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Published in:
Scientific Data

Publication date:
2020

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Citation for published version (APA):

Payton, L., Noirot, C., Hoede, C., Hüppe, L., Last, K., Wilcockson, D., Ershova, E. A., Valière, S., & Meyer, B. (2020). Daily transcriptomes of the copepod *Calanus finmarchicus* during the summer solstice at high Arctic latitudes. *Scientific Data*, 7(1). <https://doi.org/10.1038/s41597-020-00751-4>

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DATA DESCRIPTOR

Daily transcriptomes of the copepod *Calanus finmarchicus* during the summer solstice at high Arctic latitudes

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The zooplankter *Calanus finmarchicus* is a member of the so-called “Calanus Complex”, a group of copepods that constitutes a key element of the Arctic polar marine ecosystem, providing a crucial link between primary production and higher trophic levels. Climate change induces the shift of *C. finmarchicus* to higher latitudes with currently unknown impacts on its endogenous timing. Here we generated a daily transcriptome of *C. finmarchicus* at two high Arctic stations, during the more extreme time of Midnight Sun, the summer solstice. While the southern station (74.5 °N) was sea ice-free, the northern one (82.5 °N) was sea ice-covered. The mRNAs of the 42 samples have been sequenced with an average of 126 ± 5 million reads (mean \pm SE) per sample, and aligned to the reference transcriptome. We detail the quality assessment of the datasets and the complete annotation procedure, providing the possibility to investigate daily gene expression of this ecologically important species at high Arctic latitudes, and to compare gene expression according to latitude and sea ice-coverage.

Background & Summary

The copepod *Calanus finmarchicus* (Crustacea, Copepoda) is a key zooplankton species in the northern Atlantic food web as it converts sugars from algae into energy rich lipids that sustain higher consumers including marine fish larvae and seabirds^{1–3}. Its high abundance and biomass also makes it an important contributor to ocean carbon flux⁴. The species inhabits a large latitudinal range from ~40° up to 80° N⁵. However, recent findings show that *C. finmarchicus* is undergoing temperature driven geographical shifts northwards because of climate change^{6–8}, the effects of which are at their most extreme in the Northern Atlantic and Barents Sea. Therefore, the copepods will experience a change between the photoperiods they are adapted to at lower latitudes and the extreme high-latitude photoperiods. Photoperiodic variation is particularly pronounced in the Arctic with rapid change over short latitudinal ranges. The impact of such extreme photoperiods on non-endemic species is unknown, and the northward expansion of organisms at high latitudes may be limited by the adaptive capacity of their endogenous timing systems to extreme photoperiods^{8,9}.

Endogenous timing systems, or biological clocks, are ubiquitous ancient and highly adaptive mechanisms enabling organisms to track and anticipate environmental cycles and prepare biological processes accordingly^{10,11}.

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Since the identification of circadian clock genes in *C. finmarchicus*¹², studies have shown that this species possesses a functional clock that might be involved in the timing of both diel and seasonal events, such as the ecologically and biogeochemically important diel vertical migration (DVM)¹³ or diapause¹⁴. However, the Arctic environment is characterized by dramatic seasonality resulting in permanent illumination during Midnight Sun and permanent darkness during Polar Night¹⁵. As the circadian clock is entrained and synchronized by daily light/dark cycles, the persistence of daily biological processes in Arctic organisms during the absence of those remains uncertain^{16,17}, as well as the consequences for newcomer species due to global warming^{8,9}. Moreover, the Arctic is characterized by strong fluctuations in sea ice-cover, reflecting on biotic and abiotic factors, such as species communities and interactions or light penetration^{18,19}.

Copepods are among the important non-model invertebrates for which genomic resources are still limited, one barrier being that many species, including *C. finmarchicus*, have large genomes^{20,21}. The *de novo* transcriptome of *C. finmarchicus*²² represents a useful resource for assessing the impact of global warming in this species of high ecological interest. In addition to differential gene expression analyzes, RNA sequencing has increased the ability to study the expression of rhythmically expressed mRNAs^{23–25}. Indeed, at the molecular level, the endogenous clock machinery drives the rhythmic expression of downstream genes whose rhythmic translation and function ultimately underlie daily oscillations at cellular and organismal levels²⁵. Note that in the field, environmental cycles also directly generate rhythms independently from the clock. Thus, temporal transcriptomic studies allow a major breakthrough in the understanding of daily dynamics of biological processes in the field.

In this study, we performed RNA sequencing on temporally collected *in situ* samples to generate a daily transcriptome of *C. finmarchicus* in the high Arctic during summer solstice period when the sun remains high above the horizon with minimal altitude variation. Sampling of *C. finmarchicus* stage V copepodites was performed at 4 h intervals within a 24 h cycle at two ocean stations along a latitudinal gradient. The northern station (82.5 °N, Nansen Basin) was characterized by sea ice-coverage, while the southern one (74.5 °N, Barents Sea) was sea ice-free. In addition to providing the raw data, we describe its quality assessment and the alignment to the reference transcriptome to verify reliability and determine transcript quantification. Finally a complete annotation is performed and two normalized datasets are provided for further transcriptomic data exploration of this species.

Methods

Sampling design. The sampling strategy was specifically designed for the detection of rhythmic transcripts^{25,26} although it does not exclude classic differential expression analysis²⁷. Sampling design and analysis strategy are presented in Fig. 1, Table 1 and Supplementary Table 1. Sampling covered a complete 24 h cycle at 4 h intervals, resulting in seven time points per station. At each station, samplings were performed at similar time intervals of: 14–15 h, 18–19 h, 22–23 h, 2–3 h, 6–7 h, and 10–11 h (all times noted in local time (UTC + 2)). Sampling at “North” station, JR85, started on 18th June (3 days before the summer solstice) at 10–11 h and ended on 19th June at 10–11 h. Sampling at “South” station, B13, started on 30th June (9 days after the summer solstice) at 14–15 h and ended on 1st July at 14–15 h. At each timepoint the water column was sampled from 200 m to the surface with vertical hauls of a WP2 plankton net (opening \varnothing : 57 cm, net length: 236 cm, mesh size: 200 μ m) with a meshed bucket cod end (mesh size: 200 μ m) at a speed of 0.5 m*s⁻¹. Transferring the animals from the net into the stabilization solution was done within less than 12 minutes for all samplings. A 12 h period of incubation at 2–4 °C was allowed to soak the samples thoroughly with the RNAlater stabilization solution (Ambion, UK) before they were transferred to –80 °C for further transport and storage.

Sites description. Sampling has been conducted during Cruise JR17006 of the RRS *James Clark Ross* in summer 2018 at two stations along a latitudinal gradient. The station “North” was sea ice-covered and located in the Nansen Basin (JR85; 82.56°N, 30.85°E). The station “South” was sea ice-free and located in the southern Barents Sea (B13; 74.5°N, 30°E). Water depth at “North” was 3700 m and at “South” was 360 m. The sun’s altitude was always above the horizon but still showed diel oscillations of altitude above the horizon from 16 ° at midnight to 30.9 ° at midday at “North”, and from 7.7 ° at midnight to 38.6 ° at midday at “South” at the times of sampling (all times noted in local time (UTC + 2)). Sites were exposed to semidiurnal tide regimes, i.e., 2 tides per day, with a maximum amplitude of ± 0.47 m at JR85 and ± 0.36 m at B13 at times of sampling. Maps with the location of the sea ice edge at the time of sampling at “North” are available from the *meereisportal*²⁸ (https://data.meereisportal.de/gallery/index_new.php?active-tab1=method&ice-type=satellite&satellite=A®ion=n&resolution=daily&minYear=2018&minMonth=6&minDay=18&maxYear=2018&maxMonth=6&maxDay=19&show-Maps=y&dateRepeat=n&submit2=display&lang=en_US&active-tab2=satellite). Modeled data of sun altitude were obtained from the United States Naval Observatory (<https://aa.usno.navy.mil/data/docs/AltAz.php>, USNO, USA). Information on the tidal dynamics have been drawn from the TPX08 model²⁹ by using the OTPS package (Tidal Prediction Software, <http://www-po.coas.oregonstate.edu/~poa/www-po/research/po/research/tide/index.html>), via the mbotps program³⁰ (MB-System). Solar altitude, tidal height and sea-ice cover during the sampling campaign at both latitudes are detailed in Supplementary Table 2. Temperature, pressure (depth), conductivity (salinity), oxygen saturation (SBE 43, Sea-Bird Electronics) and Chlorophyll *a* (Chl *a*) fluorescence (Aquatracka III fluorometer, Chelsea Technologies Group, UK) were measured from the surface to 200 m depth and are available in Hueppe *et al.*³¹.

Copepod sorting. Copepods were sorted at 2 °C under a stereo microscope for species (*C. finmarchicus*) and stage (CV). To distinguish *C. finmarchicus* from its closely related congener *C. glacialis*, morphological indicators were used, in particular the redness of the antenna, which has been shown to be a good indicator in the regions of sampling³²; see also the molecular validation of morphological identification, below. For each timepoint and station, 3 replicates of 15 *C. finmarchicus* CV were sorted. The choice to pool 15 individuals was made to (1) get the sufficient amount of RNA required for RNA sequencing and quantitative real-time PCR analyses and (2)

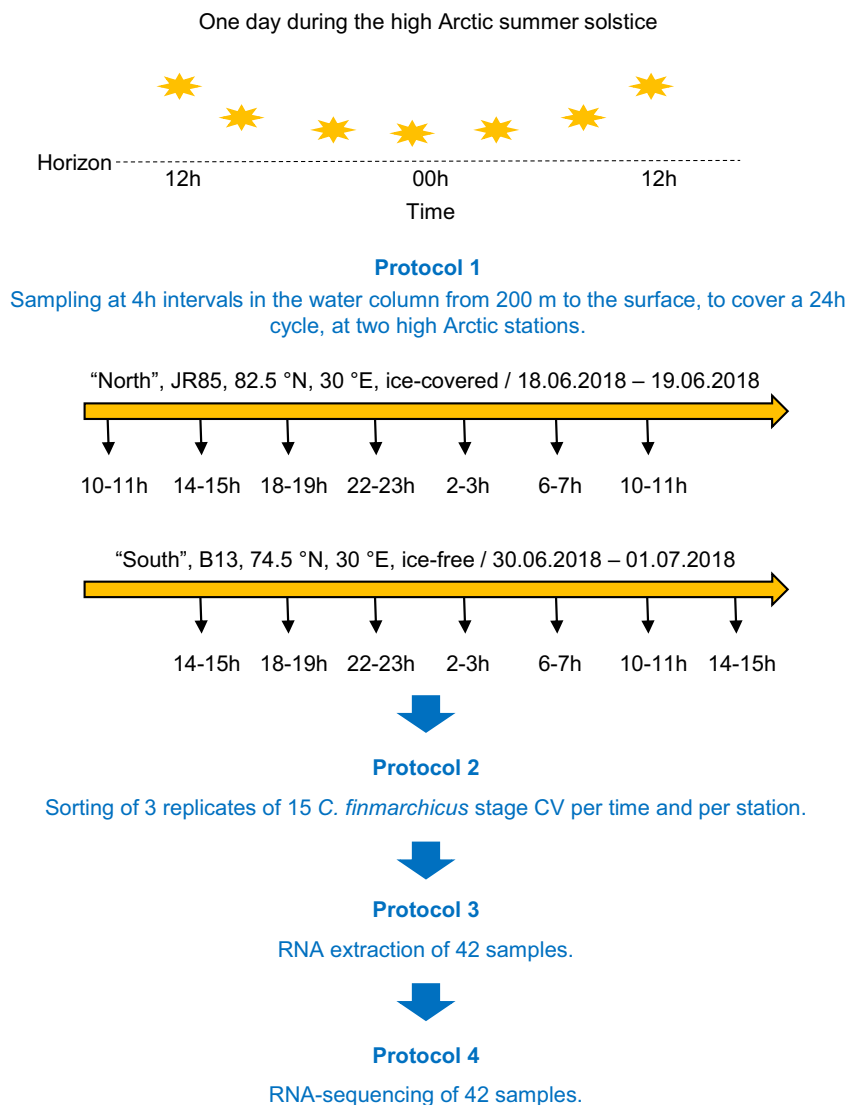


Fig. 1 Overview of the experimental workflow used to generate the transcriptomic data output (all times noted in local time (UTC + 2)). Sample details are available in Supplementary Table 1.

| Station | Number of timepoint | Sampling frequency | Total duration of sampling | Number of replicate per timepoint | Total number of samples | Sequencing strategy | Reads | Platform |
|---------|---------------------|--------------------|----------------------------|-----------------------------------|-------------------------|---------------------|--------------------------|------------------|
| “North” | 7 | 4h | 24h | 3 | 21 | RNA-seq | paired-end 2 × 150 pb | Illumina NovaSeq |
| “South” | 7 | 4h | 24h | 3 | 21 | RNA-seq | paired-end 2 × 150 pb | Illumina NovaSeq |

Table 1. Summary of sampling and sequencing strategy. Details are available in Supplementary Table 1.

increase the number of individuals (315 copepods per station in total) thereby decreasing the effect of individual variability.

RNA extraction. Each replicate was distributed to a 2 ml Precellys® homogenization tube (Bertin Instruments, France), containing a mix of 1.4 mm and 2.8 mm ceramic beads and homogenized in 600 µl of TRIzol® reagent (ThermoFisher Scientific, USA) with a Precellys® 24 Tissue Homogenizer (Bertin Instruments, France), using two times 15 sec. of homogenization at 5000 rpm with a 10 sec. break between. For RNA extraction, a Phenol/Chloroform based single-step extraction in combination with a spin column based solid phase extraction (Direct-zol™ RNA MiniPrep Kit, Zymo Research, USA) was used. Genomic DNA was removed by DNase I digestion on column as part of the RNA extraction kit and total RNA was eluted in ultra-pure water. A portion of the RNA of each of the samples was used to investigate relative expression of 8 candidate genes with SYBRGreen

| DOI | Availability | File Name | Description | File format |
|---|---|---|--|-------------|
| — | https://www.ncbi.nlm.nih.gov/bioproject/PRJNA628886 | SRR11748365.fastq.gz to SRR11748406.fastq.gz | Raw RNA sequencing data | fastq |
| https://doi.org/10.6084/m9.figshare.c.5127704 | https://doi.org/10.6084/m9.figshare.c.5127704 | PRJNA628886_raw_quantification_206K.tsv.gz | Raw count matrix of the 206,012 contigs | tsv |
| | | 76k_ids_list.txt | List of identifiers corresponding to the 76,550 contigs after filtering | txt |
| | | PRJNA628886_quantification_downsampled_76k.tsv.gz | Down-sampling normalized quantification matrix | tsv |
| | | PRJNA628886_quantification_RLE_76k.tsv.gz | RLE normalized quantification matrix | tsv |
| | | Interprot_annot_206K.gff3 | Gff3 InterProScan annotation of the 206,012 contigs | gff3 |
| | | Interprot_annot_206K.tsv | Tabulated InterProScan annotation of the 206,012 contigs | tsv |
| | | diamond_annotation_206k.tsv | Tabulated DIAMOND annotation matrix of the 206,012 contigs (NR,swissprot,trembl) | tsv |
| | | diamond_annotation_76k.tsv | Tabulated DIAMOND annotation matrix of the 76,550 contigs (NR,swissprot,trembl) | tsv |

Table 2. List of available datasets related to the study (NCBI Bioproject PRJNA628886⁴⁰ and figshare collection 5127704⁴¹). Details are available in Supplementary Table 1.

based quantitative real-time PCR (qPCR) on candidate genes, using the $2^{-\Delta Ct}$ method³² and the geometric mean of *elongation factor 1 α* and *16s rRNA* as reference, as described by Hueppe *et al.*³¹. Another portion of each sample was sent to GeT-PlaGe core facility in dried-ice for RNA sequencing.

RNA sequencing. RNA sequencing was performed at the GeT-PlaGe core facility, INRAE Toulouse. The 42 RNA sequencing libraries were prepared according to Illumina's protocols using the Illumina TruSeq Stranded mRNA sample prep kit to analyse mRNA. Briefly, mRNA were selected using poly-T beads. Then, RNA were fragmented to generate double stranded cDNA and adaptors were ligated to be sequenced. 11 cycles of PCR were applied to amplify libraries. Library quality was assessed using a Fragment Analyser (Advanced Analytical Technologies, Inc., Iowa, USA) and libraries were quantified by qPCR using the Kapa Library Quantification Kit (Roche). RNA sequencing experiments have been performed on a NovaSeq S4 lane (Illumina, California, USA) using a paired-end read length of 2×150 pb with the Illumina NovaSeq Reagent Kits.

Reads alignment and quantification. 42 RNA sequencing libraries were obtained (Fig. 1, Table 1, and Supplementary Table 1). The number of paired reads per library was between 74 million and 276 million with an average of 126 ± 5 million (mean \pm SE) reads. The RNA sequencing libraries reads quality were evaluated using FastQC³³. Contamination was checked by aligning reads against *E. coli*, Yeast and PhiX genomes.

The *Calanus finmarchicus de novo* transcriptome²², based on different life stages and deposited to Bioproject PRJNA236528, was used as the reference transcriptome. It is composed of 206,012 contigs and presents good results of quality assessment, with a nearly complete BUSCO set^{22,34}. Reads were aligned to the *de novo* transcriptome with BWA-MEM (<http://bio-bwa.sourceforge.net/bwa.shtml>). Quantification was performed with SAMtools³⁵ idxStats to generate the quantification matrix. The matrix was filtered with edgeR³⁶ and only contigs with more than 1 CPM (Count Per Million) in at least one sample were kept, providing a matrix of 76,550 contigs. Information on the datasets resulting from this study is available in Table 2.

Annotation. We provided different annotations for all further analysis. Contigs were aligned with DIAMOND³⁷ on NR (2019-09-29), Swissprot and Trembl (2018-12) to retrieve corresponding best annotations. An annotation matrix was then generated by selecting the best hit for each database if: i) the percent of the query length covered by the alignment was higher than 60%; ii) the percent of the subject length covered by the alignment was higher than 40%; iii) the percent of identity of the alignment was higher than 40%. Contigs were also processed with InterProScan³⁸ to scan InterProScan signatures. A GO was assigned to each contig with an InterProScan hit containing a GO annotation. Information on the datasets resulting from this study is available in Table 2. Note that a previous annotation of *Calanus finmarchicus* reference transcriptome²² against Non-redundant (NR) protein database is also available at <https://doi.org/10.5061/dryad.11978>.

Normalization. Two normalizations are proposed (down-sampling normalization and RLE normalization) but the choice of normalization depends on the analysis required downstream. For a rhythmic analysis, we suggest down-sampling the mapped reads to the lowest number among the 42 samples (down-sampling normalization), i.e. to 70.4 million properly mapped reads per sample for all samples (after filtering), in order to adjust for differences in sequencing depth among samples^{23,25,39}. This was performed with StreamSampler.jar (<https://github.com/shenkers/sampling>). EdgeR³⁶ was used to perform RLE normalization, since it is more appropriate for differential expression analysis. Information on the datasets resulting from this study is available in Table 2.

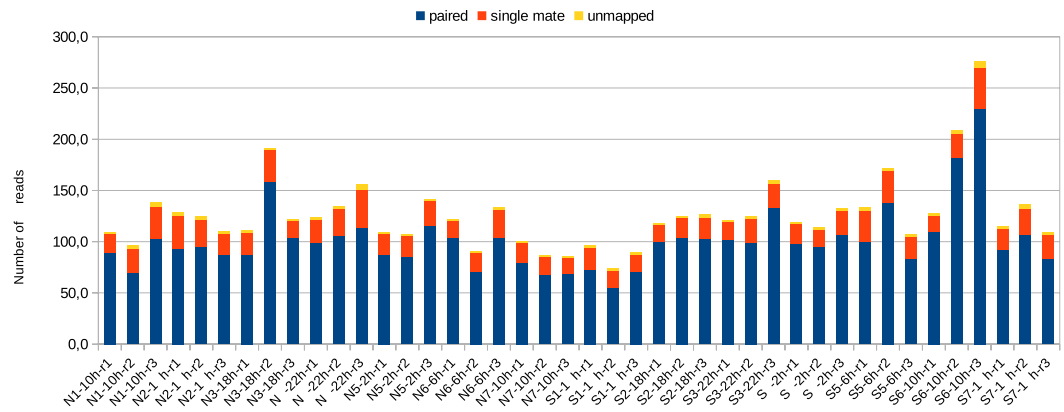


Fig. 2 Mapping statistics of the 42 samples against the set of 206,012 contigs. Number of million (M) reads: paired, single mate, and unmapped. Details are available in Supplementary Table 3.

Data Records

Raw reads were gathered in the NCBI BioProject PRJNA628886⁴⁰ which includes all BioSamples used for the study (Table 2, Supplementary table 1). We also provide the following in figshare collection 5127704⁴¹ (Table 2): the quantification matrix for the 206,012 contigs; the list of identifiers corresponding to the 76,550 contigs after filtering; the two suggested normalization matrices (down-sampling and RLE) and; the datasets annotations (DIAMOND annotation matrix, InterProScan annotation, GO association).

Technical Validation

Molecular validation of morphological identification. Since *C. finmarchicus*' Arctic congener *C. glacialis* also occurs in the region of sampling and differences between the species can be very subtle⁴², morphological identification was validated by molecular species identification on a subset of samples from the same stations^{21,43}. DNA was extracted from individual copepods using the HotShot method⁴⁴, and the species-specific nuclear insertion/deletion (InDel) marker G-150 was amplified using a modified protocol from Smolina *et al.*⁴⁵. Identification was done by accessing the size of the resulting amplicon via electrophoresis on a 2% agarose gel. Results have shown that 99% of the individuals identified as *C. finmarchicus* by the morphological identification method were also clearly identified as *C. finmarchicus* by the molecular identification method, while 0.1% were not clearly identified and 0.7% were identified as the Arctic congener *Calanus glacialis* (n = 305 individuals).

Extraction and RNA integrity. RNA extraction procedures were performed with randomization of samples to ensure reliable and unbiased data production. RNA purity was assessed by OD measurements with a NanoDrop 8000 spectrophotometer (ThermoFisher Scientific, USA), and all 260/280 and 260/230 OD ratio was superior to 1.9. RNA integrity was evaluated with a Fragment Analyzer (Advanced Analytical Technologies, Inc., Iowa, USA; RNA Kit (15nt) Standard Sensitivity, Agilent). Due to a non-conventional 28 S/18 S ribosomal ratio in this species, sample quality was evaluated on the electropherogram⁴⁶. No degradation in the inter region was observed. Total RNA samples were stored at -80°C .

Raw reads assessment and quantification overview. All samples passed the FastQC³³ "base quality control". No relevant contamination hit was found after the alignment against *E. coli*, Yeast and PhiX. The mapping rate against the reference transcriptome²² of 206,012 contigs was higher than 72.4% for properly paired reads and higher than 93.6% considering both paired and single mate reads, validating the raw reads quality (Fig. 2, Supplementary Table 3). Furthermore, over the 42 samples, the maximal percentage of multi-mapped alignment is of 3.31% (Fig. 2, Supplementary Table 3).

For an overview of the quantification matrix, a principal component analysis (PCA) was generated on the raw pseudo-count ($\log_2(\text{count} + 1)$) non-normalized matrix (Fig. 3). Results showed a clear separation between samples from "North" and "South" stations, indicating environmental variations that might be due to latitude and/or sea ice-coverage.

Filtering. Of the 206,012 transcripts, 37% (76,550) were expressed above the threshold of 1 CPM. This result corroborates previously observed results on the *C. finmarchicus* transcriptome²². Thus a large proportion of the whole contigs (63%) exhibited an extremely low level of expression, representing only $1.32 \pm 0.04\%$ of total aligned reads at "North", and $1.27 \pm 0.05\%$ at "South" (Table 3, Supplementary Table 4).

Contigs annotation. By selecting the best hit for each database, the annotation matrix generated with Diamond³⁷ has led to 36,274 and 22,527 contigs with an annotation in at least one database out of the 206,012 and 76,550 contigs respectively (Table 4). Moreover, the number of unique hits for each database is always lower than the number of contigs annotated by the respective database, highlighting the contigs' functional redundancies.

The InterProScan annotation provided annotations from many protein signature databases. The main results are presented in Table 5 and Supplementary Table 5. A GO was attributed to 65,924 contigs over the whole transcriptome (206,012 contigs), while 33,057 contigs out of the 76,550 contigs with an expression level higher than 1 CPM in at least one sample had a GO annotation (Table 5, Supplementary Table 5).

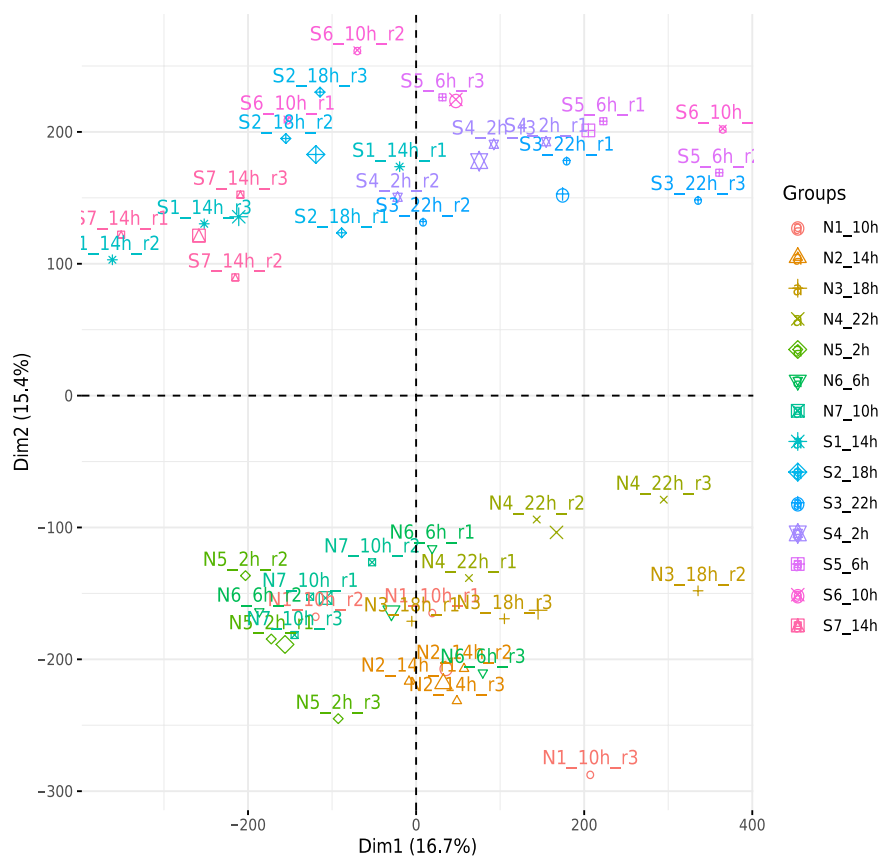


Fig. 3 Principal component analysis of the 42 samples based on expression levels of the whole transcriptome (206,012 contigs). “Groups” grouping the 3 replicates per time and per station.

| Station | Average number of aligned reads on full dataset (206,012 contigs) | Average number of aligned reads after filtering (76,550 contigs) | Percentage of aligned reads discarded by filtering |
|---------|---|--|--|
| “North” | 117.9 ± 5.4 | 116.3 ± 5.3 | 1.32 ± 0.04% |
| “South” | 129.9 ± 9.3 | 128.3 ± 9.2 | 1.27 ± 0.05% |

Table 3. Average number (million, mean ± SE) of alignments per station and percentage of alignments discarded by filtering contigs with very low expression. Details are available in Supplementary Table 4.

| Dataset | Database | Number of contigs | Number of unique hit in target database |
|-----------------|---|-------------------|---|
| 206,012 contigs | NR | 32,637 | 16,165 |
| | Trembl | 28,845 | 15,518 |
| | Swissprot | 11,668 | 5,486 |
| | With an annotation in at least one database | 36,274 | — |
| 76,550 contigs | NR | 21,171 | 11,045 |
| | Trembl | 17,988 | 10,051 |
| | Swissprot | 8,456 | 4,343 |
| | With an annotation in at least one database | 22,527 | — |

Table 4. Number of contigs annotated with DIAMOND against NR, TREMBL and Swissprot and number of unique hits in the target database.

Quantitative real-time PCR data for normalization verification. The relative expression of six core circadian clock genes (*clock*, *cycle*, *period1*, *timeless*, *cryptochrome2*, *vriille*) and 2 circadian clock-related genes (*cryptochrome1* and *doubletime2*) was investigated by quantitative real-time PCR and are available in Supplementary Table 6, allowing the verification of RNA sequencing normalization for further investigations. Regarding the two normalizations, the down-sampling normalization was selected for a rhythmic analysis based on concordant temporal expression profiles with qPCR data (using RAIN algorithm⁴⁷), while the RLE

| Dataset | Main programs | | | Extracted GO |
|-----------------|---------------|---------|---------|--------------|
| | PFAM | TMHMM | getorf | |
| 206,012 contigs | 63,661 | 123,542 | 192,688 | 65,924 |
| 76,550 contigs | 34,558 | 48,682 | 73,272 | 33,057 |

Table 5. Number of contigs with an InterProScan annotation and details on main features. Details are available in Supplementary Table 5.

normalization has been validated for differential expression analysis of the mean level of expression between stations, using the 21 samples of each stations as replicates.

Usage Notes

We present here the first *in situ* daily transcriptomes from the high Arctic, where molecular investigations of biological rhythms are exceptionally limited^{15,16}. The samplings have been realized during drastic Polar photic conditions, i.e. the summer solstice, when daily oscillations of the Sun are minimal, high in the sky and always above the horizon¹⁵. The proposed datasets are thus novel and of interest due to the unique geographical location and time of year, the ecological importance of *C. finmarchicus*, and the rigorous temporal sampling strategy. Another strength of this dataset is the high depth of the RNA sequencing, with an average of 126 ± 5 million of reads (mean \pm SE) per sample, which optimizes the detection of rhythmic transcripts²⁵ in a species with a large genome^{20,21}. Finally, the elaborate annotation of the large transcriptome is now publicly available and is thus accessible for further research.

The sampling strategy is optimized for rhythmic analysis, and particularly adapted for RAIN algorithm analysis^{23,25,47}. Moreover, dataset allows powerful differential gene expression analysis using the 21 samples per station as replicates providing time-integrated detection of differentially expressed genes in *C. finmarchicus* with latitude/sea ice-cover. With climate driven environmental changes, this dataset ultimately constitutes new insights into transcriptomic regulation in the northward migrating copepod *C. finmarchicus*.

Code availability

Parameters to software tools involved are described in the following paragraph.

FastQC: version 0.11.2, --nogroup --casava.

DIAMOND: version v0.9.22, parameters: -f 6 qseqid qlen qcovhsp pident score evaluate length sseqid slen stitle.

InterProScan: version 5.29-68.0, --goterms -t n -dp -f TSV, gff3 parameters.

BWA: version 0.7.17, standard parameters, mem algorithm.

SAMtools programs (view, sort, index and idxStats, flagstat): version 1.8, standard parameters.

EdgeR: version 3.26.5.

StreamSampler.jar: version 1.0.

Received: 5 August 2020; Accepted: 29 October 2020;

Published online: 24 November 2020

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Acknowledgements

This work was supported by the CHASE project, part of the Changing Arctic Ocean programme, jointly funded by the UKRI Natural Environment Research Council (NERC, project number: NE/R012733/1) and the German Federal Ministry of Education and Research (BMBF, project number: 03F0803A). We thank the NERC PRIZE cruise leader Professor Finlo Cottier (Scottish Association for Marine Science, UK) as well as the Captain and crew of the *RRS James Clark Ross* for their support during the cruise JR17006. Cruise time was supported by the CAO Arctic PRIZE project (NERC: NE/P006302/1). EE was supported by Arctic SIZE, a project co-funded by UiT The Arctic University of Norway and the Tromsø Research Foundation (project number 01vm/h15), and within framework of the state assignment of IO RAS (theme No. 0149- 2019- 0008). We would like to thank Vittoria Roncalli (Stazione Zoologica Anton Dohrn, Italy) for useful discussions. Open Access funding enabled and organized by Projekt DEAL.

Author contributions

L.P. designed the study, coordinated RNA extraction, contributed to data analysis and wrote the manuscript; C.N. and C.H. performed reads quality assessment, reads alignment on transcriptome, transcriptome annotation and validation and wrote the manuscript; L.H. designed the study, collected samples, extracted RNA and reviewed the manuscript; K.L. and D.W. designed the study, collected samples and reviewed the manuscript; E.E. identified the copepod species on a genetic level and reviewed the manuscript; S.V. performed the samples preparation for RNA sequencing; B.M. designed the study, reviewed the manuscript and supervised the study.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information is available for this paper at <https://doi.org/10.1038/s41597-020-00751-4>.

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