Draft Genome of the Common Snapping Turtle, *Chelydra serpentina*, a Model for Phenotypic Plasticity in Reptiles

Debojyoti Das,*1 Sunil Kumar Singh,*1 Jacob Bierstedt,* Alyssa Erickson,* Gina L. J. Galli,† Dane A. Crossley, II,* and Turk Rhen*,2

*Department of Biology, University of North Dakota, Grand Forks, North Dakota 58202, †Division of Cardiovascular Sciences, School of Medical Sciences, University of Manchester, Manchester M13 9NT, UK, and 2Department of Biological Sciences, University of North Texas, Denton, Texas 76203

**ABSTRACT** Turtles are iconic reptiles that inhabit a range of ecosystems from oceans to deserts and climates from the tropics to northern temperate regions. Yet, we have little understanding of the genetic adaptations that allow turtles to survive and reproduce in such diverse environments. Common snapping turtles, *Chelydra serpentina*, are an ideal model species for studying adaptation to climate because they are widely distributed from tropical to northern temperate zones in North America. They are also easy to maintain and breed in captivity and produce large clutch sizes, which makes them amenable to quantitative genetic and molecular genetic studies of traits like temperature-dependent sex determination. We therefore established a captive breeding colony and sequenced DNA from one female using both short and long reads. After trimming and filtering, we had 209.51Gb of Illumina reads, 25.72Gb of PacBio reads, and 21.72 Gb of Nanopore reads. The assembled genome was 2.258 Gb in size and had 13,224 scaffolds with an N50 of 5.59Mb. The longest scaffold was 27.24Mb. BUSCO analysis revealed 97.4% of core vertebrate genes in the genome. We identified 3.27 million SNPs in the reference turtle, which indicates a relatively high level of individual heterozygosity. We assembled the transcriptome using RNA-Seq data and used gene prediction software to produce 22,812 models of protein coding genes. The quality and contiguity of the snapping turtle genome is similar to or better than most published reptile genomes. The genome and genetic variants identified here provide a foundation for future studies of adaptation to climate.
on the gonads, incubation temperature has a significant impact on growth, physiology, and behavior in turtles and other reptiles (Rhen and Lang 2004, Noble et al. 2018, While et al. 2018, Singh et al. 2020).

Temperature effects are a specific example of a broader phenomenon called phenotypic plasticity in which environmental factors alter phenotype (Via and Lande 1985, Scheiner 1993, Agrawal 2001, Angilletta 2009, Warner et al. 2018). Organisms can also maintain phenotypic stability in the face of variable environments. Physiologists call this homeostasis while developmental biologists call it canalization. Although plasticity and stability appear to be distinct strategies for dealing with environmental variation, they actually represent ends of a continuum of potential responses. Plasticity/stability often has a genetic basis with different individuals being more or less responsive to environmental influences. We must decipher genome-environment interactions to understand the role plasticity/stability plays in allowing turtles to survive and reproduce in diverse climates from the tropics to temperate regions.

Genomic resources will facilitate research on the evolution of phenotypic plasticity, homeostasis, and developmental canalization in turtles. To date, genomes from six turtle species in five families have been sequenced (Shaffer et al. 2013, Wang et al. 2013, Tolls et al. 2017, Cao et al. 2019, https://www.ncbi.nlm.nih.gov/bioproject/PRJNA415469/), but each of these species lives and reproduces in a much narrower range of climates than the common snapping turtle (Chelydra serpentina). Here we assemble and annotate the first draft genome for the snapping turtle, which is the most widespread and abundant species in the family Chelydridae.

The contiguity and completeness of the snapping turtle genome is similar to or better than other reptiles and is adequate for reuse in functional and comparative genomic studies. Several characteristics make the snapping turtle a good model for turtle biology. This species is one of the most extensively studied turtles (Steyermark et al. 2008, While et al. 2018), providing a wealth of baseline information for genetic, genomic, epigenomic, and transcriptomic analyses of cell and developmental biology, physiology, behavior, ecology and evolution. This species produces large clutches (30-95 eggs/clutch) and is easy to breed and rear in captivity, making genetic studies feasible. We therefore established a captive breeding colony to study phenotypic variation in TSD (Janzen 1992, Rhen and Lang 1998, Ewert et al. 2005, Rhen et al. 2015, Schroeder et al. 2016). Controlled breeding reveals that variation in TSD within populations is highly heritable and that population differences in sex ratio at warm incubation temperatures are also heritable (K. Hilliard and T. Rhen, unpublished results). Yet, population differences in sex ratio at cool incubation temperatures are due to genetic dominance and/or non-genetic maternal effects, illustrating genome-environment interactions (K. Hilliard and T. Rhen, unpublished results). These findings provide a solid foundation for genome-wide association studies to identify specific loci that influence thermosensitivity (Schroeder et al. 2016).

The genome will also be useful for studying other ecologically important traits and characterizing population genomic variation. Such studies will provide insight into genetic adaptation to climate because snapping turtles range from tropical to northern temperate zones. For example, snapping turtles display counter-gradient variation in developmental rate with latitude: northern alleles speed embryonic developmental rate to counteract the impact of cooler soil temperatures at higher latitudes (K. Hilliard and T. Rhen, unpublished results). Another remarkable trait is their ability to tolerate hypoxic conditions. Eggs buried underground periodically experience low oxygen conditions (e.g., when soil is saturated with water after heavy rains). Hypoxia during embryogenesis programs subsequent performance in low oxygen environments: cardiomyocytes from juvenile snapping turtles exposed to hypoxia as embryos have enhanced myofilament Ca2+-sensitivity and ability to curb production of reactive oxygen species when compared to juveniles exposed to normoxic conditions as embryos (Ruhr et al. 2019). Such findings have broader implications for understanding cardiac hypoxia tolerance/susceptibility across vertebrates: i.e., most human diseases of the heart are due to insufficient oxygen supply. A contiguous, well-annotated genome is critical for epigenomic studies of developmentally plastic responses to temperature and oxygen levels as well as other abiotic factors and ecological interactions. For instance, future studies will correlate genome-wide patterns of DNA methylation with transcriptome-wide patterns of gene expression in hearts of juvenile turtles exposed to hypoxic conditions as embryos.

The genome will also be valuable for comparative studies with other Chelydridae, which are listed as vulnerable on the ICUN Red List (2017): Macrochelys temminckii in North America, Chelydra rossignoni in Central America, and Chelydra acutirostris in South America. Finally, we expect this draft will serve as a template for refinement and improvement of the snapping turtle genome assembly.

**MATERIALS AND METHODS**

**Animal husbandry**

Adult snapping turtles were captured by hand, with baited hoop nets, and during fish surveys in the state of Minnesota (MN) and transported to the University of North Dakota (UND) to establish a captive breeding colony for genetic analysis of TSD. Turtles were collected across the state of MN from the Canadian border in the north to the Iowa border in the south, which spans a 5° latitudinal range. Turtles in the colony are housed year-round in the animal quarters at UND in conditions that mimic seasonal changes in photoperiod and water temperature in MN.

Two rooms are set up with seven stock tanks per room (14 total tanks). Turtles are held in 1136-liter stock tanks (2.3 m long × 1.9 m wide × 1.6 m deep) filled with roughly 850 liters of water. One male is housed with 3 or 4 females per tank in a paternal half-sib, maternal full-sib mating design (K. Hilliard and T. Rhen, unpublished results). These tanks are 8x as long, 3.5x as wide, and 5x as deep as the average adult snapping turtle. Snapping turtles inhabit streams of similar width and depth. This provides room for the largest turtles to swim freely. Water flows continuously through tanks at a velocity similar to moving water that turtles experience naturally.

Water efflux from seven tanks passes through a multi-step filtration, sterilization, and temperature control system. The first step is mechanical filtration of solid waste as water flows into a ProFlDrum Eco 45/40 Rotary Drum Filter (RDF), which filters particles larger than 70 microns. In the second step, water passes from the RDF into a Sweetwater Low-Space Bioreactor seeded with bacteria that degrade nitrogenous wastes. In the third step, filtered water is pumped through an Emperor Aquatics SMART High Output UV Sterilizer to kill potential pathogens. In the final step, filtered and sterilized water flows through Aqua Logic Multi-Temp Chillers to control water temperature and is fed back into stock tanks. A constant turnover of 850 liters per day of fresh, de-chlorinated water is fed into the system with excess dirty water flowing out of the system into a floor drain. Water is re-circulated through the system at a rate of 2 complete water changes/tank/hour.

**Sample collection and DNA sequencing**

We extracted DNA from one adult female snapping turtle in our breeding colony. This female was captured by the MN Department of
Table 1 Summary of whole genome shotgun sequence data for *Chelydra serpentina*

| Platform          | Seq Center | Library Type | Nominal insert size | Lane or Cell | Raw Reads | Filtered Reads | Mean read length | Bases (Gb) |
|-------------------|------------|--------------|---------------------|--------------|-----------|----------------|-----------------|------------|
| HiSeq 2000        | HCI        | Paired-end   | 200 bp              | 7            | 157628596 | 148173891      | 124             | 18.37356248 |
| HiSeq 2000        | HCI        | Mate-pair    | 5.2 kb              | 7            | 169828370 | 186726281      | 124             | 23.15405884 |
| HiSeq 2000        | HCI        | Mate-pair    | 10 kb               | 7            | 217064820 | 238907655      | 124             | 29.62454922 |
| HiSeq 2000        | HCI        | Paired-end   | 200 bp              | 1            | 513299776 | 488731386      | 124             | 60.60269186 |
| HiSeq 2000        | HCI        | Paired-end   | 200 bp              | 8            | 522818714 | 480522479      | 124             | 59.58478747 |
| **total**         |            |              |                     |              | **1580640276** | **total = 191.396498** |                     |            |
| PacBio Sequel     | RTL        | SMRT         | 30 kb               | 1            | 505167   | 504425         | 124             | 4.62477795  |
| PacBio Sequel     | RTL        | SMRT         | 30 kb               | 2            | 728665   | 727435         | 124             | 5.4216822  |
| PacBio Sequel     | RTL        | SMRT         | 30 kb               | 3            | 329474   | 329001         | 124             | 2.26995261 |
| PacBio Sequel     | RTL        | SMRT         | 30 kb               | 4            | 493154   | 492682         | 124             | 4.03288831 |
| PacBio Sequel     | RTL        | SMRT         | 30 kb               | 5            | 447251   | 446685         | 124             | 3.64904182 |
| PacBio Sequel     | RTL        | SMRT         | 30 kb               | 6            | 687664   | 686769         | 124             | 5.12641628 |
| **total**         |            |              |                     |              | **3191375** | **total = 25.47005782** |                     |            |
| HiSeq 2500        | BYU        | Mate-pair    | 3kb                 | 1            | 545122952 | 255825287      | 89              | 22.76845054 |
| HiSeq 2500        | BYU        | Mate-pair    | 3kb                 | 2            | 545477596 | 255974272      | 89              | 22.78171021 |
| HiSeq 2500        | BYU        | Mate-pair    | 20kb                | 1            | 294039334 | 116881537      | 90              | 10.51933833 |
| **total**         |            |              |                     |              | **1384639882** | **total = 45.55016075** |                     |            |
| Oxford           | UND        | Nanopore     | N/A                 | Maxwell      | 560869   | N/A            | N/A             | 5.618462972 |
| Oxford           | UND        | Nanopore     | N/A                 | PC           | 391059   | N/A            | N/A             | 4.223690427 |
| Oxford           | UND        | Nanopore     | N/A                 | PC-SRE1      | 594707   | N/A            | N/A             | 3.521567316 |
| Oxford           | UND        | Nanopore     | N/A                 | PC-SRE2      | 644389   | N/A            | N/A             | 3.259168727 |
| Oxford           | UND        | Nanopore     | N/A                 | PC-SRE3      | 522053   | N/A            | N/A             | 5.103403101 |
| **total**         |            |              |                     |              | **2713077** | **total = 21.726292543** |                     |            |

Natural Resources during a fish survey of Moms Lake in central Minnesota, USA (45.9274° N, 94.7078° W) in June of 2010. We removed the female from her tank during mid-winter (water and body temperature ~3°C). Skin on the dorsum of the neck was sterilized with 70% ethanol and blood was drawn from the subcarapacial vein as described by Moon and Hernandez Foerster (2001). Whole blood was transferred to a microfuge tube and kept on ice until genomic DNA was extracted using a genomic-tip 100/G kit (Qiagen). DNA quantity was measured using Quanti-iT PicoGreen dsDNA kit and a Qubit fluorometer. DNA purity was assessed via measurement of absorbance (A230/A260 and A260/280 ratios) on a Nanodrop spectrophotometer. All DNA samples had A260/A280 ratios between 1.8 and 2.0 and A260/230 ratios between 2.0 and 2.3. DNA integrity was examined via 0.8% agarose gel electrophoresis and/or the Agilent TapeStation. Sample DNA was much longer than the 23 kb marker when examined via 0.8% agarose gel electrophoresis and/or the Agilent TapeStation. Sample DNA was also longer than the 48.5 kb marker when run on the Agilent TapeStation.

High molecular weight genomic DNA was shipped on dry ice to the High Throughput Genomics Core Facility at the Huntsman Cancer Institute, University of Utah. The facility used the Illumina TruSeq DNA PCR-Free Sample Prep protocol to make a short insert (~200 bp) library for 2 × 125 cycle paired end sequencing. The facility also used the Sage Science ELF electrophoresis system and Nextera MatePair Sample Preparation Kit to make two long insert (~5.2kb and 10kb) libraries for 2 × 125 cycle paired end sequencing. Sequencing on the Illumina HiSeq 2500 instrument produced a total of 197.58Gb of raw data (Table 1). To increase sequencing depth and augment the diversity of long insert sizes, we sent high molecular weight genomic DNA to the Sequencing Center at Brigham Young University. Two additional mate pair libraries were prepared with average insert sizes of 3kb and 20kb for 2 × 125 cycle paired end sequencing on the Illumina HiSeq 2500. This produced another 173.08Gb of raw data (Table 1). After trimming and filtering for read quality and adapters, there was a total of 236.89Gb of short read data (Table 1). Using the size of the draft genome (2.258Gb), average coverage with fully processed Illumina reads was approximately 104.9x.

We also sent high molecular weight DNA on dry ice to RTL Genomics (Lubbock Texas) for long read sequencing. The facility prepared a PacBio SMRT (Single molecule, real time) library with Sequel chemistry and sequenced the library on 6 SMRT cells. PacBio sequencing produced a total of 25.72Gb of data, with the longest reads ranging from 73.6kb to 100.6kb (Table 1). Average coverage with PacBio long reads was approximately 11.4x.

Finally, we sequenced high molecular weight DNA using the Oxford Nanopore GridION X5 system in the Genomics Core at UND. We isolated fresh DNA from whole blood of the reference turtle using three methods: the Maxwell automated nucleic acid extraction system, phenol-chloroform extraction, and phenol-chloroform extraction with size selection via the Circulomics Short Read Eliminator Kit. Libraries were made using the Ligation Sequencing Kit (SQK-LSK109) and ran on version R9.4.1 flow cells in 1D, high accuracy mode. The library prepared with DNA from the Maxwell system was sequenced on one flow cell, phenol-chloroform extracted DNA was sequenced on one flow cell, while phenol-chloroform extracted and size selected DNA was sequenced on three flow cells. Nanopore sequencing produced a total of 21.72 Gb of data (Table 1), with the longest reads ranging from 180.5 kb to 273.8 kb. Average coverage with Nanopore long reads was approximately 9.6x.

**Short and long read quality control**

Raw quality scores for reads from Illumina libraries and the Nanopore libraries are shown in Figure 1. We examined read quality using the FastQC tool and used NxTrim (v0.4.3) to filter and trim adapter sequences using default parameters and a minimum length of 25bp (O’Connell et al. 2015). This software also sorts read pairs from mate...
pair libraries into one of three categories based on the presence (or absence) and the position of junction adapters in read pairs: a mate pair bin, a paired end bin, and an unknown bin. We excluded the last category of reads from the assembly because it is impossible to tell whether reads came from one side or opposite sides of the junction adapter. A fourth bin containing single end reads is produced when one read from a pair is completely trimmed. This process of trimming and sorting reads from mate pair libraries significantly improves scaffold lengths and reduce mis-assemblies (Leggett et al. 2013, O’Connell et al. 2015). While paired end reads in the unknown category and single end reads were not used for assembly, these reads were treated as single end reads and used in later error correction and genome polishing steps.

NxTrim processed reads were then subject to another round of filtering and trimming with CLC Genomics Workbench (version 11). This was done to remove read pairs that were the result of index hopping among 200bp, 5.2kb, and 10kb libraries, which were multiplexed and run on the same lanes. The 3kb and 20kb libraries were run on separate lanes so there was no potential for index hopping. The additional round of read processing with CLC Genomics Workbench also ensured that junction and sequencing adapters were completely removed and that ambiguous sequences (limit = 2 N’s) and low-quality bases (quality limit = 0.05) were trimmed. CLC Genomics Workbench uses a modified-Mott trimming algorithm, which converts Phred (Q) scores to error probabilities and uses the quality limit as a threshold to determine stretches of low quality bases (i.e., high error probabilities) to be trimmed. We used CLC Genomics Workbench to filter phiX174 vector sequences and snapping turtle mitochondrial DNA sequences (mapping parameters; match score 1; mismatch cost 2; insertion and deletion cost 3; length fraction 0.96; similarity fraction 0.98). We discarded trimmed reads <25bp, but saved quality reads from broken pairs.

We assessed the empirical distribution of insert sizes for paired end and mate pair libraries by aligning reads to the initial assembly with Bowtie 2. We then calculated the mean and standard deviation of insert sizes to further refine input parameters for Allpaths-LG. Actual sizes of paired end and mate pair inserts were close to nominal sizes for all Illumina libraries.

We error-corrected PacBio reads using LoRDEC (v0.9) (Salmela and Rivals 2014), a hybrid error correction software that uses de Bruijn graphs constructed with trimmed and filtered Illumina reads. We used CANU (v1.8) (Koren et al. 2017) to correct and trim Nanopore sequences.

**Genome assembly and completeness**

We first estimated the size of the snapping turtle genome using k-mer frequency histograms derived from short reads and BBmap software (version 38.24) (Bushnell 2014). Genome assembly was then done in three distinct steps. In the first step, we employed ALLPATHS-LG (version 52448) (N гаранелет al. 2011), a whole-genome shotgun assembler. In the second step, we employed PBJelly (version 15.8.24) (Eнглес et al. 2012) and error-corrected PacBio reads to fill gaps and join scaffolds from the initial assembly produced by ALLPATHS-LG. After PBJelly, we used Pilon software (version 1.16) and the trimmed and filtered Illumina reads for error correction (Walker et al. 2014). In the third step, we used CANU (v1.8) (Koren et al. 2017) to produce an independent genome assembly with Nanopore sequences. We then used the intermediate assembly described above (with a very low error rate), the CANU assembly (with a higher error rate from long read technology), и quickmerge software (version 0.2)
Table 2 Statistics for the assembled *Chelydra serpentina* draft genome

| Assembly software | ALLPATHS-LG | ALLPATHS-LG + PBJELLY + PILON | ALLPATHS-LG + PBJELLY + Quickmerge + PILON |
|-------------------|-------------|-------------------------------|------------------------------------------------|
| Total Length scaffold (bp) | 2128820104 | 2314316492 | 2314078856 |
| Longest scaffold (bp) | 11970359 | 12886104 | 12890361 |
| Longest contig (bp) | 386157 | 2025513 | 2020986 |
| Number of scaffolds | 17865 | 16317 | 16317 |
| Number of contigs | 235067 | 94182 | 93330 |
| Number of gaps | 217202 | 77865 | 77013 |
| Scaffold N50 (bp) | 1191164 | 1357394 | 1358478 |
| Contig N50 (bp) | 20648 | 68275 | 68958 |
| Gaps N50 (bp) | 3590 | 3972 | 3972 |

(Chakraborty et al. 2016) to further increase the contiguity of the snapping turtle genome. In brief, quickmerge identifies high confidence overlaps between two assemblies and joins contigs and scaffolds when overlap quality surpasses user-defined thresholds. Thresholds are based on the relative length of aligned vs. unaligned regions within the entire overlapping regions to minimize the potential for spurious joining of contigs/scaffolds. We used default settings for the overlap cutoffs for selection of anchor contigs (-hco = 5.0) and extension contigs (-c = 1.5). We used the scaffold N50 from the pilon corrected CANU assembly as the length cutoff for anchor contigs (-l = 1,088,418 bases). We used the default setting for minimum alignment length to be considered for merging (-ml = 5000).

The intermediate genome assembly was used as the “reference” genome, while the CANU assembly was used as the “query” genome. The quickmerge algorithm preferentially uses the more accurate sequence from the “reference” genome in the newly joined contigs/scaffolds, while the “query” genome is used to join together higher quality contigs/scaffolds. The final draft genome was error corrected with Pilon software (version 1.16) and trimmed and filtered Illumina reads (Walker et al. 2017). Completeness of the final draft genome was assessed with Benchmarking Universal Single-Copy Orthologs (BUSCO) (Simão et al. 2015). We used Vertebrata datasets from OrthoDB V9 database containing a total of 2,586 BUSCO groups.

**Repeat annotation**

Repetitive elements in the snapping turtle genome were discerned by homology searches against known repeat databases and also by de novo prediction. We employed RepeatModeler (version open-1.0.11) to build a de novo snapping turtle repeat library (Smit and Hubley 2015). This library was subsequently used to predict, annotate and mask repeats in the snapping turtle genome using RepeatMasker (version open 4.0) (Smit et al. 2015). We used LTRharvest (GenomeTools, version 1.5.9) (Ellinghaus et al. 2008) for de novo predictions of LTR (Long Terminal Repeat) retrotransposons.

**Individual heterozygosity**

Trimmed and filtered Illumina reads were mapped to the final draft genome with CLC Genomics Workbench (no masking; match score 1; mismatch cost 2; insertion cost 2; deletion cost 3; length fraction 0.98; similarity fraction 0.98). Reads were locally realigned with multi-pass realignment (3 passes). We then called variants using the “Fixed Ploidy Variant Detector” (ploidy 2; required variant probability 95%; ignore positions with coverage above 150; ignore non-specific matches; minimum coverage 20; minimum count 4; minimum frequency 20%; base quality filter default settings). We excluded variants that were called homozygous by the software. Random variation in sequencing depth across the genome and random sequencing of alleles lead to variation from the expected allele frequency of 50% in a heterozygote so we only included variants that had allele frequencies between 25% and 75% for further analysis.

**Transcriptome assembly and gene prediction**

For transcriptome assembly, Illumina RNA-Seq reads (Table 7) were obtained from various tissues at different developmental stages (embryonic hypothalamus and pituitary gland; embryonic gonads; hatching hypothalamus and pituitary gland; hatching intestine; juvenile heart) and from dissociated embryonic gonadal cells in culture. We also sequenced RNA from embryonic gonads on the Roche 454 GS-FLX platform (Table 7). RNA quantity was measured using the Quanti-iT RNA assay kit and a Qubit fluorometer. RNA purity was assessed via absorbance measurements. All RNA samples had A260/A280 ratios between 1.75 and 2.0 and A260/230 ratios between 1.5 and 2.0. RNA integrity was examined via gel electrophoresis or Agilent TapeStation. All RNA samples had distinct 18S and 28S rRNA bands with minimal evidence of degradation (RINs were greater than 8.4).

We used the FastQC tool and CLC Genomics Workbench to trim adapter sequences and low quality bases (q-score <20) from Illumina RNA-Seq reads. Trimmed and quality filtered reads were used for transcriptome assembly using several de novo and reference-based strategies. For de novo assembly, reads from all RNA-Seq libraries were assembled together using CLC Genomics Workbench (Table 8). Reference aided assembly was performed separately for each tissue type (hypothalamus/pituitary, intestine, gonad, heart, and gonadal...)

Figure 2 Contig and scaffold N50’s for initial, intermediate, and final assemblies of the snapping turtle genome.
cells) by mapping Illumina reads to our assembled genome using Tophat (v2.1.1) and Trinity assembler with default parameters (v2.8.5) (Grabherr et al. 2011) (Table 8). Transcripts assembled using CLC Genomics Workbench and Trinity were investigated to identify potential protein-coding transcripts using TransDecoder with a minimum open reading frame of 66 amino acids (v5.5.0) (Haas et al. 2013). We also used reference-guided assembly with protein-coding transcripts from Chrysemys picta (ftp://ftp-ncbi.nlm.nih.gov/ genomes/Chrysemys_picta/RNA ), Alligator mississippiensis (ftp://ftp-ncbi.nlm.nih.gov/genomes/Alligator_mississippiensis/RNA), and Terrapene mexicana triunguis (ftp://ftp-ncbi.nlm.nih.gov/genomes/Terrapene_mexicana_triunguis/RNA) on CLC Genomic workbench (Table 8). Reads from the snapping turtle were mapped to transcripts

Figure 3 Comparison of genome assembly metrics for various reptiles. (A) Contig N50’s and scaffold N50’s and (B) number of contigs and scaffolds for Chelydra serpentina (Cs), Chrysemys picta (Cp), Terrapene mexicana triunguis (Tm), Pelodiscus sinensis (Ps), Chelonia mydas (Cm), Gopherus agassizi (Ga), Platysternon megacephalum (Pm), Anolis carolinensis (Ac), Pogona vitticeps (Pv), Gekko japonicus (Gj), Eublepharis macularius (Em), Shinisaurus crocodilus (Sc), Alligator sinensis (As), Alligator mississippiensis (Am), Crocodylus porosus (Cp), and Gavialis gangeticus (Gg).

Table 3 Comparison of the Chelydra serpentina genome to other reptile genomes

| Species | Common Name | Sequencing Technology | Coverage | Genomsize (Gb) | Contig N50 (kb) | Number of Contigs | Scaffold N50 (kb) | Number of Scaffolds | Ref. |
|---------|-------------|-----------------------|----------|----------------|----------------|-------------------|------------------|-------------------|------|
| Chelydra serpentina | Snapping Turtle | Illumina, PacBio, Nanopore | 126X | 2.26 | 872.1 | 52,731 | 5590 | 13,224 |
| Chrysemys picta | Painted Turtle | Sanger, Illumina | 18X | 2.59 | 21.3 | 262,326 | 5212 | 78,631 | 1 |
| Terrapene mexicana | Mexican Box Turtle | Illumina, 10X Genomics | 69X | 2.57 | 76.6 | 106,051 | 24249 | 52,260 | NCBI |
| Pelodiscus sinensis | Chinese Softshell Turtle | Illumina | 106X | 2.21 | 21.9 | 265,137 | 3331 | 76,151 | 2 |
| Chelonia mydas | Green Sea Turtle | Illumina | 82X | 2.24 | 20.4 | 561,968 | 3778 | 352,958 | 2 |
| Gopherus agassizi | Desert Tortoise | Illumina | 147X | 2.4 | 42.7 | 106,825 | 251 | 42,911 | 3 |
| Platysternon megacephalum | Big-headed turtle | Illumina | 208.9X | 2.32 | 41.8 | 470,184 | 7220 | 360,291 | 4 |
| Anolis carolinensis | Green anole lizard | Sanger | 7.1X | 1.8 | 79.9 | 41,986 | 4033 | 6,645 | 5 |
| Pogona vitticeps | Australian dragon lizard | Illumina | 86X | 1.77 | 31.2 | 636,524 | 2291 | 543,500 | 6 |
| Gecko japonicus | Japanese gecko | Illumina | 131X | 2.49 | 29.6 | 335,470 | 708 | 191,500 | 7 |
| Eublepharis macularius | Leopard gecko | Illumina | 136X | 2.02 | 20 | 664 | | 8 |
| Shinisaurus crocodilus | Chinese crocodile lizard | Illumina | 149X | 2.24 | 11.7 | 1470 | | 9 |
| Alligator sinensis | Chinese alligator | Illumina | 109X | 2.27 | 23.4 | 177,282 | 2188 | 9,317 | 10 |
| Alligator mississippiensis | American alligator | Illumina | 68X | 2.17 | 36 | 114,159 | 509 | 14,645 | 11 |
| Crocodylus porosus | Saltwater crocodile | Illumina | 74X | 2.12 | 32.7 | 112,407 | 204 | 23,365 | 11 |
| Gavialis gangeticus | Gharial | Illumina | 109X | 2.88 | 23.4 | 177,282 | 2188 | 9,317 | 11 |

1) Shaffer et al. 2013, 2) Wang et al. 2013, 3) Tollis et al. 2017, 4) Cao et al. 2019, 5) Alföldi et al. 2011, 6) Georges et al. 2015, 7) Liu et al. 2015, 8) Xiong et al. 2016, 9) Gao et al. 2017, 10) Wan et al. 2013, 11) Green et al. 2014.
from each species and consensus snapping turtle transcripts were extracted.

Finally, RNA from embryonic adrenal-kidney-gonad (AKG) complexes was used for direct-cDNA sequencing on the GridION system (Oxford Nanopore Technologies). Nanopore reads from direct cDNA sequencing were error-corrected using proovread (v 2.14.0) (Hackl et al. 2014) and adapter sequences removed using Porechop (v0.2.4; https://github.com/rrwick/Porechop). Cleaned Nanopore reads were assembled using CLC Genomic Workbench and CANU (v1.8) (Koren et al. 2017) (Table 8). All together, we produced 10 transcriptome assemblies using RNA from numerous tissues and sequencing platforms, as well as different assembly algorithms.

Putative protein-coding transcripts from these 10 independent assemblies were further processed with Mikado (v1.2) using default parameters (Venturini et al. 2018). Mikado uses a novel algorithm to integrate information from multiple transcriptome assemblies, splice junction detection software, and homology searches of the Swiss-Prot database to select the best-supported gene models and transcripts. We ran Mikado three times to recover as many potential protein-coding genes as possible. The first run produced 134,687 gene models, the second run produced 3,085 additional gene models, and the third run...
produced another 946 gene models for a total of 138,718 models of putative protein-coding genes.

We then used Maker (Cantarel et al. 2008) to increase the accuracy of gene models, reduce redundancy of overlapping models from the Mikado gene set, and predict new gene models that Mikado may have missed. We ran Maker basic protocol 2 (version 2.31.10), which is designed to update and combine legacy annotations (i.e., the Mikado gene models) in the light of new evidence (Campbell et al. 2014). Input for Maker included the final snapping turtle genome assembly, the 138,718 gene models from Mikado, protein evidence from the American alligator (Alligator mississippiensis), protein evidence from several turtle species (i.e., Chelonia mydas, Chrysemys picta bellii, Gopherus evgoodei, Pelodiscus sinensis, and Terrepene carolina triunguis), as well as snapping turtle transcripts (i.e., all 1,108,260 transcripts

### Table 5 Summary statistics of interspersed repeat elements in the Chelydra serpentina draft genome

| Element Type    | Number of elements | Total Length (bp) | Percentage of sequence |
|-----------------|--------------------|-------------------|------------------------|
| SINEs           | 289983             | 44174379          | 1.96                   |
| ALUs            | 1927               | 391227            | 0.02                   |
| MIRs            | 220430             | 31833398          | 1.41                   |
| LINEs:          | 687884             | 239290244         | 10.60                  |
| L1NE1           | 2095               | 697521            | 0.03                   |
| L1NE2           | 99433              | 18718432          | 0.83                   |
| L3/CR1          | 391880             | 165506806         | 7.33                   |
| LTR elements:   | 375566             | 176425159         | 7.81                   |
| ERVL            | 0                  | 0                 | 0.00                   |
| ERVL-MaLRs      | 0                  | 0                 | 0.00                   |
| ERV_classI      | 24667              | 9725298           | 0.43                   |
| ERV_classII     | 0                  | 0                 | 0.00                   |
| DNA elements:   | 1399380            | 291255572         | 12.90                  |
| hAT-Charlie     | 174802             | 39667122          | 1.76                   |
| TcMar-Tigger    | 28652              | 7023175           | 0.31                   |
| Unclassified:   | 222952             | 78727133          | 3.49                   |
| Total interspersed repeats | 829872487         |                   | 36.76                  |

**Figure 5** Comparison of repeat content among genomes for Chelydra serpentina, Chrysemys picta, and Gopherus agassizii.
assembled with CLC Genomics Workbench, but not filtered with TransDecoder. Maker produced 30,166 models for putative protein-coding genes.

We assessed Mikado and Maker gene models by blasting predicted transcripts against the painted turtle (*Chrysemys picta*) proteome. Based on BLASTX hits to *Chrysemys picta* proteins, there were 15,718 protein-coding genes in common between Mikado and Maker gene sets. However, there were also differences between gene prediction software. The Mikado gene set contained hits to 1,071 *Chrysemys picta* proteins that were not in the Maker gene set (i.e., Maker lost these genes). Conversely, the Maker gene set contained hits to 614 *Chrysemys picta* proteins that were not in the Mikado gene set (i.e., Maker discovered these genes). This comparison revealed that Mikado and Maker each produced a significant number of gene models the other software missed. To avoid losing protein-coding genes, we used both Mikado and Maker gene models in the following pipeline.

To obtain a final set of gene models that are likely to encode real proteins, we ran predicted snapping turtle proteins from Mikado and Maker through OrthoFinder with default settings (Emms and Kelly 2019). OrthoFinder classifies proteins from two or more species into sets of proteins called “orthogroups” that contain orthologs and/or paralogs. We used proteomes from mammals (*Homo sapiens*, *Mus musculus*, and *Rattus norvegicus*), archosaurs (*Gallus gallus* and *Alligator mississippiensis*), and turtles (*Chrysemys picta*, *Pelodiscus sinensis*, and *Terrapene carolina triunguis*) to identify 49,518 snapping turtle proteins that were members of “orthogroups” with proteins from at least one other vertebrate species. We then filtered exact sequence duplicates at the mRNA level to select 43,093 gene models. We further reduced redundancy by running mRNAs through CD-Hit-EST at a 98% identity level to produce a penultimate set of 25,630 gene models for protein-coding genes.

Finally, we used bedtools (v2.27.1) to cluster overlapping gene models on the same strand and remove redundant gene models that represent alternative splice variants of the same gene (Quinlan and Hall 2010). We checked both strands for gene models and removed single exon predictions with no homology to proteins in other species. We also removed single exon predictions that contained internal stop codons. This produced a final set of 22,812 gene models for protein-coding genes in the common snapping turtle.

**Gene annotation**

Many researchers simply carry out BLASTP to the Swiss-Prot database and adopt gene names and symbols from the best hit, which leads to propagation of annotation errors (Salzberg 2019). In addition, genes that are duplicated (i.e., paralogs) or lost (i.e., gene deletion) in different lineages or species may make it difficult to accurately assign gene names/symbols to orthologs. We therefore used OrthoFinder to annotate our final set of 22,812 protein-coding genes based on orthology among several amniotic vertebrates. We first assigned human gene names and symbols to 11,835 genes that displayed one-to-one orthology across snapping turtles, humans, and at least one other species (alligator, chicken, painted turtle, or box turtle). We then assigned alligator gene names and symbols to 840 genes based on one-to-one orthology across snapping turtles, alligator, and one other species (chicken, painted turtle, or box turtle). Another 236 genes were annotated with chicken gene names and symbols based on one-to-one orthology across snapping turtles, chicken, and one other turtle (painted turtle or box turtle). A fourth set of 1376 genes was annotated based on one-to-one orthology across snapping turtle, painted turtle, and the box turtle. Gene symbols from non-human databases were converted to HUGO Gene Nomenclature Committee (HGNC) gene symbols for orthologous genes. This process produced high confidence gene names and symbols for 14,287 protein-coding genes.

We used BLASTP to assign gene names and symbols to 690 more genes that had hits to the Swiss-Prot database (when all hits had the same unique name) and to 743 more genes that had hits to box turtle proteins (when there was also supporting evidence from Swiss-Prot). Gene names and symbols were assigned to 15,720 genes. Some symbols did not meet NCBI guidelines so these were replaced with locus tags (see Eukaryotic Genome Annotation Guide; https://www.ncbi.nlm.nih.gov/genbank/eukaryotic_genome_submission_annotation/#protein_id). These genes still have unique gene names, but do not have gene symbols. Another 4,930 genes were annotated with gene names based on BLASTP hits to Swiss-Prot or to box turtle proteins (i.e., these genes have locus tags, but do not have HGNC gene symbols).

**Comparative and phylogenomic analysis of protein coding genes**

We used OrthoFinder (Emms and Kelly 2019) to compare 22,812 snapping turtle proteins to proteomes from 16 other vertebrate species to assess the completeness of our gene models at a genome wide scale. We also used OrthoFinder and STRIDE (Emms and Kelly 2019) to carry out phylogenomic analyses to see whether evolutionary relationships among turtles are consistent with phylogenetic trees from prior studies. We retrieved proteomes from mammals
Table 7 Summary of whole transcriptome shotgun sequence data for *Chelydra serpentina*

| Tissue Type                                      | Sequencing Platform | Library Type | Read Length | Raw Reads | Mean read length | Bases (Gb) |
|-------------------------------------------------|---------------------|--------------|-------------|-----------|------------------|------------|
| Embryonic and Hatchling Hypothalamus/Pituitary   | Illumina            | Single-end   | 50 bp       | 17224433  | n/a              | 8.61       |
| Embryonic Gonads                                | Illumina            | Single-end   | 100 bp      | 153596329 | n/a              | 15.36      |
| Hatchling Intestine                             | Illumina            | Single-end   | 50 bp       | 31757630  | n/a              | 1.59       |
| Juvenile Heart                                  | Illumina            | Paired-end   | 150 bp      | 366536144 | n/a              | 54.98      |
| Cultured Embryonic Gonad Cells                  | Illumina            | Paired-end   | 50 bp       | 446985548 | n/a              | 22.35      |
| Embryonic Gonads                                | 454                 | Variable     | 2255133     | 387 bp    | (151-825 bp)     | 0.87       |
| Embryonic Adrenal-Kidney-Gonad Complex          | Nanopore            | Direct cDNA  | Variable    | 3164253   | 1386 bp (101-26870 bp) | 4.39  

| Total = 1176539368                               | Total = 108.15      |

Non-coding RNAs

Transfer RNAs (tRFs) were predicted using tRNAscan-SE (version 2.0) (Lowe and Eddy 1997) with a score threshold of 65. Putative tRFs that overlapped protein-coding genes were removed. Ribosomal RNAs (rRNAs) were predicted using Brrnap (Seemann and Booth 2013). Partial or shortened rRNAs were removed. Hairpin micro-RNAs (miRNAs) were predicted by aligning all hairpin and mature miRNAs sequences from miRBase (release 22) (Griffiths-Jones et al. 2008) to the snapping turtle genome using BLASTN (e-value < 1e-10 for hairpin sequences). This gave initial predictions for 16,169 hairpin sequences and 1,175,272 mature miRNA sequences. Sequences were clustered by genomic loci, returning 1,514 hairpin clusters and 989,989 mature miRNA clusters. We then selected 899 hairpin clusters that had complete overlap with a mature hairpin sequence. miRBase entries occurring in more than one cluster were removed and clusters containing hits to less than two species were removed. The consensus name for each cluster was chosen based on the most frequent mature hairpin miRNA name within the cluster. Final genomic coordinates of hairpin miRNA sequences were selected based on the lowest e-value.

RESULTS AND DISCUSSION

Genome assembly and completeness

Initial assembly of the snapping turtle genome using Illumina short reads produced a genome of 2.128Gb, which is similar to the 2.20Gb predicted by BBmap. The initial assembly with ALLPATHS-LG had a total of 17,865 scaffolds (Table 2). The intermediate genome assembly that incorporated PacBio long reads (ALLPATHS-LG, PBjelly, and Pilon) had a size of 2.314Gb with 16,317 scaffolds (Table 2). The longest scaffold was 11.97Mb for the initial assembly and 12.89Mb for the intermediate assembly (Table 2). The number of contigs decreased from 235,067 to 93,330, the longest contig increased 5.2 fold, and contig N50 increased 3.3 fold. Improvements in the intermediate assembly were largely driven by gap filling, with the number of gaps decreasing to one third of the initial assembly (Table 2).

Although improvements in assembly metrics were modest from the initial to the intermediate assembly, there were substantial improvements in assembly metrics with the final assembly (Table 2). For instance, N50 and scaffold N50 increased 12.6x fold and 3.4x fold, respectively. Improvements in assembly metrics were driven by gap filling and the incorporation of long reads (ALLPATHS-LG, PBjelly, and Pilon). The final draft genome can be found in the NCBI SRA database (SUB6351883: accession numbers SRR10270339, SRR10270340, SRR10270341, SRR10270342, SRR10270343, and SRR10270344) under BioProject PRJNA574487. Scripts are available on GitHub (https://github.com/turkrhen/snapping_turtle_genome_scripts).

Data availability

Raw data used for genome assembly, transcriptome assembly, and the final draft genome can be found in the NCBI SRA database (SUB6351883: accession numbers SRR10270339, SRR10270340, SRR10270341, SRR10270342, SRR10270343, and SRR10270344) under BioProject PRJNA574487. Scripts are available on GitHub (https://github.com/turkrhen/snapping_turtle_genome_scripts).

Table 8 Summary of intermediate transcriptome assemblies for *Chelydra serpentina*

| Assembly Type | Sequencing Platform | Tissue Type | Assembler | Total Transcripts | Transdecoder Transcripts | blast2cap3 | Mikado Input |
|---------------|---------------------|-------------|-----------|-------------------|--------------------------|------------|-------------|
| Reference aided (A. mississippiensis) | Illumina & 454 | H/P, G, I, H, C | CLC Genomics | 35436 | n/a | 35436 |
| Reference aided (C. picta) | Illumina & 454 | H/P, G, I, H, C | CLC Genomics | 38262 | n/a | 38262 |
| Reference aided (T. carolina) | Illumina & 454 | H/P, G, I, H, C | CLC Genomics | 29707 | n/a | 29707 |
| De novo | Illumina & 454 | H/P, G, I, H, C | CLC Genomics | 1161412 | 160679 | 154815 |
| De novo | Nanopore direct cDNA | AKG | Canu | 11924 | n/a | 11924 |
| De novo | Nanopore direct cDNA | AKG | CLC Genomics | 9025 | n/a | 9025 |
| De novo | Illumina | G | Trinity | 382845 | 99801 | 99801 |
| De novo | Illumina | H | Trinity | 613600 | 151273 | 151273 |
| De novo | Illumina | H/P, I | Trinity | 286368 | 75823 | 75823 |
| De novo | Illumina | C | Trinity | 837323 | 124988 | 124988 |

Key to tissue types: Embryonic and Hatchling Hypothalamus/Pituitary (H/P), Embryonic Gonads (G), Hatchling Intestine (I), Juvenile Heart (H), Cultured Embryonic Gonad Cells (C), Embryonic Adrenal-Kidney-Gonad Complexes (AKG).
draw. The final draft genome that integrated Nanopore long reads had a size of 2.258 Gb with 13,224 scaffolds (Table 2). Scaffold N50 for the final genome was 5.59 Mb while the longest scaffold was 27.24 Mb. In addition, the number of contigs and gaps dropped by half, which indicates a substantial improvement in the contiguity of the final draft genome. The GC content was estimated to be 44.34%, which is comparable to the 43–44% GC content reported in other turtle species (Shaffer et al. 2013, Wang et al. 2013, Tollis et al. 2017, Cao et al. 2019).

The snapping turtle genome displays greater contiguity than most other published reptile genomes (Figure 3; Table 3). The only exception was the Mexican box turtle, which used 10X Genomics linked reads to produce a 4.3 fold longer scaffold N50 (Figure 3A; Table 3). Yet, the snapping turtle contig N50 was 11.4 fold longer than the box turtle contig N50 (Figure 3A; Table 3). The snapping turtle genome also has half as many contigs and one quarter the scaffolds as the box turtle genome (Figure 3B; Table 3). Differences in various measures of contiguity reflect the different technologies used to acquire long-range sequence information (10X Genomics linked reads in box turtle vs. PacBio and Nanopore long reads in snapping turtle). This suggests that linked and long reads provide complementary information that could dramatically improve genome contiguity if used together.

The snapping turtle reference genome contained both complete (94.2%) and fragmented (3.2%) core vertebrate genes as assessed via BUSCO (Table 4). This estimate of completeness is comparable to the completeness of other turtle genomes (Figure 4). Only 2.6% of the BUSCO core vertebrate genes were missing from the snapping turtle genome, which is a similar level of completeness reported in other reptiles (Gao et al. 2017).

Repetitive DNA
The total length of repetitive elements accounted for 36.76% of the snapping turtle genome (Table 5). This is halfway between the repetitive DNA content of other turtle genomes: 29% in Chrysemys picta and 43% in Gopherus agassizi (Tollis et al. 2017). The greatest variation in repetitive DNA elements among species was in LTRs, DNA transposons, and unclassified repeats (Figure 5).

Individual heterozygosity
A total of 3.70 million variants were detected in the reference snapping turtle, with 3.27 million single nucleotide polymorphisms (SNPs; Table 6). In comparison, 4.99 million SNPs were reported in the big-headed turtle (Cao et al. 2019). However, the method used to identify SNPs in the big-headed turtle was much less stringent, which explains the higher number of SNPs.

Genome-wide levels of individual heterozygosity have not yet been reported for any other turtle species so we compared the snapping turtle to mammals. We found 3.27 million SNPs in the reference snapping turtle (genome size = 2.258Gb), while studies of individual humans reported 3.07, 3.21, and 3.32 million SNPs in an Asian and two Caucasians, respectively (Levy et al. 2007, Wang et al. 2008, Wheeler et al. 2008). Individual heterozygosity for SNPs in the snapping turtle, after correction for the difference in genome size, is slightly higher than observed in humans. Moreover, the 1,448 heterozygous SNPs/Mb observed in the snapping turtle falls in the upper range observed in 27 mammalian species (Abascal et al. 2016). While population genomic studies will be required to draw firm conclusions, the relatively high level of
heterozygosity in the reference snapping turtle suggests that inbreeding and/or population bottlenecks were not a common occurrence in its ancestors. The genetic variants identified here can be used as markers for studying the relationship between genotype and phenotype, as well as for analysis of genome-wide patterns of molecular evolution.

### Gene annotation

We annotated 20,650 protein-coding genes, which is very similar to the number found in the painted turtle (21,796) and desert tortoise (20,172) genomes. The remaining 2,162 models for protein-coding genes in the snapping turtle did not display homology to other known genes and are considered hypothetical proteins at this time. We assessed the accuracy of our automated annotations by conducting manual BLASTN of cDNA sequences for 2,006 gene models that were assigned HGNC gene symbols. We used GeneCards.org to crosscheck gene names/symbols that did not match manual BLASTN hits to determine whether gene names/symbols were aliases or incorrect annotations. Aliases were considered correct because they are synonyms for the same locus.

Most automated annotations with our orthology-based pipeline were correct (97.7%; n = 1960), while a small percentage (2.3%; n = 46) were incorrect. Most of the incorrectly annotated genes were assigned gene names and symbols of close paralogs (1.5%; n = 31), but some annotations were completely incorrect (0.7%; n = 15) due to propagation of annotation errors from other species. In comparison, annotation of the same genes based on the top hit to Swiss-Prot was less accurate (96.4% correct, n = 1933; 3.6% incorrect, n = 73). The rate of completely incorrect names and symbols doubled with annotation based on the top hit to Swiss-Prot (1.6%; n = 32). In addition, slightly more genes were assigned names and symbols of close paralogs rather than orthologs (2.0%; n = 41).

### Comparative analysis of protein coding genes and phylogenomic relationships

The vast majority (22,735; 99.7%) of protein-coding genes in snapping turtles were assigned to orthogroups (Figure 6; Table 9), which are gene lineages comprised of orthologs and paralogs. This is similar to the number of genes assigned to orthogroups in the painted turtle and the box turtle, but higher than the number in the big-headed turtle and Chinese softshell turtle (Table 9). In contrast, many more genes were assigned to orthogroups in the green sea turtle and the desert tortoise (Table 9), which may be due to sequence redundancy in those databases.

The number of orthogroups in snapping turtles (15,511) is very similar to the number of orthogroups in painted turtle, box turtle, green sea turtle, Chinese alligator, human, mouse, rat, and zebrafish (15,263 to 15,813 orthogroups) (Table 9). The number of orthogroups is an index of the number of gene families that are conserved across vertebrates. The median number of orthogroups (15,515) in the species we examined is very close to a prior estimate of orthogroups

### Table 10 Functional annotation of Chelydra serpentina proteins based on de novo prediction using Interproscan and evolutionary homology to human proteins (i.e., one-to-one orthologs). Total numbers are the result of merging de novo annotations with homology-based annotations and reducing redundant terms (i.e., eliminating duplicates)

| Annotation Database | Proteins Annotated | Number of Annotations | Number of Unique Terms |
|---------------------|--------------------|-----------------------|------------------------|
| Interproscan        | GO                 | 13558                 | 34064                  | 2499                   |
|                     | KEGG               | 1015                  | 2800                   | 801                    |
|                     | Reactome           | 5341                  | 19076                  | 1454                   |
| Homology            | GO                 | 12704                 | 216110                 | 17057                  |
|                     | KEGG               | 967                   | 2622                   | 866                    |
|                     | Reactome           | 7802                  | 31824                  | 1842                   |
| Total (merged)      | GO                 | 17910                 | 234877                 | 17169                  |
|                     | KEGG               | 1212                  | 3365                   | 935                    |
|                     | Reactome           | 8991                  | 34058                  | 1857                   |

### Table 11 Accession numbers for vertebrate proteomes used for comparison to the snapping turtle genome

| Proteome            | Database | Accession        | Isoforms |
|---------------------|----------|------------------|----------|
| Danio rerio         | NCBI     | GCF_000002035.6  | Yes      |
| Homo sapiens        | UniProt  | UP000005640      | No       |
| Rattus norvegicus   | UniProt  | UP000002494      | No       |
| Mus musculus        | UniProt  | UP000000589      | No       |
| Taeniopygia guttata | NCBI     | GCF_008822105.2  | Yes      |
| Meleagris gallopavo | NCBI     | GCF_000146605.3  | Yes      |
| Gallus gallus       | UniProt  | UP000000539      | No       |
| Pelodiscus sinensis | UniProt  | UP000007267      | No       |
| Platystemon megacephalum | NCBI   | GCA_003942145.1  | Yes      |
| Chrysemys picta     | NCBI     | GCF_000241765.3  | No       |
| Terrapene carolina | NCBI     | GCF_002925995.2  | No       |
| Gopherus evgooodi   | NCBI     | GCA_002896415.1  | Yes      |
| Chelonia mydas      | UniProt  | UP0000031443     | No       |
| Crocodylus porosus  | NCBI     | GCF_001723895.1  | Yes      |
| Alligator mississippi | UniProt | UP000005025     | No       |
| Alligator sinensis  | NCBI     | GCF_000455745.1  | Yes      |
Based on this index, gene prediction in the snapping turtle is as complete as the best annotated turtle, crocodilian, mammalian, and fish genomes. In contrast, big-headed turtle, desert tortoise, Chinese softshell turtle, American alligator, saltwater crocodile, and bird genomes have fewer orthogroups (13,289 to 14,880) (Table 9). This suggests gene models are incomplete (i.e., missing 700 to 2,200 genes) in those species or that genes have been lost during evolution in those species. Other turtles and Chinese alligator have the typical number of orthogroups found in well-annotated mammalian and zebrafish genomes so it is more likely that gene models are incomplete in big-headed turtle, desert tortoise, Chinese softshell turtle, American alligator, and saltwater crocodile. In support of this idea, birds are known to have fewer orthogroups (~15% less) due to poor annotation of genes in GC rich regions (Botero-Castro et al. 2017).

Relationships among turtles based on all protein coding genes (Figure 7) perfectly reflect phylogenetic relationships inferred from a smaller set of 539 nuclear genes (Shaffer et al. 2017). Snapping turtles are more closely related to sea turtles (Chelonia mydas) than to other turtles (Figure 7). This tree also shows the big-headed turtle is a sister species to eumyid turtles and that tortoises are a sister group to both the big-headed turtle and eumyid turtles. Finally, the Chinese softshell turtle is the most divergent turtle examined here. The extent of orthogroup overlap among species again suggests gene models are incomplete in the big-headed turtle, desert tortoise, Chinese softshell turtle, birds, American alligator, and saltwater crocodile (i.e., lighter colors both on and off the diagonal indicate fewer shared orthogroups; Figure 7).

**Functional annotation of protein-coding genes** Experimental annotation of protein function at a genome wide scale is impractical for new model species like the snapping turtle. However, it is possible to annotate protein function based on well-characterized structural domains and by evolutionary homology to proteins in highly curated databases. In an effort to capture both conserved and divergent structural and functional elements of snapping turtle proteins we used a combinatorial approach to annotation based on structural homology to protein domains and evolutionary homology to proteins of known function. We used InterProScan (version 5.36-75.0) to assign Gene Ontology terms, KEGG pathways, and REACTOME pathways to snapping turtle proteins (Table 10). This resulted in de novo functional annotation of 13,558 proteins based on protein architecture and functional domains. For more complete functional annotation, we also adopted Gene Ontology terms, KEGG pathways, and REACTOME pathways associated with 12,704 genes identified as one-to-one orthologs to human genes (Table 10). We merged results from these methods and reduced redundancy of functional annotations (i.e., duplicate terms). This resulted in a large set of annotations inferred from both protein signatures and evolutionary homology. As such, they should be viewed as putative rather than definitive annotations.
Non-coding RNAs

tRNAscan-SE predicted a total of 687 tRNAs and Barrnap predicted 43 rRNAs in the snapping turtle genome. Alignment and filtering of known hairpin and mature micro-RNAs (miRNA) sequences from miRBase returned a set of 204 high confidence hairpin miRNA sequences in the snapping turtle genome.

Summary assessment of genome assembly and annotation

Here we describe de novo assembly and annotation of the snapping turtle genome using both short and long read sequencing technologies and several genome assembly algorithms. The contiguity of this assembly (contig N50, scaffold N50, and number of contigs/scaffolds) is greater than most other published turtle and reptile genomes (Table 3) (Alföldi et al. 2011, Shaffer et al. 2013, Wan et al. 2013, Wang et al. 2013, Green et al. 2014, Georges et al. 2015, Liu et al. 2015, Xiong et al. 2016, Gao et al. 2017, Tolls et al. 2017, Cao et al. 2019). Gene and repeat content in the snapping turtle is very similar to other turtles. We provide the first assessment of individual heterozygosity at a genome-wide scale in a turtle and find it is at the upper end of the range of heterozygosity observed in mammals. This observation is consistent with the broad geographic range and abundance of snapping turtles across North America. The reference genome and genomic variants identified here provide a foundation for molecular genetic, quantitative genetic, and population genomic studies of adaptation to climate in the snapping turtle. An abundant species like the snapping turtle serves as a tractable model to identify specific genes underlying genome-environment interactions. Of particular interest are genes that influence thermosensitive sex determination, which can then be studied in threatened and endangered turtle species.

ACKNOWLEDGMENTS

All tissue, DNA, and RNA samples were collected using procedures approved by the Institutional Animal Care and Use Committee at the University of North Dakota. This work was supported by the National Science Foundation of the United States (grant numbers IOS-0923300, IOS-1558034, and IOS-1755282 to TR and IOS-1548562. The species range and abundance of snapping turtles across North America. The reference genome and genomic variants identified here provide a foundation for molecular genetic, quantitative genetic, and population genomic studies of adaptation to climate in the snapping turtle. An abundant species like the snapping turtle serves as a tractable model to identify specific genes underlying genome-environment interactions. Of particular interest are genes that influence thermosensitive sex determination, which can then be studied in threatened and endangered turtle species.

LITERATURE CITED

Abascal, F. A., F. Corvelo, J. L. Cruz, A. Villanueva-Cañas, M. Vlasova et al., 2016 Extreme genomic erosion after recurrent demographic bottlenecks in the highly endangered Iberian lynx. Genome Biol. 17: 251. https://doi.org/10.1186/s13059-016-1090-1

Agrawal, A. A., 2001 Ecology: Phenotypic plasticity in the interactions and evolution of species. Science 294: 321–326. https://doi.org/10.1126/science.1060701

Angilletta, M. J., 2009 Thermal Adaptation: A Theoretical and Empirical Synthesis, Ed. 1st. Oxford University Press, Oxford: UK. https://doi.org/10.1093/acprof:oso/9780198570875.001.1

Alföldi, J., F. Di Palma, M. Grabherr, C. Williams, L. Kong et al., 2011 The genome of the green anole lizard and a comparative analysis with birds and mammals. Nature 477: 587–591. https://doi.org/10.1038/nature10390

Böhm, M., B. Collen, J. E. M. Baille, P. Bowles, J. Chanson et al., 2013 The conservation status of the world’s reptiles. Biol. Conserv. 157: 372–385. https://doi.org/10.1016/j.biocon.2012.07.015

Botero-Castro, F., E. Figuet, M.-K. Tilak, B. Nabholz, and N. Galtier, 2017 Avian genomes revisited: hidden genes uncovered and the rate vs. traits paradox in birds. Mol. Biol. Evol. 34: 3123–3131. https://doi.org/10.1093/molbev/msx226

Bushnell, B., 2014 BBMap: a fast, accurate, splice-aware aligner. United States. Available online at: https://sourceforge.net/projects/bbmap/

Campbell, M. S., C. Holt, B. Moore, and M. Yandell, 2014 Genome annotation and curration using MAKER and MAKER-P. Curr. Protoc. Bioinformatics 48: 1–39. https://doi.org/10.1002/9781118114148

Cantarel, B. L., I. Korf, S. M. Robb, G. Parra, E. Ross et al., 2008 MAKER: an easy-to-use annotation pipeline designed for emerging model organism genomes. Genome Res. 18: 188–196. https://doi.org/10.1101/ gr.6743907

Cao, D., M. Wang, Y. Ge, and S. Gong, 2019 Draft genome of the big-headed turtle Platysternon megacephalum. Sci. Data 6: 60. https://doi.org/10.1038/s41597-019-0067-9

Chakraborty, M., J. G. Baldwin-Brown, A. D. Long, and J. J. Emerson, 2016 Contiguous and accurate de novo assembly of metazoan genomes with modest long read coverage. Nucleic Acids Res. 44: e147.

Ellinghaus, D., S. Kurtz, and U. Willhoft, 2008 LRTrharvest, an efficient and flexible software for de novo detection of LTR retrotransposons. BMC Bioinformatics 9: 18. https://doi.org/10.1186/1471-2105-9-18

Emms, D. M., and S. Kelly, 2019 OrthoFinder: phylogenetic orthology inference for comparative genomics. Genome Biol. 20: 238. https://doi.org/10.1186/s13059-019-1832-y

English, A. C., S. Richards, Y. Han, M. Wang, V. Vee et al., 2012 Mind the gap: upgrading genomes with Pacific Biosciences RS long-read sequencing technology. PLoS One 7: e47768. https://doi.org/10.1371/journal.pone.0047768

Ewert, M. A., C. Etchberger, and C. E. Nelson, 2004 Turtle sex-determining modes and TSD patterns, and some TSD pattern correlates, pp. 21–32 in Temperature-Dependent Sex Determination in Vertebrates, edited by Valenzuela, N., and V. Lance. Smithsonian Institution Press, Washington, D.C.

Ewert, M. A., J. W. Lang, and C. E. Nelson, 2005 Geographic variation in the pattern of temperature-dependent sex determination in the American snapping turtle (Chelydra serpentina). J. Zooi. (Lond.) 265: 81–95. https://doi.org/10.1017/S0952836905006120

Gao, J., Q. Li, Z. Wang, Y. Zhou, P. Martelli et al., 2017 Sequencing de novo assembling, and annotating the genome of the endangered Chinese crocodile lizard Shinisaurus crocodilurus. Gigascience 6: 1–6. https://doi.org/10.1093/gigascience/gix041

Georges, A., Q. Li, J. Lian, D. O’Meally, J. Deakin et al., 2015 High-coverage sequencing and annotated assembly of the genome of the Australian dragon lizard Pogona vitticeps. Gigascience 4: 45. https://doi.org/10.1093/gigascience/gix041

Gnerre, S., I. MacCallum, D. Przybylski, F. Ribeiro, J. Burton et al., 2011 High-quality draft assemblies of mammalian genomes from massively parallel sequence data. Proc. Natl. Acad. Sci. USA 108: 1513–1518. https://doi.org/10.1073/pnas.1017351108

Grabherr, M. G., B. J. Haas, M. Yassour, J. Z. Levin, D. A. Thompson et al., 2011 Full-length transcriptome assembly from RNA-seq data without a reference genome. Nat. Biotechnol. 29: 644–652. https://doi.org/10.1038/nbt.1883

Green, R. E., E. L. Braun, J. Armstrong, D. Earl, N. Nguyen et al., 2014 Three crocodilian genomes reveal ancestral patterns of evolution among archosaurs. Science 346: 1254449. https://doi.org/10.1126/science.1254449
Turtle Specialist Group, Turtle Conservancy, Turtle Survival Alliance, Turtle Conservation Fund, Chelonian Research Foundation, Conservation International, Wildlife Conservation Society, and Global Wildlife Conservation, Ojai, CA, USA, Volume 80: 1–84.
Steyermark, A. C., M. S. Finkler, and R. J. Brooks (Editors), 2008 Biology of the Snapping Turtle (Chelydra serpentina), The Johns Hopkins University Press, Baltimore.
Tollis, M., D. F. DeNardo, J. A. Cornelius, G. A. Dolby, T. Edwards et al., 2017 The Agassiz’s desert tortoise genome provides a resource for the conservation of a threatened species. PLoS One 12: e0177708. https://doi.org/10.1371/journal.pone.0177708
Valenzuela, N., and D. C. Adams, 2011 Chromosome number and sex determination coevolve in turtles. Evolution 65: 1808–1813. https://doi.org/10.1111/j.1558-5646.2011.01258.x
Venturini, L., S. Caim, G. G. Kaithakottil, and D. L. Mapleson, and D. Swarbreck, 2018 Leveraging multiple transcriptome assembly methods for improved gene structure annotation. Gigascience 7: giy093. https://doi.org/10.1093/gigascience/giy093
Via, S., and R. Lande, 1985 Genotype-environment interaction and the evolution of phenotypic plasticity. Evolution 39: 505–522. https://doi.org/10.1111/j.1558-5646.1985.tb00391.x
Walker, B. J., T. Abeel, T. Shea, M. Priest, A. Abouelliel et al., 2014 Pilon: an integrated tool for comprehensive microbial variant detection and genome assembly improvement. PLoS One 9: e112963. https://doi.org/10.1371/journal.pone.0112963
Wan, Q. H., S. K. Pan, L. Hu, Y. Zhu, P.-W. Xu et al., 2013 Genome analysis and signature discovery for diving and sensory properties of the endangered Chinese alligator. Cell Res. 23: 1091–1105. https://doi.org/10.1038/cr.2013.104
Wang, J., W. Wang, R. Li, Y. Li, G. Tian et al., 2008 The diploid genome sequence of an Asian individual. Nature 456: 60–65. https://doi.org/10.1038/nature07484
Wang, Z., J. Pascual-Anaya, A. Zadissa, W. Li, Y. Niimura et al., 2013 The draft genomes of soft-shell turtle and green sea turtle yield insights into the development and evolution of the turtle-specific body plan. Nat. Genet. 45: 701–706. https://doi.org/10.1038/ng.2615
Warner, D. A., W.-G. Du, and A. Georges, 2018 Introduction to the special issue – Developmental plasticity in reptiles: Physiological mechanisms and ecological consequences. J. Exp. Zool. 329: 153–161. https://doi.org/10.1002/jez.2199
Wheeler, D. A., M. Srinivasan, M. Egholm, Y. Shen, L. Chen et al., 2008 The complete genome of an individual by massively parallel DNA sequencing. Nature 452: 872–876. https://doi.org/10.1038/nature06884
While, G. M., D. W. A. Noble, T. Uller, D. A. Warner, J. L. Riley et al., 2018 Patterns of developmental plasticity in response to incubation temperature in reptiles. Journal of Experimental Zoology A. 329: 162–176. https://doi.org/10.1002/jez.2181
Xiong, Z., F. Li, Q. Li, L. Zhou, T. Gamble et al., 2016 Draft genome of the leopard gecko, Eublepharis macularius. Gigascience 5: 47. https://doi.org/10.1186/s13742-016-0151-4

Communicating editor: A. Sethuraman