Dormancy in Embryos: Insight from Hydrated Encysted Embryos of an Aquatic Invertebrate*

Tamar Ziv‡§§, Vered Chalifa-Caspi§§§, Nadav Denekamp¶, Inbar Plaschkes§, Sylwia Kierszniewska‖, Idit Blais**, Arie Admon‡, and Esther Lubzens‡‡‡

Numerous aquatic invertebrates remain dormant for decades in a hydrated state as encysted embryos. In search for functional pathways associated with this form of dormancy, we used label-free quantitative proteomics to compare the proteomes of hydrated encysted dormant embryos (resting eggs; RE) with nondormant embryos (amicotic eggs; AM) of the rotifer *Brachionus plicatilis*.

A total of 2631 proteins were identified in rotifer eggs. About 62% proteins showed higher abundance in AM relative to RE (Fold Change>3; p = 0.05). Proteins belonging to numerous putative functional pathways showed dramatic changes during dormancy. Most striking were changes in the mitochondria indicating an impeded metabolism. A comparison between the abundance of proteins and their corresponding transcript levels, revealed higher concordance for RE than for AM. Surprisingly, numerous highly abundant dormancy related proteins show corresponding high mRNA levels in metabolically inactive RE. As these mRNAs and proteins degrade at the time of exit from dormancy they may serve as a source of nucleotides and amino acids during the exit from dormancy. Because proteome analyses point to a similarity in functional pathways of hydrated RE and desiccated life forms, REs were dried. Similar hatching and reproductive rates were found for wet and dried REs, suggesting analogous pathways for long-term survival in wet or dry forms. Analysis by KEGG pathways revealed a few general strategies for dormancy, proposing an explanation for the low transcriptional similarity among dormancies across species, despite the resemblance in physiological phenotypes. *Molecular & Cellular Proteomics* 16: 10.1074/mcp.RA117.000109, 1746–1769, 2017.

© 2017 by The American Society for Biochemistry and Molecular Biology, Inc. This paper is available on line at http://www.mcponline.org
nondesiccated state in aquatic organisms such as rotifers, cladocerans, or copepods, in the form of resting eggs or ephippia (19–24). Most amazing is the survival of *Daphnia* ephippia in a lake sediment cores for over six centuries (25). Dormancy in these species is developmentally programmed at embryonic stages. Developmental arrest at embryonic developmental stages is also well known in insects (1), fish (26), and mammals (27–30). One of the most famous examples of diapause occurs in developing embryos of the silkworm, *Bombyx mori* (31), where a diapause hormone is excreted by the subesophageal ganglion of the mother and acts on the developing eggs, which later exhibit diapause during embryonic development. Recently, a diapause hormone receptor-like gene was identified in Artemia with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.

The resting eggs (RE) or encysted embryos formed by the rotifer *B. plicatilis* are an excellent model for investigating long-term survival in the hydrated state. Rotifers are minute rotifer *B. plicatilis* rotifers, a nonmodel aquatic invertebrate species. Embryos enter dormancy or diapause with a function in inducing and maintenance of diapause in embryos (32). The brine shrimp, *Artemia franciscana* embryos enter diapause or dormancy in the gastrula stage of embryogenesis and is one of the most studied species as it endures desiccation (5, 9). Significant advances were made in revealing functional changes during dormancy in model and nonmodel organisms (1, 6). Nevertheless, diverse experimental organisms offer additional unique advantages for investigating dormancy by contributing important insight into the wide scope of the dormancy phenomenon. Here, we aim at expanding our knowledge on dormancy in embryos, by studying resting eggs of rotifers, a nonmodel aquatic invertebrate species.
in insects, including regulation of metabolism and the cell cycle (6, 26).

Gene expression profiles linked with diapause have been studied in numerous insects (e.g. 15, 66–72 and others), but almost all of them were not performed on embryos, except for Tu et al. 2015 (73). A few studies were performed on Brachionid rotifers forming resting eggs (42, 51–53, 74). Gene expression profiles, however, are insufficient for explaining gene function, as mRNAs do not fully correlate with protein levels (75, 76). This lack of correlation was also demonstrated for prediapausing and nondiapausing locust eggs (73). Only a limited number of dormancy related proteins were revealed by proteome profiling studies of dormant or diapausing invertebrate embryos. Proteome profiling of Artemia encysted and postdiapause embryos revealed 75 proteins with differential abundance in Artemia franciscana cysts (77) and 33 unique proteins in cysts of Artemia sinica (78). Shotgun proteome analyses identified 192 diapause unique proteins in the silkworm (79) and 212 differential abundant proteins between prediapause and nondiapause locust eggs (73).

To reveal the wide scope of functional changes taking place during dormancy of encysted embryos we used a shotgun, label free proteomics approach and identified 2631 protein groups and grouped a large number of them into putative functional pathways. Because proteome profiling showed that numerous pathways in hydrated encysted embryos of rotifers are similar with those of Artemia cysts and life forms showing resistance to desiccation (6, 9), we dried REs. The viability of desiccated rotifer RE was tested by comparing the hatching percent and the reproductive rates of neonates hatching from wet and dried RE. In this study we also aim at addressing two additional topics: (1) How similar are the functional pathways in encysted dormant rotifer eggs that survive in a hydrated form for long periods of time, with those described so far for other forms of dormancy or diapause in embryos? (2) Whether proteome profiling can assist in explaining the small transcriptional similarity among dormancies across species, despite the resemblance in physiological phenotypes.

**EXPERIMENTAL PROCEDURES**

**Experimental Design and Statistical Rationale**—This study consists (Fig. 1) of: (a) proteome analyses of three rotifer egg types; amictic eggs (AM), resting eggs (RE), and resting eggs ready for hatching (E0) to discover the proteins with functions in dormancy. Triplicate samples were examined for each egg type. LC-MS/MS data was analyzed by comparing the amino acid sequences with an in silico translated newly assembled reference transcriptome. Principle Component Analysis (PCA) was used to reveal the similarities between the nine analyzed samples based on their global proteomic profiles. To discover proteins showing differential abundance in AM relative to RE or E0, 1-way ANOVA test was carried out with the contrasts of AM versus RE and AM versus E0., and in addition, proteins which were present above a certain background level in one egg type and absent in the other were identified. Gene ontology (GO) and KEGG pathway enrichment analyses were performed to identify overrepresented biological themes within the differentially abundant proteins. To ascertain the relationship between the proteome and transcriptome, the abundance of proteins was compared with gene expression levels; (b) Metabolome profiling was used to support the conclusions on differences in the metabolism of AM and RE, using t test on results of metabolites, from triplicate samples of each egg type; and (c) Analysis of functional pathways suggested a resistance of RE to desiccation. Viability of dried RE was determined by comparing (five replicates for each time point) the hatching percent of dried RE after 1–56 days of dry storage, by Tukey Post-Hoc tests, after a Chi-square test, showing a normal distribution of the percent of hatching. In addition,
reproductive output by neonates hatching from 56-day dry stored RE (n = 110) was compared with wet stored RE (n = 113), by 1-way ANOVA, after a Chi-square test, showing a normal distribution of the results. More detailed information is provided in the following sections.

**Rotifer Proteome Samples**—Rotifers were cultured as described previously (51, 53). Resting eggs (REs) were manually collected from the bottom of culture vessels of 14–25-day-old rotifer cultures, cleaned of the debris and placed in a microfuge vial and flash frozen by liquid nitrogen and stored at −80 °C. Three biological replicate samples, each consisting of 5,000 RE, were collected from three different cultures. In addition, three replicate samples of RE were collected from the same cultures, except that they were placed in vials with seawater (10 ppt) and incubated at 25 °C, in a temperature controlled room (treatment E0). After 3 months of incubation, these eggs (named E0) were flash frozen in liquid nitrogen and stored at −80 °C. For collection of amictic eggs (AM), females from the same cultures were concentrated on a 60 μm mesh plankton sieve. They were re-suspended in 10–15 ml of sterile sea water (10 ppt) and the AM eggs were removed from the females by intensive up and down aspiration with a 5-ml pipette tip. Following this treatment, AM eggs were transferred from the bottom of the vial into Petri dishes. The AM eggs were manually collected and three replicate samples from the three different cultures were placed in a microfuge vial (−5000 AM eggs per vial) and flash frozen by liquid nitrogen and stored at −80 °C.

**Metabolome Samples**—Rotifers were cultured in artificial sea water (Red Sea, Israel; 10.8 g m L−1) and fed with *Nannochloropsis* sp. (Galil Algae, Israel). RE were collected after 8–10 days of culture from three different cultures. Amictic eggs (AM) were collected from three cultures after 3–4 culture days. AM and RE were collected into microfuge vials, washed twice with Dulbecco’s phosphate buffered saline (Sigma, St. Louis, MO), frozen in LN2 and stored at −80 °C.

**Protein Extraction, Proteolysis, and Mass Spectrometry**—Proteins were extracted from samples that were also used for RNA extraction (51) with the Guanidinium thiocyanate–phenol-chloroform extract (TRI Reagent; Sigma, St. Louis, MO) following the manufacturer instructions. It was assumed that similar efficiencies in protein extraction (TRI Reagent; Sigma-Aldrich, UK), frozen in LN2 and stored at −80 °C, were obtained with the TRI Reagent for the three egg types of RE, AM, and E0 and the list of identified proteins also included minimum PEP score of 1 and at least two unique peptides for identification. The term “protein” was used in this manuscript for the identified peptides, instead of the more appropriate term of “protein groups.” The data was quantified by label free analysis using the MaxQuant software, based on extracted ion currents (XICs) of peptides enabling quantitation from each LC/MS run for each peptide identified in any of experiments (supplemental Table S1). The mass spectrometry proteomics data were deposited at the ProteomeXchange Consortium via the PRIDE partner repository with the data set identifier PXD003395.

**Transcriptome Assembly**—A reference *Brachionus plicatilis* transcriptome was constructed from several sequence collections: (1) 52,143 Expressed Sequence Tag (EST) sequences from dbEST (NCBI), after trimming of adapter sequences, low complexity sequences and terminal poly-A’s, (2) 84 mRNA sequences from RefSeq, and (3) 18,751 contigs from previous RNA-Seq analysis (51). All sequences were assembled using CAP3 with the following parameters: -c 12 -o 30 -p 75 -s 500 -y 250 -z 2. The resulting transcriptome, containing the contigs and “singlets” produced by CAP3 comprised a total of 16,680 putative transcripts. The assembled transcriptome is available at http://bioinfo.bgu.ac.il/genomes/Esther_Lubzens/public/ (Username: ester; Password: retse).

**Transcriptome Conceptual Translation and Annotation**—Transcripts longer than 60 bp were compared with the RefSeq protein database using BLASTX and were also submitted to OrfPredicator to identify the frame with the longest open reading frame. Transcripts were subsequently translated through their entire length at the frame detected by BlastX, if they had a hit with e-value < 10−5, or otherwise, they were translated at the frame detected by OrfPredicator. The resulting amino acid sequences were annotated using Blast2GO through BLASTX versus RefSeq (considering up to top 20 hits, e-value cutoff 10−5), and were assigned KEGG Orthology (KO) IDs using the KEGG KAAS server. Transcript and protein-level annotations are provided in supplemental Table S2.

**Proteins in Rotifer Eggs**—A total of 2928 proteins were identified for the three egg types of RE, AM, and E0 and the list of identified proteins is shown in supplemental Table S3A. Peptide counts were taken from the “razor + unique peptides” column in MaxQuant file. Protein intensity values from MaxQuant were Log2 transformed. A protein with zero intensity in a certain sample was regarded as “not detected” (ND) in that sample and indicated by a blank cell in supplemental Table S3A and S3B. Intensity distribution plots and Principal Component Analysis (PCA), carried out in Partek® Genomics Suite®, showed that sample E0_3 was exceptional (supplemental Fig.)
Therefore, this sample was excluded from further analyses. As shown in supplemental Fig. S1A, the intensity values in all three replicates of AM were generally higher than in RE and E0 (the distribution plot is shifted to the right). Because we could not know whether this is caused by a biological effect or by a technical bias, we thought it would be more appropriate to stay on the safe side and avoid normalization. Normalization may introduce its own bias, distort the data, and hide real changes in protein abundance among the samples. The use of unnormalized data, a stringent fold of change (FC) cutoff of 3 was used for differential abundance. As shown in supplemental Table S3B, the obtained results were like those achieved with LFQ-normalized data which underwent the same statistical analysis described below except that a more traditional FC cutoff of 1.5 was used.

For a given protein in a given egg type, the following terms were defined: (a) the protein was considered as “present” in that egg type if it was detected in at least 2 replicates with 2 or more peptides in each replicate; (b) “present above background level” if it was “present” and its average intensity value across replicates was higher than 24 (which is the 10% percentile value of all intensity values in the data set), (c) “absent” if it was not detected (ND) in all of the replicates and (d) “uncertain” if it had an intensity value in only one replicate. A total of 2342, 1751, and 1482 proteins were found to be present in AM, RE and E0 egg types, respectively (supplemental Table S3B). Altogether, 2631 proteins were found to be present in at least one of the egg types, and these proteins were submitted to subsequent analyses (supplemental Table S3B).

Identification of Proteins Showing Differential Abundance Between Dormant and Nondormant Eggs—As indicated above, prior to statistical testing, all zero intensity values were replaced by missing values. This was applied, because when one of the egg types had zero intensities and the other did not, the zeros created artificially inflated fold of change values and extremely low p values in the statistical tests. Such a phenomenon is especially severe in proteomics data, where the lowest nonzero intensities are usually around 2 to the power of 18–20. Therefore, in the current study, the statistical test neglected the zeros, and separate criteria were applied to identify proteins with apparently no expression in one egg type and apparent expression in another egg type.

A 1-Way ANOVA Test was Carried Out in Partek with the Contrasts—AM versus RE, AM versus E0, RE versus E0. Significance p values were adjusted for multiple testing using the False Discovery Rate (FDR) method. Fold of change (FC) in each contrast was expressed in linear scale, where positive and negative values indicate higher or lower abundance, respectively. A protein was regarded as showing differentially abundance between two egg types, if it had a fold change larger than 3 and FDR-adjusted p value <0.05. This resulted in 1088 proteins in the comparison between AM and RE and 1042 proteins in the comparison of AM and E0. In addition, proteins fulfilling any of the following criteria were added to the list of differentially abundant proteins: (a) It was absent in one group and present above background level (see above) in the other; (b) It was “uncertain” (see above) in one group and present above background level in the other group and the fold change was larger than 10. As shown in supplemental Table S3B, the final list of “differentially abundant proteins” included 1538 proteins for the AM versus RE contrast, and 1570 proteins for the AM versus E0 contrast. Uniprot (http://www.uniprot.org/) was used for retrieving functional information on proteins.

Analysis of RNA-Seq Data and Comparison Between Proteins and Corresponding Transcripts—RNA-Seq data from Clark et al., 2012 (51) was used for AM and RE (one replicate sample per egg type, based on the extraction of ~5000 eggs for each egg type) was reanalyzed using the current reference transcriptome. Single-end 32bp reads of AM and RE samples were aligned to the assembled transcriptome using STAR 2.3 and the number of reads per transcript were counted using BEDTools.

Normalization of the raw counts to account for library size was performed using the DESeq R package. Normalized count values were log2 transformed, after values < 1 were floored to 1. Transcripts with normalized count value < 6 in both samples were filtered out.

Protein intensity values and their corresponding RNA-Seq normalized counts (both in log2 scale) were loaded into R. For each egg type (AM, RE) a simple linear regression and a residual analysis were carried out to compare RNA and protein expression. Genes having studentized residual >2 or <-2 were considered discordant. They comprised 4.2% of the proteins. Genes having studentized residual between ~0.7 and 0.7, comprising 50% of the proteins, were considered concordant.

Gene Ontology (GO) Annotation and Enrichment Analysis—Gene Ontology (GO) annotation and enrichment analyses were carried out using Blast2GO. Enrichment was done using Fischer exact test, and the entire transcriptome was used as the background set. The list of enriched GO terms was further reduced using Blast2GO pro’s “Reduce to most specific terms” algorithm. This algorithm filters the results by removing parent terms of already existing, statistically significant, child GO terms. GO terms having enrichment FDR p value < 0.05 were reported. Further processing of the enrichment results was done using a Perl script (supplemental Tables S4, S6, and S9).

KEGG Pathway Enrichment Analysis—Enrichment analyses of KEGG pathways in selected protein sets, was carried out using Expaner (82). The sets included (supplemental Table S5A). Three sets for proteins in AM, RE or E0, (b) Two sets of differentially abundant proteins in the comparison of AM versus RE and AM versus E0, and (c) Two sets of proteins in concordance with RNA for AM and RE. All KO IDs identified in the reference transcriptome served as the background set. Drawing and coloring of KEGG pathways was done using “KEGG Mapper - Search & Color Pathway” tool.

Hierarchical Clustering—Hierarchical clustering was carried out in Partek. Prior to clustering, intensity values were z-scored so that for each protein the mean and standard deviation were 0 and 1, respectively. In the heat maps, missing values were marked in white.

Sequence Analyses of Selected Proteins—LEA proteins from selected organisms were searched against the reference transcriptome using blast and hmmsearch (supplementary File Text S1).

Lists of Proteins of Specific Interest—Proteins of specific interest were prepared by manual curation and are shown in Table I and supplemental Table S3B.

Metabolome Profiling Samples—All steps were carried out in metatSysX GmbH, Potsdam Germany. Frozen samples (from ~80 °C) of AM and RE were extracted according to a modified protocol from Giavalisco et al., 2009 (83). After extractions volumes collected for further analysis were adjusted to same number of eggs (~24,000 eggs per sample). LC-MS, MS/MS and GC-MS measurements, data processing, data annotation and data normalization, are provided in supplementary File Text S2.

Desiccation Experiments of RE—The experiments consisted of three stages: (a) preparation, collections, storage and drying of RE, (b) comparing the viability of dried and wet RE in hatching experiments, and (c) comparing the reproductive rates of neonates hatched from wet and dried RE.

(a) Preparation, Collection and Storage of RE—Resting eggs (RE) were produced by rotifers cultured in artificial sea water (Red Sea Ltd., Israel) at a salinity of 10.8 gm L-1 (equivalent to 10 ppt sea water) and fed with thawed concentrated Nanochloropsis sp. (Galil Algae, Israel). RE were collected after 9–11 days of culture, cleaned from debris and stored at room temperature (24–26 °C) in 20 ml scintilla-
Differential abundance of hallmark proteins known for their association with dormancy. The list contains proteins that significantly differed between AM and RE or E0 (FDR $p < 0.05$; FC$ > 3$), or were present only in RE or E0 but not in AM. The minus sign indicates lower abundance in the comparison between AM vs. RE or AM vs. E0 (or higher abundance in RE or E0). Numerous proteins were not detected in AM (ND AM).

| Protein ID | Majority protein IDs | Majority protein titles | Fold-change (AM vs. RE) | Fold-change (AM vs. E0) |
|------------|----------------------|-------------------------|-------------------------|-------------------------|
| Ferritin   |                      |                         |                         |                         |
| 364        | Contig 7007          | ferritin heavy chain    | $-2443.33$              | $-1743.58$              |
| 66         | Contig 2851          | ferritin heavy chain a-like | $-28.04$             | $-21.49$                |
| 116        | Contig 9535          | soma ferritin-like      | $-1238.12$              | $-1359.40$              |
| 183        | Contig 11553; Contig 717 | soma ferritin-like; soma ferritin-like | $-53.07$             | $-43.94$                |
| Glutathione s-transferase |          |                         |                         |                         |
| 1899       | Contig 7113          | glutathione s-transferase | $-23.26$             | $-32.67$                |
| 269        | Contig 9208          | glutathione s-transferase a-like | $-2.81$              | $-2.97$                 |
| 1863       | Contig 570           | glutathione s-transferase-like | $-10.38$             | $-3.22$                 |
| 127        | Contig 533           | glutathione s-transferase-like | $-3243.34$         | $-3295.31$              |
| 347        | Contig 276; Contig 7601 | hematopoietic prostaglandin d synthase; glutathione s-transferase-like | $-5.77$             | $-6.96$                 |
| Glutathione peroxidase |          |                         |                         |                         |
| 570        | Contig 3114          | glutathione peroxidase  | $-3.58$                 | $-4.62$                 |
| Peroxiredoxin |                  |                         |                         |                         |
| 214        | Contig 9623          | peroxiredoxin prdx5    | $-285.68$               | $-293.61$               |
| 106        | Contig 9620          | peroxiredoxin prdx5    | $-1020.36$              | $-850.76$               |
| Superoxide dismutase |          |                         |                         |                         |
| 145        | Contig 10586         | superoxide dismutase   | $-4.88$                 | $-4.13$                 |
| 426        | Contig 5597          | superoxide dismutase   | $-4.41$                 | $-3.36$                 |
| 1496       | Contig 1507          | superoxide dismutase   | $-8.922$                | $-3.66$                 |
| 1963       | Singlet 16655        | superoxide dismutase   | ND AM                   | ND AM                   |
| 2452       | Contig 11562         | superoxide dismutase   | ND AM                   | ND AM                   |
| 536        | Contig 7232          | copper chaperone for superoxide dismutase | ND AM             | ND AM                   |
| Thioredoxin |                  |                         |                         |                         |
| 532        | Contig 665           | thioredoxin            | $-5.00$                 | $-4.96$                 |
| 1453       | Contig 11560         | thioredoxin h-type     | $-3.06$                 | $-3.64$                 |
| Catalase   |                      |                         |                         |                         |
| 61         | Contig 3897          | catalase                | $-21.84$                | $-18.15$                |
| Oxidoreductase |              |                         |                         |                         |
| 4          | Contig 411           | uncharacterized oxidoreductase -like | $-2223.68$        | $-1861.37$              |
| 11         | Contig 448           | uncharacterized oxidoreductase -like | $-811.40$            | $-813.88$               |
| 175        | Contig 8056          | dehydrogenase reductase sdr family member 1 | $-5.52$              | $-3.24$                 |
| 566        | Contig 2703          | dehydrogenase reductase sdr family member 1 | $-10.37$             | $-5.51$                 |
| 226        | Contig 12126         | dehydrogenase reductase sdr family member 12 | $-2.56$             | $-2.37$                 |
| Selenium  |                      |                         |                         |                         |
| 25         | Contig 11382         | selenium-binding protein | $-517.75$            | $-506.53$               |
| 35         | Contig 5327          | selenium-binding protein | $-518.03$            | $-711.96$               |
| 1805       | Contig 3315          | selenoprotein o         | $-6.39$                 | $-5.95$                 |
| 508        | Contig 169           | selenoprotein o-like    | $-6.69$                 | $-7.67$                 |
| LEA proteins |                   |                         |                         |                         |
| 1          | Contig 9631          | protein lea - isoform k | $-994.73$            | $-1161.00$              |
| 2          | Contig 227; Contig 3165 | late embryogenesis abundant protein family protein; late embryogenesis abundant protein family protein | $-1399.92$        | $-1111.06$              |
| Alpha crystallin |              |                         |                         |                         |
| 9          | Contig 7561          | alpha-crystallin a chain-like | $-537.82$            | $-523.10$               |
| Trehalose  |                      |                         |                         |                         |
| 1088       | Contig 14054         | trehalose-phosphate synthase | $-15.88$             | $-23.99$                |
| Heat-shock proteins |          |                         |                         |                         |
| 46         | Contig 11379         | heat shock hsp20        | $-1260.69$              | $-1261.17$              |
| 147        | Contig 11391         | heat shock protein      | $-217.35$               | $-186.24$               |
| 349        | Contig 2953          | heat shock protein beta-1-like | $-3.49$              | $-4.16$                 |
| 16         | Contig 11373; Singlet 16498 | heat shock protein hsp-like; heat shock protein beta-1 | $-4077.68$          | $-4275.32$              |
| 521        | Contig 404           | heat shock protein hsp20 | ND AM                   | ND AM                   |
tion glass vials filled with 10 ml sea water (10 ppt; 300–1000 RE per vial). Each vial was wrapped with aluminum foil and several vials were prepared from each culture. RE were stored for 6–15 months from three different cultures (Culture 1 for 14 months, Culture 2 for 6 months, and Culture 3 for 15 months).

(b) Comparing the Viability of Dried and Wet RE in Hatching Experiments—On the first day of the experiment (Day 0), RE were removed from one vial and distributed into five 35 mm glass Petri dishes containing 10 ml sea water (10 ppt) and 50–70 RE per dish. This procedure was repeated for RE of vials from three different cultures. The Petri dishes were placed under a cold light fluorescent lamp (200–400 μEinsteins m⁻²sec⁻¹), at 24–26 °C. The number of hatched neonates was counted after 48 and 72 h. The number of unhatched RE was counted after 96 h. The percent of hatching (H) was calculated as \( \text{%H} = (H/H + UH) \times 100 \), where H is the number of hatched neonates and UH is the number of unhatched REs. In parallel, 50–70 RE from each one of three cultures were distributed to 20 Petri dishes with a small volume of seawater (50–100 μl; 10 ppt) and were dried by evaporation. The RE seemed desiccated after 15–20 min as displayed by the appearance of salt crystals. They were further dried for 24 h at room temperature (24–26 °C). After 24 h (Day 1), 10 ml of sea water (10 ppt) were added to five Petri dishes from each of three cultures (a total of 15 Petri dishes for the three replicate cultures) and the Petri dishes were subjected to hatching conditions as described above. The remaining 15 Petri dishes for each one of the three cultures (a total of 45 dishes) containing the dried RE, were wrapped in aluminum foil and stored at 24–26 °C. Hatching experiments were performed after 14, 28, and 56 days of dry storage, using five Petri dishes on each day, from each culture. The hatching percent was compared between dried RE at the different time points and wet RE of day 0.

(c) Comparing the Reproductive Rates of Neonates Hatched from Wet and Dried RE—Neonates hatching from dried RE (stored for 56 days) and neonates that hatched from the corresponding wet stored RE, were incubated in 24-well culture plates (Nunc, Denmark). Each well contained one neonate in 1 ml of seawater (10 ppt) and algae (Nannochloropsis sp. at 12–18 \( \times 10^4 \) cells). The number of rotifers and eggs were counted after 72 h of incubation, at 24–26 °C.

Statistical analyses of the hatching experiments, was performed by Tukey Post-Hoc tests (after a Chi-square test, showing a normal distribution of results). One-way ANOVA and t-tests were used to test the differences in reproductive performance of neonates from wet and dried RE (after a Chi-square test showing normal distribution of results). Main Effects ANOVA, Tukey Post-Hoc and t-tests (after a Chi-square test, showing a normal distribution) were performed by using the Dell™ Statistica™ 13.2 package for Windows software. Statistical significant differences were considered at \( p < 0.05 \).

RESULTS

General Information on the Proteome Profiles of Resting Eggs and Amictic Eggs—In this study we carried out a comparison between the proteomes of encysted dormant embryos (“resting eggs”; RE or E0) with nondormant embryos (amictic eggs; AM). Morphological differences between AM and RE are shown in Fig. 2.

A total of 2928 proteins were identified in the nine examined samples (supplemental Table S3A), using the translated and annotated B. plicatilis transcriptome as a reference (supplemental Tables S2A and S2B). As described in the Experimental Procedures, 2631 proteins displayed at least two peptides in two replicate egg samples (“Present proteins,” supplemental Table S3B). Of these proteins, 2351 proteins were annotated (89.4%) and 1492 (64%) were assigned with KEGG KO numbers. In general, most of the proteins were shared by all egg types (Fig. 3) and relatively a small number of proteins were specific to one egg type (supplemental Table S3B). Quantitative analysis showed larger differences in the proteins’ abundance between AM and RE and smaller differences between RE and E0 groups (supplemental Fig. S1A, S1B). Although E0 consists only of two biological replicates, the detailed analyses of their proteome profiles were retained as they underscore the differences between RE versus AM. In the comparison of RE versus E0, no proteins were found to have a significantly different abundance (Fig. 4). An additional selection for proteins, which occur in one egg type and not the other or have significantly different abundance among the egg types (FDR \( p \) value < 0.05, Fold change > ±3, Fig. 3), revealed 1538 proteins differing between AM and RE (AM versus RE; supplemental Table S3B). Pairwise comparisons show that ~62% of the proteins showed higher abundance (FC>3; FDR \( p \) value < 0.05) in AM versus RE, with numerous proteins displaying FC>100 (supplemental Table S3B). For compari-
son, LFQ-normalized data showed 1498 proteins differing between AM and RE, of these 1068 proteins were identical with those of the unnormalized data mentioned before (supplemental Table S3B; see explanation for unnormalized versus LFQ normalization in the Experimental Procedures). The most abundant proteins differ greatly between AM and resting eggs as seen in Fig. 5A and 5B, and these results were similar to the LFQ-normalized data (results not shown in detail but can be retrieved from data in supplemental Table S3B). AM eggs show an abundance of cytoskeleton proteins (tubulin, actin), ATP synthase, fatty acid binding protein, α-crystalline chain and the regulatory protein 14-3-3. Most of them were also found in RE. Resting eggs show high abundance of LEA proteins, ferritin, peroxiredoxin, selenium binding protein, a mammalian ependymin-related protein 1 and α-crystalline chain. Elongation factor α-2 is abundant in all egg types.

Differences Between Resting Eggs and Nondormant Eggs in Functional Pathways—

Gene Ontology and KEGG Pathways—Functional differences between the three egg types (AM, RE and E0), were highlighted by a search for enriched pathways using Gene Ontology (GO; supplemental Table S4; supplemental Fig. S2) and KEGG pathways for each egg type (supplemental Table S5; supplemental Fig. S3). In addition, we carried out enrichment analyses for KEGG pathways and GO terms of proteins showing differential abundance in the comparisons of AM versus RE and AM versus E0. (Fig. 6 and supplemental Table S5 for KEGG pathways; supplemental Fig. S4 and supplemental Table S6 for GO terms). KEGG pathways and GO terms highlight differences in energy metabolism, especially in processes with an association of the mitochondria (e.g. tricarboxylic acid (TCA or citric cycle), electron transport, oxidation phosphorylation), lipid and fatty acid metabolism, amino acid metabolic processes, transcription, translation, splicing, protein processing and remodeling, developmental processes, and proteins with a function in response to oxidative stress and environmental contaminants.

Six general patterns were identified, by comparing the protein abundance in functional pathways, between dormant and nondormant embryos. For simplicity, we present the KEGG maps for the comparison between proteins of AM and RE as there were no significant differences in the abundance between RE and E0 (supplemental Table S3B; Fig. 4).

(a) Lower Abundance of a Large Number of Functional Proteins (or Enzymes) in a Pathway—Lower abundance was observed for a large number of proteins in fatty acid degradation (Fig. 7 and supplemental Fig. S5) and in valine, leucine and isoleucine degradation pathways (supplemental Fig. S6). In these pathways, the proteins remain available but at a lower abundance and can be recruited during exit from dormancy.

(b) Lower Abundance of Proteins With a Key Function in a Pathway—Lower abundance of proteins composing the pyruvate dehydrogenase (PDH) complex. PDH is one of the central enzymes in aerobic metabolism (see role in Artemia;
serving as a gateway between glycolysis (supplemental Figs. S7 and S8) and the citrate acid cycle (TCA cycle Fig. 8A and supplemental Fig. S9). It catalyzes the overall conversion of pyruvate to acetyl-CoA and is composed of multiple copies of three enzymatic components: pyruvate dehydrogenase (E1), dihydrolipoamide acetyltransferase (E2) and lipoamide dehydrogenase (E3). All three are also found in the 2-oxoglutarate dehydrogenase complex (ID1503), which is neither detected in RE nor E0 (supplemental Table S3B). The lower abundance of PDH and the putative absence of 2-oxoglutarate dehydrogenase in RE suggest a functional impairment of the mitochondria. The lower abundance or absence of PDH is also relevant to the pyruvate pathway (supplemental Figs. S10 and S11). Another example showing lower abundance or absence of one or more proteins in a pathway includes the ribosome, where low abundance of numerous ribosomal proteins was found (Fig. 9). Notably, mitochondrial ribosomal proteins were more abundant in AM than RE and three ribosomal proteins were not detected in RE. Low abundance of proteins was also found in the cell cycle, spliceosome and mRNA surveillance pathways (supplemental Figs. S12, S13, and S14, respectively).

(c) Lower Abundance of Multitask Proteins Such as Kinases—About 65 kinases were identified and 28 showed differential abundance in the comparison of dormant and nondormant embryos, and 11 were not detected in dormant embryos (supplemental Table S3B). Most notably, MAPK (Mitogen-activated protein kinase; ID785), with a crucial regulatory function, was not detected in RE and E0. Other examples include MAP2K1 (dual specificity mitogen-activated protein kinase kinase; ID785), a key player in cell cycle regulation, showed lower abundance in RE versus AM (FC<14.9). Also, CDK2AP2 (Cyclin-dependent kinase 2-associated protein 2; ID2362), with a role in regulating the self-renewal of embryonic stem cells (ESCs) and regulation of microtubule organization of metaphase II oocytes (see http://www.uniprot.org/uniprot/075056), showed lower abundance in RE versus AM (FC<13.4). Interesting putative pathways with proteins showing differential abundance displayed in dormant versus non-dormant embryos include the Neurotrophin signaling pathway, MAPK signaling pathway, GnRH signaling pathway, Toll-like receptor signaling pathway, Wnt signaling pathway, Insulin signaling pathway, Oocyte meiosis, Progesterone-mediated and oocyte maturation (See http://www.kegg.jp; supplemental Table S7). The kinases with significantly differing abundance between AM and RE or E0, include the MAPK, JNK, and ERK 1/2.

(d) Higher Abundance of Proteins with a Putative Inhibitory Function in a Pathway—The protein PI13 (ID110) showed higher abundance in the proteasome of RE (supplemental Fig. S15). This protein was described as a proteasome inhibitor of mammalian cells and a modulator of proteasome formation. Recent studies indicate that in Drosophila melanogaster the overexpression of DmPI13 increased proteasome activity in vivo. However, in vitro studies with purified DmPI13, revealed that the protein activates 26S proteasomes but inhibits the activity of the proteasome core particle (85).

(e) Other Patterns—Mixed patterns of higher and lower abundance of proteins or occurrence of proteins only in RE in a functional pathway. This pattern was found in oxidative phosphorylation (Fig. 10 and supplemental Fig. S16) and the pentose phosphate pathway (supplemental Figs. S17 and S18). Additional examples include protein processing and endoplasmic reticulum pathways (supplemental Fig. S19).

(f) Higher Abundance of Hallmark Proteins with a Specific Protective Function—Numerous proteins with specific functions in dormant forms showed differential abundance in the comparison of AM and RE (Table I). This list includes LEA proteins, ferritin, proteins associated with cell stress such as glutathione s-transferase like proteins, peroxidredoxin, thioredoxin, superoxide dismustase, catalase, oxidoreductase, heat shock proteins, α-crystalline proteins, selenium proteins, and proteins associated with synthesis of trehalose (52, 53). Most noticeable are the high abundance of proteins in the

83)
comparison of RE versus AM to the extent of FC>500–1,000. Additional information on these proteins is found in the supplementary File Text S1.

Detailed Description of Pathways with a Known Association in Dormancy or Diapause—

(a) Energy Metabolism and the Mitochondria—Dormancy has been associated with a depression in metabolism (84) or alternative metabolic pathways (64, 86, 87), and results shown above support the contention of downregulation of metabolism in RE in association with oxidative phosphorylation, the citrate cycle, and pyruvate metabolism (Fig. 10 and supplemental Fig. S16; Fig 8A and supplemental Fig. S9; supplemental Fig. S10 and supplemental Fig. S11). Proteins associated with NADH dehydrogenase, cytochrome c oxidase, cytochrome c reductase and F-type ATPase, showed lower abundance in RE as displayed in the oxidative phosphorylation pathway (Fig 10). The putative significant down-regulation of metabolism is supported by metabolome analyses that showed the low abundance/absence of ATP in RE (Fig 11) and significantly lower abundance of metabolites in the citrate cycle (Fig 8B). Because the pentose phosphate pathway was shown to play a role in embryonic diapause of mouse blastocysts (30) we searched for the abundance of proteins in the pentose phosphate pathway (supplemental Figs. S17 and S18). Two proteins showed higher abundance in RE: glucokinase/gluconate (ID1707) and transaldolase (ID67), which are associated with phosphorylation of D-glucuronate and conversion of D-ribose-5 phosphate to D-glyceraldehyde-3-phosphate, respectively. D-glucuronate or β-D-glucose-6-phosphate could lead in RE to the synthesis of D-glyceraldehyde-3P and a participation in glycolysis. Ribose-phosphate pyrophosphokinase (ID1431), which is the gateway to nucleotide metabolism from D-ribose-5-phosphate, showed lower abundance in RE, suggesting a possible diversion from metabolism of purine, pyrimidine or histidine to the nonoxidative metabolic pathway of the Pentose Phosphate Pathway.

Another source of energy involves the degradation or metabolism of amino acids. In general, proteins of pathways associated with amino acid metabolism were more abundant in AM versus RE. One example includes proteins that were associated with valine, leucine and isoleucine degradation mentioned above (supplemental Fig. S6), with connections to the citrate cycle (through the synthesis of acetyl-CoA from amino acids). Statistically significant lower amounts of amino acids were identified in RE (Fig 12).

(b) Lipid, Fatty Acid Metabolism and Lipoproteins—Changes in the lipid metabolism were often associated with dormancy in several organisms (65, 88). In rotifers, lipid droplets were identified in resting eggs (89) and therefore these pathways were examined here in more detail. Numerous proteins with functions in lipid and fatty acid metabolism were identified in eggs but only a few showed differences in abundance between RE and AM (supplemental Table S3B). As mentioned before, proteins functioning in fatty acid degra-
tion, which are a potential source for generating energy, were less abundant in RE (Fig. 7 and supplemental Fig. S5). Of the fatty acid binding proteins, only fatty acid-binding brain protein (ID 1335) showed higher abundance in RE. Interestingly, lipid storage droplets surface-binding protein 2-like (ID49), a perilipin homolog, was significantly more abundant in RE versus AM. Similarly, lipid phosphate phosphohydrolase 1 isoform (ID1738) with a broad function in lipid metabolic processes, was found in RE but not in AM. A monoacylglycerol lipase (ID1993; abhd12-like protein) that hydrolyzes glycerol monoesters of long-chain fatty acids was not detected in RE (90). Nonspecific lipid transfer protein (ID776, ID1027) with a function in mediating the transfer of all common phospholipids, cholesterol, and gangliosides between membranes and a role in regulating steroidogenesis (91), was highly abundant in AM versus RE (FC/H11022 = 70.7, FC/H11022 = 108.2, respectively). Also, vigilin (ID297; supplemental Table S3) was highly abundant in AM in comparison with RE.

(c) DNA Synthesis, Replication and the Cell Cycle—Suspension of the cell cycle is one of the characteristics of dormancy (see the Introduction; 1, 6). A relatively small number of proteins participating in DNA replication and cell cycle were identified in rotifer eggs (supplemental Fig. S12). These include the Mcm2–7 complex, DNA ligase1-like (ID1639), DNA
polymerase delta subunit 3 (ID2467), were not detected in RE. DNA polymerase delta subunit 2 (ID2874) was more abundant in AM versus RE (FC/H11022 24.6; supplemental Table S3). In contrast DNA polymerase beta (POLB; ID75) performing base excision repair and required for DNA maintenance, replication and recombination, was higher in RE. A few regulatory proteins with an association of the cell cycle, were less abundant or were not detected in RE. PCNA (Proliferating cell nuclear antigen; ID1400) showed lower abundance by 84.7 fold in RE (versus AM). Zinc finger homeobox protein 2-like (ID2338) and homeobox-containing protein 1 isoform x2 (ID2638) were not detected in RE. PCNA is an auxiliary protein of DNA polymerase delta and is involved in the control of eukaryotic DNA replication. GO annotations for the homeobox proteins include sequence-specific DNA binding transcription factor activity and sequence-specific DNA binding. In addition, mitotic checkpoint protein (bub3-like protein; ID1923), with a putative function in regulation of chromosome segregation during oocyte meiosis in mouse (92), was not detected in RE.

(d) Transcription, Translation, Degradation and Proteome Remodeling—In general, numerous proteins associated with these functions showed lower abundance in RE, indicating an impairment of these functions. DNA-directed RNA polymerases I and III subunit rpac1 (ID667) and DNA-directed RNA polymerase II subunit rpb3 (ID1089), which catalyze the transcription of DNA to RNA, showed lower abundance in RE in comparison with AM. RNA polymerase II-associated factor 1 homolog (ID1030), mediator of RNA polymerase II transcription subunit 8 isoform x1 (ID1692) and RNA polymerase II subunit a c-terminal domain phosphatase ssu72 (ID1976), with multiple functions during transcription, were also with lower abundance in RE (supplemental Table S3). As shown before, a relatively large number of cytoplasmic ribosomal proteins were identified in rotifer eggs (Fig. 9) and numerous
with higher abundance of proteins in AM. Of interest is the elongator complex proteins 6 (ID2003) which was highly elevated in RE (FC/H11022 24). This protein is a histone acetyltransferase component of the RNA polymerase II (Pol II) holoenzyme and involved in transcriptional elongation (93). The elongator may play a role in chromatin remodeling and acetylation of histones H3 and probably H4 and functions in cell migration (see below). The proteins of the spliceosome showed differential abundance in the comparison of AM and RE (supplemental Table S13). In the mRNA Surveillance pathways (supplemental Fig. S14), many proteins were significantly higher in AM and numerous proteins were not detected in RE, suggesting lower translation and processing of mRNAs in RE. One of the most abundant proteins in the comparison of AM versus RE (FC>88) was a 116 kDa u5 small nuclear ribonucleoprotein component-like protein and part of the U4/U6-U5 tri-snRNP complex (ID2264; supplemental Table S3B), which is required for pre-mRNA splicing (94).

In contrast to the pathways discussed above, most of the proteins that were associated with protein degradation by the proteasome, showed similar abundance in AM and RE (supplemental Fig. S15). The proteasome activator complex subunit 3 (ID960) which may be involved in regulation of the cell cycle was higher in AM. Two proteins that were higher in RE include the Rpn7 (ID137), a member of the regulatory particle and the PI31, which was discussed previously. A protein annotated as proteasome activator subunit 3 (PA28 gamma; ID960) was higher in AM. Proteins associated protein processing in the endoplasmic reticulum were identified in AM and RE (supplemental Fig. S19), with differential display of a numerous proteins.

(d) Histones and Chromatin—Histones play a central role in transcription regulation, DNA repair, DNA replication and chromosomal stability. We identified proteins annotated as histone proteins (histone h1, histone h2a, histone h4, several histone-like proteins) and histone deacetylases (Supplemental Table S3B) which promote the deacetylation of lysine residues on the N-terminal part of the core histones. A histone chaperone asf1-like protein (ID1693), with a function in association with the chromatin assembly factor 1 (CAF-1) promotes replication-dependent chromatin assembly and plays a role in the formation of silent heterochromatin (95). Most of the proteins showed higher abundance in AM in comparison with RE or E0, or were not detected in RE or E0. However, one protein with significantly higher abundance in RE versus AM (FC>64), is the core histone macro-like isoform x2 (ID486). This is a variant of histone H2A which replaces conventional H2A in a subset of nucleosomes, where it represses transcription, nucleosomes wrap and compact DNA into chromatin, limiting DNA accessibility to the cellular machineries which require DNA as a template. Replication protein A (RPA) is a heterotrimeric protein complex of three polypeptides of 70, 34, and 13 kDa and is required for many DNA metabolic processes, including DNA replication, repair and DNA damage checkpoint activation. This protein (ID2067, ID1068, and ID267), showed higher abundance in AM versus AM with FC ranging 3.7 to 4.5. It is also required for the recruitment of the DNA double-strand break repair factors RAD52 and RAD51 to chromatin in response to DNA damage. Additional proteins with a lower abundance in RE and a function in modeling chromatin, are the chromobox proteins (ID1407, ID599), which are components of heterochromatin.

(e) Cytoskeleton—A relatively large number of cytoskeleton associated proteins with a diversity of functions, showed highly differentially abundance between AM and RE (supplemental Table S3B). This includes highly abundant proteins of AM versus RE, like microtubule-associated protein futsch-like (ID584; FC>110) with a neuromuscular function during embryogenesis and troponin I (ID663) that was not detected in RE and troponin t (ID796) that showed lower abundance in RE (FC<244). In contrast, a higher abundance in RE (with a FC ranging from -9 to -17 in AM versus RE) was found for tubulin alpha (ID1417, 1833), which is the major constituent of microtubules and functions in tight and gap junctions.

(f) Oocyte, Spermatozoa and Cilia Proteins—Proteins associated with oocytes and spermatozoa were generally more abundant in AM (supplemental Table S3B). Two proteins with high differences in abundance between AM and RE include
the nuclear auto-antigenic sperm protein (ID48), with a FC >334 and the oocyst wall protein (96), with a FC >179 (ID251). As proteins associated with sperm may involve proteins associated with cilia (as mentioned above), additional proteins that were annotated as cilia proteins are given in supplemental Table S3. Proteins identified as radial spoke head proteins (ID1022, ID1267, ID1409, ID2058, and ID 2472) showed lower abundance (AM versus RE, FC ranging from 34.6 to 186.2) or were not detected in RE. These proteins form a probable component of the axonemal radial spoke head. Radial spokes (97) are regularly spaced along cilia, sperm and flagella axonemes. Similarly, dynein proteins (ID1033, ID1381, ID1791, ID2168, ID2533), except for two (ID929, ID2694), showed differences in their abundance between AM and RE (FC ranging from 12 to 31) or were not detected in RE. The hu-li tai shao-like protein (adducing; ID597) with a function in the structure of egg chamber proteins (in Drosophila, 98), was more abundant in AM versus RE. Conversely, Piwi-like protein 1 ID315, ID1045, ID1138, ID2880) with a function in germline stem cell division (99) showed higher abundance in RE (FC ranging from 2.6 to 7.2).

(g) Ras Proteins—The Ras superfamily of small guanosine triphosphatases (GTPases) function as GDP/GTP-regulated molecular switches (100). Twenty-one out of the 29 identified proteins in this family (supplemental Table S3B) showed differential abundance between AM and RE and three were not detected in RE. On the other hand, ras-related protein rab-5a (ID2445) with a putative function in fusion of plasma membranes and early endosomes, was not detected in AM.

(h) Regulation—Transcription Factors, Hormones, and Signaling Pathways—in search for pathways that could be involved in regulation of dormancy in RE, we identified putative proteins that were associated with transcription factors, hor-
mones with known functions in rotifers and other organisms and proteins associated with signaling pathways (supplemental Table S3B). All of the proteins were more abundant in AM and some of them were not detected in RE or E0. In this list, we find two members of the 14-3-3 proteins, 14-3-3 protein epsilon (ID616; AM versus RE, FC/H11022 71.7) and 14-3-3 protein zeta D64; versus RE, FC/H11022 2.64). Two proteins were annotated as insulin-like growth factor 2 mRNA-binding protein1 (ID1263, ID1511), with higher abundance in AM (FC/H11022 29.1 and FC/H11022 35.6). One of them (ID1263), with a putative function as an RNA-binding factor, recruiting target transcripts to cytoplasmic protein-RNA complexes (mRNPs), allowing mRNA transport and transient storage and shields from endonuclease attacks or microRNA-mediated degradation (http://www.uniprot.org/uniprot/Q9Y6M1).

A juvenile hormone esterase binding protein (ID1562), suggesting a role for JH in rotifers, was identified in rotifer eggs. Juvenile hormone esterase degrades juvenile hormone, which acts in conjunction with ecdysteroids to control gene expression in insects (101). Proteins associated with steroids include two proteins annotated as estradiol 17-beta-dehydrogenase 8 or 12, a membrane-associated progesterone receptor component 2-like and a steroid receptor RNA activator 1. An enzyme with a function in processing of protein hormones and of neuropeptides from their precursors (ID966; neuroendocrine convertase 2-like; 102) showed highly differentially abundance in the comparison of AM and RE. A protein associated with the activation of Notch-1 (delta and notch-like epidermal growth factor-related receptor; ID2657), was identified in AM but was not detected in RE or E0. Insulin signaling pathway has been implicated in diapause regulation in insects (103) and two proteins associated with insulin were mentioned before. Many proteins could be associated with downstream functions of signaling pathways (e.g. kinases; Ras family proteins; calmodulin and others).

Calcium—Proteins annotates as calmodulin, calreticulin, calcineurin, calnexin with an association to calcium storage or binding of calcium, were more abundant in AM versus RE, or were absent in RE (supplemental Table S3B). Interestingly, one protein of the neural calcium sensor 2 (ID265), was highly more abundant in RE (FC>408 for RE versus AM) but others showed variable patterns. Moreover, mammalian ependymin-

Fig. 10. A KEGG map showing the proteins in association with Oxidative phosphorylation pathway. Boxes in gray show the proteins that were identified in rotifer eggs. Boxes in red and blue show proteins that were significantly more abundant in AM or RE, respectively. Turquoise boxes show the proteins that were detected only in RE. Statistically significant differences were indicated by 1-Way ANOVA test (FDR p value 0.05, FC>3).
related protein 1 (ID219) with a putative role in calcium-dependent cell adhesion, was one of the most abundant proteins in RE and E0 (FC>6742 in RE versus AM). Similarly, recoverin (ID54), was highly abundant in RE in comparison with AM (FC\text{H11022}/523.8 in RE versus AM). This is a neuronal calcium-binding protein with a key role in the inhibition of rhodopsin kinase, which regulates the phosphorylation of rhodopsin and is involved in the recovery phase of visual excitation and in adaptation to background light (104). This protein might play a role in light sensing, vital to hatching from RE.

Within the list of putative enzymes, a few enzymes (showing differential abundance in the comparison of AM and RE), with interesting functions were identified, such as chitinase (three proteins; ID 1770, chitinase domain-containing protein 1; ID2398 acidic mammalian chitinase-like; ID2466 probable chitinase 3-like), aurora kinase a (ID2279) with a function during meiosis and mitosis (105); where its activity peaks during the G2 phase to M phase transition in the cell cycle. Another enzyme of interest is the chorion peroxidase-like (ID543, ID1521; identified only in RE or E0) that catalyzes...
Protein cross-linking through dityrosine formation and phenol oxidase-catalyzed chorion melanization in insects (106). A protein associated with cleavage of β-carotene was identified (carotene-9 oxygenase, ID1789; beta,beta-carotene 9’,10’-dioxygenase, BCDO2), with small differences in abundance in the two egg types.

Correlation Between Proteome and Transcriptome Expression Profiles—The relationship between protein abundance and RNA expression levels show a negligible positive correlation (107) for AM (Pearson’s correlation coefficient, $R^2 = 0.2112$) and a low positive correlation (Pearson’s correlation coefficient, $R^2 = 0.4357$) for RE (Fig 13A, 13B), with $p$ value $< 2.2e-16$ for both egg types (see also supplemental Table S8). Three categories were defined in comparing the abundance levels of the proteins with the corresponding expression levels of mRNAs: (a) A group with an apparent concordance between proteins and the corresponding mRNAs (Fig 13A, 13B; in blue), (b) A group with an apparent discordance between proteins and the corresponding mRNAs (Fig 13A, 13B; in red) and (c) A middle group for which neither concordance nor discordance can be shown (Fig 13A, 13B; black). The proteins in apparent concordance with the corresponding RNAs of RE show the following enriched KEGG pathways (supplemental Table S5): ribosome, oxidative phosphorylation, valine, leucine and isoleucine degradation, RNA transport, carbon metabolism and citrate cycle. On the other hand, proteins of the AM group display an apparent concordance with mRNAs for only two KEGG enriched pathways, oxidative phosphorylation and spliceosome (supplemental Table S5F). The GO terms for the comparison of proteins and the corresponding mRNAs of AM and RE, are shown in supplemental Table S9. Interestingly, the hallmark proteins (protein lea- isoform k, LEA protein family protein, alpha-crystallin a chain-like, heat shock hsp20, soma ferritin-like; ferritin heavy chain) with high abundance in RE and/or E0 also show relatively high levels of the corresponding mRNAs (Fig 13B).

Effect of Drying on RE Hatching and Reproductive Rates of Hatched Neonates—Drying of RE had no adverse effect on the percent of hatching after 1, 14, and 56 days (Fig 14). Differences in hatching of RE from the three cultures were not statistically different, although they were kept for different times at room temperature before the drying experiments (Univariate tests of significance; $p = 0.77$). Statistically significant higher hatching was observed between wet and dried RE after 28 days of drying. Each one of the neonates that hatched from the 56-day dried RE, produced a total of $3.54 \pm 1.48$ (mean ± S.D.) rotifers and eggs ($n = 110$), after 72 h of incubation, like the number $(3.37 \pm 0.98$, mean ± S.D., $n = 113$; $p = 0.32$, t test) produced by neonates hatching from the control wet RE.

DISCUSSION

The phenomenon of dormancy is of wide interest as it occurs in a large number of organisms from prokaryotes to mammals (3, 4). However, proteomic information on dormancy is limited, particularly in nonmodel organism with dormancy exhibited during embryonic development. The intention of the current study is to reveal mechanisms of importance for maintaining dormancy in encysted rotifer embryos by quantitative proteomics, as most wide scale studies in rotifers and other organisms exhibiting dormancy, focused mostly on transcriptome profiling. The overall relatively low correlation between expression levels of mRNAs and abundance of proteins (75,76; in the comparison of pre-diapause and nondiapause locust eggs (73) and Fig 13), renders proteome profiling an important tool for providing an additional valuable view on putative functional changes during dormancy.
The differences between the RE from the three cultures were not statistically significant differences (p < 0.01) between hatching of wet RE (day 0) or RE dried for 1 day and RE after 28 days of dried storage. The differences between the RE from the three cultures were not statistically significant (p > 0.05).

A total of 2631 proteins were identified with at least two peptides in two replicate egg samples, comprising ~15% of the current available transcriptome. Assuming conservation in transcript and protein sequences, it provides information on many putative conserved functional, cellular and molecular pathways, some of which were supported by metabolome profiling. Remarkably, a relative large amount of information was obtained by quantitative proteome profiling while using an assembled transcriptome, as the genome of this species has not yet been sequenced. Proteome profiling provides here a one-time snapshot on the abundance of many proteins in three egg types. It should be noted that protein levels may not fully reflect the functional levels of a specific protein within a pathway, as this may also depend on protein turnover rates. However, low or high abundance, or detection in one group and absence (below detection level) in another group, provides insight into putative functional directions. Taking these restrictions into account, the proteome profiles were analyzed with an attempt to provide a coherent view of the functional roles of the identified proteins in supporting dormancy.

About 62% of the proteins were significantly more abundant in AM compared with RE and E0, and 47 and 42 proteins showed FC > 100 in these respective comparisons (supplemental Fig. S3). The difference between RE and E0 versus AM is also demonstrated in the list of the most highly abundant proteins of each group (Fig 5A, 5B). Differences in the abundance of proteins between resting eggs before (RE) and after (E0) the obligatory dormant period (Fig 4; supplemental Table S3B), were not significant but RE displayed 22 proteins that were not determined for E0. Although functional differences were suggested during the obligatory dormant period by E0 enriched KEGG pathways (supplemental Fig. S5), these were not statistically significant (Fig. 4).

About 16 and 28 proteins were similar to those described for cysts of Artemia sinica and Artemia franciscana, respectively (77, 78). Only one of the proteins (calcineurin), unique to the dormant silkworm, was also found as specific to RE (79). In the locust proteome (73), ten of the annotated proteins showing differential abundance between prediapause and nondiapause locust eggs, were found in rotifers. Five showed higher abundance in pre-diapause eggs versus nondiapause eggs and in RE versus AM, including glutathione S-transferase, hydroxysteroid dehydrogenase-like protein 2-like, alcohol dehydrogenase class-3, glucosamine-6-phosphate isomerase 2, isocitrate dehydrogenase. Dolichyl-Diphosphooligosaccharide—protein glycosyltransferase (which is involved in the pathway of protein glycosylation), showed lower differential abundance in the comparison in prediapause eggs versus, nondiapause eggs and in RE versus AM.

Proteome profiling revealed the large number of functional pathways that were putatively compromised in the dormant encysted RE, especially those in mitochondria (Fig 15 and Fig 16). It is usually assumed that metabolism is reduced during dormancy to conserve resources for long-term survival and resumption of life activities. We show here that metabolism is highly reduced in rotifer encysted embryos, as ATP levels were below detection levels (Fig 11). These lower ATP levels can be attributed to a significantly lower activity of the TCA cycle as demonstrated by proteome and metabolome profiling (Fig 8A, 8B; supplemental Fig. S11). In addition, proteome profiling suggests compromised function of the pyruvate metabolic pathway, the oxidative phosphorylation pathway and mitochondrial ribosomes. However, differences were not observed in the abundance of most proteins of the glycolysis or pentose phosphate pathways, suggesting that the proteins of these pathways are a source for low levels of ATP during the dormant period or can be quickly recruited to provide energy at the time of resumption of metabolism, during the exit from dormancy. A diversion from the citrate cycle to the pentose phosphate pathway is characteristic of mammalian preimplantation embryos, which may exhibit a form of dormancy (73), as this would lead to synthesis of macromolecules for creating new biomass (109).

Long-term survival of hydrated Artemia encysted embryos was attributed to a role of V-ATPases in intracellular acidification leading to metabolic depression (6). As there were no significant differences in the levels of the proteins between AM and RE (Fig 10) like the observation in Artemia (6), the role of V-type ATPases in dormant rotifer eggs remains unclear. Interestingly, F-type ATPase that transforms the energy of ATP hydrolysis to electrochemical potential differences was more abundant in RE versus AM (Fig 10). Moreover, a transcript encoding a V-type H(+)-translocating pyrophosphatase was differentially expressed in 30 min illuminated REs.
Lipids were described as an energy source during dormancy in many organisms, ranging from *C. elegans* to hibernating bears (110, 111). In the Asian tiger mosquito, however, downregulation of genes involved in lipid catabolism was observed during diapause (87). The picture for utilization of lipids in rotifer REs is far from being clear. Proteins involved in fatty acid degradation and fatty acid metabolism showed significantly lower abundance in RE and E0 (Fig. 7 and supplemental Fig. S5). Proteins associated with beta-oxidation, oxidation of long chain fatty acids, and biosynthesis of saturated and unsaturated fatty acids, were higher in AM than RE (supplemental Table S3B), possibly indicating lower utilization of lipids in encysted embryos because of an overall reduction in metabolism. Vigilin, a high-density lipoprotein, with a function in cell sterol metabolism was highly abundant in AM. On the other hand, a few proteins with an association to formation of lipid droplets, lipase activity and lipid transport, were higher in RE. The occurrence of the yolk protein vitellogenin was reported for rotifers (112), but we could not find any matches for this protein in our analyses (results not shown).

Protein and amino acid degradation is another option for generating energy. The lower abundance of amino acids in RE (Fig 12) and the lower abundance of proteins in the Valine, leucine and isoleucine metabolism pathway (supplemental Fig. S6) indicates lower availability of amino acids for metabolism and translation. In general, there were no differences between AM and RE in the level of most proteins of the proteasome but the differential abundance of three proteins, especially of PI131 may suggest regulation of the proteasome function in REs (supplemental Fig. S15). PI13 is an inhibitor of the proteasome in mammals but in *Drosophila melanogaster* it stimulates proteasome activity while being regulated by the F-box protein Nutcracker during normal development (84).

Additional functions suggested as being compromised in REs include the cell cycle progression, cytoskeleton proteins, replication, transcription, translation, amino acid metabolism, proteome remodeling among others (supplemental Table S3B). Similar general trends were reported for proteome studies in insect diapause (73, 79, 113). Most noticeable is the large number of enzymes and protein kinases that showed differentially abundant between dormant and nondormant eggs (supplemental Table S3B). Most interestingly, mitogen-activated protein kinase 1-like (MAP), regulating many functions, including cell cycle and with a role in adaptation to desiccation in nematodes (114) was not detected in RE or E0. In addition, numerous Ras family proteins showed significantly lower abundance in AM versus RE or AM versus E0. The Ras superfamily of small GTPases, act as binary molecular switches and function as modulators of a remarkably complex and diverse range of cellular processes and transcription factors (supplemental Table S3B; 100).

Additional interesting differences include lower protein abundance in RE and E0 of proteins associates with histones and chromatin, playing a central role in transcription regulation, DNA repair, DNA replication and chromosomal stability. In *Artemia*, acetylation of chromat-in-associated histone H3 lysine 56 inhibits the development of cysts (115). Hanson et al., 2013 (74), suggest a role for histones during sexual reproduction in *B. calyciflorus*. Information on the involvement of key signaling molecules, transcription factors, regulatory peptides or hormones is scarce, probably because of their lower abundance (supplemental Table S3B) but some showed higher abundance in AM versus RE or E0. They include most notably the 14-3-3 proteins with a wide array of regulatory functions (116). The 14-3-3 proteins constitute a highly conserved protein family, with more than 100 signaling proteins serving as ligands for other proteins (117). They play an important role in a wide range of cellular processes, including signal transduction, apoptosis, cell cycle progression, regulation of the Ras/Raf/MAPK cascade, the anterior-posterior axis during embryonic development and serve as checkpoint activators within eukaryotic cells. 14-3-3 proteins participate in light signaling and bind to phytochrome interacting factors in plants (118) and melatonin synthesis in mammals (119). As mentioned before, light promotes hatching of rotifer REs (44, 57). In addition, 14-3-3 showed differential abundance in the
comparison of dormant and nondormant silkworm eggs (79). Moreover, 14-3-3 proteins are also associated with the forkhead box O (FOXO) transcription factors (120). It was demonstrated recently that numerous distinct groups of genes that are targets of FOXO promote the diapause phenotype in insects. FOXO is a transcription factor functioning downstream of insulin and juvenile hormone signaling in insects and the targets include genes that are associated with metabolism, development/cell cycle, intra- and extracellular signaling, transcription, translation, protein modification, stress response, cell death and apoptosis (6, 121). The results presented here suggest a role for FOXO in dormancy of encysted rotifer eggs as a few proteins with an association to insulin and juvenile hormone metabolism were identified here, but these results require additional verification. Additional hormones may be involved in regulation as proteins with an association to estradiol synthesis and progesterone with known functions in rotifers, were also identified in the current study (43, 122).

Like other dormant organisms, a unique set of genes is evoked and highly abundant in rotifer REs (51–53). The abundance of stress tolerance related proteins was significantly altered in dormant eggs (Table I) like the changes observed in other organisms displaying dormancy (3, 8, 9). These include glutathione S- transferase (GST) with antioxidant activity, protecting cellular components from oxidative damage. Ferritin is one of the most abundant proteins in dormant RE and E0 and shows higher abundance in diapausing insects (123, 124) and copepods (125). It has been shown that the concentration of ferritin increases in response to stresses such as anoxia (126). Artemia cysts accumulate artemin, a diapause-specific ferritin homologue with chaperon activity (127). Ferritin serves as an iron storage protein and is the main component of yolk proteins in snail eggs (128). LEA proteins with an important role in facilitating desiccation (129), show high abundance in RE (Fig. 5; Table I and supplementary Text S1). There is a low correlation between the lists of the most abundant proteins in AM and resting eggs (RE or E0; Fig. 5A, 5B) and the list of most abundant proteins does not match fully with the list of most abundant transcripts (51). A detailed comparison in the level of transcripts and the corresponding proteins indicates higher concordance in RE in comparison with AM (Fig 13A, 13B), possibly because of a lower protein turnover in RE (108). Furthermore, the apparent concordant proteins of RE display a larger number of enriched KEGG pathways than those of AM proteins (supplemental Table S5). Interestingly, hallmark proteins with a specific function in RE (e.g. LEA proteins; ferritin, alpha crystalline, superoxide dismutase) also display high levels of mRNAs (Fig 13B). These proteins degrade during revival from dormancy (52–54), raising the question whether these mRNAs function in translation during the dormant period or do they serve as a reservoir for nucleic acids.

In general, rotifer dormant embryos display pathways like dormant stages that survive desiccation, such as Artemia cysts and the sleeping chironomid (8, 9, 16). We did not find differences in the hatchability of dried and wet stored RE up to 56 days of storage. Moreover, there were no differences in the reproductive output of neonates hatching from these dry or wet stored RE. Similar hatching percent of dried and wet RE after 3 months of storage were demonstrated for the freshwater rotifer species Brachionus calyciflorus (130) but most other studies (e.g. 131, 132) do not provide quantitative information on the comparative success of hatching from wet and dried field samples or quantitative information on subsequent viability. RE withstand desiccation and this trait is probably embedded already at the time of their formation, as they survive after a short desiccation period, in contrast to a slow dehydration regime required for high recovery, by bdelloid rotifers (133) and some tardigrade species that do not survive quick drying (134). Moreover, metabolism is probably reduced shortly after their formation, explaining the long-term survival in the hydrated or desiccated form. The functional events taking place during the obligatory dormant period require additional studies.

In conclusion, the current study expands our knowledge and provides insight on the wide scope of functions that are regulated in dormant encysted embryos. The interpretations from proteome profiling on a reduced metabolism and translation in dormant embryos were supported by metabolome analyses, strengthening to some extent, the conclusions, on other putative functional pathways that were identified in our analyses. The functional activity of a pathway depends on numerous enzymes or proteins involved in this pathway. In some pathways, one or more of the proteins were not detected, suggesting a malfunction of the pathway. In different organisms, the malfunction of a pathway may depend on changes in different proteins within the pathway, while leading to a similar functional effect (phenotypic function). Consequently, it can be suggested that a divergence between organisms (10) may be because of differences in the regulated genes and/or proteins within a functional pathway, while leading to the same result. Moreover, these differences may also reflect a divergence in the gene regulatory pathways (e.g. hormones, epigenetic changes etc.) between organisms. The relatively low concordance between transcriptome and proteome profiling provides another explanation for the low transcriptional similarity among dormancies across species, despite the resemblance in physiological phenotypes.

Acknowledgments—We thank Reini Malka Hamo for technical assistance. The proteomics analyses were performed at the Smoler Proteomics Center Technion and the Technical assistance of Ilana Navon in operating the mass spectrometers is much appreciated. The purchase of the Q-Exactive-Plus mass spectrometer was supported by the Israel Science Foundation (ISF), the I-CORE Program of the Planning and Budgeting Committee and the ISF (Grant 1775/12).

Molecular & Cellular Proteomics 16.10

1765
REFERENCES

1. Denlinger, D. L., Yocum, G. D., and Rinehart, J. P. (2012) Hormonal control of diapause. In: Gilbert, L. I., ed. Insect Endocrinology, pp. 430–463, Elsevier, Amsterdam.

2. Keilin, D. (1959) The problem of anabiosis or latent life: history and current concept. Proc. R. Soc. Lond. B 150, 149–191.

3. Lubzens, E., Cerda, J., and Clark, M. S. (2010) Topics in Current Genetics 113, 1–11.

4. Lubzens, E. (2015) Fre`re Jacques/Dormez vous? Dormancy, an intriguing concept. Proc. R. Soc. Lond. B 282, DOI: 10.1098/rspb.2015.1113 (Editorial).

5. Hand, S. C., Menze, M. A., Borcar, A., Patil, Y., Covi, J. A., Reynolds, J. A., and Podrabsky, J. E. (2016) Physiological strategies during diapause: insights from Artemia francicana embryos and other animals. J. Insect Physiol. 57, 584–594.

6. Hand, S. C., Denlinger, D. L., Podrabsky, J. E., and Roy, R. (2016) Mechanisms of animal diapause: recent developments from nematodes, crustaceans, insects, and fish. Am. J. Physiol. Regul. Integr. Comp. Physiol. 310, R1193–R211.

7. Kostal, V. (2006) Eco-physiological phases of insect diapause. J. Insect Physiol. 52, 113–127.

8. MacRae, T. H. (2010) Gene expression, metabolic regulation and stress tolerance during diapause. Cell Mol. Life Sci. 67, 2405–2424.

9. MacRae, T. H. (2016) Stress tolerance during diapause and quiescence of the brine shrimp, Artemia. Cur. Stress Chaperones 21, 5–18.

10. Ragland, G. J., Denlinger, D. L., and Hahn, D. A. (2010) Mechanisms of suspended animation are revealed by transcript profiling of diapause in the flesh fly. Proc. Natl. Acad. Sci. U.S.A. 107, 14490–14491.

11. Shen-Miller, J., Mudgett, M. B., Schoff, J. W., Clarke, S., and Berger, R. (1995) Exceptional seed longevity and robust growth: ancient sacred depths in lake sediments. Am. J. Bot. 82, 1367–1380.

12. Sallon, S., Solowy, E., Cohen, Y., Korinchsky, R., Egli, M., Woodhatich, I., Simchoni, O., Kislev and M. (2008) Germination, genetics, and growth of an ancient date seed. Science 320, 1464.

13. Fontenato, D., Bunnefeld, N., and Westberg, M. (2012) Long-term survival of microscopic animals under desiccation is not so long. Astrobiology 12, 853–869.

14. Hashimoto, T., Horikawa, D. D., Saito, Y., Kuswahara, H., Kozuka-Hata, H., Shin-it, T., Minakuchi, Y., Ohishi, K., Motoyama, A., Aizu, T., Enomoto, A., Kondo, K., Tanaka, S., Harai, Y., Koshikawa, K., Sagara, H., Miura, T., Yokobori, S., Miyagawa, K., Suzuki, Y., Kubo, T., Oyama, M., Kohara, Y., Fujiyama, A., Arakawa, K., Katayama, T., Toyoda, A., and Kunieda, T. (2015) Extremotolerant tardigrade genome and improved radiotolerant resistance of human cultured cells by tardigrade-unique protein. Nat. Commun. 7, 12958. doi: 10.1038/ncomms12958.

15. Clark, M. S., and Worland, M. R. (2008) How insects survival the cold: molecular mechanisms - a review. J. Comp. Physiol. B. 178, 917–933.

16. Cornette, R., and Kikawada, T. (2011) The induction of anhydrobiosis in the sleeping chironomid: current status of our knowledge. IUBMB Life 63, 419–429.

17. Gusev, O., Suetsugu, Y., Cornette, R., Kawashima, T., Logacheva, M. D., Kondrashov, A. S., Penin, A. N., Hatanaka, R., Kikuta, S., Shimura, S., Kanamori, H., Katayose, Y., Matsumo, T., Shigamardonova, E., Alexeev, D., Govorun, V., Wisecaver, J., Miyakey, A., Koyanagi, R., Fujie, M., Nishiyama, T., Shigenobu, S., Shibata, T. F., Golygina, V., Hasebe, M., Okuda, T., Satoh, N., and Kikawada, T. (2014) Comparative genome sequencing reveals genomic signature of extreme desiccation tolerance in the anhydrobiotic midge. Nat. Commun. 5, 4764.

18. Berjak, B. (2006) Unifying perspectives of some mechanisms basic to desiccation tolerance across life forms. Seed Sci. Res. 16, 1–15.

19. Carvalho, G. R., and Wolf, H. G. (1989) Resting eggs of lake- Daphnia I. Distribution, abundance and hatching of eggs collected from various depths in lake sediments. Freshwater Biol. 22, 459–470.

20. Hairston, N. G., Vanbrunt, R. A., Kearns, C. M., and Engstrom, D. R. (1995) Embryonic diapause and survivorship of diapasing eggs in a sediment egg bank. Ecology 76, 1706–1711.

21. Marcus, N. H., and Boero, F. (1998) Minireview: The importance of ben-thic-pelagic coupling and the forgotten role of life cycles in coastal systems. Limnology Oceanogr. 43, 763–768.

22. Kotani, T., Ozaki, M., Matsuoka, K., Snell, T. W., and Hagiwara, A. (2001) Reproductive isolation among genetically and temporarily isolated marine Brachionus strains. Hydrobiologia 446, 283–299.

23. Garcia-Roger, E. M., Camar, M. J., and Serra, M. (2006) Patterns in rotifer diapausing egg banks: Density and viability. J. Exp. Mar. Biol. Ecol. 336, 198–210.

24. Radzikowski, J. (2013) Resistance of dormant stages of planktonic invertebrates to adverse environmental conditions. J. Plankton Res. 35, 707–723.

25. Frisch, D., Morton, P. K., Chowdhury, P. R., Culver, B. W., Colbourne, J. K., Weider, L. J., and Jeyasingh, P. D. (2014) A millennial-scale chronicle of evolutionary responses to cultural eutrophication in Daphnia. Ecol. Lett. 17, 360–368.

26. Podrabsky, J. E., and Hand, S. C. (2015) Physiological strategies during animal diapause: lessons from brine shrimp and annual killifish. J. Exp. Biol. 218, 1897–1906.

27. Lopes, F. L., Desmarais, J. A., and Murphy, B. D. (2004) Embryonic diapause and its regulation. Reproduction 128, 669–678.

28. Ptak, G. E., Tacconi, C., Czernik, M., Toschi, P., Modliniski, J. A., and Loi, P. (2012) Embryonic diapause is conserved across mammals. PLoS ONE 7, e33027.

29. Fenelon, J. C., Banerjee, A., and Murphy, B. D. (2014) Embryonic diapause: development on hold. Int. J. Dev. Biol. 58, 163–174.

30. Fu, Z., Wang, B., Wang, S., Wu, W., Wang, Q., Chen, Y., Kong, S., Lu, J., Tong, Z., Ran, H., Tu, Z., He, B., Zhang, S., Chen, Q., Jin, W., Duan, E., Wang, H., Wang, Y. L., Li, L., Wang, F., Gao, S., and Wang, H. (2014) Integral proteome analysis of blastocysts reveals key molecular machinary governing embryonic diapause and reactivation for implantation in mice. Biol. Reprod. 90, 521–11.

31. Yamashita, O. (1996) Diapause Hormone of the Silkworm, Bombyx mori: Structure, Gene Expression and Function. J. Insect. Physiol. 42, 669–679.

32. Ye, H. L., Li, D. R., Yang, J. S., Chen, D. F., De Vos, S., Vuylsteke, M., Sorgeloos, P., Van Stappen, G., Bossier, P., Nagasawa, H., and Yang, J. W. (2017) Molecular characterization and functional analyses of a diapause hormone receptor-like gene in partenogenetic Artemia. Peptides (in press).

33. Wallace, R. L. (2002) Rotifers: Exquisite metazoan. Integ. Comp. Biol. 42, 660–667.

34. Wallace, R. L., Snell, T. R., Ricci, C., and Nogradi, T. (2006) Rotiferina. In: Segers, H., ed. Biology, Ecology and Systematics Guides to the Identification of the Microinvertebrates of the Continental Waters of the World 23 Backhuys Publishers, Leiden.

35. Walsh, E. J., Smith, H. A., and Wallace, R. L. (2014) Rotifers of temporary waters. Int. Rev. Hydrobiol. 99, 3–19.

36. Ricci, C. (2001) Dormancy patterns in rotifers. Hydrobiologia 446/447, 1–11.

37. Gilbert, J. J. (2004) Population density, sexual reproduction and diapause in monogonont rotifers: new data for Brachionus and a review. J. Limnol. 63, 32–36.

38. Gilbert, J. J. (2007) Timing of diapauses in monogonont rotifers: mechanisms and strategies. In: Alekseev, V. R., De Stasio, B., Gilbert J. J., eds.
Diapause in aquatic invertebrates: Theory and human use. Monographs in Biological Sciences, 84, pp. 11–27. Springer, Dordrecht, The Netherlands

Gilbert, J. J., and Schröder, T. (2004) Rotifiers from diapausing, fertilized eggs: unique features and emergence. Limnology Oceanogr. 49, 1341–1354

Gilbert, J. J., and Schröder, T. (2007) Intracranial variation in propensity for mixis in several rotifers: variation among females and maternal age. Hydrobiologia 593, 121–128

Schröder, T. (2005) Diapause in monogonont rotifers. Hydrobiologia 564, 291–306

Denekamp, N. Y., Suga, T., Hagiwara, A., Reinhardt, R., and Lubzens, E. (2010) A role for molecular studies in unveiling the pathways for formation of rotifer resting eggs and their survival during dormancy. In: Lubzens, E., Cerda, F., and Reinhardt, R. eds., Topics in Current Genetics 21, pp. 91–109. Springer, Heidelberg, Dordrecht, London, NY

Stout, E. P., La Clair, J. J., Snell, T. W., Shearer, T. L., and Kubanek, J. (2011) Conservation of progesterone hormone function in invertebrate reproduction. Proc. Natl. Acad. Sci. 107, 11859–11864

Kim, H.-J., Suga, K., Kim, B.-M., Rhee, J.-S., Lee, J.-S., and Hagiwara, A. (2015) Light-dependent transcriptional events during resting egg hatching of the rotifer Brachionus manjavacas. Marine Genomics 20, 25–31

Kim, R. O., Rhee, J. S., Won, E. J., Lee, K. W., Kang, C. M., Lee, Y. M., and Lee, J. S. (2011) Ultraviolet B retards growth, induces oxidative stress, and modulates DNA repair-related gene and heat shock protein gene expression in the monogonont rotifer, Brachionus sp. Aquat. Toxicol. 101, 529–539

Lee, B. Y., Kim, H. S., Hwang, D. S., Won, E. J., Choi, B. S., Choi, I. Y., Park, H. G., Rhee J. S., Lee, J. S. (2015) Whole transcriptome analysis of the monogonont rotifer Brachionus koreanus provides molecular resources for developing biomarkers of carbohydrate metabolism. Comp. Biochem. Physiol. D Genomics Proteomics 14, 33–41

Lee, J. S., Kim, R.-O., Rhee, J.-S., Han, J., Hwang, D.-S., Choi, B.-S., Lee, C.-J., Yoon, Y.-D., Lim, J.-S., Lee, Y.-M., Park, G.-S., Hagiwara, A., and Choi, I.-Y. (2011) Sequence analysis of genomic DNA (680Mb) by GS-FLX-Titanium sequencer in the monogonont rotifer, Brachionus plicatilis. BMC Genomics 12, 323–327

Burnell, A. M., Houthoofd, K., O’Hanlon, K., and Vanfleteren, J. R. (2005) Expression profiles of miRNAs and involvement of miR-100 and miR-34 in regulation of cell cycle arrest in Artemia. Biochem. Cell Biol. 287, 1392–13934

Zhao, L. L., Jin, F., Ye, X., Zhu, L., Yang, J. S., and Yang, W. J. (2015) Expression profiles of miRNAs and involvement of miR-100 and miR-34 in regulation of cell cycle arrest in Artemia. Biochem. Cell Biol. 470, 223–231

Molecular & Cellular Proteomics 16.10

Baker, D. A., and Russell, S. (2009) Gene expression during Drosophila melanogaster egg development before and after reproductive diapause. BMC Genomics 10, 242

Kankare, M., Salminen, T., Laiho, A., Vesala, L., and Heikkala, A. (2010) Changes in gene expression linked with adult reproductive diapause in a northern melt fly species: a candidate gene microarray study. BMC Ecol. 10, 3

Rinehart, J. P., Robich, R. M., and Denlinger, D. L. (2010) Isolation of diapause-regulated genes from the flesh fly, Sarcophaga crassipalpis by suppressive subtractive hybridization. J. Insect. Physiol. 56, 603–609

Yocum, G. D., Rinehart, J. P., Chirumamilla-Chapara, A., and Larson, M. L. (2009) Characterization of gene expression patterns during the initiation and maintenance phases of diapause in the Colorado potato beetle, Leptinotarsa decemlineata. J. Insect. Physiol. 55, 32–39

Clark, M. S., Thorne, M. A., Purać, J., Burns, G., Hillyard, Y., and Popović, Z. D., Grubor-Lajsić, G., and Worland, M. R. (2009) Surviving the cold: molecular analyses of insect cryoprotective dehydration in the Arctic springtail Megaphorura arctica (Tullberg). BMC Genomics 10, 328

Burnell, A. M., Thorne, M. A., Gerontoulis, G., Clark, M. S., Conway, P., and Worland, M. R. (2010) Gene expression associated with changes in cold tolerance levels of the Antarctic springtail, Cryptopygus antarcticus. Insect Mol. Biol. 19, 113–120

Poupadron, R., Schöttner, K., Korbelová, J., Prozvanková, J., Doležel, D., Pavlín, D., Beneš, V., and Koštál, V. (2015) Early transcriptional events linked to induction of diapause revealed by RNAseq in larvae of drosophilid fly, Chymomyza costata. BMC Genomics 16, 720

Tu, T., Wang, J., Hua, C., Whitman, D. W., Fan, Y., Cao, G., and Zhang, Z. (2015) Transcriptomic and proteomic analysis of pre-diapause and non-diapause eggs of migratory locust, Locusta migratoria. Pl, Ecol. 3, 223–231

Hansson, S. J., Stelzer, C. P., Welch, D. B., and Logsdon, J. M. (2013) Comparative transcriptomic analysis of obligately asexual and cyclically sexual rotifers reveals genes with putative functions in sexual reproduction, dormancy, and asexual egg production. BMC Genomics 14, 412

Griffin, T. J., Gygi, S. P., Ideker, T., Rist, B., Eng, J., Hood, L., and Aebersold, R. (2002) Complementary profiling of gene expression at the transcriptome and proteome levels in Saccharomyces cerevisiae. Mol. Cell Proteomics 1, 323–333

Ghazalpour, A., Bennett, B., Petyuk, V. A., Orozco, L., Hagopian, R., Mungri, I. N., Farber, C. R., Sinhannem, J., Kang, H. M., Furlotte, N., Park, C. C., Wen, P. Z., Brewer, H., Weitz, K., Camp, D. G., Pan, C., Yordanova, R., Neuhaus, I., Tilford, C., Siemers, N., Gargalovic, P., Eskin, E., Kirchgesner, T., Smith, D. J., Smith, R. D., and Luís, A. J. (2011) Comparative analysis of proteome and transcriptome variation in mouse. PLoS Genet. 7, e1001393

Clegg, J. S. (2001) Cryptobiosis - a peculiar state of biological organization. Comp. Biochem. Physiol. B 128, 613–624

Water, A. H., MacRae, T. H., and Bagshaw, J. C. (1989) Cell and molecular biology of Artelia development. Plenum Press, New York and London, 453pp

Abatzopoulos Th., J., Beardmore, J. A., Clegg, J. S., and Sorge los, P. (2002) Artemia: Basic and applied biology. Springer Science + Business Media Dodrecht, 286pp
Proteomes of Dormant and Nondormant Embryos

77. Wang, W., Meng, B., Chen, W., Ge, X., Liu, S., and Yu, J. (2007) A proteome study on postdiapause embryonic development of brine shrimp (Artemia franciscana). Proteomics 7, 3580–3591

78. Zhou, Q., Wu, C., D. B., Liu, F., and Xiang, J. (2008) The encoded dormant embryo proteome of Artemia sinica. Mar. Biotechnol. 10, 438–446

79. Fan, L., Lin, J., Zhong, Y., and Liu, J. (2013) Shotgun proteomic analysis on the diapause and nondiapause eggs of domesticated silkworm Bombyx mori. PLoS ONE 8, e60396

80. Cox, J., Heintz, M. Y., Lubber, C. A., Parson, I., Nagaraj, N., and Mann, M. (2014) Accurate proteome-wide label-free quantification by delayed normalization and maximal peptide ratio extraction, termed MaxLFQ. Mol. Cell Proteomics 13, 2513–2526

81. Fan, L., Lin, J., Zhong, Y., and Liu, J. (2013) Shotgun proteomic analysis on the diapause and nondiapause eggs of domesticated silkworm Bombyx mori. PLoS ONE 8, e60396

82. Ulitsky, I., Maron-Katz, A., Sagir, D., Linnar, C., Elkon, R., Tanay, A., Sharan, R., Shilo, Y., and Shamir, R. (2010) Expander: from expression microarrays to networks and functions. Nat. Prot. 5, 303–322

83. Giavalisco, P., Kohl, K., Hummel, J., Sei, W., and Willmitzer, L. (2008) Protein 2) from bovine liver. Biochim. Biophys. Acta 1783, 123–125

84. Moshkin, Y. M., Armstrong, J. A., Maeda, R. K., Tamkun, J. W., Verrijzer, C., Zhu, X., Liu, Y., and Yan, P. (2016) Radial Spokes—A Snapshot of the Westerman, J., Wirtz, K. W., Westerman, J., Wirtz, K. W., Berkhout, T., van den Dungen, J. L., and Karch, F. (2002) Histone chaperone ASF1 participates in light signaling through association with phytochrome in Drosophila melanogaster. Genes Dev. 16, 2621–2626

85. Fabrizio, P., Laggerbauer, B., Lauber, J., Lane, W. S., Lührmann, R., Lahr, M. D., and Bonning, B. C. (2000) A novel protein that binds juvenile hormone esterase in fat body tissue and pericardial cells of the tobacco hornworm Manduca sexta. J. Biol. Chem. 275, 1802–1806

86. Husson, S. J., Cluymen, E., Baggerman, G., Janssen, T., and Schoofs, L. (2005) Defective processing of neuropeptide precursors in Caenorhabditis elegans lacking proprotein convertase 2 (KPC-2/EGL-3): mutant phenotypes in the diapausing eggs of domesticated silkworm Bombyx mori. PLoS ONE 5, e89

87. Ma, D. A., Mikkat, S., Koy, C., Ringel, B., Thiesen, H. J., and Glocker, M. O. (2010) A novel class of evolutionarily conserved genes defined by piwi are essential for stem cell self-renewal. Genes Dev. 14, 3715–3727

88. Sim, C., and Denlinger, D. L. (2013) Insulin signaling and the regulation of insect diapause. Front. Physiol. 4, 189–1–10

89. Wurdak, E. S., Gilbert, J. J., and Jagels, R. (1978) Fine structure of the protein 2) from bovine liver. Biochim. Biophys. Acta 1783, 123–125

90. Li, M., Li, S., Yuan, J., Wang, Z. B., Sun, S. C., Schatten, H., and Sun, Q. Y. (2004) Molluscs. Cell Biol. Cell 10, 1794–1805

91. Cox, J., Neuhauer, N., Michalski, A., Scheltema, R. A., Olsen, J. V., and Mann, M. (2011) Andromeda – a peptide search engine integrated into MaxLFQ. Mol. Cell Proteomics 10, 1794–1805

92. Li, M., Li, S., Yuan, J., Wang, Z. B., Sun, S. C., Schatten, H., and Sun, Q. Y. (2004) Molluscs. Cell Biol. Cell 10, 1794–1805

93. Wurtz, K. W., Westerman, J., Wirtz, K. W., Berkhout, T., van den Dungen, J. L., and Karch, F. (2002) Histone chaperone ASF1 participates in light signaling through association with phytochrome in Drosophila melanogaster. Genes Dev. 16, 2621–2626

94. Shankmugavelu, M., Baytan, A. R., Chesnut, J. D., and Bonning, B. C. (2000) A novel protein that binds juvenile hormone esterase in fat body tissue and pericardial cells of the tobacco hornworm Manduca sexta. J. Biol. Chem. 275, 1802–1806

95. Fabrizio, P., Laggerbauer, B., Lauber, J., Lane, W. S., Lührmann, R., Lahr, M. D., and Bonning, B. C. (2000) A novel protein that binds juvenile hormone esterase in fat body tissue and pericardial cells of the tobacco hornworm Manduca sexta. J. Biol. Chem. 275, 1802–1806

96. Attramadal, A., Mikkat, S., Koy, C., Ringel, B., Thiesen, H. J., and Glocker, M. O. (2010) A novel class of evolutionarily conserved genes defined by piwi are essential for stem cell self-renewal. Genes Dev. 14, 3715–3727

97. Sim, C., and Denlinger, D. L. (2013) Insulin signaling and the regulation of insect diapause. Front. Physiol. 4, 189–1–10

98. Wurtz, K. W., Westerman, J., Wirtz, K. W., Berkhout, T., van den Dungen, J. L., and Karch, F. (2002) Histone chaperone ASF1 participates in light signaling through association with phytochrome in Drosophila melanogaster. Genes Dev. 16, 2621–2626

99. Cox, D. N., Chao, A., Baker, J., Chang, L., Qiao, D., and Lin, H. (1998) A novel class of evolutionarily conserved genes defined by piwi are essential for stem cell self-renewal. Genes Dev. 12, 3715–3727

100. Wilkerson, J., Rossman, K. L., and Der, C. J. (2005) The Ras superfamily: A guide to appropriate use of the Ras superfamily in molecular biology. Mol. Cell Proteomics 4, 281–315

101. Carey, H. V., Andrews, M. T., and Marin, S. L. (2003) Mammalian hibernation: cellular and molecular responses. Physiol. Rev. 83, 1153–1181

102. Hay, N. (2011) Interplay between FOXO, TOR, and Akt. Curr. Top. Dev. Biol. 95, 16–23

103. van der Goot, F. J., and Rang, R. D. (2003) SAGE surveys C. elegans carbohydrate metabolism: evidence for an anaerobic shift in the long-lived dauer larva. Mech. Ageing Dev. 124, 281–315

104. van der Goot, F. J., and Rang, R. D. (2003) SAGE surveys C. elegans carbohydrate metabolism: evidence for an anaerobic shift in the long-lived dauer larva. Mech. Ageing Dev. 124, 281–315

105. Identi-
122. Gallardo, W. G., Hagiwara, A., and Snell, T. W. (2000) Effect of juvenile hormone and serotonin (5-HT) on mixis induction of the rotifer Brachionus plicatilis Muller. J. Exp. Mar. Biol. Ecol. 252, 97–107
123. Wolschin, F., and Gadau, J. (2009) Deciphering proteomic signatures of early diapause in Nasonia. PLoS ONE 4, e6394
124. Zhang, Q., Lu, Y. X., and Xu, W. H. (2013) Proteomic and metabolomic profiles of larval hemolymph associated with diapause in the cotton bollworm, Helicoverpa armigera. BMC Genomics 14, 751
125. Tarrant, A. M., Baumgartner, M. F., Verslycke, T., and Johnson, C. L. (2008) Differential gene expression in diapausing and active Calanus finmarchicus (Copepoda). Mar. Ecol. Prog. Ser. 355, 193–207
126. Larade, K., and Storey, K. B. (2004) Accumulation and translation of ferritin heavy chain transcripts following anoxia exposure in a marine invertebrate. J. Exp. Biol 207, 1353–1360
127. Chen, T., Villeneuve, T. S., Garant, K. A., Amons, R., and MacRae, T. H. (2007) Functional characterization of artemin, a ferritin homolog synthesized in Artemia embryos during encystment and diapause. FEBS J. 274, 1093–1101
128. Bottke, W. (1982) Isolation and properties of vitellogenic ferritin from snails. J. Cell Sci. 58, 225–240
129. Tunnaciffe, A., Hincha, D. K., Leprince, O., and Macherel, D. (2010) LEA proteins: versatility of form and function. In: Lubzens, E., Cerdá, J., Clark, M. S. eds., Topics in Current Genetics 21, pp. 91–108. Dormancy and Resistance in Harsh Environments, Springer, Heidelberg, Dor- drecht, London, NY
130. Kanagasabapathi, V., and Munuswamy, N. (2011) Preservation, development, and hatching of resting eggs in the freshwater rotifer Brachionus calyciflorus Pallas. Microsc. Res. Tech. 74, 744–748
131. Gilbert, J. J. (2001) Spine development in Brachionus quadridentatus from an Australian billabong" genetic variation and induction by Asplanchna. Hydrobiologia 446/447, 19–28
132. Garcia-Roger, E. M., Carmona, M. J., and Serra, M. (2006) Hatching and viability of rotifer diapausing eggs collected from pond sediments. Freshwater Biol. 51, 1351–1358
133. Ricci, C., Melone, G., Santo, N., and Caprioli, M. (2003) Morphological response of a bdelloid rotifer to desiccation. J. Morphol. 257, 246–253
134. Boothby, T. C., Tapia, H., Brozena, A. H., Piszkiewicz, S., Smith, A. E., Giovannini, I., Rebecchi, L., Pielak, G. J., Koshland, D., and Goldstein, B. (2017) Tardigrades Use Intrinsically Disordered Proteins to Survive Desiccation. Mol. Cell 65, 975–984.e5