bioRxiv preprint doi: https://doi.org/10.1101/246173; this version posted April 18, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

Title: High quality whole genome sequence of an abundant Holarctic odontocete, the harbour
 porpoise (*Phocoena phocoena*)

3

4 Authors: Marijke Autenrieth<sup>1</sup>, Stefanie Hartmann<sup>2</sup>, Ljerka Lah<sup>1,3</sup>, Anna Roos<sup>4</sup>, Alice B.

5 Dennis<sup>1</sup>, Ralph Tiedemann<sup>1</sup>\*

6

7 <sup>1</sup> University of Potsdam, Institute of Biochemistry and Biology, Evolutionary Biology &

- 8 Systematic Zoology, 14476 Potsdam, Germany
- 9 <sup>2</sup> University of Potsdam, Institute of Biochemistry and Biology, Evolution and Adaptive
- 10 Genomics, 14476 Potsdam, Germany
- <sup>3</sup> current address: Novartis BTDM Mengeš, Kolodvorska 27, SI-1234 Mengeš, Slovenia (All
- 12 work in regards to this manuscript was performed at University of Potsdam<sup>1</sup>)

<sup>4</sup> Swedish Museum of Natural History, SE-104 05 Stockholm, Sweden

14

15 **\*Corresponding author**: Prof. Dr. Ralph Tiedemann tiedeman@uni-potsdam.de

16

bioRxiv preprint doi: https://doi.org/10.1101/246173; this version posted April 18, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.

### 18 Abstract

19

20 The harbour porpoise (Phocoena phocoena) is a highly mobile cetacean found in waters across the Northern hemisphere. It occurs in coastal water and inhabits water basins that vary 21 22 broadly in salinity, temperature, and food availability. These diverse habitats could drive 23 differentiation among populations; population structure within the north Atlantic (north of 51° 24 latitude) is not fully resolved, particularly in relation to Baltic Sea populations. Here we report 25 the first harbour porpoise genome, assembled *de novo* from a Swedish Kattegat individual. The genome is one of the most complete cetacean genomes currently available, with a total 26 27 size of 2.7 Gb, and 50% of the total length found in just 34 scaffolds. Using the largest 28 scaffolds, we were able to examine chromosome-level rearrangements relative to the genome 29 of the closest related species available, domestic cattle (Bos taurus). The draft annotation comprises 22,154 predicted gene models, which we further annotated through matches to 30 31 NCBI nucleotide database, GO categorization, and motif prediction. To infer the adaptive 32 abilities of this species, as well as their population history, we performed Bayesian skyline 33 analysis of the genome, which is concordant with the demographic history of this species, 34 including expansion and fragmentation events. Overall, this genome assembly, together with 35 the draft annotation, represents a crucial addition to the limited genetic markers currently 36 available for the study of porpoise and cetacean conservation, phylogeny, and evolution.

## 37 Introduction

As an apex predator, the harbour porpoise (Phocoena phocoena) is a key indicator for 38 39 conservation and biodiversity measurements in the Nordic Seas (Hooker & Gerber, 2004; 40 Lawrence et al., 2016; Sergio et al., 2008). Marine mammals in particular face many threats 41 from their environment (Fietz et al., 2013; Godard-Codding et al., 2011) including noise pollution (Dyndo et al., 2015; Nabe-Nielsen et al., 2014), marine debris and by-catch 42 (Scheidat et al., 2008; Unger et al., 2017), predation by grey seals (Leopold et al., 2014), and 43 infectious diseases (Siebert et al., 2001; van Beurden et al., 2017). These threats impact 44 structure, boundaries, and stability of populations. This is especially true in the 45 46 Kattegat/Baltic Sea area, where broad ecological shifts have occurred on a relatively short 47 time scale. Since forming 15,000 years ago, the Baltic has undergone periods of brackish, 48 marine, and completely fresh water, and encountered increasing and continuous humans 49 impacts including eutrophication, pollution and overharvesting (Korpinen et al., 2012; 50 Paasche et al., 2015; Ukkonen et al., 2014; Varjopuro et al., 2014). This geological history has 51 created a series of challenges to marine species, and has likely fostered local adaption and 52 population differentiation.

53 Harbour porpoises are the most abundant costal cetaceans across their wide distribution 54 from sub-polar to temperate waters in the Northern hemisphere (Fontaine et al., 2017; Gaskin, 55 1984). As one of the smallest marine mammals, they belong to the Delphinoida and are the sister group to the Monodontidae (Gatesy et al., 2013; Geisler et al., 2011; Hassanin et al., 56 57 2012). Three subspecies of harbour porpoise, P. p. vomerina (North Pacific), P. p. relicta 58 (Black Sea) and P. p. phocoena (North Atlantic), can be differentiated genetically (Rosel et al., 1999), but also by morphological traits including body size and diet (Fontaine et al., 2017; 59 60 Galatius et al., 2012).

The population size of harbour porpoises in European Atlantic Shelf waters is estimated to be 375,000 with shifts across the last decade in the exact regions they occupy (e.g. in the North Sea; (Hammond et al., 2013). Estimates of population size in the western Baltic Sea are smaller, approximately 40,000 animals (Benke et al., 2014; Scheidat et al., 2008; Viquerat et al., 2014). The Baltic Sea proper population, which is not included in the former surveys, has very low estimates (below 500 individuals; Amundin, 2016) and is considered critically endangered (Benke et al., 2014; Hammond et al., 2008; Scheidat et al., 2008).

68 As with other marine mammals in the Northern Atlantic, e.g. grey and harbour seals, subpopulations of the harbor porpoise arose during the end of the last glacial period as North 69 70 Sea populations recolonized the Baltic Sea (Fietz et al., 2016). Now these different 71 populations show shifts in habitat use based largely on food availability (Hammond et al., 72 2013) and activity patterns (Nuuttila et al., 2017), and display fine scale morphological and genetic differences (Fontaine et al., 2012, 2014; Wiemann et al., 2010) and significant 73 74 isolation by distance (Lah et al., 2016). Recent studies based on morphometric and genetic 75 data suggest that different ecotypes of harbour porpoise in the North Atlantic and Baltic Sea 76 exist and may need further conservation measures (Fontaine et al., 2014, 2017; Galatius et al., 77 2012).

These fine scale differences in morphology and behavior may constitute local adaptation, yet the genes underlying such a potentially adaptive differentiation are still unknown and would be best investigated on a whole-genome scale. To examine this, there is a need for high quality genomic resources for this species. A genome will also allow for a broader investigation of population structure, demographic history, functional, and evolutionary questions, as has been shown for other cetacean species in recent studies (Foote et al., 2016; Keane et al., 2015; Nery et al., 2013; Sun et al., 2013; Yim et al., 2013; Zhou et

al., 2013). To this end, a full genome will enable mapping of so far anonymous nuclear
microsatellite (Wiemann et al. 2010) and SNP (Lah et al. 2016) loci, thus facilitating
population genomic inference.

We present here the first *de novo* assembly of the full genome of the harbor porpoise, scaffolded with *in vitro* proximity ligation data (hereafter "Chicago" library), and draftannotated to predict its coding proteins and their functions (Deposited at NCBI as BioProject: PRJNA417595 with BioSample-ID: SAMN08000480). We demonstrate chromosome-level homology with other Cetartiodactyla (Gatesy et al., 2013), and insight into past population dynamics using a Bayesian skyline plot (Li & Durbin, 2011).

- 94
- 95

### 96 Materials and Methods

97 DNA sampling

98 Tissue for whole genome sequencing came from a single individual from the Kattegat 99 (Glommen - Falkenberg), Sweden (ID: C2009/02665). Muscle tissue was sampled in July 100 2009 from a by-caught female of probably young age (22.4kg, 110.5m), frozen, and 101 transported to Potsdam, Germany for DNA extraction. Sample preparation and Genomic DNA 102 isolation were performed following the Quiagen DNeasy Blood & Tissue Kit (Cat 69506, 103 Hilden). Successful high molecular weight DNA-isolation was confirmed by Sanger 104 sequencing of the mitochondrial control region, and visualization of fragment sizes of the 105 entire extraction using the Tape Station (Agilent 2200, Santa Clara, CA 95051). By mtDNA 106 sequencing, we verified that the analyzed specimen carried haplotype PHO7 (Tiedemann et 107 al., 1996), indicative of the separated Beltsea population of the Kattegat/Western Baltic Sea 108 region (Lah et al., 2016; Wiemann et al., 2010).

## 109 Genome sequencing and assembly

110 The draft de novo assembly was constructed from two libraries (insert sizes ca. 300 and ca. 111 500bp); sequenced in 125bp PE on the Illumina HiSeq 2500 at EUROFINS Genomics. Reads 112 were trimmed using CUTADAPT v1.10 (Martin, 2011) and an initial assembly was made using 113 SOAPDENOVO2 (Luo et al., 2015). DNA from the same sample was used by Dovetail Genomics for construction of a Chicago library (Putnam et al., 2016), and sequenced in 150bp 114 115 PE reads on an Illumina NextSeq500 at the University of Potsdam. The draft assembly was 116 then scaffolded with the Chicago library results for the final HiRise assembly, performed by 117 Dovetail Genomics.

Presence of core, single copy, and orthologous genes was measured using CEGMA and BUSCO, run in the genome mode for the Laurasiatheria database (Simão et al., 2015). BLOBTOOLS was run to examine potential contaminants, based on divergence in GC-content and read coverage variation across the assembly (Laetsch & Blaxter, 2017).

122

123 *Genome annotation* 

124 Genome annotation was performed by MAKER2 (Holt & Yandell, 2011) in two steps. MAKER2 125 makes use of different programs and draws from several lines of evidence. Prior to 126 annotation, repetitive elements were soft-masked with REPEATMASKER (Smit et al., 2013-2015) using the te protein repeat database (Smith et al., 2007). In the first MAKER2 run, three 127 128 gene predictors were used: SNAP (Bromberg et al., 2008) was ab initio trained with the 129 CEGMA results (Parra et al., 2007), GENEMARK-ES (Ter-Hovhannisyan et al., 2008) was run 130 using an HMM produced by ab initio training on the whole P. phocoena genome, and 131 AUGUSTUS was run using the presets for human, as is recommended for vertebrates (Stanke et 132 al., 2004). Protein sequences, supplied as evidence were obtained from the complete

SwissProt database (553,941 Proteins) plus NCBI entries of 184,527 proteins from eight
different cetacean groups (On 20 March 2017, all hits to following keywords: *"Balaenopteridae"*, *"Lipotes vexillifer"*, *"Neophocaena"*, *"Orcinus orca"*, *"Phocoena"*, *"Physeter catodon"*, *"Pontoporia blainvillei"*, *"Tursiops truncatus"*).

137 For the second MAKER2 run, we created a new SNAP-HMM based on the first MAKER2 output, and ran it with the same parameters as the first run, exchanging only the SNAP HMM 138 139 and excluding the protein evidence. The resulting CDS predictions were extracted from the 140 final gff file, which was created by *fathom* implemented in SNAP (Bromberg et al., 2008). 141 These gene predictions were further verified by a BLASTN search against the entire GENBANK 142 non-redundant nucleotide sequence database (date downloaded 21.07.2017). Summary 143 statistics were generated using GENOME ANNOTATION GENERATOR (Hall et al., 2014). We then 144 used all CDS and their BLAST results in BLAST2GO (Goetz et al., 2008) to identify conserved protein domains with INTERPROSCAN (including a Pfam comparison). We functional 145 146 annotated the CDS with GO terms, which are a controlled vocabulary to describe gene 147 function constantly actualized by the Gene Ontology Consortium (Ashburner et al., 2000; 148 Carbon et al., 2017).

149

### 150 *Comparative genomics*

151 The closest relative with a chromosome-level assembly currently available is the domestic 152 cattle, Bos taurus. To validate our assembly, we compared our scaffolds to the B. taurus 153 chromosomes (assembly UMD 3.1.1 downloaded from NCBI, ACCESSION 154 DAAA0000000). Specifically, the 122 P. phocoena scaffolds of at least 1Mbp were aligned 155 to the *B. taurus* chromosomes using the nucmer software of the MUMMER package v. 3.23 156 (Kurtz et al., 2004). From the coordinates of these alignments, runs of ten or more

157 consecutive matches of each at least 250bp between a given *P. phocoena* scaffold and a *B. taurus* chromosome were extracted using custom perl scripts. Their start and end positions 159 were used to generate a CIRCOS (http://circos.ca/) plot that shows regions of collinearity as 160 well as rearrangements. For the CIRCOS plot, separate ribbons are displayed between a *B. taurus* chromosome and a *P. phocoena* scaffold for consecutive hits that were each no more 162 than 20,000 bp apart. If a hit is more than 20,000bp from the next run of consecutive hits, a 163 new ribbon was started; in total 24,394 separate ribbons were constructed (Figure 1).

164

## 165 *Population genomics*

166 In using genome-wide diploid sequence data it is possible to reconstruct the population 167 history in estimating population sizes through the past (Li et al., 2011). To estimate the 168 demographic history of the individual sequenced, we used the SNP Frequency spectra based 169 on our genome assembly, which is a haploid sequence, and the PE reads used to construct the 170 de novo assembly, prior to Chicago scaffolding (described above, we used both insert sizes). 171 These reads were first mapped back to the final assembly using BWA (Li & Durbin, 2009). 172 SNP data was extracted from the resulting bam files, and variants were extracted using 173 SAMTOOLS vs.1.6. (Li, Handsaker, et al., 2009), and BCFTOOLS (Li, Handsaker, et al., 2009), 174 implemented with the script vcfutils.pl (Li, Handsaker, et al., 2009). This generated a final \*.fq.gz file, which was then used to generate the final Bayesian skyline plot in the PSMC 175 176 package, using perl scripts psmc2history.pl and psmc plot.pl (Li et al., 2011). The parameters 177 of the PSMC analysis were set following the recommendation from the authors (Li & Durbin, 178 2011, https://github.com/lh3/psmc) and we applied a generation time of 10 years (Birkun Jr. & Frantzis, 2008) and a mutation rate of  $2.2 \times 10^{-9}$  year/site (Taylor et al., 2007). 179

## 181 Results

182

## 183 De novo assembly of the P. phocoena genome

Shotgun sequencing produced a total number of 1,268M reads (Table 1), these were used to 184 185 generate a draft assembly with 2.4M scaffolds and an N50 of 33.1kb. This assembly was combined with the Chicago library data (556M read) for final scaffolding by Dovetail 186 187 Genomics (Putnam et al., 2016). The final HiRise assembly from Dovetail contains ca. 2M 188 scaffolds (Table 2) and has a total length of 2.7Gb (N50 of 23.8Mb). The greatest 189 improvements from the addition of the Chicago libraries is in building up the 34 longest 190 scaffolds, which make up approximately half of the entire assembly (Table 2). The CIRCOS 191 plot illustrates the near-completeness of these long scaffolds. We observe almost complete 192 coverage of the cow chromosomes by scaffolds bigger than 1Mb in our assembly (Figure 1). 193 The BUSCO and CEGMA analyses also suggests that we have largely reconstructed the entire 194 genome, and identified 96.9% (91.3% complete) of the 2,586 Eukaryotic and 94.2 (88.7% 195 complete) of the 6,253 Laurasiatheria BUSCO core genes and 90% of the 248 ultra-conserved 196 CEGs (54% complete).

197

## 198 Genome completeness and annotation

The MAKER2 annotation resulted in the prediction of 22,154 coding genes (Table 3). In total 21,750 CDS had a BLAST hit against the nucleotide database, which accounts for 98% of the total CDSs. Of these BLAST hits, 99% account for vertebrate, and these were dominated (90%) by hits to Cetacea (thereof 59% *Tursiops truncatus*, 27% *Orcinus orca*). Further annotation with INTERPROSCAN revealed 250,126 features of these predicted proteins. These comprise hits in several protein domain databases, e.g. 23,319 PFAM protein domains, 37,046

205 PANTHER gene families, 24,538 SUPERFAMILY annotations and 31,114 GENE3D domains.

206 Assignment of the BLAST results to Gene Ontology (GO) categories resulted in 55,143 hits

- 207 across the GO categories (
- 208 Figure 2).
- 209
- 210 Inference of Kattegat/Baltic population history

211 We inferred the population history of the harbour porpoise *P. phocoena* based on one single 212 individual (Li et al., 2011) using the PSMC algorithm, which combines all generated PE read 213 data generated. Between eight and four million years ago the inferred population size  $(N_e)$  was 214 low, around 10,000 individuals (Figure 3). It began to increase slightly at 3Myr, and rose 215 more rapidly around 2Myr, reaching an  $N_e$  of 45,000 during the following 1.5 Myr. The 216 estimated population size peaked approximates 400kyra before it dropped to a quarter of the 217 original size around 100kyrs ago, leading to a very low  $N_e$ , similar to that seen in present day 218 populations (Hammond et al., 2013).

219

## 220 Discussion

221 We present here a high quality de novo genome assembly for the harbour porpoise Pho-222 coena phocoena. With a GC-content of 41.4% and a total length of 2.7 GB, this assembly is 223 comparable to other high quality genomes (Groenen et al., 2013; Zimin et al., 2009). BUSCO 224 and CEGMA gene scans support a near completeness of core genes in the assembly, and sup-225 port that we have largely reconstructed the entire genome. For almost completely covering the 226 chromosomes of the *B. taurus* genome (Figure 1), only 122 scaffolds are needed, including 227 the 34 largest scaffolds representing 50% of the whole genome. Of these largest scaffolds 228 some completely match single *B. taurus* chromosomes, e.g., chromosome 25. Other *B. taurus* 

229 chromosomes are in only 2-3 pieces in our scaffolds, e.g. chromosomes 12, 24. Based on this 230 comparison, we infer that our assembly represents a nearly complete genome of *P. phocoena*, 231 and that our largest scaffolds are nearly-complete chromosomes. The CIRCOS plot also illus-232 trates chromosomal rearrangements between domestic cattle and the harbour porpoise, two 233 species diverged approximately 60Myrs ago within the Cetartiodactyla (Gatesy et al., 2013). 234 These chromosomal rearrangements are seen several times among distinct lineage of Cetarti-235 odactyla (Avila et al., 2015; Kulemzina et al., 2009, 2011; Pauciullo et al., 2014), e.g., com-236 parison between camel, pig and domestic cattle (Balmus et al., 2007).

The number of annotated genes (22,154) is comparable to other published cetacean genomes: 21,459 bottlenose dolphin (Lindblad-Toh et al., 2011), 20,605 minke whale (Yim et al., 2013), 22,711 grey whale (DeWoody et al., 2017). They appear to broadly span key functional gene categories, e.g. biological processes, cellular components and molecular function, both across the annotated GO terms and the INTERPROSCAN analysis. With this information we can directly search for known, respectively key genes, for further investigations, e.g. selection or adaptive traits.

244 The harbor porpoise is estimated to have split from is closest relative ca. 5Myr ago 245 (Gatesy et al., 2013). Interestingly our Bayesian skyline plot (Figure 3) coincides with this 246 date by starting a population expansion around that time point. Around 4.5 Myr ago an 247 expansion occurred, during which time the North Atlantic is known to have cooled, leading to 248 an extinction of 65% of the marine organisms (Stanley, 1995). The harbour porpoise is well 249 known in subarctic regions and some populations (e.g. Greenland) occur in areas which freeze 250 to a large extent during winter (Tolley & Rosel, 2006). Therefore, an extinction of other 251 marine species during a cold water period does not preclude that the harbour porpoise could 252 increase its population size and expand through the Atlantic. During the last interglacial 253 period, Eemian, the inferred  $N_e$  remained relatively high at around 50,000 individuals before, 254 dropping dramatically with the beginning of the last glacial period 100kya. When comparing 255 this pattern to the demographic history of other cetaceans, it is most similar to the bottlenose 256 dolphin (Tursiops truncatus), a related species with a similar North Atlantic distribution 257 (Brüniche-Olsen et al., n.d.; Foote et al., 2016; Yim et al., 2013; Zhou et al., 2013). The newly 258 forming sea ice areas, around 400kya ago, could have led to fragmentation of different 259 populations, and therefore lead to a drop in regional total effective population size in regards 260 to our sample. A potential low population size we see postulated for today would fit to the 261 history of the Baltic Sea and the population status of *P. phocoena* (Johannesson et al., 2011; 262 Johannesson & André, 2006; Ukkonen et al., 2014). Specifically, there is strong evidence for a 263 Western Baltic/Kattegat (i.e., Beltsea) population separated from the North Sea/North Atlantic 264 (Hammond et al., 2013; Lah et al., 2016), which currently counts approximately 40,000 animals (Benke et al., 2014; Scheidat et al., 2008; Viguerat et al., 2014). Our sequenced 265 266 specimen was assigned with high likelihood to this Beltsea population by mtDNA analysis (exhibiting haplotype PHO 7; cf. Tiedemann et al., 1996; Wiemann et al., 2010). 267

268 In this study we present the first whole genome assembly and annotation of the harbour 269 porpoise, at this point the most complete assembly for the Family Phocoenidae. This genome 270 adds to the Cetacean genome collection by supplying important resources for further 271 investigation within the Odontoceti as well as outside the Cetacea. This will provide an 272 invaluable resource for further genetic studies within the harbour porpoise itself, both as a 273 resource for whole-genome investigations into population structure and to identify key genes 274 associated with local adaptation. This genome represents a crucial genetic resource for further 275 investigation in the population genetics and phylogeny on other species of the *Phocoenidae* 

including the currently most rare marine mammal, the almost extinct Vaquita (Phocoena 276

277 sinus) (Taylor et al., 2017), and is hence especially important for conservation efforts.

#### 278 Acknowledgments

279 Financial support came from the Bundesamt für Naturschutz (FKZ # 3514824600), as 280 part of a larger study of population genomics. We thank Prof. Dr. Michael Hofreiter for 281 providing access to the Illumina NextSeq platform. Additional support came from the 282 University of Potsdam. Large-scale computational effort was made possible by computing 283 resources provided by the department of Genetics at University of Potsdam and the High 284 Performance Computing Cluster Orson2, managed by ZIM (Zentrum für 285 Informationstechnologie und Medienmanagement) at the University of Potsdam.

286

288

#### 287 References

- 289 Amundin, M. (2016). SAMBAH - Static Acoustic Monitoring of the Baltic Sea Harbour 290 porpoise. LIFE Project Number LIFE 08 NAT/S/000261 European Commission, 77pp. 291 available at http://www.sambah.org/SAMBAH-Final-Report-FINAL-for-website-April-292 2017.pdf.
- 293 Ashburner, M., Ball, C. A., Blake, J. A., Botstein, D., Butler, H., Cherry, J. M., ... Sherlock, 294 G. (2000). Gene ontologie: Tool for the unification of biology. *Nature Genetics*, 25(1), 295 25-29. doi:10.1038/75556.Gene
- 296 Avila, F., Baily, M. P., Merriwether, D. A., Trifonov, V. A., Rubes, J., Kutzler, M. A., ... 297 Raudsepp, T. (2015). A cytogenetic and comparative map of camelid chromosome 36 298 and the minute in alpacas. Chromosome Research, 23(2), 237-251. doi:10.1007/s10577-299 014-9463-3
- 300 Balmus, G., Trifonov, V. A., Biltueva, L. S., O'Brien, P. C. M., Alkalaeva, E. S., Fu, B., ... 301 Ferguson-Smith, M. A. (2007). Cross-species chromosome painting among camel, cattle, 302 pig and human: Further insights into the putative Cetartiodactyla ancestral karyotype. 303 Chromosome Research, 15(4), 499-515. doi:10.1007/s10577-007-1154-x
- 304 Benke, H., Bräger, S., Dähne, M., Gallus, A., Hansen, S., Honnef, C. G., ... Verfuß, U. K. 305 (2014). Baltic Sea harbour porpoise populations: Status and conservation needs derived 306 from recent survey results. Marine Ecology Progress Series, 495, 275–290. 307 doi:10.3354/meps10538
- Birkun Jr., A. A., & Frantzis, A. (2008). Phocoena phocoena ssp. relicta. The IUCN Red List 308 309
- of Threatened Species 2008: e.T17030A6737111. Downloaded on 12 October 2017.
- 310 Bromberg, Y., Yachdav, G., & Rost, B. (2008). SNAP predicts effect of mutations on protein 311 function. Bioinformatics, 24(20), 2397-2398. doi:10.1093/bioinformatics/btn435
- 312 Brüniche-Olsen, A., Westerman, R., Kazmierczyk, Z., Vertyankin, V. V., Godard-Codding, C.,

313 Bickham, J. W., & DeWoody, J. A. (n.d.). The inference of gray whale (Eschrichtius 314 robustus) population attributes from whole-genome sequences. 315 Carbon, S., Dietze, H., Lewis, S. E., Mungall, C. J., Munoz-Torres, M. C., Basu, S., ... 316 Westerfield, M. (2017). Expansion of the gene ontology knowledgebase and resources: 317 The gene ontology consortium. Nucleic Acids Research, 45(D1), D331–D338. 318 doi:10.1093/nar/gkw1108 DeWoody, J. A., Fernandez, N. B., Brüniche-Olsen, A., Antonides, J. D., Doyle, J. M., San 319 320 Miguel, P., ... Bickham, J. (2017). Characterization of the gray whale (Eschrichtius 321 robustus) genome and a genotyping array based on single nucleotide polymorphisms in 322 candidate genes. Biological Bulletin, 232(June), 186-197. 323 Dyndo, M., Wiśniewska, D. M., Rojano-Doñate, L., & Madsen, P. T. (2015). Harbour 324 porpoises react to low levels of high frequency vessel noise. Scientific Reports, 5, 11083. 325 doi:10.1038/srep11083 Fietz, K., Galatius, A., Teilmann, J., Dietz, R., Frie, A. K., Klimova, A., ... Olsen, M. T. 326 327 (2016). Shift of grey seal subspecies boundaries in response to climate, culling and 328 conservation. Molecular Ecology, 25(17), 4097-4112. doi:10.1111/mec.13748 329 Fietz, K., Graves, J. A., & Olsen, M. T. (2013). Control Control Control: A Reassessment and 330 Comparison of GenBank and Chromatogram mtDNA Sequence Variation in Baltic Grey 331 Seals (Halichoerus grypus). PLoS ONE, 8(8), 1–7. doi:10.1371/journal.pone.0072853 Fontaine, M. C., Roland, K., Calves, I., Austerlitz, F., Palstra, F. P., Tolley, K. A., ... Aguilar, 332 333 A. (2014). Postglacial climate changes and rise of three ecotypes of harbour porpoises, 334 Phocoena phocoena, in western Palearctic waters. Molecular Ecology, 23(13), 3306-335 3321. doi:10.1111/mec.12817 336 Fontaine, M. C., Snirc, A., Frantzis, A., Koutrakis, E., Oztürk, B., Oztürk, A. a, & Austerlitz, 337 F. (2012). History of expansion and anthropogenic collapse in a top marine predator of 338 the Black Sea estimated from genetic data. Proceedings of the National Academy of 339 Sciences of the USA, 109(38), E2569-76. doi:10.1073/pnas.1201258109 340 Fontaine, M. C., Thatcher, O., Ray, N., Piry, S., Brownlow, A., Davison, N. J., ... Goodman, 341 S. J. (2017). Mixing of porpoise ecotypes in southwestern UK waters revealed by genetic 342 profiling. Royal Society Open Science, 4(3), 160992. doi:10.1098/rsos.160992 343 Foote, A. D., Vijay, N., Ávila-Arcos, M. C., Baird, R. W., Durban, J. W., Fumagalli, M., ... 344 Wolf, J. B. W. (2016). Genome-culture coevolution promotes rapid divergence of killer 345 whale ecotypes. Nature Communications, 7(May), 11693. doi:10.1038/ncomms11693 346 Galatius, A., Kinze, C. C., & Teilmann, J. (2012). Population structure of harbour porpoises in 347 the Baltic region: evidence of separation based on geometric morphometric comparisons. 348 Journal of the Marine Biological Association of the United Kingdom, 92(8), 1669–1676. 349 doi:10.1017/S0025315412000513 350 Gaskin, D. (1984). The harbour porpoise *Phocoena phocoena* (L.): regional populations, 351 status, and infromation on direct and indirect catches. Reports of the International 352 Whaling Commission, 34, 569–586. 353 Gatesy, J., Geisler, J. H., Chang, J., Buell, C., Berta, A., Meredith, R. W., ... McGowen, M. R. 354 (2013). A phylogenetic blueprint for a modern whale. Molecular Phylogenetics and 355 Evolution, 66(2), 479–506. doi:10.1016/j.ympev.2012.10.012 Geisler, J. H., McGowen, M. R., Yang, G., & Gatesy, J. (2011). A supermatrix analysis of 356 357 genomic, morphological, and paleontological data from crown Cetacea. BMC 358 Evolutionary Biology, 11(1), 112. doi:10.1186/1471-2148-11-112

Godard-Codding, C. A. J., Clark, R., Fossi, M. C., Marsili, L., Maltese, S., West, A. G., ...
Stegeman, J. J. (2011). Pacific ocean-wide profile of CYP1A1 expression, stable carbon

361 and nitrogen isotope ratios, and organic contaminant burden in sperm whale skin

- biopsies. *Environmental Health Perspectives*, *119*(3), 337–343.
- 363 doi:10.1289/ehp.0901809
- Goetz, S., Garccia-Gomez, M. J., Terol, J., Williams, T. D., Nagaraj, S. H., Nueda, M. J., ...
  Conesa, A. (2008). High-throughput functional annotation and data mining with the
  Blast2GO suite. *Nucleic Acids Research*, *36*(10), 3420–3435. doi:10.1093/nar/gkn176
- Groenen, M. A. M., Archibald, A. L., Uenishi, H., Tuggle, C. K., Takeuchi, Y., Rothschild, M.
  F., ... Hunt, T. (2013). Analyses of pig genomes provide insight into porcine demography
- and evolution. *Nature*, 491(7424), 393–398. doi:10.1038/nature11622
- Hall, B., DeRego, T., & Geib, S. (2014). GAG: the Genome Annotation Generator (Version
  1.0) [Software]. doi:Available from http://genomeannotation.github.io/GAG
- Hammond, P. S., Bearzi, G., Bjørge, A., Forney, K. A., Karczmarski, L., Kasuya, T., ...
  Wilson, B. (2008). *Phocoena phocoena* (Baltic Sea subpopulation). (errata version
  published in 2016) The IUCN Red List of Threatened Species 2008:
  e.T17031A98831650. Downloaded on 10 October 2017. Retrieved from
- 376 http://www.iucnredlist.org/details/17031/0
- 377 Hammond, P. S., Macleod, K., Berggren, P., Borchers, D. L., Burt, L., Canadas, A., ...
- Vazquez, J. A. (2013). Cetacean abundance and distribution in European Atlantic shelf
  waters to inform conservation and management. *Biological Conservation*, *164*, 107–122.
  doi:10.1016/j.biocon.2013.04.010
- Hassanin, A., Delsuc, F., Ropiquet, A., Hammer, C., Jansen Van Vuuren, B., Matthee, C., ...
  Couloux, A. (2012). Pattern and timing of diversification of Cetartiodactyla (Mammalia,
  Laurasiatheria), as revealed by a comprehensive analysis of mitochondrial genomes. *Comptes Rendus Biologies*, 335(1), 32–50. doi:10.1016/j.crvi.2011.11.002
- Holt, C., & Yandell, M. (2011). MAKER2: an annotation pipeline and genome-database
  management tool for second-generation genome projects. *BMC Bioinformatics*, *12*(1),
  491. doi:10.1186/1471-2105-12-491
- Hooker, S. K., & Gerber, L. (2004). Marine Reserves as a Tool for Ecosystem-Based
  Management : The Potential Importance of Megafauna. *BioScience*, 54(1), 27–39.
  doi:10.1641/0006-3568(2004)054[0027:MRAATF]2.0.CO;2
- Johannesson, K., & André, C. (2006). Life on the margin: Genetic isolation and diversity loss
  in a peripheral marine ecosystem, the Baltic Sea. *Molecular Ecology*, *15*(8), 2013–2029.
  doi:10.1111/j.1365-294X.2006.02919.x
- Johannesson, K., Smolarz, K., Grahn, M., & André, C. (2011). The future of baltic sea
  populations: Local extinction or evolutionary rescue? *Ambio*, 40(2), 179–190.
  doi:10.1007/s13280-010-0129-x
- Keane, M., Semeiks, J., Webb, A. E., Li, Y. I., Quesada, V., Craig, T., ... deMagalhães, J. P.
  (2015). Insights into the evolution of longevity from the bowhead whale genome. *Cell Reports*, 10(1), 112–122. doi:10.1016/j.celrep.2014.12.008
- Korpinen, S., Meski, L., Andersen, J. H., & Laamanen, M. (2012). Human pressures and their
  potential impact on the Baltic Sea ecosystem. *Ecological Indicators*, 15(1), 105–114.
  doi:10.1016/j.ecolind.2011.09.023
- Kulemzina, A. I., Trifonov, V. A., Perelman, P. L., Rubtsova, N. V., Volobuev, V., FergusonSmith, M. A., ... Graphodatsky, A. S. (2009). Cross-species chromosome painting in
  Cetartiodactyla: Reconstructing the karyotype evolution in key phylogenetic lineages. *Chromosome Research*, *17*(3), 419–436. doi:10.1007/s10577-009-9032-3
- 407 Kulemzina, A. I., Yang, F., Trifonov, V. A., Ryder, O. A., Ferguson-Smith, M. A., &
- 408 Graphodatsky, A. S. (2011). Chromosome painting in Tragulidae facilitates the

reconstruction of Ruminantia ancestral karyotype. *Chromosome Research*, 19(4), 531–
539. doi:10.1007/s10577-011-9201-z
Kurtz, S., Phillippy, A., Delcher, A. L., Smoot, M., Shumway, M., Antonescu, C., & Salzberg,
S. L. (2004). Versatile and open software for comparing large genomes. *Genome*

- 413 *Biology*, 5(2), R12. doi:10.1186/gb-2004-5-2-r12
- Laetsch, D. R., & Blaxter, M. L. (2017). BlobTools: Interrogation of genome assemblies. *F1000Research*, 6(1287), 1–15. doi:10.12688/f1000research.12232.1 and *F1000Research*, 6(1287), 1–15. doi:10.12688/f1000research.12232.1 and
- 416 doi:10.5281/zenodo.845347
- Lah, L., Trense, D., Benke, H., Berggren, P., Gunnlaugsson, Þ., Lockyer, C., ... Tiedemann,
  R. (2016). Spatially Explicit Analysis of Genome-Wide SNPs Detects Subtle Population
  Structure in a Mobile Marine Mammal, the Harbor Porpoise. *PLoS ONE*, *11*(10), 1–23.
  doi:10.1371/journal.pone.0162792
- Lawrence, J. M., Armstrong, E., Gordon, J., Lusseau, S. M., & Fernandes, P. G. (2016).
  Passive and active, predator and prey: using acoustics to study interactions between
  cetaceans and forage fis. *ICES Journal of Marine Science*, *73*(8), 2075–2084.
  doi:10.1093/icesjms/fsw013
- Leopold, M. F., Begeman, L., van Bleijswijk, J. D. L., IJsseldijk, L. L., Witte, H. J., & Gröne,
  A. (2014). Exposing the grey seal as a major predator of harbour porpoises. *Proc. R. Soc. B*, 282, 20142429. doi:10.1098/rspb.2014.2429
- Li, H., & Durbin, R. (2009). Fast and accurate short read alignment with Burrows-Wheeler transform. *Bioinformatics*, *25*(14), 1754–1760. doi:10.1093/bioinformatics/btp324
- Li, H., & Durbin, R. (2011). Inference of human population history from individual wholegenome sequences. *Nature*, 475(7357), 493–496. doi:10.1038/nature10231
- Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., ... Durbin, R. (2009).
  The Sequence Alignment/Map format and SAMtools. *Bioinformatics*, 25(16), 2078–
  2079. doi:10.1093/bioinformatics/btp352
- Lindblad-Toh, K., Garber, M., Zuk, O., Lin, M. F., Parker, B. J., Washietl, S., ... Kellis, M.
  (2011). A high-resolution map of human evolutionary constraint using 29 mammals. *Nature*, 478(7370), 476–482. doi:10.1038/nature10530
- Luo, R., Liu, B., Xie, Y., Li, Z., Huang, W., Yuan, J., ... Wang, J. (2015). Erratum:
  SOAPdenovo2: an empirically improved memory-efficient short-read de novo assembler. *GigaScience*, 4(1), 30. doi:10.1186/s13742-015-0069-2
- 441 Martin, M. (2011). Cutadapt removes adapter sequences from high-throughput sequencing
  442 reads. *EMBnet.journal*, 17(1), 10. doi:10.14806/ej.17.1.200
- Nabe-Nielsen, J., Sibly, R. M., Tougaard, J., Teilmann, J., & Sveegaard, S. (2014). Effects of
  noise and by-catch on a Danish harbour porpoise population. *Ecological Modelling*, 272,
  242–251. doi:10.1016/j.ecolmodel.2013.09.025
- 446 Nery, M. F., Gonzalez, D. J., & Opazo, J. C. (2013). How to Make a Dolphin: Molecular
  447 Signature of Positive Selection in Cetacean Genome. *PLoS ONE*, 8(6), 2–8.
  448 doi:10.1371/journal.pone.0065491
- Nuuttila, H. K., Courtene-Jones, W., Baulch, S., Simon, M., & Evans, P. G. H. (2017). Don't
  forget the porpoise: acoustic monitoring reveals fine scale temporal variation between
  bottlenose dolphin and harbour porpoise in Cardigan Bay SAC. *Marine Biology*, *164*(3),
  1–16. doi:10.1007/s00227-017-3081-5
- 453 Paasche, Ø., Österblom, H., Neuenfeldt, S., Bonsdorff, E., Brander, K., Conley, D. J., ...
- 454 Stenseth, N. C. (2015). Connecting the Seas of Norden. *Nature Climate Change*, *5*(2), 455 89–92. doi:10.1038/nclimate2471
- 456 Parra, G., Bradnam, K., & Korf, I. (2007). CEGMA: A pipeline to accurately annotate core

457 genes in eukaryotic genomes. *Bioinformatics*, 23(9), 1061–1067.

- doi:10.1093/bioinformatics/btm071
- 459 Pauciullo, A., Perucatti, A., Cosenza, G., Iannuzzi, A., Incarnato, D., Genualdo, V., ...
- Iannuzzi, L. (2014). Sequential cross-species chromosome painting among river buffalo,
  cattle, sheep and goat: A useful tool for chromosome abnormalities diagnosis within the
  family bovidae. *PLoS ONE*, 9(10). doi:10.1371/journal.pone.0110297
- Putnam, N. H., Connell, B. O., Stites, J. C., Rice, B. J., Hartley, P. D., Sugnet, C. W., ...
  Rokhsar, D. S. (2016). Chromosome-scale shotgun assembly using an in vitro method for
  long-range linkage. *Genome Research*, 26, 342–350. doi:10.1101/gr.193474.115
- 466 Rosel, P. E., Tiedemann, R., & Walton, M. (1999). Genetic evidence for limited trans-Atlantic
  467 movements of the harbor porpoise, *Phocoena phocoena. Marine Biology (Berlin)*,
  468 133(4), 583–591.
- Scheidat, M., Gilles, A., Kock, K. H., & Siebert, U. (2008). Harbour porpoise *Phocoena phocoena* abundance in the southwestern Baltic Sea. *Endangered Species Research*,
  5(2–3), 215–223. doi:10.3354/esr00161
- 472 Sergio, F., Caro, T., Brown, D., Clucas, B., Hunter, J., Ketchum, J., ... Hiraldo, F. (2008). Top
  473 Predators as Conservation Tools: Ecological Rationale, Assumptions, and Efficacy.
  474 Annual Review of Ecology, Evolution, and Systematics, 39(1), 1–19.
- 475 doi:10.1146/annurev.ecolsys.39.110707.173545
- Siebert, U., Wünschmann, A., Weiss, R., Frank, H., Benke, H., & Frese, K. (2001). Postmortem findings in harbour porpoises (*phocoena phocoena*) from the German North and
  Baltic Seas. *Journal of Comparative Pathology*, *124*(2–3), 102–114.
  doi:10.1053/jcpa.2000.0436
- Simão, F. A., Waterhouse, R. M., Ioannidis, P., Kriventseva, E. V., & Zdobnov, E. M. (2015).
  BUSCO: Assessing genome assembly and annotation completeness with single-copy
  orthologs. *Bioinformatics*, *31*(19), 3210–3212. doi:10.1093/bioinformatics/btv351
- 483 Smit, A. F. A., Hubley, R., & Green, P. (n.d.). RepeatMasker Open-4.0.

484 <<u>http://www.repeatmasker.org</u>>.

- Smith, C. D., Edgar, R. C., Yandell, M. D., Smith, D. R., Celniker, S. E., Myers, E. W., &
  Karpen, G. H. (2007). Improved repeat identification and masking in Dipterans. *Gene*,
  389(1), 1–9. doi:10.1016/j.gene.2006.09.011
- 488 Stanke, M., Steinkamp, R., Waack, S., & Morgenstern, B. (2004). AUGUSTUS: A web server
  489 for gene finding in eukaryotes. *Nucleic Acids Research*, *32*(Web Server issue), W309–
  490 W312. doi:10.1093/nar/gkh379
- 491 Stanley, S. M. (1995). 7 Neogene Ice Age in the North Atlantic Region: Climatic Changes,
  492 Biotic Effects, and Forcing Factors. In *Effects of Past Global Change on Life*.
  493 Washington (DC): National Academies: National Research Council (US) Panel on
- Washington (DC): National Academies: National Research Council (US) Panel of
  Effects of Past Global Change on Life.
- Sun, Y. B., Zhou, W. P., Liu, H. Q., Irwin, D. M., Shen, Y. Y., & Zhang, Y. P. (2013). Genomewide scans for candidate genes involved in the aquatic adaptation of dolphins. *Genome Biology and Evolution*, 5(1), 130–139. doi:10.1093/gbe/evs123
- Taylor, B. L., Chivers, S. J., Larese, J., & Perrin, W. F. (2007). Generation length and percent
  mature estimates for IUCN assessments of cetaceans. *Administrative Report LJ-07-01 National Marine Fisheries*, 24. doi:10.1.1.530.4789
- 501 Taylor, B. L., Rojas-Bracho, L., Moore, J., Jaramillo-Legorreta, A., Ver Hoef, J. M.,
- 502 Cardenas-Hinojosa, G., ... Hammond, P. S. (2017). Extinction is Imminent for Mexico's
   503 Endemic Porpoise Unless Fishery Bycatch is Eliminated. *Conservation Letters*, 10(5),
- 504 588–595. doi:10.1111/conl.12331

- Ter-Hovhannisyan, V., Lomsadze, A., Chernoff, Y. O., & Borodovsky, M. (2008). Gene
   prediction in novel fungal genomes using an ab initio algorithm with unsupervised
   training, 1979–1990. doi:10.1101/gr.081612.108
- Tiedemann, R., Harder, J., Gmeiner, C., & Haase, E. (1996). Mitochondrial DNA sequence
   patterns of Harbour porpoises (*Phocoena phocoena*) from the North and the Baltic Sea.
   *Zeitschrift Für Säugetierkunde*, 61, 104–111.
- Tolley, K., & Rosel, P. (2006). Population structure and historical demography of eastern
   North Atlantic harbour porpoises inferred through mtDNA sequences. *Marine Ecology Progress Series*, 327, 297–308. doi:10.3354/meps327297
- 514 Ukkonen, P., Aaris-Sorensen, K., Arppe, L., Daugnora, L., Halkka, A., Lougas, L., ... Stora, J.
  515 (2014). An Arctic seal in temperate waters: History of the ringed seal (*Pusa hispida*) in
  516 the Baltic Sea and its adaptation to the changing environment. *The Holocene*, 24(12),
  517 1694–1706. doi:10.1177/0959683614551226
- 518 Unger, B., Herr, H., Benke, H., Böhmert, M., Burkhardt-Holm, P., Dähne, M., ... Siebert, U.
  519 (2017). Marine debris in harbour porpoises and seals from German waters. *Marine*520 *Environmental Research*, 1–8. doi:10.1016/j.marenvres.2017.07.009
- van Beurden, S. J., Ijsseldijk, L. L., van de Bildt, M. W. G., Begeman, L., Wellehan, J. F. X.,
  Waltzek, T. B., ... Penzes, J. J. (2017). A novel cetacean adenovirus in stranded harbour
  porpoises from the North Sea: detection and molecular characterization. *Archives of Virology*, *162*(7), 2035–2040. doi:10.1007/s00705-017-3310-8
- Varjopuro, R., Andrulewicz, E., Blenckner, T., Dolch, T., Heiskanen, A. S., Pihlajamäki,
  M., ... Psuty, I. (2014). Coping with persistent environmental problems: Systemic delays
  in reducing eutrophication of the Baltic Sea. *Ecology and Society*, *19*(4).
  doi:10.5751/ES-06938-190448
- Viquerat, S., Herr, H., Gilles, A., Peschko, V., Siebert, U., Sveegaard, S., & Teilmann, J.
  (2014). Abundance of harbour porpoises (*Phocoena phocoena*) in the western Baltic,
  Belt Seas and Kattegat. *Marine Biology*, 161(4), 745–754. doi:10.1007/s00227-0132374-6
- Wiemann, A., Andersen, L. W., Berggren, P., Siebert, U., Benke, H., Teilmann, J., ...
  Tiedemann, R. (2010). Mitochondrial Control Region and microsatellite analyses on
  harbour porpoise (*Phocoena phocoena*) unravel population differentiation in the Baltic
  Sea and adjacent waters. *Conservation Genetics*, 11(1), 195–211. doi:10.1007/s10592009-0023-x
- Yim, H.-S., Cho, Y. S., Guang, X., Kang, S. G., Jeong, J.-Y., Cha, S.-S., ... Lee, J.-H. (2013).
  Minke whale genome and aquatic adaptation in cetaceans. *Nature Genetics*, 46(1), 88–
  doi:10.1038/ng.2835
- 541 Zhou, X., Sun, F., Xu, S., Fan, G., Zhu, K., Liu, X., ... Yang, G. (2013). Baiji genomes reveal
  542 low genetic variability and new insights into secondary aquatic adaptations. *Nature*543 *Communications*, 4(2708), 1–6. doi:10.1038/ncomms3708
- Zimin, A. V, Delcher, A. L., Florea, L., Kelley, D. R., Schatz, M. C., Puiu, D., ... Salzberg, S.
  L. (2009). A whole-genome assembly of the domestic cow, *Bos taurus. Genome*
- 546 *Biology*, *10*(4), R42. doi:10.1186/gb-2009-10-4-r42
- 547 548

# 549 Data Accessibility

- 550 The genome assembly, finale genome sequence and the draft annotation are deposit on NCBI
- under BioProject-ID: PRJNA417595 and BioSample-ID: SAMN08000480).
- 552

# 553 Authors Contributions

- R.T. and L.L. designed the study; A.R. provided the sample and associated biological
  information. L.L. performed molecular lab work, S.H. performed initial *de novo* assembly,
- 556 M.A. executed all genome annotations and analyses, M.A., S.H., and, A.B.D. analyzed and
- 557 interpreted the results, M.A. wrote the manuscript. All authors edited and approved the final
- 558 manuscript.
- 559
- 560

# 561 **Tables and Figures**

# 562

# **Table 1** Sequencing statistics of libraries used for the two assemblies.

	Insert (bp)	n reads	Coverage
Illumina Library 300	300	794 M	74.1 X
Illumina Library 500	500	473 M	44.2 X
Chicago library	1-50kb	528 M	87.2 X

## 563

# Table 2 Assembly statistics of the harbour porpoise genome.

	SOAPdenovo assembly	Dovetail HiRise Assembly	
Total length	2,669.6 Mb	2,681.2 Mb	
Scaffolds			
N50 (number/length)	23,685 / 0.032Mb	34 / 23.8Mb	
N90 (number)	159,889	43,146	
Longest	304,733	67,078,619	
Number	2,139,681	2,025,248	

## 564

# Table 3 Genome annotation statistics

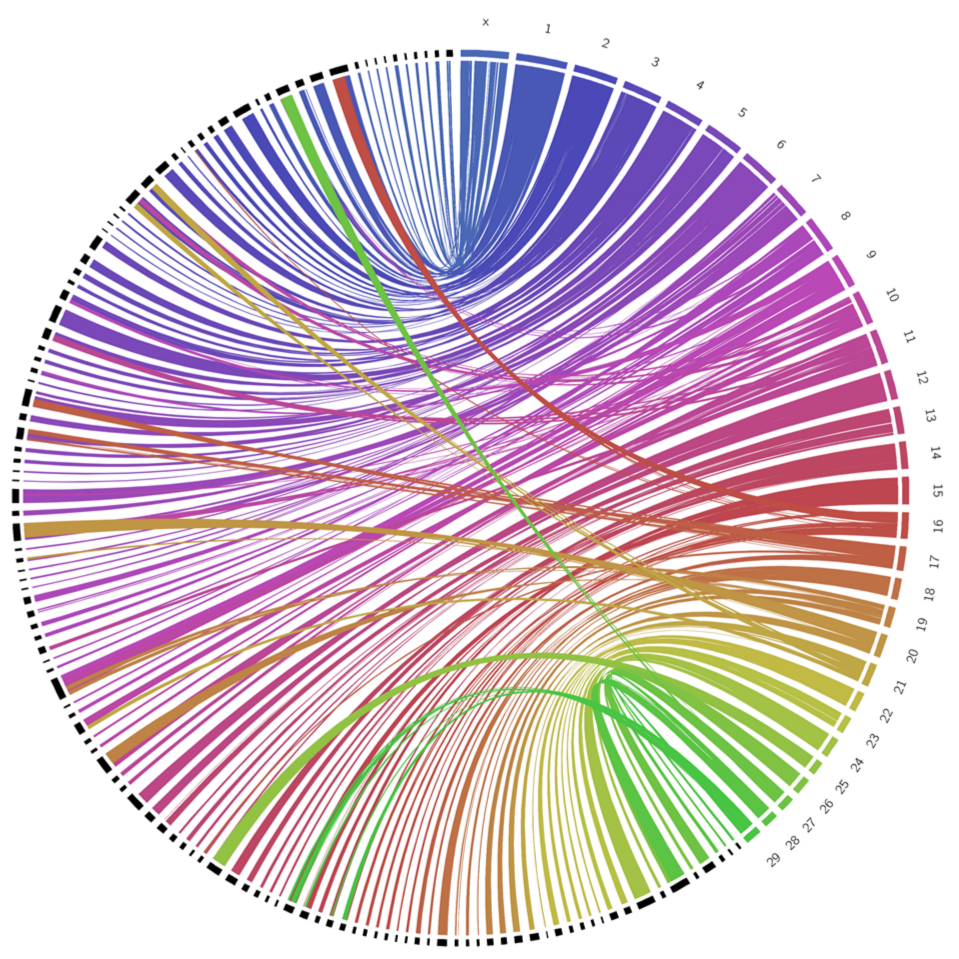
	Exons	Introns	Genes	CDS
Number	171,735	149,581	22,154	22,154
Longest	10,332	1,532,135	2,299,565	20,613
Mean length	166	3,966	28,051	1,282
% genome covered by	-	-	23,2%	1.1%
GC% in CDS	-	-	-	54.48%

565

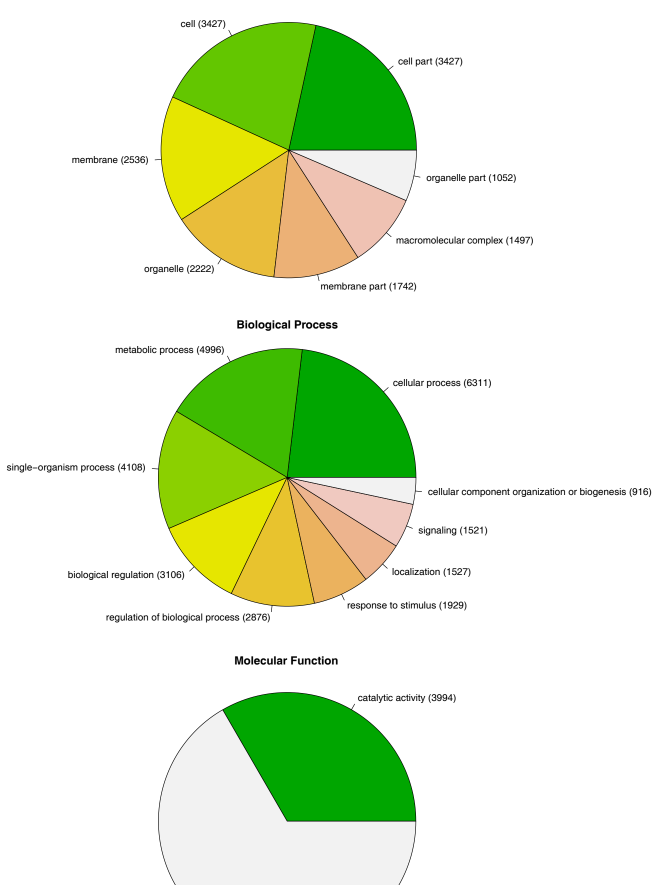
Figure 1 CIRCOS plot of harbour porpoise scaffolds (at least 1Mbp, black on the left outer rim) mapped against the cow (*Bos taurus*) chromosomes (colored bars, labeled X and 1-29).
The *B. taurus* autosomal chromosomes (1-29) as well as the X chromosome (x) are shown in different colors. The 122 largest *P. phocoena* scaffolds with consecutive MUMMER hits between a *B. taurus* chromosome of at least 250bp that are no more than 20,000bp apart are shown in black. Matches between *B. taurus* chromosomes and *P. phocoena* scaffolds are shown in the color of the *B. taurus* chromosomes.

**Figure 2** GO-Terms; GO annotation level 2, separated by gene ontology terms: "Cellular Component", "Biological Process", and "Molecular Function". Separate categories are listed, with the number of hits in parentheses.

**Figure 3** PSMC estimated harbour porpoise population size changes over time for the Baltic Sea. g = generation time;  $\mu =$  mutation rate (per site, per year). Porpoise data generated on the basis of mapping PE reads to whole genome scaffolds during SNP calling.



bioRxiv preprint doi: https://doi.org/10.1101/246173; this version posted April 18, 2018. The copyright holder for this preprint (which was not certified by peer review) is the author/funder. All rights reserved. No reuse allowed without permission.



binding (7983)

Cellular Component

