanded thornyhead (Sebastolobus macrourus), Pacific bluefin tuna (Thunnus orientalis), freshwater shrimp (Palaemon paucidens), pronghorn spiny lobster (Panulirus penicillatus), long-spined sea urchin (Diadema setosum), brown macroalgae (Sargassum horneri and Petalonia binghamiae), diatom (Phaeodactylum tricornutum), and dinoflagellate (Ceratoperidinium falcatum), in which amplification of an expected size of fragment (c.a. 550 bp) was observed in all species. Efficiency of PCR clamping was tested by adding 1 μL PNA (10 μM) to 25 μL of PCR reaction mixture using eukaryote samples mentioned above. Efficient clamping was observed in Japanese eel, broadbanded thornyhead, and Pacific bluefin tuna, while no apparent inhibition of amplification was observed in the other organisms.

Genetic analysis of the GC and BSS samples of the leptocephali

A two-step PCR employed to construct the paired-end libraries for MiSeq sequencing follows our previous study [23]. Adaptor-associated primers were used in the first PCR: 5_-ACA CTTTTTCTACAGGAAGCTCTCTTCCGATCT + 18S30F (forward) and 5_-GTGACGAGGAGGTCCAGCTCTTCCGATCT + 18S580R (reverse). The first PCR was performed in a reaction mixture (25 μL) containing 13.5 μL H₂O, 2.5 μL of 10 × PCR buffer, 2.5 μL dNTP (2 mM), 1.5 μL MgSO₄ (25 mM), 1 μL template DNA, 0.5 U KOD-Plus-ver. 2 (TOYOBO, Osaka, Japan), 1.25 μL of each primer (10 μM), and 1 μL PNA (10 μM). The reaction mixture was pre-heated at 94˚C for 3 min, followed by 30 to 40 amplification cycles (denaturation at 94˚C for 15 s, annealing at 56˚C for 30 s and extension at 68˚C for 40 s). PCR amplification was checked using 1.5% agarose gel electrophoresis. The PCR products purified using an Agencourt MPure XP (BECKMAN COULTER, Life Sciences, Brea, California, USA) were eluted in 25 μL of TE buffer following the manufacturer protocol. The second-round PCR used the first PCR products as a template and amplified the region using primers 5’–AATGATACGGCGACCAGAGATCTACAC–8 bp index –AATGATACGGCGACCAGGAG (forward) and 5’–CAAGCA GAAAGCCGATAGGAGAT–8 bp index –GTGACTGGAGTGTAAGCACT (reverse). The eight base segments represent dual-index sequences used to recognize each sample; the 5’ end-sequences are adapters that allow the final product to bind or hybridize to short oligonucleotides on the surface of the Illumina flow cell; and the 3’ end-sequences are priming sites for the MiSeq sequencing. After purification, the first PCR product was diluted 10 times using Milli-
Q water and used as a template for the second PCR. The second PCR was carried out in the same way as the first round of PCR, except the volume of the reaction mixture was 50 μL with the addition of 2.0 μL of the diluted PCR product. The PCR cycling conditions were as follows: initial denaturation at 94˚C for 3 min, followed by 10–12 cycles at 94˚C for 15 s, 5˚C for 30 s, and 68˚C for 40 s. PCR amplification was again verified checked using agarose gel electrophoresis, and the PCR products were purified using an Agencourt AMPure XP (BECKMAN COULTER, USA). The amplified PCR products were quantified and the indexed second PCR products were pooled in equal concentrations and stored at ~30˚C until use for sequencing.

A PhiX DNA spike-in control was mixed with the pooled DNA library to improve the data quality of low diversity samples, such as single PCR amplicons. DNA concentrations of the pooled library and the PhiX DNA were adjusted to 4 nM using the buffer EB (10 mM Tris-HCl pH 8.5) mixed at a ratio of 7:3.5 μL. The 4 nM library was denatured with 5 μL of fresh 0.1 N NaOH. Using the HT1 buffer (provided by the Illumina MiSeq v. 2 Reagent kit for 2 × 150 bp PE), the denatured library (10 μL; 2 nM) was diluted to a final concentration of 12 pM for sequencing on the MiSeq platform.

**MPSS data treatment processes and operational taxonomic unit picking**

Nucleotide sequences were demultiplexed based on the 5'-multiplex identifier (MID) tag and primer sequences using the default format in MiSeq. The sequences containing palindrome clips longer than 30 bp and homopolymer longer than 9 bp were trimmed from the sequences at both ends. The 3' tails with an average quality score of less than 30 at the end of the last 25-bp window were also trimmed from each sequence. The 5' and 3' tails with an average quality score of <20 at the end of the last window were also trimmed from each sequence. Sequences longer than 250 bp were truncated to 250 bp by trimming the 3' tails. The trimmed sequences shorter than 200 bp were filtered out. The demultiplexing and trimming were performed using Trimmomatic version 0.35 (http://www.usadellab.org/cms/?page=trimmomatic). The remaining sequences were merged into paired reads using Usearch version 8.0.1517 (http://www.drive5.com/usearch/). In addition, singletons were removed. Subsequently, sequences were aligned using Clustal Omega v 1.2.0. (http://www.clustal.org/omega/). Multiple sequences were aligned with each other and only sequences that were contained in more than 75% of the read positions were extracted. Filtering and a part of the multiple alignment process were performed using the screen.seqs and filter.seqs commands in Mothur, as described in the Miseq SOP (http://www.mothur.org/wiki/MiSeq_SOP) [24]. Erroneous and chimeric sequences were detected and removed using the pre.cluster (diffs = 4) and chimera.uchime (minh = 0.1; http://drive5.com/usearch/manual/uchime_algo.html) [25] commands in Mothur, respectively. Using the unique.seqs command of Mothur, the same sequences were collected into operational taxonomic units (OTUs). The contig sequences were counted as OTUs by count.seqs and used for the subsequent taxonomic identification analysis using BLASTn. Eukaryotic groups determined to be apparently of terrestrial origin were excluded, and the others were selected based on abundance.

**Statistical analysis**

Sequence read counts in the sample were converted to relative read counts per million reads, which were used for principal component analysis (PCA). The number of samples having and not having a eukaryotic group was compared between the GC and BSS samples using Fisher’s exact test. Sequence read counts converted to relative read counts per million reads were subsequently standardized to logarithm. Man–Whitney U test was used to compare the logarithms between the GC and BSS samples at α = 0.05 significance level.
Results

Molecular taxonomy of the leptocephali

The basic local-alignment search tool (BLAST) of the GenBank database was used to search the 16S rDNA sequences of 40 leptocephali (Table 1). The nucleotide sequences can be found in the DDBJ-EMBL-GenBank databases (LC439371–LC439410). According to the 16S rDNA sequences, 40 leptocephali comprised 11 *A. japonica*, 11 *A. marmorata*, two *Ariosoma major*, one *Bathyuroconger* sp., one *Conger myriaster*, one *C. jordani* (formerly *C. japonicus*), one *Eurypharynx plecanaoides*, six *Gnathophis* spp., three *Gymnothorax* spp., two *Robinsia* sp., and one *Serrivomer sector*.

Overview of eukaryotic groups in the GC and BSS samples

The GC sample was obtained from 36 leptocephali, because squeezing of GC failed in four samples (Table 1). The BSS sample was collected from 17 leptocephali (Table 1). Of OTU obtained after quality check for 18S rDNA sequences, those having low similarity (< 90%) with the top BLASTn hit sequences (10 OTUs comprising 3484 reads) were removed, resulting in 29 eukaryotic groups and 154 OTUs comprising 269185 reads (106 OTUs comprising 162897 reads in the GC sample and 101 OTUs comprising 106288 reads in the BSS sample) (Table 2). Five eukaryotic groups (asterisk in Table 2) determined to be contaminants of terrestrial origin in the laboratory and fifteen eukaryotic groups (double dagger in Table 2) occurring at low read frequency (<1%) were excluded from further analysis. Phylogenetic analysis using seven fish OTUs with their top BLASTn hits and the host 18S rDNA sequences determined using direct nucleotide sequencing (accession No. LC464077–LC464098) indicated that almost all fish 18S rDNAs obtained using NGS were of the host, revealing incomplete PNA clamping. Therefore, fish OTUs comprising 27507 reads (7 OTUs comprising 26880 reads in the GC sample and 4 OTUs comprising 627 reads in the BSS sample) were also excluded (pilcrow in Table 2). Remaining after applying these selection criteria were eight eukaryotic groups comprising jellyfish (*Cnidaria*) (21 OTUs, 99258 reads), conoid parasite (*Conoidasida*) (7 OTUs, 26272 reads), tunicate (*Chordata*) (14 OTUs, 24148 reads), copepod (*Copepoda*) (15 OTUs, 14144 reads), krill (*Euphausiacea*) (11 OTUs, 11800 reads), annelid (*Polychaeta*) (6 OTUs, 1469 reads), fungus (18 OTUs, 20452 reads), and dinoflagellate (*Dinophyceae*) (6 OTUs, 4017 reads) (Table 2). The nucleotide sequences of these 98 OTUs and seven fish OTUs generated using NGS are available in the DDBJ-EMBL-GenBank database (LC474264–LC474368). Raw read count data for the eight eukaryotic groups are available in S1 Table. One GS sample and one BSS sample having none of these eukaryotic groups were excluded from further analyses.

Eight eukaryotic groups detected in the GC samples comprised 75 OTUs and 103464 reads, and seven eukaryotic groups detected in the BSS samples comprised 64 OTUs and 97612 reads (Table 2, Fig 1A and 1B). Jellyfish was the primary component in both samples, occupying 33.0% of total reads in the GC samples and 67.5% in the BSS samples. Conoid parasite was the second-most abundant contributor (23.8%) in the GC sample but zero in the BSS sample. Less frequent eukaryotic groups in the GC and BSS samples were tunicate (10.1% and 14.0%, respectively), copepod (11.1% and 2.7%), krill (3.9% and 7.9%), annelid (< 0.1% and 1.5%), fungus (16.6% and 3.4%), and dinoflagellate (1.4% and 2.6%).

Eukaryotic compositions in each sample

Eukaryotic composition considerably varied among leptocephalus individuals (Fig 1C). A Shannon-Wiener diversity index ranged between 0 and 0.814 in the GC samples and between
Table 1. Summary of leptocephalus samples collected in 2017 and used in this study.

| ID     | Sample | BLAST top hit (% identity) | date      | coordinate (N, E) | BL (mm) |
|--------|--------|-----------------------------|-----------|-------------------|---------|
| Aj-330 | GC     | Anguilla japonica (100)      | Oct. 3    | 18.021, 130.992   | 42.0    |
| Aj-332 | GC     | Anguilla japonica (100)      | Oct. 3    | 18.021, 130.992   | 52.3    |
| Aj-343 | GC     | Anguilla japonica (100)      | Oct. 3    | 18.002, 131.003   | 45.0    |
| Aj-410 | GC     | Anguilla japonica (100)      | Oct. 3    | 18.675, 131.070   | 42.0    |
| Aj-446 | GC     | Anguilla japonica (100)      | Oct. 4    | 18.024, 131.075   | 47.0    |
| Aj-452 | GC     | Anguilla japonica (100)      | Oct. 4    | 18.069, 131.068   | 43.0    |
| Aj-453 | GC/BSS | Anguilla japonica (100)      | Oct. 4    | 18.069, 131.068   | 42.0    |
| Aj-470 | GC/BSS | Anguilla japonica (100)      | Oct. 4    | 18.069, 131.105   | 44.0    |
| Aj-475 | GC/BSS | Anguilla japonica (99)       | Oct. 4    | 18.059, 131.069   | 43.0    |
| Aj-476 | GC/BSS | Anguilla japonica (100)      | Oct. 4    | 18.059, 131.069   | 42.0    |
| Aj-664 | GC/BSS | Anguilla japonica (99)       | Oct. 10   | 18.501, 131.490   | 45.8    |
| Am-38  | GC     | Anguilla marmorata (100)     | Sep. 30   | 24.501, 130.992   | 42.2    |
| Am-571 | GC/BSS | Anguilla marmorata (100)     | Oct. 6    | 14.502, 130.988   | 36.7    |
| Am-577 | GC     | Anguilla marmorata (100)     | Oct. 6    | 14.010, 131.003   | 37.7    |
| Am-604 | GC/BSS | Anguilla marmorata (100)     | Oct. 8    | 15.490, 128.490   | 42.7    |
| Am-611 | GC     | Anguilla marmorata (100)     | Oct. 9    | 15.974, 128.502   | 41.1    |
| Am-612 | GC/BSS | Anguilla marmorata (100)     | Oct. 9    | 15.974, 128.502   | 45.2    |
| Am-697 | GC     | Anguilla marmorata (100)     | Oct. 11   | 15.601, 128.335   | 42.2    |
| Am-712 | GC     | Anguilla marmorata (100)     | Oct. 11   | 15.687, 128.369   | 37.6    |
| Am-720 | GC     | Anguilla marmorata (100)     | Oct. 12   | 15.734, 128.388   | 37.9    |
| Am-736 | GC     | Anguilla marmorata (99)      | Oct. 12   | 15.773, 128.234   | 51.0    |
| Am-909 | GC     | Anguilla marmorata (100)     | Oct. 14   | 21.810, 131.275   | 48.0    |
| CG-468 | BSS    | Ariosaoma major (99)        | Oct. 4    | 18.070, 131.124   | 65.0    |
| CG-469 | BSS    | Ariosaoma major (99)        | Oct. 4    | 18.070, 131.124   | 214.0   |
| CG-305 | GC     | Bathysuroconger vicinus (96) | Oct. 2    | 18.506, 131.006   | 57.7    |
| Cm-342 | GC     | Conger myriaster (99)       | Oct. 3    | 18.021, 130.992   | 65.0    |
| Cj-488 | GC/BSS | Conger jordani (100)        | Oct. 4    | 18.059, 131.069   | 43.1    |
| EU-758 | GC     | Eurypharynx plecanoides (99) | Oct. 12   | 15.313, 128.234   | 18.1    |
| CG-15  | GC     | Gnathophis bathytopos (99)  | Sep. 29   | 25.004, 130.985   | 62.7    |
| CG-16  | GC     | Gnathophis bathytopos (98)  | Sep. 29   | 25.004, 130.985   | 49.6    |
| CG-301 | GC     | Gnathophis bathytopos (99)  | Oct. 2    | 18.506, 131.005   | 56.3    |
| CG-303 | GC     | Gnathophis bathytopos (99)  | Oct. 2    | 18.506, 131.005   | 62.8    |
| CG-878 | GC     | Gnathophis bathytopos (99)  | Oct. 13   | 21.424, 131.143   | 67.5    |
| CG-879 | GC     | Gnathophis bathytopos (99)  | Oct. 13   | 21.424, 131.143   | 68.1    |
| MR-344 | GC     | Gymnothorax melatremus (92) | Oct. 3    | 18.002, 131.003   | 42.0    |
| MR-471 | BSS    | Gymnothorax margaritophorus (100) | Oct. 4 | 18.069, 131.101 | 34.0    |
| MR-483 | GC/BSS | Gymnothorax niphotogonus (93)| Oct. 4    | 18.059, 131.069   | 41.6    |
| CH-572 | GC/BSS | Robinsonia catherinae (96)  | Oct. 6    | 14.502, 130.988   | 69.3    |
| CH-663 | GC/BSS | Robinsonia catherinae (96)  | Oct. 9    | 18.485, 131.001   | 63.8    |
| SR-614 | BSS    | Serrivomer sector (99)      | Oct. 9    | 15.974, 128.502   | 45.5    |

*GC: only gut content sample was analyzed; GC/BSS: both gut content and body surface scraping samples were analyzed; BSS: only body surface scraping sample was analyzed.

†Based on mitochondrial 16S rDNA sequence analysis for the leptocephali.

‡Leptocephali were subjected to direct 18 rDNA sequence analysis.

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0 and 0.699 in the BSS samples with no significant difference between the samples (Mann–Whitney U test, \( p = 0.578 \)), but heterogeneity between the GC and BSS samples mentioned...
above was also the case at individual level (Fig 1C). Systematic difference in eukaryotic compositions between the GC and BSS samples was illustrated using PCA analysis (Fig 2). The GC samples were dispersed regardless of species (Fig 2, black symbols). On the other hand, the BSS samples were closely related one another (Fig 2, yellow symbols) except for three outliers (Fig 2, arrow), in which no jellyfish read was observed in these three BSS samples (see also Fig 1C, Aj-664S, Am-612S, and SR-614S).

Jellyfish reads were detected in 18 of 35 GC samples and 13 of 16 BSS samples with significant difference (Fisher’s exact test, \( p = 0.040 \)), and the standardized read counts of the jellyfish were greater in the BSS samples than in the GC samples (Mann–Whitney U test, \( p < 0.005 \)).

Table 2. Summary of eukaryotic groups detected in the gut content (GC) and body surface scraping (BSS) samples of eel leptcephali, and number of OTUs and reads of 18S rDNA.

| organism group         | Phylum          | lower taxa                  | No. OTUs | No. reads (No. larvae) | BLAST % identity |
|------------------------|-----------------|-----------------------------|----------|------------------------|------------------|
| metazoa                |                 |                             | all      | GC         | BSS               |
| jellyfish              | Cnidaria        | Anthozoa, Hydrozoa          | 21       | 18 (16)    | 16 (14)          | 96.6–100         |
| conoid parasite        | Apicomplexa     | Coccidia, Gregarinina       | 7        | 7 (0)      | 0 (0)            | 90.9–98.5        |
| tunicate               | Chordata        | Appendicularia, Thaliacea   | 14       | 12 (14)    | 13 (11)         | 95.9–100         |
| copepod                | Arthropoda      | Copepod                     | 15       | 11 (9)     | 12 (10)         | 92.3–100         |
| krill                  | Arthropoda      | Euphausia                     | 11       | 9 (10)     | 12 (11)         | 96.9–99.6        |
| annelid                | Annelida        | Polychaeta                   | 6        | 6 (1)      | 1 (1)           | 97.8–100         |
| fish*                  | Chordata        | Actinopterygii               | 7        | 7 (4)      | 5 (4)           | 91.8–99.8        |
| acorn worm#            | Hemichordata    | Enteropneusta                | 2        | 2 (2)      | 2 (2)           | 98.0–99.8        |
| shrimp#                | Arthropoda      | Decapoda                     | 4        | 1 (1)     | 2 (2)           | 99.6–100         |
| snail†                 | Mollusca        | Gastropoda                   | 9        | 2 (7)      | 1 (1)           | 94.0–99.8        |
| arrow worm‡            | Chaetognatha    | Aphragmophora                | 4        | 4 (3)      | 1 (1)           | 98.7–99.6        |
| comb jelly‡            | Ctenophora      | Tentaculata                  | 2        | 0 (2)     | 0 (2)           | 100              |
| ostracods‡             | Arthropoda      | Holocyprida                  | 1        | 1 (1)     | 1 (1)           | 100              |
| mite*                  | Arthropoda      | Archinida                    | 1        | 1 (1)     | 1 (1)           | 100              |
| public lice‡           | Arthropoda      | Insecta                      | 1        | 1 (1)     | 2 (2)           | 100              |
| silkworm‡              | Arthropoda      | Insecta                      | 1        | 1 (0)     | 0 (0)           | 99.6             |
| human†                 | Chordata        | Mammalia                     | 1        | 1 (0)     | 1 (0)           | 99.5             |
| fungi                  |                 |                             | 18       | 16 (14)    | 4 (2)           | 95.1–100         |
| dinoflagellate         | Dinoflagellata  | Gonyaulacales, Syndiniales   | 6        | 6 (5)     | 1 (1)           | 91.1–99.8        |
| radiolarias‡           | Radiozoa        | Colloidaria                  | 6        | 3 (3)     | 3 (3)           | 91.8–99.8        |
| golden algae‡          | Chrysophyceae   | Chromulinales                | 1        | 1 (0)     | 1 (0)           | 100              |
| green algae‡           | Chlorophyta     | Prasinococcales, Pyramimonadales | 2 | 0 (2) | 0 (0) | 94.9 (1) | 91.9–99.6 |
| filose amoebae‡        | Cercozoa        | Chlorarachniophycaceae       | 1        | 0 (1)     | 1 (0)           | 99.6             |
| heterokonts‡           | Bigya           | Bicoecida                    | 2        | 2 (2)     | 0 (0)           | 97.2             |
| cryptomonads‡          | Cryptophyta     | Pyrenomonaedales             | 1        | 0 (1)     | 1 (0)           | 91.7             |
| heterokont algae‡      | Dictyochophyceae| Rhizochromulinales           | 1        | 0 (1)     | 0 (0)           | 97.5             |
| apusozoa‡              | Apusozoa        | Apusomonadaid                | 1        | 0 (1)     | 0 (0)           | 95.4             |
| brown algae‡           | Phaeophyceae    | Laminariales                 | 1        | 0 (1)     | 0 (0)           | 97.7             |
| flowering plants‡      | Magnoliophyta   |                             | 7        | 6 (1)     | 1 (1)           | 94.6–100         |

\[\text{† determined to be the host 18S rDNA} \]
\[\text{‡ excluded due to the low read frequency (<1%)} \]
\[\text{§ determined to be terrestrial origin and excluded} \]
\[\text{¶ significantly greater than the other sample.} \]
Occurrences of the krill, annelid, and dinoflagellate were also greater in the BSS samples than in the GC samples (Fisher’s exact test, \( p < 0.05 \)), of which the standardized read counts of the annelid were greater in the BSS samples than in the GC samples (Mann–Whitney U test, \( p = 0.026 \)). In contrast, occurrences of the conoid parasite and fungus were significantly higher in the GC samples than in the BSS samples; specifically no conoid parasite read was detected in the BSS samples (Fisher’s exact test, \( p < 0.05 \), Mann–Whitney U test, \( p < 0.05 \)). No such significant heterogeneity between the GC and BSS samples was observed in the tunicate and copepod.

Since it has been suggested that cnidarian jellyfishes may be important diet for the leptocephalus larvae \([12, 14]\), jellyfish taxa and the read number in 13 leptocephali having both the GC and BSS samples were investigated (Fig 3). Eight families and one suborder (Calyco- phorae) of cnidarian taxa were chosen according to the abundancy (total read number larger than 100). Of 13 leptocephali, six had jellyfish reads in both the GC and BSS samples, five had those only in the BSS sample, one had those only in the GC sample, and one had no jellyfish read in both the GC and BSS samples. Of six leptocephali having jellyfish reads in both the GC...
and BSS samples, only one (Aj470) had no common jellyfish read between the GC and BSS samples.

**Discussion**

Since primer choice has been known to considerably influence quantitative estimations on the target molecules in metagenome study [26], results of our metagenomic analysis may not reflect the true eukaryote composition in the gut contents of eel leptocephali. However, our results are reliable in comparing the eukaryote composition between the GC and the BSS samples of the eel leptocephali. We observed that almost all eukaryotic groups (except for conoid parasite) highlighted were common between the GC and BSS samples. The occurrence of jellyfish, krill, annelid, and dinoflagellate and/or the standardized read counts were significantly

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Fig 3. Composition of cnidarian jellyfish taxa detected in the gut content (G) and body surface scraping (B) samples of 13 leptocephali having both the G and B samples. Sequence read counts in the sample were converted to relative read counts per million reads.

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higher in the BSS samples than in the GC samples. Abundant eukaryotic groups, such as conoid parasite and fungi, were distinctively observed in the GC samples than in the BSS samples, indicating little index hopping effect in this study. Compared with the BSS samples, conoid parasite exclusively detected in the GC samples may be indicative of very little cross contamination in the direction from the GC samples to the BSS samples. One dinoflagellate OTU assigned to *Hematodinium perezi* was observed only in the GC samples; this dinoflagellate is also known to be parasitic [27]. Therefore, the results obtained in this study indicate that just careful rinsing of the body surface of the leptocephali with sterile artificial seawater cannot prevent cross contamination of DNAs derived from the body surface and/or environment. Cnidarian planktons were suggested to be an important diet for the leptocephalus larvae, because 18S rDNA sequences of hydrozoa jellyfish predominated the GC of *A. anguilla* larvae [12,14]. However, there are several discrepancies in judging that jellyfish is the main food source for leptocephali. Data accumulating from stable isotope analysis indicate that cnidarians usually stay at relatively higher trophic position [28–36]; trophic positions of leptocephali have been observed to be apparently lower than those of cnidarian jellyfish [6, 18]. A few cnidarians may stay at low trophic positions [28,35], and it was claimed that cnidarians in oligotrophic areas, like the Sargasso Sea, may have low δ15N values [14]. However, it is unlikely that leptocephali selectively consume specific cnidarians at lower trophic position and migrate in oligotrophic water mass with oligotrophic organisms all the way down from the spawning area to nursery area. Because genes from gelatineous zooplankton have been found in the guts of lobster larvae [22,37–39], jellyfish consumption by leptocephali was further advocated owing to similarity in the highly flattened body and long larval period between the lobster and eel larvae [14]. The lobster larvae are active predator, because they have actually been observed to capture and prey upon a variety of agile zooplanktons [37,40,41]; however, such predatory behavior toward zooplankton has never been observed in leptocephali [5,42,43]. Results of the feeding experiments of leptocephali attempted till date are summarized in Table 3. Hatchery-produced *A. japonica* or *A. anguilla* leptocephali were used in all experiments except for wild-caught leptocephali of pike conger (*Muraenesox cinereus*) and white-spotted conger (*Conger myriaster*) [5]. Among the considerably wide variety of food candidates tested, all well consumed foods, except for the smallest aloricate rotifer *Proales similis*, were processed in the form of a paste. Gelatinous animals appear as an unfavorable feed, because fresh chopped Cnidaria (*Aurelia* sp.) was still capable of killing the Japanese eel leptocephali by the nematocysts and alive ctenophore even captured and preyed upon the leptocephali (unpublished laboratory experiment by Hideki Tanaka). Furthermore, Japanese eel leptocephali teeth were disadvantage for feeding on sticky jellyfish tissue (unpublished on board experiment by Seinen Chow). Therefore, we conclude that cross contamination from the body surface occurred while collecting the GC and the pronounced detection of cnidarian 18S rDNA from the body surface was due to the cnidarian nematocysts sting to the leptocephali in the plankton net. No notable difference was found in eukaryotic composition between full and empty gut individuals [14], which also supports our conclusion.

Regarding the eukaryotes that were found in the gut of leptocephali, parasites are definitive and fungi are probably positive candidates; however, it is unlikely that these can be feed for the leptocephali. Not many, but feces of zooplankton, larvacean and their houses have been occasionally observed in the gut of leptocephali [6–10], corresponding to relatively larger read counts of copepod and tunicate 18S rDNA in the gut samples observed in this study. Thus, the present study supports previous results that leptocephali utilize detritus materials, so called marine snow [6,8–10,13,17,19,44]. In the tropical and subtropical regions of the western North Pacific, copepods are reported to be the most abundant zooplankton taxa occupying at least 70% or more of total mesozooplankton biomass in mesopelagic layer, followed by
urochordates and chaetognaths but cnidarians jellyfish at much lower abundance [45–47]. Cnidarians might contribute to some part of the marine snow formation and therefore, may be a diet component for the leptocephali. However, it is unlikely that cnidarians are the main food sources for the eel leptocephali because of the higher stable isotope ratio and lower abundance.

Supporting information
S1 Table. Eukaryotic groups highlighted in this study. Raw read counts of eight eukaryotic groups detected in 36 gut contents (GC) and 17 body surface scraping (BSS) samples obtained from 40 Anguilliformes leptocephali. (XLSX)

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References

1. Alexander EC. A contribution to the life history, biology and geographical distribution of the bonefish, *Albula vulpes* (Linnaeus). Dana Rep. 1961; 53: 1–51.

2. Hulet WH. Structure and functional development of the eel leptocephalus *Ariosoma balearicum* (DeLa Roche, 1809). Phil. Trans. Roy. Soc. B 1978; 252: 107–138.

3. Moser HG. Morphological and functional aspects of marine fish larvae. In Lasker R, editor. Marine fish larvae: morphology, ecology, and relation to fisheries. Seattle: University of Washington Press; 1981. pp. 90–131.

4. Otake T. Fine structure and function of the alimentary canal in leptocephali of the Japanese eel *Anguilla japonica*. Fish. Sci. 1996; 62: 28–34.

5. Mochioka N, Iwamizu M, Kanda K. Leptocephalus eel larvae will feed in aquaria. Env. Biol. Fish. 1993; 36: 381–384.

6. Otake T, Nogami N, Maruyama K. Dissolved and particulate organic matter as possible food sources for eel leptocephali. Mar. Ecol. Prog. Ser. 1993; 92: 27–34.

7. Mochioka N, and Iwamizu M. Diet of anguiloid larvae: leptocephali feed selectively on larvacean houses and fecal pellets. Mar. Biol. 1996; 125: 446–452.

8. Miller MJ, Otake T, Aoyama J, Wouthuyzen S, Suhariti HY, Sugeha S, Tsukamoto K. Observations of gut contents of leptocephali in the North Equatorial Current and Tomini Bay, Indonesia. Coast. Mar. Sci. 2011; 35: 277–288.

9. Miller M.J, Marohn L, Wysujack K, Freese M, Pohlmann JD, Westerberg H, et al. Morphology and gut contents of anguillid and marine eel larvae in the Sargasso Sea. Zool. Anzeig. 2019; 279: 138–151. https://doi.org/10.1016/j.jcz.2019.01.008

10. Tomoda T, Chow S, Kurogi H, Okazaki M, Ambre D, Furulta H, et al. Observations of gut contents of anguilliform leptocephali collected in the western North Pacific. Nippon Suisan Gakkaishi 2017; 84: 32–44. https://doi.org/10.3989/suisan.17-00025 (in Japanese with English abstract)

11. Govoni J. Feeding on protists and particulates by the leptocephali of the worm eels *Myrophis* spp. (Teleostei, Anguilliformes, Ophichthidae), and the potential energy contribution of large aloricate protozoa. Sci. Mar. 2010; 74: 339–344. https://doi.org/10.3989/scimar.2010.74n233

12. Riemann L, Alfredsson H, Hansen MM, Als TD, Nielsen TG, Munk P, et al. Qualitative assessment of the diet of European eel larvae in the Sargasso Sea resolved by DNA barcoding. Biol. Lett. 2010; 6: 819–822. https://doi.org/10.1098/rsbl.2010.0411 PMID: 20573615

13. Terahara T, Chow S, Kurogi H, Lee S-H, Tsukamoto K, Mochioka N, et al. Efficiency of peptide nucleic acid-directed PCR clamping and its application in the investigation of natural diets of the Japanese eel leptocephali. PLoS One 2011; 6: e25715. https://doi.org/10.1371/journal.pone.0025715 PMID: 22069444
14. Ayala DJ, Munk P, Lundgreen RBC, Traving SJ, Jaspers C, Jørgensen TS, et al. Gelatinous plankton is important in the diet of European eel (Anguilla anguilla) larvae in the Sargasso Sea. Sci. Rep. 2018; 8: 6156. https://doi.org/10.1038/s41598-018-24388-x PMID: 29670123

15. Chow S, Kurogi H, Katayama S, Ambe D, Okazaki M, Watanabe T, et al. Japanese eel Anguilla japonica do not assimilate nutrition during the oceanic spawning migration: evidence from stable isotope analysis. Mar. Ecol. Prog. Ser. 2010; 402: 233–238. https://doi.org/10.3354/meps08448

16. Miyazaki S, Kim H-Y, Zenimoto K, Kitagawa T, Miller MJ, Kimura S. Stable isotope analysis of two species of anguilliform leptocephali (Anguilla japonica and Ariposa major) relative to their feeding depth in the North Equatorial Current region. Mar. Biol. 2011; 158: 2555–2564. https://doi.org/10.1007/s00227-011-1756-x

17. Miller MJ, Chikaraishi Y, Ogawa NO, Yamada Y, Tsukamoto K, Ohkouchi N. A low trophic position of Japanese eel larvae indicates feeding on marine snow. Biol. Lett. 2013; 9: 20120826. https://doi.org/10.1098/rsbl.2012.0826 PMID: 23134783

18. Feunteun E, Miller MJ, Carpentier A, Aoyama J, Dupuy C, Kuroki M, et al. Stable isotopic composition of anguilliform leptocephali and other food web components from west of the Mascarene Plateau. Prog. Oceanogr. 2015; 137: 69–83. https://doi.org/10.1016/j.pocean.2015.05.024

19. Quattrini AM, McClain-Counts J, Artabane SJ, Roa-Varón A, Mclver TC, Rhode M, Ross SW. Assessment of marine snow importance in the diet of European eel (Anguilla anguilla). Mar. Biol. 2011; 158: 2555–2564. https://doi.org/10.1007/s00227-011-1756-x

20. Palumbi S, Martin A, Romano S, McMillan WO, Stice L, Grabowski G. The Simple Fool's Guide to PCR. 2nd ed. Honolulu: University of Hawaii; 1991.

21. Chow S, Kurogi H, Yamamoto T, Tomoda T, Mochioka N, Shirotori F, et al. Reproductive isolation between sympatric Anguilla japonica and A. marmora ta. J. Fish Biol. 2017; 91: 1517–1525. https://doi.org/10.1111/jfb.13483 PMID: 28990671

22. Chow S, Suzuki S, Matsunaga T, Lavery S, Jeffs A, Takeyama H. Investigation on natural diets of larval marine animals using peptide nucleic acid (PNA)-directed PCR clamping. Mar. Biotech. 2011; 13: 305–313. https://doi.org/10.1007/s10126-010-9301-3

23. Dzhembekova N, Urushizaki S, Moncheva S, Ivanova P, Nagai S. Application of massively parallel sequencing on monitoring harmful algae at Varna Bay in the Black Sea. Harmful Algae 2017; 68: 40–51. https://doi.org/10.1016/j.hal.2017.07.004 PMID: 28962989

24. Schloss PD, Gevers D, Westcott SL. Reducing the effects of PCR amplification and sequencing artifacts on 16S rRNA-based studies. PLoS One 2011; 6: e27310. https://doi.org/10.1371/journal.pone.0027310 PMID: 22194782

25. Edgar RC, Haas BJ, Clemente JC, Quince C, Knight R. UCHIME improves sensitivity and speed of chimera detection. Bioinformatics 2011; 27: 2194–2200. https://doi.org/10.1093/bioinformatics/btr381 PMID: 21700674

26. Tremblay J, Singh K, Fern A, Kirton ES, He S, Woyke T, Lee J, Chen F, Dangl JL, Tringe SG. Primer and platform effects on 16S rRNA tag sequencing. Front. Microbiol. 2015; 6: 771. https://doi.org/10.3389/fmicb.2015.00771 PMID: 26300854

27. Newman MW, Johnson CA. A disease of blue crab (Callinectes sapidus) caused by a parasitic dinoflagella te, Hematodinium sp. J. Parasitol. 1975; 63: 554–557.

28. Cardona L, de Quevedo IA, Borrell A, Aguilar A. Massive consumption of gelatinous plankton by Mediterranean apex predators. PLoS One 2012; 7: e31329. https://doi.org/10.1371/journal.pone.0031329 PMID: 22470416

29. D’Ambra I, Carmichael RH, Graham WM. Determination of δ13C and δ15N and trophic fractionation in jellyfish: implications for food web ecology. Mar. Biol. 2014; 161: 473–480. https://doi.org/10.1007/s00227-013-2345-y

30. D’Ambra I, Graham WM, Carmichael RH, Hernandez FJ Jr. Fish rely on scyphozoan host as a primary food source: evidence from stable isotope analysis. Mar. Biol. 2015; 162: 247–252. https://doi.org/10.1007/s00227-014-2569-5

31. Fleming NEC, Harrod C, Newton J, Houghton JDR. Not all jellyfish are equal: isotopic evidence for inter- and intraspecific variation in jellyfish trophic ecology. PeerJ 2015; 3: e1110. https://doi.org/10.7717/peerj.1110 PMID: 26244116

32. Javidpour J, Cipriano-Maak AN, Mittermayr A, Dierking J. Temporal dietary shift in jellyfish revealed by stable isotope analysis. Mar. Biol. 2016; 163: 112–120. https://doi.org/10.1007/s00227-016-2892-0 PMID: 27194816

33. Ingram BA, Pitt KA, Barnes P. Stable isotopes reveal a potential kleptoparasitic relationship between an ophiuroid (Ophiocnemis marmorata) and the semaeostome jellyfish, Aurelia aurita. J. Plankt. Res. 2017; 39: 138–146. https://doi.org/10.1093/plankt/fbw088
34. MacKenzie KM, Trueman CN, Lucas CH, Bortoluzzi J. The preparation of jellyfish for stable isotope analysis. Mar. Biol. 2017; 164: 219–227. https://doi.org/10.1007/s00227-017-3242-6

35. Tilves U, Sabatés A, Blázquez M, Raya V, Fuentes VL. Associations between fish and jellyfish in the NW Mediterranean. Mar. Biol. 2018; 165: 127–140. https://doi.org/10.1007/s00227-018-3381-4

36. Zeman SM, Corrales-Ugalde M, Brodeur RD, Sutherland KR. Trophic ecology of the neustonic cnidian Velella velella in the northern California Current during an extensive bloom year: insights from gut contents and stable isotope analysis. Mar. Biol. 2018; 165: 150–162. https://doi.org/10.1007/s00227-018-3404-1

37. Suzuki N, Hoshino K, Murakami K, Takeyama H, Chow S. Molecular diet analysis of phyllosoma larvae of the Japanese spiny lobster Panulirus japonicus (Decapoda: Crustacea). Mar. Biotech. 2008; 10: 49–55. https://doi.org/10.1007/s10126-007-9038-9

38. O’Rorke R, Lavery S, Chow S, Takeyama H, Tsai P, Beckley LE, et al. Determining the diet of larvae of western rock lobster (Panulirus cygnus) using high-throughput DNA sequencing techniques. PLoS One 2012; 7: e42757. https://doi.org/10.1371/journal.pone.0042757 PMID: 22927937

39. Wangm M, Jeffs AG. Nutritional composition of potential zooplankton prey of spiny lobster larvae: a review. Rev. Aquacult. 2014; 6: 270–299. https://doi.org/10.1111/raq.12044

40. Mitchell JR. Food preferences, feeding mechanisms and related behavior in phyllosoma larvae of the California spiny lobster, Panulirus interruptus (Randall). M.Sc. Thesis, San Diego State College. 1971.

41. Saunders MI, Thompson PA, Jeffs AG, Sáwström C, Sachlikidid N, Beckley LE, Waite AM. Fussy feeders: phyllosoma larvae of the western rocklobster (Panulirus cygnus) demonstrate prey preference. PLoS One 2012; 7: e36580. https://doi.org/10.1371/journal.pone.0036580 PMID: 22586479

42. Wullur S, Yoshimatsu T, Tanaka H, Ohtani M, Sakakura Y, Kim H-J, Hagiwara A. Ingestion by Japanese eel Anguilla japonica larvae on various minute zooplanktons. Aquacult. Sci. 2013; 61: 341–347. https://doi.org/10.11233/aquaculturesci.61.341

43. Butts IAE, Sørensen SR, Politis SN, Tomkiewicz J. First-feeding by European eel larvae: A step forwards closing the life cycle in captivity. Aquaculture 2016; 464: 451–458. https://doi.org/10.1016/j.aquaculture.2016.07.028

44. Miller MJ. Ecology of anguilliform leptocephali: remarkable transparent fish larvae of the ocean surface layer. Aqua-Bio Sci. Monog. 2009; 2: 1–94. https://doi.org/10.5047/absm.2009.00204.0001

45. Landry MR, Al-Mutairi H, Selph KE, Christensen S, Nunnery S. Seasonal patterns of mesozooplankton abundance and biomass at Station ALOHA. Deep-Sea Res. II 2001; 48: 2037–2061. https://doi.org/10.1016/S0967-0645(00)00172-7

46. Sun D, Wang C. Latitudinal distribution of zooplankton communities in the Western Pacific along 160E during summer 2014. J. Mar. Sys. 2017; 169: 52–60. https://doi.org/10.1016/j.jmarsys.2017.01.011

47. Tanaka H, Kagawa H, Ohta H. Production of leptocephali of Japanese eel (Anguilla japonica) in captivity. Aquaculture 2001; 201: 51–60. https://doi.org/10.1016/S0044-8486(01)00553-1

48. Tanaka H, Kagawa H, Ohta H, Okuzawa K, Hirose K. The first report of eel larvae ingesting rotifers. Fish. Sci. 1995; 61: 171–172.

49. Chow S, Kurogi H, Watanabe S, Matsunari H, Sudo R, Nomura K, Tanaka H, Furuita H, Nishimoto A, Higuchi M, Jinbo T, Tomoda T, Onboard rearing attempts for the Japanese eel leptocephali using POM-enriched water collected in the Western North Pacific. Aquat. Liv. Resour. 2017; 30: 36. https://doi.org/10.1051/alr/2017037

50. Hagiwara A, Wullur S, Marcial HS, Hirai N, Sakakura Y. Euryhaline rotifer Proales similis as initial live food for rearing fish with small mouth. Aquaculture 2014; 432: 470474. https://doi.org/10.1016/j.aquaculture.2014.03.034

51. Tanaka H, Kagawa H, Ohta H. Production of leptocephali of Japanese eel (Anguilla japonica) in captivity. Aquaculture 2001; 201: 51–60. https://doi.org/10.1016/S0044-8486(01)00553-1

52. Tanaka H, Kagawa H, Ohta H, Unuma T, Nomura K. The first production of glass eel in captivity: fish reproduction physiology facilitates great progress in aquaculture. Fish Physiol. Biochem. 2003; 28: 493–497.

53. Tomoda T, Kurohi H, Okauchi M, Kamoshida M, Imaizumi H, Jinbo T, et al. Hatchery-reared Japanese eel Anguilla japonica larvae ingest various organic matter formed as part of marine snow. Nippon Suisan Gakkaishi 2015; 81: 715–721. https://doi.org/10.2331/suisan.81.715