1

A high-quality reference genome for the common creek chub, $Semotilus \ atromaculatus$

Amanda V. Meuser^{1,2}, Amy R. Pitura^{1,2}, Elizabeth G. Mandeville¹

¹ Department of Integrative Biology, University of Guelph, Guelph, Ontario, Canada

 2 Co-first authors

Corresponding author: Amy Pitura University of Guelph 50 Stone Road East Guelph, Ontario N1G 2W1, Canada apitura@uoguelph.ca 1-519-824-4120 ext. 52843

Running title: Semotilus atromaculatus reference genome

 $Keywords:\ Semotilus\ atromaculatus,\ reference\ genome,\ creek\ chub,\ synteny,\ cyprinid,\ leuciscid$

2

¹ Abstract

Creek chub (Semotilus atromaculatus) are a leuciscid minnow species commonly found in an-2 thropogenically disturbed environments, making them an excellent model organism to study 3 human impacts on aquatic systems. Genomic resources for creek chub and other leuciscid 4 species are currently limited. However, advancements in DNA sequencing now allow us to 5 create genomic resources at a historically low cost. Here, we present a high quality 239 con-6 tig reference genome for the common creek chub, created with PacBio HiFi sequencing. We 7 compared the assembly quality of two pipelines: Pacific Biosciences' Improved Phase Assem-8 bly (IPA; 873 contigs) and Hifasm (239 contigs). Quality and completeness of this genome 9 is comparable to the zebrafish (Danioninae) and fathead minnow (Leuciscidae) genomes. 10 The creek chub genome is highly syntenic to the zebrafish and fathead minnow genomes, 11 and while our assembly does not resolve into the expected 25 chromosomes, synteny with 12 zebrafish suggests that each creek chub chromosome is likely represented by 1-4 large contigs 13 in our assembly. This reference genome is a valuable resource that will enhance genomic bio-14 diversity studies of creek chub and other non-model leuciscid species common to disturbed 15 environments. 16

3

17 Introduction

Genomic studies of non-model species have become increasingly feasible in the past two 18 decades (Narum et al. 2013, Lou et al. 2021). Efforts are now under way to sequence 19 the tree of life (Fan et al. 2020), including organisms of no known economic importance, 20 whose genomes are likely to be used primarily for conservation or evolutionary research. 21 While genomic studies of non-model organisms are proceeding at a rapid pace, progress is 22 still limited by the lack of suitable reference genomes for many species and clades. Using 23 a reference genome from a closely related species is sometimes possible when there is no 24 available reference for focal taxa (e.g. Mandeville et al. 2019). However, this approach can 25 produce misleading results under some circumstances, including when the goal of a study is 26 to examine within-species differentiation or similar and species-specific variation may be lost 27 (as when wolf vs. dog reference genomes were used for a study of wolves; Gopalakrishnan et al. 28 2017). These concerns are especially relevant when considering clades where a substantial 29 amount of structural genomic variation exists. 30

Fish genomes are quite diverse, and vary in size from 0.5-2 pg (excluding polyploids; 31 Smith & Gregory 2009). On a locus-specific functional level, essential processes like sex de-32 termination can have an incredibly diverse genetic basis in fish (Bachtrog et al. 2014, Pennell 33 et al. 2018). Within North American teleost fish, a large proportion of fish biodiversity is 34 encompassed by the family Leuciscidae within the order Cypriniformes (Holm et al. 2022, 35 Stout et al. 2016), but previous genomic work on leuciscid minnow species has been lim-36 ited. However, being extremely numerous and geographically widespread, these species have 37 great potential for use as model species to study the effects of anthropogenic disturbance 38 and overall population genetic structure of stream fishes. Some species are quite tolerant, 39 and persist or even thrive in disturbed environments (Stammler et al. 2008). Additionally, 40 these species are known to hybridize from morphological data, but hybridization patterns 41 have not been described in detail using genetic or genomic data (Corush *et al.* 2021). 42

One major limitation for future work is that no leuciscid reference genomes have been 43 available until recently, and the highest quality available reference genome would be a ze-44 brafish (Danio rerio genome), which is not very closely related to many wild leuciscid taxa of 45 interest (Fig. 1; Schönhuth et al. (2018)). A recently published fathead minnow (Pimephales 46 promelas) genome provides one resource in the leuciscid family (Martinson et al. 2022), but 47 there is still a dearth of genomic resources for this hyper-diverse and geographically ubiqui-48 tous clade. In consequence, previous genomic studies of creek chub have relied on artificial 49 reference genomes or reference genomes from distantly related taxa (e.g. Meuser et al. 2022). 50

We sequenced the genome of the common creek chub, Semotilus atromaculatus, to pro-51 vide a genomic resource for future studies of creek chub and other leuciscid minnows in North 52 America. We chose to sequence creek chub because of the broad range, abundant popula-53 tion sizes, and generally tolerant life history of this species. Creek chub are expected to 54 have 2n=50-52 chromosomes and a genome size of 1.25pg, similar to many leuciscid species 55 (Gold & Amemiya 1987, Legendre & Steven 1969). We also compared the outcomes of two 56 assembly pipelines: Pacific Biosciences' Improved Phase Assembly (IPA) HiFi Genome As-57 sembler pipeline (github.com/PacificBiosciences/pbipa) and the software HiFiasm (Cheng 58

4

⁵⁹ *et al.* 2021). Finally, we assessed synteny between the creek chub reference genome and the ⁶⁰ zebrafish and fathead minnow genomes (Martinson *et al.* 2022).

$_{61}$ Methods

Sampling was accomplished under animal utilization protocol #4237 approved by the Univer-62 sity of Guelph Animal Care Committee, with permits from the Ontario Ministry of Natural 63 Resources and Forestry (Licence No. 1100698), and with private landowner permission. 64 One wild-caught, 10.6 cm (total length) creek chub was sampled to generate this reference 65 genome (see Fig S1 for a photo). We sampled this fish using a beach seine from Swan Creek 66 in southern Ontario, Canada, in August 2022. The sampled individual was identified mor-67 phologically by expert field personnel and later identified genetically as a creek chub with 68 DNA barcoding. Following capture, we euthanized the target individual with an overdose 69 of MS-222, then sampled and flash froze muscle tissue in liquid nitrogen within 5 minutes of 70 euthanasia to ensure preservation of high molecular weight DNA. We stored the flash-frozen 71 muscle and remainder of the specimen in a -80°C freezer, except for 2 fin clips preserved 72 in 95% ethanol. We extracted DNA from the fin clips using a DNeasy Blood & Tissue 73 kit (Qiagen) and quantified the concentration using a NanoDrop 8000 Spectrophotometer 74 (Thermo Scientific). We used this DNA to verify our phenotypic identification with DNA 75 barcoding at the University of Guelph's Advanced Analysis Center. The COI-3 region of 76 the mitochondrial genome was amplified and sequenced using thermalcycler conditions and 77 primers from Ivanova et al. (2007). The forward fasta sequence was input on BOLD's Identi-78 fication Engine (www.barcodinglife.org/index.php/IDS_OpenIdEngine; (Ratnasingham 79 & Hebert 2007)) and confirmed to belong to creek chub. 80

We sent flash frozen muscle tissue to the University of Delaware's DNA Sequencing 81 & Genotyping Center, in Newark, Delaware, USA. High molecular weight (HMW) DNA 82 extraction was completed using the MagAttract HMW DNA kit (Qiagen), then the extracted 83 DNA was quantified using a Qubit Fluorimeter and DNA fragment sizes were assessed by 84 Femto Pulse system instrument (Agilent). Next, a Megaruptor 2 (Diagenode) was used to 85 shear 3 μ g of DNA to 15kb fragments. Then, a SMRTbell DNA library was constructed 86 according to the Pacbio HiFi SMRTbell protocol using SMRTbell Express Template Prep 87 Kit 3.0 (Pacbio, 102-182-700). After BluePippin size selection (Sage Science, PAC20KB) 88 removed fragments smaller than 8 kb, the average size in the library was 18 kb based on 89 Femto Pulse System (Agilent) analysis. Finally, sequencing was performed on 2 SMRT 8m 90 cells on Sequel IIe instrument with 30 hours movie, using both the Sequel II Binding kit 2.2 91 and Sequel II Sequencing kit 2.0. 92

The initial assembly of the reference genome was performed by the University of Delaware's DNA Sequencing & Genotyping Center. They used Pacific Biosciences' Improved Phase Assembly (IPA) HiFi Genome Assembler pipeline (github.com/PacificBiosciences/pbipa). In addition, we used HiFiasm (Cheng *et al.* 2021) to create a genome assembly. This, and all subsequent computation, was performed on Digital Research Alliance of Canada's Cedar high performance computing cluster. We ran HiFiasm (v0.16.1) with 32 CPUs to create a

5

⁹⁹ HiFi-only assembly, as we did not have parental short reads or Hi-C reads to create either ¹⁰⁰ the trio-binning or Hi-C integrated assemblies (Cheng *et al.* 2021).

We assessed genome assembly quality using custom R and shell scripts to quantify distri-101 bution of assembled contig and scaffold lengths, and the number of unique assembled scaffolds 102 (Fig. 2). From these data, we calculated N50, N90, L50, and L90, as well as maximum, 103 mean, and median contig length (Table 1). As this assembly is comprised completely of long-104 read PacBio data, there are no gaps in our assembled contigs, and we hereafter refer to these 105 fragments of the genome simply as contigs. We also ran this analysis on the most recent ver-106 sions of the zebrafish (GCF_000002035.4_GRCz11) and fathead minnow (GCF_016745375, 107 Martinson *et al.* 2022). We assessed completeness of the creek chub reference genome using 108 BUSCO v5.2.2 with actinopterygii_odb10 as the database (Simão et al. 2015), as well as the 109 zebrafish and fathead minnow reference genomes for the sole purpose of comparison with the 110 same database. We used kraken 2 v2.1.2 (Wood et al. 2019) to assess contamination using a 111 custom database containing virus, plasmid, protozoa, archaea, bacteria, human, plant, and 112 fungi sequences. 113

We examined synteny between creek chub and zebrafish, using SynMap, from the plat-114 form CoGe (Comparative Genomics, genomevolution.org, Lyons & Freeling 2008). CoGe 115 DAGChainer outputs were used to create circular plots with circos (Krzywinski et al. 116 2009). We used a hard masked version of the genome, uploaded to CoGe (NCBI Window-117 Masker (Hard) (v1.0,id65989;genomic). The exact zebrafish organism used was Danio rerio 118 (zebrafish; id43752) and the genome was unmasked (v11, id66058; CDS). This is the 110 most recent zebrafish geneome (GRCz11) created by the Reference Genome Constortium, re-120 leased May 9, 2017 (ncbi.nlm.nih.gov/assembly/GCA_000002035.4). All default analysis and 121 display options were used in the Legacy Version, with the exception of Syntenic Path Assem-122 bly (SPA) being selected, contigs without synteny hidden, diagonals coloured by syntenic 123 block, contigs sorted by name, and minimum chromosome size set to 2,830,400bp, which 124 is the length of the 50th largest contig in the creek chub assembly. This SynMap analysis 125 can be generated at any time at this link: genomevolution.org/r/loxpo. We additionally 126 created a SynMap between only the 25 largest creek chub contigs and the zebrafish genome, 127 by setting the minimum chromosome length to that of the 25 largest contig in the assembly 128 (20,130,130bp, Fig S2). It can be viewed at this link: genomevolution.org/r/10xpw 129

We also used SynMap to assess synteny between creek chub and fathead minnow. The 130 zebrafish genome is more complete than the fathead minnow genome (Martinson et al. 2022). 131 However, creek chub are more closely related to fathead minnow than to zebrafish (Fig. 132 1). The version of the genome used was unmasked (v2,id66042;CDS) of GCA_016745375. 133 recently published by Martinson et al. (2022). We used the same analysis and display options 134 as mentioned above for the zebrafish SynMap. This SynMap analysis can be regenerated 135 by following this link: genomevolution.org/r/10xpx. We also created a SynMap with the 136 25 largest creek chub contigs and the fathead minnow genome (Fig S3). It can be viewed 137 at this link: genomevolution.org/r/loxq3. SynMap labels are based on the creek chub 138 and fathead minnow genomes' contig/scaffold codes from the FASTA headers and many of 139 these codes do not intuitively match the contig/scaffold's corresponding number. See Tables 140 S1 and S2 for a breakdown of the creek chub and fathead minnow genome's contig/scaffold 141 codes and corresponding contig/scaffold numbers. 142

6

We created a simple phylogeny to display the relationship between the creek chub, zebrafish, fathead minnow, and several other model teleost fish (Fig. 1). We created this phylogeny using the R package fishtree (Chang *et al.* 2019), which pulls phylogenetic data from its pre-assembled online database. We added the fish photo with Adobe Photoshop (v22.0.0).

148 Results

PacBio HiFi sequencing on two SMRT cells produced 133GB of raw data in FASTQ for-149 mat. This corresponded to 4,313,794 raw reads with a mean length of 16,406 base pairs, 150 corresponding to a coverage of 64x. An initial genome assembly was constructed by the 151 University of Delaware sequencing facility's bioinformatics team using Pacific Biosciences's 152 IPA pipeline, and resulted in an assembly that consisted of 873 contigs, with mean contig 153 length of 1.257,076, and an N50 of 5.722,762 (Table 1). We then improved upon this initial 154 assembly using the HiFiasm pipeline (Cheng et al. 2021), resulting in an assembly with 239 155 contigs, with a mean contig length of 4,599,676, and an N50 of 30,568,897, which is halfway 156 between the N50 of the zebrafish and fathead minnow genomes (Table 1). BUSCO analy-157 sis indicates that in addition to being highly contiguous, this genome is largely complete. 158 with a score of 98.0% and 97.9% respectively for the HiFiasm and IPA assemblies (Table 159 2). BUSCO values were similar between the two assemblies with the exception of a higher 160 proportion of genes designated as complete and duplicated in the HiFiasm assembly relative 161 to the IPA assembly (2.5% verses 1.6%, respectively), however both values are similar to the 162 fathead minnow and lower than the zebrafish (Table. 2). Kraken2 analysis did not find any 163 contigs to be entirely contaminated with non-creek chub DNA. As the HiFiasm assembly 164 was comparable to or improved over the IPA assembly in all respects - high completeness 165 and low contamination, but fewer contigs, higher N50, and larger mean contig size - we used 166 the HiFiasm assembly for all subsequent analyses. 167

Comparative Genomics (CoGe)'s SynMap (Lyons & Freeling 2008) analysis produced 168 2274 syntenic blocks and 24532 syntenic matches with zebrafish. We see few major chromo-169 somal rearrangements in creek chub relative to zebrafish (Fig. 3). The haploid chromosome 170 number is expected to be the same (n=25) for zebrafish, fathead minnow, and creek chub 171 (Gold & Amemiya 1987), and most contigs in our assembly corresponded in part or whole 172 to zebrafish chromosomes (Fig. 4). While our assembly is less contiguous than the zebrafish 173 genome, in many cases zebrafish chromosomes map to 1–4 larger assembled contigs of the 174 creek chub genome (Fig. 4) and the 50 largest contigs in the creek chub assembly contain 175 just over 95% of the total assembly content (Table 1). 176

Although the large scale pattern is synteny with zebrafish, there are a few regions of the creek chub genome that appear more sharply divergent. In particular, the creek chub contig 9 (the largest complete contig) showed synteny with both chromosomes 4 and 7 in the zebrafish genome (Fig. 4), suggesting there may have been an interchromosomal fusion event in creek chub. Contig 9 also shows two small inversions on the zebrafish chromosome 7 (Fig. 3), indicating intrachromosomal rearrangements. However, as Fig. 3 is coloured by syntenic block, it is obvious that there have been many minor chromosomal rearrangements

7

or mutations. There are no stretches of synteny along any chromosome or contig that are greater than a few dots – with each dot representing a window of 20 genes in which at least 5 genes are syntenic between species. Quantified in a different way, no stretches of synteny along any creek chub contig are greater than 12000 nucleotides, with the average being 407 nucleotides.

In our SynMap analysis between creek chub and fathead minnow (Fig. 5), which produced 189 19230 syntenic matches in 2042 blocks, we can see that there have also been few major 190 chromosomal rearrangements since these species diverged. One obvious rearrangement is 191 potentially fission or fusion events, whereby scaffold 1 of the fathead minnow genome is 192 split between contigs 28 and 42 of the creek chub genome and scaffold 2 is split between 193 contigs 3, 36, and 44 (Fig. 6). The syntenic matches are less continuous and consistent 194 with some of the fathead minnow scaffolds when compared to zebrafish (Fig. 6 versus Fig. 195 4). However, this likely reflects the quality of the fathead minnow and creek chub genomes 196 compared to the zebrafish genome, rather than a closer phylogenetic relationship between 197 zebrafish and creek chub than fathead minnow and creek chub. Indeed, creek chub and 198 fathead minnow are more closely related to one another than to zebrafish (Fig. 1). The 199 three species are contained within the order Cypriniformes, with zebrafish in the family 200 Danioninae and fathead minnow and creek chub in the family Leuciscidae (Schönhuth et al. 201 2018, Stout et al. 2016). The increased number of syntenic matches over numerous different 202 contigs of each species (Fig. 6), as opposed to contained between a few as we see with 203 zebrafish and creek chub (Fig. 4), are most likely due to the lower continuity and quality of 204 annotation of the fathead minnow genome compared to the zebrafish genome. CoGe predicts 205 syntenic genes based off of sequence similarity, but with a lower quality annotation, is more 206 likely to identify transposable elements or repetitive regions as syntenic between genomes, 207 increasing background noise in the SynMap (Lyons & Freeling 2008). 208

²⁰⁹ Discussion

Our de novo sequencing approach relied entirely on PacBio data, which allowed us to suc-210 cessfully assemble sequence data into a relatively small number of longer contigs (n=239 for)211 the HiFiasm assembly; Table 1). While this assembly is not quite chromosome scale, as the 212 expected haploid chromosome number is 25, there are larger scaffolds which likely approach 213 full chromosomes (Fig. 2), and synteny analyses with zebrafish suggest that each creek chub 214 chromosome is likely covered by 1–4 large contigs (Fig. 4). Analyses of completeness with 215 BUSCO confirm that a high proportion of expected genes are included (about 98% for both 216 assemblies), reinforcing that sequencing produced a high quality reference genome. The con-217 tiguity and completeness of this assembly makes it a valuable resource for genomic studies 218 of non-model leuciscid fish. The high contiguity of sequence enabled by PacBio will allow 219 recovery of genetic architecture of traits where linkage of multiple loci might be extremely 220 relevant (for example, examining the genetic basis of sex determination; Meuser et al. 2022). 221 A high quality reference genome will also enable analyses that require whole-genome data. 222 such as identifying inversions between closely related species (Faria et al. 2019), or demo-223 graphic inference (MSMC and PSMC; SFS; ABC; Li & Durbin 2011, Schiffels & Durbin 224

²²⁵ 2014, Beichman *et al.* 2018).

In the interest of constructing the most complete and continuous assembly possible from our data, we used two different assembly pipelines, IPA and HiFiasm (Cheng *et al.* 2021). Using HiFiasm, we successfully reduced the number of contigs from 873 to 239, and increased the N50 roughly six-fold (Table 1, Fig. 2). Much of the improvement in N50 and contig number likely resulted from the linking of multiple long contigs to form contigs that approach chromosome length, which enabled better understanding of synteny with other related species (Fig. 4 and Fig. 6).

As expected, much of the creek chub genome is syntenic with previously published 233 genomes of model organisms, namely zebrafish. However, there are also some rearrange-234 ments, including a number of inversions and regions that are not syntenic with the zebrafish 235 genome. We do not vet know what functions are encoded by those particular regions of 236 the genome, but structural genome changes, especially of the sex determining region, are 237 likely to play a major role in diversification of species-rich clades of fish like the Cyprini-238 formes (Payseur et al. 2018, Huang et al. 2020). One particular region to note in the syntemy 239 analysis was that approximately half of zebrafish chromosome 4 was not conserved between 240 species. This region is a sex determining region in zebrafish, which has been shown to exist 241 in wild – but not lab raised – strains of zebrafish (Wilson et al. 2014). Sex determination 242 systems vary widely across teleost fish species (Bachtrog et al. 2014, Pennell et al. 2018). 243 Creek chub are not known to have a large sex determining region (Meuser et al. 2022) or 244 heteromorphic sex chromosomes (Gold *et al.* 1979), which could be part of the reason why 245 there are no other large regions lacking synteny between the two genomes. 246

While our creek chub genome assembly does not quite have one large contig per chromo-247 some, for each zebrafish chromosome there are 1–4 larger contigs in our genome assembly 248 that are highly syntenic and likely together comprise the creek chub chromosome (Fig. 4). 249 This tells us that our genome is nearly chromosome-resolution, less a few joins between 250 large contigs. This is especially apparent when comparing to the fathead minnow reference 251 genome; our creek chub reference genome has fewer and larger contigs than the fathead 252 minnow genome (Table 1, Fig. 6). While our creek chub genome is not yet annotated, it 253 is certainly nearly complete and of similar quality to other recently published fish genomes 254 (Martinson et al. 2022). 255

The high-quality creek chub reference genome presented in this paper will enable new 256 insights about the evolutionary history and genome function of leuciscid fish species. Initially, 257 we intend to use this reference genome to investigate the effects of anthropogenic disturbance 258 on a suite of leuciscid fish species. Creek chub and a number of closely related species are 259 widely distributed in North America, and are found in disturbed environments, which makes 260 them an ideal study species for assessing impacts of urbanization and agricultural land use 261 on fish species (similar to previous work in other taxa; Miles et al. 2019, Wei et al. 2021). 262 A future goal is to produce a genome annotation, which would allow analysis of functional 263 patterns of genomic variation and gene expression in a more meaningful way. More broadly, 264 we are now entering a new and exciting era for genomics of non-model organisms, when it is 265 possible to move beyond using genomes of model organisms as reference, and gain the more 266 fine-grain insights that can only be obtained with a conspecific or closely related reference 267 genome (Gopalakrishnan et al. 2017). Generating high quality reference genomes is essential 268

9

for quantifying genomic variation across the incredible biodiversity of fishes (Fan *et al.* 2020), and will lead to new insights about the evolution of this species-rich group of vertebrates.

271 Acknowledgements

Computing was accomplished through an allocation from the Digital Research Alliance of 272 Canada to EGM. We would like to thank T. Frauley for assistance with fieldwork, B. Schultz 273 for productive discussions about analysis of synteny, and S.E. McFarlane for manuscript com-274 ments and discussion of what should be in a genome paper. This manuscript was improved 275 by comments from the entire Mandeville lab at University of Guelph. We also thank B. 276 Kingham and O. Shevchenko of the University of Delaware DNA Sequencing & Genotyping 277 Center for coordinating the DNA extraction, library preparation, sequencing, and initial 278 assembly with the IPA pipeline. Finally, we would like to thank to E. Lyons and A. Nelson 279 from CoGe for assistance with creating the SynMap analyses, and the landowners of the 280 property bordering Swan Creek for allowing us access. 281

²⁸² Author contributions

EGM, AVM, and ARP planned the project. ARP and AVM completed field sampling and tissue dissections. AVM and ARP completed the analyses and made the figures, with assistance from EGM. All authors contributed to writing and revising the manuscript.

²⁶⁶ Data Availability Statement

Supplemental files are available at FigShare. File S1 contains a photo of the creek chub used 287 to create the reference genome. File S2 contains a syntenic dot plot between zebrafish and 288 the creek chub assembly's largest 25 contigs. File S3 contains a syntenic dot plot between 289 fathead minnow and the creek chub assembly's largest 25 contigs. File S4 contains a table 290 of the creek chub assembly's contig headers, the associated contig number, and length of 291 the contig in base pairs. File S5 contains a table of the fathead minnow assembly's scaffold 292 code, the associated scaffold number, and length of the scaffold in base pairs. Upon the 293 acceptance of this manuscript, data and scripts used for analysis will be made publicly 294 available in Data Dryad. The genome is available on the NCBI genomes repository, under 295 accession number PRJNA994924. Custom scripts used in this work will be available on 296 Github: github.com/amanda-meuser/CreekChubGenome. 297

²⁹⁸ Conflict of Interest

²⁹⁹ The authors declare no conflict of interest.

300 Funder Information

This research was undertaken using a Resources for Research Groups (RRG) computing allocation from the Digital Research Alliance of Canada. Sequencing for this project was funded by the Canada First Research Excellence Fund, specifically, University of Guelph's Food From Thought Research Support grant.

11

305 **References**

- Bachtrog D, Mank JE, Peichel CL, et al. (2014) Sex Determination: Why So Many Ways of
 Doing It? PLoS Biology, 12, e1001899.
- ³⁰⁸ Beichman AC, Huerta-Sanchez E, Lohmueller KE (2018) Using Genomic Data to Infer His-
- toric Population Dynamics of Nonmodel Organisms. Annual Review of Ecology, Evolution,
 and Systematics, 49, 433–456.
- ³¹¹ Chang J, Rabosky DL, Smith SA, Alfaro ME (2019) An R package and online resource for
 ³¹² macroevolutionary studies using the ray-finned fish tree of life. *Methods in Ecology and* ³¹³ Evolution, 10, 1118–1124.
- ³¹⁴ Cheng H, Concepcion GT, Feng X, Zhang H, Li H (2021) Haplotype-resolved de novo as-³¹⁵ sembly using phased assembly graphs with hifiasm. *Nature Methods*, **18**, 170–175.
- Corush JB, Fitzpatrick BM, Wolfe EL, Keck BP (2021) Breeding behaviour predicts patterns
 of natural hybridization in North American minnows (Cyprinidae). Journal of Evolution ary Biology, 34, 486–500.
- Fan G, Song Y, Yang L, et al. (2020) Initial data release and announcement of the 10,000
 Fish Genomes Project (Fish10K). GigaScience, 9, giaa080.
- Faria R, Johannesson K, Butlin RK, Westram AM (2019) Evolving Inversions. Trends in
 Ecology and Evolution, 34, 239–248, publisher: Elsevier Ltd.
- Gold JR, Amemiya CT (1987) Genome size variation in North American minnows
 (Cyprinidae). II. Variation among 20 species. Genome / National Research Council
 Canada, 29, 481–489.
- Gold JR, Whitlock CW, Karel WJ, Barlow JA (1979) Cytogenetic studies in North American minnows (Cyprinidae). VI. Karyotypes of thirteen species in the genus Notropis.: VI. Karyotypes of thirteen species in the genus *Notropis*. *CYTOLOGIA*, **44**, 457–466.
- Gopalakrishnan S, Samaniego Castruita JA, Sinding MHS, et al. (2017) The wolf reference
 genome sequence (Canis lupus lupus) and its implications for Canis spp. population genomics. BMC Genomics, 18, 495.
- Holm E, Mandrak NE, Burridge ME (2022) A Field Guide to Freshwater Fishes of Ontario.
 3rd edn., Royal Ontario Museum Press.
- Huang K, Andrew RL, Owens GL, Ostevik KL, Rieseberg LH (2020) Multiple chromoso mal inversions contribute to adaptive divergence of a dune sunflower ecotype. *Molecular Ecology*, pp. 1–15.
- Ivanova NV, Zemlak TS, Hanner RH, Hebert PDN (2007) Universal primer cocktails for fish
 DNA barcoding: BARCODING. *Molecular Ecology Notes*, 7, 544–548.
- Krzywinski M, Schein J, Birol *et al.* (2009) Circos: An information aesthetic for comparative
 genomics. *Genome Research*, **19**, 1639–1645.

12

- Legendre P, Steven M D (1969) Denombrement des chromosomes chez quelques cyprins. Le
 Naturaliste Canadien, 96, 913–918.
- Li H, Durbin R (2011) Inference of human population history from individual whole-genome sequences. *Nature*, **475**, 493–496.
- Lou RN, Jacobs A, Wilder AP, Therkildsen NO (2021) A beginner's guide to low-coverage whole genome sequencing for population genomics. *Molecular Ecology*, **30**, 5966–5993.
- Lyons E, Freeling M (2008) How to usefully compare homologous plant genes and chromosomes as DNA sequences: How to usefully compare plant genomes. *The Plant Journal*, **53**, 661–673.
- Mandeville EG, Walters AW, Nordberg BJ, Higgins KH, Burckhardt JC, Wagner CE (2019)
 Variable hybridization outcomes in trout are predicted by historical fish stocking and
 environmental context. *Molecular Ecology*, 28, 3738–3755.
- Martinson JW, Bencic DC, Toth GP, *et al.* (2022) De Novo Assembly of the Nearly Complete Fathead Minnow Reference Genome Reveals a Repetitive but Compact Genome.
- Environmental Toxicology and Chemistry, **41**, 448–461.
- Meuser AV, Pyne CB, Mandeville EG (2022) Limited evidence of a genetic basis for sex de termination in the common creek chub, Semotilus atromaculatus. Journal of Evolutionary
 Biology, 35, 1635–1645.
- Miles LS, Rivkin LR, Johnson MTJ, Munshi-South J, Verrelli BC (2019) Gene flow and genetic drift in urban environments. *Molecular Ecology*, **28**, 4138–4151.
- Narum SR, Buerkle CA, Davey JW, Miller MR, Hohenlohe PA (2013) Genotyping-by sequencing in ecological and conservation genomics. *Molecular Ecology*, 22, 2841–2847,
 iSBN: 1365-294X _eprint: NIHMS150003.
- Payseur BA, Presgraves DC, Filatov DA (2018) Introduction: Sex chromosomes and speci ation. Molecular Ecology, 27, 3745–3748.
- Pennell MW, Mank JE, Peichel CL (2018) Transitions in sex determination and sex chro mosomes across vertebrate species. *Molecular Ecology*, 27, 3950–3963.
- Ratnasingham S, Hebert PDN (2007) BARCODING: bold: The Barcode of Life Data System
 (http://www.barcodinglife.org): BARCODING. Molecular Ecology Notes, 7, 355–364.
- Schiffels S, Durbin R (2014) Inferring human population size and separation history from
 multiple genome sequences. *Nature Genetics*, 46, 919–925.
- Schönhuth S, Vukić J, Šanda R, Yang L, Mayden RL (2018) Phylogenetic relationships and
- classification of the Holarctic family Leuciscidae (Cypriniformes: Cyprinoidei). Molecular
 Phylogenetics and Evolution, 127, 781–799.

13

- ³⁷⁵ Simão FA, Waterhouse RM, Ioannidis P, Kriventseva EV, Zdobnov EM (2015) BUSCO: as-
- sessing genome assembly and annotation completeness with single-copy orthologs. *Bioinformatics*, **31**, 3210–3212.
- Smith EM, Gregory TR (2009) Patterns of genome size diversity in the ray-finned fishes.
 Hydrobiologia, 625, 1–25.
- Stammler KL, McLaughlin RL, Mandrak NE (2008) Streams modified for drainage provide
 fish habitat in agricultural areas. *Canadian Journal of Fisheries and Aquatic Sciences*,
 65, 509–522.
- Stout CC, Tan M, Lemmon AR, Lemmon EM, Armbruster JW (2016) Resolving Cyprini formes relationships using an anchored enrichment approach. *BMC Evolutionary Biology*,
 16, 244.
- Wei X, Huang M, Yue Q, et al. (2021) Long-term urbanization impacts the eastern golden
 frog (*Pelophylax plancyi*) in Shanghai City: Demographic history, genetic structure, and
 implications for amphibian conservation in intensively urbanizing environments. *Evolu- tionary Applications*, 14, 117–135.
- ³⁹⁰ Wilson CA, High SK, McCluskey BM, *et al.* (2014) Wild Sex in Zebrafish: Loss of the ³⁹¹ Natural Sex Determinant in Domesticated Strains. *Genetics*, **198**, 1291–1308.
- Wood DE, Lu J, Langmead B (2019) Improved metagenomic analysis with Kraken 2. Genome
 Biology, 20, 257.

14

³⁹⁴ Tables and Figures

Genome Statistics	Creek Chub (HiFiasm)	Creek Chub (IPA)	Zebrafish (GRCz11)	Fathead Minnow (GCF_016745375)
Number of contigs	239	873	NA	NA
Number of scaffolds	NA	NA	25 (1897 un- placed)	910 *
Longest contig or scaffold (bp)	58,351,558	23,528,990	78,093,715	59,790,976
Mean contig or scaffold length (bp)	4,599,676	1,257,076	873,221	1,170,614
Median contig/scaffold length (bp)	119,920	206,534	146,921	47,256
N50	30,568,897	5,722,762	52,186,027	11,952,773 *
N90	6,569,117	798,807	$339,\!135$	1,205,132
L50	15	49	14	23
L90	39	255	405	126
Percent of total genome as- sembly in 50 largest contigs	95.13	50.77	93.90	73.18
Percent of total genome as- sembly in 25 largest contigs	74.67	32.49	92.69	54.84

Table 1: Genome statistics for both the HiFiasm and IPA genome assemblies and the most recent versions of the zebrafish and fathead minnow genomes. Statistics for each assembly were generated using a custom script written in a combination of both shell and R. NA = not applicable. * Denotes fathead minnow statistics from Martinson *et al.* (2022).

BUSCO v5.2.2 (actinoptery-	Creek Chub	Creek Chub	Zebrafish	Fathead Minnow
gii_odb10)	(HiFiasm)	(IPA)	(GRCz11)	$(GCF_016745375)$
Complete	3566 (98.0%)	3562 (97.9%)	3483 (95.6%)	3524 (96.9%)
Complete and single-copy	3476 (95.5%)	3505~(96.3%)	3434 (94.3%)	3431 (94.3%)
Complete and duplicated	90~(2.5%)	57~(1.6%)	49(1.3%)	93~(2.6%)
Fragmented	25 (0.7%)	28~(0.8%)	57 (1.6%)	52 (1.4%)
Missing	49 (1.3%)	50~(1.3%)	100 (2.8%)	64 (1.7%)

Table 2: BUSCO (benchmarking universal single-copy orthologs) scores for both the HiFiasm and IPA genome assemblies and the most recent version of the zebrafish and fathead minnow genomes. Generated using BUSCO v5.2.2 (database: actinopterygii_odb10). Total number of BUSCO groups searched for each genome: 3640.

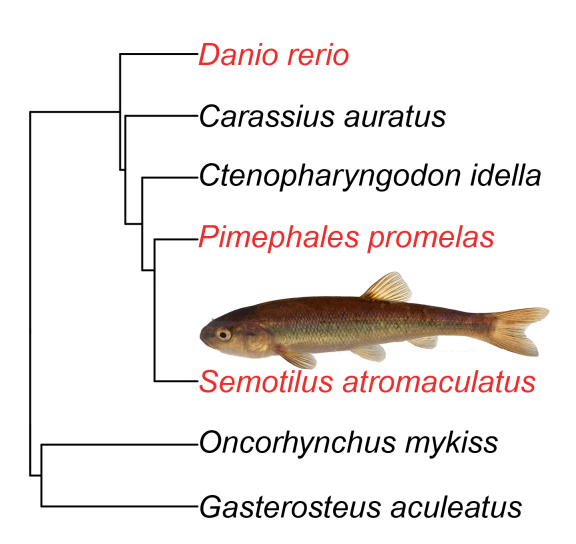


Figure 1: Phylogeny showing the relationship between zebrafish, fathead minnow, creek chub, and other fish commonly used as model species. The inset photo shows a creek chub individual. The phylogeny was created using data from the fishtree package in RStudio (Chang *et al.* 2019).

16

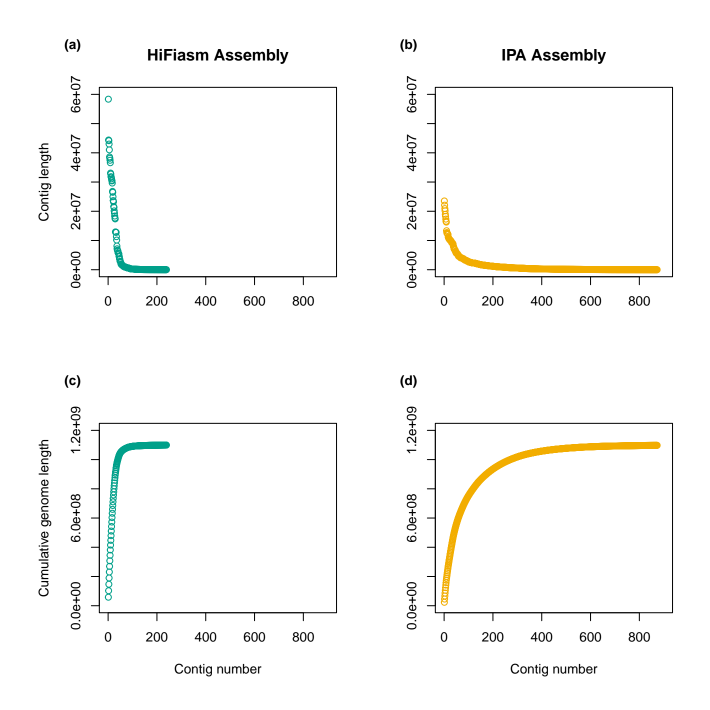


Figure 2: Visual comparison of the number of contigs, contig lengths, and cumulative genome lengths for both the HiFiasm and IPA genome assemblies. (a) Length of each contig in the HiFiasm assembly. (b) Length of each contig in the IPA assembly. (c) Cumulative genome length of HiFiasm assembly. (d) Cumulative genome length of IPA assembly.

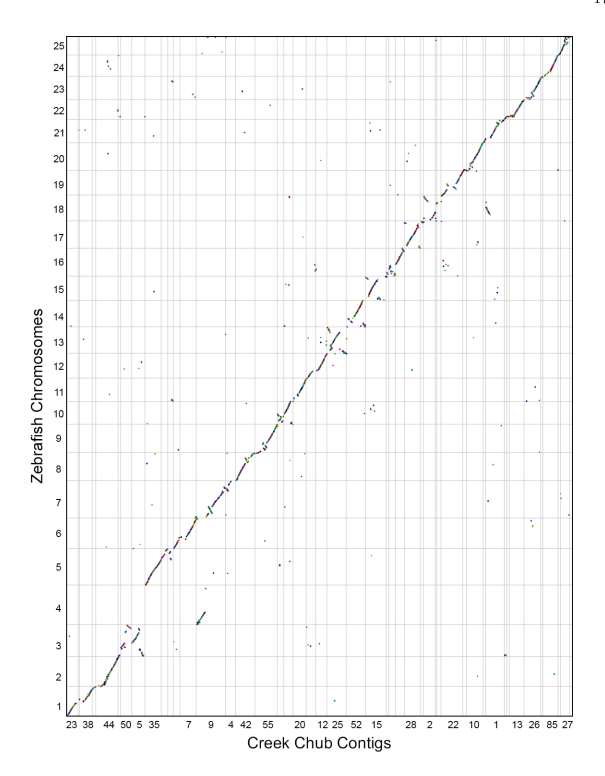


Figure 3: Dot plot showing synteny between creek chub (x-axis) and zebrafish (y-axis). All 25 zebrafish chromosomes from the GRCz11 version of the genome are present, while only the 50 largest contigs from the creek chub have been displayed, by setting the minimum contig length to 2,830,400 base pairs. The dot plot was made using CoGe's SynMap (Lyons & Freeling 2008). Each colour represents a different syntenic block. The figure can be regenerated at any time by following this link: genomevolution.org/r/10xpo

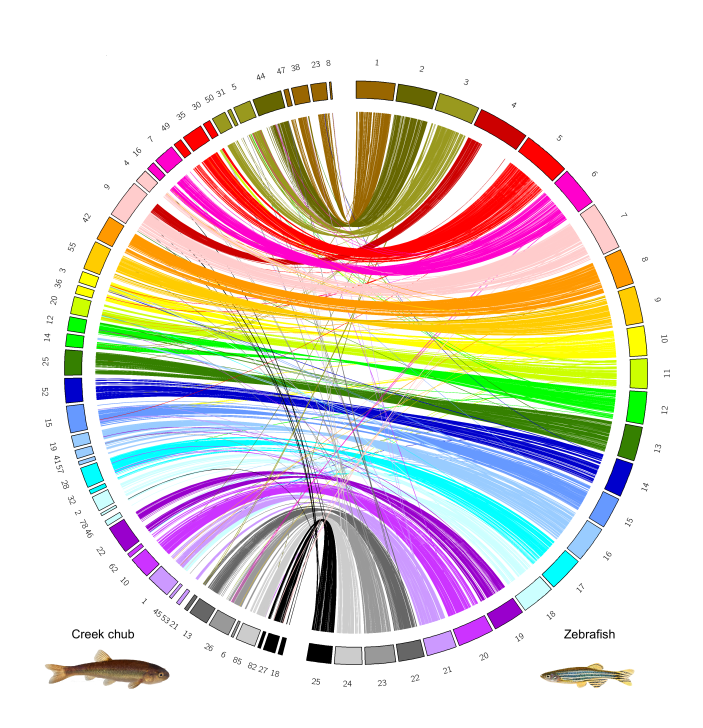


Figure 4: circos plot of syntenic matches between creek chub (left) and zebrafish (right). Creek chub contigs are coloured and mapped to reflect the zebrafish chromosome it has the majority of synetic matches with. Zebrafish photo credit: Mirko_Rosenau.

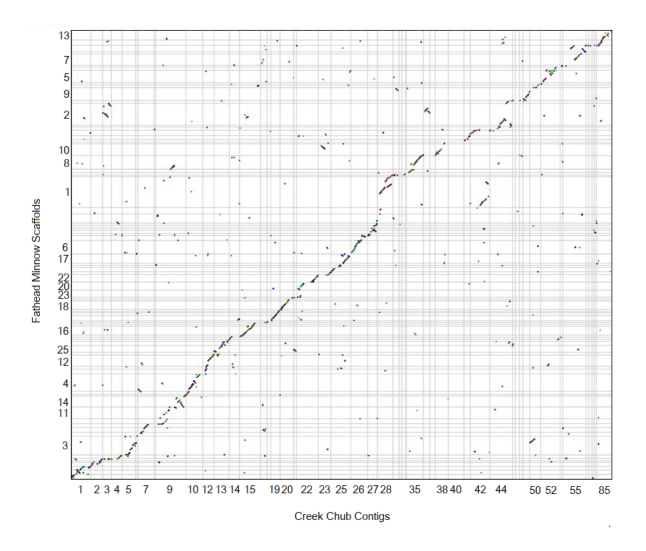


Figure 5: Dot plot made using CoGe's SynMap (Lyons & Freeling 2008) showing synteny between creek chub (x-axis) and fathead minnow (y-axis). Only the 50 largest contigs from the creek chub genome have been displayed, by setting the minimum chromosomes length to 2,830,400 base pairs. Each colour represents a different syntenic block. The figure can be regenerated at any time by following this link: genomevolution.org/r/10xpx

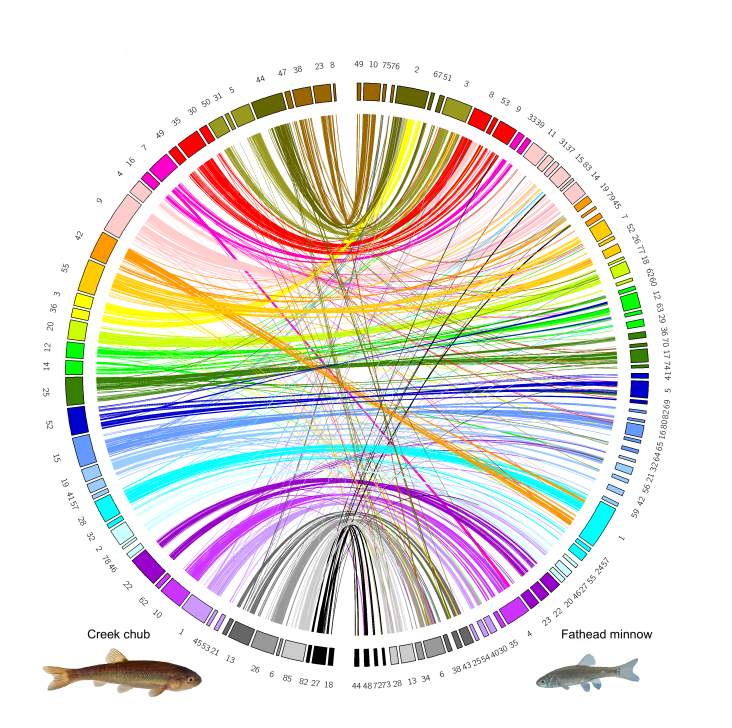


Figure 6: circos plot of syntenic matches between creek chub (left) and fathead minnow (right). Fathead minnow scaffolds are coloured and mapped to reflect the creek chub contig they have the majority of syntenic matches with.

21

³⁹⁵ Supplemental Figures



Figure S1: The creek chub individual used to create the reference genome. This fish was sampled from Swan Creek, Ontario, Canada. Note the dot present at the base of the dorsal fin and intermediate scale size compared to similar species. Not visible in the photo are the small barbels in the groove of each side of the mouth and minimally visible is the large mouth and black "moustache".

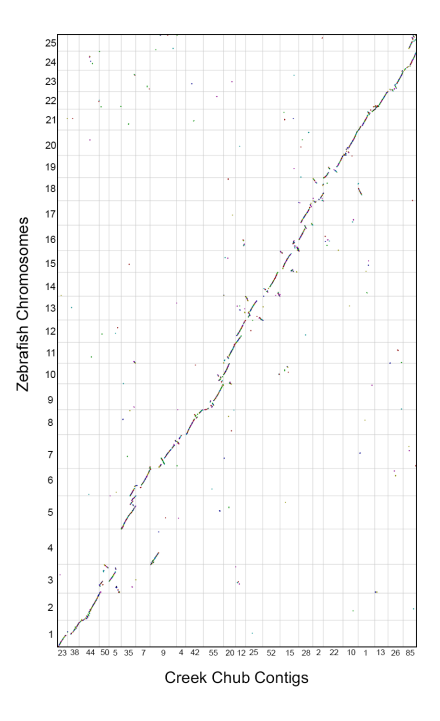


Figure S2: Dot plot made using CoGe's SynMap (Lyons & Freeling 2008) showing synteny between creek chub (x-axis) and zebrafish (y-axis). All 25 zebrafish chromosomes are present, while only the 25 largest contigs from the creek chub have been displayed, by setting the minimum chromosome length to 20,130,130 base pairs. Each colour represents a different syntenic block. The figure can be regenerated at any time by following this link: genomevolution.org/r/10xpw

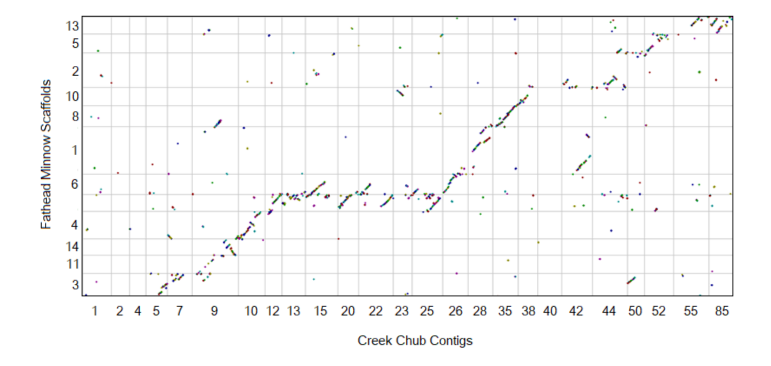


Figure S3: Dot plot made using CoGe's SynMap (Lyons & Freeling 2008) showing synteny between creek chub (x-axis) and fathead minnow (y-axis). Only the 25 largest contigs from the creek chub genome have been displayed, by setting the minimum contig length to 20,130,130 base pairs. Each colour represents a different syntenic block. The figure can be regenerated at any time by following this link: genomevolution.org/r/10xq3