DATA NOTE

The genome sequence of the orange-striped anemone, *Diadumene lineata* (Verrill, 1869) [version 1; peer review: 3 approved]

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**Abstract**

We present a genome assembly from an individual *Diadumene lineata* (the orange-striped anemone; Cnidaria; Anthozoa; Actiniaria; Diadumenidae). The genome sequence is 313 megabases in span. The majority of the assembly (96.03%) is scaffolded into 16 chromosomal pseudomolecules. The complete mitochondrial genome was also assembled and is 17.6 kilobases in length.

**Keywords**

Diadumene lineata, orange-striped anemone, genome sequence, chromosomal, Cnidaria

This article is included in the Tree of Life gateway.

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Species taxonomy
Eukaryota; Metazoa; Cnidaria; Anthozoa; Hexacorallia; Actiniaria; Nynantheae; Diadumenidae; Diadumene; Diadumene lineata (Verrill, 1869) (NCBI:txid1789172).

Background
The Orange-striped anemone, Diadumene lineata (Verrill, 1869), is believed to be the world’s most widely distributed sea anemone. Native to the Northwest Pacific, it is now established on almost every temperate and tropical coast worldwide, and is a remarkable colonising species that serves as a model by which to address invasion hypotheses (Flenniken, 2017). In the UK, it has been recorded all along the south coast of England, around the Welsh coast, and from a few sites in Northern Ireland and Scotland. In these areas, it is typically found in sheltered estuaries attached to artificial structures in marinas and harbours, often in association with oysters and mussels, but also on sheltered natural shores, on stones, shells and seaweeds.

Diadumene lineata is a small, delicate anemone, with a smooth column up to 20 mm in diameter (in the UK, but larger in its native range). Generally, it is olive green or brown with contrasting orange vertical stripes. It has 25–100 slender, smooth tentacles, which are all of one type and usually colourless, but can be reddish. Thread-like defensive organs (acontia) can extend through pores in the column. It preys mainly on small crustaceans but may also consume larvae of commercially important species such as oysters and mussels. Under suitable conditions, it can quickly form large clonal aggregations.

In its native range D. lineata reproduces both asexually by fission and sexually (Ryan & Miller, 2019). However, outside its native range it is presumed that only asexual reproduction occurs, as no populations with both males and females together have been reported, except for a recently discovered population with both males and females in Coos Bay, Oregon, USA (Newcomer et al., 2019).

Genome sequence report
The genome was sequenced from a single D. lineata of unknown sex collected from Queen Anne’s Battery Marina visitors’ pontoon, Plymouth, UK (Figure 1). A total of 27-fold coverage in Pacific Biosciences single-molecule HiFi long reads and 82-fold coverage in 10X Genomics read clouds were generated. Primary assembly contigs were scaffolded with chromosome conformation Hi-C data. Manual assembly curation corrected 113 missing/misjoins and removed 43 haplotypic duplications, reducing the assembly size by 1.34% and the scaffold number by 41.95%, and increasing the scaffold N50 by 101.80%.

The final assembly has a total length of 313 Mb in 137 sequence scaffolds with a scaffold N50 of 17.7 Mb (Table 1). The majority, 96.03%, of the assembly sequence was assigned to 11 chromosomal-level scaffolds, representing 16 autosomes (numbered by sequence length) (Figure 2–Figure 5; Table 2). Two 3-Mbp sub-chromosome sized scaffolds were added as S17 and S18 to the unlocalised sequences. S17 and S18 are part of the host, as evidenced by SSU markers and coverage. Parts of

Table 1. Genome data for Diadumene lineata, jaDiaLine6.1.

| Project accession data                  |
|-----------------------------------------|
| Assembly identifier                     | jaDiaLine6.1                         |
| Species                                | Diadumene lineata                    |
| Specimen                                | jaDiaLine6 (genome assembly); jaDiaLine7 (Hi-C, RNA-Seq) |
| NCBI taxonomy ID                        | 1789172                              |
| BioProject                              | PRJEB46855                            |
| BioSample ID                            | SAMEA7536572                         |
| Isolate information                     | Whole organism (jaDiaLine6); other somatic tissue (jaDiaLine7) |

| Raw data accessions                     |
|-----------------------------------------|
| PacificBiosciences SEQUEL II            | ERR6808024                            |
| 10X Genomics Illumina                   | ERR6688656-ERR6688659                 |
| Hi-C Illumina                          | ERR6688655                            |
| PolyA RNA-Seq Illumina                  | ERR6688660                            |

| Genome assembly                         |
|-----------------------------------------|
| Assembly accession                      | GCA_918843875.1                       |
| Accession of alternate haplotype        | GCA_918843945.1                       |
| Span (Mb)                               | 313                                  |
| Number of contigs                       | 320                                  |
| Contig N50 length (Mb)                  | 2.7                                  |
| Number of scaffolds                     | 137                                  |
| Scaffold N50 length (Mb)                | 17.7                                 |
| Longest scaffold (Mb)                   | 42.0                                 |
| BUSCO* genome score                     | C:96.1%(S:95.6%,D:0.5%), F:2.0%, M:1.9%, n:954 |

*BUSCO scores based on the metazoa_odb10 BUSCO set using v5.1.2. C= complete [S= single copy, D=duplicated], F=fragmented, M=missing, n=number of orthologues in comparison. A full set of BUSCO scores is available at https://blobtoolkit.genomeshubs.org/view/jaDiaLine6.1/dataset/CAKXNV01/busco.
the centromere could not be uniquely assigned to chromosomes and are part of the unlocalised sequence.

The assembly has a BUSCO v5.1.2 (Manni et al., 2021) completeness of 96.1% (single 95.6%, duplicated 0.5%) using the metazoa_odb10 reference set (n=954). While not fully phased, the assembly deposited is of one haplotype. Contigs corresponding to the second haplotype have also been deposited.

**Methods**

**Sample acquisition and DNA extraction**

Two *D. lineata* specimens (jaDiaLine6 and jaDiaLine7) were collected by hand from Queen Anne’s Battery Marina visitors’ pontoon, Plymouth, UK (latitude 50.3644, longitude -4.1320) by John Bishop, Joanna Harley (both Marine Biological Association) and Rob Mrowicki (Natural History Museum). The specimens were identified by Chris Wood (Marine Biological Association) and John Bishop and snap-frozen in liquid nitrogen.

DNA was extracted at the Tree of Life laboratory, Wellcome Sanger Institute. The jaDiaLine6 sample was weighed and dissected on dry ice with tissue set aside for Hi-C sequencing. Whole organism tissue was disrupted using a Nippi Power-masher fitted with a BioMasher pestle. Fragment size analysis of 0.01–0.5 ng of DNA was then performed using an Agilent FemtoPulse. High molecular weight (HMW) DNA was extracted using the Qiagen MagAttract HMW DNA extraction kit. Low molecular weight DNA was removed from a 200-ng aliquot of extracted DNA using 0.8X AMPure XP purification kit prior to 10X Chromium sequencing; a minimum of 50 ng DNA was submitted for 10X sequencing. HMW DNA was sheared into an average fragment size between 12–20 kb in a Megaruptor 3 system with speed setting 30. Sheared DNA was purified by solid-phase reversible immobilisation using AMPure PB beads with a 1.8X ratio of beads to sample to remove the shorter fragments and concentrate the DNA sample. The concentration of the sheared and purified DNA was assessed using a Nanodrop spectrophotometer and Qubit Fluorometer and Qubit dsDNA High Sensitivity Assay kit. Fragment size distribution was evaluated by running the sample on the FemtoPulse system.
RNA was extracted from jaDiaLine7 in the Tree of Life Laboratory at the WSI using TRIzol, according to the manufacturer’s instructions. RNA was then eluted in 50 μl RNAs-free water and its concentration RNA assessed using a Nanodrop spectrophotometer and Qubit Fluorometer using the Qubit RNA Broad-Range (BR) Assay kit. Analysis of the integrity of the RNA was done using Agilent RNA 6000 Pico Kit and Eukaryotic Total RNA assay.

**Sequencing**

Pacific Biosciences HiFi circular consensus and 10X Genomics Chromium read cloud sequencing libraries were constructed according to the manufacturers’ instructions. Sequencing was performed by the Scientific Operations core at the Wellcome Sanger Institute on Pacific Biosciences SEQUEL II (HiFi), Illumina NovaSeq 6000 (10X) and Illumina HiSeq 4000 (RNA-Seq) instruments. Hi-C data were generated in the Tree of Life laboratory from remaining tissue of jaDiaLine7 using the Arima v2 kit and sequenced on a NovaSeq 6000 instrument.

**Genome assembly**

Assembly was carried out with Hifiasm (Cheng et al., 2021); haplotypic duplication was identified and removed with purge_dups (Guan et al., 2020). One round of polishing was performed by aligning 10X Genomics read data to the assembly with longranger align, calling variants with freebayes (Garrison & Marth, 2012). The assembly was then scaffolded with Hi-C data (Rao et al., 2014) using SALSA2 (Ghurye et al., 2019). The assembly was checked for contamination as described previously (Howe et al., 2021). Manual curation was performed.
Figure 4. Genome assembly of *Diadumene lineata*, jaDiaLine6.1: cumulative sequence. BlobToolKit cumulative sequence plot. The grey line shows cumulative length for all scaffolds. Coloured lines show cumulative lengths of scaffolds assigned to each phylum using the buscogenes taxrule. An interactive version of this figure is available at https://blobtoolkit.genomehubs.org/view/jaDiaLine6.1/dataset/CAKKNV01/cumulative.

Figure 5. Genome assembly of *Diadumene lineata*, jaDiaLine6.1: Hi-C contact map. Hi-C contact map of the jaDiaLine6.1 assembly, visualised in HiGlass. Chromosomes are arranged in size order from left to right and top to bottom. The interactive Hi-C map can be viewed at https://genome-note-higlass.tol.sanger.ac.uk/?d=bZPq5k_oTFuM-wZyPAZdXg under track jaDiaLine6.
using HiGlass (Kerpedjiev et al., 2018) and Pretext. The mitochondrial genome was assembled using MitoHiFi (Uliano-Silva et al., 2021), which performs annotation using MitoFinder (Allio et al., 2020). The genome was analysed and BUSCO scores generated within the BlobToolKit environment (Challis et al., 2020). Table 3 contains a list of all software tool versions used, where appropriate.

**Ethics/compliance issues**
The materials that have contributed to this genome note have been supplied by a Darwin Tree of Life Partner. The submission of materials by a Darwin Tree of Life Partner is subject to the Darwin Tree of Life Project Sampling Code of Practice. By agreeing with and signing up to the Sampling Code of Practice, the Darwin Tree of Life Partner agrees they will meet the legal and ethical requirements and standards set out within this document in respect of all samples acquired for, and supplied to, the Darwin Tree of Life Project. Each transfer of samples is further undertaken according to a Research Collaboration Agreement or Material Transfer Agreement entered into by the Darwin Tree of Life Partner, Genome Research Limited (operating as the Wellcome Sanger Institute), and in some circumstances other Darwin Tree of Life collaborators.

**Table 2. Chromosomal pseudomolecules in the genome assembly of Diadumene lineata, jaDiaLine6.1.**

| INSDC accession | Chromosome | Size (Mb) | GC% |
|-----------------|------------|-----------|-----|
| OU974069.1      | 1          | 42.01     | 35.3|
| OU974070.1      | 2          | 33.73     | 35.3|
| OU974071.1      | 3          | 23.14     | 35.6|
| OU974072.1      | 4          | 19.48     | 35.2|
| OU974073.1      | 5          | 18.71     | 35.4|
| OU974074.1      | 6          | 18.65     | 35.3|
| OU974075.1      | 7          | 17.67     | 35.4|
| OU974076.1      | 8          | 15.82     | 35.3|
| OU974077.1      | 9          | 15.30     | 35.3|
| OU974078.1      | 10         | 14.93     | 35.3|
| OU974079.1      | 11         | 14.14     | 35.2|
| OU974080.1      | 12         | 14.10     | 35.2|
| OU974083.1      | 13         | 13.13     | 35.1|
| OU974081.1      | 14         | 13.25     | 35.3|
| OU974082.1      | 15         | 13.16     | 35.4|
| OU974084.1      | 16         | 12.63     | 35.0|
| OU974085.1      | MT         | 0.02      | 37.4|
| -                | Unplaced   | 13.14     | 33.4|

**Table 3. Software tools used.**

| Software tool | Version | Source |
|---------------|---------|--------|
| Hifiasm       | 0.15.3-r339 | Cheng et al., 2021 |
| purge_dups    | 1.2.3   | Guan et al., 2020 |
| SALSA2        | 2.2     | Ghury et al., 2019 |
| longranger    | 2.2.2   | https://support.10xgenomics.com/genome-exome/software/pipelines/latest/advanced/other-pipelines |
| freebayes     | 1.3.1-17-gaa2ace8 | Garrison & Marth, 2012 |
| MitoHiFi      | 2.0     | Uliano-Silva et al., 2021 |
| HiGlass       | 1.11.6   | Kerpedjiev et al., 2018 |
| PretextView   | 0.2.x   | https://github.com/wtsi-hpag/PretextView |
| BlobToolKit   | 3.0.5   | Challis et al., 2020 |

**Data availability**
European Nucleotide Archive: Diadumene lineata (orange-striped anemone). Accession number PRJEB46855; https://identifiers.org/ena.embl/PRJEB46855. The genome sequence is released openly for reuse. The *D. lineata* genome sequencing initiative is part of the Darwin Tree of Life (DTol) project. All raw sequence data and the assembly have been deposited in INSDC databases. The genome will be annotated using the RNA-Seq data and presented through the Ensembl pipeline at the European Bioinformatics Institute. Raw data and assembly accession identifiers are reported in Table 1.

**Author information**
Members of the Marine Biological Association Genome Acquisition Lab are listed here: https://doi.org/10.5281/zenodo.5913830.

Members of the Darwin Tree of Life Barcoding collective are listed here: https://doi.org/10.5281/zenodo.5744972.

Members of the Wellcome Sanger Institute Tree of Life programme are listed here: https://doi.org/10.5281/zenodo.6125027.

Members of Wellcome Sanger Institute Scientific Operations: DNA Pipelines collective are listed here: https://doi.org/10.5281/zenodo.5746904.

Members of the Tree of Life Core Informatics collective are listed here: https://doi.org/10.5281/zenodo.5638618.

Members of the Darwin Tree of Life Consortium are listed here: https://doi.org/10.5281/zenodo.5638618.
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Uliano-Silva M, Nunes JGF, Krasheninnikova K, et al.: marcelauliano/MitoHiFi: mitohifi_v2.0. 2021. Publisher Full Text
Kun Wang  
School of Ecology and Environment, Northwestern Polytechnical University, Xi’an, China

1. "it is now established on almost every temperate and tropical coast worldwide" might be overstated?  
2. "Under suitable conditions, it can quickly form large clonal aggregations." Should be more specified.  
3. "of the assembly sequence was assigned to 11 chromosomal-level scaffolds, representing 16 autosomes" is confusing.  
4. The range of DNA amount (0.01–0.5 ng) is unusually low for fragment size analysis. Should be explained.

**Is the rationale for creating the dataset(s) clearly described?**  
Yes

**Are the protocols appropriate and is the work technically sound?**  
Yes

**Are sufficient details of methods and materials provided to allow replication by others?**  
Yes

**Are the datasets clearly presented in a useable and accessible format?**  
Yes

*Competing Interests*: No competing interests were disclosed.

*Reviewer Expertise*: Evolutionary genetics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Overall response: Generating genomic resources for this species is a great step forward, which will facilitate a range of research for *D. lineata*, specifically, and sea anemones more broadly as genomic resources are not widespread at present.

We provide some feedback below on the background, and to a certain extent the justification for sequencing this specific taxon, and on the methodology.

First, we provide some feedback that is geared toward improving the background information so that interested readers have a roadmap to the current literature on this species.

1. Both Flenniken (2017) and Glon et al. (2021) provided excellent background information on the invasion history and invasion-facilitating characteristics of *D. lineata*.

2. The link connected with the statement “Under suitable conditions, it can quickly form large clonal aggregations” leads to a non-functioning page. However, this statement could be supported with primary citations, such as:

   Shick (1976) which documented the occurrence of a high-density population, presumably of a single clone, in Blue Hills, Maine.

   Shick and Lamb (1977) which discussed the general tendency of *D. lineata* (under the former name *Haliplanella luciae*) to form clonal aggregations and provided allozyme evidence of clonal aggregations from a collection of sites.

   Ryan et al. (2021) which provided evidence with microsatellite markers of highly clonal population genetic structures from eight sites across the Atlantic Coast of the US.

3. Gamete production and mixed sex populations are now well established in the non-native range, though fertilization and larval settlement have not been confirmed in the field:

   Ryan and Miller (2019) demonstrated that gamete production is common across the range of *D. lineata* on the Atlantic Coast of the United States, including many populations with fertile males and females present.
Newcomer et al. (2019) also found a population with fertile males and females coexisting on the Pacific Coast of the United States.

4. Gamete production in *D. lineata* has been documented in Japan (within the presumed native range) by Fukui (1991), Fukui (1995), and Ryan and Kubota (2016), but not Ryan and Miller (2019).

5. Asexual reproduction in *D. lineata* has been documented in Japan by several authors, including Uchida (1932, 1936) and Atoda (1973), but not Ryan and Miller (2019).

6. It might also be useful to acknowledge that literature on this species has been published under many species names over time. Hancock et al. (2017) gives a particularly nice overview of the taxonomic history and documented occurrences.

**Second, we provide some feedback on the methodology to generate the genomic data.**

1. In Figure 1, perhaps circle the specific specimen to which you are referring as there are multiple individual anemones in the photograph. Perhaps even cropping the image would be useful.

2. More information on the manual curation would be nice to determine exactly what was done. For example, was there any bacterial removal from the assembly?

3. Was RNA extracted from the whole organism?

4. Were both anemones – jaDialine6 and jaDialine7 – sterile? Also, when were they collected (date?). The authors mention ‘unknown sex’ in the Genome sequence report section, but a little bit more information would be useful. Also, this is presumably from the sample used for HiC rather than RNA? Two anemones were used for all sequencing, so clarifying this is also useful.

5. Continuing from the point above, more information on what specimens were sequenced would be useful.

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Is the rationale for creating the dataset(s) clearly described?
Partly

Are the protocols appropriate and is the work technically sound?
Yes

Are sufficient details of methods and materials provided to allow replication by others?
Partly

Are the datasets clearly presented in a useable and accessible format?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Evolutionary ecology, population genetics, sea anemone biology

We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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This Genome Note provides a useful review of the genome assembly and annotation efforts for
**Diadumene lineata.** As there is currently a dearth of genome assemblies from the Cnidaria and particularly chromosome-level assemblies (perhaps only for the recent assemblies of *Nematostella vectensis* and *Scolanthus callimorphus* (Zimmerman et al. 2022),¹ the addition of this assembly from *Diadumene lineata* is welcome. The genome size (313Mb) and chromosome number (N=16) is consistent with those of *N. vectensis* (244Mb, 15 chromosomes) and *S. callimorphus* (414Mb, 15 chromosomes) (Zimmerman et al. 2022).¹ It would be interesting to have further details of the manual curation that led to the welcome improvement in scaffold number and scaffold N50, and also useful to know what tissues were sampled for the RNASeq dataset. Presumably, this is the whole organism? However, the resources detailed within will be extremely useful for the study of cnidarian genomics and beyond.

**References**
1. Zimmermann B, Robb S, Genikhovich G, Fropf W, et al.: Sea anemone genomes reveal ancestral metazoan chromosomal macrosynteny. *bioRxiv*. 2020. Publisher Full Text

**Is the rationale for creating the dataset(s) clearly described?**
Yes

**Are the protocols appropriate and is the work technically sound?**
Yes

**Are sufficient details of methods and materials provided to allow replication by others?**
Partly

**Are the datasets clearly presented in a useable and accessible format?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Genomics (invertebrates)

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.