Transcriptomic and genomic changes associated with radioadaptation in Exophiala dermatitidis

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ABSTRACT

Melanized fungi have been isolated from some of the harshest radioactive environments, and their ability to thrive in these locations is in part due to the pigment melanin. Melanin imparts a selective advantage to fungi by providing a physical shield, a chemical shield, and possibly a signaling mechanism. In previous work we demonstrated that protracted exposure of the melanized yeast Exophiala dermatitidis to mixed alpha-, beta-, and gamma-emitting radiation resulted in an adapted strain able to mount a unique response to ionizing radiation in the environment in a melanin-dependent fashion. By exploring the genome and transcriptome of this adapted melanized strain relative to a non-irradiated control we determined the altered response was transcriptomic in nature, as whole genome sequencing revealed limited variation. Transcriptomic analysis indicated that of the adapted isolates analyzed, two lineages existed: one like the naïve, non-adapted strain, and one with a unique transcriptomic signature that exhibited downregulation of metabolic processes, and upregulation of translation-associated genes. Analysis of differential gene expression in the adapted strain showed an overlap in response between the control conditions and reactive oxygen species conditions, whereas exposure to an alpha particle source resulted in a robust downregulation of metabolic processes and upregulation of DNA replication and repair genes, and RNA metabolic processes. This suggest previous exposure to radiation primes the fungus to respond to subsequent exposures in a unique way. By exploring this unique response, we have expanded our knowledge of how melanized fungi interact with and respond to ionizing radiation in their environment.

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1. Introduction

Numerous fungal species can be considered as some of the most radiation resistant organisms. They have been isolated from several extreme environments, such as Antarctic deserts, the Chernobyl Atomic Energy Station (including within reactor 4 where the disaster actually initiated), and the International Space Station (ISS) [37,46,35,34,44,5,43,20,12], and so have proven not only capable of surviving both acute and chronic exposure to various forms of radiation whilst enduring a variety of other environmental stresses, but also thriving in these unique conditions [47].

In the search to identify what makes fungi so impervious to radiation, the pigment melanin has been identified as an interesting candidate. Melanin has been detected in ancient fossils dating back to the Cretaceous and Jurassic periods, and is present in most kingdoms, suggesting strong evolutionary preservation [55,31,16]. The presence of melanin, moreover, is associated with enhanced survival and improved fitness in extremely radioactive environments, as indicated by the prevalence of melanized fungal strains isolated from these locations [28]. This indicates that melanin has played an important role in Earth's history, and still imparts some important attributes to many organisms, particularly with regards to stress resistance.

In previous works, we and others have demonstrated that melanized fungal species, including Cryptococcus neoformans, Exophiala...
E. dermatitidis, and Cryomyces antarcticus, display enhanced growth and improved survival when exposed to ionizing radiation when compared to their non-melanized counterparts [29,30,27,36,11]. Melanin appears to enhance resistance to radioactive environments by providing a physical shield via Compton scattering, a chemical shield by quenching reactive oxygen species (ROS), altering growth kinetics and energy use, and possibly as a signaling mechanism by communicating conditions in the environment via its action as a redox capacitor, thereby initiating a transcriptomic response [11,10,50,7,19,25,41]. Work by Zhdanova and Tugay on fungal isolates from the Chernobyl “exclusion zone” demonstrated that a number of isolates were 1) resistant to, 2) stimulated by, and 3) adapted for growth in radioactive environments [49,12,48,56]. The fungal isolates demonstrated enhanced hyphal growth in the direction of a radioactive source, as well as enhanced spore germination, whereas control strains that had never been exposed to radiation experienced growth inhibition when exposed to radiation. The molecular details of how fungi detect, interact with, and respond to ionizing radiation in the environment, however, are unknown.

To investigate this issue further, we recently developed radiation adapted lab strains that could be used to further our understanding of this phenomenon [30] through the protracted exposure of melanized and non-melanized E. dermatitidis strains to Actinium-225 (225Ac), which is a mixed alpha, beta and gamma-emitter. A subsequent phenotypic characterization of these “radioadapted” strains (Fig. 1) determined that the melanized, adapted strain showed increased colony size relative to naïve and albino control strains when grown adjacent to an alpha particle emitter, had enhanced electron transfer capacity, and displayed improved resistance to ROS. We interpreted these results to mean that previous exposure to ionizing radiation enhanced the capacity of our fungal cultures to detect and respond to radiation in the environment.

Our goal with the current study is to further understand the changes that occur in this adapted, melanized strain that enable it to detect and interact with ionizing radiation in the environment.

To that end, we set out to explore the genomic and transcriptomic landscape of several of these radioadapted strains. We found that, while few permanent mutations occurred during the adaptation process, one lineage of the melanized (wild type) adapted strain exhibited extensive transcriptomic changes, as well as the ability to respond to subsequent ionizing radiation exposure more robustly. Details of this response are provided below, which provide insight into how and why fungi grow and thrive in these uniquely harsh conditions.

2. Materials and methods

2.1. Growth conditions

The strains that were used in this experiment were the wild type (WT) strain of E. dermatitidis 8656 (aka ATCC34100, Exophiala dermatitidis CBS 525.76) and the albino mutant strain, wdpks1 [14], that were either radioadapted by exposure to alpha radiation containing 1 mCi/ml 225Ac or were passaged under background conditions and served as naïve controls, all of which were developed by our group in a previous study [30]. The protracted dose of 183 Gy/5 weeks delivered at a low dose rate of 0.02 mGy/min is a non-lethal dose for a melanized fungus, as melanized fungi can survive doses of >200 Gy delivered at high dose rates of >10000 mGy/min [28]. Single colonies from these naïve control and 225Ac-adapted strains were isolated, cultured, and preserved as frozen glycerol stocks to be utilized for all experiments in this study (Table 1). Isolates were maintained in a modified Sabouraud Emmons Broth (SAB; 2% dextrose, 1% peptone) at 30°C with shaking at 200 rpm until they reached an approximate concentration of 10^6 cells/ml. Cells were then transferred into minimal media (MM; 2 g/L KH2PO4, 1.25 g/L MgSO4 7H2O, 0.5 g/L glycine, 0.5 g/L dextrose, 0.003 g/L thiamine) and cultured at 30°C.

2.2. Polonium-210 exposure

Isolates grown in minimal media where dextrose was replaced with 0.01% sucrose (MMS) were used in alpha (α) particle radiation stimulation studies as previously described [30]. In brief, a 500 μCi Polonium-210 (210Po) source (NRD, Grand Island, NY, USA) housed in a metal holder was placed within 3 mm of solid media. The source within the metal holder produced a collimated 0.5 μCi/hr beam of radiation that interacted with the solid growth media. Fungal cultures were diluted to a concentration of 10^4 cells/ml and approximately 10 cells were plated on solid MMS media such that they were outside of the collimated beam of radiation generated by the 210Po source.

2.3. DNA and RNA sequencing

To obtain nucleic acids for genome and transcriptomic analysis, samples were prepared as follows. Isolate cultures were grown in MM as described above until the wildtype strains developed visible melanin, which was approximately 2 days, and were still in logarithmic growth. The concentration of each culture was measured and samples containing 3 x 10^7 cells/ml were collected through centrifugation, the supernatant removed, and the total genomic DNA extracted using OmniPrep™ for Yeast (G-Biosciences, St. Louis, MO).

For RNA collection, MM cultures of two wildtype naïve and four wildtype, melanized adapted strains were transferred to MMS and cultured for an additional 24 hrs to simulate starvation conditions.

Table 1

| Table 1 | Strains used in this study. |
|---------|-----------------------------|
| Original Strain | Adaptation conditions [30] | Total Dose (over 5 weeks/35d) | Isolate ID |
| ATCC34100, Exophiala dermatitidis CBS 525.76 | Wildtype Naive (WTN) | Background | 1, 2 |
| ATCC34100, Exophiala dermatitidis CBS 525.76 | Wildtype 225Ac Adapted (WTA) | 183 Gy | 3–6 |
| wdpks1 | Albino Naive (pksΔ) | Background | 7, 8 |
| wdpks1 | Albino 225Ac Adapted (pksA) | 183 Gy | 9, 12 |
Cultures were then and adjusted to $10^7$ cells/ml and transferred to 6-well plates. Each plate was then exposed to one of the following conditions:

1) Background radiation
2) Ionizing radiation exposure
   The $^{210}$Po strip (500 μCi) was placed directly above of the wells and generated a 0.5 μGy/hr dose rate. The fungus settled to the bottom, and the growth media was sufficient to shield alpha-particles from direct interaction with the fungal cells.
3) ROS exposure (through addition of 0.1 mM H$_2$O$_2$ to the medium)

Exposures took placed at 30 °C in the dark for 1 week, after which total RNA was extracted from the samples with the Ribopure RNA purification Kit for Yeast (ThermoFisher Scientific). Libraries were constructed and sequenced on the Illumina NovaSeq S4 Sequencer by the Yale Center for Genome Analysis (YCGA, Yale School of Medicine, West Haven, CT).

### 2.4. Genomic analysis

For genome sequencing 2 wildtype naïve, 3 wildtype adapted, 2 wdpks1 naïve, and 2 wdpks1 adapted strains were analyzed (Table 1). The reference genome and annotation files for the raw paired-end 150 base reads were trimmed using Trimmomatic v0.36 [6] and aligned to the publicly available *E. dermatitidis* NIH/UT8656 reference genome obtained from the EnsembFungi web portal (https://fungi.ensembl.org/). Specifically, the Burrows-Wheeler Aligner (BWA) software package v 0.7.17 [24] was used to map reads and the sequence data was further processed using SAMtools v 1.9. PCR artifacts were removed using Picard tools MarkDuplicates (https://broadinstitute.github.io/picard/) and variants were identified using GATK v 3.8.7 [32]. GATK’s IndelRealigner was used to realign reads containing putative INDELS and GATK’s Haplotype Caller was used to call variants. GATK’s GenotypeGVCFs was used to combine the resulting Variant Call Format (VCF) files, which generated one VCF file for all *E. dermatitidis* WT samples and a second VCF files for *pks* samples, which were filtered using GATK’s VariantFilteration based on stringent cutoffs for quality and coverage (SNPs: QD $< 2.0$, MQ $< 40.0$, QUAL $< 100$, FS $> 60.0$, MQRankSum $< -12.5$,SOR $> 4.0$, ReadPosRankSum $< -8.0$; INDELS: QD $< 2.0$, FS $> 200.0$, MQRankSum $< -12.5$, SOR $> 4$, InbreedingCoeff $< -0.8$, ReadPosRankSum $< -20.0$). The resulting high-quality variants were then filtered such that any variant present in the naive strain was removed from that strain’s sample set. The remaining variants were annotated using annotation files obtained from EnsembFungi and functional effects were predicted using snpEff [9].

### 2.5. Transcriptomic analysis

Analysis of RNA-seq data was performed on samples outlined in Table 2 in the following manner, as previously described [45]: transcript lengths and counts were determined using Salmon v 1.3.0 software [38] with raw FASTQ reads as input and an index built from the *E. dermatitidis* NIH/8656 transcriptome that was assembled for prior experiments and is available on NCBI (Accession No. PRJNA224192) [45,8,41]. The resulting quant.sf files were used for pairwise comparisons with the DESeq2 package from Bioconductor version 3.0 [13] or R Studio version 3.1 [1]. Genes were differentially expressed between two conditions if their adjusted p-value (also referred to here as FDR) was <0.05. The resulting lists of differentially expressed genes were compared between conditions using Venny [33], and were analyzed for enrichment of certain Gene Ontology – Biological Process categories using FungiFun2 (https://elbe.hki-jena.de/fungifun/) [39].

### Table 2

| Condition                     | WT Melanized Strain Analyzed | Isolate ID |
|-------------------------------|------------------------------|------------|
| Background Radiation          | WT Naïve Control (WTN-C)     | 1          |
|                               | WT Adapted Control (WTA-C)   | 2          |
| Ionizing Radiation Exposure   | WT Naïve Radiation (WTN-R)   | 3          |
|                               | WT Adapted Radiation (WTA-R) | 4          |
| Reactive Oxygen Species Exposure | WT Naïve Oxidation (WTN-O) | 5          |
|                               | WT Adapted Oxidation (WTA-O) | 6          |

### 3. Results and discussion

#### 3.1. Genomic changes in radioadapted *E. dermatitidis* strains

Because ionizing radiation is a potent mutagen, we first assessed the genomic change that occurred within the adapted isolates from both the wildtype (WT) and wdpks1 (*pks*) genetic backgrounds to understand how protracted exposure affected their DNA sequence.

An overview of the genetic variants identified in WTA and *pks* strains relative to the naïve strains (WT and *pks*) are displayed in Table 3. Interestingly, very few mutations were observed in the WTA radioadapted strains, with the number of identified SNPs ranging from 0 to 2 and the number of INDELS ranging from 2 to 9. Observed variants included a missense mutation in strain WTA5 in the HMPREF1120_01028 gene, which encodes an ATP-dependent RNA helicase, a conservative iframe insertion in WTA3 in the HMPREF1120_07349 gene, which encodes a flagellar motor protein domain, and a disruptive iframe deletion in WTA6 in the HMPREF1120_00159 gene, which encodes a kinesin family member (Table 4, Supporting Table 1). Notably, two frameshift mutations (which often lead to a truncated or nonfunctional protein products) were observed in the WTA3 radioadapted strain. These variants occurred in stress-response nuclear envelope protein-encoding HMPREF1120_01736 and alpha/beta-hydrolase-encoding HMPREF1120_03360.

Among the wdpks1 radioadapted strains, *pks*A9 featured 2 SNPs and 7 INDELS, which was comparable to the number of variants observed in the WT radioadapted strains (Table 3). These variants included frameshift mutations in HMPREF1120_06854, which encodes phytanoyl-CoA hydroxylase, and HMPREF1120_04304, which encodes ubiquitin-conjugating enzyme E2 (Table 5, Supporting Table 2). Notably, the latter frameshift insertion was also observed in *pks*A12. An increased number of SNPs was observed in *pks*A12 relative to all other radioadapted strains (Table 5, Supporting Table 2). Most of these SNPs (17), and well as 2 INDELS, occurred in a 500 bp intergenic region approximately 2.5 kbs upstream from the coding region of HMPREF1120_04917, which encodes ubiquitin C-terminal hydrolase. The *pks*A12 strain also harbored frameshift mutations that occurred in heat shock protein domain-encoding HMPREF1120_00517, arginase-encoding HMPREF1120_06649, and non-ribosomal peptide synthetase (NRPS) SidC-encoding HMPREF1120_07636, which is involved in siderophore biosynthesis. Another notable variant was an inter-
mutation type, which was predicted using snpEff. For intergenic variants, the gene name and annotation refer to the gene that is closest to that mutation.

### Table 4

| CHROM | POSITION | MUTATION TYPE                  | GENE/NEAREST GENE                  | ANNOTATION                             | OCCURRENCE |
|-------|----------|--------------------------------|------------------------------------|----------------------------------------|------------|
| 1     | 433,055  | Disruptive inframe deletion    | HMPREF1120_00159                   | Kinesin family member                   | WTA6       |
| 2     | 2,836,819| Missense                        | HMPREF1120_01028                   | ATP-dependent RNA helicase              | WTA5       |
| 3     | 631,490  | Frameshift                      | HMPREF1120_01736                   | Stress-response nuclear envelope         | WTA3       |
| 4     | 2,527,760| Intergenic                      | HMPREF1120_02837                   | UDP-glucose 4-epimerase                 | WTA3       |
| 5     | 1,006,101| Frameshift                      | HMPREF1120_03360                   | Alpha-beta-hydrolase                     | WTA3       |
| 6     | 1,268,882| 3’ UTR variant                  | HMPREF1120_03462                   | Phosphatidylinositol glycan             | WTA3       |
| 7     | 1,576,609| Intergenic                      | HMPREF1120_03561                   | Alanine transaminase                     | WTA6       |
| 8     | 3,187,642| Intergenic                      | HMPREF1120_04157                   | MFS transporter                         | WTA3, WTA5, WTA6 |
| 9     | 1,158,832| Intergenic                      | HMPREF1120_04747                   | Carbonic anhydrase                      | WTA5       |
| 10    | 6,226,494| Intergenic                      | HMPREF1120_08594                   | SS family solute:Na + symporter         | WTA3, WTA6 |
| 11    | 1,418,001| Conservative inframe insertion  | HMPREF1120_07349                   | Flagellar motor protein MobB domain     | WTA3       |
| 12    | 1,971,753| Intergenic                      | HMPREF1120_07545                   | Redox-sensitive bicupin: pirin superfamily | WTA3, WTA5 |
| 13    | 1,180,436| Intergenic                      | HMPREF1120_08242                   | Hypothetical protein                    | WTA3       |

### Table 5

| CHROM | POSITION | MUTATION TYPE                  | GENE/NEAREST GENE                  | ANNOTATION                             | OCCURRENCE |
|-------|----------|--------------------------------|------------------------------------|----------------------------------------|------------|
| 1     | 4,126,160| Frameshift                      | HMPREF1120_00517                   | DnaJ-domain (heat shock protein)        | pksA12     |
| 2     | 3,069,904| Intergenic                      | HMPREF1120_01028                   | DNA Pol 3 subunit                       | pksA9, pksA12 |
| 3     | 843,500  | Intergenic                      | HMPREF1120_03030                   | Endo-1,3(4)-beta-glucanase              | pksA9      |
| 4     | 2,568,652| 3’ UTR variant                  | HMPREF1120_03921                   | Hypothetical protein                    | pksA12     |
| 5     | 3,187,642| Intergenic                      | HMPREF1120_04157                   | MFS transporter                         | pksA9, pksA12 |
| 6     | 3,617,951| Frameshift                      | HMPREF1120_04304                   | Ubiquitin-conjugating enzyme E2         | pksA9, pksA12 |
| 7     | 1,631,962| Intergenic                      | HMPREF1120_04917                   | Ubiquitin C-terminal hydrolase          | pksA12     |
| 8     | 1,631,965| Intergenic                      | HMPREF1120_04917                   | Ubiquitin C-terminal hydrolase          | pksA12     |
| 9     | 1,631,762–1632254| Intergenic variants | HMPREF1120_04917 | Ubiquitin C-terminal hydrolase | pksA12     |
| 10    | 998,035  | Conservative inframe deletion   | HMPREF1120_09594                   | Transducin (beta)-like 1                | pksA12     |
| 11    | 3,033,533| Frameshift                      | HMPREF1120_06649                   | Arginase                               | pksA12     |
| 12    | 48,768   | Frameshift                      | HMPREF1120_06854                   | Phytanoyl-CoA hydroxylase               | pksA9      |
| 13    | 1,198,245| Missense                        | HMPREF1120_07268                   | Alcohol dehydrogenase (NADP+)           | pksA12     |
| 14    | 1,666,136| Disruptive inframe deletion     | HMPREF1120_07440                   | MFS transporter, SP family, sugar:H + symporter | pksA12 |
| 15    | 1,971,753| Intergenic                      | HMPREF1120_07545                   | Redox-sensitive bicupin: pirin superfamily | pksA9      |
| 16    | 2,002,737| Intergenic                      | HMPREF1120_07545                   | Redox-sensitive bicupin: pirin superfamily | pksA12     |
| 17    | 2,252,182| Frameshift                      | HMPREF1120_07636                   | NRF2 Sod2 (siderophore biosynthesis)     | pksA12     |
| 18    | 70,963   | Intergenic                      | HMPREF1120_07867                   | Multi-sensor signal transduction histidine kinase | pksA12 |
| 19    | 1,099,400| Intergenic                      | HMPREF1120_08213                   | Transcription factor                    | pksA9      |
| 20    | 1,142,537| Missense                        | HMPREF1120_08228                   | Hypothetical protein                    | pksA12     |
| 21    | 1,503,364| Intergenic                      | HMPREF1120_08366                   | Autophagy-related protein               | pksA9, pksA12 |

We previously demonstrated that radioadaptation resulted in increased colony growth in radioadapted strains in comparison with naïve strains when both strains were exposed to ionizing radiation from an α particle emitting Polonium-210 source, as well as increased electron transfer capacity, and improved resistance to the toxic effects of ROS [30].

To understand what was occurring to produce this altered biological response, we set out to characterize the transcriptome of the adapted, melanized strain (WTA) relative to the naïve, melanized strain (WT). For this experiment, we used wild type strains (Table 2) that were also characterized by genome sequencing, including four strains isolated from the wildtype 225Ac adapted culture (WTA3-6) were used, along with two control strains.
(WTN1-2) that were isolated from a naïve, wildtype culture not previously exposed to this radionuclide. We did not perform this analysis on the non-melanized strains, as they did not exhibit radiation-associated phenotypes in our initial experiments [30].

The overall transcript reads and expression changes between each strain and condition are given in Supporting Tables 3 and 4. The first analysis on this data that we performed on this data was to view the correlation between Transcripts Per Million (TPM) values produced by the Salmon alignment and transcript quantification program, to determine how each dataset was correlated across replicates, strains, and conditions. Notably, Fig. 2 shows that this comparison revealed that the TPM values from control sample 1 (wildtype, naïve sample 1C) highly correlated ($R^2 > 0.95$) not only with sample 2 (wildtype, naïve sample 2C), but with also samples 3 (wildtype, adapted 3C) and 6 (wild type, adapted 6C), whereas samples 4 (wildtype, adapted 4C) and 5 (wildtype, adapted 5C) were closely correlated with each other under this condition ($R^2 = 0.9735$) but substantially different than the other four samples ($R^2 < 0.73$). This suggested to us that the adapted culture from which the four strains were isolated included a mixture of at least two strains – one with a transcriptome that we observed was similar to the naïve strain, and one that had a unique transcriptomic signature. The second observation we made from this analysis, as demonstrated in Fig. 3, was that the subsequent stress exposures (to $^{210}$Po and $H_2O_2$) had extremely minor effects on the transcriptomes of both sets of strains (represented here by WTN and WTA). Overall, the number of differentially expressed genes between each condition and sample can be observed in Fig. 4.

3.3. Differentially expressed genes in the adapted strains

It is clear from the total number of differentially transcribed genes, that the major differences observed were associated with the adaptation process in isolates 4 and 5, and not the exposure conditions. These strains demonstrated that, although not essential, there was a possibility for a marked change in gene expression in cells that were incubated in such an environment, so we focused on the differences between these (hereafter WTA) and the naïve (hereafter WTN) strains. We did this by analyzing the transcripts that changed in abundance between these two pairs of strains either in background conditions or after exposure to $^{210}$Po or to 0.1 mM $H_2O_2$ under conditions of starvation. Such differences under background conditions would suggest more stable changes between the two strains that could potentially be used as signatures of previous exposure in E. dermatitidis, while differences in the response to $^{210}$Po, for example, would reflect the responses of radioadapted strains to future radiation challenge. Fig. 5 attempts to unravel these possibilities by observing the overlap between these three sets of up- and down-regulated genes. Interestingly, although the exposure did not greatly affect the transcriptome in any given strain lineage, less than half of the differentially expressed genes were shared among the three conditions, suggesting that the adapted strains had a unique response to these distinct environments. This was especially notable for $^{210}$Po exposure, as 696/1713 of the genes differentially expressed between the two strains were unique to this condition. We view this result as evidence that the adapted strain had the ability to respond more robustly to a second exposure to ionizing radiation. The differences between the WTN and WTA strains with varied transcriptomic signatures become even clearer after gene enrichment analysis, where one considers the function of these sets of significantly regulated transcripts (Table 6). We observed that in all cases, genes encoding proteins involved in oxidation–reduction processes were significantly downregulated in the adapted compared to the naïve. The oxidation–reduction process title is some-what misleading, as what we actually see are genes involved in catalyzing redox reactions in the context of the electron transport chain and metabolism (Supporting Table 8), and include enzymes such as alcohol dehydrogenase, acyl-CoA dehydrogenase, cytochrome P450 monoxygenase, glucose-1 dehydrogenase [17]. When you look at how the genes in this process overlap, a substantial number were unique to the radioactive environment conditions (Fig. 5 and Supporting Tables 5–7), yet these unique genes were still primarily involved in catalyzing redox reactions in the context of the electron transport chain and metabolism. There is a down regulation in catalase (HMMPREF1120_07713) and superoxide dismutase (HMMPREF1120_05007,) which are enzymes involved in response to oxidative stress [54] but they are downregulated in the adapted strains across all conditions. This suggest that it is part of the adaptive process, but not involved in the mechanisms involved in improved fitness and ROS resistance observed under subsequent exposure to radiation.

Transmembrane transport and the general category of “metabolic process” were also downregulated in two out of the three gene sets. These are relatively vague functional classes, but overall, they suggest a suppression down of several primary metabolic and redox reactions in the adapted strains.

The patterns observed in the upregulated group were easier to interpret. For example, functions related to translation were enriched in all three sets. This suggests an increase in protein production, which was, interestingly, also observed when E. dermatitidis was grown in the presence of low dose $\gamma$-radiation [41]. Another interesting observation is the enrichment of genes involved in DNA replication in the adapted cells exposed to $^{210}$Po, a pattern which was even clearer when only the genes unique to this condition were analyzed (Fig. 5). In this case, not only DNA replication but DNA recombination was enriched, as well as the obsolete ATP catabolic processes, which upon review included 27 genes, including 13 helicases, 2 DNA topoisomerases, and an elongation factor (EF-3), many of which are involved in RNA remodeling, DNA repair, [21,12,15,3]. Overall, there were many genes that were unique between the WTR and the AR set compared to the other two conditions (Supporting Tables 5–7). This points to both the possible presence of DNA damage even under these low dose conditions as well as the ability of the adapted strain to respond more quickly and robustly to such damage. Genes uniquely upregulated in the adapted strain in response to irradiation included several DNA helicases, DNA ligase, and the DNA repair proteins rad52, ERCC-6, rhp54, rad9, uvsE, among others, while downregulated genes, which showed a greater magnitude of regulation, included transporters (10 major facilitator superfamily proteins), but mostly included proteins with only general predictions, suggesting that this regulatory response was subtle and complex, and involved in several biosynthetic pathways (e.g. biotin and glycogen synthesis, Supporting Tables 5 and 7). In all, although we did not observe large responses to exposure to oxidative stress or irradiation, we did observe a strong and consistent change in one lineage of E. dermatitidis that was adapted to long-term radionuclide presence, and we observed evidence that this adaptation process allowed for this strain to become more responsive to future ionizing radiation exposure.

3.4. Adaption in E. dermatitidis

The purpose of the study was to identify the underlying genomic and transcriptomic events that are the result of radioadaptation in E. dermatitidis cultures and to further understand how those changes impact the fungal response to subsequent radiation exposure. These changes, we believe, would allow us to identify and understand the adaptation that is taking place.
Ionizing radiation induces the production of free radicals, including reactive oxygen and nitrogen species. Our previous data demonstrated that radioadapted cells presented similar biological responses to both subsequent H₂O₂ and ionizing radiation exposure, so we could not determine whether adapted strains could distinguish these stresses [30]. In our previous work we assessed response of the adapted strains to strontium-90 (beta) and cesium-137 (gamma) [30]. We found that while there was a stimulation of growth with a gamma emitter in the naïve strain, the adapted strain showed no enhanced growth. Neither the naïve nor the adapted strain showed enhanced growth with a beta source.

Here, what we observed was enhanced colony growth in our melanized adapted strain (Fig. 1) only when using ²¹⁰Po, an alpha-particle emitter. Relative to gamma and beta radiation, alpha particles have with a high linear energy transfer (LET) and are capable of generating more ROS through radiolysis when interacting with the growth media that the fungus would be plated on [22]. This is why alpha radiation is densely-ionizing, while gamma and beta radiation are sparsely-ionizing. We did not observe any

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**Fig. 2.** Comparison of control RNAseq samples using Transcripts Per Million (TPM) values observed for each predicted transcript encoded by the *E. dermatitidis* genome. Samples 1 and 2 represent naïve strains, while samples 3–6 represent adapted strains.

**Fig. 3.** Comparison of the responses of naïve (Strain 1) and adapted (Strain 4) to H₂O₂ (O) and Polonium-210 (R) exposure, visualized with RNAseq data using Transcripts Per Million (TPM) values observed for each predicted transcript encoded by the *E. dermatitidis* genome.
enhanced growth response when these strains were grown near radioactive sources with lower LET, so we hypothesized that the difference in biological response could be due to enhanced resistance to the toxic effects of the ROS generated by the alpha particles. This hypothesis seemed plausible, as our melanized adapted strain presented enhanced resistance to H$_2$O$_2$.

Our initial analysis using whole genome sequencing was completed to compare the naïve and adapted strains of the melanized wildtype and albino wpdks1 mutants (Table 1). There was not a significant difference in the number or type of mutations when comparing the adapted strains of the WTA to the pkSA mutants (Table 3–5) which indicates the phenotypic changes observed in the melanized adapted strain were not genomic in nature. In fact, only 4 mutations were observed in the one of the strains that presented an altered transcriptomic signature (Table 5).

The transcriptomic analysis, unsurprisingly, provided us with more information to explore. With this data, we were able to make two major conclusions. First, we observed that the stresses that we challenged these strains with ($^{210}$Po and H$_2$O$_2$) did not dramatically affect the transcriptome, at least at the time point that we characterized (1 week). This suggests that E. dermatitidis can return its transcriptome to a steady state after an extended period under a moderate stress. Second, we identified two lineages with varied transcriptomic signatures in the radioadapted, melanized E. dermatitidis strain. One lineage aligned with the naïve strain (Figs. 2 and 3), while the other presented a unique signature. It is not surprising that we observed different lineages within the adapted strain, as adaptation was performed on a pool of stationary phase cells where individual cells would have gone through radiation exposure and age-related mutagenesis, which would have resulted in some variance [18,42].

We then focused on the lineage that showed a distinct expression pattern, which provided us with several interesting results. First, we observed a general upregulation of translation genes, which could suggest that the adapted strains enhance their protein synthesis machinery for resilience to environmental stresses, such as ionizing radiation and ROS. We previously observed a similar expression pattern upon low dose gamma irradiation in wildtype E. dermatitidis [41] while in response to acute, high dose gamma irradiation we see the opposite pattern – ribosomal genes being strongly downregulated [45]. Ribosomal biosynthesis is tightly associated with growth rate, so the upregulation of this set of genes may be a mark of the apparent increase in growth rate we observe in the adapted strains [53]. The enrichment of genes involved in glycolysis (glycolytic process) among the upregulated transcripts in the adapted strains may also be indicative of faster growth. On the other hand, several genes in the GO categories of oxidation–reduction processes and metabolic processes were significantly downregulated. We consider this to signify two potential situations. First, because these enzymes were generally involved in a wide variety of diverse and vague metabolic pathways, it is possible that their downregulation is due to an overall streamlining of the cell toward protein synthesis. Second, these changes could indicate an overall global change in the redox balance of the cell, which is to be expected in strains adapted to high ROS environments such as chronic irradiation. Electron transport and metabolism, in fact, themselves are a source of ROS [17,54] so a down regulation in these processes could reduce the internal production of ROS, and thereby position the fungi to withstand a more robust external load of ROS. Respiration in fungi is not a linear process but is in fact significantly branched, with numerous alternative enzymes [17]. The more substantial response in downregulation of more of these enzymes in the $^{210}$Po conditions (Fig. 5B) thereby reducing the redundancy in the metabolic processes and could be a unique response to the $^{210}$Po conditions to further reduce the internal ROS load.

Finally, by comparing the pairwise analysis of wild type and adapted strains responding to each condition, another subset of genes, specifically upregulated in response to $^{210}$Po exposure was revealed. These included genes traditionally associated with ionizing radiation damage, including those involved in DNA replication and recombination. This finding is interesting considering the extensive literature on "inducible repair" in fungi, where cells recovering from an initial exposure to ionizing radiation are more resilient to a subsequent exposure [23,51]. This is presumably due to the cell's ability to mount and therefore more efficiently respond to the second round of damage, but this phenomenon has not generally been characterized at the transcriptomic level. It is also particularly interesting because this response is distinct from the response to H$_2$O$_2$, suggesting that the cellular responses to general oxidative stress do not necessarily overlap with the response to ionizing radiation exposure. It will be interesting, at this juncture, to understand whether these changes result in increased resistance to acute ionizing radiation or other DNA damage [52] as it is still controversial whether the inducible repair response is mediated at the transcript or protein level, and most experiments on inducible repair do not initiate with exposure to the low doses that were used with these adapted strains.

In our previous work melanized E. dermatitidis presented an altered biological response following the adaptation process in the face of subsequent radiation and ROS exposures while the non-melanized strain did not [30]. This observation led us to hypothesize that we would observe a robust transcriptomic response under these same conditions in the melanin biosynthetic pathway, represented by the polyketide synthase (PKS) gene (HMPREP1120_03173). Analysis of the transcriptome of the adapted strains showed a slight down regulation of the PKS gene under all conditions when compared to the naïve strain.
ing Table 4) with log2fold change values ranging from −0.49 to −0.63. While melanin plays an essential role in the phenotypic response of adapted *E. dermatitidis*, transcriptomic regulation of the biosynthesis of the pigment does not appear to be substantially altered by the adaption process. We also looked at alterations in carotenoid genes (HMPREF1120_03263, HMPREF1120_02864), which are known for their antioxidant and photoprotective properties [26,40] and found no substantial changes (Supporting Table 4) due to the adaptation process in these genes.

4. Conclusion

In conclusion, we observed several things when completing a genomic and transcriptomic study on *E. Dermatitidis* strains
adapted to $^{223}$Ac exposure. First, the adaptation response appeared to be mediated at the transcriptomic level, as the adaptation process resulted in very few, and likely insubstantial genetic mutations. This also supports the extremely stable genome in the face of protracted exposure to ionizing radiation. Second, we found that even a low dose of radiation with a high LET in the environment, or ROS in the form of H$_2$O$_2$, elicited a transcriptomic response that in general resulted in down regulation of metabolic transcripts and an up regulation in translation in the adapted versus naïve melanized *E. dermatitidis*. Downregulation of metabolic processes in general would reduce the internal ROS load, thereby positioning the adapted fungi to better respond to an external load of ROS, while an upregulation of translation would prime the fungi for growth. Finally, we noted, that despite a similar number of up/down regulated transcripts, the adapted cultures, when exposed to radiation in their environment, presented a unique transcriptional profile, whereas the transcriptomic profile of the culture exposed to ROS significantly overlapped with the control culture. While several of the differentially regulated processes where similar between the conditions, the scale of transcripts that were unique to the $^{210}$Po condition suggests a more robust response. Also unique to this condition was a general upregulation of DNA replication and repair enzymes, suggesting that previous exposure to radiation positions the adapted strain to better respond to damage caused by an alpha-particle emitting source in its environment. From this we can conclude that the adaptation process resulted in an altered transcriptomic response from fungus grown in the same conditions without radiation, and that this previous exposure altered how this fungus responded to subsequent interactions, even indirect.

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**CRediT authorship contribution statement**

Mackenzie E. Malo: Conceptualization, Data curation, Formal analysis, Investigation, Visualization, Writing - original draft, Writing - review & editing. Zachary Schultzhaus: Data curation, Formal analysis, Writing - original draft, Writing - review & editing. Connor Frank: Investigation. Jillian Romsdahl: Data curation, Formal analysis, Writing - original draft, Writing - review & editing. Zheng Wang: Funding acquisition, Supervision, Writing - review & editing. Ekatrina Dadachova: Funding acquisition, Project administration, Resources, Supervision, Writing - review & editing.

**Appendix A. Supplementary data**

**Supplemental Table 1: SNPs and INDELS identified in WTA radioadapted strains.** *Supplemental Table 2: SNPs and INDELS identified in pksA radioadapted strains.** *Supplemental Table 3:* Salmon output for analysis of transcriptomic data including transcripts per million (TPM) and transcript read lengths. **Supplemental Table 4:** Output from DESeq2 analysis between given conditions demonstrating Log2fold changes and significance information. Genes with a padj < 0.05 were determined to be differentially expressed. **Supplemental Table 5:** Genes that were uniquely upregulated/downregulated in adapted strains compared to naïve strains, in the presence of ionizing radiation. **Supplemental Table 6:** Uniprot annotations of upregulated genes from *Supplemental Table 5*. **Supplemental Table 7:** Uniprot annotations of downregulated genes from *Supplemental Table 5*. **Supplemental Table 8:** Enriched genes in the GO:005514 Oxidation-Reduction Process category. Supplementary data to this article can be found online at https://doi.org/10.1016/j.csbj.2020.12.013.

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