Nematodes can survive in a suspended form of life for indefinite time

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Abstract

When environmental conditions are unfavorable, such as the complete absence of water or oxygen, high temperature, freezing or extreme salinity, some organisms can enter suspended animation (cryptobiosis)¹. This reversible transition is preceded by execution of complex genetic and biochemical programs (preconditioning)^{2,3,4}. Under laboratory conditions, however, animals have only been maintained in a viable cryptobiotic state for a short time. Here we show that desiccation followed by freezing allows C. elegans dauer larvae to retain full viability over very long periods (around 500 days). Consistent with this finding, recently nematode individuals have been reanimated from the Siberian permafrost⁵, that according to precise radiocarbon dating shows that they remained in cryptobiosis since the late Pleistocene, for about 46,000 years. Phylogenomic inference based on our high-quality genome assembly and morphological analysis demonstrate that these nematodes belong to a novel parthenogenetic species, which we named Panagrolaimus kolymaensis. Genome analysis revealed that the core of the molecular toolkit for cryptobiosis in P. kolymaensis and C. elegans is orthologous. To survive desiccation and freezing under laboratory conditions these two species display similar biochemical responses. Thus, nematodes possess extraordinarily robust adaptive mechanisms that potentially allow them to remain in suspended animation over geological time scales.

Main

Organisms from various taxonomic groups can survive extreme and otherwise lethal conditions such as the complete absence of water or oxygen, high temperature, freezing or extreme salinity. The survival strategies of such organisms include a state known as suspended animation, or cryptobiosis, in which they reduce their metabolism to an undetectable level¹. Spectacular examples of long-term cryptobiosis include a *Bacillus* spore that was preserved in the abdomen of bees buried in amber for 25 to 40 million years⁶, and a 1000 to 1500 years-old *Lotus* seed found in an ancient lake that was subsequently able to germinate⁷. Metazoans such as tardigrades, rotifers and nematodes are also known for remaining in cryptobiosis for prolonged periods^{8,9}. In 2018, we discovered that nematodes from the Siberian permafrost could be revived to life after dwelling in permafrost for many tens of millenia⁵.

Investigation of the molecular mechanisms that allow species to enter suspended animation was greatly facilitated by the finding that *C. elegans*, a well-established genetic model organism, can employ a metabolically quiescent stage known as a dauer larva to enter cryptobiosis²⁴. It has been shown that *C. elegans* possess a general program to survive different kinds of abiotic stress^{3,4}. However, so far, it has only remained possible to maintain larvae in this state for relatively short periods in the laboratory without significant reductions in viability.

Here, we show that desiccation followed by freezing allows dauer larvae to retain full viability after prolonged periods in cryptobiosis. In parallel, we further investigated reanimated animals from permafrost. Precise radiocarbon dating indicates that they remained in cryptobiosis for about 46,000 years, since late Pleistocene. Genome assembly and morphological analysis define that these nematodes belong to new species (*Panagrolaimus kolymaensis*). We show that *P. kolymaensis* utilise similar mechanisms as *C. elegans* to survive desiccation and freezing under laboratory conditions. Our findings suggest nematodes possess adaptive mechanisms to survive in suspended animation for an indefinite time.

C. elegans dauer larvae can remain in a cryptobiotic state for prolonged periods of time Our previous work on *C. elegans* dauer larvae led to the identification of several biochemical pathways that are required for entry into cryptobiosis^{3,4,10}. This process is facilitated by exposure to mild desiccation (98% RH, relative humidity, 4 days) known as preconditioning^{10, ¹¹ in which the transcriptome, proteome, and metabolic pathways undergo a specific remodelling that enhances survival^{3, 4, 10, 12,}. Despite of preconditioning, the survival ability of desiccated dauer larvae at room temperature declines very rapidly, with most larvae dead after around 10 days (Fig.1a). Direct freezing at -80°C, on the other hand, leads to almost instant death of the animals⁴. To test whether combining these conditions could extend the viability of} dauer larvae, we transferred the desiccated larvae to -80°C. Remarkably, under these conditions, there was no significant decline in viability even after 480 days (Fig. 1a). Moreover, after thawing, the animals resumed reproductive growth and produced progeny in numbers similar to those of animals kept under control conditions (Fig. 1b). Since no reduction in survival has been observed at any point in extended experiments, these results suggest that the combination of anhydrobiosis and freezing can prolong the survival and reproductive ability of dauer larvae indefinitely.

Discovery of reanimated nematodes from Permafrost: site and radiocarbon dating

Our finding that *C. elegans* dauer larvae can remain viable for apparently indefinite periods following desiccation and freezing is surprising given that these nematodes are normally found in mild climatic zones. This ability, however, must be present in species that are exposed to harsher environment, for instance in polar regions or permafrost (perennially frozen sediments). Intensive research during the last decades has demonstrated that permafrosts are indeed unique ecosystems that preserve life forms at sub-zero temperatures over thousands of years^{13,14,15,1617,18}. Interestingly, the possibility to exploit permafrost as a source for reanimating multicellular animals was already recognized in 1930s. P.N. Kapterev, as a GULAG prisoner, found a viable cladoceran crustacean *Chydorus sphaericus* preserved in the Transbaikalian permafrost for presumably several thousand years^{19,20}. Unfortunately, this observation remained unnoticed for many decades.

It is well known that several species of *Panagrolaimus* can undergo several forms of cryptobiosis, namely anhydrobiosis (life without water) and cryobiosis (life at freezing)^{10,11, 21,22,23,24}. As mentioned above, in 2018 we showed that nematodes from the Siberian permafrost with morphologies consistent with the genera *Panagrolaimus* and *Plectus* could be reanimated thousands of years after they had been frozen in the permafrost⁵. Several viable nematode individuals were found in two out of more than 300 permafrost samples of different ages and genesis collected by researchers from the Soil Cryology Lab, Pushchino, Russia, in the course of perennial paleo-ecological expeditions carried out in the coastal sector of the northeastern Arctic¹⁷. Detailed descriptions of the study site (outcrop Duvanny Yar, Kolyma River, Fig.2a), sampling and revitalizing procedures are provided in Supplementary Information. Like other late Pleistocene permafrost formations in the northeastern Arctic, Duvanny Yar comprises permanently frozen ice-rich silt deposits riddled with large polygonal ice wedges and divided by them into mineral blocks^{25,26} (Fig.2b). The sediments include sandy alluvial layers, peat lenses, buried paleosols and Pleistocene rodent burrows (Fig.2c). The burrow (P-1320), in which *Panagrolaimus* nematodes were found (Fig.2d), was taken from the frozen outcrop wall

about 40 m below the surface and about 11 m above river water level in undisturbed and never thawed late Pleistocene permafrost deposits. The fossil burrow left by arctic gophers of the genus *Citellus* consisted of an entrance tunnel and a large nesting chamber up to 25 cm in diameter²⁵.

The sterility of permafrost sampling and age of cultivated biota were discussed in detail in several reviews^{27,13,28}. Based on previous reports, the age of the organisms found in a burrow is equal to the initial freezing time and corresponds to the age of organic matter conserved in the syncryogenic sediments. This makes it possible to use radiocarbon dating of organic matter to establish the age of organisms. We performed accelerator mass spectrometry (AMS) radiocarbon analysis of plant material obtained from studied borrow P-1320, and determined a direct ¹⁴C age of 44,315±405 BP (Institute of Geography, RAS; sample IGAN_{AMS} 9137). Calibrated age range is 45,839 – 47,769 cal BP (95.4% probability) (Fig.S1).

Reanimated animals belong to a novel parthenogenetic species *Panagrolaimus* kolymaensis sp.nov.

The revived animal was cultivated in the laboratory for over 100 generations and initially described as *Panagrolaimus* aff. detritophagus⁵ based on morphology. We conducted a morphological analysis of the revived animal (Fig.3, S2, Table S1; See details of analysis in SI), which confirmed unambiguously that the animal belongs to the genus of *Panagrolaimus*, in agreement with a previous phylogenetic analysis of the 18S ribosomal RNA sequence⁵. However, due to the morphological uniformity of *Panagrolaimus*, unusual even for nematodes, morphology and molecular analysis of a single ribosomal RNA sequence is insufficient to describe a species. We found the species to be parthenogenetic, which further complicates description under many species concepts.

To obtain comprehensive molecular data for determining the species using phylogenomics, we generated a high-quality genome assembly. Using PacBio HiFi sequencing, we generated 84X coverage in long reads (average length 14,425 bp). Our analysis of repeat and gene content is described in Supplementary Table 3. K-mer analysis of the reads clearly indicated that this animal has a triploid genome (Fig.4a), similar to other parthenogenetic *Panagrolaimus* species²⁹. Despite the challenges that a triploid genome poses for assembly, we obtained a highly contiguous contig assembly of the three pseudohaplotypes that comprise a total of ~266 Mb and thus have a similar genome size as other parthenongenetic *Panagrolaimus* species²⁹. The contig N50 value of all three pseudohaplotypes is 3.8 Mb. Since the three pseudohaplotypes exhibited a noticeable degree of divergence, we further investigated their homology by using the apparent homeologs in our gene predictions to align long continuous

contigs based on micro synteny (Fig. 4b). Links between the contigs clearly shows the triploid pattern of the genome.

To place the species in the genus *Panagrolaimus*, we generated an initial phylogeny using the 18S and 28S genes using maximum likelihood before conducting a broader multigene phylogenomic analysis using a concatenated, partitioned alignment of 60 genes and a coalescence-based approach using a broader set of 12,295 gene trees (see supplementary methods, Supplementary Table 4 and 5). These analyses retrieve the novel species as sister to all other sequenced *Panagrolaimus* species, but as an ingroup to *Propanagrolaimus*³⁰ (Fig.4c; Fig.S3b-c). Thus, the phylogenetic placement provides strong evidence that this animal belongs to a novel species. Supporting this, there is substantial sequence divergence between this novel species, *Panagrolaimus* sp. PS1159 and *Panagrolaimus* sp. ES5, estimated in our concatenated alignment to be on average 2.06 and 2.11 amino acid substitutions per site, respectively. This divergence is consistent with previous data on ages of *Panagrolaimus* nematodes²⁹, and more broadly with the divergence of nematodes in general, which can be hyper-diverse^{31,32}. Our data also contradicts the assumption that parthenogenesis is monophyletic in the *Panagrolaimus* genus²⁹ (Fig.3c). Based on the Kolyma River location where the animal was unearthed, we propose the following taxonomic classification and species name:

Phylum Nematoda Potts, 1932

Class Chromadorea Inglis 1983

Suborder Tylenchina Thorne, 1949

Family Panagrolaimidae Thorne, 1937

Panagrolaimus kolymaensis sp. nov.

C. elegans dauer larvae and P. kolymaensis utilize similar mechanisms to enter and remain in cryptobiotic state for prolonged periods

The high-quality genome of *P. kolymaensis* allowed us to compare its molecular toolkit for cryptobiosis with that of *C. elegans*^{3,4,10,12}. We used orthology clustering and phylogenetics to investigate whether the genome of *P. kolymaensis* contains genes previously implicated in cryptobiosis in *C. elegans*. Our analysis showed that the *P. kolymaensis* genome encodes orthologs to a *C. elegans* trehalose phosphate synthase gene (tps-2) and to a trehalose phosphatase gene (gob-1) (Fig.S4c, supplementary file Orthology analysis). Furthermore, we found orthologs to all *C. elegans* enzymes required for polyamine biosynthesis, the TCA cycle, glycolysis, gluconeogenesis, and glyoxylate shunt (Fig.S4c, supplementary file Orthology analysis) suggesting that *P. kolymaensis* possesses a similar but not identical molecular tool kit as *C. elegans* to survive cryptobiosis.

To survive extreme desiccation, C. elegans dauer larvae need to be first preconditioned at high relative humidity (98% RH) for 4 days¹⁰. During preconditioning, dauer larvae upregulate trehalose biosynthesis that ensures their survival to harsh desiccation^{10,12}. We therefore tested whether preconditioning also facilitates survival of *P. kolymaensis*. Although a small proportion of P. kolymaensis individuals survive harsh desiccation and freezing without preconditioning (Fig.5a), similar to other panagrolaimid species¹¹, mixed populations of P. kolymaensis survive harsh desiccation significantly longer (p<0.0001) following preconditioning (Fig.5a). Similarly, preconditioning and desiccation further enhanced survival of P. kolymaensis following freezing (-80°C). Furthermore, like C. elegans, P. kolymaensis upregulate trehalose levels up to 20-fold upon preconditioning (Fig.5b). We previously reported that, to upregulate trehalose levels upon preconditioning, C. elegans dauer larva dissipate their fat reserves (triacylglycerols) by activating the glyoxylate shunt and gluconeogenic pathway²⁵. Upon preconditioning, we found that triacylglyceride (TAG) levels are significantly decreased in P. kolymaensis (Fig.S5a&b). To further investigate whether the acetyl-CoA derived from degradation of TAGs culminate in trehalose, we applied the previously developed method of metabolic labelling with ¹⁴C-acetate in combination with 2D-TLC^{4,25}. As shown in Fig.5c, preconditioning led to a huge increase of radioactivity in trehalose and also to a lesser extent in some amino acids (glycine/serine, phenylalanine; panels c and d). Interestingly, P. kolymaensis displayed an additional spot (Fig.5d, enumerated as 7), that was not found in C. elegans²⁵, which we identified as trehalose-6-phosphate (Fig.S5c-h), an immediate precursor of trehalose, using mass spectrometry. Thus, in order to resist harsh desiccation, similar to C. elegans, P. kolymaensis utilizes glyoxylate shunt and consequently acetate derived from TAGs to synthesize trehalose. Detection of the immediate precursor (trehalose-6-phosphate) suggests the flux of metabolites is intense in the latter.

Our findings indicate that by adapting to survival of the cryptobiotic state for short time frames in environments like permafrost, some nematode species gained the potential for individual worms to remain in the state for geological timeframes. This raises the question whether there is an upper limit to the length of time an individual can remain in the cryptobiotic state, potentially only limited by drastic changes to the environment such as strong fluctuations in ambient temperature, natural radioactivity or other abiotic factors. These findings have implications for our understanding of evolutionary processes, since generation times could be stretched from days to millennia, species ages might be much longer than anticipated, and longterm survival of individuals of species can lead to the refoundation of otherwise extinct lineages. Finally, understanding the precise mechanisms of long-term cryptobiosis and cues that lead to successful revivals can inform new methods for long term storage of cells and tissues and their transportation in space.

Methods

Materials and C. elegans strains

[1-14C] -acetate (sodium salt) from Hartmann Analytic (Braunschweig, Germany). All other chemicals were purchased from Sigma-Aldrich (Taufkirchen, Germany). The *Caenorhabditis* Genetic Centre (CGC) provided *daf-2(e1370)* and *E. coli NA22* strains.

Genomic DNA isolation from Panagrolaimus kolymaensis nematodes

P. kolymaensis nematodes were grown on several plates of NGM agar plated with E. Coli NA22 bacteria at 20°C. Worms were collected from the plates, washed with water at least three to five times by centrifugation at 1000 g to remove any residual bacteria and any debris. The worm pellet was dissolved in 5 volumes of worm lysis buffer (0.1M Tris-HCl pH=8.5, 0.1M NaCl, 50mM EDTA pH=8.0) and distributed in 1.5 ml of microcentrifuge tubes. These tubes are incubated at -80 C for 20 minutes. 100 ul of Proteinase 'K' (20 mg/ml) was added to each tube and they are incubated at 60 C overnight. 625 ul of cold GTC buffer (4M Guanidinium Thiocynate, 25mM Sodium citrate, 0.5% (v/v) N-lauroylsarcosine, 7%(v/v) Beta Mercaptoethanol) was added to the tube, incubated on ice 30 min and mixed by inverting every 10 min. 1 volume of phenol-chloroform-isoamyl alcohol (pH=8) was added to the lysate and mixed by inverting the tube 10-15 times. Tubes were centrifuged for 5 min at 10,000 g at 4 °C to separate the phases. The upper aqueous phase was carefully collected into a fresh tube. One volume of fresh chloroform was added and mixed by inverting the tubes for 10-15 times and centrifuged for 5 min at 10,000 g at 4°C to separate the phases. One volume of cold 5 M NaCl was added, mixed by inverting the tubes and incubated on ice for 15 min. After incubation these tubes were centrifuged for 15 min at 12,000–16,000 g at 4 °C. The supernatant containing the nucleic acids were slowly transferred into a fresh tube. One volume of isopropanol was added to the tube, inverted few times and incubated on ice for 30 minutes. After incubation, the tubes were centrifuged at 3000 g for 30-45 min at 25°C and the supernatant was discarded without disturbing the pellet. The pellet was washed twice with 1 ml of 70% ethanol, tubes were centrifuged at 3000 g for 5 min and supernatant was discarded and incubated at 37°C for 10-15 min to dry the pellet. The pellet was resuspended carefully in TE buffer. The quality and length of the high-molecular weight (HMW) genomic DNA (gDNA) was analyzed with pulse field gel electrophoresis.

Genome sequencing and assembly

Long insert library has been prepared as recommended by Pacific Bioscienes according to the protocol 'Procedure & Checklist-Preparing gDNA Libraries Using the SMRTbell® Express Template Preparation Kit 2.0'. In summary, RNAse treated HMW gDNA has been sheared to 20Kb fragments on the MegaRuptorTM device (Diagenode) and 10 µg sheared gDNA have been used for library preparation. The PacBio SMRTbellTM library was size selected in two fractions (9-13kb, > 13kb) using the BluePippinTM device with cassette definition of 0.75% DF MarkerS1 3-10 kb improved Recovery. The second fraction of the size selected library has been loaded with 95 pM on plate on one Sequel SMRT cell (8M). Sequel polymerase 2.0 has been used in combination with the v2 PacBio sequencing primer and the Sequel sequencing kit 2.0EA, runtime was 30 hours. We created PacBio CCS reads from the subreads.bam file using PacBio's ccs command linetool (version4.2.0). We obtained 8.5Gb high quality CCS reads (HiFi reads) with a N50 of 14.4 Kb. We ran HiCanu (version 2.2)³³ to create the contig assembly. Blobtools (version 1.1.1,CITE)³⁴ was used to identify and remove bacterial contigs. The final triploid contig assembly consists of 856 contigs has a N50 of 3.82 Mb and a size of 266Mb. The mitochondrial genome was created with the mitoHifi pipeline (version 2^{35}) based on the assembled contigs and the closely related reference mitochondrial genome of Panagrellus redivivus (strain: PS2298/MT8872, ENAaccession: AP017464). The mitoHifi pipeline identified 49 mitochondrial contigs ranging from 13-32Kb. The final annotated circular mitochondrial genome has length of 17467 bp.

To identify pseudohaplotypes in the *P. kolymanensis* sp. nov genome assembly, we selected the longest isoform of each predicted protein-coding gene in our assembly and in the *C. elegans* genome (downloaded from WormBase Parasite, release WBPS15) using AGAT (version 0.4.0) and clustered them into orthologous groups (OGs) using OrthoFinder (version 2.5.2). We identified OGs that contained three *Panagrolaimus* sequences (i.e. groups that were present as single-copy in all three pseudohaplotypes) and used these to identify trios of multi-megabase size contigs derived from the three pseudohaplotypes. We visualized synteny between the three pseudohaplotypes using Circos to plot the positions of each homeolog (version 0.69-8).

Genome annotation

We used RepeatModeler 1.0.8 (<u>http://www.repeatmasker.org/</u>) with parameter '-engine ncbi' to create a library of repeat families. This library was used with RepeatMasker 4.0.9 to softmask the *Panagrolaimus* genome. To annotate genes, we cross mapped protein models from

an existing *Panagrolaimus* as external evidence in the Augustus based pipeline³⁶. We evaluated the completeness of our predictions using BUSCO on the gVolante web interface.

Orthology analysis

We conducted a gene orthology analysis using genomic data from Panagrolaimus kolymaensis, the plectid nematode species from the permafrost, as well as genomic data from WormBase Parasite (https://parasite.wormbase.org; accessed 17/12/2020): Caenorhabditis elegans, Diploscapter coronatus, Diploscapter pachys, Halicephalobus mephisto, Panagrellus redivivus, Panagrolaimus davidi, Panagrolaimus sp. ES5, Panagrolaimus sp. PS1159, Panagrolaimus superbus, Plectus sambesii, and Propanagrolaimus sp. JU765. For plectids, genomic resources are scarce. We therefore added transcriptome data of *Plectus murrayi*, Anaplectus granulosus, Neocamacolaimus parasiticus, and Stephanolaimus elegans. The latter three transcriptomes were kindly provided by Dr. Oleksandr Holovachov (Swedish museum of natural history). The Anaplectus granulosus, and Neocamacolaimus parasiticus transcriptomes have been published ^{37,38}. All three transcriptomes were assembled de novo with Trinity³⁹. The exact procedures are described in the respective publications^{37,38}. The *Stephanolaimus elegans* transcriptome was assembled in the same way as described for *Neocamacolaimus parasiticus*. The Plectus murrayi transcriptome was built from raw reads deposited at NCBI (https://sradownloadb.be-md.ncbi.nlm.nih.gov/sos2/sra-pub-run-13/SRR6827978/SRR6827978.1; accessed 22.12.2020). The transcriptome was assembled using Galaxy Trinity version 2.9.1^{40,39} using all default options and including in silico normalization of reads before assembly. Transdecoder (conda version 5.5.0)⁴¹ was used to translate to amino acid sequence. Identical

version $4.8.1^{42,43}$. were removed with cd-hit and reads the Trinity get longest isoform seq per trinity gene.pl command⁴¹ (Trinity conda version 2.8.5) (Anaconda Software Distribution, Conda, Version 4.9.2, Anaconda, Nov. 2020) was used, to remove shorter isoforms. Amino acid translated longest isoforms from genomic data were extracted with AGAT (Dainat, https://www.doi.org/10.5281/zenodo.3552717) from genome assembly FASTA files and genome annotation GFF3 files, using the agat convert sp gxf2gxf.pl command to convert all GFF3 files to the required GFF3 format, and the agat sp keep longest isoform.pl as well as the agat sp extract sequences.pl commands to extract amino acid translated FASTA files. The headers and names of all FASTA files were modified to allow for simple species assignment of each sequence in subsequent analysis. The orthology analysis was conducted with OrthoFinder v. 2.5.^{144, 45} using default settings. For genes of interest, we constructed alignments with MAFFT v. 7.475⁴⁶ using the

localpair and maxiterate (1000) functions. We removed spurious sequences and areas that were not well aligned with Trimal v. 1.4.rev22⁴⁷ (procedure stated in supplementary file Orthology analysis below each phylogeny). We then ran phylogenetic analysis with Iqtree2 v. 2.0.6⁴⁸, with -bb 1000 option, testing the model for each analysis (models eventually used stated in supplementary file Orthology analysis). We also checked for PFAM domains using Interproscan v. 5.50-84.0⁴⁹. Part of the analysis was performed on the HPC RRZK CHEOPS of the Regional Computing Centre (RRZK) of the University of Cologne. The phylogenies were visualized with Dendroscope 3.7.6⁵⁰ and figures were created with Inkscape {https://inkscape.org}.

Phylogenomics

Sequences of 18S and 28S genes from 44 taxa across the *Propanagrolaimus*, *Panagrolaimus*, *Panagrellus* and *Halicephalobus* genera were aligned (MAFFT L-INS-I v7.475)⁴⁶, concatenated⁵¹ and used to infer a species tree using maximum likelihood via (IQTREE)⁵² and partitioned by best-fit models of sequence evolution for both⁵³. Nodal support was determined using 1000 bootstrap pseudoreplicates. A further 60 genes from 101 taxa were used to confirm the taxonomic position using the supermatrix concatenation methods outlined above. Given the limitations of differential gene sampling, we expanded our phylogenomic analyses to include a coalescence approach using 12,295 ML gene trees inferred for orthogroups containing the target animal. Instances of multiple genes per species per group were treated as paralogs/orthologs and analysed using ASTRAL-Pro⁵⁴.

Desiccation survival assay

C. elegans dauer larvae desiccation assays were performed as described in¹⁰ *Panagrolaimus kolymaensis* desiccation assays were performed with mixed populations of the worms.

Exposure of nematodes to extreme environments

Dauer larvae or *Panagrolaimus kolymaensis* nematodes were preconditioned and desiccated as described²⁴ and then transferred to elevated temperature of 34 C, freezing (-80 C) and anoxia. Anoxic environment was generated in a desiccation chamber at 60%RH by flushing the Nitrogen gas into the chamber. The concentration of oxygen inside the chamber was monitored. After each timepoint they were rehydrated with 500 μ l of water for 2-3 hours. Rehydrated worms were transferred to NGM agar plates with *E. Coli NA22* as food. Survivors

were counted after overnight recovery. Each experiment was performed on two different days with at least two technical replicates.

Trehalose quantification from nematode lysates

Trehalose measurements were performed as described in¹².

Radiolabeling, metabolite extraction and 2D-TLC

The above-mentioned procedures were performed according to previous reports^{4,12}.

Identification of trehalose-6-phosphate from TLC plates

Normalised aqueous fractions from the non-preconditioned and preconditioned samples were separated by high performance thin layer chromatography (HPTLC), using 1-propanolmethanol-ammonia (32%)-water (28:8:7:7 v/v/v/v) as first, dried for 15 min and 1-butanolacetone-glacial acetic acid–water (35:35:7:23 v/v/v/v) second dimension respectively. Using the trehalose as a standard on both dimensions of the TLC, the regions of interest were scrapped out from the TLCs. The scraped-out silica was extracted with 10 ml of 50% methanol twice. The fractions were combined, dried under vacuum and dissolved in 100 ul of Ms mix solution containing 4:2:1 (Isopropanol:Methanol:Chloroform) with 7.5 mM ammonium formate. Mass spectrometric analysis was performed on a Q Exactive instrument (Thermo Fischer Scientific, Bremen, DE) equipped with a robotic nanoflow ion source TriVersa NanoMate (Advion BioSciences, Ithaca, USA) using nanoelectrospray chips with a diameter of 4.1 µm. The ion source was controlled by the Chipsoft 8.3.1 software (Advion BioSciences). Ionization voltage was + 0.96 kV in negative mode; backpressure was set at 1.25 psi. The temperature of the ion transfer capillary was 200°C; S-lens RF level was set to 50%. FT MS spectra were acquired within the range of m/z 50–750 at the mass resolution of R m/z 200 = 140000; automated gain control (AGC) of 3×10⁶ and with the maximal injection time of 3000 ms. FT MS/MS spectra were acquired within the range of $m/z \ 50-750$ at the mass resolution of R $m/z \ 200 = 140000$; automated gain control (AGC) of 3×10^4 and with the maximal injection time of 30 s.

Triacylglycerols measurement from Panagrolaimus kolymaensis lysates

Non-preconditioned and preconditioned pellets were lysed in 200 ul of isopropanol with 0.5 mm Zircornium beads twice for 15 min. The lysates were centrifuged at 1300 g for 5 min and 20 ul of the lysate was used for protein estimation. The supernatant was carefully collected

without any debris and was dried in the desiccator. To dried samples 700 ul of 10:3 (v/v) MTBE/MeOH solution with internal standard (TG-d5 50:0) were added and left on a shaker for 1 hour at 1400 rpm and 4°C. 140 ul of water was added and left on shaker for 15 min at 4°C. These samples were centrifuged at 13400 rpm for 15 min at 4°C. The upper organic fraction was collected and transferred to 1.5 ml glass vial and left for drying in the desiccator. The dried samples were reconstituted in appropriate volume of 300 ul of 4:2:1 (Isopropanol: Methanol: Chloroform) and volume corresponding to 1 μ g protein was used for injection.

LC-MS/MS analysis was performed on a high-performance liquid chromatography system (Agilent 1200 HPLC) coupled to a Xevo G2-S QTof (Waters). The samples were resolved on a reverse phase C18 column (Cortecs C18 2.7um from Waters) with 50:50:0.1:1% (Water:methanol:formic acid: 1M Ammonium formate) and 25:85:0.1:1% (Acetonitrile:Isopropanol:Formic acid:1M Ammonium formate) as mobile phase. The following gradient program was used: Eluent B from 0 % to 100 % within 12 min; 100 % from 12 min to 17min; 0 % from 17 min to 25 min. The flow rate was set at 0.3 ml/min. The samples were normalised according to the total protein concentration and the worm numbers. The TG species were extracted from Lipidmaps database and Skyline software was used to analyse the TG profile.

Author contributions

AS, VG, PS, TK conceived and designed the study. AS, VG, TH, MP, AT, GH, MH, ER, PS and TK contributed to the original draft. AS performed isolation and cultivation of nematodes. VG performed desiccation survival assays, trehalose measurement, 2D-TLC of metabolites, trehalose-6-phosphate detection, combination of cryptobiosis experiments, genomic DNA isolation from *Panagrolaimus kolymaensis* and prepared the figures. MP performed genome assembly. TH conducted orthology analyses and single gene phylogenies, and worked on figures. AT provided morphological description, light and scanning electron microscopy. LS analyzed the genome assembly and created figures. GMH performed the phylogenetic analyses. ST performed triacylglycerols measurements, trehalose-6-phosphate detection with suggestions from Andrej Shevchenko. MH annotated supplementary table 3. PS performed genome annotation and supervised TH. EWM supervised MP and had overall responsibility for sequencing and assembly including funding it.

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Figure 1. C. elegans dauer larvae can remain in a cryptobiotic state for prolonged periods

of time. a) Desiccated *daf-2 (e1370)* dauer larvae survive to freezing (-80°C) for an extremely long period. Error bars indicate standard error of mean of two independent experiments with two technical replicates performed on two different days. b) Brood size of desiccated dauer larvae exposed to freezing remain similar to that non- desiccated dauer larvae. Average brood size is the mean of seven dauer larvae per each condition. Statistical comparison was performed by using non-parametric Kolmogorov-smirnov test n.s p>0.05.

Figure 2. Study site.

a) Location of the Duvanny Yar outcrope on the Kolyma River, northeastern Siberia, Russia.
b) view of the upper part of outcrop composed of ice wedges and permafrost silty deposits. c)
lithostratigraphic scheme of deposits, showing location of studied rodent borrow (red circle).
d) Fossil rodent burrow with herbaceous litter and seeds buried in permafrost deposits. m a.r.l.
= meters above river level.

Figure 3. General morphology of Panagrolaimus kolymaensis sp. n., female.

Scanning electron pictures (a, c), light microscopy photographs (e, f) and graphic presentations (b, d, g) of holotype: a, b) entire body, c, d) anterior ends, e) anterior body, f) perivulvar body region, g) tail. Abbreviations: l.f. – lateral field, ov – ovary, pro – procorpus of the pharynx, t.b. – terminal bulb of the pharynx, u – uterus with eggs, v – vulva, v.p. – ventral pore. Scale bars: a, d, e, f, g – 20 μ m, b – 100 μ m, c – 2 μ m.

Figure 4. Genome assembly and phylogenomics reveals that the newly discovered *Panagrolaimus kolymaensis* species is triploid.

a) Kmer spectra of the *Panagrolaimus kolymaensis* PacBio HiFi data. Kmers of length 19 were counted using Jellyfish. b) Circos plot showing the triploid structure of the *Panagrolaimus kolymensis* genome. The relationship between eight contigs, representing 39.9 Mb (15%) of the genome, is shown. Contig IDs are shown. Scale is in megabases (Mb). c) Inferred species tree for all taxa. The maximum likelihood tree inferred using a concatenated supermatrix (18S and 28S genes) with bootstrap support values is displayed. All genera are represented as monophyletic clades. *Panagrolaimus kolymaensis* is highlighted in red and basal to all other *Panagrolaimus* taxa. Internal nodes, where all subsequent branches represent identical sequences, are displayed with a black star.

Figure 5. *C. elegans* dauer larvae and *P. kolymaensis* utilize similar mechanisms to survive cryptobiosis.

a) Survival rate of *P. kolymaensis* nematodes to desiccation and freezing (-80°C). Error bars indicate standard error of mean of two independent experiments with two technical replicates performed on two different days. Statistical comparison was performed using unpaired t test with Welch correction. n.s. p > 0.05, ****p < 0.0001. For desiccation (non-preconditioned) n = 289, freezing (non-preconditioned) n = 675, desiccation (preconditioned at 98%RH) n = 953 and freezing (preconditioned at 98%RH) n = 1295. b) *P. kolymaensis* nematodes and *daf-2(e1370)* dauer larvae upregulate trehalose levels upon preconditioning at 98%RH. Error bars indicate standard error of mean of two independent experiments with three replicates performed on two different days. Statistical comparison was performed using two-way ANOVA with Holm-Sidak's multiple comparison test, ****p < 0.0001. c-d) 2D-thin layer chromatography of ¹⁴C-acetate labelled metabolites from *P. kolymaensis* that were non-preconditioned at 98%RH. Enumerated spots indicate trehalose (1), glucose (2), glutamate (3), glutamine (4), serine/glycine (5) and phenylalanine (6). Representative images from at least two independent experiments performed on two different days.

Figure S1. Calibration of a radiocarbon (¹⁴C) date

Radiocarbon date (44,315±405 BP) and calibrated age (45,839 - 47,769 cal BP) of plant material collected from buried borrow P-1320. Radiocarbon ages were converted to calendar age equivalents with the OxCal V.4.4 program using the IntCal20 calibration curve⁵⁵. Pink-shaded area — radiocarbon date with standard deviation; grey-shaded area — radiocarbon date projection on the calibration curve with 95.4% probability.

Figure S2. Morphology of *Panagrolaimus kolymaensis* sp. n., female.

Graphic presentations of holotype (a,b) and SEM pictures (c-i): a) anterior body, b) female reproductive branch, c–e) anterior end of three different female specimens, f) anterior part of the lateral ridge, g) vulva, h) ventral excretory/secretory pore, i) posterior body with anus and lateral ridge. Scale bars: a,b - 50 μ m, c-e - 3 μ m, f, g, i - 10 μ m, h - 5 μ m.

Figure S3. Phylogenies inferred for genes sets using both concatenation and coalescence approaches.

a) The ploidy level was analysed with *Smudgeplot* v0.2.1⁵⁶. KMC version 3.1.0⁵⁷ was used to count the 21-mers in the PacBio CCS reads. Then we ran *smudgeplot.py* to determine the lower and upper coverage cut-offs. These were determined to be 14 and 380. 21-mers with a coverage between 14 and 380 were filtered with kmc_tools. Then, we computed the k-mer pairs from the filtered 21-mers by running *smudgeplot.py hetkmers*. Finally, the produced a smudgeplot shows an estimated ploidy of 3. b) Species tree inferred for 12,295 gene trees using a coalescence approach. The species tree implemented using the coalescence approach with orthogroup gene trees is displayed. Novel species in this study are displayed in red. All nodes have a posterior probability of 1. c) Species tree inferred for 102 taxa. The maximum likelihood tree inferred using a concatenated supermatrix of 60 genes is displayed. Bootstrap values are only displayed for nodes with less than 100% support. The platyhelminth *Macrostum lignano* serves as an outgroup for rooting. The Ancestral panagrolaimus, basal to all others within the *Panagrolaimus* genera, is highlighted in red.

Figure S4. Combination of cryptobiotic states enhances survival of *C. elegans* dauer larvae.

a) Desiccated dauer larvae manifest enhanced survival rate to heat stress (34°C). Error bars indicate standard deviation of two independent experiments with two technical replicates. b) Desiccated dauer larvae display enhances survival rate to anoxia. Error bars indicate standard deviation of two independent experiments with two technical replicates. Statistical comparison was performed by paired two tailed t-test. *p<0.05. c) *P. kolymaensis* possesses *C. elegans* gene orthologs to enzymes required for TCA cycle, glyoxylate shunt, glycolysis, gluconeogenesis, trehalose synthesis, and polyamine synthesis. Black filled circles: Ortholog presence suggested by orthogroup clustering, phylogenetic analysis, and domain architecture. White filled circles: No ortholog found via current analysis. Coloured filled circles: Presence of *P. kolymaensis* gene(s), related to a number of *C. elegans* genes (all genes of same colour) that are all co-orthologous to that gene (those genes). Label: *C. elegans* enzyme names and orthogroup that contains that gene according to our orthogroup clustering

Figure S5. *P. kolymaensis* reduces triacylglycerols (TAGs) levels and accumulates trehalose-6-phosphate upon preconditioning at 98%RH.

a) 1D-Thin layer chromatography of acetate labelled organic fractions of non-preconditioned (1) and preconditioned (2) *P. kolymaensis*. b) Mass spectrometric quantification of TAG levels of non-preconditioned (1) and preconditioned (2) *P. kolymaensis*. Error bars indicate standard deviation of two independent experiments with two technical replicates. Statistical analysis was performed using unpaired t-test with Welch correction **p<0.001. c-d) non-preconditioned and preconditioned mass spectrum of an empty region, e-f) spot 1 (trehalose), g-h) spot 7 (trehalose-6-phosphate) scraped out and extracted from the 2D-TLC.

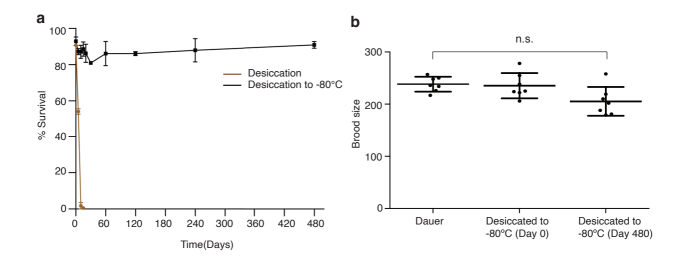


Figure 1 Shatilovich *et al*

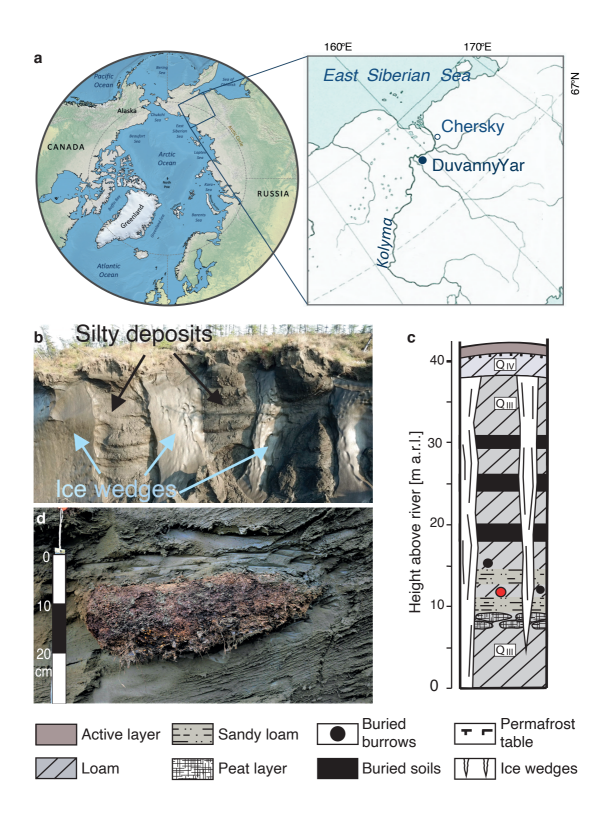


Figure 2 Shatilovich *et al*

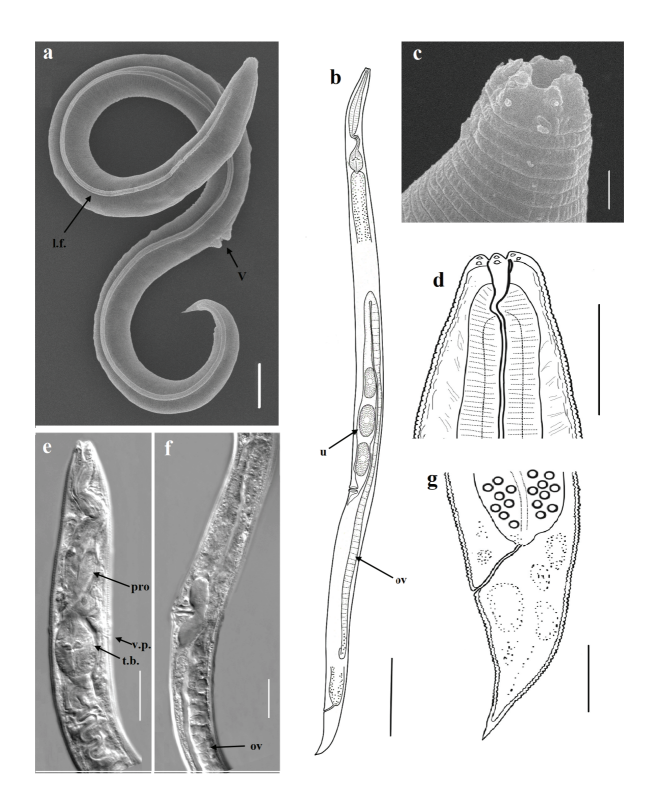


Figure 3 Shatilovich *et al*

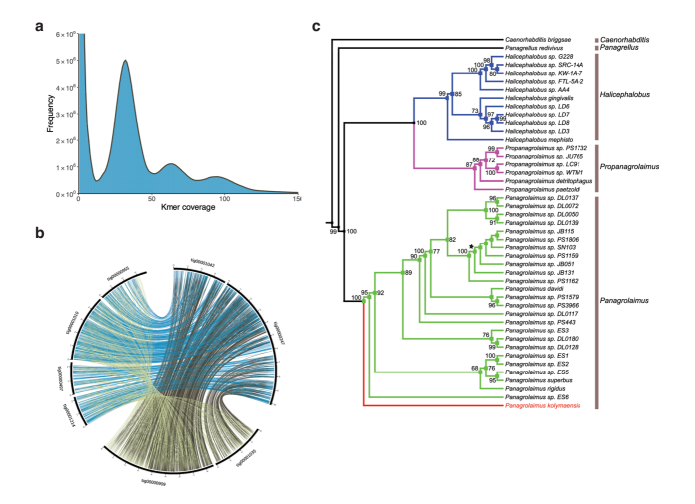
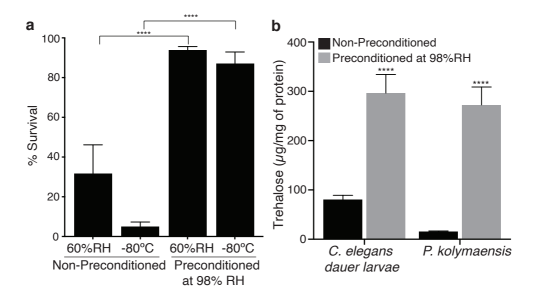


Figure 4 Shatilovich *et al*



Non-preconditioned

Preconditioned at 98%RH

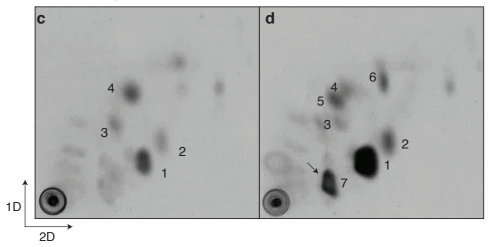


Figure 5 Shatilovich *et al*

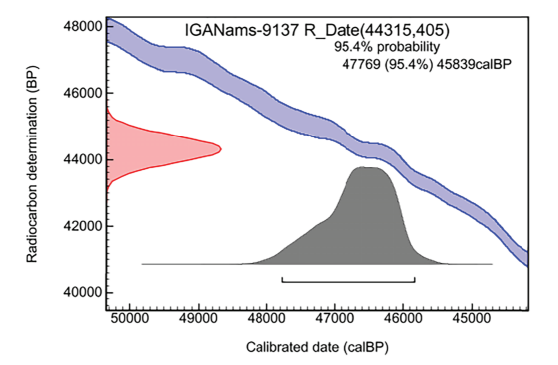


Figure S1 Shatilovich *et al*

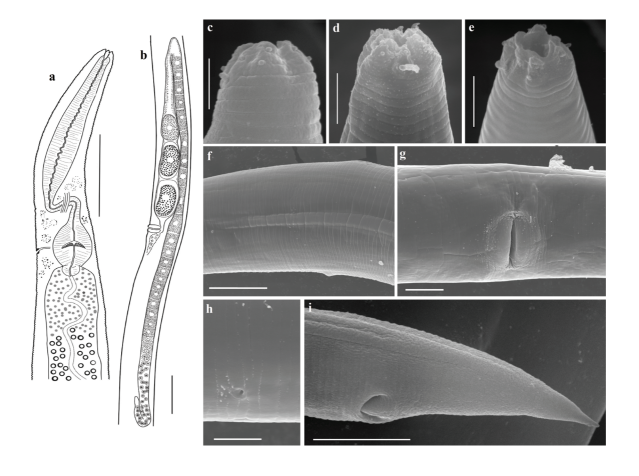


Figure S2 Shatilovich *et al*

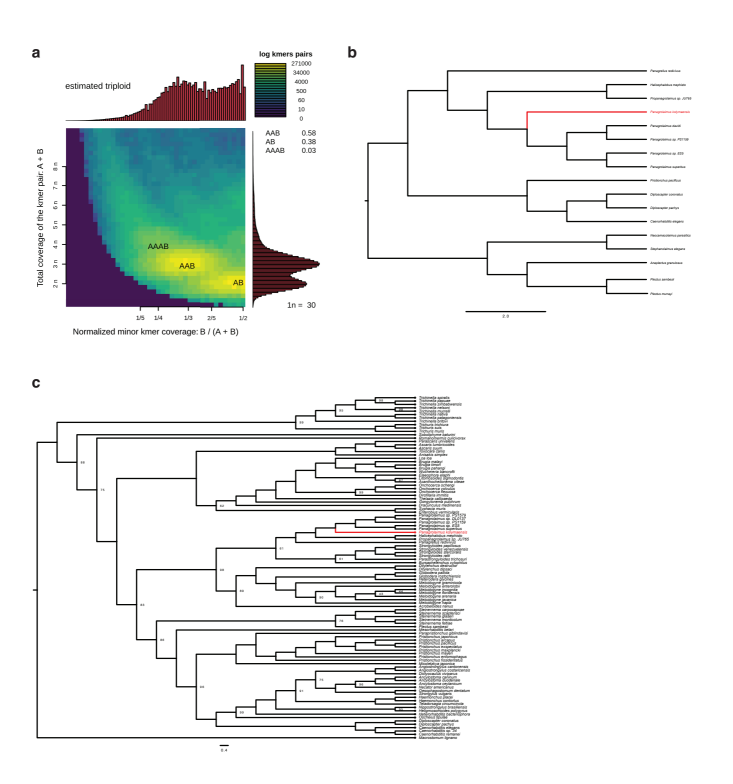


Figure S3 Shatilovich *et al*

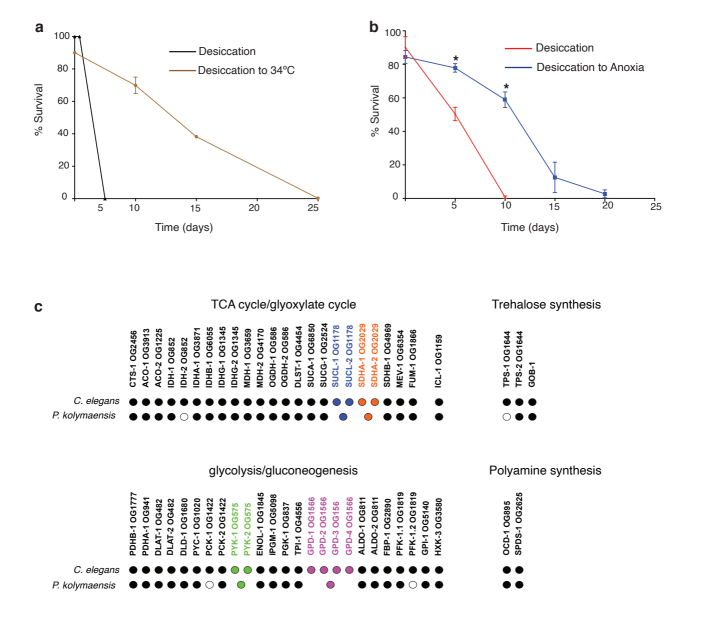


Figure S4 Shatilovich *et al*

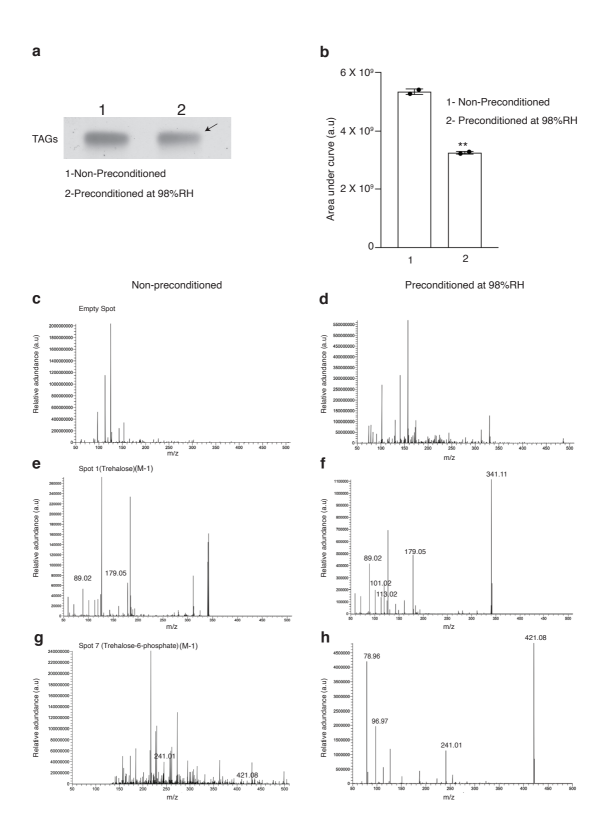


Figure S5 Shatilovich *et al*