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Practical low-coverage genomewide sequencing of hundreds of individually barcoded samples for population and evolutionary genomics in nonmodel species

NINA OVERGAARD THERKILDSEN1 and STEPHEN R. PALUMBI

Hopkins Marine Station, Department of Biology, Stanford University, 120 Oceanview Blvd., Pacific Grove, CA 93950, USA

Abstract

Today most population genomic studies of nonmodel organisms either sequence a subset of the genome deeply in each individual or sequence pools of unlabelled individuals. With a step-by-step workflow, we illustrate how low-coverage whole-genome sequencing of hundreds of individually barcoded samples is now a practical alternative strategy for obtaining genomewide data on a population scale. We used a highly efficient protocol to generate high-quality libraries for ~6.5 USD from each of 876 Atlantic silversides (a teleost fish with a genome size ~730 Mb) that we sequenced to 1–4× genome coverage. In the absence of a reference genome, we developed a bioinformatic pipeline for mapping the genomic reads to a de novo assembled reference transcriptome. This provides an 'in silico' method for exome capture that avoids the complexities and expenses of using wet chemistry for target isolation. Using novel tools for analysis of low-coverage data, we extracted population allele frequencies, individual genotype likelihoods and polymorphism data for 2 504 335 SNPs across the exome for the 876 fish. To illustrate the use of the resulting data, we present a preliminary analysis of geographical patterns in the exome data and a comparison of complete mitochondrial genome sequences for each individual (constructed from the low-coverage data) that show population colonization patterns along the US east coast. With a total cost per sample of less than 50 USD (including sequencing) and ability to prepare 96 libraries in only 5 h, our approach adds a viable new option to the population genomics toolbox.

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Introduction

DNA sequencing costs have decreased more than 300 000-fold since the turn of the century (Wetterstrand 2015). Yet, despite the increasing affordability of sequencing data, researchers remain faced with decisions about how to distribute sequencing effort between breadth of genomic coverage, depth of coverage per individual and the total number of individuals analysed (Sims *et al.* 2014).

Studies focusing on intraspecific genetic variation often need large sample sizes to accurately estimate population allele frequencies. This need has spurred the development of reduced-representation techniques that focus sequencing on a small fraction of the genome, which can then be sequenced to a sufficient depth for

Correspondence: Nina Overgaard Therkildsen, Fax: 607-255-0349; E-mail: nt246@cornell.edu

¹Present address: Department of Natural Resources, Cornell University, 208 Fernow Hall, Ithaca, NY 14853, USA

reliable genotype calls across many individuals. Approaches targeting the sequence flanking restriction cut sites (such as RAD-seq (Davey *et al.* 2011)) have become the method of choice for many studies on non-model organisms because they provide a relatively cheap and fast way to generate genomewide population genomic data and do not require prior knowledge about the genome sequence, enabling important insights in a diversity of systems (e.g. Davey *et al.* 2011; Narum *et al.* 2013; Andrews *et al.* 2014).

However, a large proportion of RAD markers are typically located outside protein-coding sequence, precluding functional analysis of polymorphism patterns if markers cannot be anchored to an annotated reference genome (Jones & Good 2016). For studies focusing on selection and adaptation, it may be preferable to target protein-coding or promotor regions of the genome (Pespeni *et al.* 2012; De Wit *et al.* 2015). RNA-seq (sequencing of expressed transcripts (Wang *et al.* 2009; De Wit *et al.* 2012)) offers an efficient way to target sequencing to protein-coding regions and obtain

transcriptome-wide polymorphism data, but requires high-quality RNA (De Wit et al. 2012, 2015). A more versatile alternative is targeted sequence capture with hybridization probes (Grover et al. 2012; Jones & Good 2016). This method requires prior knowledge about the target sequence for probe design, but several effective workflows have been developed for nonmodel species (e.g. Cosart et al. 2011; Bi et al. 2012). The key drawback is the high cost of synthesizing custom capture probes, as well as logistics surrounding probe design and laboratory protocols.

As an alternative to sequencing a subset of the genome deeply enough to reliably call individual genotypes, an increasingly popular option is to spread the sequencing over greater parts of the genome or across more individuals. Simulation studies have demonstrated that sampling many individuals at low read depth provides more precise estimates of population parameters than higher read depth for fewer individuals (Buerkle & Gompert 2013; Fumagalli 2013; Nevado et al. 2014). In fact, these studies have suggested that spreading sequencing depth to 1-2 reads per locus and individual (1-2× coverage or less) maximizes the information gained about a population. At such low read depth, individual genotype calls are highly uncertain, so this sequencing design is not suitable for analysis requiring accurate individual genotypes. However, even though most population genetic software packages currently require individual genotypes as input, there is no inherent need for genotypes to be accurately known for many types of analysis of selection, population structure or demographic history. In fact, many commonly used software packages simply collapse genotype data into allele counts for populations (Buerkle & Gompert 2013). New methods and software packages that estimate population-level statistics directly from genotype likelihoods without calling genotypes (and without genotype imputation) offer powerful opportunities for taking advantage of the full information contained in low-coverage sequence data for more accurate population parameter estimates (e.g. Li 2011; Nielsen et al. 2011; Buerkle & Gompert 2013; Fumagalli et al. 2014). Such methods can also estimate individual-based parameters like admixture coefficients and parentage probabilities by combining information across loci to compensate for low coverage at individual loci (e.g. Buerkle & Gompert 2013; Fumagalli et al. 2014; Lindtke et al. 2014; Korneliussen & Moltke 2015).

With decreasing sequencing costs and improved analytical tools, whole-genome resequencing of population samples is becoming a viable strategy for many studies (e.g. Jones et al. 2012; Liu et al. 2014; Xia et al. 2015). Yet so far, most population-level resequencing has been conducted on pools of unlabelled individuals (Pool-seq),

rather than on individually barcoded samples (reviewed by Schlötterer et al. 2014). Undoubtedly, Pool-seg offers the most cost-effective approach for estimating population allele frequencies across the entire genome. Both theoretical and empirical studies have demonstrated that it can generate reliable estimates when the number of individuals in a pool and the sequencing depth are sufficiently high (Futschik & Schlötterer 2010; Zhu et al. 2012; Schlötterer et al. 2014). The obvious downside is that all information about individuals is lost, making it difficult to control for uneven contribution to the pool and precluding any individual-level analysis. Although barcoded sequencing adapters make it possible to multiplex hundreds of labelled samples in a sequencing run, Poolseq has become popular because with traditional methods, it would be very labour-intensive and costly to prepare separate libraries for hundreds of individuals (the cost could easily exceed the costs of sequencing).

Here, we demonstrate a workflow that overcomes these limitations with a library preparation protocol that is sufficiently rapid and inexpensive to allow highthroughput individual sample processing, eliminating the need for pooling unbarcoded samples in many studies. The library preparation protocol was originally developed for small microbial genomes (<15 Mb; Kryazhimskiy et al. 2014; Baym et al. 2015). Our contribution is to demonstrate that it also produces high-quality libraries for a teleost fish with a genome size several orders of magnitude larger (~730 Mb) and that the library fragment lengths can easily be tuned to suit different sequencing applications. With a sequencing depth of only $1-2\times$ per individual, we are able to recover the complete mitochondrial genome sequence for each individual and obtain individual genotype likelihoods across the nuclear genome.

The libraries can be used for whole-genome resequencing analysis when a relevant reference is available for the species under study. However, even though we can now sequence the full genome of any organism, assembling a high-quality reference genome still remains a nontrivial, costly and time-consuming task, especially for highly polymorphic species (Voskoboynik et al. 2013; Ellegren 2014). We therefore also demonstrate an 'in silico' exome capture approach that is a more straightforward starting point for organisms with no prior genomic resources. The idea is to take advantage of our simple sample preparation procedure for shotgun whole-genome sequencing, but then focus the analysis on reads that map to a reference transcriptome, which is easier to assemble de novo than a full genome (see also Lamichhaney et al. 2012 for a Pool-seq version of this approach).

In this article, we describe our step-by-step workflow (Fig. 1) for using this approach to generate exomewide data for the Atlantic silverside, a small estuarine fish

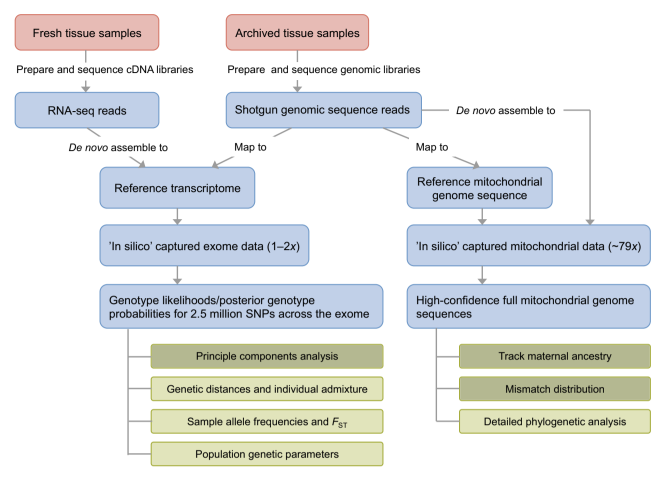


Fig. 1 Diagram illustrating our workflow. The sample input material is marked in red, the procedure for getting from RNA and DNA to the final data sets is marked in blue, and examples of the types of information that can be extracted from the different datasets are shown in green (applications demonstrated in this article are highlighted in darker green).

(730 Mb estimated genome size). With a total of cost of 50 USD per sample (including both library preparation and $1\times$ genome sequencing) and the ability to process 96 samples in only 5 h, our study illustrates that this method will be a viable strategy for obtaining individual-level genomewide data for many organisms.

Materials and methods

Study organism and samples

Our study species is the Atlantic silverside *Menidia menidia*, a small estuarine fish with almost no prior genomic resources. The most closely related species with fully annotated reference genomes in the Ensembl browser are the medaka (*Oryzias latipes*), the platyfish (*Xiphophorus maculatus*) and the tilapia (*Oreochromis niloticus*) with estimated genome sizes ranging from 730 to 927 Mb (golden path length in Ensembl release 82; Cunningham *et al.* 2015). These species all diverged from the silverside more than 100 million years ago (Setiamarga *et al.* 2009;

Near *et al.* 2012), so are unlikely to be good references for mapping genomic silverside reads, hence the need for a species-specific reference.

Our target samples had been stored whole in a $-20\,^{\circ}$ C freezer for between 8 and 17 years. These fish were either collected directly from the wild at one of four different locations along the East coast of North America (from Hice *et al.* 2012) or were generation F1–F11 from a selection experiment started with wild-caught parents collected in New York (from Conover & Munch 2002). Because RNA was unrecoverable from these archived individuals, we collected eight fresh specimens in New York for RNA-seq to generate a reference transcriptome.

Generating a reference transcriptome

To capture a broad diversity of transcripts expressed at different life stages and in different tissue types, we prepared cDNA libraries with RNA from five whole silverside larvae and multiple tissues from three adults, sequenced them in one Illumina HiSeq lane with 100-bp paired-end reads and merged de novo assemblies created from the combined read set with two different programs (CLC Genomics Workbench (http://www. clcbio.com) and Trinity (Haas et al. 2013)). To reduce redundancy in our merged assembly, we metaassembled transcripts based on sequence similarity and used a sequential reciprocal best-hit-BLAST approach to select a single best representative for each nonredundant protein from related fish species with available reference genomes (platyfish, medaka and tilapia). We also added transcripts that contained full open reading frames but had low similarity to the reciprocal-best-BLAST hit contigs.

DNA extraction and library preparation

For the genomic analysis, we used the Qiagen DNeasy Blood and Tissue kit to extract DNA from muscle tissue. We evaluated the degradation level of each extract through 1.5% agarose gel electrophoresis: only samples that showed clear high molecular weight bands and limited smearing were retained for library preparation. To ensure DNA integrity in the retained samples, we removed fragments shorter than ~1000 bp from each extract using Agencourt AMPure XP beads in a 0.4:1 AMPure to sample ratio and eluted the DNA in 10 mm Tris-Cl, pH 8.5. DNA yield and degradation varied between samples, but we were able to obtain good quality DNA extracts from a total of 876 individuals (at least 50 individuals from each population sample).

We measured DNA concentrations with a Quant-iT high-sensitivity assay (Invitrogen) and prepared a separate barcoded library for each individual with Illumina's Nextera kit according to the protocol developed by Kryazhimskiy et al. (2014) (see Baym et al. 2015 for a slightly modified version). Briefly, the tagmentation reaction, which simultaneously fragments the DNA and incorporates partial adapters, was carried out in a 2.5 µL volume with 1.6-7.9 ng of input DNA for each library (this is 1/20 of the reaction volume and DNA input suggested by the manufacturer). We then used a two-step PCR procedure with a total of 12 cycles (8 + 4) to add the remaining Illumina adapter sequence with dual index barcodes and amplify the libraries. The PCR was conducted with the KAPA Library Amplification Kit and the Illumina Nextera index kit with primers N501-N508 + S511 and N701–N712 + N714. As a final step, we purified and size-selected the amplification products with Agencourt AMPure XP beads and quantified the concentration of the final libraries with the Invitrogen Quant-iT high-sensitivity assay (see Table S1, Supporting information for reagent catalogue numbers). We also

examined the fragment size distribution of multiple libraries from each plate on an Agilent Bioanalyzer instrument. Similar to what was shown in the original implementation by Baym et al. (2015), the entire library preparation protocol can be completed for 96 samples in less than 5 h for about 6.50 USD per sample (Table S1, Supporting information), reducing the cost more than 10-fold compared to the regular protocol for Illumina's Nextera kit.

In addition to slightly increasing the amount of input DNA for some samples, our only modifications to the Kryazhimskiy et al. (2014) protocol were to extend the elongation step to 3 min in the initial PCR and to 2 min in the reconditioning PCR (to promote amplification of longer fragments) and to change the size selectivity in the final library purification step. We made these changes because the majority of fragments in our first batch of 76 libraries prepared with the original protocol were shorter than the combined length of the paired reads and the adapter sequence $(2 \times 125 \text{ bp})$ read length + 138 bp sequence = 388 bp, see Results). Using a 0.6:1 AMPure XP beads to library ratio (rather than the 1:1 ratio used in the original protocol), we increased average fragment lengths by almost 100 bp.

We combined equimolar amounts of 56-76 libraries into separate pools for sequencing in 13.5 lanes of paired-end 125-bp reads on an Illumina HiSeq 2000 (v4 chemistry) at the University of Utah's Bioinformatics Core Facility. To even out the data yield among samples, we repooled libraries that initially had obtained the lowest read output for supplementary sequencing in 4.5 additional HiSeq lanes.

To assess the effect of our preparation method on library bias and complexity, we compared our results to two libraries that we had previously prepared with Illumina's TruSeq DNA PCR-Free Sample Preparation Kit. Each of these libraries was prepared according to the manufacturer's instructions for the 550-bp insert size workflow (except using a Branson Sonifier Cell Disruptor 200 for fragmentation instead of a Covaris instrument) with a total of 2 µg input DNA pooled in equimolar amounts from 50 of the same silverside individuals included in the low-coverage set. The two pooled PCR-free libraries were sequenced in 1.5 lanes of paired-end 125-bp reads on an Illumina HiSeq instrument.

Data filtering

Prior to any downstream analysis, we filtered the raw reads to remove potential artefacts and low-quality data. We first removed exact duplicate read pairs (likely caused by PCR amplification) with the program FASTUNIQ v1.1 (Xu *et al.* 2012) and then used TRIMMOMATIC v0.32 (Bolger *et al.* 2014) run in both palindrome and simple mode to trim off adapter sequence. We also used the Trimmomatic sliding window approach to trim off the rest of the read if the average sequence quality over any four bases fell below 15. We used the program FLASH v. 1.2.9 (Magoč & Salzberg 2011) to merge overlapping paired-end reads into single consensus reads and removed sequence that mapped to potential contaminant sources including human, bacterial and viral genomes (between 0.4% and 1.4% of reads per library, Appendix S1, Supporting information).

Mapping

Through a benchmarking test with the program TEASER (Smolka 2015), we identified BOWTIE2 v2.2.3 (Langmead & Salzberg 2012) in preset mode 'very sensitive local' as one of the best-performing mappers for our data set (in a comparison of five commonly used mappers). We therefore used this program and preset mode to map all the genomic reads to the reference transcriptome. We used SAMTOOLS v1.2 (Li et al. 2009) to filter the alignment files, retaining both unpaired, orphaned and concordantly paired reads with a mapping quality >20 but discarding discordantly mapped pairs. The inferred fragment lengths (based on mapping position) revealed a residual presence of overlapping read ends not merged by FLASH for 4–16% of the mapped pairs. To avoid double-counting the sequencing support during SNP calling, we used the clipOverlap program in the BAMUTIL package v1.0.14 (Breese & Liu 2013) to soft clip overlapping read ends (maintaining only the read with the highest quality score in overlapping regions). We removed duplicate reads with the MarkDuplicates module of PICARD Tools v1.139 (http://broadinstitute.github.io/picard/) and summarized the coverage and mapping depth across the reference transcriptome for each individual with the BEDTOOLS COVERAGEBED program v2.19.1 (Quinlan & Hall 2010) and the samtools mpileup module (Li et al. 2009). The outputs from cov-ERAGEBED and MPILEUP were processed with custom scripts to compute average mapping depths after excluding positions with $>4\times$ the mean mapping depth (likely repetitive sequence). We estimated the genome size based on the relationship between the observed mapping depth and the amount of qualityfiltered sequence for each sample. The extent of GC bias was evaluated by computing the average depth of coverage for nonoverlapping 200-bp windows along the entire transcriptome with a script by R. V. Panday (available at http://www.popoolation.at/mauritiana_ genome/). We compared patterns between the merged read set from 50 Nextera-style libraries and the pooled

PCR-free libraries, both downsampled to 50 million mapped reads.

SNP calling and posterior genotype probabilities

Prior to calling variants, we realigned reads around indels with the GATK INDELREALIGNER (McKenna et al. 2010). We then used the program ANGSD v0.910 (Korneliussen et al. 2014) to call single-nucleotide polymorphisms (SNPs) at sites with a probability <1e-6 of being monomorphic based on the mapped reads for all 876 individuals (excluding sites with a total read depth <300 and >3028 (mean depth +2 standard deviations) and bases with a quality score <20). We also used ANGSD to estimate allele frequencies in each population sample of 50 individuals and computed posterior genotype probabilities, which incorporate the uncertainty about true genotypes for each individual at each SNP. The posterior genotype probabilities for all SNPs with a global minor allele frequency >1% were used to estimate the covariance matrix among individuals with NGSTOOLS (Fumagalli et al. 2014), and we used the base package in R for eigen decomposition to summarize the covariance patterns with a principle components analysis.

Mitochondrial genome reconstruction

We mapped the genomic reads to the mitochondrial genome sequence for M. menidia (GenBank Acc. GI: 197311199; Setiamarga et al. 2008) with the same procedure and filtering criteria as described above. We then used freebayes v0.9.21-5-g018c661 (Garrison & Marth 2012) in haploid mode to call variants for each sample, requiring at least three observations of alternative alleles and retaining only nonreference genotype calls with phred-scale quality >20. Based on the haploid genotype calls, we extracted a consensus mitochondrial genome sequence for each individual with the VCF2-FASTA tools from the VCFLIB package (https://github. com/ekg/vcflib). We generated a multiple sequence alignment of these sequences with Clustal Omega (Sievers et al. 2011) and used the R-package haplotypes (Aktas 2015) to compute a pairwise mismatch distribution and quantify the number of unique mitochondrial haplotypes observed within each population. To illustrate the applicability to species without a pre-existing mitochondrial genome reference, we in parallel de novo assembled the mitochondrial genome for our 200 samples collected along the geographical cline with the baiting and iterative mapping approach implemented in the program MITOBIM v.1.8 (Hahn et al. 2013). We used only the commonly available COI barcoding gene sequence as a starting seed and used both the 'denovo' and the 'mapping' mode to iteratively

extend to the full sequence based on matching reads from the quality-filtered read pool from each individual.

Results

Reference transcriptome

The final nuclear transcriptome assembly contained 20 998 contigs with a minimum length of 200 bp, an N50 of 3347 bp, and a combined length of 53.3 Mb. The average GC content was 49.7% (compared to ~40% across all the genomic reads). Overall, 74-78% of RNA-seq reads mapped uniquely onto the assembly and only 1-2% of reads mapped to multiple locations, indicating a low level of redundancy. The contig set shows significant BLASTx hits to 84% of gene models in the related platyfish genome. For 74% of these genes, the top HSP covers >90% of the total length of the reference protein, indicating complete or nearly complete transcripts.

Genomic sequence data quality

The first round of sequencing yielded 0.17-4.16 Gb of raw sequence per library (757 Gb in total). Our followup sequencing added another 229 Gb, resulting in an average of 1.13 Gb raw sequence data (~9 million read pairs) per individual (see Appendix S2 and Fig. S1, Supporting information for details on how we evened out the sequencing effort across individuals). Quality trimming removed on average only 3% of bases, except from one lane with notably lower quality scores, which had an average of >9% of bases removed in the filtering.

The two size selection protocols had substantial impact on the sequence usability. With the original protocol (Kryazhimskiy et al. 2014), the two ends overlapped for 74% of read pairs, with an average overlap of 58 bp, making 22% of the read data redundant after merging. The short length of our inserts also caused adapter read-through in the shortest constructs, so another 8% of the raw data were adapter sequence. The modified protocol substantially increased the mean insert length in our libraries, resulting in a much lower proportion of paired reads overlapping (11-64%, average of 25.4%). This reduced overlap redundancy to 6.3% and the level of discarded adapter sequence to 1.9% (Fig. S2, Supporting information), so that on average, 87% of bases were retained for mapping after all quality filtering. One aspect of our results that bodes well for future use of this approach is that the amount of input DNA did not appear to systematically affect the mean insert length or the total amount of quality-filtered data generated per library (Figs S3–S5, Supporting information).

Mapping

On average, 14% of the reads mapped uniquely to the reference transcriptome with a mapping quality >20. Almost half of these (46%) were mapped in concordant pairs (both ends mapped to the same contig in the expected orientation and with an inferred insert length <1500 bp). Less than 3% of the mapped reads were in discordant pairs with the two ends mapped to different contigs, and these were filtered out. Because we mapped genomic reads to a transcriptome reference that spans intron-exon boundaries, ~37% of all the mapped reads were orphans from pairs where only one end mapped (the other end likely originated from intronic or intergenic sequence that is ignored for the present analysis). Mapping depth tended to only decrease moderately around intron-exon boundaries, however, because the applied mapping algorithm allows local mapping of reads that only match the reference for part of their length (Fig. 2). Indeed, distinct break positions where a large number of reads either start or end their local alignment can be observed in the alignment files (Fig. 2). As also highlighted by Montes et al. (2013), localizing these break positions represents a novel way to infer the location of exon-intron boundaries within the transcriptome reference.

Library complexity, mapping depth and GC bias

Ideal sequencing libraries should contain a large number of unique DNA fragments so that most molecules are sequenced only once, and these fragments should be randomly sampled from the genome for homogenous sequencing coverage. The MarkDuplicates algorithm flagged 3-14% (on average 6%) of our mapped reads as potential duplicates based on identical mapping positions. However, the duplication rates estimated for single-end reads (for which only the mapping position of one end of a fragment is known) were up to 28× greater than the duplication rates estimated for paired reads (for which the mapping position of both ends of a fragment can be taken into account, Fig. 3), indicating that the overall estimates are inflated by false positives (see Appendix S3 and Fig. S6, Supporting information). As expected, the total amount of sequence generated for each library correlated strongly with the duplication rate estimates (Fig. 3). Yet, even in the libraries for which we had 5× genome coverage, the highest paired-read duplication rate estimated was only 2.4% (the inflated estimate including single-end reads was 12%), indicating a high level of complexity and very limited PCR duplication in the libraries.

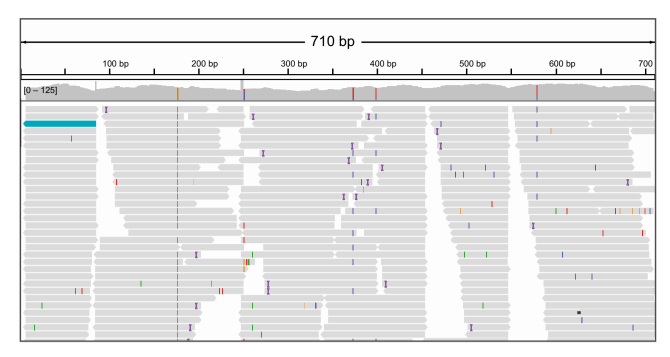


Fig. 2 Example illustration of how the genomic reads from 50 individuals map to a transcriptome contig from a protein-coding gene. The barplot in the top panel shows the total sequencing depth along the 710-bp contig, and each horizontal grey bar underneath shows the mapped portion of a sequencing read. Discontinuities in the mapping boundaries indicate positions where the reads contain data from introns that do not align to the transcriptome contig, thereby denoting intron—exon boundaries. Figure generated with the Integrative Genomics Viewer (Thorvaldsdóttir *et al.* 2013). [Colour figure can be viewed at wileyonlinelibrary.com]

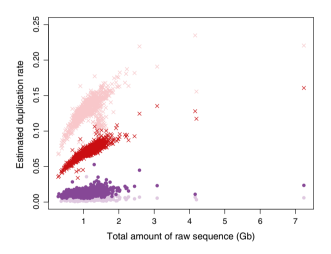


Fig. 3 Scatter plot showing estimated levels of duplication and the total amount of raw sequence generated for each individual. The dark red crosses show the overall duplication rate estimates from MarkDuplicates (paired and single-end reads), the pink crosses show MarkDuplicate estimates based on single-end reads only, the light purple dots show estimates of identical sequence duplicates (from FastUniq), and the dark purple dots show the sum of exact sequence duplicates (FastUniq) and the paired duplicates flagged by MarkDuplicates. We believe that the latter category (dark purple) provides the most reliable estimates of true duplication rates (see text).

About 1.6% of positions in the transcriptome sequence showed excessive mapping depth (greater than four times the mean depth) – likely because of repetitive sequence or assembly errors - and these positions were excluded from all further analysis. For the rest of the transcriptome, the average mapping depth was 1.3× per individual (Fig. S7, Supporting information). Due to the stochastic sampling process involved in sequencing, the reads did not cover the entire reference sequence for each individual. However, within individuals, an average of 66% of positions in the transcriptome were covered by at least one read. In samples with greater than $3.5 \times$ coverage, >90% of positions were covered (Fig. 4). The number of years (8-17) a sample had been stored frozen did not seem to affect the average mapping depth or transcriptome coverage, indicating robustness to variable levels of degradation. Across samples, we observed the highest depth of coverage in regions with GC content between 30 and 50% and reduced coverage in regions with extreme GC content. Because an almost equally strong bias was seen for the PCR-free libraries (Fig. 5b), a large part of this bias most likely arose during sequencing and not in library preparation. Within the range of GC content observed across 99% of the transcriptome (26.5–73.6% GC, Fig 5a,b), the average coverage only varied twofold. The fraction of the reference not covered by

any reads was identical for the PCR-free and the Nextera library sets (2.1%), and although the slightly higher GC bias resulted in a slightly less even distribution of reads than for the PCR-free libraries (Fig. 5c), we consistently saw an average coverage >60× across the population groups of 50 individuals with >82% of bases covered by reads from at least 25 of the 50 different individuals (Fig. S8, Supporting information).

Genome size estimation and per sample sequencing costs

We did not sequence any samples deeply enough to robustly estimate the silverside genome size with k-mer based approaches (Marçais & Kingsford 2011; Chikhi & Medvedev 2014). However, comparisons of the amount of cleaned sequence per individual and the average mapping depth achieved across the reference transcriptome (excluding highly repetitive sites) showed that we needed about 730 Mb of cleaned sequence reads to get an average of 1× mapping depth, indicating that this is the approximate size of the silverside genome (similar to genome sizes reported for related species). Given that our conservative quality filtering discarded about 13% of the raw data, we would need about 840 Mb of raw reads per individual to achieve 1× genome coverage, which means that 65 individuals with a silverside-sized genome can be pooled in a single Illumina lane (assuming 54 Gb

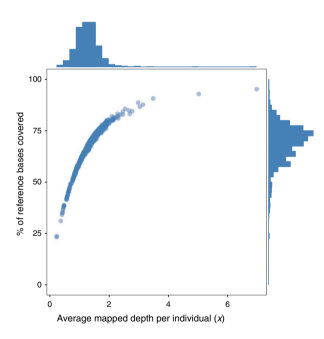
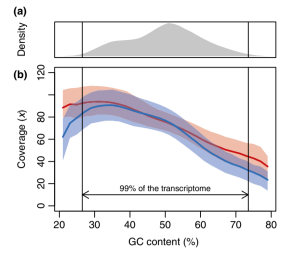


Fig. 4 Scatter plot showing the percentage of the transcriptome reference covered by at least one read as a function of the average mapped depth for each individual. The marginal histograms show the overall distributions of mapped depth per individual (x-axis), and the percentage of the reference covered by each individual (y-axis). [Colour figure can be viewed at wileyonlinelibrary.com]



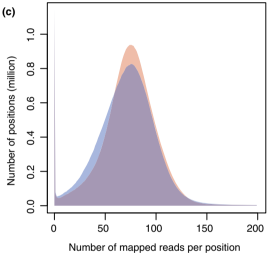


Fig. 5 Comparison of mapping patterns in the PCR-free (red) and the Nextera-based (blue) libraries downsampled to 50 million mapped reads. Panel (a) shows a density plot of GC content in 200-bp windows across the transcriptome reference. Panel (b) shows the mean depth of coverage plotted against GC content in 200-bp windows across the transcriptome for the two library types. The shading represents the 25th and 75th inner quartile regions for each library type, and the vertical black lines delimit the range of GC content observed across 99% of the transcriptome. Panel (c) shows density plots of the depth of coverage per position in the transcriptome for the two kinds of libraries.

of raw sequence output, as we saw on average). This brings the current cost of sequencing to ~40 USD/individual (for 1× genome coverage; see Table S2, Supporting information for a comparison of sequencing costs for organisms with different genome sizes).

SNP calling and posterior genotype probabilities

ANGSD detected a total of 2 504 335 SNPs with a minor allele frequency >1% across all samples. The populationspecific allele frequencies for each of these SNPs estimated from the genotype likelihoods can be used for a suite of downstream applications including scans for signatures of selection and reconstruction of demographic history (N.O. Therkildsen and S.R. Palumbi, in preparation). Despite the low certainty about individual genotypes for each individual SNP (given the only $1-2\times$ coverage), the PCA that integrated genotype probability information across the entire exome clearly grouped individuals into distinct clusters that largely corresponded to geographical origin (Fig. 6). The first principle component separates the geographical samples along a north-south continuum, while PC 2 shows the distinctness of silversides from the Gulf of St. Lawrence, as has been reported from D-loop sequence data (Mach et al. 2010). In addition, the PCA also reveals that a few individuals cluster with a different group than most other individuals collected at the same locations, indicating that they may be migrants. The effects of such divergent individuals would be impossible to quantify in pooled sequencing designs.

Mitochondrial genome reconstruction

On average, 0.2% of the quality-filtered reads mapped to the mitochondrial genome with a mapping quality >20, resulting in an average mapping depth of $79 \times$ along the mitochondrial genome for each individual (range $2 \times -740 \times$). The ratio of mitochondrial to transcriptome mapping depths depended strongly on the amount of time a sample had been stored frozen prior to sequencing (Fig. 7). Samples that had been stored for 17 years showed only about six times greater mapping depth for the mitochondrial genome compared to the transcriptome, while the enrichment was on average 99 times for

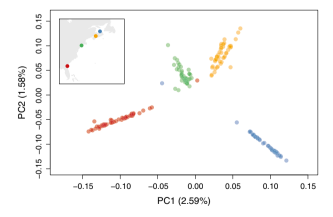


Fig. 6 Clustering of individuals from different sampling locations along the first two principle components (generated from the exome SNP data). Each transparent dot represents an individual sampled from the colour-coded locations indicated on the inset map.

samples stored for only 8 years. This large and consistent difference cannot be attributed to library preparation or sequencing lane effects because samples had been randomized between batches. As mentioned above, we also did not see a similar effect of sample age on the mapping rate for the transcriptome. These results therefore indicate differential rates of degradation of mitochondrial and nuclear DNA in storage.

Among our samples, 784 (98%) of the samples stored for fewer than 17 years had a mapping depth of at least $5\times$ throughout >95% of positions in the mitochondrial genome, allowing high-confidence haploid genotype calls. The multiple alignment of all consensus sequences revealed that 1287 of the 16 458 positions in the mitochondrial genome were variable (651 variants were singletons; that is, the alternative allele was only observed in a single individual). All individuals differed by at least four positions from the reference sequence, but many individuals had completely identical sequences, resulting in only 272 unique haplotypes among the samples. The unique sequences differed by on average 40.7 positions providing strong resolution for tracking of individual lineages.

Among the F1 offspring from wild parents, 89% of haplotypes were unique. By contrast, only 13% of haplotypes were unique in F6 samples after five generations of rearing in the laboratory, and 7% were unique in

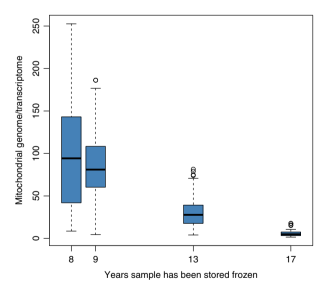


Fig. 7 Boxplot summarizing the ratio between mapping depth at the mitochondrial genome and mapping depth at the transcriptome for samples stored frozen for different lengths of time. The horizontal band in each box represents the median, the top and bottom of the boxes represent the 25th and 75th percentiles, and the error bars define the 5th and the 95th percentiles (data points falling outside these percentiles are marked by dots). [Colour figure can be viewed at wileyonline-library.com]

samples collected after 10 generations of laboratory rearing, illustrating strong reduction of maternal lineages and potential inbreeding.

For the field-collected individuals from the four geographical locations, pairwise mismatch distributions show that the southernmost samples have the most divergent mitochondrial genomes (largest number of polymorphisms in pairwise comparisons), while the mitochondrial genomes sampled at northern locations differ at fewer sites (Fig. 8). This pattern is consistent with northward migration from southern refugia since the last glaciation, as has been previously suggested for this species based on D-loop sequences (Mach et al. 2010). The full-length mitochondrial genomes provide a much richer data set for detailed reconstruction of demographic history of this species.

We also tested whether we could reconstruct mitochondrial genomes without a full reference sequence. Using only COI as a seed and MITObim's default parameters, we were able to de novo assemble the full mitochondrial genome in a single contig for 166 of the 200 geographical samples over 14-157 iterations. Parameter optimization or using a mitochondrial genome sequence from a related species could probably improve the success rate substantially, showing that the low-coverage data make it possible to recover full mitochondrial genomes even without a pre-assembled reference.

Discussion

This study demonstrates that whole-genome sequencing of hundreds of individuals now represents a practical approach for obtaining population genomic data from model and nonmodel species alike. We illustrate an

efficient workflow for preparing high-quality individually barcoded libraries that enable more robust and versatile data analysis than pooled sample sequencing, and we show how 'in silico' capture of genomic reads can be used to generate exomewide polymorphism data for organisms without a reference genome sequence. With a total cost per sample of less than 50 USD (including both library preparation and 1× genome sequencing), our approach offers a viable alternative to RAD-seq, physical target enrichment and Pool-seq for many studies.

High-quality libraries despite low DNA input amounts

The Nextera-based libraries showed only slightly greater GC bias and a slightly less even genomic distribution of reads than previous libraries generated from a subset of the same samples with a PCR-free method, indicating that the cost reduction and simplified workflow do not substantially compromise library quality. We also saw that DNA input amounts as low as 2 ng did not significantly reduce the diversity of sequenced molecules in libraries prepared from the ~730 Mb silverside genome, as we consistently saw duplication rates <2.4%, even in libraries sequenced to $>4\times$ genome coverage. The ability to generate high-complexity libraries with ng amounts of input DNA makes whole-genome sequencing available for a broad set of samples that may not yield sufficient DNA for alternative methods that require as much as 1–2 μg of DNA.

Robustness to DNA degradation

The efficiency with which DNA is incorporated into the library probably depends to some degree on DNA

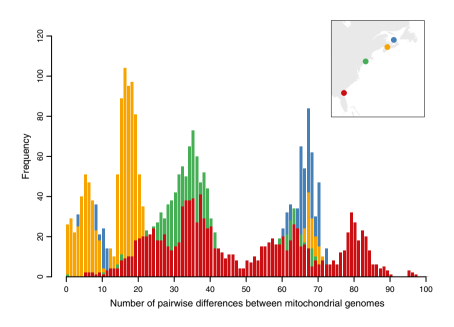


Fig. 8 Mismatch distributions showing the number of pairwise nucleotide differences between the full mitochondrial genome sequences of silversides samples from four different locations along North America's east coast (the colour-coded sampling locations are shown on the inset map).

quality. For nuclear DNA, we saw no consistent differences in the quality of libraries generated with samples stored in a minus 20 °C freezer for 8-17 years. We did, however, only use DNA extracts that contained high molecular weight DNA, and used a precleaning procedure to remove fragments shorter than 1000 bp prior to DNA quantification. The notable decrease in enrichment for mitochondrial DNA with sample age suggests faster decay of mitochondrial vs. nuclear DNA under our sample storage conditions (whole fish frozen at −20 °C). This faster decay is surprising considering that mitochondrial fragments are often the only recoverable DNA from ancient samples and have been reported to decay at slower rates than nuclear DNA over long time scales (hundreds to thousands of years; Schwarz et al. 2009; Allentoft et al. 2012). Several other studies have reported faster decay of mitochondrial than nuclear DNA in experimental comparisons, however, indicating that different processes may affect short- and long-term DNA preservation (Foran 2006; Higgins et al. 2015).

Efficient workflow with easy tuning for different sequencing applications

The entire library preparation protocol can be completed for 96 samples in less than 5 h total time, and without expensive specialized laboratory equipment. We found that simply altering the ratio of AMPure XP beads to library volume caused consistent shifts in the size distribution of fragments in the final library, thereby allowing flexible tuning to different sequencing applications. To minimize redundancy from sequence overlap and adapter read-through, library fragments should be longer than the total read length (the sum of both ends for paired reads) plus the adapters. However, fragments larger than 600 bp tend to cluster much less efficiently than fragments in the 250- to 500-bp range on Illumina flow cells (Bronner et al. 2013). This resulted in lower sequence yields for libraries with longer fragments, so the optimal balance between data loss and data yield was reached for libraries with inserts between 250 and 350 bp (Fig. S5, Supporting information).

Contrary to several previous studies (Adey *et al.* 2010; Baym *et al.* 2015) and Illumina's Nextera Technical Note, we did not find that variation in the amount of input DNA (in the range 1.6–7.9 ng) predictably affected the fragment length distribution of the libraries. A similar result has been reported in another study optimizing the Nextera protocol for a range of different organisms (Lamble *et al.* 2013), suggesting that exact normalization prior to library preparation – currently the most time-consuming part of the protocol – may not always be necessary and can perhaps be

replaced with a bead normalization (Lamble *et al.* 2013). However, as the effect of input concentration is likely to vary between organisms (depending, for example, on genome composition, GC content and sample quality), we echo the recommendation of Baym *et al.* (2015) to always calibrate input amounts and size selection conditions on dilution series of a few representative samples before scaling up library preparation for novel DNA sources.

Individual barcoding vs. pooled sequencing

Our study shows that the price difference between pooling unlabelled DNA into a single library and preparing separate barcoded libraries for each individual is diminishing. Our library preparations cost ~6.5 USD per sample, so the total reagent cost for individual barcoding of 50 individuals is about ~325 USD, a modest portion of an overall NGS sequencing budget. Considering furthermore that we can prepare 96 individual libraries in the same amount of time it takes to make a pooled library, we believe that our approach is practical for broad adoption. The slightly higher cost compared to Pool-seq should for many studies be well worth the much increased versatility of individual-level low-coverage data compared to an anonymous pool of sequences.

A key advantage of individually barcoded data is the ability to account for uneven sequencing depth among individuals. This is particularly important when degradation levels vary between samples, as was the case for our silversides. Perhaps more important, however, is the suite of additional individual-level analysis made possible. Novel tools that integrate signals from low-confidence genotype likelihoods across thousands of nuclear loci now allow accurate inference about relatedness and genetic similarities among individuals even at 1-2× coverage (e.g. Fumagalli 2013; Gompert et al. 2014; Korneliussen et al. 2014; Korneliussen & Moltke 2015; Vieira et al. 2015; Snyder-Mackler et al. 2016). As illustrated here with the PCA, this individual-level analysis can help detect cryptic differentiation within a population sample. When undetected in Pool-seq data, such heterogeneity among individuals can dramatically bias population allele frequency estimates. Estimation of individual admixture proportions also makes certain types of individual clustering possible with low-coverage data (Skotte et al. 2013), so that analyses of population structure are not limited to predefined sample groupings. In a similar vein, individually barcoded samples are not limited to a single pool. Samples can be regrouped in many different configurations, for example according to different phenotypic traits for association studies (Kim et al. 2011; Skotte et al. 2012). This allows for flexible analysis of both allele frequencies and patterns of linkage disequilibrium across

the genome (Li 2011; Maruki & Lynch 2014). We foresee a rapid development in tools for analysing sequencing data in a probabilistic framework without calling genotypes over the next years, which should enable even more powerful analysis from low-coverage sequencing.

Another 'added bonus' of barcoding individuals is the high-confidence full mitochondrial genome sequence we were able to recover for each individual, allowing powerful inference about maternal ancestry and evolutionary history. As utilized in genome skimming projects (Straub et al. 2012; Dodsworth 2015), high copy numbers per cell typically ensure deep sequencing of the mitochondrial genome even when the nuclear genomewide coverage is shallow. The average of 79× coverage we observed across the mitochondrial genome for each individual allowed for both de novo assembly of the full mitochondrial sequence and polymorphism detection based on mapping to a pre-existing reference. The resulting full-length mitochondrial genome sequences, representing nonrecombining, maternally inherited markers, provide valuable complementary information to nuclear data at a resolution not achievable with earlier techniques. A previous study that examined a 340-bp fragment of the D-loop in 1029 silversides found the same haplotype in 66% of their samples and most other haplotypes differing only by a single substitution (Mach et al. 2010). The full mitochondrial genomes of our 200 individuals collected over the same geographical cline differed on average by >40 nucleotides (Fig. 8). At a divergence rate of 2% per million years, two 16 500-bp mitochondrial genomes will differ by 1 change per 3000 years. As a result, full mtDNA data sets have the possibility of capturing demographic events that have occurred during the last several glacial cycles. Such changes may be important in temperate species affected by swings in latitudinal range. In our data set, there is a strong peak in northern populations at an average of 6bp differences, representing perhaps invasion of these northern populations over the last 18 000 years as glaciers receded. Peaks in southern populations at 18, 32 and 66 differences perhaps suggest lineage divergence that occurred 50 000, 90 000 and 200 000 years ago. More detailed recent historical demography might be mined from these kinds of data sets.

Expanding the tool box for both model and nonmodel species genomics

As discussed above, dividing a given sequencing effort over individually barcoded samples should provide substantially more robust and versatile data than sequencing a pooled sample to the same combined depth. Our cheap and efficient library preparation method should therefore be useful for many population whole-genome

resequencing studies of species with full reference genome sequences.

Our 'brute force' approach to obtaining exome and mitochondrial sequence for a species without a reference genome may seem inefficient as we only end up using ~8% of the data. However, the cost of nontarget sequencing must be weighed against the cost and effort associated with isolating the target parts of the genome prior to sequencing. Although several workflows have been developed for designing capture probes for nonmodel species (reviewed by Gasc et al. 2016 and Jones & Good 2016) and methodological improvements along with new commercial providers have made capture technology more affordable, the cost of custom-synthesizing probes the entire transcriptome for hundreds of individuals is still relatively high. Although it is currently somewhat more expensive to capture 'in silico' rather than with physical baits for many applications, the benefits include a much simpler and faster high-throughput sample preparation method that introduces less bias, and, importantly, the availability of all the nontarget genomic reads for later analysis. Because our approach is based on shotgun genome sequencing, the sequencing cost per sample will scale with the genome size of the study organism, so the relative advantages of 'in silico' vs. physical capture will depend strongly on the genome size of the organism under study (see Table S2, Supporting information for an comparison of the per individual cost of our approach for organisms with different genome sizes). However, as sequencing costs further decrease, the set of organisms for which our approach is cost-effective will expand.

For some time to come, our method will be more expensive than RAD-seq for many studies. However, the ability to focus on specific parts of the genome adds a valuable functional dimension to data sets. Our approach also does not suffer from the technical artefacts associated with polymorphisms in restriction cut sites that in RAD-seq can reduce overlap in the loci sequenced in different individuals and cause allele dropout that may complicate and bias inferences, especially when the genetic diversity within species is high (Gautier et al. 2012; Arnold et al. 2013). We therefore believe that 'in silico' capture adds a new promising tool to the nonmodel species population genomics toolbox. We have shown here for the silverside how it can be incorporated into a workflow for species with no prior genomic resources and can powerfully address both individual-level and population-level questions. For applications where high-confidence individual genotypes are required, however, other methods will be more appropriate, so method

selection should always depend on particular study objectives.

Future potential

We here demonstrate a workflow focused on 'in silico' exome capture with a de novo assembled transcriptome reference as a tractable and straightforward starting point for population genomic analysis. As the raw data cover the whole genome, however, many other regions of interest can be targeted (e.g. microRNAs, transcription factors or transposable elements). An alternative strategy could also be to de novo assemble the genomic reads to the extent possible with only low-coverage sequencing of polymorphic individuals, potentially using the transcriptome reference contigs as seeds for assembling genomic regions flanking the (Lamichhaney et al. 2012; Ruby et al. 2013). The resulting longer contigs can then be used as a more extensive reference for mapping, covering a larger part of the genome (but also complicating the analysis pipeline). Reference-free variant calling methods also offer promising prospects (Iqbal et al. 2012; Leggett & MacLean 2014; Uricaru et al. 2015), and with the increasing availability of long sequence reads and methods for improving assembly contiguity, it will become easier and faster to develop full genome reference sequences for new species, making low-coverage whole-genome resequencing data even more powerful. Our exome-focused approach allows a short cut to start exploring genomic patterns in versatile sequence data that will have a long-lasting value and enduring applicability as reference sequences improve.

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The authors conceived of and designed the study together. N.O.T. performed the experiments and analysed the data with input from S.R.P., and the authors wrote the manuscript together.

Data accessibility

The Atlantic silverside transcriptome is deposited in the NCBI GenBank Transcriptome Shotgun Assembly Sequence Database (TSA) under Accession no. GEVY00000000.

A table with detailed statistics on all steps in the quality filtering and mapping procedures is available from the Dryad Digital Repository: doi:10.5061/dryad.ft596. This repository also includes a list of the command line arguments used for running each step in the pipeline.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Appendix S1 Contamination filtering.

Appendix S2 Strategy for evening out sequencing effort among individuals.

Appendix S3 Interpreting duplication rate estimates.

Table S1 Breakdown of reagent costs for the library preparation protocol.

Table S2 Estimated per sample cost of our 'in silico' exome capture protocol for organisms with different genome sizes.

Fig. S1 Scatter plot showing the targeted percentage of the total data yield in a lane for each re-sequenced library (x-axis) vs. the observed percentage of the total data yield generated from this library (y-axis).

Fig. S2 Scatter plot showing the mean insert length (inferred from mapping positions) and the percentage of raw data lost to read overlap and adapter read-through for libraries prepared with the original Kryazhimskiy *et al.* protocol (orange), and our modified protocol (green).

Fig. S3 Scatter plots showing the DNA input amount plotted against the mean insert length (a), the percentage of the raw data retained after quality filtering (b), and the amount of raw sequence data generated in the first round of sequencing (c) for each library.

Fig. S4 Scatter plot showing the input concentration plotted against the total amount of quality-filtered data generated from each library in the first round of sequencing.

Fig. S5 Scatter plots showing the mean insert length plotted against the amount of raw sequence data generated in the first round of sequencing (a) and the percentage of the raw data retained after quality filtering (b) for each library.

Fig. S6 Scatterplot showing duplication rate estimates plotted against the DNA input amount for each library.

Fig. S7 A density plot of the average mapped depth across the transcriptome (excluding sites with >4x the mean depth) per sample achieved in the first round of sequencing (yellow) and after the supplemental sequence had been added (blue).

Fig. S8 Typical distributions of combined mapping statistics for population samples of 50 individuals.