The rising tide of high-quality genomic resources

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Few images are more iconic of coral reef ecosystems than an orange clownfish (Amphiprion percula) nestled among the tentacles of its mutualistic partner, the sea anemone (Figure 1a). Popularized as the Disney character, "Nemo," clownfish are more than a charismatic on-screen presence. Among biologists, they are an ecological and evolutionary research model, shedding light on everything from social organization (Wong, Uppaluri, Medina, Seymour, & Buston, 2016) to mutualisms (Schmiege, D’Aloia, & Buston, 2017). Now, clownfish have yet another reason to be in the spotlight. In this issue, Lehmann et al. (2018) present a chromosome-level genome assembly for A. percula with 908.8 Megabases (Mb) of the assembled sequence placed into 24 chromosomes. The A. percula genome is the third, and most contiguous, in a flurry of clownfish genomes published in 2018 (see also Amphiprion frenatus, Marcionetti, Rossier, Bertrand, Litsios, & Salamin, 2018; Amphiprion ocellaris, Tan et al., 2018). Beyond strengthening future research efforts and allowing for intragenus comparisons, what is most striking about the A. percula genome is its completeness. With a scaffold N50 of 38.4 Mb and 98% of the assembly in chromosomal scaffolds, the A. percula genome is one of the most contiguous fish genomes published thus far, surpassing the high water marks of many model fishes that preceded it. Notably absent from the efforts of Lehmann et al. (2018), however, was the need for a large-scale consortium effort or an assembly approach that required unusually outsized financial resources. Instead, Lehmann et al. (2018) took advantage of this technology, typically by generating large amounts of short-read (~250 base pairs or less) data produced on an Illumina platform and overlaying lower coverage (e.g., ~10–20x) long-read data to form “hybrid” assemblies. Lehmann et al. (2018) flipped this common script by completely focusing their initial efforts on long reads produced on the Pacific Biosciences (PacBio) RS II platform. In total, they generated ~121x coverage of the A. percula genome with PacBio reads which corresponded to ~114 Gigabases (Gb) of sequence data. At an approximate current rate of ~1 Gb of output per run for ~$400 USD, the financial resources needed for this portion of the effort were not trivial, likely

Earlier this decade, reference genomes for the threespine stickleback (Jones et al., 2012) and zebrafish (Howe et al., 2013) were published in Nature, heralding in a new era of genome biology in fishes. As models of evolutionary, ecological and medical research, the assemblies were highly contiguous (scaffold N50, stickleback = ~10.8 Mb, Jones et al., 2012; scaffold N50, zebrafish = ~1.6 Mb, Howe et al., 2013) and empowered an array of studies. (Both genomes have been cited more than 900 times according to Google Scholar.) Of the 27 published chromosome-level fish genomes, the A. percula genome stands alone with ~98% of the assembled genome ordered into chromosomes (Figure 1b, Lehmann et al., 2018). This impressive feat highlights the power of modern genome sequencing and assembly tools when paired with appropriate resources for the task, a thoughtful approach, and a bit of genomic luck (e.g., few difficult to assemble regions).

This raises an important question: How exactly did Lehmann et al. (2018) construct such a high-quality, contiguous genome? First, they took full advantage of sequencing technology that was in its infancy when the zebrafish and stickleback genome projects were underway, namely long-read sequencing [i.e., read lengths in excess of ~10 kilobases; also referred to as “third-generation sequencing,” Hayden (2009)]. Most modern genome sequencing efforts take advantage of this technology, typically by generating large amounts (e.g., >100x coverage) of short-read (~250 base pairs or less) data produced on an Illumina platform and overlaying lower coverage (e.g., ~10–20x) long-read data to form “hybrid” assemblies. Lehmann et al. (2018) flipped this common script by completely focusing their initial efforts on long reads produced on the Pacific Biosciences (PacBio) RS II platform. In total, they generated ~121x coverage of the A. percula genome with PacBio reads which corresponded to ~114 Gigabases (Gb) of sequence data. At an approximate current rate of ~1 Gb of output per run for ~$400 USD, the financial resources needed for this portion of the effort were not trivial, likely
approaching $50,000. However, with the introduction of the PacBio Sequel platform, this scale of long-read data is attainable for a fraction of the cost, perhaps as little as $10,000.

Using a combination of high-coverage PacBio data and a dedicated assembly algorithm, Lehmann et al. (2018) generated 12 "preliminary" assemblies for a range of algorithm settings. Given the contiguity of these initial assemblies—contig N50s ranging from 1.02 to 1.80 Mb with ~97% of benchmarking universal single-copy orthologs included (Simão, Waterhouse, Ioannidis, Kriventseva, & Zdobnov, 2015)—the authors could have stopped there and still offered a new, high-quality assembly for the community. Instead, they went further by incorporating chromatin contact maps generated by Phase Genomics with their ProximoTM Hi-C technology (see Lieberman-Aiden et al., 2009) to organize the *A. percula* genome into a chromosome-scale reference assembly. Alternatively, those seeking a chromosome-scale genome assembly could employ Dovetail’s similar Hi-C/HiRise approach. Either way, chromatin contact mapping takes advantage of the inverse relationship between proximity of nuclear DNA and genomic distance. Physically close stretches of DNA are crosslinked and fragmented, and the ends of the resulting fragments are ligated together. The fragments are then paired-end sequenced with short-read technology. Captured within the sequence data is a frequency distribution of how often two fragments of the genome interact, and thus, how physically close they are to one another. This information allows contigs to be clustered into chromosomal groups and then oriented within chromosomes (see Burton et al., 2013). The contiguity of the *A. percula* assembly dramatically increased following this process, with scaffold N50 rising from 1.9 to 38.1 Mb, or a > 20-fold improvement. In total, >1,000 contigs comprising ~98% of the total assembly length were placed into 24 chromosomal scaffolds. To make the *A. percula* genome easier to use, Lehmann et al. (2018) mirrored larger-scale genome consortia efforts and created the aptly named genome browser, Nemo Genome DB (http://nemogenome.org/).

Beyond methodological curiosity, the *A. percula* assembly adds to a growing body of high-quality eukaryote genomes, unlocking the potential for new insight into general or lineage-specific patterns of genome evolution in fishes. Indeed, with the addition of *A. percula*, there are now six orders of fishes with more than one chromosome-level genome assembly (2–6 species per order; Figure 1b). Such an extensive resource sets the stage for new understanding of how genomic architecture (e.g., gene duplications, rearrangements) has evolved in a globally important and diverse group. For instance, the evolution of novel genes with important adaptive functions can greatly influence the evolutionary trajectory of species and comparisons of larger syntenic regions of interest among species will no doubt clarify the evolutionary processes underlying them.

As genome sequencing and assembly technology have steadily marched forward, it is difficult to specify exactly when generating high-quality genomes became more widely approachable. In their study, Lehmann et al. (2018) clearly showed that we have entered a new realm of eukaryotic genome biology. The authors took a species with no previous genome-scale data and constructed the most contiguous fish genome ever sequenced. In a world of sticklebacks and zebrafish, this was not a trivial feat. While the authors’ efforts and methodological approach should be commended, a portion of their success stemmed from a nexus of technological advances, namely the maturation of long-read sequencing, the rise of methods for placing existing scaffolds into chromosomes and the decreasing costs of both. Indeed, the *A. percula* genome is indicative of a larger shift...
in eukaryote genomics where more and more chromosome-scale assemblies will be offered. And with this rising genomic tide will come new insight into the nature and evolution of genome structure across increasingly large swaths of the tree of life.

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REFERENCES


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