Title: Ionizing radiation responses are incidental to desiccation responses in the bdelloid rotifer Adineta vaga

V.C. Moris**1,2, L. Bruneau1, J. Berthe1, A-C. Heuskin3, S. Penninckx4, S. Ritter5, U. Weber5, M. Durante5,6, E. G. J. Danchin7, B. Hespeels*1, K. Van Doninck*1,2

Affiliations
1 Laboratory of Evolutionary Genetics and Ecology (LEGE), Department of Biology - URBE University of Namur, Rue de Bruxelles, 61, B-5000 Namur, Belgium
2 Laboratory of Molecular Biology & Evolution (MBE), Department of Biology, Université Libre de Bruxelles, 1000 Brussels, Belgium
3 Namur Research Institute for Life Sciences (NARILIS), Laboratory of Analysis by Nuclear Reactions (LARN), University of Namur, Rue de Bruxelles, 61, B-5000 Namur, Belgium
4 Medical Physics Department, Institut Jules Bordet – Université Libre de Bruxelles, 90 rue Meylemeersch, 1070 Brussels, Belgium
5 Biophysics Department, GSI Helmholtzzentrum für Schwerionenforschung, Darmstadt, Germany
6 Institute of Condensed Matter Physics, Technische Universität Darmstadt, Darmstadt, Germany
7 Institut Sophia Agrobiotech, INRAE, Université Côte d’Azur, CNRS, 06903, Sophia Antipolis, France.
* equally co-last authors
** corresponding author: victoria.carla.moris@gmail.com

ORCID IDs
Victoria C. Moris: 0000-0002-7454-0400
Jérémy Berthe: 0000-0002-7061-2083
Sébastien Penninckx: 0000-0002-3016-778X
Anne-Catherine Heuskin: 0000-0002-4960-5284
M. Durante: 0000-0002-4615-553X
Boris Hespeels: 0000-0003-2678-4352
Karine Van Doninck: 0000-0002-4790-8951
Abstract

Background: The remarkable resistance to ionizing radiation found in anhydrobiotic organisms, such as some bacteria, tardigrades, and bdelloid rotifers has been hypothesized to be incidental to the desiccation resistance. Both stresses produce reactive oxygen species and cause damage to DNA and other macromolecules, including DNA. However, this hypothesis has only been investigated in a few species and it is so far not possible to determine how universal it is.

Results: In this study, we analyzed the transcriptomic response of the bdelloid rotifer Adineta vaga to desiccation and to low- (X-rays) and high- (Fe) LET radiation in order to highlight the molecular and genetic mechanisms triggered by both stresses. We detected a transcriptomic response common to desiccation and both radiations with the over-expression of highly expressed genes mainly involved in DNA repair, protein modifications, or coding for Heat Shock Proteins, histones variants, enhancers but also many genes with unknown functions. We also discovered a distinct transcriptomic response specific to rehydration, which involved the over-expression of genes encoding Late Embryogenesis Abundant proteins, Heat Shock Proteins, and glucose repressive proteins. Moreover, we identified numerous genes encoding antioxidants that are constitutively highly expressed, which may contribute to the bdelloid rotifer resistance.

Conclusions: These results suggest that the extreme resistance of bdelloid rotifers to radiation might indeed be inherited from and a consequence of their capacity to resist complete desiccation. This study paves the way for functional experiments targeting promising candidate proteins playing central roles in radiation and desiccation resistance.

Keywords: desiccation, radiation, antioxidants, DNA repair, bdelloid rotifers
Background

Living in extreme environments requires effective mechanisms of cellular and molecular protection and of repair. One of the most severe abiotic stresses for cells is desiccation. Yet, some organisms, such as some bacteria, tardigrades, some nematodes, some insect larvae, and bdelloid rotifers, are capable of anhydrobiosis at any stage of their life-cycle and survive extreme water loss entering a dormant or ametabolic state (Ricci & Pagani, 1997; Rebecchi et al., 2006; Gusev et al., 2014). Anhydrobiotic organisms were also shown to be able to survive extreme doses of ionizing radiation (IR). According to Mattimore and Battista (1996), resistance to high doses of IR is incidental to the desiccation stress resistance since both stresses are known to cause similar damages (Mattimore and Battista 1996; Gladyshev and Meselson 2008; Gladyshev and Arkhipova 2010; Hespeels et al., 2014). However, this hypothesis has been supported by only limited data from a few species such as the bacterium Deinococcus radiodurans (proteomics: Ujaoney et al., 2017), larvae of the anhydrobiotic insect Polypedilum vanderplanki (qPCR, transcriptomics: Ryabova et al., 2017) and in the tardigrade species Richtersius cf. coronifer (qPCR of Hsp70: Jönsson and Schill, 2007). In this study, we used the bdelloid rotifer Adineta vaga to shed light on the genes and biological processes activated by both stresses, desiccation and IR.

Since their discovery (van Leeuwenhoek 1702), bdelloid rotifers have intrigued scientists because of their capacity to survive complete desiccation at any stage of their life-cycle and their resistance to a variety of other stresses including freezing, exposure to deep vacuum or high doses of IR (i.e., >500 Gy) (Ricci and Caprioli 2005; Gladyshev and Meselson 2008; Fischer et al. 2013; Jönsson and Wojcik 2017; Ricci 2017; Hespeels et al. 2020). Moreover, these eutelic (i.e., a fixed number of somatic cells at maturity) metazoans are among the smallest animals on Earth, being notorious for their parthenogenetic mode of reproduction in the absence of males (Hsu 1956; Birky 2010. Terwagne et al 2022). Their ability to re-start a new population from one individual and to survive complete desiccation or freezing, enables these organisms to inhabit environments where few animals can thrive, including semi-terrestrial habitats subjected to frequent episodes of drought (Fontaneto and Melone, 2003) such as mosses, lichens, rain gutters, but also soil, deserts and the arctic (Ricci, 1987; Fontaneto and Melone, 2003; Fontaneto et al., 2005; Robeson et al. 2009; Iakovenko et al. 2015; Shmakova et al., 2021). Among the highly radiation and desiccation-resistant bdelloid rotifer species, Adineta vaga has been the most extensively studied (Krisko et al., 2012; Flot et al., 2013; Hespeels et al. 2014, Hespeels et al. 2015, Hecox-Lea and Mark Welch, 2018; Hespeels 2020; Terwagne et al., 2022) and its genome has been recently re-sequenced and assembled at a chromosome-scale resolution level (Simion et al., 2021).

Both desiccation and IR are known to be associated with cellular membrane damage and the generation of Reactive Oxygen Species (ROS), inducing lipid peroxidation and DNA and protein
damage (Potts 1994; Franca 2007, Daly 2012). Survival to desiccation and IR therefore requires adjustments that maintain the function of macromolecules (i.e., proteins and DNA) despite the production of ROS. This can be achieved through two nonexclusive ways: either by preserving the integrity of these molecules or by repairing the damage accumulated. The loss of DNA integrity through DNA double strand breaks (DSBs) was shown to be repaired in A. vaga within 24-48 hours after desiccation or IR (Hespeels et al. 2014; 2020; Terwagne et al 2022). Moreover, previous studies highlighted an efficient antioxidant system in A. vaga protecting cellular constituents from oxidative damage (Krisko et al. 2012; Flot et al. 2013), and a wide expansion of gene families involved in oxidative stress resistance and DNA repair (Krisko et al. 2012; Flot et al. 2013; Hespeels et al. 2014; Hecox Lea and Mark Welch 2018; Simion et al. 2021). Genes coding for proteins involved in the six main DNA repair pathways conserved for the most part from prokaryotes to eukaryotes (Sancar et al. 2004; Hakem 2008; Pearl et al. 2015; White and Allers 2018) have been found in the A. vaga genome and transcriptome: Homologous Recombination (HR) and Non-Homologous End-Joining (NHEJ) repairing DSBs, Base Excision Repair (BER), Nucleotide Excision Repair (NER), Mismatch Repair (MR), and Alternative Excision Repair (AER) known to correct damaged or mismatched bases (Flot et al. 2013, Hecox Lea and Mark Welch 2018). While DNA repair pathways have been studied in A. vaga during and post desiccation (Hecox Lea and Mark Welch 2018), their contribution to repair DSBs and damages due to high doses of radiation remains to be characterized. Moreover, some proteins highly expressed in A. vaga following exposure to IR seem to have been acquired horizontally (Nicolas et al., unpublised). The genome of the bdelloid rotifers indeed contains the highest proportion of genes of non-metazoan origin within the studied animal genomes (Flot et al. 2013; Nowell et al. 2018; Simion et al 2021), many being expressed following desiccation and contributing to the metabolism of bdelloid rotifers (Boschetti et al. 2011).

Even though some damages and responses, including oxidative stress resistance and DNA repair, are hypothesized to be shared between desiccation and radiation, others might be specific to each stress factor. Indeed, anhydrobiosis, is a complex phenomenon that requires adaptations at the molecular and cellular level in order to face severe constraints associated with the loss of water (i.e., <10% of dried mass) and causing the complete cessation of any biological activity (Billi and Potts 2002; Alpert 2005; Hespeels et al. 2014). Desiccation in bdelloid rotifers also induces a change of their whole body which adopts a “tun” shape, via muscle contraction, allowing an optimal resistance to stress (Marotta et al. 2010). Desiccation resistance has been associated, in plants, bacteria, fungi, extremophilic bacteria and some tardigrades, with the accumulation of bioprotectants like non-reducing disaccharides (e.g., trehalose) that vitrify the cellular matrix (Rebecchi et al., 2007; Hengherr et al., 2008). Since trehalose has not been found in bdelloid rotifers while the trehalase gene was detected (Lapinski and Tunnacliffe, 2003; Hespeels et al. 2015), it was suggested that bdelloid rotifers might protect themselves against desiccation or osmotic stress by promoting the synthesis of proteins like Late Embryogenesis Abundant (LEA) proteins or Heat Shock Proteins (HSP), also found in plants and tardigrades, believed to act as...
molecular shields (Lapinski and McGee 2005; Reuner et al., 2010; Hand et al., 2011; Tripathi et al. 2012).

IR resistance might also involve distinct, additional responses which can differ according to the type of radiation being distinguished by their Linear Energy Transfer (LET). LET is the average energy (in keV) delivered locally to the traversed matter by a charged particle over a distance of 1 μm (Joiner and van der Kogel 2018; Hagiwara et al. 2019). Different types of IR are known to induce distinct ionization patterns. Sparse ionization (low-LET such as X-rays) produces a uniform pattern of damages throughout its target while dense ionization (high-LET, e.g., protons, Fe-ions) generates complex DNA damages including clustered DSBs (Lehnert 2007; Semenenko and Stewart 2004; Penninckx et al., 2019; Penninckx et al., 2021). As a consequence of these complex lesions difficult to repair, cell killing is higher after exposure to the same dose of high-compare to low-LET radiation. The maximum effectiveness for cell killing is between 100 and 200 keV/micron in water (Friedrich et al., 2013). Iron (Fe) ions at 1 GeV/n have a LET in water of 147 Kev/micron, and represent therefore a highly effective radiation quality (Durante & Cucinotta, 2011). Accordingly, the survival and fertility rates were observed to be significantly reduced in A. vaga individuals exposed to a given dose of protons and Fe-ions in comparison with X-rays (Hespeels et al. 2020). Only a few recent studies started to analyze biological processes and genes involved in responses to X-rays and charged particles (Cekanaviciute et al., 2023). However, the role of these processes and genes in bdelloid rotifer resistance and how they differ between different types of radiation has never been investigated.

Identifying these genes and processes involved in the desiccation and IR resistance of the bdelloid rotifer A. vaga allows us to determine the main actors of their extreme resistance and whether they are common to both stresses (Fig 1A). We therefore analyzed here the transcriptomic response of A. vaga individuals exposed to desiccation and low- and high-LET radiation (respectively X-ray and Fe-ions). Our analysis allowed identifying the desiccation, post-rehydration and post-radiation core responses and highlight numerous candidate genes. Some of these genes are likely to be responsible for resistance to both stresses and could be targeted in future functional studies.

Results

1. Fertility of A. vaga decreases after exposure to high-LET radiation

We observed that longer desiccation periods (i.e., 14 days) in absence of radiation were not associated with a decrease in survival and fertility in comparison to 1-day desiccated samples (Fig S1). Radiation also did not impact the survival rates of desiccated and irradiated individuals compared to non-irradiated controls (X-rays: 96±3%, vs. 96±3%; Fe: 92±2% vs. 91±3%; Fig S1). However, the fertility rate after 500 Gy Fe-ions (9±5%) was lower than after 500 Gy X-ray radiation (92±7%). Indeed, 72±17% of 500 Gy Fe-ions irradiated samples were characterized by the production of sterile eggs (Fig 1B in blue), unable to complete full embryological development.
2. **Transcriptomic response in A. vaga are highest 2.5 hours post-radiation and 1.5 hours post rehydration**

With the Differential Gene Expression (DGE) analysis of A. vaga individuals under different treatments, including “entering desiccation” (in grey), 14 days desiccation (yellow), 500 Gy low-LET (orange or blue) and high-LET radiation (purple) ([Fig 2A](#)), we observed that the highest variance in gene expression (PC1=56%) correlates with the radiation treatment (irradiated vs. non-irradiated) and the time post radiation ([Fig 2B](#)). The maximal distance observed on PCA, is between control samples and the timepoint 2.5h after low- and high-LET radiation ([Fig 2B](#)). At this timepoint, we also measured a high percentage of genes under-expressed (UE) (in red, 6.6-13.4%) and over-expressed (OE) (in light green, 6.1-6.6%) with a percentage even slightly higher at 8h following Fe-ions irradiation (8.3%) ([Fig 2C](#)). PC2, encompassing 16% of the variance, correlates with the desiccation treatment (desiccated vs. hydrated) ([Fig 2B; Supplementary results](#)). Samples rehydrated for 1.5 hours (h) after 14 days of desiccation show the highest number of UE and OE genes ([Fig 2C](#)). We also observed that rotifers entering desiccation or post-desiccation (1.5h or 2.5h post desiccation/radiation) had more UE genes than OE genes, while it was the opposite 2.5h post radiation without previous desiccation ([Fig 2C](#)). At 24h post-radiation, the difference between control and irradiated samples is the lowest ([Fig 2B, pentagons](#)) with a lower percentage of genes being differentially expressed (DE) ([Fig 2C](#)). We noticed that three hydrated controls ([Fig 2B, yellow-green](#)) had a positive PC2 and were more distant to the other controls. These rotifers were sampled with slightly different conditions (see Material section). The replicates from the experiments with hydrated rotifers irradiated with X-rays were also grouped two by two ([Fig 2B, blue triangles, circles and squares](#)) which is likely due to the sampling for this treatment (see Material section).

3. **Biological processes associated to DE genes following desiccation and radiation**

In order to identify the biological processes triggered by desiccation and by IR, GO enrichment analyses of DE genes were performed for both desiccation and radiation separately ([Supplementary Table S1](#)). Only half of the DE genes have GO ids attributed and could be considered for the analysis ([Figs S3A, S3B, S3C, S3D](#)).

Some enriched biological processes ([Figs 3A, 3B](#)) and genes ([Fig 3C](#)) were found to be shared between A. vaga individuals entering desiccation (in grey) and individuals rehydrated after 14 days of desiccation (in yellow): 915 OE genes and 1885 UE genes in both conditions ([Fig 3C, Supplementary Table S1](#)). Amongst under-expressed biological processes found significantly enriched in both conditions were processes mainly involved in signaling pathways and transport while amongst OE genes they were mostly involved in oxidation-reduction processes, ADP-ribosylation and microtubule-based processes ([Figs 3A, 3B](#)). Under-expressed biological
processed being significantly enriched specifically to the entry in desiccation included mainly cell communication and transmembrane transport (Fig 3A) while carbohydrate and trehalose metabolic processes were enriched within OE genes (Fig 3B). Finally, we found significant enrichment specific to the rehydration timepoint following 14 days desiccation in UE genes involved in signaling, DNA replication and transcription, while protein modifications and signal transduction processes were enriched in OE genes (Figs 3A, 3B). Signaling was enriched within UE genes at the entry of desiccation and during rehydration, as well as within OE genes during rehydration suggesting some reactivation (Figs 3A,3 B).

A similar approach was applied to define the biological processes enriched in response to radiation. Similar biological processes were found to be enriched in UE genes as those found UE in desiccation treatment: signal transduction, cell communication, and ion transport (Supplementary Table S2, Fig S4). At 2.5h and 8h post-radiation, the timepoints where most genes are differentially expressed compared to the controls (Fig 2B, 2C), respectively 906 and 724 genes were over-expressed in the three radiation conditions (Fig 3D). After GO enrichment analysis of these genes, we measured a significant enrichment of biological processes involved in distinct DNA repair processes, protein modification processes, transcription and transport both at 2.5h and 8h post-radiation (Fig 3E). We also found a significant enrichment in oxidation-reduction processes 8h post-radiation, also detected following desiccation (Fig 3E).

4. Highly over-expressed genes following desiccation and radiation

Because half of the DE genes had no GO annotation, we used a second approach to identify the potential function of proteins encoded specifically by OE genes both after desiccation and radiation, using Pfam domains, Kegg/GO ids, and blast hits.

Selecting the timepoints with the most differential responses (more distant to the controls on the PCA; Fig 2B), including 1.5h rehydration after 14 days of desiccation, 2.5h post X-ray radiation with and without desiccation and 2.5h post-desiccation and Fe-ions irradiation, a total of 258 OE genes was found in common to both rehydration after desiccation and radiation (Fig 4A; Supplementary Table S3). In addition to these genes, we found 260 OE genes in all rehydration response (with or without radiation, Fig 4A), and 648 OE genes common to all radiation treatments (with or without previous desiccation, Fig 4A). Amongst those genes, we were unable to get any cues (blast hits or pfam of orthologs having known function) regarding the function of 58 genes in the common response (22.5%), 86 within the rehydration response (33%), and 178 within the radiation core response (27.5%). Below, we focused specifically on differentially expressed genes (log2fc>0.5) with high expression levels in the treatments (TPMs>150) to describe the key genes and processes involved in the resistances to desiccation and radiation, as presented in Fig 4B.

1. Key OE genes common to both rehydration after desiccation and radiation: we found 10 OE genes with high TPMs (>150) 2.5h post radiation (all types) and 1.5h post rehydration (without
radiation) amongst the 258 OE genes, of the common response to rehydration and radiation. These genes likely code for glucose-repressible gene-1 protein Grg1, and an endothelial differentiation-related factor, as well as proteins involved in ubiquitination (2 genes), transcription regulation (a CCAAT/enhancer, a transcription factor), transport, and 3 genes with unknown functions (Fig 4B; Supplementary Table S3). The gene coding for CCAAT/enhancer likely involved in transcription regulation is the gene with the highest TPM values (TPMs>1000, 2.5h post radiation (Fe-ions or X-rays) and 1.5h post 14 days desiccation; Supplementary Table S3). We decided to investigate in addition the 19 genes (l2fc>0.5 2.5h post radiation (all types) and 1.5h post rehydration (without radiation)), also reaching TPMs>150 2.5h post radiation (all types) but not post rehydration without radiation. Indeed, desiccation is known to produce less damages such as DNA damages compared to IR. Therefore, the potential genes of resistance even though being over-expressed, might not reach these levels of expression (TPMs>150) after desiccation only. The 19 additional genes were found to be likely involved in: DNA repair (2 PARP, Ligase E, 2 polymerases beta), an exoribonuclease, an aldehyde dehydrogenase (antioxidant), a 2OG-Fe(II) oxygenase, a protein phosphatase, and one more gene involved with ubiquitination, a major vault protein, and 8 genes with unknown function.

2. Key OE genes specific to rehydration post desiccation: we found 52 OE genes with high TPMs (>150) amongst the 260 OE genes of the rehydration response following desiccation (with or without radiation) coding for two heat shock proteins (HSPs), LEA, Grg1, a ribosomal protein, a lipase/acylhydrolase, a protease (trypsin pfam domain PF00089), proteins with a calcium binding domain, proteins involved in signaling and microtubules, as well as 29 genes with unknown functions (Fig 4B; Supplementary Table S4).

3. Key OE genes specific to the radiation response: we found 58 OE genes with high TPMs (>150) post radiation amongst the 648 genes of the radiation response, regardless of the radiation or desiccation treatment. These 58 genes are coding for proteins involved in protein modifications (ubiquitination, proteolysis, phosphorylation, homeostasis), in DNA repair/damage or synthesis, in transcription regulation (CCAAT/enhancer), in apoptosis inhibition, and in vesicle transport. We also found genes coding for HSPs, Major Vault proteins, histones and histone variants (two H2B variants, one H2A variant H2Abd2, one core histone H3, and one core histone macro- isoform X2), an RNA ligase, a nucleoredoxin, a beta lactamase, a hydrolase, a microsomal signal peptidase, von Willebrand factors and 15 genes with unknown functions (Fig 4B; Supplementary Table S5).

Based on the 8.3% genes of non-metazoan origin (Horizontal Gene Transfer : HGTs) detected in the genome of A. vaga (Simion et al., 2020), we found a significant enrichment in HGTs in the common response (37 HGTs amongst the 258 genes (14.3%); \(\chi^2=12.4, df=1, alpha=0.01\); Fig 4B and Supplementary Table S3) and in the rehydration response following desiccation (37 HGTs amongst 260 genes (14.3%); \(\chi^2=12.01, df=1, alpha=0.01\); Fig 4B and Supplementary Table S4).
Although we found 67 HGTs amongst the 648 genes (10.3%) in the radiation response, this did not correspond to a significant enrichment ($\chi^2=3.54$, df=1, alpha=0.01; Fig 4B and Supplementary Table S5). Amongst these genes, 2OG-Fe(II) oxygenase, Ligase E, two RNA ligases, and genes belonging to Endonuclease/Exonuclease phosphatase particularly retained our attention because of their high TPM values (TPMs>150). For instance, the ligase E (FUN_003353) (previously ligase K in Hecox Lea and Mark Welch 2018) is one of the genes found in the shared response to rehydration and radiation with the highest log2 foldchange (log2fc = 7.32, 2.5h post-desiccation and X-ray radiation; log2fc = 2.3, 1.5h post 14 days of desiccation) and highest expression levels in irradiated A. vaga rotifers (TPMs=862, 2.5h post desiccation and X-ray radiation; TPMs=1522, 2.5h post desiccation and Fe-ions irradiation; TPMs=23, 1.5h post 15 days of desiccation) compared to the controls (1.88<TPMs<7.83) (Supplementary Table S3).

5. **High constitutive expression of A. vaga antioxidant system enhanced by desiccation and rehydration processes.**

Since we observed a significant enrichment of oxidation-reduction processes amongst OE genes in A. vaga individuals entering desiccation, 1.5h post-rehydration after 14 days of desiccation, and 8h post-radiation with or without desiccation (Figs 3B, 3E), we annotated, based on Pfam domains and blast search, in total 217 genes coding for antioxidants in the entire A. vaga genome (Simion et al., 2020): 65 encoding glutathione S-transferases, 59 encoding aldo/keto reductases, 26 encoding thioredoxins, 18 encoding aldehyde dehydrogenases, 16 encoding peroxiredoxins, 11 encoding superoxide dismutases (SOD), 8 encoding isocitrate dehydrogenases, 5 encoding catalases, 5 encoding glutathione peroxidases, and 4 encoding glutathione reductases (Supplementary Table S6; Fig 5A).

We observed that many antioxidant genes were already highly expressed in the controls (TPMs>150; Fig 5A). Indeed, 26 genes are highly constitutively expressed in all conditions including controls (except 2 peroxiredoxins and 1 isocitrate dehydrogenase having TPM values slightly lower than 150 post-radiation): three aldehyde dehydrogenases (different than the ones which are OE), one aldo/keto reductase, one catalase, one glutathione peroxidase, five glutathione s-transferases, two isocitrate dehydrogenases, four peroxiredoxins, four SOD (2 Fe/Mn, 2 Cu/Zn) with 2 of them being also OE at some time points post-radiation treatments and five thioredoxins. Amongst these 26 genes, four show even higher levels of expression both in controls and desiccation/radiation treatments (TPMs>700): one superoxide dismutase (SOD Cu/Zn), two thioredoxins, one glutathione S-transferase (Fig 5A, 4C, Supplementary Table S6).

After stress, the number of highly expressed (TPMs>150) antioxidant genes did not increase (Fig 5A). Still, we found 69 genes being OE in at least one condition compared to the control but which did not reach high TPMs post stress (Fig 5B). In accordance with the observed significant enrichment of oxidation-reduction processes (Fig 3B), the entry in desiccation and the rehydration
after long periods of desiccation led to more OE antioxidant genes than radiation only (Fig 5B). Indeed, 1.5h post 14 days desiccated condition had the highest number of antioxidant OE genes, 48 in total (Supplementary Table S7), with 30 genes found to be OE exclusively in this condition (indicated as number on Fig 5B). Amongst the 30 genes found OE exclusively post rehydration without radiation, we notably found two other aldehyde dehydrogenases (FUN_007530, FUN_008548) and one catalase (FUN_019206) highly expressed in the controls (respectively 150>TPMs>200; 250>TPMs>300; 190<TPMs<270 Fig 5A, 5C) but reach even higher TPMs 1.5h post rehydration (respectively, TPMs=303; TPMs=400, TPMs=489; Supplementary Table S7). Within the remaining of the 48 genes, four genes were found both only in A. vaga individuals entering desiccation and in those being rehydrated for 1.5h after 14 days of desiccation, and 14 genes were found in common with at least one condition of radiation (all time points included; Fig 5B). Only two of these genes were found in common to all the different conditions 2.5h post stress: one aldehyde dehydrogenase (already mentioned in the common response), and one aldo/keto reductase (Supplementary results for more details).

We found a common response to rehydration with six antioxidant genes (Fig 4A): 4 coding for aldo/keto reductases, one for a catalase, and one for a SOD (Cu/Zn) (Fig 5C). However, only 2 antioxidant genes were found in the core radiation response (2.5h post radiation): another gene coding for an aldehyde dehydrogenase and a glutathione s-transferase. A SOD (Cu/Zn) was also found in the core radiation response but 8h post-radiation (Fig 5C).

This lack of common genes seems to be mainly due to the difference in the responses to the irradiation and desiccation treatments. Less antioxidant genes were OE in hydrated conditions. Indeed, only 7 antioxidant genes were OE 2.5h post X-ray radiation without pre-desiccation, versus 12 genes 2.5h post X-ray radiation and desiccation. One gene, though, a SOD Fe/Mn seems to be specific to irradiated hydrated conditions as it was OE 2.5h and 8h post X-rays, even if this gene is part of those being constitutively highly expressed in all conditions (Fig 5B, 5C). Moreover, Fe-ions radiation seems to trigger the OE of more antioxidant genes than X-rays: 27 versus 19 genes (in total all time points included), with nine additional genes: 5 glutathione s-transferases, 1 peroxiredoxin, 2 thioredoxins, 1 aldo/keto reductase (Fig 5B).

6. Genes involved in NHEJ and BER are highly expressed post-rehydration and post-radiation

Since we found a significant GO enrichment in DNA repair biological processes in the core response to radiation, we characterized genes coding for proteins involved in DNA repair within the genome of A. vaga. We identified 222 DNA repair genes in A. vaga based on KEGG ids, Pfam domains, and genes previously identified in Hecox-Lea et al., 2018, and grouped them by repair pathway: 23 genes in NHEJ, 29 genes in HR, 46 genes in BER, 49 genes in NER, 6 genes in MR, 3 genes in alternative-NHEJ, 19 genes in DNA replication, 26 genes in cross pathways, and 21 genes with functions related to DNA repair (Fig 6A; Supplementary Table S8).

Genes involved in NHEJ and BER are highly expressed post-rehydration and post-radiation
Unlike numerous genes encoding antioxidants that are constitutively expressed in the controls, only six genes involved in DNA repair have a high expression level (TPMs>150) in all controls: four genes coding for high mobility group proteins potentially involved in BER (Prasad et al., 2010), MSH2 involved in MR, and a phospholipase involved in related DNA repair processes (Fig 6A; 6C).

We found 17 DNA repair genes in the common response to rehydration and radiation and 30 additional genes in the core response to radiation (genes indicated in blue in Fig 3A; Fig 6B, Fig 7, Figs S5, S6, S7, S8, Supplementary Table S8). Most genes being OE 2.5h post-radiation belong to BER (21/53) and NHEJ (12/23) pathways, unlike genes from HR (6/40), NER (8/63), or MR (4/31), taking into account genes involved in cross-paths (Figs 6A, 6B, 7, Figs S5, S6, S7, S8). The few genes with high log.foldchange values in HR, NER and MR are those coding for proteins involved in cross-paths (e.g., PCNA; Ligase E), whereas genes coding for proteins exclusively involved in BER (e.g., Polymerases β, PARPs) and NHEJ (e.g., KU80; Artemis; DNA-PKcs; APLF; APTX) reach high log.foldchange values (Fig 6B, Fig 7, Figs S5, S6, S7, S8). Most of these genes reach high TPMs (>150), in particular, one gene coding for PARP reached TPM values >1000 2.5h post-radiation. We noticed that some actors in BER (e.g., glycosylase, APEX, PCNA and Ligase 3) and in NHEJ (e.g., KU70, KU80, Artemis, Polymerase λ) had only one copy over-expressed after radiation, whereas other actors had multiple copies over-expressed (e.g., XRCC4, APLF, PARP, Polymerase β, FEN1) (Figs 6B and 7).

Most of the genes found OE post rehydration without radiation were found OE post radiation, such as KU80, Artemis, Lig4, glycosylase, PARP, polβ, Ligase E (Fig 6B, 7) indicating similar DNA repair mechanisms for both stresses. Moreover, many genes involved in DNA repair which were OE post-radiation were slightly OE post rehydration without desiccation but were below the established thresholds for log2foldchange and FDR.

We observed for the different types of radiation, the same genes involved in DNA repair being OE, with similar log.foldchange values and with, for most of them, the highest values 2.5h post-radiation (Fig 6B, Figs S5, S6, S7, S8). Therefore, we represented TPMs only 2.5h post-desiccation and X-ray radiation in Figs 6A, and 6C, and 6D. The only difference we noticed was that some genes OE post X-rays radiation were significantly OE 8h post Fe-ions irradiation, and not 2.5h (e.g., TDP1, Ligase 1 in BER, Fig 6B; Ligase 4, Ligase D in NHEJ, Fig S5; XPC and CSB in NER, Fig S7).

**Discussion**

Transcriptomic analyses of the bdelloid rotifer A. vaga facing desiccation and IR allowed us to determine which genes and biological processes are likely involved in response to these stresses. Moreover, this allowed identifying genes commonly responsive to both stresses and some candidates that appeared specific to a peculiar response (summarized on Fig 8). We mainly
discuss below enriched biological processes, and overrepresented gene families with OE genes (log2fc>0.5, FDR<0.01) reaching high expression levels (TPMs>150) after IR or desiccation.

1. **Common transcriptomic response of **A. vaga** to rehydration and radiation

Both desiccation and radiation lead to similar challenges to face such as DNA/protein damages and ROS formation (*Mattimore and Battista 1996; Gladyshev and Meselson 2008; Gladyshev and Arkhipova 2010; Hespeels et al., 2014*). These damages can lead to compromised cellular function and even cell death, emphasizing the need for a swift and rapid expression of protection and repair mechanisms.

Unlike the radio-sensitive monogonont rotifer *Brachionus koreanus* in which the p53 gene, trigger of apoptosis in mammalian cells (*Lane, 1992; Vousden and Prives, 2009*), is OE post gamma radiation (150 and 200 Gy; *Han et al., 2014*), p53 seems was not found in *A. vaga* genome. Similarly, this gene seems to also be missing in the strong cryptobiont heterotardigrade *Echiniscoides* cf. *sigismundi* (*Kalimari et al., 2019*). The absence of p53 gene in *A. vaga* does not definitely rule out the potential presence of apoptosis. In addition, we found the over-expression of some genes suggested to be involved with the prevention of apoptosis such as Hsp70 (*Gibbons et al., 2000*). Because of these results and the fact that adult somatic cells in *A. vaga* are not engaged in mitosis (eutelic), we hypothesize that apoptosis is unlikely to occur the bdelloid rotifer exposed to radiation or desiccation. Instead of inducing programmed cell death when damages accumulate, regulated by p53, bdelloid rotifers seem to possess an arsenal of genes that we describe below required to prevent and repair the damages incurred.

The similar damages caused both by desiccation and IR likely explain why we found a common transcriptomic response in the bdelloid rotifer *A. vaga* (*Fig. 8* in pink) both post rehydration (1.5h) and radiation (2.5h) with the oxidoreduction and protein modification processes being enriched (*Figs 3B and 3E*) and 258 genes being commonly OE including 17 genes involved in DNA repair (*Figs 4A and 4B*). For most of the OE genes, the over-expression was at the highest early after stress exposure (timepoints 1.5h or 2.5h post rehydration or radiation), indicating a fast transcriptomic response from the bdelloid rotifer *A. vaga* upon exposure, probably contributing to its high resistance and resilience.

At protein level, cells can either scavenge ROS with antioxidants to prevent oxidative damage in general (*Gill and Tuteja, 2010*), or protect proteins with chaperones like HSPs (*Bukau et al., 2006; Kampinga and Craig, 2010*). In general, molecular chaperones play a crucial role in protein homeostasis (proteostasis) by regulating protein quality control, folding and turnover, preventing newly synthesized proteins to aggregate into non-functional structures (*Bukau et al., 2006; Stengel et al., 2010*). Two genes coding for antioxidants, one aldehyde dehydrogenase and one aldo/keto reductase, were also found in the common response (*Fig 5B*), while antioxidant genes appear constitutively expressed in *A. vaga* (*Fig. 5A, 4-5C, Fig 8* in green), suggesting the
presence of a general mechanism of oxidative stress prevention. Moreover, we found evidence of several HSP (e.g., HSP70) coding gene constitutively highly expressed (TPMs>150; Fig 8 in green; Supplementary Table 1) and some genes over-expressed in the common response (Fig 4B), although more genes coding for HSPs were identified in the rehydration post-desiccation response.

When antioxidant and chaperones systems fail to respectively protect the proteins or to refold a denatured protein, proteins can be modified. Many genes and enriched biological processes found OE after both stressed in A. vaga is protein modifications including protein involved in ubiquitination (Fig 3E; Fig 4B). These protein modifications can cause change of a protein activity/function (dephosphorylation) or mark targeted proteins for degradation (Kubota, 2009). Ubiquitination often mark misfold proteins for proteolysis involving the 26S proteasome, a large multiprotein complex (Römisch, 2005; Kubota, 2009; Ciechanover et al., 2000). Such proteins involved in protein ubiquitination were also found OE in other desiccation resistant species in response to rehydration (e.g., in tardigrade: Wang et al. 2014). Interestingly, ubiquitin-dependent signaling processes have also been shown to play a role in the choice of the double strand breaks (DSB) repair pathway (HR or NHEJ) (Schwertman et al., 2016) and in the re-establishment of genome integrity after DSBs (Jackson and Durocher, 2013).

While proteins appear to be protected from oxidative damage in A. vaga during radiation (see also Krisko et al., 2012), DNA is not protected and damages accumulate with prolonged desiccation and higher doses of radiation (Hespeels et al. 2014). DNA double strand breaks (DSBs), but also single strand break (SSBs), and base damages are known to be caused by direct or indirect damages following radiation (O’Neill et al., 1993; Wallace 1988; Lehnert, 2007; Semenenko and Stewart, 2004) and were indeed observed in A. vaga (Hespeels et al., 2014, 2020). The high radiation doses were chosen in this study in order to improve the detection actors involved in radiation resistance. However, DNA DSBs are also caused by prolonged desiccation in A. vaga but with less DSBs being detected on pulsed-field gel electrophoresis (PFGE) following desiccation than post-radiation doses reaching 500 Gy (Hespeels et al., 2014, 2020). Here, we found 17 genes to be involved with DNA repair in the common transcriptomic response (Fig 4A). The high radiation levels (500 Gy) inducing more DSBs than desiccation (Hespeels et al., 2020) probably explains why we found more genes in the core response to radiation (30 additional genes; Fig 4A; Fig 8 in blue) and why one of the most enriched biological processes post-radiations was DNA repair (Fig 3E). It is also probably the reason why many of these 30 additional genes were slightly OE post rehydration without radiation but with log2fc and FDR values close to the chosen thresholds (Supplementary Table S8). Finally, it also explains why not many genes have been found to be OE post desiccation in rotifers in other studies (Hecox Lea and Mark Welch, 2018).
Unlike antioxidant for which many genes are already highly expressed in the controls (see dedicated section), we found only six genes involved in DNA repair which are highly constitutively expressed (TPMs>150 in all conditions, including controls), notably four high mobility proteins (Fig. 5C; Fig 8 in green). High mobility proteins have a role in the chromatin structure, transcriptional regulation but also seem to enhance DNA repair (Lange et al., 2008). Indeed, the absence of High mobility group protein B1 (HMGB1) was shown to lead to increased mutagenesis, decreased cell survival, and altered chromatin reorganization after DNA damage (Lange et al., 2008). It was also shown the HMGB1 was involved with telomere homeostasis and prevent the DNA damage induced by radiation in human breast cancer cells (Ke et al., 2015). Perhaps in the same way as antioxidants being highly expressed also in the control conditions, the high constitutive expression of genes coding for high mobility proteins could be due to the numerous episodes of desiccation A. vaga might face in its natural environment leading to DNA damages and regulation of many genes involved in its desiccation resistance.

Intriguingly, we found that the DNA repair proteins encoded by genes OE post desiccation and/or radiation were mostly involved in BER and NHEJ DNA repair pathways. Some actors seem particularly important (as they reached particularly high expression levels) such as PARP and two polymerases beta in BER pathway (Fig 6B), APLF and APTX in NHEJ pathway (Fig 7), and DNA ligase E and PCNA involved in cross pathways (Fig 6B and Fig 7). Specifically, the PARP gene family may play a key role for the bdelloid rotifer resistance as we found a high number of over-expressed copies: 17 genes are OE 2.5h post all type of radiations and 9 genes are OE post-rehydration after 14 days of desiccation (Fig 6B). PARPs are involved in BER pathway, notably for recruiting other proteins that are critical in this pathway (de Murcia et al, 1994). It has already been shown that PARPs are important to deal with radiation DNA damages as their expression level increased in carcinogen-treated cells or in cells exposed to radiation or DNA damaging agents (Juarez-Salinas et al., 1979; Skidmore et al., 1979; Chalmers et al., 2004). Besides DNA repair, PARPs appear to be also involved in other cellular processes such as the modulation of chromatin structure, transcription, replication, recombination, regulation of membrane structures, cell viability, cell division, and actin cytoskeleton (Vyas et al., 2013), explaining perhaps their particular over-expression.

These results shed light on DNA repair pathways and main actors used to repair the damaged DNA in A. vaga. We found similar results than in Hecox Lea and Mark Welch (2018) where A. vaga individuals being desiccated for 7 days and rehydrated for 1h. The ligase E (previously named ligase K) was already found to be OE (l2fc= 1) in A. vaga individuals being desiccated for 7 days and rehydrated for 1h (Hecox Lea and Mark Welch, 2018). However, it was not possible at that time to determine if one pathway was more over-expressed than another to repair the damages. Indeed, they found to be OE 1h post rehydration genes coding for APLF (found in this study but 0.01<FDR<0.05 1.5h post-rehydration), polymerase lambda (also found OE in this study). However, they found BLM (HR) to be OE (not the case in our study), whereas we found
KU80 or Artemis to be OE 1.5h post-rehydration (Fig S5). With to the study of irradiated individuals with high IR doses causing more DNA damages than desiccation only, we were able to amplify the response and found that BER and NHEJ pathways are more likely to be used to repair DNA in A. vaga as most actors from these pathways were OE post radiation. The small differences between the two studies are likely due to the length of the desiccation (7 versus 14 days) and time of rehydration (1h version 1.5h). We hypothesize that more DNA damages might be cause with longer desiccation periods. Moreover, the l2fc values might have been bigger if we measured the gene expression levels of rehydrated rotifers at other time points (e.g., 2.5h, and 8h).

The genes found OE in bdelloid rotifers in this study associated with specific DNA repair pathways were different from the repair genes reported in other radiation and desiccation resistant species. In the bacterium D. radiodurans proteins involved in NER and HR were upregulated post-desiccation and post-radiation (Uljanow et al. 2017). In P. vanderplanki insect larvae proteins of HR (Rad23 and Rad51) were upregulated both after desiccation and high- and low-LET radiation (Gusev et al., 2010), while in tardigrades all major repair pathways appear expressed (Förster et al., 2012). This suggest that various strategies to repair DNA damages evolved in different desiccation resistant species.

DSB repair is known to require in many eukaryotes the conserved histone variant of canonical H2A, H2AX, which is absent from studied bdelloid rotifer species (Van Doninck et al., 2009). In response to DSBs, H2AX becomes phosphorylated in its C-terminus, triggering the retention and accumulation of repair and checkpoint proteins to the DNA breaks (Van Doninck et al., 2009). Amongst other H2A variants in the genome, some were OE after both stresses (e.g., H2Abd1; TPMs>100 2.5h post radiation; Supplementary Table S3; Fig 8). In addition to H2Abd1, we found an increased expression of other histones following radiation, such as two H2B variants, the other identified H2A variant (H2Abd2), one core histone H3, and one core histone macro-isoform X2 (TPMs>200, 2.5h post-radiation; Supplementary Tables S3 and S4). We do not know, so far, the exact role of these histone variants in the resistances of A. vaga. In general, core histones and its variants play a role in chromatin (de)compaction and in the maintenance, replication and expression of the genome, which is important when numerous genes are transcribed in response to desiccation and/or radiation stress (with over 1000 genes over-expressed post-stress). Consequently, it is logical that we also identified actors involved in DNA replication, RNA synthesis, and RNA maturation processes (e.g., 40S ribosomal protein, nucleolin-like isoform protein found OE and highly expressed in the core response to radiation, Fig 4B, Fig 8 in blue).

Besides DNA damages, RNA damages, likely also occur during desiccation and radiation but are so far less studied (Malotti et al., 2021). However, RNA might be even more prone to modifications and degradation compared to DNA notably because of their single stranded structure (Li et al., 2020), the non-interaction with proteins or nucleosomes protecting the DNA (Simms and Zaher, 2016), and their high abundance in the cell (80–90 % of the total nucleic acid
composition) of the cell (Li et al., 2020). While less lethal than genome mutations, their non-repair could result in delayed or faulty translation and inactive proteins, or dysregulation of gene expression (Nunomura et al., 2007). Indeed, oxidative RNA damage has been associated with neurodegenerative disorders (Nunomura et al., 2007). This probably explains why we detected the OE in the common response genes coding for proteins described to be involved in RNA repair, RNA silencing and degradation of RNA substrates (Fig 8 in pink), such as RNA ligases (Burroughs and Aravind, 2016) or ERI1 exoribonuclease (Asikainen et al., 2005; Thomas et al., 2014).

Finally, we also identified OE genes belonging to gene families whose roles in A. vaga resistance to desiccation and IR remains to be shown. However, their high log2-foldchange and/or TPM values compared to the controls, or the number of over-expressed genes, suggest they may play a role in A. vaga resistance and warrant further investigation. Among those genes, we first noticed the 2OG-Fe(II) oxygenase superfamily in the transcriptomic responses. Of the 49 annotated genes in the A. vaga genome, six are over-expressed in the common response to rehydration and radiation (Supplementary Table S3), and seven others are over-expressed under all radiation conditions (Supplementary Table S5). This gene family is involved in various processes essential for stress resistance, such as protein modification, DNA and mRNA repair, and synthesis of secondary metabolites (Loenarz and Schofield, 2011). Second, we found the major vault proteins (MVPs) with five OE genes: one found in the common response to rehydration and radiation (TPMs>150, 2.5h post radiation), and four in the common response to the different radiation types (two with TPMs>150, 2.5h post radiation). Their precise role in bdelloid rotifer resistance remain unclear even though such proteins have been linked to stress response in other studies, such as drug resistance, DNA repair (Lara et al., 2011), and response to DNA-damaging agents like IR (Shimamoto et al., 2006). Finally, two OE genes in the shared response and reaching high TPMs coded for proteins characterized by the domain “bZIP transcription factor” (PF00170). This domain is found in proteins regulating different physiological processes (e.g., adipogenesis, response to oxidative stress, in surveillance immunity, or cancer progression; Reddy et al. 2016). One of these genes, likely coding for a CCAAT/enhancer-binding protein, reached the highest TPM values in the common response (TPMs>1000, 2.5h post radiation or 1.5h post-rehydration) (Supplementary Table S3). We hypothesize that these putative transcription factors likely trigger the downstream upregulation of several genes involved in the response to desiccation and/or radiation in A. vaga.

2. Genes specific to desiccation and/or rehydration process

Although numerous genes were identified in the common transcriptomic response to rehydration following desiccation and radiation, some genes appeared to be specific to desiccation stress (Fig 8 in yellow). Desiccation not only induces damages and generates reactive oxygen species (ROS) similar to ionizing radiation (IR) (as shown in tardigrade, Giovannini et al., 2022), but also affects
the physiological state of the organism (hydrated vs. desiccated). Here, we showed that the dehydration process leading to the complete desiccation of A. vaga was characterized by a decrease of the metabolic activity (many under-expressed genes) with significant enrichment amongst UE genes of signal transduction, transport, and cell communication processes (Fig 3A). Cell communication and transport processes were also significantly enriched post radiation, however the number of UE genes involved in these processes was more than a two third smaller (Fig S4), suggesting not such reduction of metabolism post radiation. Similar patterns of gene under-expression were also observed during anhydrobiosis in the limno-terrestrial tardigrade Milnesium tardigradum (Wang et al., 2014).

Despite a general trend of reduced expression during dehydration, desiccation survival requires complex adaptations at morphological, physiological, and molecular levels to protect cellular components from desiccation-induced damage or facilitate their repair upon rehydration. Specific proteins seem particularly important in desiccation entry and the rehydration process following desiccation as we found 52 OE genes with high expression levels (TPMs>150) specific to post rehydration (Fig 4B) code for cell-adhesion proteins and microtubules, chaperones such as Late Embryogenesis Abundant (LEA) proteins and Heat Shock Proteins (HSPs), Glucose-repressible proteins, lipase and proteins involved in signaling or characterized by a calcium binding domain (Fig 4B), in addition to what is already described for the shared response. We described below the genes for which we can interpret their potential functions regarding desiccation resistance.

Genes coding for cell-adhesion proteins and microtubules (e.g., tubulin) that were over-expressed during desiccation entry and post-rehydration likely play a role in cell shape reorganization during water loss (desiccation) and rehydration (Fig 8 in yellow). Previous studies have reported alterations in the cytoskeleton and cell adhesion complexes in desiccated and rehydrated bdelloid rotifers (Marotta et al., 2010). Similar genes were also found to be over-expressed after rehydration in the tardigrade Milnesium tardigradum (Wang et al. 2014). In agreement with these findings, multiple homologs of such genes were reported in the genome of the bdelloid rotifer A. ricciae, supporting their importance in desiccation-resistant bdelloids (Eyres et al., 2012).

Several anhydrobiotic species produce disaccharides, such as trehalose, which are known to replace water and vitrify the cytoplasm (Erkut et al., 2011, Sakurai et al., 2008, Tapia and Koshland, 2014, Hengherr et al., 2008, Jönsson and Persson, 2010, Westh and Ramløv, 1991, Boothby et al., 2017). Surprisingly, trehalose has never been detected in bdelloid rotifers. In our study, we confirmed the over-expression and high abundance of trehalase genes (involved in the trehalose metabolic pathway, enriched during desiccation entry) potentially correlated with the absence or low abundance of trehalose in desiccated bdelloids (Lapinski and Tunnacliffe, 2003; Hespeels et al. 2015). However, we revealed that carbohydrate metabolic processes were enriched during the entry in desiccation (and only then) in bdelloid rotifers. Genes involved in carbohydrate metabolic processes could play a role in the production of carbohydrates instead of
trehalose to vitrify the cytoplasm, stabilizing the membranes and macromolecules upon desiccation (Carpenter et al., 1987; Crow and Crowe, 1992; Sun and Leopold, 1997). In addition, the over-expression of such genes in bdelloid rotifers entering desiccation may mediate their metabolism during desiccation-induced shutdown. Indeed, carbohydrates serve as a major source of energy and carbon. Consequently, the expansion of carbohydrate-related genes in bdelloids has been attributed to adaptations to diverse food sources, which in turn broadens their ecological niches, as observed in human gut microorganisms (Flot et al., 2013; Hespeels et al., 2012; Boschetti et al., 2011).

We found that genes coding for chaperones are also highly OE following desiccation (Fig 8 in yellow). First, hydrophobic LEA proteins are highly expressed and likely play a similar role as trehalose during desiccation and rehydration processes in bdelloid rotifers. These proteins were already detected to be upregulated in the bdelloid rotifer Philodina roseola during desiccation (Tunnacliffe et al. 2005), in desiccation resistant plants (Galau et al., 1986), in bacteria (Battista et al., 2001; Ujaoney et al., 2017), and in other desiccation-resistant metazoans (Browne et al., 2004; Adhikari et al., 2009; Bahrndorff et al., 2009; Schokraie et al. 2010; Hand et al., 2011; Hand and Menze, 2015; Ryabova et al., 2017). LEA proteins likely stabilize the cellular membranes in the absence of water (as suggested by Pouchkina-Stantcheva et al. 2007; Tunnacliffe and Wise, 2007; Tolleter et al., 2007; Tunnacliffe et al., 2010; Hand et al., 2011), protect against protein aggregation (Goyal et al., 2005), and have similar functions as non-reductive sugars in vitrification of the cytoplasm (Hand et al., 2011; Shimizu et al., 2010; Hibshman and Goldstein, 2021). LEA proteins have also been shown to protect crucial metabolic enzymes and were suggested to act as chaperone-like (Grelet et al., 2005; Reyes et al., 2005; Amara et al., 2014; Paul et al., 2014). Second, other types of chaperones highly expressed during the rehydration process of A. vaga (not OE during desiccation entry) were the heat shock proteins (HSPs), a few of them were also found in the common response to radiation and rehydration. Some of the OE HSPs already reached high expression levels in control condition (TPMs>300, Supplementary Table S4; e.g., FUN_27450 coding for a HSP70, TPMs= 2239), but even higher expression levels post-rehydration (e.g., TPMs= 3350 1.5h post rehydration following 14 days desiccation). Radiation only, although triggering over-expression of genes coding for HSPs, did not induce such a high expression level. HSPs are known to act as protein chaperones and therefore likely help nascent and misfolded proteins to gain their correct conformation (Kampinga and Craig, 2010; Stengel et al., 2010; Kubota, 2009). The high number of HSP genes (74 according to Pfam domain annotations) detected in A. vaga genome, their differential expression upon desiccation/rehydration processes and the high level of transcripts reported in this study, may support a central role of HSPs in their ability to deal with desiccation and radiation stresses. Indeed, such proteins were found to be over-expressed in other desiccation-resistant species to deal with this stress (Katoh et al., 2004; Cornette et al., 2010; Mizrahi et al., 2010; Reuner et al., 2010; Gechev et al., 2013).
Upon rehydration, one gene family particularly retained our attention because several copy numbers were over-expressed and reached high levels of expression (TPMs>150). These were annotated with the Pfam domain Glucose-repressible protein Grg1 (PF11034). Amongst the eight genes annotated with this Pfam domain Grg1 (PF11034) in A. vaga genome, six of them were OE with high TPMs (TPMS>150) post-rehydration (five also at desiccation entry), with one being found in the common response to rehydration and radiation (Fig 8). One gene was not OE but showed high constitutive TPMs in the controls (Fig 8 in green; Supplementary Table S3). This gene family was already found over-expressed in the bdelloid rotifer A. ricciae entering desiccation (Boschetti et al., 2011), but was never reported in other anhydrobiotic metazoans, including tardigrades, nor in the desiccation-resistant bacterium D. radiodurans. Although these genes were not predicted as acquired via HGTs in A. vaga using the Alienomics pipeline (Simion et al., 2021), the best blast matches were among fungi (e.g., Microbotryum lycnhdis-dioicae, Cladophialophora carrionii with e-value between 1e-12 and 5e-4) which was concordant with the gene tree inferred from Boschetti et al., 2011. It is interesting to note that genes coding for Grg1 are expressed in fungi during conidiation process when there is nutrient deprivation and desiccation (Kothe et al., 1998). We hypothesize that these Grg1 genes could have been horizontally acquired from fungi and retained for their plausible role in desiccation resistance (and glucose absence) characterized by a metabolic arrest in bdelloid rotifers.

3. Antioxidant role in desiccation and radiation resistances

Both desiccation and IR result in ROS production (Krisko 2010; Franca 2007; Krisko 2012; Daly 2012) that induces damages if not mitigated by antioxidants. Our dataset reveals that major antioxidant genes are highly expressed in both control and desiccated/irradiated samples (Fig 8 in green), with 26 genes showing high expression levels (TPMs>150; Fig 5A, 5C). This suggests that the constitutive expression of these antioxidants in A. vaga individuals may help them cope with oxidative stress, as they often encounter unpredictable desiccation events in their semi-terrestrial environments. Constitutive expression of protective proteins was also suggested in tardigrades (Hashimoto et al., 2016).

Desiccation seems to require more antioxidants than radiation (Fig 8 in yellow). Indeed, more antioxidant genes were OE during desiccation and rehydration following 14 days of desiccation (47 OE genes; yellow condition in Fig 5B) than after radiation only (9 OE genes when combining all time points; blue condition in Fig 5B). This link between anhydrobiosis and the antioxidant response was already emphasized by the many gene copy numbers of antioxidants found in the genome of A. vaga (Flot et al., 2013). This observation suggests either that the ROS concentration increases following desiccation or that the number of antioxidants decreased during desiccation and were re-expressed following rehydration.

Within genes specific to the core rehydration response (indicated in green in Fig. 5B), we found four aldo/keto reductases, one catalase, and one Copper/Zinc superoxide dismutase (SOD).
While this Cu/Zn SOD (FUN_013448) and another one (FUN_007045) were found OE post desiccation and after long rehydration, an Iron Manganese (FUN_017932) SOD was found to be OE only in hydrated irradiated rotifers (Fig 5B), even if it is also highly constitutively expressed (Fig 5C). This observation indicates specific role of SOD in hydrated or desiccated condition. An over-expression of genes coding for catalase, glutathione reductase, glutathione peroxidase, and SOD have also been reported for tardigrades during desiccation (Rizzo et al., 2010; Giovannini et al., 2022).

Two genes however seem to be key actors in A. vaga to deal with ROS and oxidative stress following radiation as they are highly OE post radiation: one aldehyde dehydrogenase found in the common response to rehydration and radiation, and a nucleoredoxin found in the core response to radiation (not considered in this study and in Flot et al., 2013 as an antioxidant gene family). These genes reaching higher expression level post radiation might be involved in additional processes other than their protective role against ROS. Indeed, the nucleoredoxin (TPMs>1000, 2.5h post radiation) might be required to keep the integrity of the antioxidant system, as it has been shown in plants (Kneeshaw et al., 2017). It was also shown in the tomato that nucleoredoxin can act as a positive regulator increasing antioxidant capacity and inducing HSPs to protect cells against oxidative damage and protein denaturation during heat stress (Cha et al., 2022). The aldehyde dehydrogenase seems to be important for multiple organisms in response to different stresses as it has been observed to be upregulated in bacteria (environmental and chemical stress), plants (dehydration, salinity, and oxidative stress), yeast (ethanol exposure and oxidative stress), Caenorhabditis elegans (lipid peroxidation), and mammals (oxidative stress and lipid peroxidation) (Singh et al., 2013). Its higher expression post radiation (TPMs>200) might be due to its potential role in the activation of DSBs resistance and DNA repair (Cojoc et al., 2015; Clark and Palle, 2016). This could explain why this gene was highly over-expressed post-radiation compared to other antioxidant gene in the bdelloid rotifer A. vaga, as X-rays lead to more DSBs than desiccation (Hespeels et al., 2014 & 2020). In addition, we found four other aldehyde dehydrogenases (Fig 5B) which were over-expressed only post rehydration after long desiccation. These genes might be involved more specifically in damages caused by desiccation only.

4. **HGTs have contributed to the evolution of resistance mechanisms in bdelloid rotifers**

Genomes of bdelloid rotifers such as A. vaga were described to contain an unusually high proportion of HGTs (Gladyshev et al., 2008; Flot et al., 2013; Eyres et al., 2015; Hespeels et al., 2015; Nowell et al., 2018; Simion et al., 2021), lastly determined to be 8.3% of protein-coding genes in case of A. vaga genome (Simion et al., 2021). Frequent desiccation events might explain this high percentage making the membranes leakier and the DNA fragmented, facilitating horizontal gene transfer events. It was previously hypothesized that HGTs in bdelloid rotifers may play a role in their extreme resistance to various stress (Boschetti et al., 2011, 2012; Eyres et
al., 2015, Szydlowski et al. 2015), unlike other species resistant to desiccation and/or radiation (e.g., Yoshida et al., 2017; Nowell et al., 2018; Kamilari et al., 2019). However, this hypothesis needs to be further evaluated as high proportion of HGTs was also found in the non-desiccating Rotaria rotifers (R. macrura, and R. magnacalcarata; Nowell et al., 2018). We hypothesize that genes important for resistance to extreme stresses and acquired via HGT have provided a selective advantage to the individuals that acquired those HGTs, being retained. If anhydrobiosis was the ancestral state, it could potentially explain the high proportion of HGTs which could have been acquired priorly to the loss of anhydrobiosis in some non-desiccating rotifers.

In this study, we found a significant enrichment of HGTs in the common transcriptomic response to rehydration following desiccation and radiation, and in the response to rehydration with ca. 14% of the OE genes (Fig 4B). Similar results (14.2%) have been found in response to rehydration (Nowell et al., 2018). Genes involved with desiccation resistance have also been described previously in rotifers (Boscetti et al., 2011, 2012, Hecox-Lea et al., 2018). Some genes detected by the pipeline Alienomics (Simion et al., 2021), such as 2OG-Fe(II) oxygenases, Ligase E, two RNA ligases, and genes belonging to Endonuclease/Exonuclease/phosphatase gene family particularly retained our attention because of their high TPM values and their potential role as key actors to resist desiccation and radiation. For instance, the ligase (FUN_003353) characterized as ligase E by Nicolas et al. (unpublished) (previously ligase K in Hecox Lea and Mark Welch 2018) is one of the genes found in the shared response to rehydration and radiation with the highest log2 foldchange (l2fc=7.32, 2.5h post desiccation and X-ray radiation; l2fc =2.3 1.5h post 15 days of desiccation) and highest expression levels in irradiated rotifers (TPMs>800 2.5h post-radiation; Supplementary Table S3). This gene, responsible for ligation, last step of the all repair pathways, appears to play a critical role in their resistance (Nicolas et al., unpublished). It is likely involved in BER but could potentially be also used for the ligation step of other repair pathways. Two other ligases (FUN_027119/FUN_021735), RNA ligases predicted as resulting from HGT, were found OE only in the core response to radiation and with high TPMs (TPMs>100, 2.5h post-radiation, Supplementary Table S5). These RNA ligases may play a role in nucleic acid repair as it was suggested for the RNA ligase DraRnl, upregulated in Deinococcus radiodurans after radiation (Schmier et al., 2017). Horizontal gene transfers have already been shown to play important roles in the evolution of 2OG-Fe(II) oxygenase genes in bacteria (Jia et al., 2017), which seems to have also played important roles for resistance to extreme conditions in bdelloid rotifers.

5. A. vaga exhibits similar molecular responses to both high- and low-LET radiation

This study allowed to compare the biological and molecular responses to low- (X-rays) versus high- LET (Fe-ions) radiations, known to induce different type of DNA damages (base lesions, abasic sites, SSBs, DSBs; Lehnert 2007; Semenenko and Stewart 2004; Penninckx et al.,
2019, Penninckx et al., 2021). Damages induced by low LET are more homogeneously distributed in the cells and along the DNA than when exposed with Fe-ions which induce concentrated clusters of DNA damages (Hespeels et al., 2020; Hespeels et al. unpublished). The more harmful impact of high-LET radiation exposure on reproduction activity of A. vaga confirmed what was already observed by Hespeels et al., 2020, as Fe-ions irradiation was associated with a severe sterilization of irradiated population (9±5% of fertile individuals post 500 Gy of Fe-ions vs 92±7% after 500 Gy X-ray radiation). Our findings revealed that the molecular actors involved in the radiation response of A. vaga seem to be independent of radiation type. The overall transcriptomic response observed was strikingly similar between high- and low-LET-radiation, with log2foldchange and TPM values for specific genes (antioxidant, DNA repair-associated genes, HSPs) falling within the same range (Fig 5B, 6B) and 648 genes found OE in the core response to radiation (Fig 4A, 4B). We found some genes coding for antioxidants which seem specific to Fe-ions radiation, however these genes did not reach high TPMs or l2fc values.

We mostly noticed one small difference: a time shift in the transcriptomic response between X-ray and Fe-exposed samples. While the major pattern of differentially expressed genes for A. vaga individuals exposed to 500 Gy occurred 2.5h post-rehydration, the response 8h post-Fe-ions irradiation was consistently larger than the response reported 8h post-X-ray radiation (e.g., Fig 3D). This shift could be due to a delayed response in Fe-exposed animals, which are expected to deal with more complex damages, resulting in slower recovery and a postponed transcriptomic response compared to those irradiated with X-rays. The delayed reactivation of rehydrated A. vaga individuals might also be attributed to a longer desiccation period (5 days) compared to X-ray irradiated samples (2 days) due to technical reasons for Fe-ions irradiation (see the Materials and Methods section). Extended desiccation periods were indeed correlated with a more prolonged recovery time in comparison to shorter periods (B. Hespeels, pers. comm.).

We hypothesize that the similarity in the transcriptomic responses following high- and low-LET radiation might be because A. vaga is almost never exposed to these high-LET radiation (Fe-ions). Consequently, a specific response to the more complex damage caused by high-LET radiation has not been selected for. A. vaga can probably use the same actors to face damages due to desiccation or radiation, even if high-LET radiation induce more clustered DNA damages than desiccation or low-LET radiation. This potentially less-adapted response to high-LET radiation might account for the reduced fertility observed following Fe-ion irradiation compared to X-ray exposure.

The difference in fertility rates observed with high-LET radiation triggers the question of how the germline cells deal with damages from radiation and if the mechanisms differ compared to somatic cells. We found in this study that most actors being OE post IR and desiccation belonged to NHEJ and BER pathways. NHEJ is known to be less faithful than HR when repairing DSBs and could cause some genetic modifications. Since bdelloid rotifers are known to be eutelic with a fix number of somatic cells once in the adult stage, they might be able to survive even if a few genetic
modifications remain in their somatic genomes. However, cellular division and DNA replication were shown to still occur in the germline of A. vaga (Terwagne et al., 2022). Therefore, the production of viable offspring requires the complete and faithful repair of damaged germline nuclei. Since 72% of the released eggs were unable to successfully complete embryological development after 500 Gy of Fe-ions, we hypothesized that the DNA repair process acting in the germlinal cells of bdellooids was overtaken by the complexity of damage and was associated with structural genomic change leading ultimately to the stop of embryological development. Transcriptomes used in this study represent mostly somatic cells (oocytes being a small proportion of the rotifer’s body), and we therefore cannot conclude on the DNA repair pathways acting in the germline of A. vaga. Future studies should establish alternative methods (e.g., single cell sequencing) allowing the investigation of the DNA repair mechanisms in developing oocytes after radiation. Future experiments will contribute to our understanding of how rotifers can survive despite the presence of residual DSBs documented in earlier studies (Hespeels et al., 2020).

Conclusions

The common transcriptomic response found in bdelloid rotifers Adineta vaga after rehydration post-desiccation and radiation (summarized on Fig 8 in pink) provides evidence that A. vaga resistance to high doses of ionizing radiation (IR) is likely derived from an evolutionary adaptation to anhydrobiosis, a stress-resistant state they routinely encounter in their semi-terrestrial environments, consistent with the hypothesis of Mattimore and Battista (1996). A. vaga bdelloid rotifers appear to be well-equipped to mitigate damages caused by reactive oxygen species (ROS) promptly and protect their proteome, as antioxidant genes and chaperones (HSPs) are constitutively highly expressed (Fig 8 in green). These genes and those involved in proteolysis via ubiquitination, over-expressed after both stresses, are critical actors in their resistance to desiccation and radiation as they preserve protein integrity including of those essential to repair damages (e.g., double strand breaks). If many DNA repair genes (NHEJ, BER) were found over-expressed both post rehydration and radiation, IR seems to induce the over-expression of more genes (Fig 8 in blue), as it probably causes more DNA damages. However, most essential genes for managing IR-DNA damages seem to have already evolved for withstanding desiccation, and have been co-opted to counteract radiation stress as well. In addition to these genes, desiccation resistance requires specific actors (Fig 8 in yellow) to deal with the modifications in the organism’s morphology or cells and damaged due related to water loss (more chaperones, HSPs and LEA, antioxidants, glucose repressive proteins, cytoskeleton proteins), and with metabolic reduction as seen in anhydrobiosis.

While our research has identified a common response of desiccation and IR resistance, it is necessary to investigate these resistences across different species with different resistance to radiation and desiccation, including desiccation sensitive species (Jönsson 2019; Boris et al., unpublished). We could have then a broader picture of how the resistance to the two stresses are connected. Finally, this study paves the road for reverse genetics experiments (e.g.,
knockouts) targeting promising candidate genes involved in DNA repair and coding for antioxidants but especially orphan genes with unknown functions (e.g., 58 genes found in the common response, Supplementary Table S9) which seem critical and unique of bdelloid rotifer radiation and desiccation resistance.

Methods

Bdelloid rotifer cultures

Experiments were performed using isogenic Adineta vaga clones originated from a single individual from the laboratory of Matthew Meselson at Harvard University. The cultures have been maintained hydrated at 21°C in 150 × 20 mm Petri dishes supplemented with natural spring water (Spa®) and fed weekly with sterile lettuce juice. A. vaga individuals used for the experiment aiming to evaluate the impact of 14 days of desiccation were cultivated with the same condition in the exception of food source which was Escherichia coli 1655MG.

Desiccation protocol

A. vaga individuals were dried following the optimized protocol previously described in Hespeels et al. 2014. Briefly, cultures were washed with 15 mL Spa® water the day before their collection. Individuals were detached from the petri dish with a cell scraper followed by a short round of vortex. Animals were transferred to a 15 mL Falcon tubes for centrifugation. Pellets, from each petri dish, were pooled in a final tube. The concentrated pellet containing rotifers was resuspended in Spa® water to a concentration of 80,000 individuals per mL. Then, for each desiccation sample, 0.5 mL of liquid was placed in the center of Petri dishes containing 30 mL of 3% Low Melting Point agarose (LMP agarose, Invitrogen). LMP agarose plates with hydrated individuals (approx. 40,000 per plate) were placed in a climatic chamber (WEKK 0028) for dehydration with the following parameters: (1) linear decrease in relative humidity (RH) from 70% to 55% for 17 h (Temperature: 21-23°C), (2) linear decrease in relative humidity from 55% to 41% for 1 h (Temperature: 21-23°C), and (3) maintenance at 41% RH and 21°C for the desiccated period. Based on this protocol, it took approx. 37 hours after the start of the dehydration process to dry all A. vaga individuals.

X-rays radiation

40,000 hydrated or one day desiccated bdelloids were irradiated with 225 kVp X-rays at a dose rate of ~7.8 Gy/min (using X-ray irradiator PXi X-RAD 225 XL) up to a final dose of 500 Gy. During irradiation, samples were maintained on a refrigerated water bag to mitigate heating due to soft X-rays.

Fe-ions irradiation

Irradiations with high-LET 56Fe-ions were performed at the GSI Helmholtz Center for Heavy Ion Research in Darmstadt, Germany. The iron ion beam was accelerated up to 1 GeV/n in the SIS18...
synchrotron and extracted in air in Cave A. Dosimetry and beam monitoring is described in Luoni et al. (doi: 10.3389/fphy.2020.568145). The beam has an LET of 147 keV/micron in water, to be compared to around 2 keV/micron for the 225 kVp X-rays. The preparation of desiccated samples took place in Namur University according to the protocol described above. Samples containing 40,000 individuals were transported under refrigerated environment to GSI the day before the exposure. Two sets of 3 days old desiccated bdelloid rotifers were assembled inside a sample holder allowing the exposition of multiple samples during a single beam time. After the exposure, samples were stored at 4°C and retrieved to Belgium for analysis. Samples were rehydrated 5 days after their entry in desiccation.

Survival and fertility

After desiccation or radiation, bdelloid individuals were cultivated (and potentially rehydrated using 15 mL Spa® water) at 21°C for 48 h. Bdeloids were considered alive when they had fully recovered motility or when the mastax moved in contracted individuals. Living bdelloids were separated from dead ones by transferring the supernatant, containing the latter, to new plates. Subsequently, the survival rate was manually calculated by tallying the number of living and dead specimens in each Petri dish under a binocular microscope.

In instances where individual count surpassed approximately 1,000 individuals, extrapolation was employed. This was achieved by counting sixfold the number of living or dead specimens observed in a 2 μL sample drawn from a homogenously mixed culture, with a final volume of 2–5 mL.

The reproductive capacity was defined as the ability for each isolated A. vaga individual to lay eggs and to develop clonal populations after being desiccated and potentially irradiated. We tested the fertility of 1 day desiccated and irradiated individuals by randomly selecting and isolating a minimum of 60 successfully rehydrated individuals per condition; each isolated female was deposited in a well of a 12-well petri plate. Each well was filled with 2 mL of Spa water and 50 μL of sterilized lettuce juice. After 30 days, wells were observed under a binocular stereoscope checking for: (1) the presence of a population (minimum 2 adults and 1 egg per well), (2) the presence of only eggs that did not hatch (+ eventually the single adult from the start defined as a sterile individual), and (3) the presence of only dead individual(s).

RNA extraction and RNA sequencing

Total RNAs were extracted using the RNAqueous-4PCR Kit (Ambion, Austin) following manufacturer instructions. Minimum 500 ng RNA of each sample was delivered to Genomicscore (Leuven, Belgium) or Genoscope (Paris, France). X-ray and Fe-ions irradiated RNA samples were treated as follow: RNA libraries were prepared using the TruSeq Stranded mRNA protocol (Illumina, San Diego, USA). Libraries were purified and evaluated using an Agilent 2100
bioanalyzer. RNA libraries were sequenced with an Illumina NextSeq 500 (Illumina, San Diego, CA USA) as paired-end 2 × 75 base pair reads using the NextSeq version 2.5 mid or high-output 150 cycle kit (Illumina). Samples studying the expression following 14 days of desiccation were treated as followed: total RNA was enriched in mRNA based on polyA tails, chemically fragmented and converted into single-stranded cDNA using random hexamer priming to then generate double-stranded cDNA. Next, paired-end libraries were prepared following the Illumina 222s protocol (Illumina DNA sample kit): briefly, fragments were end-repaired, 3'-adenylated, and ligated with Illumina adapters. DNA fragments ranging in size from 300 to 600 bp (including the adapters) were PCR-amplified using adapter-specific primers. Libraries were purified, quantified (Qubit Fluorometer; Life technologies), and library profiles were evaluated using an Agilent 2100 bioanalyzer. A paired-end flow cell of 101-bp reads was sequenced for each library on an Illumina HiSeq2000 platform.

The fastq files of each transcriptome are accessible on NCBI under the Bioproject PRJNA962496 under the biosamples SAMN34396008-SAMN34396072.

**Differential gene expression analysis (DGE)**

The whole bioinformatic workflow described below was performed in Galaxy Europe platform (usegalaxy.eu). Sequencing adapters and the first ten nucleotides were removed with Trim Galore (version 0.4.3.0) ([http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/](http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/)). The quality of the transcriptomes was checked by using FASTQC evaluation software FastQC (version 0.67) ([http://www.bioinformatics.babraham.ac.uk/projects/fastqc/](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)) before and after trimming the adapters. Trimmed-reads were mapped onto the genome assembly 2020 of *A. vaga* ([Simion et al., 2020](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)) using RNA-Star (version 2.5.2b0) ([Dobin et al., 2013](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)). We transformed for the mapping the gff file into a gtf file sing gffread (version 2.2.1.1) ([Trapnell et al., 2010](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)). A matrix of normalized counts per gene was assembled by htseq-count (version 0.6.1) ([Anders et al., 2015](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)).

We conducted the differential gene expression analysis (DGE) with all the data together, combining 15 control samples together, and combining 11 desiccated samples, using DESeq2 (version 2.11.39, [Anders and Huber, 2010](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/); [Love et al., 2014](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)) and EdgeR (version 3.28.0) ([Robinson et al., 2010](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/); [McCarthy et al., 2012](http://www.bioinformatics.babraham.ac.uk/projects/fastqc/)) in the R statistical software (version 3.4.1) ([http://www.R-project.org](http://www.R-project.org)) (not performed in galaxy platform).

For subsequent analyses, genes were considered significantly differentially expressed in the treated samples compared to the control samples if they have both adjusted p-value < 0.01 for DESeq2 analyses and an adjusted false discovery rate (FDR) < 0.01 for EdgeR results and the absolute values of their log₂-fold-change ≥ 0.5 with both tools. Those with a log₂-fold-change ≥ 0.5 were more expressed in the treated samples than in the controls, whereas those with a log₂-fold-change ≤ -0.5 were less expressed in the treated samples than in the controls. In this study we used the values obtained with DESeq2 for the graphs and tables.
DGE comparisons

We analyzed the transcriptomic responses of *A. vaga* individuals exposed to desiccation and/or high-/low-LET radiation compared to hydrated controls (n=15) (Fig 2A). In order to capture the transcriptomic pattern resulting from the dehydration response, we focused our analysis on *A. vaga* individuals entering desiccation after 48 hours (n=11). In order to characterize the radiation response of *A. vaga* individuals to X-rays, hydrated and desiccated animals were exposed to 500 Gy. The transcriptomic response was investigated post radiation exposure and optional rehydration at different time points: 2.5h, 8h and 24h (this latter timepoint was only investigated in previously desiccated rotifers) (n=4 for each time point). Comparison of radiation response in hydrated and rehydrated animals ensured the discrimination of radiation specific response from genes impacted by the dehydration/rehydration process. A similar experiment was performed by exposing only desiccated *A. vaga* individuals to 500 Gy Fe-ions radiation (n=4 for time points 2.5h, 8h, and 24h). This second kinetic was performed in order to compare the biological and transcriptomic response of irradiated *A. vaga* against low- or high-LET radiation. Finally, we analyzed the transcriptomic response in *A. vaga* individuals rehydrated for 1.5h after being desiccated during 14 days (n=3).

Search for genes coding for enzymes involved in DNA repair pathways

KO (KEGG Ontology) annotations were performed using bi-directional best hit with the KAAS annotation server (Moriya et al., 2015). Eleven supplemental organisms enriched the KAAS default gene dataset in order to increase the annotation quality. Based on KO numbers, pathway maps were designed using the KEGG pathway reconstruction tool (Kanehisa and Goto, 2000; Kanehisa, 2019; Kanehisa et al., 2023).

Annotated genes were validated by blast and *A. vaga* genome was manually curated in order to complete DNA repair pathways and antioxidant gene families. We searched for putative protein functions by similarity searches using BLASTP in the non-redundant database (NRdb) using a threshold expect value E < 10^-5. Horizontal gene transfer (HGT) acquisition was evaluated based on Alienomics as previously calculated for the *A. vaga* gene set (Simion et al. 2020). This approach only identifies putative horizontal gene transfers (HGTs) from non-metazoans (e.g., bacteria, plants or fungi).

Transcripts per millions

Transcripts per million (TPMs) for each gene was computed with a custom Python script (given in the supplements).

Go enrichment analysis
Go ids were annotated using InterProScan (Zdobnov and Apweiler, 2001) in galaxy. GO enrichment analysis was performed in R, using the package “topGo” from Bioconductor (Alexa et al., 2016), (script available under request).

**Pathway and heatmap**

KEGG ids were annotated by InterProScan in galaxy software at the same time as GO ids. DNA repair pathways (NHEJ, BER, NER, HR, MR) were represented based on the KEGG pathway map, respectively, map03450, map03410, map03420, map03440, map03430. The log2-foldchanges of genes significantly differentially expressed (confirmed with EdgeR) obtained with Deseq2 were represented for each gene of the pathways, using ComplexHeatmap package from Bioconductor (Gu et al. 2016). Additionally, TPMs were also represented using the same package (script available under request).

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and materials**

All data generated or analyzed during this study are published on zenodo (available when manuscript published), and transcriptomic reads can be found on NCBI under the Bioproject PRJNA962496 under the biosamples SAMN34396008-SAMN34396072.

**Competing interests**

The authors declare that they have no competing interests

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**Author contributions**

VCM: Transcriptomic analyses, figures, tables, core manuscript writing
LB: Preparation of samples for X-ray and Fe-ions irradiation, lab cultures, RNA extraction and purification
JB: preparation of samples for Fe-ion irradiation, lab cultures, survival and fertility assays
ACH: Radiation experiment support
SP: Radiation experiment support
SR: Fe-ion irradiation experiment
UW: Fe-ion irradiation experiment
MD: Fe-ion irradiation experiment
EGJD: Support for preliminary transcriptomic analyses
BH: lab cultures, preparation of samples for X-ray and Fe-ion irradiation, survival and fertility rate, RNA extraction and purification, preliminary transcriptomic analyses, experimental design conception, manuscript writing, fund acquisitions
KVD: experimental design conception, follow-up on the results, manuscript writing, fund acquisitions, project supervisor

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References in order of apparition


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Figures, tables and supplementary files

Figures

A) Hypothesis investigated in this study: are they common molecular mechanisms (pink) involved in the resistance to desiccation (yellow) and radiation (blue) in the bdelloid rotifer *Adineta vaga* since both stresses are known to induce similar damages including DNA double strand breaks (DSBs), are the DNA repair mechanisms the same following both stresses, and do high and low LET induce similar transcriptomic response since they are known to lead different types of damages and that high-LET impacts more the fertility of *A. vaga* compared to low-LET. B) Percentage of survival (dark blue) and fertility rate of *A. vaga* exposed to desiccation, rehydration post 14 days desiccation, and 500 Gy of X-rays (low-LET) or Fe-ions (high-LET). The fertility is represented by three different histograms: fraction of individuals able to produce viable offspring (green), fraction of individuals unable to produce viable offspring but only sterile egg(s) (blue), fraction of individuals unable to lay egg or with premature dead (red).

Fig 1
**Fig 2 Design of the comparative transcriptomic approach and responses**

A) Experimental design of the comparative transcriptomic analyses of *Adineta vaga* individuals exposed to desiccation and/or radiation: hydrated controls (in various greens), individuals entering desiccation (48 hours desiccated: in gray), rehydrated after 14 days of desiccation (in yellow), irradiated with 500 Gy X-rays (low-LET) after desiccation (in orange), irradiated with 500 Gy Fe-ions (high-LET) after desiccation (in purple), hydrated individuals irradiated with 500 Gy X-rays (in blue). The different investigated time points post-rehydration or radiation are represented as follows: 0 hours (triangles), 1.5 and 2.5 hours (circles), 8 hours (squares), 24 hours (pentagons). N = the number of replicates analyzed per condition. B) Principal component analysis of the transcriptomes analyzed (with 3 to 4 replicates per condition) with PC1 representing 56% of the variance in gene expression between samples and PC2 16% of the variance. C) Histogram of the number of genes (first y axis) and the percentage of genes within the genome (second y axis) being significantly (FDR<0.01 with both DESeq2 and EdgeR) over- (log₂foldchange>0.5 in green) or under-expressed (log₂foldchange<-0.5 in red) in the different conditions post-desiccation and/or radiation.
**Fig 3 Gene Ontology enrichment analyses** A) GO biological processes significantly enriched (chi-square test p-value<0.05, min. 3 OE genes with the GO id) with genes being significantly under-expressed (framed in red, FDR<0.01 and log2foldchange < -0.5 in DESeq2 and EdgeR) in *A. vaga* individuals entering desiccation (in gray) and 1.5 hours post-rehydration after 14 days of desiccation (in yellow). B) GO biological processes significantly enriched (chi-square test p-value<0.05, min. 3 OE genes with the GO id) with genes being significantly over-expressed (framed in green, FDR<0.01 and log2foldchange > 0.5 in DESeq2 and EdgeR) in *A. vaga* individuals entering desiccation (in gray) and 1.5 hours post-rehydration after 14 days of desiccation (in yellow). C) Venn diagrams with number of genes significantly over-expressed and under-expressed (red outlines) in *A. vaga* individuals entering desiccation (gray) or after rehydration and 14 days of desiccation (yellow), showing 915 genes commonly over-expressed and 1885 genes commonly under-expressed. D) Venn diagrams of genes significantly over-expressed in *A. vaga* individuals under three different conditions of radiation at 2.5h and 8h post-irradiation and/or rehydration: hydrated *A. vaga* individuals irradiated with X-rays (blue), desiccated *A. vaga* individuals irradiated with X-rays post rehydration (orange), desiccated *A. vaga* individuals irradiated with Fe post rehydration (purple). At 2.5h and 8h post-irradiation and/or rehydration, respectively 906 and 724 genes are over-expressed in the three conditions. E) GO biological processes significantly enriched (chi-square test p-value<0.05, min. 3 OE genes with the GO id) with genes found to be over-expressed in the core response to radiation (906 genes 2.5 hours post-radiation and 724 genes 8 hours post-radiation).
Fig 4 Core transcriptomic response of A. vaga to desiccation and radiation. A) Venn diagram of genes significantly over-expressed (OE: FDR<0.01 and log2foldchange>0.5 with DESeq2 and EdgeR) in A. vaga individuals in different studied conditions: 1.5 hours post-rehydration following 14 days of desiccation (yellow), 2.5 hours post-radiation and desiccation (Fe radiation in purple, X-ray radiation in orange), and 2.5 hours post-radiation (X-rays, blue). Genes specifically coding for DNA repair are written in dark blue and those coding for antioxidants are written in pink. B) OE genes with high level of expression (TPMs>150) and the potential function of the protein for which they code or the biological processes in which they are likely involved in the core response to rehydration (encircled in dark yellow in the Venn diagram, TPMs>150 in the condition 1.5 hours post-rehydration after 14 days of desiccation), in the core response to radiation (encircled in blue in the Venn diagram, TPMs>150 in all condition 2.5 hours post irradiation) and the genes found in both responses (common, TPMs>150 in the condition 2.5 hours post-desiccation and X-ray irradiation).
A

Number of genes

<table>
<thead>
<tr>
<th>Aldehyde dehydrogenase</th>
<th>Aldo keto reductase</th>
<th>Catalase</th>
<th>Glut per</th>
<th>Glut red</th>
<th>Glut-transf</th>
<th>Isocitrate dehydrogenase</th>
<th>Peroxiredoxin</th>
<th>SOD</th>
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<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Legend
- Control
- Dess. + x-rays (2.5h)

TPMs
- <10
- 10-150
- >150
- >700

B

69 Genes over-expressed
(L2fc=0.5) in at least one condition

<table>
<thead>
<tr>
<th>L2fc</th>
<th>x-rays</th>
<th>Fe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dess 14d</td>
<td>Dess 48h</td>
<td></td>
</tr>
</tbody>
</table>

TPMs

C

26 Genes highly expressed
(TPMs>150) in controls

Legend
- 2 genes in common response: rehydration & radiation
- 6 genes specific to common rehydration response
- 2 genes specific to common radiation response
- 3 genes specific to hydrated irradiated rollers
- 34 genes specific: rehydration post long desiccation/desiccation entry
Fig 5 Transcriptomic response in *A. vaga* of genes coding for antioxidants. A) Number of genes coding for the different antioxidant gene families (x axis). According to the level of expression: transcripts per millions (TPMs), the number of genes is represented in different color: TPMs < 10 (gray), 10<TPMs<150 (light orange), >150 (red), >700 (dark red). The number of genes in each expression category is represented for *A. vaga* hydrated controls (green) and *A. vaga* individuals 2.5 hours post X-ray radiation and rehydration (orange). B) Heatmap of change of expression of 39 genes coding for antioxidant gene families which are at least over-expressed in one condition (log₂foldchange > 0.5, FDR < 0.01 with both DESeq2 and EdgeR). The 30 genes found OE exclusively after 14 days of desiccation are indicated as numbers in the corresponding column. Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different color code representing the level of expression. The gene IDs coding for the different kinds of antioxidants are written next to the TPMs, and Horizontal transfer genes (HGT) are represented by the violet stars. C) Heatmap representing the level of expression (TPMs) of 26 genes showing constitutive high expression (TPMs>150) in all conditions coding for antioxidants. After desiccation and radiation, the number of genes coding for antioxidants showing high TPMs does not change. The ID of the genes found in the common response to rehydration and radiation are indicated in pink (2 genes), in the common rehydration response (in green: 6 genes), in the common radiation response (in dark blue: 2 genes), specific to hydrated irradiated rotifers (in light blue). The four genes specific to desiccation and 1.5h post rehydration are indicated in yellow, same color as the 30 genes specific to 1.5h post rehydration (indicated as numbers).
Fig 6 Transcriptomic response in *A. vaga* of genes involved in DNA repair pathways. A) Number of genes coding for genes involved in DNA repair pathways: Alternative Excision Repair (Alt ER), Base Excision Repair (BER), Homologous Recombination (HR), Mismatch Repair (MR), Nucleotide Excision Repair (NER), Non-Homologous End-Joining (NHEJ), DNA Replication (REP), Cross-Pathways (Cross), Damages-DNA Repair related pathways (DDR-related). According to the level of expression (transcripts per millions (TPMs)), the number of genes is represented in different colors: TPMs<10 (gray), 10<TPMs<150 (light orange), >150 (red), >700 (dark red). The number of genes in each expression category is represented for *A. vaga* hydrated controls (green) and *A. vaga* individuals 2.5 hours post X-ray radiation and rehydration (orange). B) BER pathway showing numerous genes being over-expressed (L2fc > 0.5) and some with high TPMs after desiccation and/or radiation. This figure represents the different actors of the BER pathway and their interaction, actors being represented by different symbols. The heatmap of the log₂foldchange shows the level of expression of BER genes change compared to the expression level in the controls. The color is not white when the gene is significantly (FDR < 0.01 with both DESeq2 and EdgeR) over- (log₂foldchange>0.5 in green) or under-expressed (log₂foldchange<-0.5 in orange or red). Genes potentially involved in this pathway are indicated in the bottom of the figure. The investigated conditions are: individuals entering desiccation (gray), after 14 days desiccation and 1.5 hours rehydration (yellow), after desiccation and X-rays radiation (orange), after desiccation and Fe radiation (purple), after X-ray radiation without desiccation (blue). The investigated time points are represented as follow: 0 hours (triangles), 1.5 or 2.5 hours (circles), 8 hours (squares), 24 hours (pentagons). Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different color code representing the level of expression. The gene IDs coding for the different kinds of BER genes are written next to the TPMs, and Horizontal transfer genes (HGT) are represented by the violet stars. C) Heatmap representing the level of expression (TPMs) of 6 genes coding for proteins involved in DNA repair pathways (BER, MR, DDR-related) showing constitutive high expression (TPMs>150) in all conditions. D) Heatmap representing the level of expression (TPMs) of 15 additional genes coding for proteins involved in DNA repair which have a high gene expression level after radiation and desiccation.
**Fig 7** Expression of genes involved in the Non-Homologous End-Joining (NHEJ) and Homologous Recombination (HR) pathways for double strand break (DSB) repair. The figure represents the different actors act of these 2 DNA repair pathways being represented by different symbols. The heatmap of the log2 foldchange shows the change level of expression of genes coding for these actors in the different conditions in *A. vaga* individuals compared to the gene expression level in the controls. The color is not white when the gene is significantly (FDR < 0.01 with both DESeq2 and EdgeR) over- (log2foldchange>0.5 in green) or under-expressed (log2foldchange<-0.5 in orange or red). In this figure are represented only the following conditions: individuals entering desiccation (gray), after 14 days desiccation and 1.5 hours rehydration (yellow), after 2.5-, 8- and 24-hours post desiccation and X-rays radiation (orange), since the response was similar after desiccation and Fe radiation and after X-ray radiation without desiccation (all conditions represented in Supplementary Figures S4 and S5). Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different color code representing the level of expression. The gene IDs coding for the different DNA repair genes are written next to the TPMs, and Horizontal transfer genes (HGT) are represented by the violet stars. This figure illustrates that DSB are likely repaired in *A. vaga* individuals by NHEJ rather than by HR.
**Fig 8 Summary of common and specific responses to desiccation and ionizing radiation.**

Common damages known to be induced by desiccation and ionizing radiation: membrane damages, ROS production leading to phospholipid peroxidation and de-esterification, protein damages (oxidation, disulfide bridge, denaturation, aggregation, misfolding), and DNA damages (Double Strand Break: DSB, Single Strand Break: SSB, chemical modifications). The major molecular responses observed in *Adineta vaga* post-rehydration (with or without radiation; in yellow) and post-radiation (in blue), or common to both (in pink). Genes coding for some antioxidants, HSPs, high mobility proteins, and one CCAAT enhancer were found to be constitutively highly expressed (in green) in all conditions including controls. We found genes coding for HSPs, Major Vault proteins (MVP), Glucose repressive proteins (Grg1), aldehyde dehydrogenase, regulation factors such as CCAAT enhancer, histone variants genes involved in protein modifications like ubiquitination, in proteolysis, DNA repair (mostly NHEJ and BER pathways) and RNA repair, being OE in the common response to rehydration and irradiation. Specific to rehydration, we found OE genes coding for Late Embryogenesis Abundant proteins (LEA), Heat Shock Proteins (HSPs), tubulin, Glucose repressive proteins (Grg1), and more antioxidants. Radiation increases the number of OE genes coding for histone variants, RNA/DNA repair proteins, and MVPs.
Fig S1 Survival rate of *A. vaga* individuals exposed to desiccation and radiation stress. Survival rate was evaluated 2 days post rehydration or post radiation. To ensure reliable results, survival data were obtained from at least three replicates. Samples exposed to 500 Gy of iron ions were transported from Belgium to GSI and then returned to the authors' laboratory for analysis. To evaluate the impact of transportation on the samples, a control group labeled "Ctl transport" was also sent to GSI. This control group experienced similar conditions as the exposed samples but did not undergo radiation exposure.

Fig S2 Volcano plots of differential genes over-expressed (OE) and under-expressed (UE) in A) *A. vaga* individuals entering desiccation, B) 2.5 hours post desiccation and x-rays radiation. The percentage of the genome over- and under-expressed are written. Although a higher percentage of genes are under-expressed in B) the log₂-foldchange values are bigger in OE than UE genes.
**Fig S3** Number of genes with GO ids for a specific log<sub>2</sub>foldchange values identified as A) over-expressed (OE) genes in *A. vaga* rotifers entering desiccation, B) under-expressed (UE) genes in *A. vaga* rotifers entering desiccation, C) 906 OE genes in the core response to radiation 2.5 hours post irradiation, D) 724 OE genes in the core response to irradiation 8 hours post radiation.
Fig S4 Gene Ontology enrichment analyses GO biological processes significantly enriched (chi-square test p-value<0.05, min. 3 OE genes with the GO id) with genes being significantly A) under-expressed (framed in red, FDR<0.01 and log₂foldchange < -0.5 in DESeq2 and EdgeR) and B) over-expressed (framed in green, FDR<0.01 and log₂foldchange > 0.5 in DESeq2 and EdgeR) in A. vaga individuals entering desiccation (in gray) and 1.5 hours post-rehydration after 14 days of desiccation (in yellow), and in the core response to radiation 2.5h (pink) and 8h (purple) post radiation.
**Fig S5** Genes coding for proteins involved in Non-Homologous End-Joining (NHEJ) pathway differentially expressed post desiccation and/or radiation. The figure represents how the different actors act in this DNA repair pathway and in which order. They are represented by different symbols. How the level of expression of genes (gene ids written on the right side of the figure) coding for these actors change in the different conditions is represented by a heatmap of the log2 foldchange. The color is not white when the gene is significantly (FDR < 0.01 with both DESeq2 and EdgeR) over-expressed (log2 foldchange > 0.5 in green) or under-expressed (log2 foldchange < -0.5 in orange or red). The investigated conditions are: individuals entering desiccation (gray), after 14 days desiccation and 1.5 hours rehydration (yellow), after desiccation and x-rays radiation (orange), after desiccation and Fe radiation (purple), after x-ray radiation without desiccation (blue). The investigated time points are represented as follow: 0 hours (triangles), 1.5 or 2.5 hours (circles), 8 hours (squares), 24 hours (pentagons). Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different code representing the level of expression. Horizontal transfer genes (HGT) are represented by the violet stars.
Fig S6 Genes coding for proteins involved in Homologous Recombination (HR) pathway differentially expressed post desiccation and/or radiation. The figure represents how the different actors act in this DNA repair pathway and in which order. They are represented by different symbols. How the level of expression of genes (gene ids written on the right side of the figure) coding for these actors change in the different conditions is represented by a heatmap of the log2-foldchange. The color is not white when the gene is significantly (FDR < 0.01 with both DESeq2 and EdgeR) over- (log2-foldchange>0.5 in green) or under-expressed (log2-foldchange<-0.5 in orange or red). The investigated conditions are: individuals entering desiccation (gray), after 14 days desiccation and 1.5 hours rehydration (yellow), after desiccation and x-rays radiation (orange), after desiccation and Fe radiation (purple), after x-ray radiation without desiccation (blue). The investigated time points are represented as follow: 0 hours (triangles), 1.5 or 2.5 hours (circles), 8 hours (squares), 24 hours (pentagons). Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different code representing the level of expression. Horizontal transfer genes (HGT) are represented by the violet stars.
Fig S7 Genes coding for proteins involved in Nucleotide excision repair (NER) pathway differentially expressed post desiccation and/or radiation. The figure represents how the different actors act in this DNA repair pathway and in which order. They are represented by...
different symbols. How the level of expression of genes (gene ids written on the right side of the figure) coding for these actors change in the different conditions is represented by a heatmap of the log-fold change. The color is not white when the gene is significantly (FDR < 0.01 with both DESeq2 and EdgeR) over-expressed (log-foldchange > 0.5 in green) or under-expressed (log-foldchange < -0.5 in orange or red). Genes potentially involved in this pathway are indicated in the bottom of the figure. The investigated conditions are: individuals entering desiccation (gray), after 14 days desiccation and 1.5 hours rehydration (yellow), after desiccation and x-rays radiation (orange), after desiccation and Fe radiation (purple), after x-ray radiation without desiccation (blue). The investigated time points are represented as follow: 0 hours (triangles), 1.5 or 2.5 hours (circles), 8 hours (squares), 24 hours (pentagons). Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different code representing the level of expression. Horizontal transfer genes (HGT) are represented by the violet stars.

**Fig S8** Genes coding for proteins involved in Mismatch repair pathway (MR) differentially expressed post desiccation and/or radiation. The figure represents how the different actors act in this DNA repair pathway and in which order. They are represented by different symbols. How the level of expression of genes (gene ids written on the right side of the figure) coding for these...
actors change in the different conditions is represented by a heatmap of the log.foldchange. The color is not white when the gene is significantly (FDR < 0.01 with both DESeq2 and EdgeR) over- (log.foldchange>0.5 in green) or under-expressed (log.foldchange<-0.5 in orange or red). Genes potentially involved in this pathway are indicated in the bottom of the figure. The investigated conditions are: individuals entering desiccation (gray), after 14 days desiccation and 1.5 hours rehydration (yellow), after desiccation and x-rays radiation (orange), after desiccation and Fe radiation (purple), after x-ray radiation without desiccation (blue). The investigated time points are represented as follow: 0 hours (triangles), 1.5 or 2.5 hours (circles), 8 hours (squares), 24 hours (pentagons). Heatmap of transcripts per millions (TPMs) for each gene is represented on the right side and with a different code representing the level of expression. Horizontal transfer genes (HGT) are represented by the violet stars.

**Supplementary Tables (on zenodo)**

**Supplementary Table S1 Genes within Adineta vaga genome.** The columns give the gene ids from the gene assembly in Simion et al. (2020), the correspondence gene ids in the genome published by Simion et al. (2021), if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S2 GO analysis of Under-Expressed genes in the core response.**

**Supplementary Table S3 Genes found over-expressed in the core common response to radiation and desiccation.** The columns give the gene ids, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S4 Genes found over-expressed in the core rehydration response.** The columns give the gene ids, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S5 Genes found over-expressed in the core irradiation response.** The columns give the gene ids, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S6 List of genes coding for the defined antioxidant gene families.** The columns give the gene ids, the antioxidant family, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S7 34 Genes coding for antioxidants exclusively significantly over-expressed in A. vaga individuals 1.5 hours post rehydration after 14 days of desiccation.**
The columns give the gene ids, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S8** List of genes coding for proteins involved in DNA repair. The columns give the gene ids, the DNA repair pathway in which the gene is involved, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.

**Supplementary Table S9** List of genes with unknown functions but high log₂ foldchange and TPMs post radiation and/or desiccation. The columns give the gene ids, if the gene is annotated as HGT, the blast hits, the KEGG ids, GO ids, pfam domains, the results for all analyses with DESeq2 and EdgeR, and the TPM values for all the different analyzed conditions.